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PHYLOGENETICS OF THE BORAGE FAMILY: DELIMITING BORAGINALES AND ASSESSING CLOSEST RELATIVES

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ABSTRACT

The placement of Boraginales, and relationships within the family have remained elusive in modern, broad phylogenetic studies. In order to assess the phylogeny of Boraginales, and specifically to test the sister lineage of the order, a data matrix of the chloroplast markers *rbcL*, *ndhF*, and *trnL-trnF* was assembled from GenBank and *de novo* sequences (representing 132 new GenBank accessions). Phylogenies inferred using Maximum Likelihood and Bayesian frameworks resulted in identical topologies. Tests for alternative topologies were used to assess whether any of the candidates for sister (Solanales, Gentianales, Lamiales, or *Vahlia*) to Boraginales could be ruled out with this dataset. Gentianales was eliminated as the possible closest relative to Boraginales. Additionally, SH tests were used to test topological results within Boraginales: monophyly of Hydrophyllaceae cannot be rejected and paraphyly of Ehretiaceae with respect to the parasitic Lennoaceae is supported. Taxonomic implications are discussed within the context of these phylogenetic results.

Key words: Boraginaceae, Boraginales, Hydrophyllaceae, Lamiids, Lennoaceae, phylogenetics, systematics.

INTRODUCTION

Boraginales, as currently recognized by taxonomic experts, includes approximately 2500 species (Mabberley 2008; Weigend et al. 2014). The family has a worldwide distribution, with many species occupying seasonally dry and xeric habitats in both tropical and temperate biomes (Gottschling et al. 2001; Weigend et al. 2014). The name derives from the Latin *burra*, which references the often hirsute or hispid leaves of plants within the family (Simpson 2010). Inflorescences are generally a monochasial, scorpioid cyme or composed of cymose units (Buys and Hilger 2003) and flowers are bisexual, actinomorphic, and pentamerous. Fruit morphology has historically been important in circumscribing higher taxa such as genera (de Candolle 1846; Cohen 2013), as well as to delimit lower-level relationships including species boundaries (Hasenstab-Lehman and Simpson 2012). Several taxonomic and systematic questions still remain unanswered for this group, and evidence from DNA has both clarified and confounded aspects of relationships. Despite recent attempts to achieve resolution, confident identification of the closest phylogenetic neighbors to Boraginaceae have remained elusive (Albach et al. 2001; Bremer et al. 2002; Moore et al. 2010; Soltis et al. 2011; Weigend et al. 2014; Refulio-Rodriguez and Olmstead 2014).

De Candolle (1846) was the first to treat Boraginaceae as comprising four subfamilies: Boraginoideae, Heliotropioideae, Ehretioideae, and Cordioideae. This circumscription of Boraginaceae (“sensu DC.”) remained unchanged for nearly 150 years, with new species described and assigned to each of the subfamilies by specialists in the group (e.g., Johnston 1927). Boraginaceae sensu DC. have been placed in a variety of or-

ders (Table 1) under several classification systems. Takhtajan’s (1987) classification differed from those of contemporaries because it recognized the order Boraginales Juss. ex Bercht. & J. Presl, which comprised Boraginaceae sensu DC., Hoplestigmataceae Gilg, Hydrophyllaceae R. Br., Lennoaceae Solms and Tetrachondraceae Skotts. ex Wettst. Subsequent molecular evidence indicated that Tetrachondraceae are instead embedded in Lamiales, but the positions of the other families of Boraginales were unresolved (Wagstaff et al. 2000). In the most recent angiosperm-wide classification, the circumscription of Boraginaceae was expanded to include two previously distinct families, Hydrophyllaceae and Lennoaceae (APG III 2009), while taxonomic experts recognize several smaller families with a distinct order (Cohen 2013; Weigend et al. 2013, 2014). The phylogenetic position of Boraginales remains uncertain with respect to Lamiales, Solanales, and Gentianales, but the order has been placed both phylogenetically and taxonomically in Lamiidae (APG III 2009; Refulio-Rodriguez and Olmstead 2014).

APG III (2009) proposed an expanded familial concept of Boraginaceae in part based on molecular evidence clearly demonstrating that Boraginaceae sensu DC. and Hydrophyllaceae are paraphyletic with respect to each other (Olmstead et al. 1993; Ferguson 1999; Gottschling et al. 2001; Moore and Jansen 2006; Nazaire and Hufford 2012; Cohen 2013; Weigend et al. 2013, 2014; Refulio-Rodriguez and Olmstead 2014). Molecular evidence also supports the removal of several genera from de Candolle’s Hydrophyllaceae to different orders and families. For example, *Hydrolea* L. is now placed in Solanales (Soltis et al. 2000) and *Pteleocarpa* Oliv. is now considered to be in Gentianales (Mabberley 2008). *Codon* L., a small African genus, is of particular note because it is sister to all taxa of de Candolle’s Boraginaceae (Luebert and Wen 2008; Nazaire and Hufford 2012; Weigend et al. 2014). These results have contributed to taxonomic discord. Some authors place *Codon* in its own family, Codonaceae (Weigend and Hilger 2010; Weigend et al. 2013, 2014), whereas others place it in its own tribe within Boraginaceae (Nazaire and Hufford 2012). Table 1 compares

Table 1. Classifications of Boraginaceae s.l. and relatives. The APG III (2009) system is adopted here as a starting point for the present research.

