Phylogenetic placement of two enigmatic genera, *Borthwickia* and *Stixis*, based on molecular and pollen data, and the description of a new family of Brassicales, Borthwickiaceae

Jun-Xia Su,^{1,2,3} Wei Wang,¹ Li-Bing Zhang^{4,5} & Zhi-Duan Chen¹

1 State Key Laboratory of Systematic and Evolutionary Botany, Institute of Botany, Chinese Academy of Sciences, Beijing 100093, China

2 Graduate School of the Chinese Academy of Sciences, Beijing 100039, China

3 College of Life Science, Shanxi Normal University, Linfen, Shanxi 041004, China

4 Chengdu Institute of Biology, Chinese Academy of Sciences, P.O. Box 416, Chengdu, Sichuan 610041, China

5 Missouri Botanical Garden, P.O. Box 299, St. Louis, Missouri 63166-0299, U.S.A.

Jun-Xia Su and Wei Wang contributed equally to this paper.

Author for correspondence: Zhi-Duan Chen, zhiduan@ibcas.ac.cn

Abstract Capparaceae (Brassicales) as traditionally circumscribed is heterogeneous, and several genera have been segregated from it based on molecular and/or morphological data. However, *Borthwickia* and *Stixis*, two Southeast Asian endemic genera of Capparaceae with controversial positions, have not previously been evaluated in a molecular phylogenetic study. Here, we used four plastid DNA regions (*matK*, *ndhF*, *rbcL*, *trnL-trnF*) and pollen data to determine their phylogenetic relationships within core Brassicales. Our results showed that neither *Borthwickia* nor *Stixis* is a member of Capparaceae. The two genera, together with *Forchhammeria*, Gyrostemonaceae, Resedaceae, and *Tirania*, formed a clade with strong support. *Stixis* is closely related to *Tirania*, a relationship that is also supported by morphological characters, such as six sepals and three- or four-locular ovaries. Most interestingly, *Borthwickia* was resolved as sister to the *Forchhammeria*-Resedaceae-*Stixis-Tirania* clade with moderate to strong support. However, *Borthwickia* differs markedly from its sister group in having opposite leaves, one indistinct stigma, more than four carpels and locules, a linear ovary with ridges, and pollen grains with perforate exine sculpturing. Thus, we describe a new family, Borthwickiaceae, for the genus.

Keywords Borthwickia; Borthwickiaceae; Brassicales; Capparaceae; phylogeny; pollen morphology; Stixis

Supplementary Material The alignment file is available in the Supplementary Data section of the online version of this article (http://www.ingentaconnect.com/content/iapt/tax).

■ INTRODUCTION

During the past two decades, tremendous progress has been made in understanding relationships among angiosperms by using DNA sequence data (e.g., Soltis & al., 2000, 2011; Wang & al., 2009; Moore & al., 2010). With the basic phylogenetic framework of angiosperms established, a DNA phylogeny-based angiosperm classification system was proposed (APG, 1998) and has been repeatedly updated (APG II, 2003; APG III, 2009). However, due to relatively limited taxon sampling within families, those higher-level phylogenetic analyses of angiosperms did not clarify the circumscriptions of some heterogeneous families. These required more focused work, such as done in Celastraceae (Simmons & al., 2001a, b; Zhang & Simmons, 2006), Capparaceae (Rodman & al., 1993, 1996, 1998; Karol & al., 1999; Chandler & Bayer, 2000; Hall & al., 2002, 2004), Scrophulariaceae (Olmstead & Reeves, 1995; De Pamphilis & al., 1997; Olmstead & al., 2001; Oxelman & al., 2005), and Simaroubaceae (Fernando & al., 1995).

Capparaceae sensu Pax & Hoffmann (1936), comprising ca. 45 genera with 800 species, are primarily restricted to seasonally dry tropical forests. The family has been regarded as a heterogeneous assemblage within Brassicales and has been considered to include many unrelated taxa. Based on phylogenetic analyses, some taxa have been segregated from Capparaceae, including Cleomoideae (Hall & al., 2002), *Calyptrotheca* Gilg (Applequist & Wallace, 2000), *Emblingia* F. Muell. (Chandler & Bayer, 2000), *Forchhammeria* Liebm. (Hall & al., 2002), *Koeberlinia* Zucc. (Rodman & al., 1993), *Pentadiplandra* Baill. (Rodman & al., 1996), *Physena* Noronha ex Thouars (Morton & al., 1997), *Setchellanthus* Brandegee (Karol & al., 1999), and *Tirania* Pierre (Hall & al., 2004). To date, other incertae sedis genera, such as *Borthwickia* W.W. Smith, *Keithia* Spreng., *Neothorelia* Gagnep., *Poilanedora* Gagnep., and *Stixis* Lour., have not been sampled in a molecular phylogenetic study because of the difficulty in obtaining plant material. The systematic positions and allies particularly of *Borthwickia* and *Stixis* have been controversial.

Borthwickia contains one species, *B. trifoliata* (Fig. 1), which is restricted to wet valleys, forests and ravines in southern to southeastern Yunnan (China) and eastern to northern Myanmar (Sun, 1999). Since it was first described by Smith (1911), *Borthwickia* has usually been placed in Capparaceae (e.g., Pax & Hoffmann, 1936; Jacobs, 1968; Brummitt, 1992; Sun, 1999; Wu & al., 2004; Zhang & Tucker, 2008). Smith (1911) considered *Borthwickia* to be close to *Ritchiea* R. Br.

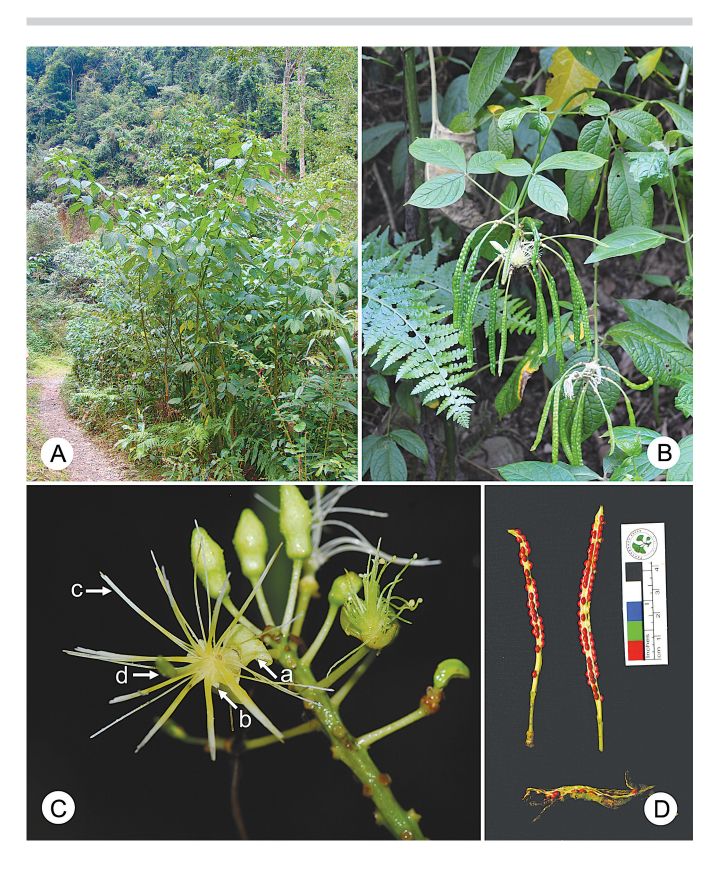


Fig. 1. *Borthwickia trifoliata* in Yunnan, China. **A**, Habitat; **B**, plant with leaves, inflorescences, and fruits; **C**, flower with sepal (a), petal (b), stamen (c), and ovary (d); **D**, fruits and seeds. — Photos: A, Jun-Xia Su; B, Jiang-Hai He; C, Xiao-Hua Jin; D, Tuo Yang.

ex G. Don. Nevertheless, Jacobs (1968) first described the fruit characters of *Borthwickia* and suggested that the genus had an affinity with *Maerua* Forssk. based on the presence of an androgynophore, many stamens, and a torulose fruit (Fig. 1B–D). Based on its unique morphological characters, such as a flower with two sepals and six petals, Pax & Hoffmann (1936) questioned the position of *Borthwickia* within Capparaceae. Kers (2003) further suggested that *Borthwickia* does not belong to Capparaceae, an idea that was followed by Mabberley (2008). Nevertheless, neither Kers (2003) nor Mabberley (2008) assigned the genus to any family.

