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A COMPLETE GENERIC PHYLOGENY OF MALPIGHIACEAE INFERRED FROM NUCLEOTIDE SEQUENCE DATA AND MORPHOLOGY¹

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- Premise of the study: The Malpighiaceae include ~1300 tropical flowering plant species in which generic definitions and intergeneric relationships have long been problematic. The goals of our study were to resolve relationships among the 11 generic segregates from the New World genus Mascagnia, test the monophyly of the largest remaining Malpighiaceae genera, and clarify the placement of Old World Malpighiaceae.
- Methods: We combined DNA sequence data for four genes (plastid ndhF, matK, and rbcL and nuclear PHYC) from 338 ingroup accessions that represented all 77 currently recognized genera with morphological data from 144 ingroup species to produce a complete generic phylogeny of the family.
- *Key results and conclusions*: The genera are distributed among 14 mostly well-supported clades. The interrelationships of these major subclades have strong support, except for the clade comprising the wing-fruited genera (i.e., the malpighioid+*Amorimia*, *Ectopopterys*, hiraeoid, stigmaphylloid, and tetrapteroid clades). These results resolve numerous systematic problems, while others have emerged and constitute opportunities for future study. Malpighiaceae migrated from the New to Old World nine times, with two of those migrants being very recent arrivals from the New World. The seven other Old World clades dispersed much earlier, likely during the Tertiary. Comparison of floral morphology in Old World Malpighiaceae with their closest New World relatives suggests that morphological stasis in the New World likely results from selection by neotropical oil-bee pollinators and that the morphological diversity found in Old World flowers has evolved following their release from selection by those bees.

Key words: biogeography; floral symmetry; Malpighiaceae; *matK*; mutualism; *ndhF*; *PHYC*; phylogeny; pollination biology; *rbcL*; systematics.

The Malpighiaceae are an angiosperm family of trees, shrubs, and vines in the tropical and subtropical forests and savannas of the New and Old Worlds. They comprise ~1300 species in 77 genera, with ~150 species belonging to 17 exclusively Old World genera. The majority of the genera and species are found in the New World, and the only members of the family found in both hemispheres are two species of New World genera that also occur in coastal western Africa. New World Malpighiaceae typically have unicellular, 2-branched hairs, simple opposite leaves, bilaterally symmetrical flowers with two large oil glands on the abaxial surface of four or all five sepals, five clawed petals, 10 stamens, and a tricarpellate gynoecium with one ovule

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per locule (Anderson, 2004). Some of these characteristics have been lost in Old World members of the family. Malpighiaceae produce a variety of dry or fleshy, indehiscent or dehiscent fruits that are smooth-walled, bristly, or winged (Davis et al., 2001). This diversity in fruits, coupled with the relative uniformity of the flowers of New World Malpighiaceae (Anderson, 1979), led earlier workers to define genera and tribes principally on the basis of fruit morphology (Niedenzu, 1928). Published phylogenetic studies (Cameron et al., 2001; Davis et al., 2001, 2002b, 2004) have highlighted the problematic nature of this single-character taxonomy. Some of the largest traditionally recognized genera have been shown to be polyphyletic [e.g., Mascagnia (DC.) Colla, Cameron et al., 2001; Davis et al., 2001; and Discussion later], and Davis et al. (2001) concluded that all tribes previously recognized for the family (Hutchinson, 1967) were polyphyletic except Gaudichaudieae, whose members ironically bear diverse fruit types, but exhibit distinctive floral features. These data suggest that there have been several origins of very similar fruit types within the family. Reconstructions of fruit evolution (Davis et al., 2001) established that winged fruits evolved early in Malpighiaceae from a smooth-walled ancestor and that there were at least three reversals to smooth-walled fruits from within the large heterogeneous wing-fruited clade, which constitutes the majority of Malpighiaceae species. Bristly fruits are similarly homoplastic and evolved at least three (and more likely four) times in clades with wing-fruited ancestors.

A major goal of our work since 2001 has been to improve the resolution of the Malpighiaceae phylogeny by sequencing

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representatives of the genera that were missing from our previous studies plus a much broader sampling of the larger and more problematic genera. Our aims for this study were to (1) resolve relationships among the 11 segregates from the New World genus Mascagnia sensu lato (s.l.), (2) test the monophyly of the largest remaining Malpighiaceae genera, and (3) clarify the placement of Old World Malpighiaceae. To achieve this goal, we included 309 species of Malpighiaceae, representing about a fourth of the total species diversity in this large tropical clade. We sampled these species across four gene regions that have previously been shown to be phylogenetically informative in the family but have never been analyzed simultaneously (i.e., plastid [pt] matK, ndhF, rbcL, and the low-copy nuclear phytochrome gene PHYC]. We also added a morphological data set for representatives of all the genera and integrated those data into our molecular phylogenetic analyses. This expanded sampling has enabled us to present here a greatly improved phylogeny of the family, which has made it possible to further untangle some of the problems caused by the traditional classifications based mainly on fruit morphology. Working back and forth between molecular and morphological approaches, we are achieving a synergy that is leading not only to new phylogenetic insights for this group, but also to the translation of those insights into a revised classification for the family. The progress of those revisions is summarized in the discussion and on our website for the family (Anderson et al., 2006 onward).

In addition to fruit morphology, biogeography has also been important for circumscribing major groups within the family. Niedenzu (1928) placed the Old World Malpighiaceae (not including the two more recently introduced species from the New World) in subtribe Aspidopteryginae within his lateral-winged tribe Hiraeeae and in subtribe Sphedamnocarpinae within his dorsal-winged tribe Banisterieae. While each of these wingfruited tribes has previously been shown to be polyphyletic (Cameron et al., 2001; Davis et al., 2001), the precise New World relatives of these Old World clades have remained elusive. In previous studies, the Old World genera were shown to represent six to nine different clades, each of which was more closely related to a New World clade than to other Old World clades. Alternative topologies in which these Old World species were forced to form a single clade were strongly rejected. More recent investigations of the biogeographic history of Malpighiaceae (Davis et al., 2002a, b, 2004) have better clarified the placement of many of these Old World lineages, but the identity of the closest New World relatives of several Old World clades remains problematic due to low support and/or incomplete taxon sampling. These difficulties are magnified by our subsequent discovery of several previously unsampled, mostly New World, phylogenetic lineages identified here. Resolving the number of Old World lineages and discerning their placement among the many newly recognized New World members of the family will aid in future biogeographic studies of the family.

Finally, establishing the placement of most Malpighiaceae genera will help to clarify patterns of floral evolution within the family. Malpighiaceae exhibit a highly stereotyped floral morphology characterized by an elegant suite of floral characters that appears to be closely tied to their oil-bee pollinators (Vogel, 1974, 1990; Anderson, 1990). These bees, however, do not exist in the Old World where most of these unique floral features have been lost (Davis, 2002). Malpighiaceae thus provide an excellent opportunity to examine the origin and maintenance of morphological stasis in the Tree of Life.

MATERIALS AND METHODS

Taxon sampling-Three hundred thirty-eight accessions representing all 77 currently recognized genera of Malpighiaceae (Anderson et al., 2006) plus 19 outgroup accessions were included in our analyses (see Appendix 1). Our sampling builds greatly on our past efforts (Cameron et al., 2001; Davis et al., 2001, 2002b, 2004). In particular, the following genera have been added since our last studies: Aenigmatanthera W. R. Anderson, Amorimia W. R. Anderson, Bronwenia W. R. Anderson & C. Davis, Banisteriopsis C. B. Rob. s.s., Calcicola W. R. Anderson & C. Davis, Camarea A. St.-Hil., Christianella W. R. Anderson, Clonodia Griseb., Digoniopterys Arènes, Henleophytum H. Karst., Madagasikaria C. Davis, Microsteira Baker, Philgamia Baill., and Verrucularia Adr. Juss. In addition, we have greatly expanded our sampling in the largest and most heterogeneous genera, including Banisteriopsis, Bunchosia H.B.K., Byrsonima H.B.K., Heteropterys H.B.K., Janusia Adr. Juss. and its close relatives (herein referred to as the aspicarpoid clade), Malpighia L., Mascagnia s.l., and Tetrapterys Cav. These taxa were targeted primarily because they were either shown to be nonmonophyletic in previous studies (e.g., Mascagnia s.l., and several aspicarpoid genera) or were suspected of being nonmonophyletic on the basis of recent morphological investigations (e.g., Banisteriopsis, Heteropterys, Malpighia, and Tetrapterys). Bunchosia and Byrsonima were not suspected of being nonmonophyletic but are two of the largest genera in the family, with 68 and 130 species, respectively. We included more species of these two genera to make our sampling more reflective of Malpighiaceae diversity, which will be important for future evolutionary studies in the group.

Broader phylogenetic analyses based on several genes have independently confirmed the sister group relationship of Malpighiaceae and Elatinaceae (Davis and Chase, 2004; Tokuoka and Tobe, 2006; Wurdack and Davis, 2009), and more recent analyses that include denser taxon and character sampling across the order Malpighiales indicate that the newly recognized family Centroplacaceae (Zhang and Simmons, 2006; Wurdack and Davis, 2009) is sister to Elatinaceae-Malpighiaceae (Wurdack and Davis, 2009; Xi et al., 2010). Members of Elatinaceae and Centroplacaceae were, therefore, included as outgroups. In addition, we included several more distant outgroups to help stabilize the ingroup topology, including other Malpighiales (Chrysobalanaceae, Dichapetalaceae, Euphorbiaceae, Goupiaceae, Ochnaceae, Phyllanthaceae, Picrodendraceae, Putranjivaceae, and Violaceae), Celastrales (Celastraceae), and Saxifragales (Peridiscaceae). *Peridiscus lucidus* Benth. (Peridiscaceae) was used for rooting purposes.