DE CANDOLLE (1846)	CRONQUIST (1988)	TAKHTAJAN (1987)	THORNE (1992)	GOTTSCHLING ET AL. (2001)	APG III (2009) (followed by Nazaire and Hufford 2012)	WEIGEND ET AL. (2014)
Subclassis	Asteridae	Boraginales	Solanales	Boraginales	Lamiids	Boraginales
Corolliflorae	Lamiales	Hydrophyllaceae	Solanaceae	Boraginaceae	Garryales	Boraginaceae
(not a complete list)	Lennoaceae	Boraginaceae ¹	Duckeodendraceae	Cordiaceae	Gentianales	Codonaceae
Hydrophy- llaceae	Boraginaceae	Tetrachondraceae	Goetzeaceae	Ehretiaceae	Lamiales	Wellstediaceae
Boraginaceae ¹	Verbenaceae	Hoplostigmataceae	Nolanaceae	Hydrophyllaceae	Solanales	Hydrophy- llaceae I
Lennoaceae	Lamiaceae	Lennoaceae	Convolvulaceae	Heliotropiaceae	Families: ²	Hydrophy- llaceae II
(<i>incertae sedis</i>)	Solanales		Hydrophyllaceae	Lennoaceae	Boraginaceae s.l. ³	Heliotropiaceae
	Hydrophyllaceae		Boraginaceae ¹		Icacinaeae	Cordiaceae
	Gentianales		Hoplostigmataceae		Metteniusaceae	Ehretiaceae
	Callitrichales		Tetrachondraceae		Oncothecaceae	Lennoaceae
	Scrophulariales		Polemoniaceae		Vahliaceae	
	Campanulales					
	Rubiales					
	Dipsacales					
	Calycerales					
	Asterales					

¹ Includes four subfamilies: Boraginoideae, Heliotropioideae, Ehretioideae, Cordioideae

² Families not placed into orders

³ Includes Hydrophyllaceae, Lennoaceae, and all subfamilies of de Candolle

the taxonomic treatment of this group of plants and its relatives over the last nearly 170 years.

APG III (2009) proposed an expanded Boraginaceae for the additional reason that molecular evidence places previously distinct Lennoaceae within Boraginaceae sensu DC. Lennoaceae as treated in the most recent monograph of the group, consists of two genera and four species (Yatskievych et al. 1986). This group of holoparasites has a disjunct distribution, occurring in southwestern North America and Colombia (Yatskievych et al. 1986). Classification of Lennoaceae has been notoriously difficult using morphology or molecular evidence because of the highly autapomorphic morphology of these holoparasites, and sequencing of chloroplast loci often yields what appear to be pseudogenes (pers. obs.). As shown in Table 1, the group has been placed in Lamiales (Cronquist 1988) and Solanales (Thorne 1992), whereas specialists on parasitic plants instead suspected an affiliation with borages (Yatskievych et al. 1986). Indeed, it is now clear that these parasites are most closely related to recently derived woody borages (Gottschling et al. 2001, 2004, 2005; Moore and Jansen 2006; Nazaire and Hufford 2012; Gottschling et al. 2014). However, their exact phylogenetic position varies in these studies, and the current study seeks to resolve their position within Boraginales.

As reflected in Table 1, both the taxonomy and classification of Boraginales and relatives have been in flux for the last twenty years because of varied phylogenetic results and differing conclusions as to how best to treat the group based on inferred evolutionary history. Gottschling et al. (2001) were the first to conduct a molecular phylogenetic study of Boraginales and to reveal the major lineages. Based on ITS1 sequences, the authors proposed a classification in which order Boraginales was adopted, each of de Candolle's subfamilies of Boraginaceae was elevated to the rank of family, and Hydrophyllaceae and Lennoaceae were recognized. Relationships inferred

from ITS1 were (Boraginaceae (Hydrophyllaceae (Heliotropiaceae Schrad. (Cordiaceae R. Br. ex Dumort. (Ehretiaceae C. Mart. ex Lindl. + Lennoaceae))))). Based on their taxon and nucleotide sampling, many of the constituent lineages received strong support as monophyletic. However, much of the backbone of the tree was either unresolved or not strongly supported, and *Codon* was not sampled. Gottschling et al. (2001) began to reveal the major lineages within Boraginales, but outgroup sampling was inadequate to assess placement of the group among asterids.

Following Gottschling et al. (2001), other studies recovered a similar topology for Boraginales using different molecular markers, although taxon sampling was sparser in most (Moore and Jansen 2006; Cohen 2013; Weigend et al. 2013). None of these studies explored the closest relative to Boraginales. A study by Nazaire and Hufford (2012) is of note because of important novel findings, and their taxonomic treatment highlighted the continued debate about how to best classify Boraginales. The study generated new sequence data, supplemented by Genbank data, and assembled a data set that included nrITS and several chloroplast loci and a broad sampling of Boraginales. This broad taxon sample included *Codon*, which was resolved as sister to what they treated as subfamily Boraginoideae. Contra Weigend and Hilger (2010), Nazaire and Hufford (2012) advocated recognizing *Codon* as a tribe of Boraginoideae rather than as a separate, monogeneric family. The authors supported the APG III (2009) classification, adopting a broadly circumscribed Boraginaceae with constituent clades treated as subfamilies (Table 1). However, none of the foregoing studies sampled broadly enough to identify the closest relatives of Boraginales.