Stixis, consisting of seven species, occurs from Sikkim (India) to Malaysia, with the Indochinese peninsula as a primary center of diversity (Jacobs, 1963; Hutchinson, 1967; Chen & al., 2003). It was described as a member of Capparaceae by De Loureiro (1790: 295), but was subsequently placed in different tribes of the family by different authors. Pax & Hoffmann (1936) placed Stixis together with Forchhammeria, Neothorelia, Physena, and Tirania in tribe Stixeae based on receptacle and flower characters. Hutchinson (1967) placed Stixis in tribe Cadabeae together with Bachmannia Pax, Boscia Lam., Buchholzia Engl., Cadaba Forssk., Courbonia Brongn., Hypselandra Pax & K. Hoffm., Maerua, and Thylachium Lour., based on their bisexual flowers without petals. Wu & al. (2004) placed Stixis together with Forchhammeria in tribe Stixeae and Tirania in tribe Cappareae, because petals are present in the former two genera, but absent in Tirania. However, Kers (2003) excluded tribe Stixeae of Pax & Hoffmann (1936) from Capparaceae; this treatment was followed by Mabberley (2008) and supported by recent molecular phylogenetic analyses (by sampling Forchhammeria and Tirania; Hall & al., 2002, 2004; Hall, 2008; Martín-Bravo & al., 2007, 2009, 2010).

Pollen characters are potentially of systematic significance in Brassicales because different families sometimes have different pollen characters, such as Capparaceae (Mitra, 1975), Resedaceae (Perveen & Qaiser, 2001; El Naggar, 2002), and Gyrostemonaceae (Tobe & Takahashi, 1995). So far, the pollen morphology of most families of Brassicales has been investigated (e.g., Erdtman, 1952; Mitra, 1975; Perveen & Qaiser, 2001), but that of *Borthwickia* has never been surveyed. Additionally, the pollen characters of *Stixis* have not been used to explore its systematic position although they are different from those of some other members of Capparaceae (Mitra, 1975).

Previous studies have placed Capparaceae, *Forchhammeria*, and *Tirania* in the core Brassicales, which also contain Brassicaceae, Cleomaceae, Emblingiaceae, Gyrostemonaceae, Pentadiplandraceae, Resedaceae, and Tovariaceae (Hall & al., 2002, 2004). The aim of our study was to investigate the phylogenetic position of *Borthwickia* and *Stixis* in the core Brassicales using evidence from molecular analysis and pollen morphology.

MATERIALS AND METHODS

Taxon sampling and molecular data. — Our molecular analysis included four plastid DNA markers (*matK*, *ndhF*, *rbcL*, *trnL-trnF*) which were previously used in phylogenetic studies

of Brassicales (e.g., Hall & al., 2002, 2004; Hall, 2008; Martín-Bravo & al., 2007, 2009, 2010). In this study, the delimitation of Capparaceae followed the treatments of Hall (2008) and Iltis & al. (2011). A total of 66 samples of 57 species from 39 genera in 12 families, representing all families of the core Brassicales and four orphan genera of uncertain position (Borthwickia, Forchhammeria, Stixis, Tirania), were included. Three individuals from two populations of monotypic Borthwickia and ten individuals representing three of seven species of Stixis were newly sampled in this study. Based on previous studies (Rodman & al., 1993, 1996, 1998; Hall & al., 2002, 2004), we selected Bataceae (Batis maritima), Caricaceae (Carica papava), Koeberliniaceae (Koeberlinia spinosa), and Tropaeolaceae (Tropaeolum majus) as outgroups because all of these fall outside core Brassicales. Taxa and GenBank accession numbers for all samples included in this study and voucher information for newly generated sequences are listed in the Appendix.

DNA extraction, PCR amplification, and sequencing. -Total genomic DNA was extracted from silica gel-dried leaf material or herbarium specimens using the Plant Genomic DNA Kit (Tiangen Biotech, Beijing, China) following the manufacturer's protocol. The matK, rbcL, and trnL-trnF regions were amplified following the PCR protocols described by Li & al. (2004), and the ndhF region was amplified following that of Bohs & Olmstead (1997). The matK region was amplified with the primer pair AF/8R of Ooi & al. (1995), the ndhF region with the 972F/2110R primers of Olmstead & al. (1993), the *rbcL* region with the 1F/1494R primers of Chen & al. (1998), and the *trnL-trnF* region with the c/f primers of Taberlet & al. (1991). PCR products were purified using the Tian quick Midi Purification Kit (Tiangen Biotech) following the manufacturer's protocol and were directly sequenced. Additional primers 06F forward (Li & al., 2004) and 06R reverse (Xiang, 2010) for matK and 991R for rbcL (Chen & al., 1998) were used for sequencing. Sequencing reactions were conducted using the ABI Prism BigDye Terminator Cycle Sequencing Kit version 3.1 (Applied Biosystems, Foster City, California, U.S.A.). Sequences were analyzed using an ABI 3730x1 DNA sequencer. Fifty-one sequences were newly obtained from four plastid regions (12 matK, 13 ndhF, 13 rbcL, 13 trnL-trnF).

Phylogenetic analysis. — Sequences were aligned using Clustal X v.1.83 (Thompson & al., 1997) and manually adjusted with BioEdit v.5.0.9 (Hall, 1999). Two difficult-to-align regions in the *trnL-trnF* region, representing 89 nucleotides, were removed from the analyses. Gap characters were scored using "simple indel coding rules" (Simmons & Ochoterena, 2000) with GapCoder (Young & Healy, 2003). Phylogenetic analyses were initially performed for individual regions using maximum parsimony (MP). Because no strongly supported conflicting nodes were found among the trees from different regions, the combined dataset of four loci was analyzed with MP and Bayesian inference (BI) methods as implemented in PAUP* v.4.0b10 (Swofford, 2003) and MrBayes v.3.0b4 (Ronquist & Huelsenbeck, 2003), respectively.

For the MP analysis, heuristic searches were performed with 1000 random-addition replicates, with one tree held at each step during stepwise addition, and tree-bisection-reconnection branch swapping. Multrees was in effect and steepest descent off. Nonparametric bootstrap support (BS; Felsenstein, 1985) for each clade was estimated from 1000 replicates with 10 random taxon additions and heuristic search options.

For the BI analysis, the best-fitting substitution model for each DNA region (matK, ndhF, rbcL, trnL-trnF) was selected by ModelTest v.3.8 (Posada & Crandall, 1998) and using the Akaike information criterion (AIC). The chosen models were TVM+G (nst = 6, rate = gamma) for both *matK* and *trnL-trnF*, and TVM+I+G (nst = 6, rate = gamma) for both ndhF and rbcL. Indel characters of four DNA regions were divided into four independent partitions. The indel data were run under the datatype = standard option. Two independent Markov Chain Monte Carlo (MCMC) runs were conducted simultaneously, each with four linked chains, for 10,000,000 generations, sampling one tree every 1000 generations. Convergence of runs was indicated when the average standard deviation of split frequencies dropped below 0.01 after 2,730,000 generations. In addition, the log-likelihood values of the cold chain of the two simultaneous runs and graphical plots of -lnL generated by the sump command were checked to confirm that stationarity had been reached. For each run, the first 3000 trees were discarded as burn-in; the consensus tree and posterior probability (PP) values were calculated using the remaining 7001 trees.

