

Combined phylogenetic analysis of the subclass Marchantiidae (Marchantiophyta): towards a robustly diagnosed classification

Jorge R. Flores^{a,b,*}, Santiago A. Catalano^{a,b}, Jesus Muñoz^c and Guillermo M. Suárez^{a,b}

^aUnidad Ejecutora Lillo (UEL; FML-CONICET), Miguel Lillo 251, S.M. de Tucumán 4000, Argentina; ^bFacultad de Ciencias Naturales e Instituto Miguel Lillo, Universidad Nacional de Tucumán, Miguel Lillo 205, S.M. de Tucumán 4000, Argentina; ^cReal Jardín Botánico (RJB – CSIC), Plaza de Murillo 2 Madrid, 28014, Spain

Accepted 22 August 2017

Abstract

The most extensive combined phylogenetic analyses of the subclass Marchantiidae yet undertaken was conducted on the basis of morphological and molecular data. The morphological data comprised 126 characters and 56 species. Taxonomic sampling included 35 ingroup species with all genera and orders of Marchantiidae sampled, and 21 outgroup species with two genera of Blasiidae (Marchantiopsida), 15 species of Jungermanniopsida (the three subclasses represented) and the three genera of Haplomitriopsida. *Takakia ceratophylla* (Bryophyta) was employed to root the trees. Character sampling involved 92 gametophytic and 34 sporophytic traits, supplemented with ten continuous characters. Molecular data included 11 molecular markers: one nuclear ribosomal (26S), three mitochondrial genes (*nad1*, *nad5*, *rps3*) and seven chloroplast regions (*atpB*, *psbT-psbH*, *rbcL*, *ITS*, *rpoC1*, *rps4*, *psbA*). Searches were performed under extended implied weighting, weighting the character blocks against the average homoplasy. Clade stability was assessed across three additional weighting schemes (implied weighting corrected for missing entries, standard implied weighting and equal weighting) in three datasets (molecular, morphological and combined). The contribution from different biological phases regarding node recovery and diagnosis was evaluated. Our results agree with many of the previous studies but cast doubt on some relationships, mainly at the family and interfamily level. The combined analyses underlined the fact that, by combining data, taxonomic enhancements could be achieved regarding taxon delimitation and quality of diagnosis. Support values for many clades of previous molecular studies were improved by the addition of morphological data. The long-held assumption that morphology may render spurious or low-quality results in this taxonomic group is challenged. The morphological trends previously proposed are re-evaluated in light of the new phylogenetic scheme.

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Introduction

Among embryophytes (land plants), liverworts (Marchantiophyta) are recognized as the basal group (Qiu et al., 1998; Nickrent et al., 2000; Karol et al., 2001). The complex thalloid liverworts (Marchantiidae) encompass about 345 species and 36 genera (Villarreal et al., 2016) which are remarkably different from other embryophytes. Complex thalloid liverworts are characterized by a combination of both plesiomorphic characters and novelties in their structure. On the one hand, several of these morphological features, such as a

haplodiploid life cycle and the absence of vascular tissue, are widespread among bryophytes. On the other, highly specialized fertile branches (gametangiophores), internal schizogenous air chambers and pegged rhizoids are traits exclusive to the complex thalloid liverworts (Schuster, 1984a,b; Bischler, 1998; Crandall-Stotler and Stotler, 2000). Given their position within embryophytes and the morphological peculiarities, Marchantiidae stand as crucial taxa to disentangle the evolutionary history of land plants.

The first classifications within Marchantiidae, based on phylogenetic systematic principles, were presented based on morphological data. However, the subsequent classifications relied exclusively on molecular data. The most comprehensive morphological phylogeny (Bischler,

*Corresponding author.

E-mail address: jrflores@conicet.gov.ar

1998) supported the distinction of a basal Ricciineae and a derived group comprising Corsiineae + Targioniineae + Marchantiineae. One of the few combined analyses was published by Boisselier-Dubayle et al. (2002) and included 27 out of 35 genera of Marchantiidae. The combined trees obtained in that study rendered a completely different pattern of relationships from that published by Bischler (1998). Subsequent molecular analyses largely agree with the relationships obtained by Boisselier-Dubayle et al. (2002), except in the relationships among morphologically simple families (Wheeler, 2000; Crandall-Stotler et al., 2005; Forrest et al., 2006; He-Nygrén et al., 2006).

Boisselier-Dubayle et al. (2002) analysed the conflict between morphological data and molecular data, showing that both sources of evidence produced very different trees, but also that support values could be improved by concatenating different data types. Crandall-Stotler et al. (2005) reported a high degree of homoplasy within morphological data. Nevertheless, they were able to find synapomorphies for most of the nodes. Subsequently, many diagnoses were modified and several more diagnostic characters were found to define higher taxonomic groups (e.g. lenticular apical cell for Metzgeriidae; Crandall-Stotler et al., 2009) than in previous classifications. Contrasting with Boisselier-Dubayle et al. (2002) and Crandall-Stotler et al. (2005), He-Nygrén et al. (2006) did not carry out an explicit survey on character conflict. Even so, they performed the most extensive combined analysis in terms of morphological data inclusion. In addition, the results obtained by He-Nygrén et al. (2006) displayed no significant differences when different optimality criteria were considered. Although He-Nygrén et al.'s study was the first attempt to reach a reliable diagnosed classification, Marchantiidae taxonomic sampling included only 13 genera (37% of the accepted genera in the subclass).

The current accepted classification (Crandall-Stotler et al., 2009) was achieved as a result of several multi-locus phylogenies produced during the last decade. Compared to the previous morphology-based classification, several taxonomic categories were omitted. In addition, many genera of Marchantiidae were relocated in completely different categories (e.g. *Monocarpus* D.J. Carr (Forrest et al., 2015) or *Neohodgsonia* (Perss.) Perss. (Crandall-Stotler et al., 2009)). As Crandall-Stotler et al. (2009) pointed out, diagnosing these unexpected groups—from the morphological point of view—represented a real challenge.

Despite the advances in the morphological knowledge of Marchantiidae accomplished in the last decade (e.g. Duckett et al., 2014), there was no attempt to include this new information into quantitative phylogenetic analyses. Although recently performed phylogenies sampled a considerable number of molecular markers (Forrest et al., 2006, 2015; Villarreal et al.,

2016), support values for several nodes were still low. On the one hand, some taxonomic changes have been recently proposed based upon these clades with low support (Long et al., 2016a,b). For example, given that the genera *Exormotheca* Mitt. and *Stephensiella* Kashyap were found to be the sister taxa of *Cronisia* Berk., Exormothecaceae was merged with Corsiniaceae (Long et al., 2016a). However, the node constituted by *Cronisia*, *Exormotheca* and *Stephensiella* lacked significant support values (fig. 2 in Villarreal et al., 2016). On the other hand, taxonomic proposals based on nodes with high support were controversial from a morphological perspective (Forrest et al., 2015; Long et al., 2016b). Even if the average support at the family level was high in recent molecular studies (Forrest et al., 2006; Villarreal et al., 2016), deep nodes remained ambiguous (Wheeler, 2000; Boisselier-Dubayle et al., 2002; Forrest et al., 2006; Villarreal et al., 2016). A large number of the inclusive nodes within Marchantiales, the crown group of the subclass (Crandall-Stotler et al., 2009), still had low support values (Villarreal et al., 2016). In particular, derived nodes (Ricciaceae, Oxymitraceae, Monocleaceae, etc.) were hard to resolve even in combined analyses (Boisselier-Dubayle et al., 2002). Therefore, with few exceptions, interfamilial relationships persisted as a taxonomic challenge (Crandall-Stotler et al., 2009).

The strategies to overcome environmental stress and their significance to bryophytes evolution were regular issues of interest in previous studies (Bischler and Jovet-Ast, 1981; Longton, 1988a; Hedderson and Longton, 1996; Bischler, 1998). Bischler (1998) defined four different groups based on different morphological and ecological characteristics, postulating a strong correspondence of those groups with morphological characters and taxonomic/phylogenetic membership. Similar morphologies were explained as consequences of common ancestry rather than similar selective forces (Bischler, 1998; Bischler et al., 2005). Concerning mosses, Longton (1988b) stated that the basic life strategy was being perennial, considering ephemeral strategies as derived. In contrast, Bischler (1998) considered ephemeral life traits as ancestral for complex thalloid liverworts. Although character mapping has been considered to elucidate the evolutionary pattern of some traditional characters (Villarreal et al., 2016), Bischler's hypothesis on the concerted evolution of life-history traits has not been challenged analytically.

The recent modifications within the phylogeny of Marchantiidae have involved both new scenarios of evolutionary trends and changes in morphological diagnosis of different clades within this group. Here we present the results of the largest combined phylogenetic analysis in Marchantiidae in terms of taxon and character sampling. This study allows the solving of previous uncertainties in some nodes, updating group

diagnoses and testing previous ideas about character evolution. In addition, the contribution to and conflict between morphological characters from the different life phases were evaluated in depth.

Materials and methods

Taxonomic and morphological character sampling

Despite taxonomic changes having been recently proposed for some groups, the main taxonomic scheme followed throughout this study was that of Crandall-Stotler et al. (2009). After Long et al. (2016a, b), the genera *Exormotheca* and *Aitchisoniella* Kayshap were considered members of Corsiniaceae and Cleveaceae, respectively. However, in order to test recently proposed synonyms (Long et al., 2016a), the following genera were scored as independent entities: *Stephensiella* and *Exormotheca*, and *Preissia* Corda, *Bucegia* Radian and *Marchantia* L. In the combined dataset, ingroup sampling consisted of 37 genera of Marchantiopsida (35 genera of Marchantiidae plus two genera of Blasiidae), which represented all of the orders, families and genera within the class. Outgroups included 15 species of Jungermanniopsida, with the three major subclasses of this group being represented (Jungermanniidae, Pelliidae and Metzgeriidae). In addition, the dataset included species from the three genera of Haplomitriopsida. *Takakia ceratophylla* (Mitt.) Grolle (Bryophyta) was employed to root the tree. Five genera were absent in the morphological partition: *Cavicularia* Steph. (Blasiidae), *Riella* Mont. (Marchantiidae), *Apotreubia* S. Hatt. & Mizut. (Haplomitriopsida), *Pleurozia* Dumort and *Pallavicinia* Gray (Jungermanniopsida). All but five species were scored mainly from observed specimens: *Athalamia pinguis* Falc., *Austroriella salta* J. Milne & Cargill, *Geothallus tuberosus* Campb. (Marchantiidae), *Pellia epiphylla* L. Corda and *Lejeunea cavifolia* (Ehrh.) Lindb. (Jungermanniopsida). In total, the combined dataset comprised 56 species; 20 more taxa than the last combined analysis (Boisselier-Dubayle et al., 2002). Taxa and vouchers are listed on Table 1.

The morphological dataset comprised 126 characters from different sources. The present morphological matrix was initially constructed by extending that of Bischler (1998). Some characters scored at the phylum level by Crandall-Stotler and Stotler (2000) were also included. The original coding of nine of these “traditional” characters were modified (see *Character Definition* in the Supplementary material). Along with these traits, 66 completely novel characters were incorporated. Pegged rhizoids, for instance, were recently studied in depth by Duckett et al. (2014). Most of these characters, especially those that were now re-interpreted, were

included in a phylogenetic matrix for the first time. Similarly, characters derived from spores have been studied extensively in several taxa (Gupta and Udar, 1986) but infrequently used for phylogenetic analyses within this group; novel data from spores of Marchantiidae were considered in this study. Features representing continuous traits have seldom been considered in phylogenetic analyses of bryophytes. In cases where this sort of information was included, characters were commonly discretized losing valuable phylogenetic information (Farris, 1990; Goloboff et al., 2006). In the present study, 10 different variables were analysed as continuous characters (Goloboff et al., 2006). In order to incorporate the morphological variation among populations, a minimum of 10 specimens per species was examined. The final range of variation for each character was established as the mean value \pm standard deviation (SD). Ninety-two characters were scored from the gametophytic phase and 34 from the sporophyte. In conjunction, these included 12 cytological characters, nine developmental features, 18 sexual traits and 87 macroscopic characters. The morphological dataset presented 4540 more cells than the most extensive morphological dataset at the subclass level to date (1886 cells; Bischler, 1998). A detailed description of character, state definition and images can be found in the Supplementary material. The final dataset is available at Morphobank (Project: 2674; morphobank.org/permalink/?P2674).

Molecular data

The previous most comprehensive molecular sampling was carried out by Villarreal et al. (2016). The present study focused on extending outgroup sampling and filling gaps within the former analysis. Therefore, *psbA* and *rbcL* markers were sequenced for three species (*Plagiochila* sp. (Dumort) Dumort; *Radula voluta* Taylor ex Gottsche, Lindenb. & Nees and *Riccia* sp. L.). Amplifications were carried out at the Real Jardín Botánico de Madrid (Spain) following the conditions described by Forrest and Crandall-Stotler (2004; primers described in Table S1). Sequencing was conducted by MacroGen Inc. (Seoul). Sequences by Villarreal et al. (2016) were downloaded from GenBank. Nucleotides were aligned with MAFFT (Katoh and Toh, 2008) using default penalty settings. Ambiguously aligned positions were excluded from the analysis. Data were compiled using GB2TNT (Goloboff and Catalano, 2012), a pipeline to build molecular datasets from GenBank.