In the most recent phylogenetic analysis of Boraginales, Weigend et al. (2014) supported Gottschling et al.'s (2001) classification (Table 1). In this study, four chloroplast loci were sequenced for a number of in-group and asterid taxa. The

authors recognized Boraginales with two major clades, I and II. Boraginales I comprised the monogeneric families Codonaceae and Wellstediaceae Pilg., as well as Boraginaceae. Boraginales II contained two major clades of Hydrophyllaceae along with Heliotropiaceae, Cordiaceae, Ehretiaceae, and Lennoaceae. The overall topology inferred was (Codonaceae (Wellstediaceae + Boraginaceae)) (Hydrophyllaceae I (Hydrophyllaceae II (Heliotropiaceae (Cordiaceae (Ehretiaceae + Lennoaceae))))). Of the outgroups sampled, Solanales are sister to Boraginales but without strong support.

Similarly, larger-scale studies of angiosperms have not definitively identified the sister of Boraginales (Albach et al. 2001; Bremer et al. 2002; Moore et al. 2010; Soltis et al. 2011; Weigend et al. 2014; Refulio-Rodriguez and Olmstead 2014). Vahliaceae (Bremer et al. 2002), Garryaceae, Lamiales + Solanales and Vahliaceae (Albach et al. 2001), and Lamiidae (Moore et al. 2010; Soltis et al. 2011) have all variously resolved as sister to Boraginales, but never with strong support. In Weigend et al. (2014), Vahliaceae resolved as sister to Lamiales, relatively distant from Boraginales.

Generally, as reviewed above, classification of borages and relatives has suffered from instability especially over the past 20 years. The biggest controversy arises from whether to treat the clade at the ordinal level with constituent clades as families or to treat the entire clade as a family. Many of the taxonomic changes may have been made prematurely, based on phylogenetic studies with poor taxon sampling or poorly resolved phylogenies. This problem is exacerbated by the uncertain position of Boraginales among lamiids.

In this study, I sought to (1) test relationships among Boraginales reported in previous studies and (2) determine the closest relatives of Boraginales using data from select chloroplast genic regions. Taxonomy is discussed in light of the phylogenetic analyses.

MATERIALS AND METHODS

Plant Samples

Samples were field collected or obtained from dried herbarium specimens at the Rancho Santa Ana Botanic Garden (RSA-POM) herbarium. All specimen identifications were verified before inclusion in the study.

Taxon Sampling

DNA sequence data were generated for 41 taxa, representing all major lineages identified to date within Boraginales except *Wellstedia* Baf. and *Hoplostigma* Pierre; material of these two genera was not available. Within Boraginales, taxon sampling included major lineages found in previous molecular studies as follows: Boraginaceae (Weigend et al. 2014), Hydrophyllaceae (Ferguson et al. 1999), Heliotropiaceae (Diane et al. 2002), Ehretiaceae (Gottschling et al. 2004), and Cordiaceae (Gottschling et al. 2005). A combination of de novo sequencing (*Metteniusa*, *Oncotheca*) and GenBank data was used to achieve wide sampling across potential sister lineages among lamiids. The goal was to sample taxa from all lineages that have been placed sister to or part of a polytomy with Boraginales in previous molecular studies, including Solanales, Lamiales, Gentianales, and Vahliaceae. In some cases, terminals are combinations of two or more species from the same genus. Although this assumes that each of

these genera is monophyletic, this is unlikely to be problematic given the goals of this study. Trees were rooted with *Campanula*, which repeatedly has been shown to be well outside the lamiids (Albach et al. 2001; Bremer et al. 2002; Moore et al. 2010; Soltis et al. 2011). Table 2 provides a complete list of taxa, voucher specimens, and genic regions represented in the data matrix, as well as GenBank accession numbers for de novo sequences generated for this study. Appendix 1 provides species and numbers for accessions downloaded from GenBank.

Choice of Genic Regions

Chloroplast loci were selected for phylogeny reconstruction based on two criteria: (1) availability either by de novo sequencing or from GenBank for all taxa sampled including the wide range of putative relatives of Boraginales and (2) rates of evolution ranging from relatively fast to slow. The latter is important given that this study seeks to infer relationships both at the family and ordinal levels: *rbcL* evolves slowly and is easily aligned across all taxa including parasitic plants of *Pholisma* Nutt. ex. Hook.; *ndhF* and especially the *trnL-F* spacer region evolve more rapidly and have a history of resolving species and generic level relationships with high statistical support (Shaw et al. 2005).

DNA Isolation and Sequencing

DNA was isolated using a three-day, modified version of the CTAB (cetyl trimethyl ammonium bromide) protocol (Doyle and Doyle 1987; Friar 2005). Amplifications were done in 25 μ l volumes containing: 2.5 μ l 10 \times standard Mg-free Buffer, 1.25 μ l 1.5 μ M MgCl₂, 0.125 μ l 5000 U/ml TAQ polymerase, 1.2 μ l 10 μ M forward and reverse primers, 1.25 μ l 200 μ M dNTPs, 16.375 μ l H₂O, and 1.0 μ l total of 1–10 μ g/ml genomic DNA. Amplifications were carried out on an Applied Biosystems 2720 thermal cycler, using the following conditions: 5 min 95°C, 35 cycles of 1 min 95°C, 40 seconds 53°C, 1 min 72°C, with a final extension of 7 min at 72°C. The PCR amplicons were precipitated with 20% polyethylene glycol 8000 (PEG) in 2.5 M NaCl using equal volumes of PEG to PCR product; the mixture was incubated at 37°C for 15 minutes. DNA was pelleted by centrifugation for 15 min at 14,000 rpm. The pellet was washed with 80% ethanol. Sequencing was done on an ABI 3130xl at Rancho Santa Ana Botanic Garden using the same primers as for amplification.