Pollen morphology. — Pollen samples of two species, Borthwickia trifoliata (Jin & al. YNET125, Qimaba, Lüchun, Honghe, Yunnan, China) and Stixis suaveolens (Qinghua Li 737, XTBG, Menglun, Mengla, Xishuangbanna, Yunnan, China), were collected from herbarium specimens in PE. In addition, pollen samples of Borthwickia trifoliata were also taken from flower specimens pickled in FAA of two individuals collected in Qimaba, Lüchun, Yunnan, China, on 22 May 2011. The material was acetolyzed for light microscopy according to Erdtman (1960). The acetolyzed pollen grains were mounted on glass slides in silicone oil. Observations and measurements were made with a Zeiss Imager A1 microscope. Pollen grain size was based on the average of 20 measurements. For Scanning Electron Microscope (SEM), acetolyzed pollen grains were suspended in a drop of pure alcohol and directly transferred with a fine pipette to a clean stub with double-sided tape. The stubs were coated with gold palladium for 100-130 s in a Hitachi Ion Sputter E-1010 (Hitachi Science Systems, Tokyo, Japan) at 15 mA. Pollen grains were observed and photographed using a Hitachi S-4800 SEM at 10.0 kV. The terminology used here for the description of pollen followed Punt & al. (2007), which is available online (http://www3.bio.uu.nl/palaeo/glossary/glos-int.htm). The pollen morphology of *Borthwickia* and *Stixis* was compared with that of other Brassicales taxa.

Molecular phylogenetics. — The aligned *matK*, *ndhF*, *rbcL*, and *trnL-trnF* datasets had 1346, 940, 1270, and 1297 characters, respectively. Table 1 summarizes the number of variable and parsimony-informative sites and tree statistics for the various datasets. The combined dataset consisted of 4853 characters, of which 2043 were variable and 1240 were potentially parsimonious trees of 4441 steps with a consistency index (CI) of 0.63 and a retention index (RI) of 0.83. The trees from the BI analyses had similar topologies as the MP strict consensus tree (Fig. 2).

Our analyses indicate that the core Brassicales comprise three monotypic families (Emblingiaceae, Pentadiplandraceae, Tovariaceae) and two clades, i.e., Brassicaceae-Capparaceae-Cleomaceae and GRFT (sensu Hall & al., 2004; represented by Gyrostemonaceae, Resedaceae, *Forchhammeria*, and *Tirania*). Within the GRFT clade, Gyrostemonaceae is the earliest-diverging lineage, followed by *Borthwickia* and a well-supported *Forchhammeria*-Resedaceae-*Stixis-Tirania* clade (MP-BS = 100%, PP = 100%). *Borthwickia* is resolved as sister to the latter clade with moderate to strong support (MP-BS = 82%, PP = 100%). *Stixis* is monophyletic (MP-BS = 100%, PP = 100%) and sister to *Tirania* (MP-BS = 100%, PP = 100%).

Pollen morphology. — Pollen grains of *Stixis suaveolens* are tricolporate, mainly spheroidal in equatorial view, and trilobate-circular in polar view. Endoapertures are subcircular. Grains are 14.8 (12.9–17.0) × 11.0 (9.5–13.6) µm. Exine sculpturing is coarsely reticulate with heterobrochus. Colpi with distinct margins are nearly equal to the polar axis in length and taper toward the poles. Colpus membranes are psilate (Fig. 3A–C).

Our pollen morphological observations of three individuals from two populations of *Borthwickia trifoliata* indicated

Table 1. Statistics from maximum parsimony analyses of the various datasets.

Dataset	No. taxa	Total length	Variable characters (incl. gaps)	Informative characters (incl. gaps)	Gaps scored	No. trees	Length of trees	CI	RI	RC
matK	62	1346	712	447	26	14,160	1684	0.61	0.83	0.51
ndhF	60	940	468	303	17	14,985	1183	0.60	0.82	0.49
rbcL	53	1270	301	179	0	184	610	0.62	0.79	0.49
trnL-F	54	1297	562	311	175	13,710	941	0.74	0.87	0.65
Four loci	66	4853	2043	1240	218	528	4441	0.63	0.83	0.53

Abbreviations: CI, consistency index; RI, retention index; RC, rescaled consistency index.

that pollen grains of the species are tricolporate, subprolate in equatorial view, and trilobate-circular in polar view. Endoapertures are circular. Grains are 29.7 (26.6–33.2) \times 22.3 (18.4–25.1) µm. Exine sculpturing is perforate. Colpi are nearly equal to the polar axis in length and have inconspicuous margins and acute ends. Colpus membranes have small tubercles and holes (Fig. 3D–F).

DISCUSSION

Neither Stixis nor Borthwickia is a member of Capparaceae. — Our analyses indicate that the core Brassicales include three monotypic families (Emblingiaceae, Pentadiplandraceae, Tovariaceae) and two clades, Brassicaceae-Capparaceae-Cleomaceae and GRFT, which is congruent with previous molecular studies (e.g., Rodman & al., 1993, 1996, 1998; Hall & al., 2002, 2004; Martín-Bravo & al., 2009, 2010). Traditionally, Borthwickia and Stixis have been placed in Capparaceae sensu Pax & Hoffmann (1936), and have been considered to have a close relationship with Ritchiea or Maerua (Smith, 1911; Jacobs, 1968) and tribe Cadabeae (Hutchinson, 1967), respectively. Our results indicate that both Borthwickia and Stixis were distantly related to Capparaceae, supported by having one 5-bp insertion (indel V) in the *trnL-trnF* region (vs. absent) and lacking one 6-bp insertion (indel VI) in the ndhF gene (vs. present) (Fig. 2). These two genera also differ markedly from Capparaceae in some morphological characters, including axile placentation (vs. parietal), five or more sepals (vs. usually four), and three or more carpels (vs. usually two) (Table 2). Importantly, our observations in Stixis suaveolens indicate that pollen grains are spheroidal and less than 15 µm in polar axis diameter, which is similar to that of the other five species of Stixis, whereas pollen grains of Capparaceae are

Table 2. A comparison of nine morphological characters of Borthwickia and Stixis and their putatively related families or genera.

Taxon	Phyllotaxis	No. sepals	No. petals	No. carpels	No. locules	No. stamens	No. stigmas	Exine sculpture	Placentation
Capparaceae	Alternate	(3-)4(-7)	(0-)4(-8)	2(-8)	1(-2)	4-8(>10)	1	Perforate, reticu- late, scabrate, etc.	Parietal
Boscia	Alternate	4	0	2	1	>10	1	Echinulate	Parietal
Buchholzia	Alternate	4	0	2	1	>10	1	Reticulate	Parietal
Cadaba	Alternate	4	2-4(0)	2(4)	1–2	4-8	1	Reticulate, striate	Parietal
Maerua	Alternate	4	0, 4	2(3)	1–2	>10	1	Reticulate, per- forate	Parietal
Ritchiea	Alternate	4	4	2–4	1	>6	1	Columella, reticu- late, rugulose	Parietal
Thylachium	Alternate	4	0	6–10	1	>10	1	?	Parietal
Gyrostemonaceae	Alternate	48	0	≥1	≥1	>10	≥2	Scabrate-spinulate	Axile
Borthwickia	Opposite	5-8	5-8	4–6	4–6	>10	1	Perforate	Axile
Resedaceae	Alternate	(2-)4-6(-8)	(2-)4-6(-8)	(2-)3-8	1	>10	≥2	Reticulate, rugu- late, striate	Axile (basal-central) marginal, parietal
Forchhammeria	Alternate	4-8	0	2	2	>10	≥ 2	Reticulate	Axile
Neothorelia	Alternate	6	6	3	3	>10	≥2	Reticulate	Axile
Stixis	Alternate	6(5)	0	3(4)	3	>10	≥2	Reticulate	Axile
Tirania	Alternate	6	6	4	4	>10	≥ 2	?	Axile

Characters for Capparaceae and six genera of Capparaceae (*Boscia, Buchholzia, Cadaba, Maerua, Ritchiea, Thylachium*) from Erdtman (1952), Hutchinson (1967), Mitra (1975), and Kers (2003); for Gyrostemonaceae from Tobe & Takahashi (1995) and George (2003); for Resedaceae from Erdtman (1952), Abdallah & de Wit (1978), Perveen & Qaiser (2001), El Naggar (2002), Kubitzki (2003), Martín-Bravo & al. (2007), and Heywood & al. (2007); for *Borthwickia* and *Stixis* from Sun (1999), Wu & al. (2004), Zhang & Tucker (2008), and this study; for *Forchhammeria* from Erdtman (1952) and Hansen (1977); and for *Neothorelia* from Mitra (1975).