Phylogenetic analyses, constrained searches and topology comparisons

Although molecular data are commonly analysed by following model-based methods, the study of

Table 1
Taxa included in the analysis

Species	Locality: <i>Voucher</i>	Support literature	Genbank accession number														
			26S	<i>rbcL</i>	<i>rps4</i>	<i>psbA</i>	<i>psbT</i>	<i>atpB</i>	<i>rpoC1</i>	<i>cpITS</i>	<i>nad1</i>	<i>nad5</i>	<i>rps3</i>				
<i>Atichisoniella himalayensis</i> Kashyap	India: <i>Abnrad 346 (PC)</i>	(Bischler et al., 1994; Long et al., 2016b)	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–
<i>Aneura pinguis</i> (L.) Dumort	Argentina: <i>JR Flores 43</i>	(Hattori et al., 1966)	AY608195	AY688744	DQ983852	AY607921	JF513424	AY507346	–	AF033631	JF513358	AY688744	–	–	–	–	–
<i>Aporoebia nana</i> (S.Hatt. & Inoue) S.Hatt. & Mizut.			DQ460030	AB476552	DQ268983	AY877397	KJ590878	KJ590840	KJ590840	KJ590799	–	DQ268907	KJ590920	–	–	–	–
<i>Asterella tenella</i> (L.) P.Beauv.	Mexico: <i>Orricut 1562</i>	(Evans, 1920; Little, 1936; Sharp, 1939; Wittlake, 1954; Bischler, 1998)	KT793344	KT793556	KT793706	KT793458	KT793496	KT793374	KT793594	KT793739	–	KT793799	KT793650	–	–	–	–
<i>Athalamia pinguis</i> Falc.		(Sokhi and Mehra, 1973; Bischler, 1998; Boisselier-Dubayle et al., 2002; Crandall-Stotler et al., 2009; Rubasinghe, 2011)	DQ265774	DQ286003	DQ220677	DQ265747	KJ590880	KJ590842	KT793596	KT793741	KJ590801	DQ268912	KJ590922	–	–	–	–
<i>Austrorella salta</i> Milne & Cargill		(Cargill and Milne, 2013)	KT356898	KT356969	KT356979	KT356959	KT793498	KT356898	KT356990	KT356911	KT356946	–	–	–	–	–	–
<i>Blastia pusilla</i> L.	France: <i>Fesaloniez 123; Pierron 24057; Cuyvet 18486; Poirion s/n; Gardet s/n; Coppey s/n.</i>	(Duckett and Renzaglia, 1993)	AY688683	AF536232	AY507436	AY507477	KJ590881	AY507349	–	KT793742	KJ590802	AY688746	KJ590923	–	–	–	–
<i>Bucegia romanica</i> Radian	Hungary: <i>Boros s/n</i> Poland: <i>Szewykowski & Chudzinska 1002</i>		KJ590831	KJ590910	KJ590953	KJ590867	KJ590882	KJ590843	KT793598	KT793744	KJ590803	KT793801	KJ590924	–	–	–	–
<i>Cavicularia densa</i> Steph.			–	DQ268968	DQ268984	AY507479	KJ590883	KJ590844	KT356991	KT356914	KJ590804	DQ268913	KJ590925	–	–	–	–
<i>Clevea nana</i> (= <i>Athalamia hyalina</i>)	France: <i>Bischler et Lecerf 93614; Bonot & Pierron 604; Pierron, 3410; Skrzypczak & Skrzypczak 1226</i>		DQ265773.1	DQ286002	DQ220676	DQ265746	KJ590884	KJ590845	KT356992	KT356915	KJ590805	DQ268911	KJ590926	–	–	–	–
<i>Conocephalum conicum</i> (L.) Dumort	France: <i>Bischler et Baudoin 75137, 75175; Jover-Asi et Bischler 71734, 71738, 71264, 71747, 71882; Castelli s/n; Viteq s/n; Dismer s/n</i>		KT356888	KT356971	KT356981	KT356961	KT793502	KT356899	KT356993	KT356916	KT356948	KT793804	KT793654	–	–	–	–

Table 1
(Continued)

Species	Locality: Voucher	Support literature	Genbank accession number												
			26S	rbcL	rps4	psbA	psbT	atpB	rpoCl	cpITS	naal	nad5	rps3		
<i>Corsinia coriandrina</i> (Spreng.) Lindb.	France: Jover-Asi et Bischler, 71136-7, 71155, 71377, 71343, 71354, 71387, 71422-3, 71454, 71492, 71499, 71545, 74328; Bischler 74058, 74067, 74089, 74106, 74108, 74113, 74124, 74170, 74142, 74148, 74162, 74178	(Hassel de Menéndez, 1963; Bischler, 1998; Bischler et al., 2005)	KT793348	KT793560	KT793710	KT793462	KT793506	KT793381	KT793604	KT793750	KT793423	KT793806	KT793658		
<i>Cronisia fimbriata</i> (Nees) Whittier. & Bischl.	Bolivia: Fuentes 2841	(Bischler et al., 2005)	-	KT793562	-	KT793464	-	KT793383	-	KT793752	-	-	-		
<i>Cryptomerium tenerum</i> (Hook.) Underw.	Argentina: Hassel de Menéndez	(Hassel de Menéndez, 1963)	KT356889	KT356972	KT356982	KT356962	KT793507	KT356900	FJ173789	KT356918	KT356949	KT793807	KT793659		
<i>Cyathodium cavernarum</i> Kunze	Cabo Verde: Bolle s/n	(Bischler et al., 2005)	KT793349	KT793565	KT793712	KT793467	KT793509	-	KT793607	KT793756	KT793425	KT793809	KT793661		
<i>Dunortiera hirsuta</i> (Sw.) Nees	Argentina: JR Flores	(Hassel de Menéndez, 1963; Bischler et al., 2005)	DQ265780	DQ286009	DQ220683	DQ265753	KJ590888	KJ590849	KT356996	KT356920	KJ590809	DQ268920	KJ590931		
<i>Exornotheca pustulosa</i> Mitt.	France: Bischler 74077; Jover-Asi & Bischler 74609	(Bischler et al., 2005)	DQ265781	DQ286010	DQ220684	DQ265754	KJ590889	KJ590850	KT356997	KT356921	KJ590810	DQ268921	KJ590932		
<i>Fossombronia foveolata</i> Lindb.	Argentina: JR Flores s/n	(Haupt, 1920; Renzaglia, 1982; Crandall-Stotler et al., 2005)	AF226029	HQ446985	HQ447037	AY507482	-	AY507354	-	-	-	-	-		
<i>Frullania asagrayana</i> Mont.	Bolivia: Curchill et al. 23373 (mixed with <i>F. beauverdii</i>)	(Schuster, 1984a,b, 1992)	KX493338	FJ380822	-	FJ380657	-	-	-	-	-	-	-		
<i>Geothallus tuberosus</i> Camp.	German: Lehman s/n	(Campbell, 1896)	KJ590833	KJ59091	KJ590955	KJ590869	KJ590890	KJ590851	KT356998	KT356922	KJ590811	KT793817	KJ590933		
<i>Haplomitrium hookeri</i> (Sm.) Nees	Argentina: JR Flores 58	(Grubb, 1970; Crandall-Stotler et al., 2005)	DQ268882	-	AY608068	-	-	HQ412996	KT793615	-	KT793431	-	-		
<i>Jungmannia</i> sp. L.	Argentina: JR Flores 58	(Crandall-Stotler et al., 2005)	AY316351	AY507409	AY507493	AY149816	-	-	-	X00667	KF852564	AY688756	KF851659		
<i>Lejeunea confolia</i> (Ehrh.) Lindb.	Spain: Mañoz et al. s/n	(Schuster, 1980; Crandall-Stotler and Stotler, 2000)	DQ026545	-	DQ787470	AM396279	AY312077	DQ646043	-	-	-	AJ000701	-		
<i>Lepidoczia reptans</i> (L.) Dumort	Indonesia: Richards, P. s/n	(Schuster, 1969; Crandall-Stotler and Stotler, 2000)	AY608229	U87075	AY608083	AY607962	-	DQ646025	-	-	JF513380	AY608293	JF316404		
<i>Lophocolea heterophylla</i> (Schrad.) Dumort	Spain: Mañoz et al. s/n	(Schuster, 1969; Crandall-Stotler and Stotler, 2000)	DQ268889	DQ268932	DQ268987	DQ268999	-	DQ646034	-	AF033625	AY354959	DQ268932	KF942740		
<i>Lunularia cruciata</i> (L.) Lindb.	Argentina: JR Flores 11	(Hassel de Menéndez, 1963)	DQ265782	DQ286011	DQ220685	DQ265755	KT793519	KT356902	KT357000	KT356923	KT356951	DQ268933	KT793671		

Table 1
(Continued)

Species	Locality: <i>Voucher</i>	Support literature	Genbank accession number										
			26S	<i>rbcL</i>	<i>rps4</i>	<i>psbA</i>	<i>psbT</i>	<i>atpB</i>	<i>rpoC1</i>	<i>cpITS</i>	<i>nad1</i>	<i>nad5</i>	<i>rps3</i>
<i>Makinoa crispate</i> (Steph.) Miyake	Japan: <i>U. Faurie s/n</i>		AY877374	AY877390	AY877393	AY877399	KF852178	AY877395	–	–	JF513370	AY877386	KF851588
<i>Mammia californica</i> (Gottsche) L.C. Wheeler	Mexico: <i>McGregor & Rosario s/n</i>		GQ910726	KJ590913	KJ590956	KJ590870	KJ590893	KJ590852	KT357001	KT356924	KJ590814	KT793818	KJ590936
<i>Marchantia polymorpha</i> L. <i>Metzgeria furcata</i> (L.) Corda	Argentina: <i>JR Flores 18, 28</i> Bolivia: <i>Marko Lewis s/n</i>	(Renzaglia, 1982; Kawahara, 1986; Crandall-Stotler and Stotler, 2000)	KT356891	KT356974	KT356984	KT356964	KT793528	KT356903	KT357002	KT356925	KT356952	KT793822	KT793677
<i>Monocarpus sphaerocarpius</i> Carr.	Australia: <i>Fowcett s/n</i>		KT356886	KT356975	KT356985	KT356965	–	KT356904	KT357003	KT356926	KT356953	–	–
<i>Monoclea gottschei</i> Lindb.	Argentina: <i>JR Flores 8, 23</i>	(Hassel de Menéndez, 1963)	DQ265787	DQ286016	DQ220690	DQ265760	KJ590895	KJ590854	HQ026346	KT793777	KJ590816	DQ268943	KJ590938
<i>Monosolenium tenerum</i> Griffl. <i>Neohodgsonia mirabilis</i> (Pers.) Pers.	Japan: <i>Inoue 915; Deguchi 962</i> New Zealand: <i>Strethmann, H 51072</i>	(Crandall-Stotler, 1980; Bischler, 1998) (Bischler, 1998)	DQ265788	DQ286017	DQ220691	KJ590871	KJ590896	KJ590855	KT357004	KT356928	KJ590817	DQ268944	KJ590939
<i>Oxymitra incrassata</i> (Brot.) Sörgo & Sim-Sim	France: <i>Soitiaux et Soitiaux 1234</i>	(Hassel de Menéndez, 1963; Bischler, 1998; Bischler et al., 2005)	AY688692	AY507415	AY507456	AY507499	–	AY877396	–	KT793778	–	AY688759	–
<i>Pollavicinia lyellii</i> (Hook.) Gray	France: <i>Soitiaux et Soitiaux 1234</i>		KJ590834	KJ590914	KJ590957	KJ590872	KJ590898	KJ590856	KT357005	KT356930	KJ590819	KT793826	KJ590941
<i>Pellia epiphylla</i> L. Corda	AY734742	(Renzaglia, 1982; Crandall-Stotler and Stotler, 2000; Crandall- Stotler et al., 2005)	AY688693	AY688787	DQ463120	AY507502	–	DQ646048	–	–	JF513374	AY688761	KF851591
<i>Peltolopsis quadrata</i> (Saut.) Müll. Frib.	France: <i>Castelli 65001a, 66001, 67001</i>		AY688693	AY688787	AY507457	AY507502	KT793534	AY688823	–	JF513375	–	–	–
<i>Plagiochasma rupestre</i> (J.R. Forst. & G. Forst.) Steph.	Argentina: <i>JR Flores 27; Schiavone, 633</i>		KJ590835	KJ590915	KJ590958	KJ590873	KJ590900	KJ590858	FJ173796	KT793779	KJ590821	KT793828	KJ590943
<i>Plagiotechila</i> sp. (Dumort)	Argentina: <i>JR Flores 54</i>		–	KY985367	–	MF036173	–	–	–	–	–	–	–
<i>Pleurozia purpurea</i> Lindb.	France: <i>Alorge s/n; Jeanperi s/n; Durand s/n</i>		DQ268899	AY877391	AY608100	AY877401	–	HQ412997	–	–	AY607889	DQ268951	–
<i>Preissia quadrata</i> (Scop.) Nees	Argentina: <i>JR Flores 51</i>		DQ265793	KJ590916	KJ590959	KJ590874	KJ590901	KJ590859	KT357008	KT356933	KJ590822	DQ268953	KJ590944
<i>Raddia voluta</i> Taylor ex Gottsche, Lindenb. & Nees	Argentina: <i>JR Flores 51</i>		–	–	HQ447037	MF036172	–	–	–	–	–	–	–

Table 1
(Continued)