Molecular methods—Total cellular DNAs were prepared using the protocol of Davis et al. (2002a) or were obtained from other sources (see Acknowledgments). Voucher information is listed in Appendix 1.

Amplification and sequencing protocols for obtaining *matK* followed Cameron et al. (2001), using their primers 400F, *trnK*-2R, and 842F; *ndhF* followed Davis et al. (2001); *rbcL* followed Cameron et al. (2001); and *PHYC* followed Davis et al. (2002b) with the addition of forward primer int-1F (5'-CCAGCTACTGATATACCWCARGCTTC-3'), which when paired with reverse primer 623r/cdo produced an ~800 base-pair (bp) amplicon.

Double-stranded polymerase chain reaction (PCR) products were primarily gel-extracted and purified using the QIAquick Gel Extraction Kit. PCR products were sequenced in both directions using dye terminators and sequencing protocols at the University of Michigan DNA facility (Ann Arbor, Michigan, USA) and at MWG Biotechnology (High Point, North Carolina, USA). Chromatograms were assembled into contiguous sequences and checked for accuracy using the software program Sequencher 4.7 (Gene Codes Corp., Ann Arbor, Michigan, USA). All newly generated sequences were submitted to GenBank (see Appendix 1).

Morphological scoring—Morphological data were recorded for 144 ingroup species that represented all genera of Malpighiaceae, plus six outgroup taxa. Seventy-five discrete (binary or multistate) morphological characters were scored, representing vegetative and reproductive structures (see Appendix 2). We have indicated representative voucher specimens for each of the species that were scored in Appendix 1, but in many cases additional specimens were needed to adequately score all of the relevant data.

Phylogenetic analyses—Nucleotide sequences were aligned by eye with the program MacClade 4.0 (Maddison and Maddison, 2000); the ends of sequences were trimmed from each data set to maintain complementary data between taxa. The aligned pt *matK*, *ndhF*, *rbcL*, and nu *PHYC* data sets included 1194, 867, 1414, and 1180 bp, respectively; 75 morphological characters were also included as part of the alignment. Data matrices are available in Treebase (no. 10998, http://www.treebase.org) and from C. C. Davis.

Maximum likelihood (ML) bootstrap consensus trees and Bayesian posterior probabilities from all individual analyses of the five data partitions revealed no strongly supported incongruent clades (i.e., >80 ML bootstrap percentage [BP]) and were thus analyzed simultaneously using the search strategies described below. Searches using the combined data were conducted using a data set that included 357 accessions (338 ingroup and 19 outgroup).

The optimal model of molecular evolution for the individual and combined analyses was determined by the Akaike information criterion (AIC) using the program Modeltest ver. 3.7 (Posada and Crandall, 1998; Posada and Buckley, 2004). In each case, the optimal model was the general time reversible model, with rate heterogeneity modeled by assuming that some sites are invariable and that the rate of evolution at other sites is modeled using a discrete approximation to a gamma distribution [GTR+I+Г]). A mixed model was applied to our combined analyses to accommodate the morphological data: GTR+I+F for the molecular data and the Mk1 model for the morphological data (Lewis, 2001). Maximum likelihood (ML) analyses of the individual and combined matrices were implemented in the program RAxML 7.1.0 (Stamatakis, 2006) (distributed by A. Stamatakis at http:// icwww.epfl.ch/~stamatak/index-Dateien/Page443.htm) starting from random trees and using the default parameters. Analyses were rerun at least three times. ML BP values were estimated from 100 bootstrap replicates. We subsequently conducted analyses not presented here using RAxML 7.2.6 in which $GTR+I+\Gamma$ was enforced for all partitions, including morphology. Those topologies were very similar to the ones presented here, but were slightly better resolved. Bayesian analyses were similarly implemented in the parallelized version of MrBayes ver. 3.1.2 (Huelsenbeck and Ronquist, 2001) following Davis et al. (2005). Bayesian posterior probabilities (BPP) were calculated from five restarts with a burn-in period of 400 000 generations.

RESULTS

We increased the ingroup taxonomic sampling by a factor of ~4.3 over that used in preparing our earlier phylogeny of the family (Davis et al., 2001). This sampling includes representatives of all the genera currently recognized in the family and many of the species groups in the most diverse genera. For the 45 genera for which we sampled multiple species, 35 (or 78%) are monophyletic. For the sake of space, we present the ML results from the combined data here (Fig. 1, but see Appendix S1 online at http://www.amjbot. org/cgi/content/full/ajb.1000146/DC1 for BPP). A tree summarizing the major ingroup clades is also presented for simplicity in Fig. 2.

Several broad generalizations can be made about our new phylogeny of Malpighiaceae. We have obtained far greater resolution of the family than in our previous studies (Cameron et al., 2001; Davis et al., 2001). In addition to identifying several novel relationships, our findings largely confirm these previous results, but often with greatly increased support (e.g., the placements of Barnebya W. R. Anderson & B. Gates and Lophopterys Adr. Juss.). Bootstrap support along the spine of the tree (Fig. 2), and for all of the major subclades, is mostly very (>90 BP) to moderately (\geq 71 BP) strong. The exception is that the tetrapteroid, stigmaphylloid, and malpighioid clades, as well as Ectopopterys W. R. Anderson, are not well resolved with respect to one another. Furthermore, within each of the 14 major subclades identified (Figs. 1, 2), there is generally strong support for internal clades, with the exception of the christianelloids, cordobioids, hiraeoids, and madagasikarioids.

DISCUSSION

This is the first complete generic-level analysis of Malpighiaceae and one of the largest to date for a major tropical angiosperm clade. Furthermore, it is one of the largest phylogenetic studies to include a companion, morphological data set. The morphological data set will be invaluable for ongoing taxonomic revisions of the family and for future analyses of floral evolution (see below). Importantly, the results presented here meet the goals of our study to (1) resolve relationships among the 11 segregates from the New World genus *Mascagnia* s.l., (2) test the monophyly of the largest remaining Malpighiaceae genera, and (3) clarify the placement of Old World Malpighiaceae. We shall summarize the most novel developments here and highlight areas where new challenges have arisen.

I. Mascagnia *s.l.*—*Mascagnia s.l.* contains all Malpighiaceae species with lateral-winged samaras, 10 fertile stamens, and internal stigmas that do not readily fit among other taxa with similar morphologies, including *Hiraea* Jacq., *Jubelina* Adr. Juss., *Mezia* Nied., *Tetrapterys*, and *Triopterys* L. (Anderson, 2006). It has been known that *Mascagnia s.l.* is not monophyletic (Cameron et al., 2001; Davis et al., 2001), but the number and placement of its constituent subclades have been unclear, and until very recently morphology has similarly not provided a clear direction for dividing this group into monophyletic sub-units (Anderson, 2006).

On the basis of our preliminary phylogenetic results and morphology, W. Anderson (2006) and his colleagues (Anderson and Davis, 2007; Anderson and Corso, 2007) recognized 10 generic segregates from Mascagnia: Adelphia W. R. Anderson, Aenigmatanthera W. R. Anderson, Alicia W. R. Anderson, Amorimia, Calcicola W. R. Anderson & C. Davis, Carolus W. R. Anderson, Christianella W. R. Anderson, Malpighiodes Nied., Niedenzuella W. R. Anderson, and Psychopterys W. R. Anderson & S. Corso. We show here for the first time that these former members of Mascagnia, plus Malpighia leticiana W. R. Anderson and Callaeum Small, which have previously been included in Mascagnia, belong to 11 clades variously scattered across Malpighiaceae (in Fig. 1, *Callaeum* and *Alicia* form a clade that has little support [51 BP], and so we are considering them separately in this calculation). All of the new generic segregates of *Mascagnia* s.l. that have been sampled with two or more species are monophyletic with >78 BP, with the exception of *Niedenzuella* (see below). Mascagnia s.s. is now greatly reduced and consists of about 37 species. These species are vines with interpetiolar stipules, and they possess glands impressed in the abaxial surface of the lamina, glabrous petals exposed in the bud, orbicular membranous samaras with arching and anastomosing veins in the lateral wing, and a three-lobed disc that subtends the fruit (Anderson and Davis, 2005a). There is not space to discuss all of these many new generic segregates here. Instead, we refer the reader to recently published morphological and taxonomic accounts of these groups (Anderson and Davis, 2005a, b; Anderson, 2006; Anderson and Corso, 2007; Anderson and Davis, 2007).