Sequence Editing and Alignment

Sequences were edited using Sequencher[®] version 5.2. Annotated sequences were deposited in GenBank (see Appendix 1 for accession numbers). The initial alignment was done in ClustalW (Thompson et al. 1997) and the final alignment was achieved manually in MacClade 4.08 (Maddison and Maddison 2008).

Maximum Likelihood Analysis

Phylogenetic inference using a maximum likelihood optimality criterion (ML; Felsenstein 1981) was implemented utilizing the RAXML 7.2.8 (Stamatakis 2006) plugin in Geneious version 7.1 (Kearse et al. 2012). The GTR + I + G model of nucleotide evolution was selected for both *ndhF* and *trnL-F*, and

Table 2. List of taxa for newly generated data with voucher information and GenBank accession numbers. All vouchers are deposited at RSABG-POM Herbarium.

Taxon Name	Voucher information	<i>rbcL</i>	<i>ndhF</i>	<i>trnL-trnF</i>
<i>Antiphytum floribundum</i> (Torr.) A. Gray	Mexico: 2001, Porter 13096	KX929168	KX929211	KX929254
<i>Anchusa azurea</i> Mill.	Italy: Carle & Kurschner 79-201	KX929167	KX929210	KX929253
<i>Bourreria baccata</i> Raf.	Cabrera 10463	KX929169	KX929212	KX929255
<i>Bourreria spathulata</i> (Miers) Hemsl.	Mexico: 1987, Salinas & Ramos s.n.	KX929170	KX929213	KX929256
<i>Buglossoides calabra</i> I.M. Johnst.	Norris 5045	KX929171	KX929214	KX929257
<i>Codon royenii</i> L.	South Africa: 1950, Mougine 304	KX929172	KX929215	KX929258
<i>Coldenia procumbens</i> L.	Sri Lanka: 1969, Wirawan 939	KX929173	KX929216	KX929259
<i>Cordia boissieri</i> A. DC.	USA: 2010, Hasenstab-Lehman HBG2	KX929174	KX929217	KX929260
<i>Cordia brevispicata</i> M. Martens & Galeotti	USA: 2010, Hasenstab-Lehman HBG5	KX929175	KX929218	KX929261
<i>Cordia elaeagnoides</i> DC.	Mexico: 2011, Hasenstab-Lehman 389	KX929176	KX929219	KX929262
<i>Cryptantha intermedia</i> (A. Gray) Greene	USA: 2007, Hasenstab-Lehman 3	KX929209	KX929252	KX929295
<i>Cynoglossum clandestinum</i> Desf.	Russia: Podlech 50757	KX929177	KX929220	KX929263
<i>Draperia systyla</i> (A. Gray) Torr.	USA: Tillforth 1125	KX929178	KX929221	KX929264
<i>Ehretia acuminata</i> R. Br.	China: Hai-ning 890747	KX929183	KX929226	KX929269
<i>Ehretia anacua</i> (Terán & Berland.) I.M. Johnst.	USA: 2010, Hasenstab-Lehman HBG4	KX929181	KX929224	KX929267
<i>Ehretia latifolia</i> Loisel.	USA: 2010, Hasenstab-Lehman HBG3	KX929179	KX929222	KX929265
<i>Ehretia microcalyx</i> Vaupel	Crosby 1027	KX929180	KX929223	KX929266
<i>Ehretia rigida</i> (Thunb.) Druce	South Africa: 1992, Balkwill 7431	KX929182	KX929225	KX929268
<i>Eriodictyon crassifolium</i> Benth.	Ross 6738	KX929184	KX929227	KX929270
<i>Eriodictyon trichocalyx</i> A. Heller	USA: 2008, Fraga 2209	KX929185	KX929228	KX929271
<i>Halgania erecta</i> Ewart & Rees	Australia: 1990, Smith 1359	KX929186	KX929229	KX929272
<i>Heliotropium amplexicaule</i> Vahl	Argentina: McKinnon et al. s.n.	KX929187	KX929230	KX929273
<i>Heliotropium arborescens</i> L.	Peru: 2012, Hasenstab-Lehman 393	KX929188	KX929231	KX929274
<i>Heliotropium bacciferum</i> Forssk.	Madagascar: Podlech 53258	KX929189	KX929232	KX929275
<i>Hydrophyllum canadense</i> L.	Halse 6969	KX929190	KX929233	KX929276
<i>Lithodora zahnii</i> (Heldr. ex Halácsy) I.M. Johnst.	Strid et al. 28331	KX929191	KX929234	KX929277
<i>Nama hispida</i> A. Gray	Halse 4439	KX929192	KX929235	KX929278
<i>Oncotheca balansae</i> Baill.	New Caledonia: 1980, Morat 6579	KX929193	KX929236	KX929279
<i>Patagonula americana</i> L.	Brazil: Zardini 6921	KX929194	KX929237	KX929280
<i>Phacelia secunda</i> J.F. Gmel.	Chile: 2006, Tepe 2146	KX929195	KX929238	KX929281
<i>Pholisma arenarium</i> (2) Nutt. ex Hook.	USA: 2011, Grummer s.n.	KX929197	KX929240	KX929283
<i>Pholisma arenarium</i> Nutt. ex Hook.	Mexico: 1997, Raz 180	KX929196	KX929239	KX929282
<i>Pholisma culiacanum</i> (Dressler & Kuijt) Yatsk.	Mexico: 1981, Beetle & Yatskievych 81-335	KX929198	KX929241	KX929284
<i>Pholisma sonora</i> (Torr. ex A. Gray) Yatsk.	USA: 1982, Wilson & Yatskievych s.n.	KX929199	KX929242	KX929285
<i>Solenanthus</i> Ledeb.	Russia: 1972, Korzhenevsky 16.7	KX929200	KX929243	KX929286
<i>Tiquilia canescens</i> (A. DC.) A.T. Richardson	USA: 2011, Swanson & Bell 4113	KX929201	KX929244	KX929287
<i>Tournefortia calycosa</i> (Donn. Sm.) D.L. Nash	Hansen & Nee 7564	KX929203	KX929246	KX929289
<i>Tournefortia volubilis</i> L.	USA: 1981, Brumbach 9750	KX929202	KX929245	KX929288
<i>Trichodesma stocksii</i> Boiss.	USA: 2010, Hasenstab-Lehman HBG6	KX929204	KX929247	KX929290
<i>Vahlia capensis</i> (L. f.) Thunb.	Namibia: 1988, Goldblatt 8853	KX929205	KX929248	KX929291
<i>Varronia spinescens</i> (L.) Borhidi	Mexico: 2011, Hasenstab-Lehman 384	KX929206	KX929249	KX929292
<i>Varronia stellata</i> (Greenm.) Borhidi	Mexico: 1990, RSA528715	KX929207	KX929250	KX929293
<i>Wigandia urens</i> (Ruiz & Pav.) Kunth	Mexico: 2011, Hasenstab-Lehman 376	KX929208	KX929251	KX929294