Note: Both Smith (1911) and Pax & Hoffmann (1936) considered that flowers of *Borthwickia* had two sepals and six petals, but sepals and petals were each five to eight based on our observations as well as the description of Sun (1999) and Zhang & Tucker (2008).

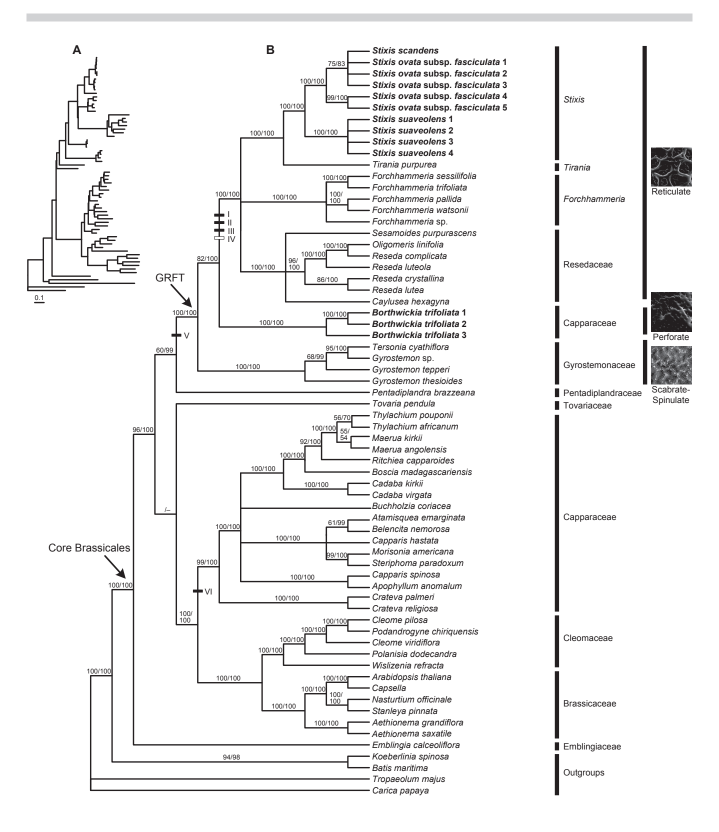


Fig. 2. Phylogenetic relationships of the core Brassicales based on the combined *matK*, *ndhF*, *rbcL*, and *trnL-trnF* datasets. **A**, Bayesian phylogram; **B**, strict consensus tree of 528 most parsimonious trees. — Numbers above branches are MP bootstrap values and Bayesian posterior probabilities as percentages (>50%). The dash (–) indicates a node that does not appear in the BI trees. Molecular synapomorphies (indels) are indicated by boxes on the branches and Roman numerals. Filled boxes represent non-homoplasious synapomorphies, and empty boxes indicate homoplasious changes. Familial classification is based on APG III (2009). The delimitation of Capparaceae is based on Hall (2008) and Iltis & al. (2011).

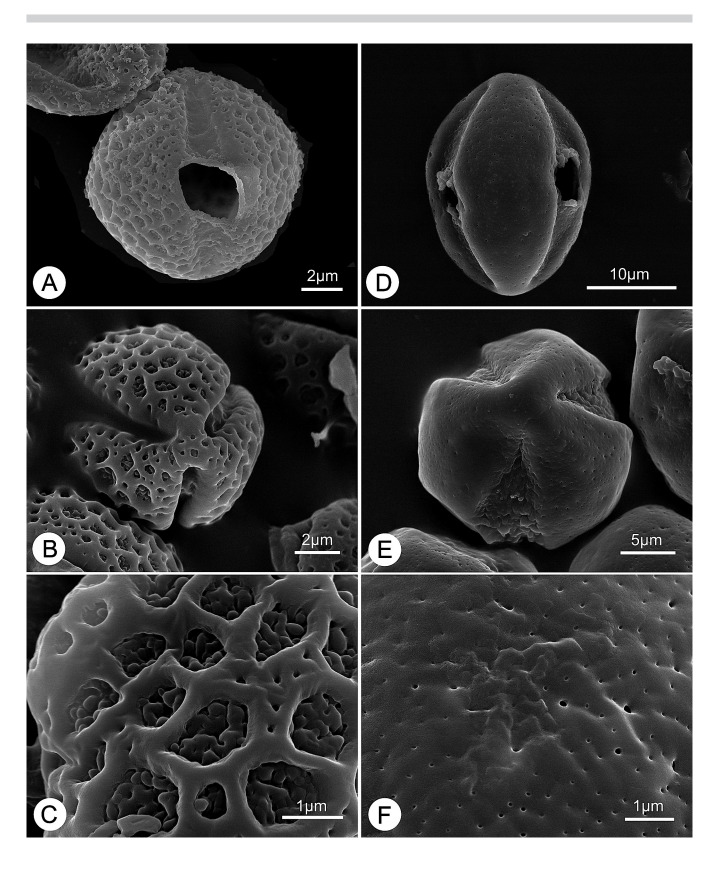


Fig. 3. SEM photographs of pollen grains. **A–C:** *Stixis suaveolens*: **A**, equatorial view; **B**, polar view; **C**, exine pattern. **D–F:** *Borthwickia trifoliata*: **D**, equatorial view; **E**, polar view; **F**, exine pattern.

usually prolate and more than 20 µm in polar axis diameter (Erdtman, 1952; Mitra, 1975) (Fig. 3).

Our results show that both *Borthwickia* and *Stixis* are members of the GRFT clade, which is characterized by an exclusive 5-bp insertion (indel V) in the *trnL-trnF* region (Fig. 2; also see Hall & al., 2002). In addition, *Borthwickia* and *Stixis* also shared several morphological characters, such as an irregular number of sepals, numerous stamens, and axile placentation, with some other members of the GRFT clade (Table 2), providing further support for the new positions of *Borthwickia* and *Stixis*.

Stixis is closely related to Tirania. — Stixis was placed in tribe Stixeae of Capparaceae, with Forchhammeria, Neothorelia, and Tirania, and was considered close to Forchhammeria based on having flowers without petals (Pax & Hoffmann, 1936). However, Stixis is remarkably different from Forchhammeria in having bisexual flowers (vs. dioecious; Hansen, 1977) and an ovary usually with three carpels and locules (vs. an ovary with two carpels and locules; Table 2). Additionally, Stixis is endemic to Southeast Asia, but Forchhammeria is limited to Mexico, Central America, and the West Indies (Hansen, 1977).

Our results strongly support that *Stixis* is sister to *Tirania*, and the two genera together with *Forchhammeria* and Resedaceae form a monophyletic group within the GRFT clade. This group is characterized by one 9-bp deletion (indel I) in the *matK* gene, two 8-bp deletions (indels II, III) in the *trnL-trnF* region, and one 3-bp deletion (indel IV) in the *ndhF* gene (Fig. 2). The close relationship between *Stixis* and *Tirania* is supported by morphological characters, such as the presence of six sepals and an ovary with more than two locules (Table 2). Although closely related to *Tirania*, *Stixis* differs from *Tirania* in having racemes and panicles (vs. solitary flowers) and lacking stipules and petals (vs. present; Hutchinson, 1967; Sun, 1999; Zhang & Tucker, 2008). Therefore, our data support retaining *Stixis* and *Tirania* as separate genera.