Niedenzuella—*Niedenzuella* comprises former members of *Mascagnia* and *Tetrapterys* (Anderson, 2006) and forms a wellsupported (91 BP) clade with *Aenigmatanthera*. This is supported by their many shared morphological traits, including decussate pseudoracemes, marginal leaf glands, tiny epipetiolar stipules, and yellow petals (Anderson, 2006). *Niedenzuella*,



Fig. 1. One of the maximum likelihood (ML) topologies based on the four-gene-plus-morphology data set. ML bootstrap percentages >50 are indicated at the nodes. Generic names in parentheses reflect the previous taxonomic placement for those species. Former members of *Mascagnia* are highlighted in gray. The seven ancient, Old World clades are numbered; Old World species are shown in boldface. For Bayesian posterior probabilities see Appendix S1 in the Supplemental Data accompanying the online version of this article. Three hundred fifty-seven accessions were included for this analysis, but only 356 are shown; *Peridiscus lucidus* Benth. was used for rooting, but has been removed for brevity.

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Fig. 2. Summary of major clades of Malpighiaceae emerging from this study derived from Fig. 1. Maximum likelihood bootstrap percentages >50 are indicated at nodes.

however, is not monophyletic—*N. stannea* is more closely related to *Aenigmatanthera* with 91 BP than it is to the remaining species of *Niedenzuella*. To avoid a proliferation of generic names, we intend to expand *Niedenzuella* and reduce *Aenigmatanthera* to a synonym of the former.

Triopterys—*Triopterys* consists of five species restricted to the West Indies. It has been distinguished from other genera principally on the basis of its Y-shaped samaroid fruit, vs. the butterfly-shaped, X-shaped, or orbicular samaras of *Hiraea*, *Tetrapterys*, and *Mascagnia* s.s., respectively. Our tree (Fig. 1) shows that *Triopterys* is well placed (86 BP) in the *Mascagnia* s.s. clade. We intend to combine the two genera under the name *Mascagnia* and propose conserving that name against the older name *Triopterys* in the near future.

II. Testing the monophyly of the largest remaining genera of Malpighiaceae-Beyond Mascagnia s.l., all of the other large genera of Malpighiaceae, with the exception of Bunchosia and Byrsonima, are not monophyletic as traditionally circumscribed (Fig. 1). These include Aspicarpa Rich. (and its related genera, in the aspicarpoid clade), Banisteriopsis, Heteropterys, Malpighia, Sphedamnocarpus Hook. f., Stigmaphyllon Adr. Juss., and Tetrapterys. Three of these genera, all of which are distantly related members of the large wing-fruited banisterioid clade, merit discussion together: Heteropterys, Banisteriopsis, and Sphedamnocarpus. These large genera bear a single prominent dorsal wing on each of their mericarps but otherwise resemble small genera whose mericarps bear a minimal dorsal crest or none at all, such that the mericarp resembles a nutlet instead of a samara. Our results demonstrate that these putative sister genera are not reciprocally monophyletic. Instead, each genus with wingless species is nested within a genus with winged fruits, rendering the latter nonmonophyletic: Clonodia is nested within Heteropterys with 100 BP, Diplopterys Adr. Juss. is nested within one of the major subclades of Banisteriopsis sensu Gates (1982) with 100 BP, and Philgamia is nested within Sphedamnocarpus with 99 BP. In each of these cases, it appears that the principal wing has been secondarily lost or greatly reduced. We will consider each of these instances below and follow with discussion of the remaining large or problematic genera of Malpighiaceae.

Heteropterys—The sampling of this large genus was greatly expanded here, with the result that the monophyly of the genus receives 98 BP. This contrasts with previous results that indicated very low support for the monophyly of Heteropterys (Cameron et al., 2001; Davis et al., 2001). In addition, for the first time, we were able to include the wingless Clonodia, whose phylogenetic affinities have long been problematic (Anderson, 1981). Species of *Clonodia* resemble some species of *Mascagnia*, most notably in their alate pink and white petals and racemose inflorescences. However, the same features are found in some species of *Heteropterys*, which also share the lenticels and subdistinct carpels of *Clonodia*. On this basis, Anderson (1981) thought that an origin of *Clonodia* from a *Heteropterys*-like ancestor was more likely. Our analyses indicate that *Clonodia* is indeed sister to two pink-flowered members of Heteropterys series Rhodopetalis Nied., H. brachiata and H. rufula (Fig. 1). Although support for the placement of *Clonodia* with the mostly pink-flowered *Heteropterys* is not strong (<50 BP), it is well nested (100 BP) within Heteropterys, and as such, its continued recognition would make Heteropterys nonmonophyletic. To remedy this problem, we have treated Clonodia as a synonym of *Heteropterys* (Anderson and Davis, 2007).

Banisteriopsis-Banisteriopsis sensu Gates (1982) is not monophyletic-its members belong to three distinct and wellsupported (>80 BP) clades within the stigmaphylloids (Fig. 1), which correspond to the three subgenera recognized by Gates: subg. Banisteriopsis, subg. Hemiramma (Griseb.) B. Gates, and subg. Pleiopterys (Nied.) B. Gates. The wingless Diplopterys (represented in Fig. 1 by D. cabrerana) is nested within Banisteriopsis subg. Pleiopterys. To remedy this situation, we transferred members of subgenus Pleiopterys to an expanded Diplopterys, which now includes both wingless and wing-fruited species (Anderson and Davis, 2006). Members of Gates's subg. Hemiramma are now recognized as the genus Bronwenia (Anderson and Davis, 2007), whose species are characterized by marginal leaf glands (if any), flowers borne in short, dense pseudoracemes, calyx glands attached below the free part of the sepals, yellow, usually glabrous petals, and samaras with the nut smooth or bearing a single ridge or winglet, and the carpophore absent or short. Members of subg. Banisteriopsis s.s. comprise the largest number of species of Banisteriopsis s.l. and

Sphedamnocarpus—Sphedamnocarpus is disjunctly distributed between Africa and Madagascar; the wingless Philgamia is wholly Malagasy. The African species [Sphedamnocarpus angolensis (Adr. Juss.) Oliv., S. galphimiifolius (Adr. Juss.) Szyszyl., and S. pruriens (Adr. Juss.) Szyszyl. in Fig. 1] form a well-supported subclade (100 BP), the Malagasy species of Sphedamnocarpus and Philgamia form a sister subclade with similar support (99 BP), and the two subclades together form a clade with similarly high support (100 BP). These results are reinforced by floral morphology: Malagasy Sphedamnocarpus and Philgamia have white, radial flowers, while those of the African Sphedamnocarpus are yellow or cream-colored and zygomorphic. Biogeography is obviously an important indicator of relationships in this clade. Under the circumstances, the later name Philgamia will have to be reduced to synonymy under Sphedamnocarpus unless all the Malagasy species of Sphedamnocarpus, including Philgamia, are determined to merit recognition at the level of genus. In that case, the Malagasy genus will bear the name Philgamia.

Stigmaphyllon-Members of the Old World Australasian clade Ryssopterys are well nested (86 BP) within the large, wellsupported (100 BP) New World clade Stigmaphyllon (Fig. 1). Ryssopterys species resemble Stigmaphyllon in most aspects of their morphology: the two groups have nearly identical leaves, usually consisting of a long petiole bearing a pair of conspicuous glands at the apex and a large, often cordate lamina. In both, the flowers are borne in umbels or pseudoracemes disposed in a dichasial inflorescence, and the samaras are identical (Anderson, 1997). The principal difference between them is that in Ryssopterys the sepals lack abaxial glands, the corolla is radial, and the styles have terminal stigmas, whereas in Stigmaphyllon the lateral sepals bear large paired abaxial glands, the corolla is strongly bilateral, and the styles are stigmatic on the internal angle of the apex and often bear an abaxial foliole that gives the genus its name (Anderson, 1997). Ryssopterys is also notable for having in each species some plants with staminate flowers and others with morphologically bisexual flowers (Yampolsky and Yampolsky, 1922; Christiane Anderson, personal communication, University of Michigan), while species of Stigmaphyllon bear only bisexual flowers (Anderson, 1997). Christiane Anderson has recently revised Ryssopterys and will soon publish a treatment that will resolve both the generic status and the species taxonomy of this group (Anderson, in press).

The aspicarpoid clade—Members of this well-supported (100 BP) stigmaphylloid subclade include the genera *Aspicarpa*, *Gaudichaudia* H.B.K., *Janusia*, and *Peregrina* W. R. Anderson. The aspicarpoid clade is well supported by morphology and is distinguished by the loss of four or all five of the stamens opposite the petals and the possession of a single style borne on the anterior carpel.

Janusia s.l. is disjunctly distributed between North and South America (Anderson, 1982, 1987). However, the North American species of Janusia are not most closely related to the South American species of Janusia. Instead, Janusia in North America is sister to all of the remaining aspicarpoids. To correct this problem, we resurrected the generic name *Cottsia* Dubard & Dop for the North American species of *Janusia* (Anderson and Davis, 2007). In addition, we have adopted here the already-published combination *Janusia linearifolia* (A. St.-Hil.) Adr. Juss. for *Peregrina linearifolia* (A. St.-Hil.) W. R. Anderson. Despite these taxonomic improvements, however, *Aspicarpa, Gaudichaudia*, and the remaining members of *Janusia* s.s. are not monophyletic. The best course of treatment of this group remains to be determined and will be a focus of our future research.