Species	Locality: Voucher	Support literature	Genbank accession number											
			26S	rbcL	rps4	psbA	psbT	atpB	rpoC1	cpITS	naaI	naa5	rps3	
<i>Reboulia hemisphaerica</i> (L.) Raddi	France: Bischler74132, 74128, 74110, 74626; Jovet-Ast et Bischler 74562	(Hassel de Menéndez, 1963)	KJ590836	KJ590917	KJ590960	KJ590875	KJ590902	KJ590860	KT357010	KT356935	KJ590823	–	–	KJ590945
<i>Riccia</i> sp. (cf. <i>paraguayensis</i>) L.	Argentina: JR Flores 49	(Hassel de Menéndez, 1963)	–	KY985366	–	MF036171	–	–	–	–	–	–	–	–
<i>Riccia fluitans</i> L.	France: D Gabriel s/n	(Underswood, 1894; Sharp, 1939; Jovet-Ast, 1986; Singh, 2014)	DQ265795	DQ286023	DQ220696	DQ265766	KJ590903	KJ590861	KT793634	KT356936	KJ590824	DQ268956	DQ268956	KJ590946
<i>Ricciocarpos natans</i> (L.) Corda	Uruguay: Suarez et al. 1317	(Hassel de Menéndez, 1963)	DQ265796	DQ286024	DQ220698	DQ265767	KJ590904	KJ590862	KT793637	KT356937	KJ590825	DQ268958	DQ268958	KJ590947
<i>Riccia helicophylla</i> (Bory & Mont.) Mont.	Algeria: <i>Trabat</i> s/n; <i>Durieu de Maisonneuve</i> s/n	(Montagne, 1852)	KJ590837	KJ590918	KJ590961	KJ590876	KJ590905	KJ590863	KT357011	KT356938	KJ590826	KT793832	KT793832	KJ590948
<i>Sauteria alpina</i> (Nees) Nees	Austria: P Geissler 17087 Italy: Bischler 866 Poland: Siewekowski & Chudzinska 1001	(Rubasinghe, 2011; Borovichev et al., 2012)	DQ265797	DQ286025	DQ220699	DQ265768	KJ590906	KJ590864	KT357012	KT356939	KJ590827	DQ268960	DQ268960	KJ590949
<i>Sphaerocarpos texanus</i> Aust.	Morocco: Jovet-Ast a,b s/n	(Haynes, 1910)	AY688697	AY507425	AY507466	AY507511	KT793545	AY507382	KT357015	KT356942	KT356956	AY688771	AY688771	KT793696
<i>Stephensoniella brevipedunculata</i> Kashyap	Himalaya: Chopra 500	(Kashyap, 1914a,b, 1915)	–	KT793584	–	KT793486	–	KT793410	–	KT793789	–	–	–	–
<i>Symphogyna undulata</i> Colenso	Chile: JR Flores 26, 50	(Crandall-Stotler et al., 2005)	AY877380	AY688790	AY688804	AY688835	–	AY688825	–	–	–	AY688773	AY688773	–
<i>Takakia ceratophylla</i> (Mitt.) Grolle	Nepal: Long 17247	(Grubb, 1970; Renzaglia et al., 1997)	HM751509	GU295867	AY908023	JF513404	AB254135	DQ646093	–	AF426188	AY354937.1	DQ268963	DQ268963	–
<i>Targionia hypophylla</i> L.	Argentina: JR Flores 56, 48; Schiavone 2417	(Hassel de Menéndez, 1963; Bischler et al., 2005)	KJ590839	AY507427	AY688805	AY507514	KJ590908	AY507384	FJ173802	KT793790	KJ590829	DQ268965	DQ268965	KJ590951
<i>Trenbia</i> sp. K.I. Goebel	Samoa: F. Reinecke 17	(Schuster and Scott, 1969; Schuster, 1984a, b,a)	AY688699	AY507428	AY507468	AY507515	KT793547	HQ412995	KT793639	–	JF973315	AY688774	AY688774	JF973315
<i>Wiesnerella demodata</i> (Mitt.) Stephani	Japan: Akiyama & Hiraoka 1; Akiyama 13013, Inoue 875, Saire & Toyota 3 Indonesia, Java: Verdoorn s/n Nepal: Long 17247 La Reunion: Bischler 89021; Gimalac 1703, 1706, 12569		DQ265799	DQ286027	DQ220701	–	KJ590909	KJ590866	KT793640	KT356944	KJ590830	DQ268966	DQ268966	KJ590952

Voucher and supporting literature refer to the taxa included in the morphological dataset. New sequences are marked in bold type.

morphology has been based largely on parsimony. In contrast with molecular data, the use of probabilistic models to evaluate morphology has continually led to undesirable results (Goloboff and Pol, 2005; Beck and Lee, 2014; Goloboff et al., 2017). For instance, Mk models have recently been shown to be particularly sensitive to the high heterogeneity rate present in morphological datasets, resulting in spurious groupings and imprecise dating (Steiper and Seiffert, 2012; Beck and Lee, 2014). These outcomes still generate a great reluctance amongst many authors to adopt models as a mean to analyse morphological data. Weighted parsimony, however, has been shown to improve the analysis of morphological data (Goloboff et al., 2008a,b) and has acquired a key role in combined phylogenetic analyses (e.g. Mirande, 2017). As the principal premise of this paper is to achieve a phylogenetic hypothesis based on a total-evidence approach, searches were performed under weighted parsimony as the optimality criterion.

Therefore, the main systematic and taxonomic conclusions of the present study are derived from the analysis of the combined dataset under extended implied weighing, weighting blocks in accordance to their average homoplasy (Goloboff, 2014). The blocks represented each gene and the morphological dataset.

Group sensitivity and searches. In order to assess group stability, searches were conducted across several weighting schemes for individual partitions and the combined dataset. In addition to the searches under block weighting (BW), an alternative search strategy involved weighting each character according to its homoplasy but taking into account the proportion of missing entries (IW_M). In IW_M , each missing entry was assumed to have half of the homoplasy of the observed entries ($P = 0.5$). In searches under both IW_M and BW, the TNT default concavity value was used ($K = 3$; $K3$). Three concavity values for standard implied weighting were also explored ($K3$, $K5$ and $K7$; Goloboff, 1993). Finally, the data also were analysed under equal weighting (EqW). All weighting settings were applied to the individual partitions and the combined data (except BW in the case of the morphological dataset). Hence, a total of 17 different analyses were conducted (six for combined data, six for the molecular dataset and five for morphological dataset). In all cases, analyses were carried out in TNT 1.5 (Goloboff et al., 2008a,b; Goloboff and Catalano, 2016) with new technology searches using 10 RAS as starting point. For each replicate, 10 iterations of Ratchet (Nixon, 1999) and Tree Drifting (Goloboff, 1999) were performed, ending the search when after the best score was hit seven times. Support values were estimated by Symmetric Resampling (SR) considering

the difference between the most frequent group and the most frequent contradictory group (GC; Goloboff et al., 2003). Additional estimates of support (Bootstrapping and Bremer) were also calculated and are available in Fig. S1.

Topological comparisons. In order to assess the contribution of each source of information to the final result, the topologies recovered from different data types were compared in terms of (i) SPR (subtree pruning and regrafting) distance (Goloboff, 2008) and (ii) number of shared nodes. As a single SPR rearrangement may involve a movement of a few or several nodes, each movement was weighted by a constant value of 5 (Goloboff, 2008). The morphological trees obtained under each weighting condition were compared to all the molecular trees. In turn, the molecular trees were compared to the combined tree. The number of SPR moves and the number of shared nodes were also calculated for the morphological trees obtained by Bischler (1998) and Boisselier-Dubayle et al. (2002). In addition, the same comparison was performed to discern the relative contribution of gametophyte and sporophyte characters. To compare our morphological tree with Bischler (1998) and Boisselier-Dubayle et al. (2002) trees, evaluations involved two different trees as reference: (a) our molecular tree obtained under BW and (b) Villarreal et al.'s (2016) molecular tree. Regarding our two sets of morphological characters, comparisons only used our molecular tree as reference.

Constrained searches. In order to quantify the suboptimality of the groups not retrieved in the best trees, constrained searches were carried out in the combined data under BW. Understanding suboptimality in terms of fit (F) is hardly intuitive. Therefore, the fit difference between the constrained and the unconstrained trees was translated into the *number of mean homoplastic characters with an extra step* required to explain the suboptimal tree (\bar{x}_{+1} ; Carrizo and Catalano, 2015). The estimation of character with mean homoplasy (\bar{x}) was obtained as the tree length divided by the number of characters. The ratio between the topology fit differences ($F_{\text{unconstrained}}$, $F_{\text{constrained}}$) and mean character fit differences (\bar{x} , \bar{x}_{+1}) allowed suboptimality to be conceived in terms of the number of \bar{x}_{+1} [$(F_{\text{unconstrained}} - F_{\text{constrained}}) / (F_{\bar{x}} - F_{\bar{x}_{+1}})$].

Character reconstruction, synapomorphies and diagnosis

Group diagnoses were determined by mapping morphological characters onto the combined tree obtained under BW. In addition, the role of different sets of characters to diagnose specific clades was addressed.

Because sporophytic characters have commonly been viewed as having no taxonomic information at lower levels (Schuster, 1984a,b; Crandall-Stotler and Stotler, 2000; Crandall-Stotler et al., 2009), special attention was paid to those characters. In particular, the proportion of synapomorphies derived from sporophytic characters was quantified for each node. To assess whether sporophytic characters diagnose taxonomic groups below the class level, the number of sporophytic synapomorphies per taxonomic category was also measured.

Phylogenetic searches, sensitivity assessment and diagnosis evaluation were implemented in scripts using TNT macro language. These are available upon request.

Life history patterns and taxonomic membership

Bischler (1998) hypothesized that life-history traits were phylogenetically correlated in Marchantiidae and, consequently, morphological similarity was caused by common ancestry. In order to test these hypotheses in light of the new data, the four patterns found by Bischler (named Groups I–IV) and the putative associated characters were mapped onto the final tree. Given their relevance in liverwort biology (Bischler et al., 2005), four features were optimized: branch length, number of spores per capsule, spore size and capsule dehiscence. Branch length and spore size were scored as continuous characters; capsule dehiscence was scored as a binary character (*cleistocarpous/noncleistocarpous*). The number of spores per capsule was scored as a multistate character: *less than 500 spores* (0), *more or equal to 1000 and less than 10000* (1) and *more than 10000 spores* (2).

Results

Combined Tree under BW and Constrained Searches

The combined dataset analysed under extended implied weighting (BW) produced a single fully resolved tree (Fig. 1). It was largely congruent with recent phylogenetic analyses (Forrest et al., 2015; Long et al., 2016a,b).

The major groups (Haplomitriopsida, Jungermanniopsida and Marchantiopsida) were recovered as monophyletic with high support values (≈ 100 ; Fig. 1), as well as the sister relationship among the four orders within Marchantiidae. Some of these nodes had already been found in previous studies but have remained without a formal recognition (Crandall-Stotler et al., 2009; Forrest et al., 2015). To facilitate descriptions, some of these nodes are referred to as clades A–F. Clade A was rendered as the association between the genus *Monocarpus* and the order Sphaerocarpaceae (*Sphaerocarpus*, *Geothalpus*, *Austriella* and *Riella*) with strong evidence (SR 98,

Fig. 1); the monophyly of Sphaerocarpaceae s.s. received more moderate support (SR 72). As in recent molecular phylogenies (Forrest et al., 2015; Villarreal et al., 2016), the previous definition of Marchantiales was not supported. Clade B was established upon the node constituted by *Lunularia* (Lunulariales) and the remaining genera of Marchantiales except *Monocarpus*. This clade with a moderate (to high) support value (SR 88), was recovered as the sister group to Clade A.

Mid-level nodes within Clade B were recovered in most cases with low support. The node F (*Riccia* + *Ricciocarpos* + *Oxymitra*) + E (*Monoclea* + *Conoccephalum*) was the sister clade to the monophyletic Aytoniaceae (*Asterella* + *Cryptomitrium* + *Mannia* + *Plagiochasma* + *Reboulia*). The poorly supported clade *Monosolenium* + *Cyathodium* was related to the highly supported Clade D (*Exormotheca* + *Aitchisoniella* + *Stephensiella* + *Corsinia* + *Cronisia*). Clade C (*Wiesnerella* + *Targionia*) was sister to both the remaining morphologically simple groups and Aytoniaceae (Fig. 1). In contrast, the less inclusive nodes between families tended to be well supported. Clades C, D, F and E received support values from 82 to 100 (Fig. 1).

At the family level, Cleveaceae and Corsiniaceae (Long et al., 2016a) were both nonmonophyletic because *Aitchisoniella* was nested within the latter group. The genera *Cronisia* and *Corsinia* formed a moderately supported clade (SR 76; Fig. 1) with *Exormotheca* as their sister taxa (SR 64; Fig. 1). *Stephensiella* constituted a highly supported clade with *Aitchisoniella* (SR 100; Fig. 1). The clade formed by *Sauteria*, *Clevea*, *Peltolepis* and *Athalamia* was also highly supported (SR 100; Fig. 1). The other nonmonotypic families were retrieved as monophyletic with general high support.

Phylogenetic searches forcing the monophyly of groups absent from the best trees led to widely different suboptimality values depending on the considered taxa. On the one hand, ingroup families were highly suboptimal regarding the BW combined tree. The monophyly of Corsiniaceae entailed a difference of 167.1 mean homoplastic characters gaining a further step. As for Cleveaceae, its monophyly involved 172.2 mean characters gaining an extra step. Comparatively, outgroup taxonomic groups implied less than one step in a character with mean homoplasy (0.2–0.6). Therefore, there was no evidence to discard the monophyletic status of outgroup taxa whereas the monophyly of the ingroup families was considerably questioned.