Forchhammeria, Neothorelia, Stixis, and Tirania were placed in tribe Stixeae by Pax & Hoffmann (1936). Doweld & Reveal (2008) proposed a familial rank for the tribe, Stixaceae. However, it did not be accepted or used by subsequent authors (APG III, 2009; Martín-Bravo & al., 2009, 2010). To date, three of these four genera have been sampled in molecular phylogenetic analyses, but the monophyly of this group (represented by Forchhammeria, Stixis, and Tirania) remains not to be resolved (Fig. 2). Neothorelia shares several morphological features with Stixis and Tirania, such as six-merous flowers and an ovary with more than two locules (Table 2). Moreover, Neothorelia has pollen grains almost identical to those of Stixis (Mitra, 1975; this study). Thus, Neothorelia seems to be closely related to Stixis and Tirania, but needs to be evaluated in a phylogenetic context. However, the family assignments of members of tribe Stixeae are still premature, for which further study by sampling more markers and more taxa, especially from Neothorelia, is needed.

Borthwickia belongs to a monotypic family. — In our phylogenetic tree (Fig. 2), *Borthwickia* was segregated from Capparaceae and embedded within the GRFT clade with strong

support. *Borthwickia* is morphologically distinct from both other members of the GRFT clade and Capparaceae in having opposite leaves (vs. alternate) and an ovary with more than four locules (vs. less than four; Fig. 1A, B; Table 2).

Our analyses resolved Borthwickia as sister to the Forchhammeria-Resedaceae-Stixis-Tirania clade, but Borthwickia is remarkably different from all members of this clade in having a single stigma (vs. more than one; Table 2) and a linear ovary with ridges (vs. globose, ovoid, or elliptic; Hutchinson, 1967; Kers, 2003; Kubitzki, 2003). Significantly, our pollen observations showed that the exine sculpture of Borthwickia pollen grains is perforate, unlike the reticulate ornamentation in Forchhammeria, Resedaceae, and Stixis (Fig. 3; Table 2). Additionally, Borthwickia is distinguished from its sister clade in lacking one 9-bp deletion in the matK dataset, two deletions (8-bp and 9-bp) in the trnL-trnF dataset, and one 3-bp deletion in the ndhF dataset (Fig. 2). Because Borthwickia has the autapomorphies and other distinct morphological and molecular characters mentioned above, we propose a new monotypic family for it and describe Borthwickiaceae below.

TAXONOMIC TREATMENT

Borthwickiaceae J.X. Su, Wei Wang, Li Bing Zhang & Z.D. Chen, fam. nov. – Type: *Borthwickia* W.W. Smith in Trans. & Proc. Bot. Soc. Edinburgh 24: 175. 1911.

Frutex vel arbuscula. Rami juniores quadrangulati, vetustiores cylindrici. Folia trifoliata, opposita. Inflorescentiae terminales, racemosae. Flores bisexuales. Sepala 5–8, omnino connata, in bilobos rumpentia sub anthesi. Petala 5–8, distincta, sepali tubo manifesto breviora. Stamina 60–70, in androgynophori brevis crassi apice inserta; pollinis granula tricolpora, exiniis punctis. Nectarium conicum, androgynophorum cingens. Ovarium lineare porcatum, 4–6 loculis, placentis axialibus. Stigma simplex sessile obsoletum indivisum. Capsula torulosa, ab basi ad apicem dehiscens in siccitate. Semina reniformia; embryo flexus.

Shrubs or small trees, 1–6 m tall, unarmed, evergreen. Twigs quadrangular with dense, short, white pubescence, later glabrescent; larger branches cylindrical. Stipules absent. Leaves opposite, palmately ternate compound; petiole (3-)5-13(-20) cm, petiolules ca. 1 cm; leaflet blade membranous, margins entire, abaxially with white short pubescence on veins, adaxially glabrous, lateral veins 7-9 pairs, reticulate veins impressed adaxially and prominent abaxially; terminal leaflet oblong, elliptic-lanceolate, sometimes obovate-lanceolate, (5-)8-20(-30) \times (1.5–)4–10(–16) cm; lateral leaflets ovate-lanceolate, slightly smaller than terminal one, base asymmetric. Racemes terminal, 8-20 cm, sessile, rachis with dense short white pubescence; bracts 1.0-1.5 cm, usually linear, deciduous. Flower parts spirally arranged on receptacle; pedicel 1.2-1.5 cm, trichomes like those on axis. Sepals 5-8, whitish, connected into a galericulate tube with short white pubescence on both surfaces, splitting into two lobes at anthesis, deciduous. Petals 5-8, whitish, oblong or spatulate, 1.5-1.8 cm, 1/3-1/2 as long as calyx tube, subequal, distinct, erect, proximally valvate and distally imbricate, membranous and glabrous above middle, thick and with dense pubescence abaxially and sparse pubescence adaxially below middle. Androgynophore ca. 5 mm. Stamens 60-70, 1.4-2.0 cm, fertile, free, at summit of androgynophore; anthers ovate, 2-locular, longitudinally dehiscent, dorsifixed; pollen grains 29.7 (26.6–33.2) × 22.3 (18.4–25.1) µm, tricolporate, subprolate, exine perforate. Nectary conical, ascending from petal base to stamen base, surrounding androgynophore. Ovary linear, 1.0-1.5 cm, distally with 4-6 vertical grooves and ridges, 4-6-locular, with axile placentation, each locule with ovules in two rows; stigma indistinct, sessile. Fruit a capsule, $6-9 \text{ cm} \times 4-6 \text{ mm}$, linear, terete, torulose, base attenuate, apex with a 3-5 mm beak; dehiscent along ventral suture from base to apex leaving a persistent axis with 4-6 ridges; pericarp thin, black to brown after drying. Seeds numerous, 2-3 mm, red when fresh and reddish brown after drying, reniform; embryo bent, scarcely differentiated. Fl. Apr.-Jun., fr. Jul.-Oct. (Smith, 1911; Sun, 1999; Zhang & Tucker, 2008; own observations)

Etymology. – Borthwickia was named in honor of A.W. Borthwick, an English doctor of science and botanist (Smith, 1911).

Distribution and habitat. – Borthwickia contains one species: *B. trifoliata* W.W. Smith. It occurs in wet valleys, forests, and ravines at altitudes of 300–1400 m in southern to southeastern Yunnan, China, and eastern to northern Myanmar.

Representative specimens examined. – BURMA. Shan State, 15 May 1909, MacGregor 714 (E), alt. 500 m, habitat not specified. CHINA. Yunnan: Honghe, Hekou, Laohuazhai, 19 Aug. 1993 (young fruit), Shui & al. 003469 (PE), alt. 1160 m, in forests; Honghe, Hekou, the way from Dawei Mt. to Yaoshan, 09 Oct. 1999 (mature fruit), Shui & al. 11932 (KUN), alt. 900 m, in forests; Honghe, Lüchun, Qimaba, 25 Jun. 2009 (flower, young fruit), Jin & al. YNET125 (PE), alt. 400 m, in forests; Honghe, Jinping, Mengla, Tuomazhai, 28 Jun. 2009 (young fruit), Jin & al. YNET381 (PE), alt. 900 m, in forests; Xishuangbanna, Mengla, Yiwu, Guafengzhai, Ren & Su RS002 (PE), 17 Jan. 2011, alt. 800 m, in ravines; Honghe, Jinping, Mengla, Tuomazhai, 09 Oct. 2011, Yang & al. 00499 (PE), alt. 963 m, in ravines; Honghe, Lüchun, the way from Huanglian Mt. to Qimaba, 10 Oct. 2011 (mature fruit), Yang & al. 00536 (PE), alt. 1074 m, in forests. - DNA sequences of Borthwickia were extracted from three specimens (YNET125, YNET381, RS002).

ACKNOWLEDGMENTS

We sincerely thank Joe Miller and Pamela S. Soltis for carefully reading an early draft of the manuscript and Xiao-Hua Jin, Bao-Qing Ren, Chun-Ce Guo, Bing Liu, Bo Liu, Xu Quan, Miao Sun, Tuo Yang, Jin-Bo Zhang, Yi Zhong as well as Jiang-Hai He, Zhi-Guo Yang, and Jian-Wu Li for their great help in field collection. We also thank Yu-Ming Shui for providing plant material, Yi-Zhen Xi and Zhao-Chen Kong for their advice on the description of pollen grains, and Yin-Hou Xiao for assistance with SEM observations. This research was financially supported by the National Natural Science Foundation of China (grant nos. 40830209 and 31061160184), and CAS Visiting Professorship for Senior International Scientists (2011T1S24).