Malpighia—Our results demonstrate that the peculiar species Mascagnia leticiana W. R. Anderson is nested (76 BP) within Malpighia (Fig. 1). Mascagnia leticiana shares a combination of characters that link it either to Malpighia or to Mascagnia s.s., including similar stipules, leaf glands, petal color, androecium, and gynoecium (Anderson and Davis, 2005b). On the other hand, the needle-like hairs of M. leticiana and its tree habit make it a bad fit with Mascagnia s.s. and a better fit with Malpighia. The difficulty of placing Mascagnia leticiana in *Malpighia*, however, is that its fruits are strikingly different from those of Malpighia. Whereas the fruit is indehiscent and fleshy in Malpighia (i.e., drupelike) and dry, dehiscent, and samaroid in Mascagnia, the fruit of Mascagnia leticiana is intermediate, i.e., it is small and rather fleshy, but conspicuously winged. Given the phylogenetic placement of M. leticiana, it would appear that this represents a striking reversal to a presumably dehiscent Mascagnia-like fruit within a lineage of indehiscent fleshy-fruited species. Anderson and Davis (2005b) transferred M. leticiana to Malpighia, on the basis of these results.

Tetrapterys—Even after the segregation of *Niedenzuella* (discussed above) from *Tetrapterys*, the genus may not be monophyletic: its two constituent well-supported (100 BP) subclades may not be each other's closest relatives (Fig. 1). This is not entirely surprising given the morphological diversity of the species that have traditionally been assigned to this genus. Resolution of this problem will have high priority among the things we do next and should be done carefully with additional data from morphology and molecular sequences.

Lophanthera—Lophanthera Adr. Juss. and Spachea Adr. Juss. are sister genera and present an interesting problem. Lophanthera comprises five species, four in Amazonian South America and one in the lowlands of Costa Rica (Anderson et al., 2006). Spachea comprises six species: five in northern South America, Panama, and the lowlands of Costa Rica and Nicaragua and one in Cuba (Anderson et al., 2006). The two genera are both distinguished by putative morphological synapomorphies, so it was a surprise to find the Central American species of Lophanthera, L. hammelii W. R. Anderson, more closely related to Spachea than to other species of Lophanthera (Fig. 1). It is intriguing that *L. hammelii* and *S. correae* Cuatrec. & Croat are sympatric in Costa Rica (W. Anderson, 2007), but we do not want to read too much into those distributions or into the fact that L. hammelii is disjunct from its Amazonian congeners. Other similar disjunctions involving monophyletic taxa are known within the family (e.g., Dicella Griseb. in Costa Rica, W. Anderson, 2007). Moreover, it is very rare for us to encounter cases where there is such a disagreement between evidence from morphology and molecular sequences, so we do not intend to propose any taxonomic adjustments until this group receives more thorough study.

III. Phylogenetic placement of Old World Malpighiaceae— One of our most significant findings is the placement of the Old World clades of Malpighiaceae. We identified seven Old World clades and their closest New World relatives, in nearly all instances with 85 BP or greater (Fig. 1). Except for the acridocarpoid clade, all of these Old World clades are scattered deep within the predominantly New World banisterioid clade. If we also include the New World species *Heteropterys leona* (Cav.) Exell and *Stigmaphyllon bannisterioides* (L.) C. Anderson, which are native to the New World but have representative populations in coastal West Africa, the total number of Old World clades increases to nine. These findings will greatly help to produce an improved knowledge of the biogeography and floral evolution within the family, which we discuss below, and will be critical for future studies.

Biogeography of Malpighiaceae—Anderson (1979, 1990) argued on morphological grounds that the Malpighiaceae originated in South America in isolation from Africa. Molecular analyses (Cameron et al., 2001; Davis et al., 2001, 2002b, 2004) have similarly supported a likely New World origin for the family. In our previous studies (Cameron et al., 2001; Davis et al., 2001), there was still uncertainty about how many times the Malpighiaceae migrated from the New World to the Old World, because the placement of the Old World clades was not fully resolved.

It is now clear from the results presented here that the Malpighiaceae reached the Old World a total of nine times (Fig. 1). Two of those migrants are the species *Heteropterys leona* and *Stigmaphyllon bannisterioides*, both members of large New World genera and both found on the Atlantic coasts of Central America, the Caribbean, and South America as well as on the coast of West Africa. Those species have fruits that appear to be adapted for dispersal by water (e.g., arenchymatous fruit walls and/or dissected outgrowths on the fruit wall that presumably trap air and promote buoyancy; Anderson, 1997, 2001). These two species may have reached Africa via water dispersal across the Atlantic relatively recently, but long enough ago for both to have established extensive populations in coastal Africa (for a map showing the African distribution of *H. leona*, see Anderson et al., 2006).

The seven other Old World clades have diverged significantly from their New World relatives, suggesting much earlier arrivals in the Old World, likely during the Cenozoic (Davis et al., 2002b, 2004). Six of these clades, all consisting of only one or two genera, receive 99–100 BP. The madagasikarioids are the seventh and by the far the largest Old World clade, comprising eight genera. That clade is moderately supported (65 BP) here, but well supported elsewhere with increased taxon sampling and more rapidly evolving gene regions (Davis, 2002). Six of these Old World clades are well placed (\geq 86 BP) with their closest New World relatives (Fig. 1), while the placement of the seventh, the hiptageoid clade, has less support (59 BP) but is likely to be closely related to the New World genera *Carolus, Dicella*, and *Tricomaria*.

Anderson (1990) has suggested that the Malpighiaceae reached the Old World by dispersing across the southern Atlantic (without suggesting when that occurred or how great a gap existed at the time). Davis et al. (2002b, 2004) have argued that long-distance dispersal across the southern Atlantic is unlikely. Instead, they suggested that Malpighiaceae dispersed into North America from South America via the Caribbean Basin, crossed the North Atlantic into Eurasia, and subsequently reached the Old World tropics when paleoland and paleoclimatic configurations could have facilitated this migration. These ideas are testable, and the way forward lies in combining additional information from DNA sequences, morphology, fossils, and past and present-day distributions of the many New and Old World clades identified here. This study will allow us to better focus our ongoing efforts to obtain a much more thoughtful geographic sampling of species, especially from within those predominantly New World clades that also include Old World taxa. Such an approach will allow us to make better inferences on the routes followed by Malpighiaceae following their departure from South America.

IV. Floral evolution—Specialist pollination systems are often invoked as key factors underlying floral diversification as suggested, for example, in studies of orchids (Darwin, 1862; van der Pijl and Dodson, 1966) and Polemoniaceae (Grant and Grant, 1965; Stebbins, 1974). In some cases, however, specialized pollination systems may constrain floral morphological evolution. Neotropical species of Malpighiaceae may illustrate this pattern. Most are pollinated by specialist oil-collecting bees (Vogel, 1974), a relationship that appears to be ancient (Taylor and Crepet, 1987) and show relatively little floral diversity in contrast to diverse fruit structure and habit type (Anderson, 1979). A major question of evolutionary biologists is how such conserved morphologies are maintained over long periods of time, even in the face of diversification into different ecological roles (Beldade et al., 2002; Brakefield and Roskam, 2006). Are they the result of intrinsic genetic and developmental constraints, or are they labile and actively maintained by extrinsic factors (in this case, their oil-bee pollinators) over millions of years?

Malpighiaceae are a natural laboratory for testing these hypotheses because they comprise multiple sister-pairs of New World lineages that have maintained their oil-bee pollinators and Old World lineages that have lost them (the oil-bees that pollinate most New World Malpighiaceace are not present in the Old World; Vogel, 1990; Michener, 2000). New World Malpighiaceace are especially species-rich, yet they exhibit a conserved floral morphology (Fig. 3), which appears to be tightly associated with their oil-bee pollinators (Vogel, 1974, 1990; Anderson, 1979, 1990; Sigrist and Sazima, 2004; Gaglianone, 2005). In contrast, the Old World genera are relatively speciespoor, yet they display a diverse array of floral morphologies (Fig. 3). The majority of these Old World species lack most of the characteristic floral features critical to the pollination syndrome of New World Malpighiaceae, including especially the unique bilateral symmetry of the corolla (one dorsal petal functioning as a flag and the four lateral petals forming mirror-image pairs) and paired oil glands on the abaxial side of four or all five sepals. In the few Old World species that maintain some calyx glands (viz., African Acridocarpus Guill. & Perr. and Australasian Hiptage Gaertn.), the glands have shifted from producing oil to sugar (Lobreau-Callen, 1989). In most Old World Malpighiaceace, the only obvious reward for pollinators is pollen.