Partitioned analyses

Molecular dataset. The strict consensus of the molecular trees recovered across the entire analytical conditions had 34 nodes (out of 54; Fig. 2). Molecular trees recovered Jungermanniopsida and Marchantiopsida as monophyletic. Inside

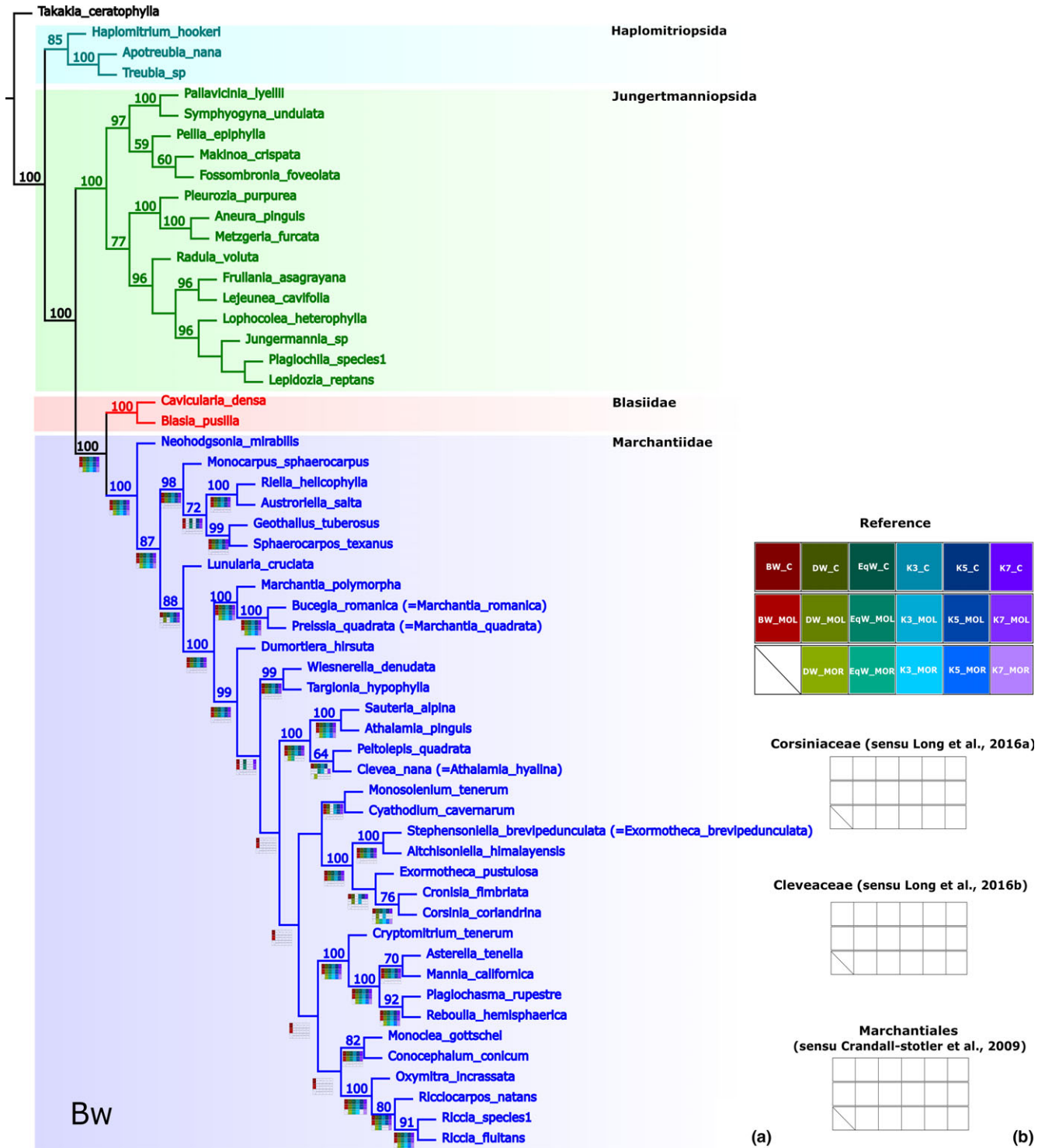


Fig. 1. (a) Combined tree under extended implied weighting (Block weighting). Symmetric Resampling values are indicated above branches (only > 50 values are shown; see additional support measures in the Fig. S1 in the Supplementary material). Sensitivity plots for ingroup nodes monophyly are shown below branches. (b) Sensitivity plots for Cleveaceae, Corsiniaceae and Marchantiales [as defined by Long et al. (2016a,b) and Crandall-Stotler et al. (2009)], two groups that were not monophyletic in the combined analysis under block weighting. Sensitivity plot reference: coloured squares represents monophyly. C = combined data, MOL = molecular data, MOR = morphological data. Extended implied weighting settings: block weighting (BW) and character weighted considering their homoplasy and number of missing entries (IWM). K3, K5 and K7 are concavities values explored under standard implied weighting. EqW refers to equal weighting. [Colour figure can be viewed at wileyonlinelibrary.com].

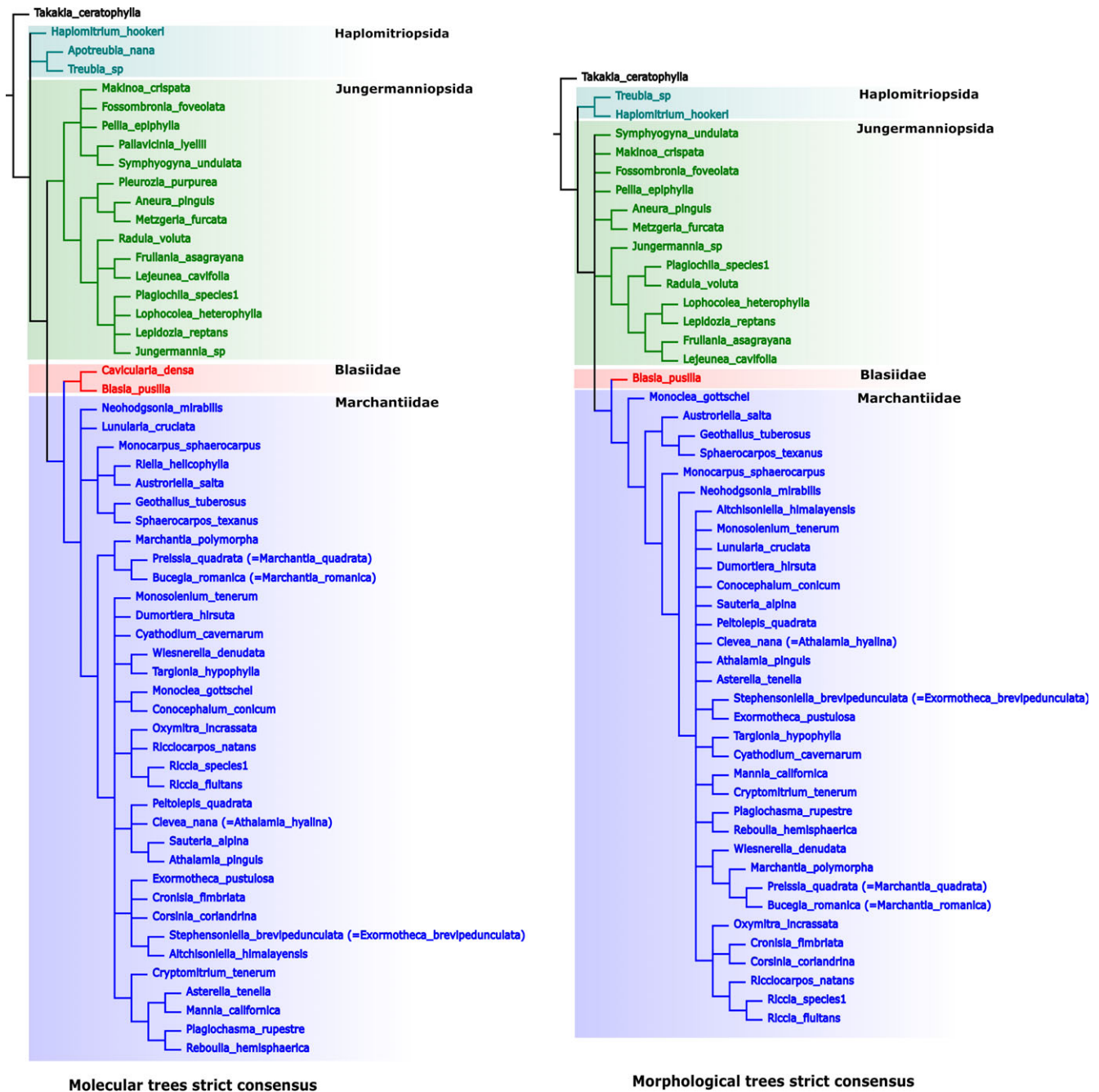


Fig. 2. Strict consensus of the molecular (left) and morphological trees (right) obtained under the different weighting schemes. [Colour figure can be viewed at wileyonlinelibrary.com].

Marchantiopsida, relationships were poorly resolved. The association between the orders Lunulariales or Neohodgsoniales, and the remaining orders was unresolved. The family Marchantiaceae was sister to the remaining Marchantiales. The rest of the families within Marchantiales were placed in a nine-way polytomy. Clades with high support values in the combined tree under BW were also recovered in the analysis of the molecular partition. The genera *Aitichisoniella* and

Stephensoniella were sister taxa within a clade constituted by the remaining genera of Corsiniaceae. Cleveaceae was not recovered as monophyletic. Searches under BW concluded in a single fully resolved tree, highly similar to the combined tree (see below under Relative data contribution).

Morphological dataset. The strict consensus of the morphological trees obtained across the entire

analytical conditions had 29 nodes (out of 49; Fig. 2). The order Marchantiales (in the sense of Crandall-Stotler et al., 2009) was not recovered as monophyletic. *Monoclea* was found to be closely related to Sphaerocarpaceae and *Lunularia* was nested within Marchantiales. *Monocarpus* was the sister taxon of the remaining Marchantiales. *Neohodgsonia* was not recovered inside Marchantiaceae. The rest of the taxa were placed within a 16-way polytomy. The genus *Aitchisoniella* was not grouped with the rest of the family Cleveaceae although its association with other families was unclear. The other members of Cleveaceae did not constitute a monophyletic group. Corsiniaceae was also nonmonophyletic. The genera *Exormotheca* and *Stephensiella* were sister taxa whereas *Corsinia* + *Cronisia* were related to Ricciaceae and Oxymitraceae. Searches under standard implied weighting (K7) led to a single fully resolved tree which maximized congruence with both the combined data and the molecular data (see below under Relative data contribution).

Sensitivity analysis

About two-thirds of the ingroup clades (22 of 36) were not affected by different weighting schemes (Fig. 1). Although it was not a strict pattern, clades with high support values tended to be less sensitive to parameter variation. Unsupported nodes (SR < 50) were nonmonophyletic in 11 to 15 weighting schemes. Weakly to moderately supported clades (SR 50–90) were not monophyletic in two to seven weighting schemes. Highly supported groups (SR > 90) were not monophyletic in five weighting conditions or less. Two exceptions to this pattern were Sphaerocarpaceae (SR 72) and Oxymitraceae + Ricciaceae (SR 100), which were nonmonophyletic in nine analytical conditions. The genus *Exormotheca* behaved as a wildcard taxon, alternating its position with members of Corsiniaceae. The most inclusive nodes within Marchantiales (i.e. excluding Marchantiaceae and Dumortieraceae) were extremely sensitive. Moreover, such nodes were only recovered in two analytical conditions (molecular and combined data under BW; SR < 50). However, lower level clades within Marchantiales were recovered under different weighting schemes. The clades Monocleaceae + Conocephalaceae, Exormothecaceae + Corsiniaceae and Monosoleniaceae + Cyathodiaceae were markedly stable (10 to 12 schemes). The genus *Aitchisoniella* constituted a stable clade with *Stephensiella* in 12 weighting schemes.

Nodes of the morphological trees were more sensitive to parameter variation than molecular trees (sensitivity plots in Fig. 1). Five families, and relationships inside them, were resolved by the morphological data (Ricciaceae, Aytoniaceae, Corsiniaceae, Cleveaceae

and Marchantiaceae). Taxa relationships inside Sphaerocarpaceae were considerably more stable in the molecular data.

Relative data type contribution

Trees derived from molecular data were highly similar to the BW combined tree. In particular, the BW molecular tree was the most similar to the BW combined cladogram (1.2 SPR; Fig. 3). The molecular tree obtained under BW also shared 48 nodes with the combined tree. Among morphological trees, the tree obtained considering a concavity value of seven (K7) was the most similar to the BW molecular tree (8.1 SPR moves and 17 shared nodes).

Regardless of the reference tree (our BW molecular tree or Villarreal et al.'s (2016) tree), the comparisons between the K7 morphological tree and Bischler (1998) and Boisselier-Dubayle et al. (2002) trees yielded lower SPR values for our morphological tree (Table 2). Bischler's (1998) morphological tree involved 9.1 and 9.2 SPR moves to the BW molecular tree and Villarreal et al.'s (2016) tree, respectively. Boisselier-Dubayle et al.'s (2002) tree entailed 9.5 SPR moves to the BW molecular tree and 10.7 SPR moves to Villarreal et al.'s (2016) tree.