TMV + I + G was selected for *rbcL* using the Akaike information criterion (AIC; Akaike 1974; Posada and Crandall 2001) implemented in jModeltest version 3.7 (Posada 2008). Statistical support was assessed with a maximum likelihood bootstrap (bs) analysis implemented in RAxML (Stamatakis 2006), with bs support values estimated from 10,000 replicates.

Bayesian Inference

Analyses were done using MrBayes version 3.1.2 (Huelsenbeck and Ronquist 2001) implemented on the CIPRES portal (Miller et al. 2010.) Models for each dataset were determined as in the maximum likelihood analyses. The loci were run as

separate process partitions in the mcmc algorithm. All Bayesian analyses ran for 10,000,000 generations, with sampling every 100 generations. Consensus trees were produced from trees sampled after the standard deviation of split frequencies reached a value of 0.01, with all trees prior to this discarded as burn-in trees. Posterior probabilities (pp) were calculated on post burn-in trees.

Alternative Hypothesis Testing

Paraphyly of Hydrophyllaceae and Ehretiaceae (see below) was unexpected and called for testing of whether monophyly of either family can be refuted by the data. To accomplish this, the

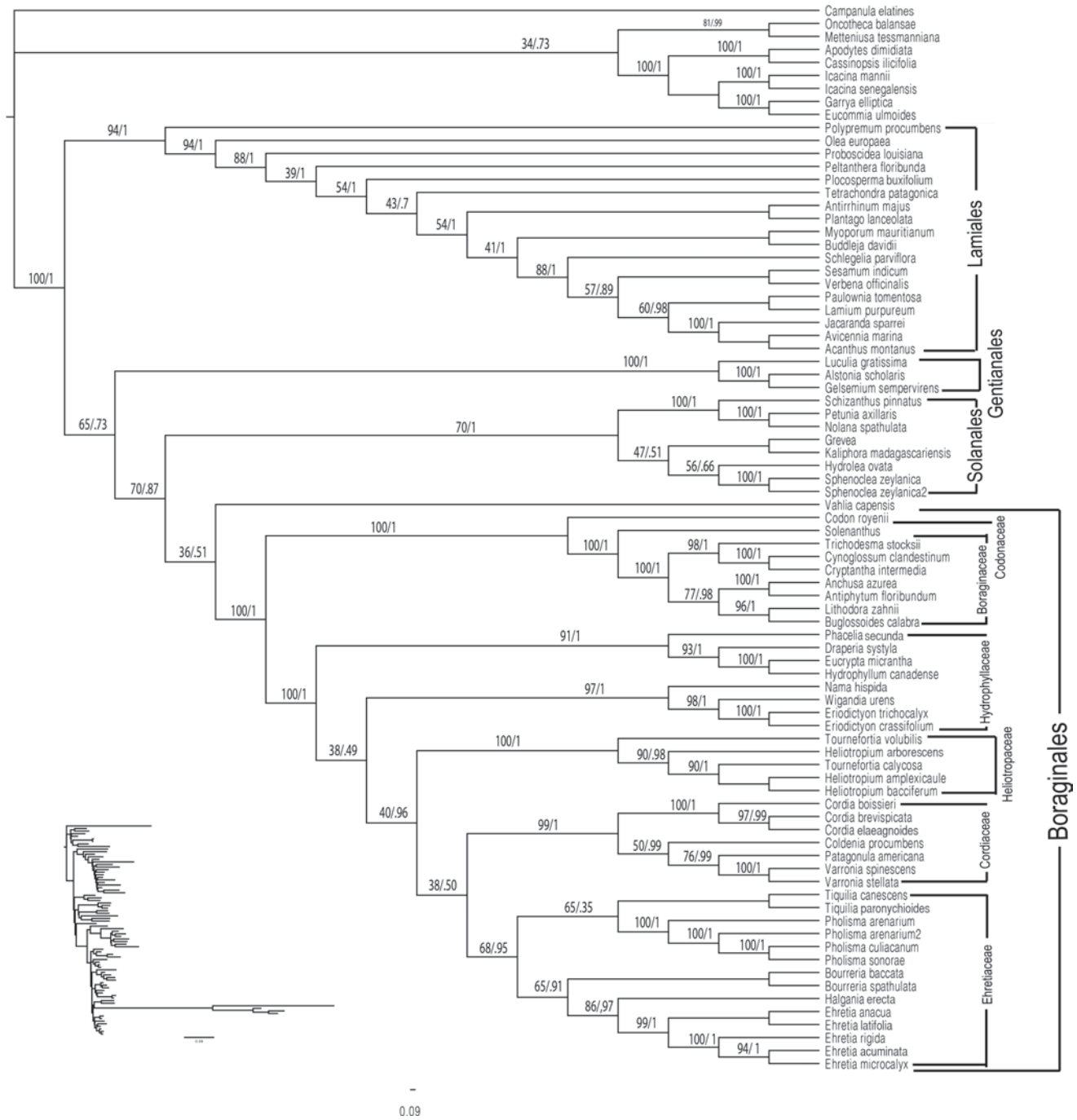


Fig. 1. Maximum likelihood cladogram of Boraginales inferred in RAxML. Support values to the left of the slash (/) are maximum likelihood bootstraps, to the right are posterior probabilities from Bayesian inference implemented in MrBayes. The inset depicts the phylogram from the maximum likelihood analysis to show branch lengths. The extremely long branch is the parasitic *Pholisma*.

Shimodaira-Hasegawa test (Shimodaira and Hasegawa 1999) was implemented in PAUP* (Swofford 2003) to compare tree topologies yielded by the ML and Bayesian analyses to trees constraining these groups to monophyly. The best-fit model for each locus as determined above under the AIC criterion was used. Constrained topologies were Ehretiaceae monophyletic with respect to *Pholisma*, and Hydrophyllaceae I and II forming a single clade. Additionally, to investigate the sister taxon to Boraginales, constrained topologies placing *Vahlia* Thunb.,

Lamiales, Gentianales, and Solanales sister to Boraginales were analyzed.

RESULTS

Phylogenetic Inference: Relationships within Boraginales

Topologies from the likelihood and Bayesian analyses were identical. The maximum likelihood cladogram is presented in Fig. 1 with bs and pp support values. Boraginales formed a