These observations suggest that the stereotyped New World flower may be maintained by selection by their oil-bee pollinators, not by intrinsic genetic constraints, and that when lineages migrated to the Old World where those pollinators were not found their floral morphology soon departed from that of their ancestors. This conclusion is also supported by (1) recent developmental and genetic data indicating that these changes may be relatively labile at the genetic level (Zhang et al., 2010) and (2) by those few New World Malpighiaceae species that have apparently abandoned pollination by oil-bees. An excellent example is *Psychopterys*, a Mexican/Central American genus of



Fig. 3. Corolla morphology of Malpighiaceae of the seven New and Old World sister clades highlighted in Fig. 1. Seven panels (all but that at lower right) illustrate the New World flower morphology contrasted with a representative from its Old World sister clade on the right. None of those Old World clades have maintained the stereotypical corolla symmetry of their New World sisters, which is ancestral in the family (Zhang et al., 2010). The three Old World genera that have bilaterally symmetrical corollas (*Acridocarpus, Tristellateia* Thouars, and *Sphedamnocarpus*) possess two dorsal petals (vs. only one in the New World) and one ventral petal, so the whole appearance of the flower is radically different from what a pollinator sees on approaching a flower in the New World. In the other four Old World genera (*Flabellariopsis* Wilczek, *Flabellaria* Cav., *Ryssopterys*, and *Madagasikaria*), the corolla is radially symmetrical. The eighth panel (lower right) compares two New World genera in the same clade: *Hiraea*, with a bilaterally symmetrical corolla and calyx glands for pollination by oil-bees, and *Psychopterys*, with a radially symmetrical corolla and eglandular sepals.

the hiraeoid clade that completely lacks calyx glands and has radially symmetrical corollas, quite different from those of other genera of that clade, which have the typical New World morphology (Fig. 3; Anderson and Corso, 2007). In *Galphimia* Cav., C. Anderson (2007) found something similar: the calyx glands are rudimentary or absent, and while it is often still possible to identify which is the flag petal, the overall symmetry of the corolla is much closer to radial than that of most New World Malpighiaceace. A final example is the Mexican genus *Lasiocarpus* Liebm. and its South American sister, *Ptilochaeta* Turcz. In this case, both genera lack calyx glands and possess radially symmetrical corollas (W. Anderson and C. Davis, unpublished data).

To more thoroughly address the question of developmental and genetic constraints vs. active maintenance, we need to integrate these morphological and phylogenetic insights into a strong analytical framework. Furthermore, it will be crucial to further examine the underlying genetic architecture of the floral morphology in Malpighiaceace (cf. Zhang et al., 2010). The phylogenetic results we present here represent an essential step toward making these kinds of evolutionary analyses possible and are a major focus of inquiry in Davis's laboratory.

Future directions—The analyses reported here have done a great deal to resolve infrafamilial relationships in the Malpighiaceae, but as described above, there remains a need for better resolution in several parts of the tree, especially in the tetrapteroids and stigmaphylloids. We are hopeful that progress can be achieved using two approaches to resolve these problems. The first will be to continue to expand character sampling across many members of the family using additional low copy nuclear genes, which have been shown to be very informative for resolving infrafamilial relationships in Malpighiaceae (Davis, 2002; Davis et al., 2002b) and relationships within the larger clade Malpighiales (Davis and Chase, 2004; Wurdack and Davis, 2009). One candidate is exon 9 of EMB2765, which has been useful for resolving relationships across the order Malpighiales (Wurdack and Davis, 2009). The second will be to greatly expand the number of characters using a phylogenomic approach, i.e., sampling a very large number of characters across a carefully selected subset of taxa representing all major clades within the family. These placeholder taxa can then be combined as a "scaffold" (Wiens et al., 2005; Wiens, 2006; Xi et al., 2010) with data sets that include greater taxon sampling (but far fewer characters) and analyzed simultaneously as a large supermatrix. The addition of a large number of more slowly evolving gene regions has been especially helpful in resolving rapid radiations deeper in Malpighiales (Davis et al., 2007), as well as in Saxifragales (Jian et al., 2008).

LITERATURE CITED

- ANDERSON, C. 1997. Monograph of Stigmaphyllon (Malpighiaceae). Systematic Botany Monographs 51: 1–313.
- ANDERSON, C. 2001. The identities of two water-dispersed species of *Heteropterys* (Malpighiaceae): *H. leona* and *H. platyptera. Contributions* from the University of Michigan Herbarium 23: 35–47.
- ANDERSON, C. 2007. Revision of *Galphimia* (Malpighiaceae). *Contributions* from the University of Michigan Herbarium 25: 1–82.
- ANDERSON, C. In press. Revision of *Ryssopterys* and transfer to *Stigmaphyllon* (*Malpighiaceae*). Blumea.
- ANDERSON, W. R. 1979. Floral conservatism in neotropical Malpighiaceae. *Biotropica* 11: 219–223.

- ANDERSON, W. R. 1981. Malpighiaceae in the botany of the Guayana Highland—Part XI. Memoirs of the New York Botanical Garden 32: 21–305.
- ANDERSON, W. R. 1982. Notes on neotropical Malpighiaceae—I. Contributions from the University of Michigan Herbarium 15: 93–136.
- ANDERSON, W. R. 1987. Notes on neotropical Malpighiacecae—II. Contributions from the University of Michigan Herbarium 16: 55–108.
- ANDERSON, W. R. 1990. The origin of the Malpighiaceae—The evidence from morphology. *Memoirs of the New York Botanical Garden* 64: 210–224.
- ANDERSON, W. R. 2004. Malpighiaceae (Malpighia family). *In* N. Smith, S. A. Mori, A. Henderson, D. W. Stevenson, and S. V. Heald [eds.], Flowering plants of the neotropics, 229–232. Princeton University Press, Princeton, New Jersey, USA, in association with The New York Botanical Garden, Bronx, New York, USA.
- ANDERSON, W. R. 2006. Eight segregates from the neotropical genus Mascagnia (Malpighiaceae). Novon 16: 168–204.
- ANDERSON, W. R. 2007. Malpighiaceae. In B. E. Hammel, M. H. Grayum, C. Herrera, and N. Zamora [eds.], Manual de plantas de Costa Rica, vol. 6, 253–312. Monographs in Systematic Botany from the Missouri Botanical Garden, vol. 111. Missouri Botanical Garden, St. Louis, Missouri, USA.
- ANDERSON, W. R., C. ANDERSON, AND C. C. DAVIS. 2006 onward [continuously updated]. Malpighiaceae [online]. Website http://herbarium. lsa.umich.edu/malpigh [accessed 05 November 2009].
- ANDERSON, W. R., AND S. CORSO. 2007. Psychopterys, a new genus of Malpighiaceae from Mexico and Central America. Contributions from the University of Michigan Herbarium 25: 113–135.
- ANDERSON, W. R., AND C. C. DAVIS. 2005a. The Mascagnia cordifolia group (Malpighiaceae). Contributions from the University of Michigan Herbarium 24: 33–44.
- ANDERSON, W. R., AND C. C. DAVIS. 2005b. Transfer of Mascagnia leticiana to Malpighia (Malpighiaceae). Contributions from the University of Michigan Herbarium 24: 45–49.
- ANDERSON, W. R., AND C. C. DAVIS. 2006. Expansion of *Diplopterys* at the expense of *Banisteriopsis* (Malpighiaceae). *Harvard Papers in Botany* 11: 1–16.
- ANDERSON, W. R., AND C. C. DAVIS. 2007. Generic adjustments in neotropical Malpighiaceae. Contributions from the University of Michigan Herbarium 25: 137–166.
- BELDADE, P., K. KOOPS, AND P. M. BRAKEFIELD. 2002. Developmental constraints versus flexibility in morphological evolution. *Nature* 416: 844–847.
- BRAKEFIELD, P. M., AND J. C. ROSKAM. 2006. Exploring evolutionary constraints is a task for an integrative evolutionary biology. *American Naturalist* 168: S4–S13.
- CAMERON, K. M., M. W. CHASE, W. R. ANDERSON, AND H. G. HILLS. 2001. Molecular systematics of Malpighiaceae: Evidence from plastid *rbcL* and *matK* sequences. *American Journal of Botany* 88: 1847–1862.
- DARWIN, C. 1862. On the various contrivances by which orchids are fertilized by insects. John Murray, London, UK.
- DAVIS, C. C. 2002. Madagasikaria (Malpighiaceae): A new genus from Madagascar with implications for floral evolution in Malpighiaceae. American Journal of Botany 89: 699–706.
- DAVIS, C. C., W. R. ANDERSON, AND M. J. DONOGHUE. 2001. Phylogeny of Malpighiaceae: Evidence from chloroplast *ndhF* and *trnL-F* nucleotide sequences. *American Journal of Botany* 88: 1830–1846.
- DAVIS, C. C., C. D. BELL, P. W. FRITSCH, AND S. MATHEWS. 2002a. Phylogeny of *Acridocarpus-Brachylophon* (Malpighiaceae): Implications for Tertiary tropical floras and Afroasian biogeography. *Evolution* 56: 2395–2405.
- DAVIS, C. C., C. D. BELL, S. MATHEWS, AND M. J. DONOGHUE. 2002b. Laurasian migration explains Gondwanan disjunctions: Evidence from Malpighiaceae. *Proceedings of the National Academy of Sciences*, USA 99: 6833–6837.
- DAVIS, C. C., AND M. W. CHASE. 2004. Elatinaceae are sister to Malpighiaceae; Peridiscaceae belong to Saxifragales. *American Journal of Botany* 91: 262–273.