In terms of shared nodes, values varied depending on the reference tree (Table 2). The number of shared nodes was slightly higher for Bischler's tree when our BW molecular tree was used as reference (18 shared nodes for Bischler's tree; 17 shared nodes for the K7 tree and Boisselier-Dubayle et al.'s tree). In contrast, the number of shared nodes was higher for the K7 morphological tree when Villarreal et al.'s tree was used as reference (18 shared nodes for our K7 tree; 17 shared nodes for Bischler's and Boisselier-Dubayle et al.'s trees).

The gametophyte tree was more similar to the BW molecular tree than the sporophyte tree. In terms of shared nodes, the single gametophytic tree had 15 common nodes with the molecular tree and an SPR distance of 8.38 weighted movements (Fig. 4). The analysis including only sporophytic characters produced 100 optimal trees which shared eight nodes with the molecular tree and had an average SPR distance of 12.8 weighted movements.

Conflict and contribution of data sources were also reflected in the support values. Despite the wide differences among molecular trees and morphological trees, average support values were improved in the combined analysis in comparison with the molecular-only analysis. In particular, five clades of the ingroup had higher support values (Figs 2 and 4). The mean SR support value was 70 for the BW combined tree, whereas it was 65.7 for the BW molecular topology.

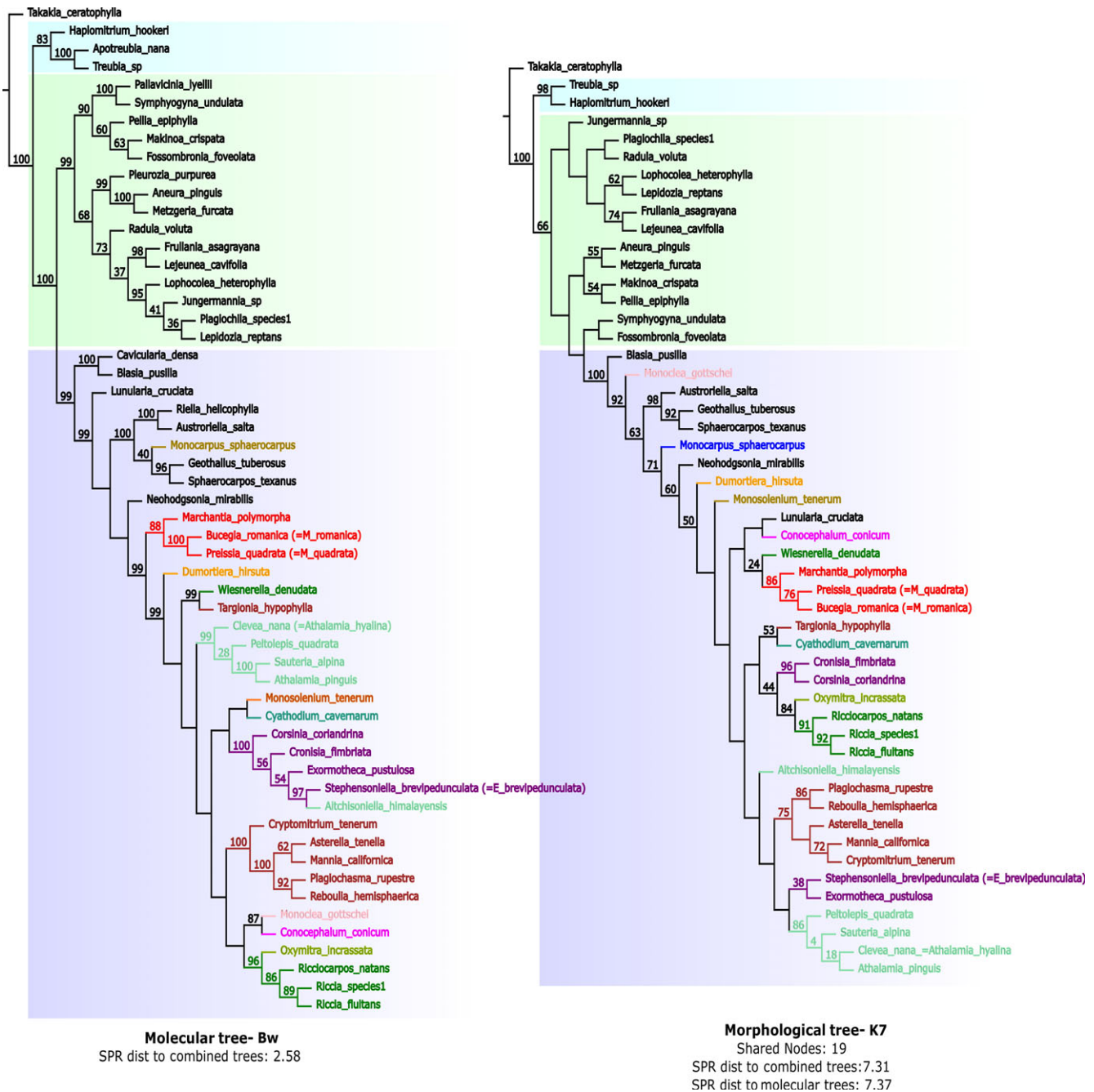


Fig. 3. Molecular tree under extended implied weighting (Block weighting) and morphological tree under implied weighting ($K = 7$) which maximised similarity among data types. Weighted SPR movements to the reference tree and shared nodes are indicated below each topology. [Colour figure can be viewed at wileyonlinelibrary.com].

Synapomorphy mapping and diagnosis assessment

Mapping morphological characters on the combined tree allowed determination of synapomorphies for 26 out of 35 ingroup nodes. The class Marchantiopsida and seven high-level clades inside it had high to moderate support values and morphological apomorphic characters (Fig. 5; Table 3). Marchantiopsida and Marchantiidae were both supported by nine

synapomorphies whereas node B had 10 synapomorphic characters (being diagnosed for the first time). Clade F was supported by seven synapomorphies; two continuous characters and five discrete characters. The Clade E was supported by a single spore trait, germ tube absent. Two characters were diagnostic for Clade D. Three characters were synapomorphies for both clades C and A. Descriptions of these specific groups are in Table 3, a detailed list of

Table 2

Comparison among different morphology-based trees (our morphological tree obtained under implied weighting, and Bischler’s and Boisselier-Dubayle et al.’s trees). The tree obtained in the present study under K7 implied weighting maximized similarity with reference trees in three out of four cases (bold). Bischler’s tree maximized the number of shared nodes with the BW molecular tree (bold)

	SPR moves to the BW molecular tree	SPR moves to Villarreal et al.’s tree	Shared nodes with the BW molecular tree	Shared nodes with Villarreal et al.’s tree
K7 morphological tree	8.1	7.8	17	18
Bischler’s morphological tree	9.1	9.2	18	17
Boisselier-Dubayle et al.’s morphological tree	9.5	10.7	17	17

BW, block weighting; SPR, subtree pruning and regrafting.

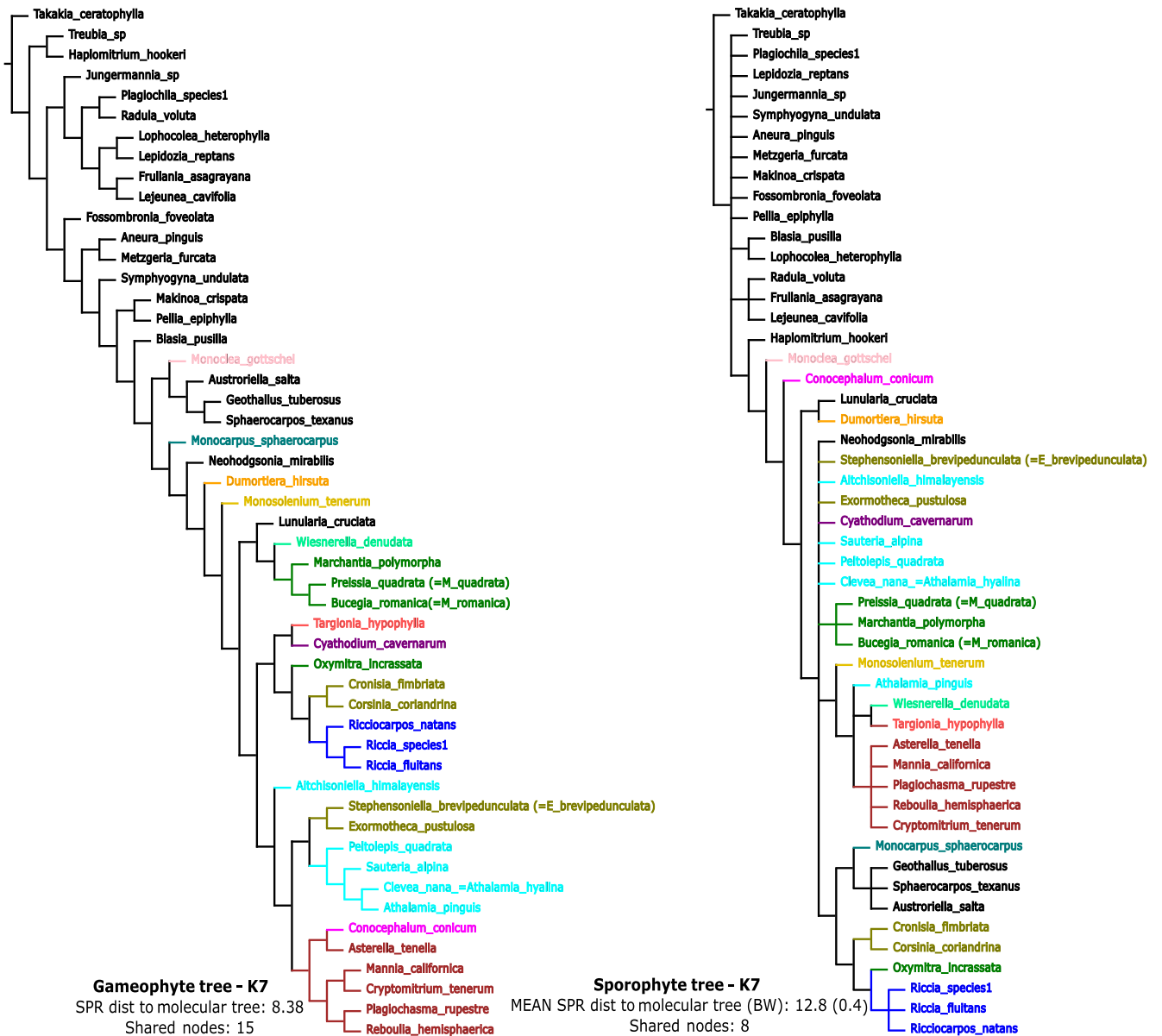


Fig. 4. Morphological trees derived from gametophytic characters and sporophytic characters under implied weighting ($K = 7$). Families within Marchantiales are highlighted. SPR values and shared nodes with the molecular tree are indicated below each tree. For sporophytic tree, the average SPR value is provided (SD in parenthesis). [Colour figure can be viewed at wileyonlinelibrary.com].

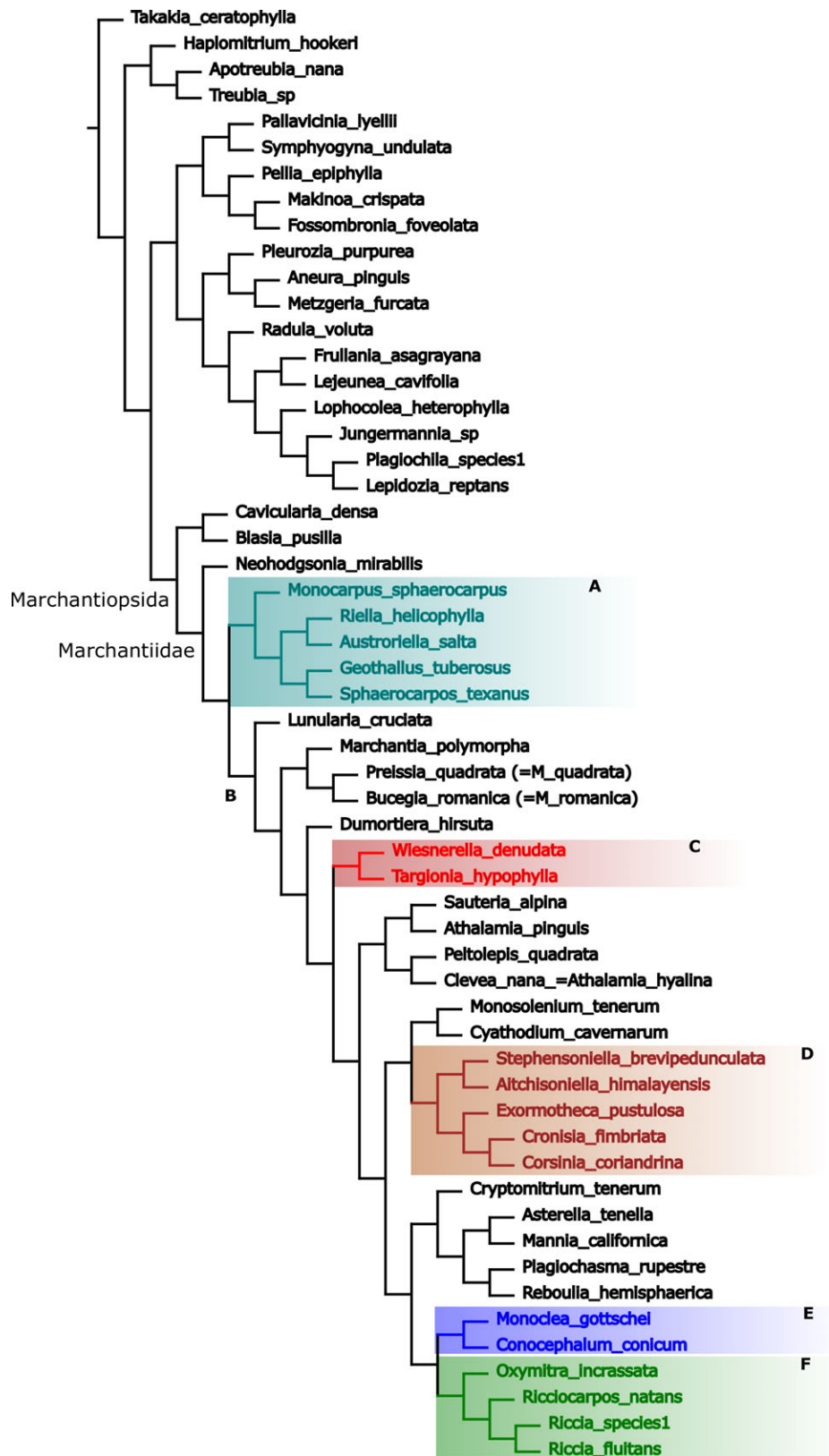


Fig. 5. Groups with improved diagnoses (Combined tree under BW). A detailed diagnosis for each group is in Table 3. The selected inter-family level nodes were those groups simultaneously well supported and diagnosed. [Colour figure can be viewed at wileyonlinelibrary.com].