strongly supported clade (bs = 100, pp = 1.00). Boraginaceae + *Codon* were recovered as sister with strong support (bs = 100, pp = 1.00) and as the earliest diverging clade within the family; together they were sister to a clade comprising the remaining lineages. The majority of sampled hydrophylls formed a grade, albeit without strong support for the branching order (bs = 38, pp = 0.49). One clade of Hydrophyllaceae formed a strongly supported clade (bs = 91, pp = 1.00) composed of *Phacelia* Juss., *Draperia* Torr., *Eucrypta* Nutt., and *Hydrophyllum* L. Another clade of Hydrophyllaceae was strongly supported as monophyletic, and comprised the annual *Nama* L. and long-lived shrub genera *Eriodictyon* Benth. and *Wigandia* Kunth. The remaining three lineages, Heliotropiaceae, Cordiaceae, and Ehretiaceae, formed a clade (bs = 40, pp = 0.96), but their interrelationships were not resolved with certainty by this dataset. A topology of Heliotropiaceae sister to Cordiaceae + Ehretiaceae (Fig. 1) was inferred consistently by both analytical approaches, but never with strong statistical support. The monophyly of Heliotropiaceae was strongly supported (bs = 100, pp = 1.00), as was that of Cordiaceae (bs = 99, pp = 1.00). Ehretiaceae were monophyletic with inclusion of the parasitic *Pholisma* (Lennoaceae) but with strong support only from pp (bs = 68, pp = 0.95).

Phylogenetic Inference: Relationships to Other Lamiids

Vahlia is recovered as sister to Boraginaceae, but this relationship lacks statistical support (bs = 36, pp = 0.51). Boraginaceae + *Vahlia* are placed in Lamiidae with strong support (bs = 100, pp = 1.00), along with Solanales, Gentianales, and Lamiales. Each of these orders was recovered as monophyletic with strong support. Lamiales was inferred as the first to diverge, followed by Gentianales (pp = 0.73). Solanales was recovered as sister to the *Vahlia* + Boraginales clade but not with strong support (bs = 70, pp = 0.87). The remaining taxa formed an unresolved clade: Garryales, Icacinaceae s.l., *Metteniusa* H. Karst., and *Oncotheca* Baill. Note that *Metteniusa* + *Oncotheca* resolved as sister taxa with especially strong support from bs (bs = 81, pp = 0.99), a novel result.

Alternative Hypothesis Testing

SH tests implemented in PAUP* returned mixed results for several of the topologies tested. Constraining Hydrophyllaceae as a clade, monophyly could not be rejected by the data ($p = 0.093$), indicating that both hypotheses (monophyly and paraphyly) remain viable. The alternative hypothesis that forced Ehretiaceae to be monophyletic to the exclusion of Lennoaceae was rejected ($p = 0.002$). With respect to other Lamiids, Gentianales were rejected as sister to Boraginales ($p = 0.0003$), but sister relationships involving *Vahlia* ($p = 0.065$), Lamiales ($p = 0.059$) and Solanales ($p = 0.072$) could not be rejected. Thus, all of these taxa remain candidates as the closest living relative of Boraginales.