- DAVIS, C. C., P. W. FRITSCH, C. D. BELL, AND S. MATHEWS. 2004. High latitude Tertiary migrations of an exclusively tropical clade: Evidence from Malpighiaceae. *International Journal of Plant Sciences* 165: S107–S121.
- DAVIS, C. C., C. O. WEBB, K. J. WURDACK, C. A. JARAMILLO, AND M. J. DONOGHUE. 2005. Explosive radiation of Malpighiales supports a mid-Cretaceous origin of tropical rain forests. *American Naturalist* 165: E36–E65.
- DAVIS, C. C., M. LATVIS, D. L. NICKRENT, K. J. WURDACK, AND D. A. BAUM. 2007. Floral gigantism in Rafflesiaceae. *Science* 315: 1812. FRENOPOULO, C. 2005. The ritual use of ayahuasca. *Journal of*
- FRENOPOULO, C. 2005. The ritual use of ayahuasca. Journal of Psychoactive Drugs 37: 237–239.
- GAGLIANONE, M. C. 2005. Nesting biology, seasonality, and flower hosts of *Epicharis nigrita* (Friese, 1900) (Hymenoptera: Apidae: Centridini), with a comparative analysis for the genus. *Studies on Neotropical Fauna and Environment* 40: 191–200.
- GATES, B. 1982. Banisteriopsis, Diplopterys (Malpighiaceae). Flora Neotropica Monograph 30: 1–237.
- GRANT, V., AND K. A. GRANT. 1965. Flower pollination in the phlox family. Columbia University Press, New York, New York, USA.
- HOLMGREN, P. K., N. H. HOLMGREN, AND L. C. BARNETT. 1990. Index herbariorum, part 1: The herbaria of the world. New York Botanical Garden, Bronx, New York, USA.
- HUELSENBECK, J. P., AND F. RONQUIST. 2001. MRBAYES: Bayesian inference of phylogenetic trees. *Bioinformatics* 17: 754–755.
- HUTCHINSON, J. 1967. The genera of flowering plants. Clarendon Press, Oxford, UK.
- JIAN, S. G., P. S. SOLTIS, M. A. GITZENDANNER, M. J. MOORE, R. LI, T. A. HENDRY, Y. L. QIU, ET AL. 2008. Resolving an ancient, rapid radiation in Saxifragales. *Systematic Biology* 57: 38–57.
- LEWIS, P. O. 2001. A likelihood approach to estimating phylogeny from discrete morphological character data. *Systematic Biology* 50: 913–925.
- LOBREAU-CALLEN, D. 1989. Les Malpighiaceae et leurs pollinisateurs: Coadaptation ou coévolution. *Bulletin du Muséum National d'Histoire Naturelle, série 4, B, Adansonia* 11: 79–94.
- MADDISON, W. P., AND D. R. MADDISON. 2000. MacClade version 4: Analysis of phylogeny and character evolution. Sinauer, Sunderland, Massachusetts, USA.
- MICHENER, C. D. 2000. The bees of the world. Johns Hopkins University Press, Baltimore, Maryland, USA.
- NIEDENZU, F. 1928. Malpighiacecae. Das Pflanzenreich IV 141: 1-870.
- POSADA, D., AND T. R. BUCKLEY. 2004. Model selection and model averaging in phylogenetics: Advantages of Akaike information criterion and Bayesian approaches over likelihood ratio tests. *Systematic Biology* 53: 793–808.

- POSADA, D., AND K. A. CRANDALL. 1998. MODELTEST: Testing the model of DNA substitution. *Bioinformatics* 14: 817–818.
- SIGRIST, M. R., AND M. SAZIMA. 2004. Pollination and reproductive biology of twelve species of neotropical Malpighiaceae: Stigma morphology and its implications for the breeding system. *Annals of Botany* 94: 33–41.
- STAMATAKIS, A. 2006. RAXML-VI-HPC: Maximum likelihood-based phylogenetic analyses with thousands of taxa and mixed models. *Bioinformatics* 22: 2688–2690.
- STEBBINS, G. L. 1974. Flowering plants: Evolution above the species level. Harvard University Press, Cambridge, Massachusetts, USA.
- TAYLOR, D. W., AND W. L. CREPET. 1987. Fossil floral evidence of Malpighiaceae and an early plant–pollinator relationship. *American Journal of Botany* 74: 274–286.
- TOKUOKA, T., AND H. TOBE. 2006. Phylogenetic analyses of Malpighiales using plastid and nuclear DNA sequences, with particular reference to the embryology of Euphorbiaceae sens. str. *Journal of Plant Research* 119: 599–616.
- VAN DER PIJL, L., AND H. DODSON. 1966. Orchid flowers: Their pollination and evolution. University of Miami Press, Coral Gables, Florida, USA.
- VOGEL, S. 1974. Ölblumen und ölsammelnde Bienen. Tropische und Subtropische Pflanzenwelt 7: 283–547.
- VOGEL, S. 1990. History of the Malpighiaceae in the light of pollination ecology. *Memoirs of the New York Botanical Garden* 55: 130–142.
- WIENS, J. J. 2006. Missing data and the design of phylogenetic analyses. *Journal of Biomedical Informatics* 39: 34–42.
- WIENS, J. J., J. W. FETZNER, C. L. PARKINSON, AND T. W. REEDER. 2005. Hylid frog phylogeny and sampling strategies for speciose clades. *Systematic Biology* 54: 778–807.
- WURDACK, K. J., AND C. C. DAVIS. 2009. Malpighiales phylogenetics: Gaining ground on one of the most recalcitrant clades in the angiosperm tree of life. *American Journal of Botany* 96: 1551–1570.
- XI, Z., B. R. RUHFEL, K. J. WURDACK, A. AMORIM, AND C. C. DAVIS. 2010. Complete plastid genomes do not resolve deep relationships in Malpighiales. Botany 2010: Annual meeting of the Botanical Society of America, Providence, Rhode Island, USA [online abstract, website http://2010.botanyconference.org/engine/search/index. php?func=detail&aid=581].
- YAMPOLSKY, C., AND H. YAMPOLSKY. 1922. Distribution of sex forms in the phanerogamic flora. *Bibliotheca Genetica* 3: 1–62.
- ZHANG, L.-B., AND M. P. SIMMONS. 2006. Phylogeny and delimitation of the Celastrales inferred from nuclear and plastid genes. *Systematic Botany* 31: 122–137.
- ZHANG, W., E. M. KRAMER, AND C. C. DAVIS. 2010. Floral symmetry genes and the origin and maintenance of zygomorphy in a plant-pollinator mutualism. *Proceedings of the National Academy of Sciences, USA* 107: 6388–6393.

APPENDIX 1. Taxa sequenced with	th distribution, voucher inf	formation, and GenBan	k accession numbers	. Herbarium acronyi	ns follow <i>Index</i> I	herbariorum (Holmgren
et al., 1990).							