Table 3
Synapomorphies for mid- and high-level groups within Marchantiidae and Marchantiopsida

Clade name	Taxa included	Character number: synapomorphies
F	Ricciaceae [<i>Riccia</i> + <i>Ricciocarpos</i>]+ Oxymitraceae [<i>Oxymitra</i>]	2: ratio apex width/base width: 1.1–1.5 -> 1.8–2.2 3: pegged rhizoids/smooth rhizoids rate: 0.6–1.0 -> 0.35–0.45 108: capsule dehiscence: irregular -> cleistocarpous 111: elaters: present -> absent 53: perigonia position: embedded in receptacles -> in anacrogynous clusters 95: embryo ontogeny: foot and seta from hypobasal cell -> sporangium from all of it 69: antheridia stalk anatomy: four or more seriate -> uniseriate 125: germ tube: present -> absent
E	Monocleaceae [<i>Monoclea</i>] + Conocephalaceae [<i>Conocephalum</i>]	
C	Wiesnerellaceae [<i>Wiesnerella</i>] + Targioniaceae [<i>Targionia</i>]	2: ratio apex width/base width: 1.0–1.1 -> 0.7–0.9 7: diameter pegged rhizoids/smooth rhizoids: 0.7 -> 0.76–0.77 105: capsule internal wall thickenings: annular -> semi-annular
D	Corsiniaceae [<i>Cronisia</i> + <i>Corsinia</i>] + Exormothecaceae [<i>Exormotheca</i> + <i>Aitchisoniella</i> + <i>Stephensiella</i>]	53: perigonia position: embedded in receptacles -> in anacrogynous clusters 27: drought tolerance: no -> yes
A	Monocarpaceae [<i>Monocarpus</i>] + Sphaerocarpaceae [<i>Sphaerocarpus</i> + <i>Austroriella</i> + <i>Riella</i> + <i>Geothallus</i>]	108: capsule dehiscence: into four valves -> cleistocarpous 111: elaters: present -> absent 120: spore, proximal gross morphology: as in distal face -> different from distal face
Marchantiopsida	Blasiidae [<i>Blasia</i> + <i>Cavicularia</i>] + Marchantiidae [Neohodgsoniales + Sphaerocarpaceae + Lunulariales + Marchantiales]	17: phyllotaxi: one-third/one half -> none 34: ventral appendages: absent -> two rows 117: polarized spores: no -> yes 89: outer perichaetium: leaf-like scales -> marchantioid involucre 92: calyptra: shoot calyptra -> true calyptra 90: Position of involucre: none -> dorsal 75: spermatid spline aperture: 1-tubule aperture- -> 3-tubule aperture or more 77: spermatid, basal body dimorphism: no -> yes 82: spermatid, notch presence: no -> yes
Marchantiidae	Sphaerocarpaceae(including <i>Monocarpus</i>) + Lunulariales + Marchantiales + Neohodgsoniales	104: capsule wall: multistratose -> unistratose 116: spore mother cell lobed: present -> absent 125: germ tube: absent -> present 11: protonema: globose -> flattened 28: air chambers: absent -> Marchantia type 60: archegoniophore, stalk anatomy: none -> homogeneous 62: female receptacle anatomy: none -> homogeneous 106: capsule wall thickenings: absent -> present 64: archegonia neck cells (CT): five-cells row -> six-cells row
B	Lunulariales [<i>Lunularia</i>] + Marchantiales	9: female branch length: 0.4–0.5 -> 1.2 36: ventral appendage shape: acute -> lanceolate 43: rhizoids type: unicellular, smooth only -> unicellular, dimorphic 44: typed of pegged rhizoids: none -> slightly pegged rhizoids only 45: pegs: nonpegged -> blunt to pointed 47: pegged rhizoids, distribution throughout the plant: none-pegged -> ventrally distributed 94: embryo type: filamentous -> octant 38: scales appendage number: without scales -> one 39: scales margin: no scales -> entire 42:appendage cell form: no scale -> hexagonal or pentagonal

synapomorphies for each node in the phylogeny is in Table S2).

Within the entire phylogeny, continuous characters diagnosed 18 clades (16 within Marchantiopsida). Additionally, 18 clades were diagnosed by at least one sporophytic feature. In ten out of these 18 nodes, 50% or more of the diagnostic traits were derived from the sporophyte (Fig. 6; Fig. S2).

The distribution of this proportion per taxonomic range was observed to be higher for orders than for classes or subclasses. The inclusion of sporophytic characters into the diagnosis at the interfamilial level was also considerably high. In fact, order and interfamilial nodes received the highest sporophyte contribution in terms of synapomorphies (Fig. 6; Fig. S2).

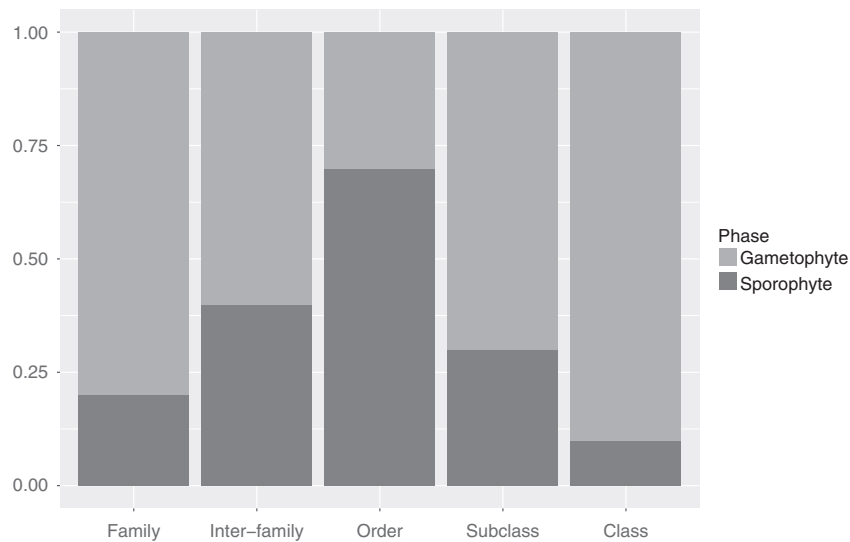


Fig. 6. Percentage of synapomorphies per ontogenetic phase per taxonomic category. C = class; SC = subclass; O = order; F = family and X = inter-family. Proportion of sporophytic synapomorphies mapped along the combined tree is included as Fig. S2 in the Supplementary material.

Mapping life-history traits

None of the ecological groups defined by Bischler (1998) had a single origin (Fig. 7). Group II was the basal group along Marchantiidae, whereas groups III and IV characterized distant nodes to Sphaerocarpaceae. Group I was polyphyletic in three different clades (Fig. 7). Capsule dehiscence was reconstructed with 13 steps; the cleistocarpous state diagnosed Sphaerocarpaceae + *Monocarpus*, Oxymitracaceae + Ricciaceae and *Corsinia* + *Cronisia*. The optimization of the number of spores per capsule required nine steps. The continuous characters (spore size and branch length), had 1.5 and 2.15 steps each. Spore size was largest among morphologically simple groups (Oxymitracaceae + Ricciaceae + Aytoniaceae + Conocephalaceae + Monocleaceae), whereas smaller sizes were concentrated in the basal clades. Conversely, gametophyte branch length showed lower values in younger clades and higher values among deep nodes. The number of spores per capsule tended to be lower at distant groups and in the basally placed group A (Fig. 7).

Discussion

In the current work, we present the results of the largest combined analysis for the complex thalloid liverworts. Novel sources of morphological information were evaluated including continuous and structural characters; features used in previous studies were reinterpreted. The results obtained from the combined data corroborate many of the recent proposals but cast doubt on the monophyly of some families. The

relative contribution of different types of morphological characters indicate that sporophytic characters provide considerable information for low taxonomic levels. The exhaustive morphological character sampling allows improvement of the analysis in terms of data congruence, sheds light on phylogenetic relationships among families, and improves the diagnoses of high-level groups. The main taxonomic and systematics conclusions presented below are derived from the combined tree under BW. A discussion on group stability and character contribution derives from the results of the partitioned analyses.

Monophyly and stability of groups

Most of the deep relationships within the subclass Marchantiidae are in agreement with previous studies (Wheeler, 2000; Boisselier-Dubayle et al., 2002; Forrest et al., 2006; Villarreal et al., 2016). The order Marchantiales, as conceived by Crandall-Stotler et al. (2009) or Bischler (1998), is not retrieved as monophyletic in any of our analyses (Fig. 1). After finding a close association between *Monocarpus* and Sphaerocarpaceae, Forrest et al. (2015) and Villarreal et al. (2016) rejected Marchantiales *sensu* Crandall-Stotler et al. (2009), which agrees with our finding of a well-diagnosed Clade A (*Monocarpus* + Sphaerocarpaceae; Figs 1 and 5).

The present analyses recovered several interesting groups above the family level (Figs 1 and 5). The earlier removal of the genus *Lunularia* from Marchantiales led to the proposal of the order Lunulariales (Long, 2006; Crandall-Stotler et al., 2009). Relationships among high-level taxa remained unresolved

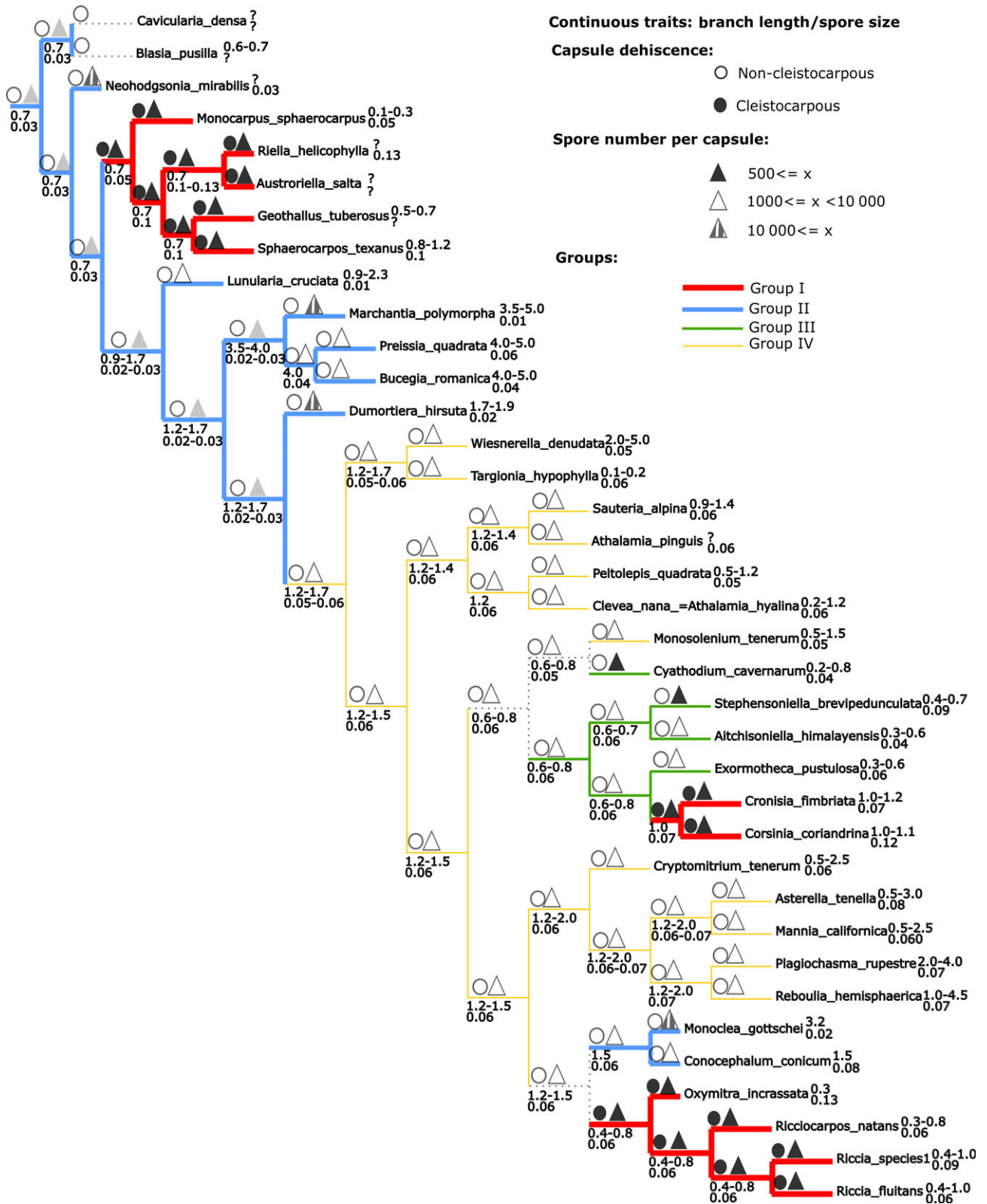


Fig. 7. Character mapping of selected adaptive features across ingroup (Combined tree under BW). Branches coloured according Bischler's life-history groups. Selected characters shown as continuous characters below branches [branch length (cm)/spore size (mm)] and coloured symbols. Grey symbols and dotted branches represents ambiguous optimisations. [Colour figure can be viewed at wileyonlinelibrary.com].