■ LITERATURE CITED

- Abdallah, M.S. & de Wit, H.C.D. 1978. The Resedaceae: A taxonomical revision of the family (final installment). *Meded. Landbouwhoogeschool* 78: 1–416.
- APG (Angiosperm Phylogeny Group). 1998. An ordinal classification for the families of flowering plants. <u>Ann. Missouri Bot. Gard.</u> 85: 531–553.
- APG II (Angiosperm Phylogeny Group II). 2003. An update of the angiosperm phylogeny group classification for the orders and families of flowering plants: APG II. Bot. J. Linn. Soc. 141: 399–436.
- APG III (Angiosperm Phylogeny Group III). 2009. An update of the angiosperm phylogeny group classification for the orders and families of flowering plants: APG III. Bot. J. Linn. Soc. 161: 105– 121.
- Applequist, W.L. & Wallace, R.S. 2000. Phylogeny of the Madagascan endemic family Didiereaceae. *Pl. Syst. Evol.* 221: 157–166.
- Bohs, L. & Olmstead, R.G. 1997. Phylogenetic relationships in Solanum (Solanaceae) based on ndhF sequences. Syst. Bot. 22: 5–17.
- Brummitt, R.K. 1992. Vascular plant families and genera. Kew: Royal Botanic Gardens.
- Chandler, G.T. & Bayer, R.J. 2000. Phylogenetic placement of the enigmatic Western Australian genus *Emblingia* based on *rbcL* sequences. *Pl. Spec. Biol.* 15: 67–72.
- Chen, W.H., Shui, Y.M., Yang, Z.G. & Cheng, X. 2003. New records of *Stixis* Lour. (Capparaceae) from China. *Acta Phytotax. Sin.* 41: 89–90.
- Chen, Z.D., Wang, X.Q., Sun, H.Y., Han, Y., Zhang, Z.X., Zou, Y.P. & Lu, A.M. 1998. Systematic position of the Rhoipteleaceae: Evidence from nucleotide sequences of the *rbcL* gene. *Acta Phytotax. Sin.* 36: 1–7.
- **De Loureiro, J.** 1790. *Flora cochinchinensis*, vol. 2. Lisbon: Ulyssipone, Typis, et Expensis Academicis.
- De Pamphilis, C.W., Young, N.D. & Wolfe, A.D. 1997. Evolution of plastid gene *rps2* in a lineage of hemiparasitic and holoparasitic plants: Many losses of photosynthesis and complex patterns of rate variation. *Proc. Natl. Acad. Sci. U.S.A.* 94: 7367–7372.
- Doweld, A. & Reveal, J.L. 2008. New suprageneric names for vascular plants. *Phytologia* 90: 416–417.
- El Naggar, S.M. 2002. Taxonomic significance of pollen morphology in some taxa of Resedaceae. *Feddes Repert*. 113: 518–527.
- Erdtman, G. 1952. Pollen morphology and plant taxonomy: Angiosperms. Stockholm: Almquist & Wiksell.
- Erdtman, G. 1960. The acetolysis method, a revised description. Svensk Bot. Tidskr. 54: 561–564.
- Felsenstein, J. 1985. Confidence limits on phylogenies: An approach using the bootstrap. *Evolution* 39: 783–791.
- Fernando, E.S., Gadek, P.A. & Quinn, C.J. 1995. Simaroubaceae, an artificial construct: Evidence from *rbcL* sequence variation. *Amer. J. Bot.* 82: 92–103.
- George, A.S. 2003. Gyrostemonaceae. Pp. 213–217 in: Kubitzki, K. & Bayer, C. (eds.), *The families and genera of vascular plants*, vol. 5, *Flowering plants: Dicotyledons; Malvales, Capparales, and non-betalain Caryophyllales*. Berlin: Springer.
- Hall, J.C. 2008. Systematics of Capparaceae and Cleomaceae: An evaluation of the generic delimitations of *Capparis* and *Cleome* using plastid DNA sequence data. *Botany* 86: 682–696.
- Hall, J.C., Iltis, H.H. & Sytsma, K.J. 2004. Molecular phylogenetics of core Brassicales, placement of orphan genera *Emblingia*, *Forchhammeria*, *Tirania*, and character evolution. <u>Syst. Bot. 29</u>: 654–669.
- Hall, J.C., Sytsma, K.J. & Iltis, H.H. 2002. Phylogeny of Capparaceae and Brassicaceae based on chloroplast sequence data. *Amer. J. Bot.* 89: 1826–1842.
- Hall, T.A. 1999. BioEdit: A user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. *Nucl. Acids Symp. Ser.* 41: 95–98.

- Hansen, B.F. 1977. A monograph of Forchhammeria (Capparidaceae). M.Sc. Dissertation, University of Wisconsin, Madison, Wisonsin, U.S.A.
- Heywood, V.H., Brummitt, R.K., Culham, A. & Seberg, O. 2007. Flowering plant families of the world. Kew: Royal Botanic Gardens.
- Hutchinson, J. 1967. The genera of flowering plants (Angiospermae), vol. 2, Dicotyledons. Oxford: Clarendon Press.
- Iltis, H.H., Hall, J.C., Cochrane, T.S. & Sytsma, K.J. 2011. Studies in the Cleomaceae I. on the separate recognition of Capparaceae, Cleomaceae, and Brassicaceae. Ann. Missouri Bot. Gard. 98: 28–36.
- Jacobs, M. 1963. The genus *Stixis* (Capparaceae). A census. *Blumea* 12: 5–12.
- Jacobs, M. 1968. *Borthwickia* (Capparaceae) from Yunnan and in fruit. *Blumea* 16: 360.
- Karol, K.G., Rodman, J.E., Conti, E. & Sytsma, K.J. 1999. Nucleotide sequence of *rbcL* and phylogenetic relationships of *Setchellanthus caeruleus* (Setchellanthaceae). *Taxon* 48: 303–315.
- Kers, L.E. 2003. Capparaceae. Pp. 36–56 in: Kubitzki, K. & Bayer, C. (eds.), The families and genera of vascular plants, vol. 5, Flowering plants: Dicotyledons; Malvales, Capparales, and non-betalain Caryophyllales. Berlin: Springer.
- Kubitzki, K. 2003. Resedaceae. Pp. 334–338 in: Kubitzki, K. & Bayer, C. (eds.), *The families and genera of vascular plants*, vol. 5, *Flowering plants: Dicotyledons; Malvales, Capparales, and non-betalain Caryophyllales.* Berlin: Springer.
- Li, R.Q., Chen, Z.D., Lu, A.M., Soltis, D.E., Soltis, P.S. & Manos, P.S. 2004. Phylogenetic relationships in Fagales based on DNA sequences from three genomes. *Int. J. Pl. Sci.* 165: 311–324.
- Mabberley, D.J. 2008. *Mabberley's plant-book*, ed. 3. Cambridge: Cambridge University Press.
- Martín-Bravo, S., Meimberg, H., Luceño, M., Märkl, W., Valcárcel, V., Bräuchler, C., Vargas, P. & Heubl, G. 2007. Molecular systematics and biogeography of Resedaceae based on ITS and *trnL-F* sequences. *Molec. Phylogenet. Evol.* 44: 1105–1120.
- Martín-Bravo, S., Valcárcel, V., Vargas, P. & Luceño, M. 2010. Geographical speciation related to Pleistocene range shifts in the western Mediterranean mountains (*Reseda* sect. *Glaucoreseda*, Resedaceae). *Taxon* 59: 466–482.
- Martín-Bravo, S., Vargas, P. & Luceño, M. 2009. Is Oligomeris (Resedaceae) indigenous to North America? Molecular evidence for a natural colonization from the old world. Amer. J. Bot. 96: 507–518.
- Mitra, K. 1975. Contribution to the pollen morphology of the family Capparaceae. *Bull. Bot. Surv. India* 17: 7–31.
- Moore, M.J., Soltis, P.S., Bell, C.D., Burleigh, J.G. & Soltis, D.E. 2010. Phylogenetic analysis of 83 plastid genes further resolves the early diversification of eudicots. *Proc. Natl. Acad. Sci. U.S.A.* 107: 4623–4628.
- Morton, C.M., Karol, K.G. & Chase, M.W. 1997. Taxonomic affinities of *Physena* (Physenaceae) and *Asteropeia* (Theaceae). *Bot. Rev.* 63: 231–239.
- Olmstead, R.G. & Reeves, P.A. 1995. Evidence for the polyphyly of the Scrophulariaceae based on chloroplast *rbcL* and *ndhF* sequences. *Ann. Missouri Bot. Gard.* 82: 176–193.
- Olmstead, R.G., De Pamphilis, C.W., Wolfe, A.D., Young, N.D., Elisons, W.J. & Reeves, P.A. 2001. Disintegration of the Scrophulariaceae. Amer. J. Bot. 88: 348–361.
- **Olmstead, R.G., Sweere, J.A. & Wolfe, K.H.** 1993. Ninety extra nucleotide in *ndhF* gene of tobacco chloroplast DNA: A summary of revisions to the 1986 genome sequence. *Pl. Molec. Biol.* 22: 1191–1193.
- Ooi, K., Endo, Y., Yokoyama, J. & Murakami, N. 1995. Useful primer designs to amplify DNA fragments of the plastid gene *matK* from angiosperm plants. J. Jap. Bot. 70: 328–331.
- Oxelman, B., Kornhall, P., Olmstead, R.G. & Bremer, B. 2005. Further disintegration of Scrophulariaceae. *Taxon* 54: 411–425.