				Plastid		Nuclear
Taxon	Distr. ^a	Voucher	ndhF	matK	rbcL	РНҮС
Celastraceae*						
Denhamia celastroides (F. Muell.) Jessup	OW	<i>Chase 2050</i> (K)	AY425043	EF135526	AJ402941	AY425097
Centroplacaceae*	OW		FI(700/5			
Bhesa paniculata Arn ‡	OW	Iakeuchi & Ama 10450 (GH) Sugumaran 118 (US)	FJ6/0065			EI660883
Bhesa robusta (Roxb.) Ding Hou	OW	Laman 283 (L)		FJ670001	AY935723	1.1009885
Centroplacus glaucinus Pierre†	OW	White 128, ser. 1 (MO)	FJ670066	FJ670002	AY663646	FJ669884
Chrysobalanaceae*						
Atuna racemosa Raf.	OW	<i>Chase 2118</i> (K)	AY425030	EF135503	AF089758	AY425088
Dichapetalaceae*	OW	$E_{incom} = 10/8/02 (V)$	AV425044	EE125527	A E080764	AV125008
Euphorbiaceae*	0	<i>Fisson s.n.</i> 10/8/95 (K)	AI 423044	EF133327	AF089704	A1423096
Acalypha californica Benth.	NW	Levin 2192 (SD)	AY425027	EF13549	AY380341	AY425087
Endospermum moluccanum (Teijsm. & Binn.) Kurz	OW	<i>Chase 1258</i> (K)	AY425051	EF135533	AJ402950	AY425106
Elatinaceae*						
Bergia pedicellaris (F. Muell.) Benth.†	OW	Harwood 1285 (A)	HQ246766	HQ247213	HQ247446	HQ246984
<i>Elating minima</i> (Nutt.) Fisch & C. A. Mey :	NW	Oldham 16691 (MICH)	HO246824	EF155500	AI 360344	HO247048
Elatine triandra Schkuhr	NW	Burton et al. 13384 (MICH)	AY425049	EF135532	AY380348	AY425104
<i>Elatine</i> sp.	OW	Qiu 99051 (Z)	HQ246825	HQ247273		HQ247049
Goupiaceae*						
Goupia glabra Aubl.	NW	Prevost 3031 (CAY)	AY425054	EF135544	AJ235780	AY425108
Ochnaceae*	OW	$D_{\text{residential}} 21.01(\Lambda)$	11/10/072	EI(70021	137290254	137405100
Ocnna sp. Peridiscaceae*	Uw	Davis 31-01 (A)	AI 425075	FJ0/0031	AI 380334	A1425125
Peridiscus lucidus Benth.	NW	Soares 205 (CEPEC)	AY425076	DO411570	AY380356	AY425125
Phyllanthaceae*				- (
Phyllanthus calycinus Labill.	OW	<i>Chase 2163</i> (K)	EU002250	EF135580	AY663603	AY579869
Picrodendraceae*						
Androstachys johnsonii Prain	OW	<i>Chase 1904</i> (K)	AF500495		AJ402922	AF500522
Putranjiva rozhurahij Wall	OW	Wurdack D57 (US)	AY425048	FF135530	M95757	AY425102
Violaceae*	011	Waldack D57 (05)	A1+230+6	LI 155550	W1/5757	A1423102
Hymenanthera alpina (Kirk) W. R. B. Oliv.	OW	<i>Chase 501</i> (K)	AY425059	EF135552	Z75692	AY425112
Malpighiaceae						
Acmanthera latifolia (Adr. Juss.) Griseb.†	NW	Anderson 13771 (MICH)	AF351013	AF344524	AF344454	AF500528
Acridocarpus adenophorus Adr. Juss.†	OW	McPherson 14406 (MICH)	AY 137243	HQ247173	HQ247415	AY499049
Acridocarpus alternifolius (Schumach. & Thonn.) Nied.	OW	Davis 99-10 (A)	HQ246730	HQ24/1/4	HQ24/416	HQ246944
Acridocarpus excelsus Adr Juss	OW	Rakotomalaza et al. 378 (MO)	AT 137247 AY 137248	HQ247175	HQ247417 HQ247418	AY499050
Acridocarpus macrocalyx Engl	OW	Davis 99-09 (A)	AF351017	HQ247177	HQ247419	AF500532
Acridocarpus natalitius Adr. Juss.†	OW	Goldblatt s.n. (PRE)	AF351016	AF344525	AF344455	AF500531
Acridocarpus orientalis Adr. Juss.	OW	Popov 85 (MO)	AY137255	HQ247178	HQ247420	AY499052
Acridocarpus scheffleri Engl.	OW	Mlangwa et al. 1417 (MO)	HQ246731	-	HQ247421	HQ246945
Acridocarpus smeathmannii (DC.) Guill. & Perr.	OW	Davis 99-13 (A)	AY137261	HQ247179	HQ247422	AY499053
Acridocarpus sp.	OW	Bourobou 939 (MO)	HQ246732	HQ247180		HQ246946
Acridocarpus spectabilis (Nied.) Doorn-Hoekm.	OW	<i>GC</i> 47679 (MO)	HQ246733	HQ247181	HQ247423	HQ246947
Acridocarpus staudtii (Engl.) Hutch. & Dalziel	OW	Davis 99-7 (A)	HQ246734	HQ247182	HQ247424	HQ246948
Acridocarpus zanzibaricus Adr. Juss.	OW	Davis 99-23 (A)	AY137264	HQ247183	HQ247425	AY499054
Adelphia hiraea (Gaertn.) W. R. Anderson [†]	NW	Anderson 13802 (MICH)	AF351059	AF344566	AF344498	AF500563
Aenigmatanthera doniana (Griseb.) W. R. Anderson	IN W	Prance & Henriques 29888 (MICH)	HQ246735		HQ247426	HQ246949
Aenigmatanthera lasiandra (Adr. Juss.) W. R. Anderson	IN W	Dubs 1400 (MICH)	HQ240737	HO247184	HQ247428	HQ240931
Achigmatanthera lasiandra (Adr. Juss.) W. R. Anderson		Equation of al. 1012 (MICH)	HQ240730	HQ247184	HQ247427	HQ240930
Alicia anisopatala (Adr. Juss.) W. R. Anderson [†]	NW	Anderson 13598 (MICH)	AF351053	AE3//163	AF3///0/	AE500557
Amorimia amazonica (Nied.) W.R. Anderson ⁺	NW	Steinbach 339 (MICH)	HO246739	HO247186	HO247430	HO246953
Amorimia camporum W. R. Anderson	NW	Campos & Díaz 2490 (MICH)	HQ246740	HQ247187	HQ247431	HQ246954
Amorimia exotropica (Griseb.) W. R. Anderson	NW	Vanni et al. 2903 (MICH)	HQ246741	HQ247188		HQ246955
Amorimia kariniana W. R. Anderson	NW	Asplund 16617 (S)	HQ246742	HQ247189	HQ247432	HQ246956
Amorimia rigida (Adr. Juss.) W. R. Anderson [†]	NW	Anderson 13691 (MICH)	AY499101	HQ247190	HQ247433	AY499075
Amorimia velutina W. R. Anderson	NW	Shepherd et al. 4409 (NY)	HQ246743	HQ247191		HQ246957
Aspicarpa brevipes (DC.) W. R. Anderson [†]	NW	Cházaro 6295 (MICH)	AF351082	AF344526	AF344456	AF500581
Aspicarpa harleyi W. R. Anderson†	NW	Anderson 13696 (MICH)	HQ246744	HQ247192	HQ247434	HQ246958
Aspicarpa hirtella Rich.†	NW	Anderson 13525 (MICH)	AF351033	AF344527	AF344457	AF500541
Aspicarpa hyssopifolia A. Gray	IN W	Anderson 13521 (MICH)	HQ246745	HQ24/193	4F211150	HQ246959
Aspicarpa paienena (Onseo.) O Donen & Lourleig	TN NA	лиин son 13014 (МПСП)	AF551020	AF344328	AF 344438	nQ240900

				Plastid		Nuclear
Taxon	Distr. ^a	Voucher	ndhF	matK	rbcL	РНҮС
Aspicarpa sericea Griseb. [†]	NW	Anderson 13586 (MICH)	HQ246747		HQ247436	HQ246962
Aspicarpa sericea Griseb.†	NW	Krapovickas & Schinini 39191 (MICH)	HQ246746	HQ247194	HQ247435	HQ246961
Aspidopterys sp.†	OW	Larsen & Larsen 34311 (P)	HQ263253	HQ263254		HQ263255
Aspidopterys sp.	OW	Bogor XVI.G.110 (BO) [<i>Chase M239</i> (K)]	AF351019	AF344529	AF344459	AF436796
Aspidopterys sp.	OW	Srisanga 1137 (P)	HQ246748	HQ247195	HQ263252	HQ246963
Aspidopterys tomentosa (Blume) Adr. Juss.†	OW	Tsi Zhanhuo 92-121 (MO)	HQ246749	HQ247196	HQ247437	HQ246964
Banisteriopsis acerosa (Nied.) B. Gates	NW	Guala 1370 (MICH)		HQ247197		HQ246965
Banisteriopsis adenopoda (Adr. Juss.) B. Gates	NW	Silva 1109 (MICH)	HQ246750		HQ247438	HQ246966
Banisteriopsis angustifolia (Adr. Juss.) B. Gates	NW	Carvalho 80 (MICH)	HQ246751	HQ247198	110247420	HQ246967
Banisteriopsis argyrophylla (Adr. Juss.) B. Gates	NW	Silva et al. 3548 (MICH)	HQ246752	HQ24/199	HQ247439	HQ246968
Banisteriopsis caapi (Griseb.) C. V. Morton	IN W	Anderson 13/91 (MICH)	HQ240753	HQ247200	HQ247440	HQ240909
Banisteriopsis calcicolu B. Gates	NW	Kranovickas & Cristóbal 44854	HQ240754	HQ247201		HQ240970
Bunisteriopsis conjusti B. Gates	19 99	(MICH)	11Q240755	11Q247202		11Q240971
Banisteriopsis gardneriana (Adr. Juss.) W. R. Anderson & B. Gates	NW	Azevedo & Lopes 313 (MICH)		HQ247203		HQ246972
Banisteriopsis harleyi B. Gates	NW	Carvalho 87 (MICH)	HQ246756	HQ247204		HQ246973
Banisteriopsis laevifolia (Adr. Juss.) B. Gates†	NW	Anderson 13631 (MICH)	HQ246757	HQ247205	HQ247441	HQ246974
Banisteriopsis latifolia (Adr. Juss.) B. Gates	NW	Azevedo 698 (MICH)	HQ246758	HQ247206		HQ246975
Banisteriopsis martiniana (Adr. Juss.) Cuatrec.†	NW	Chase 90161 (MICH)	HQ246759	HQ247207	HQ247442	HQ246976
Banisteriopsis muricata (Cav.) Cuatrec.†	NW	Anderson 13799 (MICH)	HQ246760	HQ247208	HQ247443	HQ246977
Banisteriopsis nummifera (Adr. Juss.) B. Gates	NW	Harley 54681 (MICH)	HQ246761	HQ247209	HQ247444	HQ246978
Banisteriopsis paraguariensis B. Gates	NW	Ferrucci et al. 1624 (MICH)	HQ246762	HQ247210		HQ246979
Banisteriopsis prancei B. Gates	NW	Anderson 9910 (MICH)	11001/17/2	110017011		HQ246980
Banisteriopsis pulchra B. Gates	NW	Dubs 2258 (MICH)	HQ246763	HQ24/211	110247445	HQ246981
Banisteriopsis schwanniolaes (Griseb.) B. Gates	NW	Morendro 278 (MICH)	HQ240704	ПQ247212	ПQ247443	HQ240982
Barnebya dispar (Griseb.) W. R. Anderson & B. Gates†	NW	Leoni 1473 (MICH)	AF351073	AF344531	AJ402924/	AF500574
Blepharandra fimbriata MacBryde‡	NW	Anderson 13882 (MICH)	HO246767		HO247447	HO246985
Blepharandra heteropetala W. R. Anderson†	NW	Anderson 13782 (MICH)	AF351008	AF344532	AF344461	AF500523
Blepharandra hypoleuca (Benth.) Griseb.	NW	Anderson 13844 (MICH)	HQ246768	HQ247214	HQ247448	HQ246986
Brachylophon curtisii Oliv.†	OW	Middleton et al. 387 (A)	AF351018	HQ247215	HQ247449	AY137337
Bronwenia cinerascens (Benth.) W. R. Anderson & C.	NW	Nee 48570 (MICH)	HQ246769	HQ247216		HQ246987
Davis						
Bronwenia cinerascens (Benth.) W. R. Anderson & C. Davis	NW	Nee 48324 (MICH)			HQ247450	
Bronwenia cornifolia (H. B. K.) W. R. Anderson & C. Davis†	NW	Clark 1 (MICH)	HQ246770	HQ247217	HQ247451	HQ246988
Bronwenia ferruginea (Cav.) W. R. Anderson & C. Davis	NW	Amorim et al. 3222 (MICH)	HQ246771	HQ247218	HQ247452	HQ246989
Bronwenia mathiasiae (W. R. Anderson) W. R. Anderson & C. Davis†	NW	Nelson et al. 518 (MICH)	HQ246772	HQ247219	HQ247453	HQ246990
Bunchosia angustifolia Adr. Juss.	NW	Combra S. 766 (MICH)	HQ246773	HQ247220		HQ246991
Bunchosia apiculata Huber	IN W	Gentry et al. 50215 (MICH)	HQ240774	HQ247221	775274	HQ240992
Bunchosia armeniaca (Cav.) DC.	NW	Nag 41005 (MICH)	HO246775	HO247222	213214	HO2/6003
Bunchosia decussiflora W R Anderson	NW	de Granville 4905 (MICH)	HQ246776	HQ247222		HQ246994
Bunchosia deflexa Triana & Planch.	NW	Gentry et al. 54765 (MICH)	HO246777	HQ247224		HO246995
Bunchosia glandulifera (Jacq.) H. B. K.	NW	Anderson 13761 (MICH)	HO246778	HO247225	HO247454	HO246996
Bunchosia glandulosa (Cav.) DC.	NW	Correll 50038 (MICH)				HQ246998
Bunchosia glandulosa (Cav.) DC.	NW	Goldman 3697 (GH)	HQ246779	HQ247226		HQ246997
Bunchosia macrophylla Rose	NW	Hammel 17747 (CR)	HQ246780	HQ247227	HQ247455	HQ246999
Bunchosia mcvaughii W. R. Anderson	NW	Evans s.n. (MICH)	HQ246781	HQ247228	HQ247456	HQ247000
Bunchosia mollis Benth.	NW	Jansen-Jacobs et al. 3958 (MICH)	HQ246782	HQ247229		HQ247001
Bunchosia nitida (Jacq.) DC.	NW	Silverstone-Sopkin & Paz 7096 (MICH)	HQ246783			HQ247002
Bunchosia odorata (Jacq.) DC.	NW	Flora Falcón 791 (MICH)	HQ246784	HQ247230		HQ247003
Bunchosia pilocarpa Rusby	NW	Smith et al. 14020 (MICH)	HQ246785	HQ247231		HQ247004
Bunchosia polystachia (Andrews) DC.	IN W	Ernst 1900 (MICH)	HQ246/86	HQ247232		HQ24/005
Dunchosia swartziana Griseb.	IN W	webster & Wilson 4820 (MICH)	U0216707	ПQ24/233 ПО247224	U02/7/57	HQ247007
Bunchosia veluticarna W.P. Anderson	NW	Anderson 13805 (MICH)	11Q240787 HO246789	HQ24/234	HQ24/43/	HO247007
Burdachia sphaerocarna Adr. Juss +	NW	Anderson 13774 (MICH)	AF351071	AF344534	AF344462	AF500572
Byrsonima basiloba Adr. Juss	NW	Anderson 13671 (MICH)	HO246789	HO247236	HO247459	HO247009
Byrsonima coccolobifolia H. B. K.†	NW	Anderson 13661 (MICH)	HQ246790	HQ247237	HQ247460	HQ247010
Byrsonima crassifolia (L.) H. B. K.†	NW	FTG 81-680A (MICH)	AF351011	AF344535	L01892	AF500526