DISCUSSION

Phylogeny of Boraginales

This study advances our knowledge of relationships among lineages of Boraginales in several ways. The first clade to diverge consists of Boraginaceae + *Codon*. This is consistent with two

recently published studies (Nazaire and Hufford 2012; Weigend et al. 2014). *Wellstedia*, which was not sampled for the present study, belongs in this clade as well based on previous analyses of chloroplast sequences (Weigend et al. 2014). Other studies of Boraginales focused on lower level relationships (Långström and Chase 2002; Nazaire and Hufford 2012; Weigend et al. 2013; Cohen 2013) or on relationships among species and genera within tribes (Luebert and Wen 2008; Cohen and Davis 2012).

The second major clade within Boraginales consists of four lineages, here referred to as Hydrophyllaceae, Heliotropiaceae, Cordiaceae, and Ehretiaceae. To the degree that taxon sampling is comparable, paraphyly of Hydrophyllaceae is consistent with previous phylogenetic analyses (Weigend et al. 2014), but this split is not strongly supported statistically. Further, alternative hypothesis testing demonstrates that these data cannot refute monophyly of Hydrophyllaceae. This phylogenetic uncertainty illustrates the need for both additional taxon sampling and especially more sequence data. Taxonomic changes cannot be recommended at this time, especially since morphological synapomorphies of the clades have yet to be identified. Placement of taxa in Hydrophyllaceae by de Candolle (1846) and subsequent authors seems to have been largely based on symplesiomorphies. A well-resolved and supported phylogeny will allow specialists in the group to identify morphological apomorphies that clearly delimit clades and will pave the way for a stable classification.

A shared common ancestry of Heliotropiaceae, Cordiaceae, and Ehretiaceae has long been posited taxonomically (de Candolle 1846), with evidence from morphology (Johnston 1927; Cohen 2013) and DNA sequence data (Ferguson 1999; Weigend et al. 2001; Moore and Jansen 2006; Luebert and Wen 2008; Nazaire and Hufford 2012; Weigend et al. 2013, 2014). In this study, Heliotropiaceae are the first to diverge, consistent with other studies, and are strongly supported. Diane et al. (2002), Hilger and Diane (2003) and Luebert et al. (2011a,b) provide detailed discussions of relationships within Heliotropiaceae.

Cordiaceae and Ehretiaceae resolve as sister clades, but with poor support (bs = 38, pp = 0.50), consistent with other studies (i.e., Nazaire and Hufford 2012; Weigend et al. 2014). *Cordia* L. is monophyletic and sister to the remaining Cordiaceae, which contrasts with Gottschling et al.'s (2005) study wherein *Patagonula* L. is embedded within *Cordia*. Additional research is needed to fully elucidate relationships within Cordiaceae.

Ehretiaceae is rendered paraphyletic by inclusion of three *Pholisma* species sampled from Lennoaceae. The present analysis places *Tiquilia* Pers. sister to *Pholisma* with weak support (pp = 0.65), but together with the remaining Ehretiaceae sampled, form a clade (pp = 0.95) indicating that the parasitic clade should be subsumed into Ehretiaceae in order to establish monophyletic taxa. Further basis for the placement of *Pholisma* within Ehretiaceae comes from the SH test which rejected monophyly of Ehretiaceae ($p = 0.002$), given the data. This contrasts with Weigend et al. (2014) wherein *Pholisma* forms a polytomy with several other taxa from Ehretiaceae, and is still not placed with certainty in Gottschling et al. 2014. The alternative—splitting Ehretiaceae into smaller units—is left up to taxonomic specialists in the group.

Relationships with Other Lamiids

Vahlia is recovered as sister to Boraginales, consistent with Nazaire and Hufford (2012) and Bremer et al. (2002). This

finding contrasts with Weigend et al. (2014) wherein *Vahlia* is sister to Lamiales. In the present study, *Vahlia* + Boraginales are sister to Solanales, followed successively by Gentianales, then Lamiales. However, none of these relationships is statistically supported. The Shimodaira-Hasegawa tests eliminate Gentianales as sister to Boraginales but retain *Vahlia* ($p = 0.065$), Solanales ($p = 0.072$), or Lamiales ($p = 0.059$) in contention. Other studies such as Soltis et al. (2011), based on 17 genes from across the mitochondrion, ribosomal cistron, and plastome, resolved relationships in Lamiidae as (Vahliaceae ((Boraginaceae + Lamiales) (Gentianales + Solanales))), but also without strong support. With the advent of high-throughput sequencing these relationships may be clarified by sampling of a number of independent loci from the nucleus.

Notes on Classification

The advantages of creating classifications that reflect phylogeny and recognize monophyletic taxa have been well documented. Once phylogenetic patterns and monophyletic groups are identified and strongly supported, resulting classifications should remain stable unless new data, whether molecular or morphological, alter our understanding of phylogenetic patterns.

Over the last 15 years, APG has consistently recognized a broad Boraginaceae, inclusive of Hydrophyllaceae and Lennoaceae. Recent textbooks and field guides reflect this change. However, recognition of Boraginales at the ordinal level with as many as nine families (some monogeneric) has been advocated by taxonomic specialists in the group (Gottschling et al. 2001; Cohen 2013; Weigend et al. 2013, 2014).

The constituent genera and species of Hydrophyllaceae remain incompletely known, and it is still unclear whether hydrophylls represent one lineage or two. Considerably expanded taxon and nucleotide sampling will be required to clearly delineate clade membership.