throughout different studies (Crandall-Stotler et al., 2009), and thus the proposal of four different orders within Marchantiidae seemed appropriate (Sphaerocarpaceae, Neohodgsoniales, Lunulariales and Marchantiales). However, Villarreal et al. (2016) found a close association between Lunulariales and Marchantiales (except *Monocarpus*). Our analyses recover the same group, Clade B (Figs 1 and 5), which differs from others definitions of Marchantiales (Bischler, 1998; Crandall-Stotler and Stotler, 2000; Long, 2006; Crandall-Stotler et al., 2009) in excluding *Monocarpus* and including *Lunularia* (as Lunulariales). Although the distinction of two different orders (Lunulariales and Marchantiales) is not necessarily contradicted, it does not seem to be completely suitable given the present results. This is strengthened by the fact that the node comprising Marchantiales (except *Monocarpus*) lacks diagnostic characters and *Lunularia* only shows a low number of autapomorphic changes (Fig S1 and Table S2).

Many of the clades recovered at mid-taxonomic levels by Villarreal et al. (2016; i.e. those which grouped two families) are also found in the present study. The Clade F (*Riccia* + *Ricciocarpos* + *Oxymitra*; Fig 5) was formerly considered as an order (Crandall-Stotler and Stotler, 2000) or suborder (Bischler, 1998). This clade was previously recovered with high support value (Villarreal et al., 2016), and our results agreed in the three analyses (combined, molecular and morphological data; Fig. 1). The rest of the interfamilial groups have not been recognized formally in modern classifications. Clades E (*Conocephalum* + *Monoclea*) and C (*Wiesnerella* + *Targionia*; Fig. 5) are both highly supported and stable in the combined and molecular data. Clade D (*Stephensiella* + *Aitchisoniella* + *Exormotheca* + *Corsinia* + *Cronisia*) is exclusively recovered by our combined and molecular data (Figs 1 and 5), contradicting recent changes (see below).

The outcomes of the present study show discrepancies with the recently made nomenclatural changes at the family level and below (Long and Crandall-Stotler, 2016; Long et al., 2016a,b). The genus *Aitchisoniella* was recently transferred to Cleveaceae (Long et al., 2016b) on the basis of its sister relationship with the remaining genera of the family (Villarreal et al., 2016). In contrast to the results of Villarreal et al. (2016), the position of *Aitchisoniella* within Cleveaceae is not supported by our data (Fig. 1; see Sensitivity plots). In our analyses, on the one hand, *Aitchisoniella* was resolved as sister to *Stephensiella* with high support (SR 100) and clearly nested in Corsiniaceae (Fig. 1; Clade D in Fig. 5). *Exormotheca*, on the other hand, was recovered as sister to *Corsinia* and *Cronisia* in the analyses based on both the combined and molecular data (Fig. 1). Morphology, nonetheless, produces puzzling results regarding Corsiniaceae. As in Villarreal

et al. (2016), morphological trees recovered *Exormotheca* and *Stephensiella* as sister taxa but these were unrelated to the remaining Corsiniaceae (Fig. 3). Hence, the nomenclatural changes proposed to both Cleveaceae and the genus *Stephensiella* (Long et al., 2016a,b) are not supported by the results of our analyses. Instead, our results suggest that Corsiniaceae should be reviewed to accommodate *Aitchisoniella*, and that *Stephensiella* and *Exormotheca* are actually independent taxa.

The genera *Corsinia* and *Cronisia* were found to be sister taxa by the first time in a combined analysis (Fig. 1). Although this clade was resolved here with a moderate support value, it was found in most of the analytical conditions (13 out of 17); showing a high stability (Fig. 1). The fact that Villarreal et al. (2016) recovered a clade consisting of paraphyletic “traditional” families (Corsiniaceae and Exormothecaceae; fig. 2 in Villarreal et al., 2016), led to a logical rearrangement of these groups (Long et al., 2016a,b). Compared to previous studies (Boisselier-Dubayle et al., 2002; Forrest et al., 2006; Villarreal et al., 2016), the inclusion of extensive morphological data turned over many relationships at the genus level. Unlike *Stephensiella* and *Exormotheca* (Villarreal et al., 2016), the novel link between *Cronisia* and *Corsinia* is supported by a large number of morphological characters. In addition, both genera have few autapomorphic characters (Table S2). Altogether, this makes *Cronisia* and *Corsinia* suitable taxa for being merged under a single generic name.

As Crandall-Stotler et al. (2009) pointed out, trying to solve the deep relationships inside Marchantiidae has commonly challenged researchers. Our results show Clade B (Marchantiales and Lunulariales; Fig. 5) is an unstable clade in the molecular dataset and absent in the morphological trees (sensitivity plots in Fig. 1). By contrast, the Clade B showed up as a stable group in the combined analysis. These outcomes suggest that adding morphological data to molecular datasets improves the stability of the clade regardless of the overall incongruence between partitions. Additionally, Clade B received a higher support value when morphological data was included. Similarly, support values of other less-inclusive groups (Sphaerocarpaceae, Ricciaceae and Corsiniaceae) were significantly improved after the addition of morphology (Table S3). These results highlight the fact that extensive morphological data can successfully capture similar patterns to those obtained with molecular data. Even more, unstable or poorly supported results derived from molecular data can be significantly improved after the addition of morphology.

The differences in the results as compared with previous studies may be a consequence of both the addition of new data and the different methods employed

to derive the phylogenetic hypotheses. It has been pointed out that nodes with low support may be sensitive to changes in analytical conditions (Giribet, 2003; Miller and Hormiga, 2004). Under this reasoning, a clade founded on scarce evidence could be unstable to the addition of new conflicting characters. The differences between our results and previous phylogenies focused mainly on Corsiniaceae and Cleveaceae. In Villarreal et al.'s (2016) phylogeny, the groups Corsiniaceae + *Exormotheca* and *Exormotheca* + *Stephensoniella* had low to moderate support (BS <50 and 79, respectively). In our study, these taxa are highly unstable, especially *Exormotheca* (see sensitivity plots in Fig. 1). Additionally, recent phylogenies were mostly conducted under maximum-likelihood or Bayesian analyses (Forrest et al., 2015; Villarreal et al., 2016), but implied weighting was never used for analysing data of this group. Thus, the differences in the results might be at least partially attributed to the different analytical methods.

Contribution of data, conflict and agreement

Boisselier-Dubayle et al. (2002) and Crandall-Stotler et al. (2005) have previously analysed the incongruence in results based on different data types in Marchantiales. Even if exhaustive, such comparisons did not quantify the degree of conflict and contribution of specific groups of morphological characters to recover clades. The topological comparisons (SPR distance and shared nodes) carried out in the present analysis showed the high similarity of the molecular trees with the combined tree. Boisselier-Dubayle et al. (2002) stated that given a large amount of molecular data, such an outcome should be expected. However, the extension to which morphological results depart from molecular trees has been rarely evaluated. In this study, the SPR measures indicated that our morphological trees were similar to the molecular trees (8.1 SPR movements; Fig. 3; Table 2). The comparisons between the molecular trees and the morphological trees of Bischler (1998) and Boisselier-Dubayle et al. (2002) yielded SPR values above 9 (Table 2). In terms of shared nodes, Bischler's tree is similar to our BW molecular tree (Table 2). This similarity was explained by the monophyly of the traditional Exormothecaceae in both Bischler's tree and our BW molecular tree. The alternative comparison, using Villarreal et al.'s (2016) tree as reference, retrieved the K7 morphological tree as the most similar (Table 2). Consequently, our K7 morphological tree maximized similarity with molecular hypotheses in three out of four comparisons.

It has been formerly stressed that morphology-based trees and molecular-based trees rendered markedly different topologies (Boisselier-Dubayle et al., 2002).

Although this conflict is also reflected in our trees (Fig. 3), the K7 morphology-based tree has both dissimilarities with previous morphological trees and common aspects with molecular trees (Bischler, 1998; Crandall-Stotler and Stotler, 2000; Boisselier-Dubayle et al., 2002). As previous authors have pointed out, morphological trees tended to place morphologically complex species in more distant positions regarding Sphaerocarpaceae (Boisselier-Dubayle et al., 2002; Forrest et al., 2006; Crandall-Stotler et al., 2009). In contrast, our morphological trees somewhat resemble the topology of molecular trees (Fig. 3; see online supplemental material). In the K7 morphological tree, the morphologically simple clade F was recovered in a distant position regarding Sphaerocarpaceae (Fig. 3); contrasting with earlier morphological phylogenies (Boisselier-Dubayle et al., 1997, 2002; Bischler, 1998; Crandall-Stotler and Stotler, 2000). The genera *Corsinia* and *Cronisia* were closely related to group F, as in Forrest et al. (2006). The "traditional" Exormothecaceae was rejected (*Aitchisoniella* + *Stephensoniella* + *Exormotheca*; as in Villarreal et al., 2016) and *Neohodgsonia* was excluded from Marchantiaceae (as in Forrest et al., 2006). Nonetheless, our morphological trees also have some coincidences with former morphological trees. For example, they exclude *Monoctlea* from Marchantiales and do not support Lunulariales (as in Crandall-Stotler and Stotler, 2000). In this sense, results from our morphological data can be interpreted as intermediate between the equally-weighted morphological trees of previous studies (Bischler, 1998; Crandall-Stotler and Stotler, 2000; Boisselier-Dubayle et al., 2002) and various molecular hypotheses (Boisselier-Dubayle et al., 2002; Forrest et al., 2006; Villarreal et al., 2016; and this study).

The most in-depth analysis to elucidate the extension of data conflict between sets of morphological characters was performed by Crandall-Stotler et al. (2005). They concluded that sporophytic characters are not more informative than gametophytic characters. In our study, the topological similarity with the molecular tree is maximized by the tree derived from the gametophytic characters (Fig. 4). Nevertheless, this does not imply that sporophytic characters are uninformative. Although some groups are in conflict compared with trees derived from other kind of characters (e.g. Sphaerocarpaceae's nested position into Marchantiales), several taxonomic groups in the molecular tree are also found in the sporophyte tree but not in the gametophyte tree (Fig. 4). Clade F was found in the combined analysis and also in the sporophytic tree, in agreement with previous phylogenies (Bischler, 1998; Crandall-Stotler and Stotler, 2000; Wheeler, 2000; Boisselier-Dubayle et al., 2002; Forrest et al., 2006; Villarreal et al., 2016; Fig. 4). Similarly, the family Aytoniaceae is supported by the sporophytic data but

rejected by gametophytic characters (Fig. 4). Sporophytic traits also resolved two additional clades in accordance with molecular hypotheses (Fig. 5): groups C (*Wiesnerella* and *Targionia*) and A (*Monocarpus* and Sphaerocarpaceae; Wheeler, 2000; Boisselier-Dubayle et al., 2002; Forrest et al., 2006; Villarreal et al., 2016).

Our data show that trees based on morphological data have a certain level of agreement with the molecular topology in terms of SPR distance, shared nodes and groups placement. The increased similarity compared to previous morphological datasets is probably related to the inclusion of new information and the re-interpretation of previous characters. Until now, morphological datasets were relatively small (43 discrete characters) and none of the larger matrices was concentrated on Marchantiidae (Crandall-Stotler and Stotler, 2000; He-Nygrén et al., 2006). In addition, it is worth noting that Boisselier-Dubayle et al. (1997) previously suggested using weighting approaches for diminishing the effect of incongruence between data types. Homoplasy reported by earlier works (Boisselier-Dubayle et al., 2002; Crandall-Stotler et al., 2005) was now downweighted by the use of implied weighting (Goloboff, 1993). Altogether, these factors can be considered as improvements in the analysis of morphological data.

Synapomorphies and Diagnosis

Crandall-Stotler et al. (2009) remarked on the fact that some characters could not be confidently assigned to several groups within the current classification. Many of the groups diagnosed for the first time in the present study were already recovered in previous analyses (Forrest et al., 2006; Villarreal et al., 2016), yet they could not be fully evaluated on the basis of morphological characters. Consequently, their morphological definition remained dubious. In the present combined study, the diagnosis of several groups was clarified.