- Pax, F. & Hoffmann, K. 1936. Capparidaceae. Pp. 146–233 in: Engler, A. & Prantl, K. (eds.), *Die natürlichen Pflanzenfamilien*, vol. 17b. Leipzig: Engelmann.
- Perveen, A. & Qaiser, M. 2001. Pollen flora of Pakistan XXVIII: Resedaceae. *Turkish J. Bot.* 25: 39–42.
- Posada, D. & Crandall, K.A. 1998. Modeltest: Testing the model of DNA substitution. *Bioinformatics* 14: 817–818.
- Punt, W., Hoen, P.P., Blackmore, S., Nilsson, S. & Le Thomas, A. 2007. Glossary of pollen and spore terminology. *Rev. Palaeobot. Palynol.* 143: 1–81.
- Rodman, J.E., Karol, K.G., Price, R.A. & Sytsma, K.J. 1996. Molecules, morphology, and Dahlgren's expanded order Capparales. *Syst. Bot.* 21: 289–307.
- Rodman, J.E., Price, R.A., Karol, K., Conti, E., Sytsma, K.J. & Palmer, J.D. 1993. Nucleotide-sequences of the *rbcL* gene indicate monophyly of mustard oil plants. *Ann. Missouri Bot. Gard.* 80: 686–699.
- Rodman, J.E., Soltis, P.S., Soltis, D.E., Sytsma, K.J. & Karol, K.G. 1998. Parallel evolution of glucosinolate biosynthesis inferred from congruent nuclear and plastid gene phylogenies. <u>Amer. J. Bot. 85</u>: 997–1006.
- Ronquist, F. & Huelsenbeck, J.P. 2003. MrBayes 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics* 19: 1572– 1574.
- Simmons, M.P. & Ochoterena, H. 2000. Gaps as characters in sequence-based phylogenetic analyses. Syst. Biol. 49: 369–381.
- Simmons, M.P., Clevinger, C.C., Savolainen, V., Archer, R.H., Mathews, S. & Doyle, J.J. 2001a. Phylogeny of the Celastraceae inferred from phytochrome B and morphology. *Amer. J. Bot.* 88: 313–325.
- Simmons, M.P., Savolainen, V., Clevinger, C.C., Archer, R.H. & Davis, J.I. 2001b. Phylogeny of the Celastraceae inferred from 26S nrDNA, phytochrome B, *atpB*, *rbcL*, and morphology. <u>Molec.</u> <u>Phylogenet. Evol.</u> 19: 353–366.
- Smith, W.W. 1911. *Borthwickia*, a new genera of Capparidaceae. *Trans.* & *Proc. Bot. Soc. Edinburgh* 24: 175–176.
- Soltis, D.E., Smith, S.A., Cellinese, N., Wurdack, K.J., Tank, D.C., Brockington, S.F., Refulio-Rodriguez, N.F., Walker, J.B., Moore, M.J. & Carlsward, B.S. 2011. Angiosperm phylogeny: 17 genes, 640 taxa. Amer. J. Bot. 98: 704–730.
- Soltis, D.E., Soltis, P.S., Chase, M.W., Mort, M.E., Albach, D.C., Zanis, M., Savolainen, V., Hahn, W.H., Hoot, S.B., Fay, M.F., Axtell, M., Swensen, S.M., Prince, L.M., Kress, W.J., Nixon, K.C. & Farris, J.S. 2000. Angiosperm phylogeny inferred from 18S rDNA, *rbcL*, and *atpB* sequences. *Bot. J. Linn. Soc.* 133: 381–461.
- Sun, B.X. 1999. Capparaceae. Pp. 484–540 in: Wu, Z.Y. (ed.), Flora Reipublicae Popularis Sinicae, vol. 32. Beijing: Science Press.
- Swofford, D.L. 2003. PAUP*: Phylogenetic analysis using parsimony (*and other methods), version 4.0b10. Sunderland, Massachusetts: Sinauer.
- Taberlet, P., Gielly, L., Pautou, G. & Bouvet, J. 1991. Universal primers for amplification of three non-coding regions of chloroplast DNA. *Pl. Molec. Biol.* 17: 1105–1109.
- Thompson, J.D., Gibson, T.J., Plewniak, F., Jeanmougin, F. & Higgins, D.G. 1997. The CLUSTAL X windows interface: Flexible strategies for multiple sequence alignment aided by quality analysis tools. *Nucl. Acids Res.* 24: 4876–4882.
- Tobe, H. & Takahashi, M. 1995. Pollen morphology of Gyrostemonaceae, Bataceae, and Koeberlinia. J. Pl. Res. 108: 283–288.
- Wang, H.C., Moore, M.J., Soltis, P.S., Bell, C.D., Brockington, S.F., Alexandre, R., Davis, C.C., Latvis, M., Manchester, S.R. & Soltis, D.E. 2009. Rosid radiation and the rapid rise of angiospermdominated forests. *Proc. Natl. Acad. Sci. U.S.A.* 106: 3853–3858.
- Wu, Z.Y., Lu, A.M., Tang, Y.C., Chen, Z.D. & Li, D.Z. 2004. The families and genera of angiosperms in China: A comprehensive analysis. Beijing: Science Press.

- Xiang, X.G. 2010. *Phylogeny and biogeography of Fagales*. Ph.D. Dissertation, Institute of Botany, Chinese Academy of Sciences, Beijing, China.
- Young, N.D. & Healy, J. 2003. GapCoder automates the use of indel characters in phylogenetic analysis. <u>B.M.C. Bioinf.</u> 4: 6, doi:10.1186/1471-2105-4-6.
- Zhang, L.B. & Simmons, M.P. 2006. Phylogeny and delimitation of the Celastrales inferred from nuclear and plastid genes. *Syst. Bot.* 31: 122–137.
- Zhang, M.L. & Tucker, G.C. 2008. Capparaceae. Pp. 433–450 in: Wu, Z.Y., Raven, P.H. & Hong, D.Y. (eds.), *Flora of China*, vol. 7. Beijing: Science Press; St. Louis: Missouri Botanical Garden.

Appendix. Taxa and GenBank accession numbers for all samples included in this study; voucher information is given for newly generated sequences. An asterisk after the accession number indicates sequences reported here for the first time.