Additionally, relationships as reconstructed here and by Nazaire and Hufford (2012) and Gottschling et al. (2014) show that the parasitic taxa placed in de Candolle's Lennoaceae are embedded within Ehretiaceae. A classification recognizing Lennoaceae is inconsistent with phylogenetic relationships and should be revised to either split groups within Ehretiaceae further or to subsume the parasites into a taxonomic group inclusive of their autotrophic relatives. It would be best to sample *Lennoa* before formally making taxonomic changes. As well, it would be wise to test the placement of the parasitic plants shown here with data from other genomes, as chloroplast loci may be strongly impacted by the plant's life history strategy (Krause 2008). The chloroplasts of holoparasites may be subject to high rates of evolution and mutations resulting in loss of gene function, evolutionary patterns that can be misleading when these loci are used for phylogenetic inference (Bromham et al. 2013). For stability in naming systems, morphological characters that have been discussed by other authors, and recognition of taxonomy advocated by specialists in Boraginales, this study supports recognition of Boraginales sensu Weigend et al. (2014).

CONCLUSIONS

It is likely that the chloroplast genome has a different evolutionary history than that of nrITS, as suggested by the different branching order among families of Boraginales in this study as

compared to Gottschling et al. (2001), as well as by the close affiliation of *Pholisma* with *Tiquilia* within Ehretiaceae. More sampling of the nuclear and possibly the mitochondrial genome may reveal alternate hypotheses of relationships within Boraginales. Circumscribing Hydrophyllaceae as monophyletic or forming two distinct clades is a matter for future studies by other authors (Weigend et al. 2014) as data analyzed here cannot refute either hypothesis of relationships. Until these relationships are more fully investigated with additional sequence data and taxa it would be premature to make drastic changes to the classification.

The closest relatives of Boraginales remain ambiguous. Again, additional sampling from the other genomes using high-throughput sequencing may be necessary to resolve this relationship. However, given that whole-genome sampling did not resolve relationships among lamiids with strong support (Moore et al. 2010), these relationships may remain elusive, perhaps reflecting rapid evolutionary divergence within Lamiidae.

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APPENDIX 1

Data from GenBank included in the study. Formatted as follows: taxon name, *rbcl* accession, *ndhF* accession, *trnL-trnF* accession.

Acanthus montanus (Nees) T. Anderson, L12592, AJ429115, AJ430912; *Alstonia scholaris* (L.) R. Br., X91760, AJ011982, AJ430907; *Antirrhinum majus* L., L11688, L36392, AJ430929; *Apodytes dimidiata* E. Mey. ex Arn., AJ428895, AJ429109, AJ430899; *Avicennia marina* (Forssk.) Vierh., U28868, AJ429116, AJ430913; *Buddleja davidii* Franch., AJ001757, L36394, AF380861.1; *Campanula elatines* L., EU713438.1, L39387, AJ430970; *Cassinopsis ilicifolia* Kuntze, AJ428896, AJ429110, AJ430900; *Eucommia ulmoides* Oliv., L01917, AJ429113, AJ430905; *Garrya elliptica* Douglas ex Lindl., L01919, AF147714, JN234723.1; *Gelsemium sempervirens* (L.) J. St.-Hil., L14397, AJ011984, AJ430908; *Grevea* Baill. sp., AJ428898, AJ430426, AJ430944; *Hydrolea ovata* Nutt. ex Choisy, L14397, AF013999, AJ430943; *Icacina manni* Oliv., Q384929.1, AJ400888.1, HQ412985.1; *Icacina senegalensis* A. Juss., AJ428897, AJ429111, AJ430901; *Jacaranda sparrei* A.H. Gentry, AF102647, AF102631, FJ870040.1; *Kaliphora madagascariensis* Hook.

f., AJ402963, AJ431206, AJ430945; *Lamium purpureum* L., U78694, JF780029, JF780029.1; *Luculia gratissima* (Wall.) Sweet, HQ384914.1, AJ011987, AJ430911; *Myoporum mauritianum* A. DC., L36445, L36403, AJ430934; *Nolana spathulata* Ruiz & Pav., U08616.1, FJ914007.1, EU581036.1; *Olea europaea* L., AJ001766, AF027288, AJ430529; *Paulownia tomentosa* (Thunb.) Steud., L36447, L36406, AJ430926; *Peltanthera floribunda* Benth., AJ001762, AF027281, AJ430916; *Petunia axillaris* (Lam.) Britton, Sterns & Poggenb., HQ384915.1, U08926.1, HQ412970.1; *Plantago lanceolata* L., L36454, L36408, AY101952.1; *Plocosperma buxifolium* Benth., Z68829, AJ011985, AJ430903; *Polypremum procumbens* L., AJ011989, AJ011986, AJ430938; *Probovidea louisiana* (Mill.) Thell., L01946, AJ236267, KJ743191.1; *Schlegelia parviflora* (Oerst.) Monach., L36448, L36410, AJ430932; *Sesamum indicum* L., L14408, L36413, AF479010.1; *Sphenoclea zeylanica* Gaertn., L18798, AJ429119, AJ430947; *Sphenoclea zeylanica* (2) KT740859, KT738658, AJ430947; *Schizanthus pinnatus* Ruiz & Pav., AY101063.1, U08929.1, AY101172.1; *Tetrachondra patagonica* Skottsb., AF254787.1, AF027272, AJ430939; *Verbena officinalis* L., Z37473, HM216789.1, JN408589.1.