In the contemporary classification (Crandall-Stotler et al., 2009), Marchantiopsida and Marchantiidae were described in general terms. That is, some characters were actually apomorphic (e.g. *cuneate apical cell* or *uni-stratose capsule wall*; Crandall-Stotler et al., 2005) whereas others were non-apomorphic traits (e.g. *plants thalloid*, *rarely leafy* or *dehiscence by valves, lid or cleistocarpous*). In our BW combined tree, these groups are supported by nine morphological characters each, mainly gametophytic features (Fig. 6; Table 3; Fig. S2). Clade B (Marchantiales and Lunulariales; Fig. 5) has not been diagnosed since its original recovery (Wheeler, 2000; Boisselier-Dubayle et al., 2002; Forrest et al., 2006). In the present combined analysis, such a node is delimited by 15 synapomorphies, most of these scored from the gametophytic phase.

Hexagonal/pentagonal scale cells, lanceolate shaped scales and thin pegged rhizoids are examples of diagnostic characters for this group.

Most of the previous analyses (Wheeler, 1998, 2000; Forrest et al., 2006; He-Nygrén et al., 2006; Villarreal et al., 2016) recovered Clade F (*Oxymitra*, *Ricciocarpos* and *Riccia*; Fig. 5). However, it was not recognized within the contemporary classification (Crandall-Stotler et al., 2009). This group was defined almost exclusively by sporophytic characters in previous morphological analyses (Bischler, 1998; Boisselier-Dubayle et al., 2002). In our study, this node is diagnosed by seven morphological synapomorphies, four of them being gametophytic traits. Thus, the number of synapomorphies not only increased but also new gametophyte-related characters are added. A distinctive diagnostic trait is the ratio between apex and base width of 1.8–2.2; indicating an obcordate thallus shape. A 0.3–0.4 proportion of pegged rhizoids/smooth rhizoids is likewise diagnostic for this clade. Antheridia (male gametangia) primarily positioned in anacrogynous clusters (not derived from an apical cell; as opposed to being gathered in receptacles or specialized branches) is a further gametophytic synapomorphy of this node.

Molecular evidence has consistently recovered both Clades E (*Monoclea* and *Conocephalum*; Forrest et al., 2015; Villarreal et al., 2016) and C (*Wiesnerella* and *Targionia*; Boisselier-Dubayle et al., 2002; Forrest et al., 2006; Villarreal et al., 2016); but none of these studies could provide synapomorphic morphological characters diagnosing both clades, as found here. Clade E is diagnosed by the absence of a germinal tube, whereas C is defined by three synapomorphies: a ratio of 0.7–0.9 between apex and base width, a 0.76–0.77 proportion of pegged rhizoids and semi-annular capsule thickenings (as opposed to annular thickenings in remaining clades).

The results obtained in the present analysis share many nodes with previous phylogenetic hypotheses (Forrest et al., 2006, 2015; Villarreal et al., 2016) although the diagnoses of some nodes differ from that previously proposed. Many characters were potentially linked to the recently proposed Corsiniaceae (*Exormotheca* + *Corsinia* + *Cronisia*; Long et al., 2016a). However, only two characters are synapomorphies of this clade: pegged rhizoids originating near the thallus apex and a multiple-of-8 chromosome number. No spore-related character supported the link between *Exormotheca* and *Cronisia* + *Corsinia* as suggested (Long et al., 2016a). The novel Clade D, which includes *Aitchisoniella* and Corsiniaceae (*Exormotheca*, *Stephensiella*, *Cronisia* and *Corsinia*), was diagnosed by two vegetative traits: antheridia in anacrogynous clusters and drought tolerance. Nevertheless, it must be noted that the diagnosis of Clade D is not comparable to the new Corsiniaceae (Long et al., 2016a)

because it comprises different taxa (i.e., includes *Aitchisoniella*). Clade A (*Monocarpus* and Sphaerocarpaceae; Fig. 5) was recently found by Forrest et al. (2015) who proposed that this group could be described on the basis of sporophytic trait reductions. Our results confirmed Forrest et al.'s proposal by mapping reversions as synapomorphic character changes (cleistocarpous capsules, no elaters and distinctive ornamentation of the spore proximal face).

Diagnoses of the remaining families with more than one genus were also modified relative to Crandall-Stotler et al.'s (2009) classification, as a result of Aytoniaceae and Cleveaceae (excluding *Aitchisoniella*) constraining their original delimitations. Nevertheless, 10 and eight synapomorphies are found for each taxon, respectively. Many of these apomorphies were already proposed by Crandall-Stotler et al. (2009) whereas others are completely new characters. For instance, a 1.09–1.1 pegged rhizoids/smooth rhizoids proportion and simple pegged rhizoids distributed in stalk furrows are new synapomorphies for Aytoniaceae. Cleveaceae (excluding *Aitchisoniella*) is fairly well diagnosed by a high peg density (9.4–11.0), mid to long pegs (4.5–4.6) and tuber presence, among others. Marchantiaceae rendered as apomorphies many of the characters previously proposed by Crandall-Stotler et al. (2009). However, novel characters related to pegged rhizoids are also added. Ricciaceae, as was the case in several classifications (Bischler, 1998; Crandall-Stotler et al., 2009), is circumscribed mainly by reversals (absences).

Two additional features characterize the new diagnoses presented in this study: a high proportion of sporophytic characters and input of continuous characters. Bischler (1998) documented an important number of synapomorphies related to sporophytic characters. Namely, 52% of her sampled sporophytic characters presented apomorphic states (Bischler, 1998). That percentage was informative neither on the ubiquity of changes nor the proportion of sporophytic apomorphies per node. After a scrutiny of the synapomorphies reported by Bischler (1998), it turned out that the sporophytic phase contributed 46 out of 126 changes (36%; mainly concentrated at deep nodes). In our analysis, synapomorphies are dominated by gametophytic features at the family level (Fig. 6; Fig. S2). These quantities may coincide with the notion that sporophytic traits provide scarce information at low taxonomic levels (Schuster, 1984a,b; Crandall-Stotler and Stotler, 2000). However, this idea is not supported when the synapomorphies presented at the level of orders and nodes that grouped families are considered. These nodes were diagnosed by a large number of sporophytic traits (Fig. 6; Fig. S2). Conversely, sporophytic features are poorly represented in the diagnosis at the level of subclasses and classes. The relatively high number of sporophytic synapomorphies at low taxonomic levels compared to that obtained in

Bischler (1998) could be related to the higher number of sporophytic characters included in our dataset. Indeed, Bischler's (1998) dataset included 12 sporophytic characters whereas our matrix scored 34 sporophytic traits. These new findings provide a basis upon which new groups could be proposed relying on both high support values and a clearly stated diagnosis.

Several synapomorphies retrieved in the present analysis involve continuous characters, a novel result given that previous phylogenetic analyses in this group did not include this type of character. The lack of continuous characters in earlier studies (Bischler, 1998; Boisselier-Dubayle et al., 2002) can be explained by the absence of a method that could deal with continuous variation, because Goloboff et al. published their approach later, in 2006. However, recent studies have explicitly suggested that continuous characters be excluded. Oyston et al. (2016) argued that at higher taxonomic categories (\approx deeper nodes), taxa tend to differ more contrastingly than at lower levels (\approx shallow nodes). Consequently, Oyston et al. claimed that classical morphometrics (continuous characters) are not suitable for deep levels. Discretized characters were considered better because they exhibit a gap in the continuous trait being evaluated (Oyston et al., 2016). The analysis of continuous characters analysed as such (Goloboff et al., 2006) led us to diagnose 16 out of 35 ingroup nodes using this type of data. Indeed, even if the consensus is poorly resolved, some nodes at the family level and at middle level are recovered when continuous characters are used to infer phylogenetic relationships (Fig. S4). Thus, our findings contradict the notion that continuous traits are appropriate only for shallow taxonomic levels. Further surveys on continuous characters should be conducted in order to completely elucidate their relevance in liverwort phylogeny.

Life history traits, morphological resemblance and evolutionary scenarios

Although some morphological traits were slightly decoupled regarding life-history groups, a general correlated pattern is clearly recovered (Fig. 7). Bischler (1998) established such life-history groups on the grounds of a statistical analysis of environmental factors and morphological features. By mapping these groups onto her morphological tree, she suggested that morphology and life strategies co-variation is phylogenetically structured.¹ Therefore, morphological

¹Note that Bischler (1998) changed group labelling in her phylogenetic mapping (fig. 15 in Bischler, 1998). However, the groups remained the same [e.g. Marchantiaceae and Conocephalaceae were first placed in group 2 (Bischler, 1998, p. 135) and then in group 4 (p. 136)].

resemblance was argued to be caused by ancestry rather than by environmental pressure. The character reconstruction in our combined tree is in partial agreement with Bischler's hypothesis. Explaining the morphological diversity in terms of common ancestry would require each life-history group to be monophyletic or paraphyletic. Because groups II–IV had a unique origin, morphological resemblance within each group could be attributed to common ancestry. Similarly, within each clade of Group I taxa tended to be highly similar with regard to morphological characters (Fig. 7). Environmental pressure could be considered to explain morphological diversity among unrelated taxa. Unlike members of the same clade, similarity between unrelated taxa could not be caused by common ancestry. The resemblance between, say, the genera *Riccia* and *Monocarpus* is easier to explain in terms of similar environmental pressure (Fig. 7). Our findings, therefore, allow Bischler's hypothesis to be reformulated. That is, based on the current phylogenetic patterns, morphological resemblance is partially explained by common ancestry (inside each clade) and partially by convergence (between unrelated members of the group I).

Multiple independent character reductions occurred in taxa of Group I, commonly limited to open dry habitats (cleistocarpous capsules, small gametophytes, few and large spores). These features are favoured under environmental stress (Bischler, 1998; Bischler et al., 2005). Conversely, members of groups II–IV tended to develop morphologically complex features (larger gametophytes, noncleistocarpous capsules, numerous and small spores), appropriate for mesic habitats (Bischler, 1998). Thus, the correlation between morphological traits and life strategies found by Bischler (1998) is confirmed.

Deviations from the expected evolutionary trends (Bischler, 1998) were a consequence of both dissimilar topologies and methodological issues. The statistical survey of Bischler (1998) was conducted without considering phylogeny as a source of variation constraint. Rather, species were treated as independent statistical entities. In order to evaluate the concerted evolution of potentially adaptive characters, the effect of the phylogeny should be eliminated. By doing so, species could be treated as statistically independent units (Felsenstein, 1985). An exhaustive quantitative evaluation of putative adaptive characters is still needed. At the moment, the adaptive value of morphology remains a poorly investigated area.

Final remarks

The first in-depth combined analysis for the complex thalloid liverworts was conducted. An improved

character sampling led to the construction of the largest morphological dataset. Key findings were achieved regarding morphological contribution and diagnosis improvement. In sum, the well-established assumption that morphology may produce completely incongruent patterns with molecular data was discredited. Indeed, many nodes recovered with low support values in previous studies were retrieved by our combined data with increased support. This suggests that the combination of apparently conflicting data types may reveal hidden support for most of the groups.

It is argued that common weaknesses of morphological data are challenging character circumscription and, for this specific group, no informative changes (Boisselier-Dubayle et al., 2002). Morphological data are often described as highly homoplastic in plants (Buck et al., 2000; Ranker et al., 2004; Liu et al., 2012; Yu et al., 2013; Wu et al., 2015; among others). Such a characterization became a widespread notion for different groups of organisms. Scotland et al. (2003) even suggested that, regardless of the taxonomic group, morphological characters should not be evaluated given their problematic nature. Our results, as well as others' (e.g. Wahlberg et al., 2005), strongly reject such a statement. The inclusion of more and novel characters produced topologies which differ from small-matrix-derived morphological phylogenies and are more similar to molecular trees. Likewise, it was shown not only that the number of diagnosed groups increased, but also that most of the previously proposed diagnoses were imprecise.

Several synapomorphies at the interfamily level were found for the first time and some unsupported taxonomic changes were questioned. Although deep relationships among derived families are still dubious, diagnoses and support values of many interfamily nodes were improved. Subsequently, this could be translated into new groups gathering derived families. The taxonomy of the families Cleveaceae and Corsiniaceae, as well as the genera *Aitchisoniella* and *Stephensoniella*, shall be reviewed. At the present, finding a proper set of synapomorphies for the order Marchantiales is still a major problem. A reasonable approach would be to redefine Marchantiales so as to include its sister taxon Lunulariales. By doing this, a supported and accurately diagnosed category could be proposed. The fact that morphology-derived trees improved their congruence with molecular evidence (increased values of shared nodes and SPR distance) will encourage the survey of more and new morphological characters.

Acknowledgments

We thank Emilio Cano (Real Jardín Botánico) for his technical assistance in extracting and amplifying

plant material. We are also indebted to the curators of the PC herbarium for providing invaluable specimens. Several colleagues also contributed fresh material collected in northern Argentina. J.R.F. is supported by a Doctoral Fellowship of the National Scientific and Technical Research Council (CONICET). This study was funded by FONCyT (PICT-1838, PICT-1930 and PICT -0810), the National University of Tucuman (PIUNT-G524) and CONICET (PUE 0070). TNT is freely available thanks to the Willi Hennig Society.

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

Fig. S1. Additional support measures.

Fig. S2. Proportion of synapomorphies derived from the sporophyte at each diagnosed branch.

Fig. S3. Strict consensus of trees derived from the ten continuous characters under K3.

Table S1. Primers for sequenced genes.

Table S2. Synapomorphies for each node of the combined tree under block weighting (node numbers in Figure S1).

Table S3. Support values for selected groups.