Taxon, voucher, accession number of matK, ndhF, rbcL, trnL-trnF. - missing data; * newly generated sequences.

INGROUPS: Aethionema grandiflora L., AF144354, AF064657, AY167983, -. Aethionema saxatile R. Br., EU371817, AY483250, AY483262, AY122451. Apophyllum anomalum F. Muell., AY483227, AY122356, AY483264, AY122409. Arabidopsis thaliana (L.) Heynh., AF144348, AY122394, D88901, AY122452. Atamisquea emarginata Miers, EU371745, AY122357, -, AY122410. Belencita nemorosa (Jacq.) Dugand, EU371746, AY122358, -, AY122411. Borthwickia *trifoliata* 1 W.W. Smith, China, Yunnan, Honghe, Lüchun, Qimaba, *Jin & al. YNET 125* (PE), JQ733089*, JQ733101*, JQ733114*, JQ733127*. *B. trifoliata* 2 W.W. Smith, China, Yunnan, Honghe, Jinping, Mengla, Tuomazhai, *Jin & al. YNET 381* (PE), JQ733091*, JQ733103*, JQ733116*, JQ733129*. *B. trifoliata* 3 W.W. Smith, China, Yunnan, Xishuangbanna, Mengla, Yiwu, Guafengzhai, Ren B.Q. & Su J.X. RS002 (PE), JQ733090*, JQ733102*, JQ733115*, JQ733128*. Boscia madagascariensis (DC.) Hadj-Moust., EU371749, AY122359, -, AY122412. Buchholzia coriacea Engl., EU371750, AY122360, -, AY122413. Cadaba kirkii Oliv., EU371752, AY122361, -, AY122414. Cadaba virgata Bojer, EU371753, AY122362, -, AY122415. Capparis hastata Jacq., AY483228, AY122366, M95754, AY122420. Capparis spinosa L., EU371772, EU373694, AY167985, AY122422. Capsella bursa-astoris (L.) Medic., -, AY122396, D88904, AY122454. Capsella rubella Reut., AF144334, -, -, -. Caylusea hexagyna (Forssk.) M.L. Green, FJ212207, -, FJ212220, DQ987069. Cleome pilosa Benth., AY483231, AY122385, AY483267, -. Cleome viridiflora Schreb., AY483232, AY122386, AY483268, AY122441. Crateva palmeri Rose, AY483229, AY122370, AY483265, AY122427. Crateva religiosa G. Forst., EU371780, AY122371, -, AY122428. Emblingia calceoliflora F. Muell., -, AY483256, AF146014, -. Forchhammeria pallida Liebm., -, AY122381, AY483274, AY122437. Forchhammeria sessilifolia Standl., AY483243, AY483257, AY483275, -. Forchhammeria sp., AY483244, AY483258, AY483276, -. Forchhammeria trifoliata Radlk., AY483245, AY483259, AY483277, -. Forchhammeria watsonii Rose, AY483246, AY483260, AY483278, -. Gyrostemon sp., AY483236, AY483252, L22439, -. Gyrostemon tepperi (F. Muell. ex H. Walter) A.S. George, AY483237, AY483253, L22440, -. Gyrostemon thesioides (Hook. f.) A.S. George, FJ212199, -, FJ212210, DQ986975. Maerua angolensis DC., EU371783, AY122377, -, AY122433. Maerua kirkii (Oliv.) F. White, AY483230, AY122378, AY483266, AY122434. Morisonia americana L., EU371784, AY122374, -, AY122430. Nasturtium officinale R. Br., AY483225, AY122399, AF020325, AY122457. Oligomeris linifolia (Vahl) J.F. Macbr., AY483240, AY483255, AY483272, FJ212256. Pentadiplandra brazzeana Baill., AY483239, AY483254, U38533, AY122463. Podandrogyne chiriquensis (Standl.) Woodson, AY483233, AY122393, AY483269, AY122450. Polanisia dodecandra DC., AY483234, AY483251, AY167984, AY122447. Reseda complicata Bory, FJ212205, -, FJ212218, DQ987046. Reseda crystallina Webb & Berthel., FJ212200, -, FJ212212, FJ212283. Reseda lutea L., AY483241, AY122406, AY483273, AY122464. Reseda luteola L., FJ212206, -, FJ212219, DQ987050. Ritchiea capparoides (Andr.) Britten, EU371785, AY122375, -, AY122431. Sesamoides purpurascens (L.) G. López., FJ212208, -, FJ212221, DQ987064. Stanleya pinnata (Pursh) Britton, AY483226, AY122401, AY483263, AY122459. Steriphoma paradoxum Endl., EU371786, AY122376, -, AY122432. Stixis ovata subsp. fasciculata 4 (King) Jacobs, China, Yunnan, Xishuangbanna, Mengla, Ren B.Q. & Su J.X. RS001 (PE), JQ733093*, JQ733105*, JQ733118*, JQ733134*. S. ovata subsp. fasciculata 5 (King) Jacobs, China, Yunnan, Xishuangbanna, Mengla, Gongbing Mt., Ren B.Q. & Su J.X. RS013 (PE), JQ733095*, JQ733107*, JQ733120*, JQ733135*. S. ovata subsp. fasciculata 1 (King) Jacobs, China, Yunnan, Honghe, Hekou, Yaoshan, Taiyangzhai, Baiquanchong, Ren B.Q. & Su J.X. RS017 (PE), JQ733097*, JQ733109*, JQ733122*, JQ733136*. S. ovata subsp. fasciculata 2 (King) Jacobs, China, Yunnan, Honghe, Hekou, Yaoshan, Liangzizhai, *Ren B.Q. & Su J.X. RS019* (PE), JQ733098*, JQ733110*, JQ733123*, JQ733137*. *S. ovata* subsp. *fasciculata* 3 (King) Jacobs, China, Yunnan, Honghe, Hekou, Yaoshan, Geniao, *Yang Z.G. 008* (KUN), –, JQ733112*, JQ733125*, JQ733139*. *S. scandens* Lour. China, Yunnan, Honghe, Hekou, Yaoshan, Taiyangzhai, Baiquanchong, Shui Y.M. & al. 12443 (KUN), JQ733099*, JQ733111*, JQ733124*, JQ733138*. S. suaveolens 1 (Roxb.) Pierre, China, Hainan, Chen Z.D. HN171 (PE), JQ733100*, JQ733113*, JQ733126*, JQ733133*. S. suaveolens 3 (Roxb.) Pierre, China, Yunnan, Xishuangbanna, Mengla, Menglun, XTBG, Guo C.C. BN04 (PE), JQ733092*, JQ733104*, JQ733117*, JQ733130*. S. suaveolens 4 (Roxb.) Pierre, China, Yunnan, Xishuangbanna, Mengla, Menglun, XTBG, Ren B.Q. & Su J.X. RS012 (PE), JQ733094*, JQ733106*, JQ733119*, JQ733131*. S. suaveolens 2 (Roxb.) Pierre, China, Yunnan, Honghe, Hekou, Yaoshan, Taiyangzhai, Dudian, Ren B.O. & Su J.X. RS015 (PE), JQ733096*, JQ733108*, JQ733121*, JQ733132*. Tersonia cyathiflora (Fenzl) A.S. George ex J.W. Green, AY483238, AY122404, L22441, AY122462. Thylachium africanum Lour., EU371788, AY122379, -, AY122435. Thylachium pouponii Aubrév. & Pellegr., EU371789, AY122380, -, AY122436. Tirania purpurea Pierre, -, AY483261, AY483279, -. Tovaria pendula Ruiz & Pav., AY483242, AY122407, M95758, AY122465. Wislizenia refracta Engelm., AY483235, AY122391, AY483271, AY122448. OUTGROUPS: Batis maritima L., AY483219, AY122403, L22438, -. Carica papaya L., AY483221, AY483248, M95671, DQ061124. Koeberlinia spinosa Zucc., AY483222, AY483249, L14600, -. Tropaeolum majus L., AY483224, AY122408, L14706, AB043665.