				Plastid		Nuclear
Taxon	Distr. ^a	Voucher	ndhF	matK	rbcL	PHYC
Byrsonima crispa Adr. Juss.	NW	Anderson 13755 (MICH)	HQ246791	HQ247238	HQ247461	HQ247011
Byrsonima duckeana W. R. Anderson	NW	Anderson 13765 (MICH)	HQ246792	HQ247239	-	HQ247012
Byrsonima lucida (Mill.) DC.	NW	FTG 85-139 (MICH)	HQ246793	HQ247240	HQ247462	HQ247013
Byrsonima macrophylla (Pers.) W. R. Anderson	NW	Anderson 13647 (MICH)	HQ246794	HQ247241		HQ247014
Byrsonima morii W. R. Anderson	NW	Harley 26462 (MICH)	HQ246795		HQ247463	HQ247015
Byrsonima triopterifolia Adr. Juss.	NW	Anderson 13702 (MICH)	HQ246796	HQ247242	HQ247464	HQ247016
Calcicola parvifolia (Adr. Juss.) W. R. Anderson & C.	NW	Salinas T. 5444 (MICH)	HQ246797	HQ247243	HQ247465	HQ247017
Davis [†] Callagum antifabrila (Grisch) D. M. Johnson	NW	Morquetz & Wallnöfer 117 161085	HO246708	H0247244	H0247466	HO247018
Canaeum anitgebrite (Grisco.) D. M. Johnson	1 4 4 4	(MICH)	11Q240798	11Q247244	11Q247400	11Q247018
Callaeum clavipetalum D. M. Johnson	NW	Acosta C. 812 (MICH)	HO246799	HO247245	HO247467	HO247019
Callaeum malnighioides (Turcz.) D. M. Johnson	NW	Ventura A. 19165 (MICH)	HO246800	HO247246	HO247468	HO247020
Callaeum nicaraguense (Griseb.) Small	NW	Moreno 25336 (MICH)	HQ246801	HQ247247	HQ247469	HQ247021
Callaeum psilophyllum (Adr. Juss.) D. M. Johnson†	NW	FTG 61-291 (FTG)	-	HQ247248	-	HQ247023
Callaeum psilophyllum (Adr. Juss.) D. M. Johnson†	NW	Krapovickas & Cristóbal 44674	HQ246802	HQ247249	HQ247470	HQ247022
		(MICH)				
Callaeum septentrionale (Adr. Juss.) D. M. Johnson†	NW	Anderson 13491 (MICH)	AF351058	AF344536	AF344463	AF500562
Camarea axillaris A. StHil. [†]	NW	Amorim 3659 (CEPEC)	AY 499081	HQ247250	HQ247471	AY499055
Carolus anaerssonii (W. R. Anderson) W. R. Anderson	IN W	Anderson 12710 (MICH)	HQ240803	HQ247251	A E244406	HQ247024
Carolus chlorocarnus (Adr. Juss.) W. R. Anderson	NW	Argevedo et al. 1147 (MICH)	HO246804	HO247252	AF344490	HO247025
Carolus sinemariensis (Aubl.) W.R. Anderson ⁺	NW	Cochrane et al 12331 (MICH)	HQ240804	HQ247252	HO247472	HQ247023
Carolus sinemariensis (Aubl.) W. R. Anderson ⁺	NW	Henkel 1832 (MICH)	HQ246805	11Q2+7255	11Q2+7+72	HQ247027
Caucanthus auriculatus (Radlk.) Nied.†	OW	Kindeketa et al. 1527 (MO)	HO246807	HO247254	HO247473	HO247028
Caucanthus auriculatus (Radlk.) Nied.	OW	Knox 2128 (MICH)	AF351020	AF344537	AF344464	AF436797
Christianella multiglandulosa (Nied.) W. R. Anderson†	NW	Hahn 1736 (MICH)	HQ246808	HQ247255	HQ247474	HQ247029
Christianella surinamensis (Kosterm.) W. R. Anderson	NW	Killeen et al. 7674 (MICH)	HQ246809	HQ247256	HQ247475	HQ247030
Coleostachys genipifolia Adr. Juss. [†]	NW	Anderson 13752 (MICH)	AF351012	AF344538	AF344465	AF500527
Cordobia argentea (Griseb.) Nied.†	NW	Anderson 13583 (MICH)	AF351081	AF344539	AF344466	HQ247031
Cordobia argentea (Griseb.) Nied.†	NW	Zardini & Duarte 49837A (MICH)	HQ246810	1 52 4 4 5 5 4	1 22 4 4 40 4	1 5500520
Cottsia californica (Benth.) W. R. Anderson & C. Davist	NW	Anderson 12539 (MICH)	AF351031	AF344554	AF344486	AF500539
Diggidig formuging (Maguiro & K. D. Pholos) W. P.	IN W NIW	Anderson 12540 (MICH)	AF351030	AF344333	AF344487	AF500538
Anderson ⁺	IN W	Berry et al. 4987 (MICH)	AF551010	AF344340	AF544407	AF300323
Diacidia galphimioides (Maguire) W R Anderson	NW	Berry et al 5275 (MICH)	HO246811	HO247257	HO247476	HO247032
Diasnis albida Nied.†	OW	De Wilde 5920 (MO)	HO246813	HQ247259	HO247478	HQ247034
Diaspis albida Nied.†	OW	Robertson 5045 (MO)	HQ246812	HQ247258	HQ247477	HQ247033
Dicella bracteosa (Adr. Juss.) Griseb.	NW	Kallunki et al. 576 (MICH)	HQ246814	HQ247260	HQ247479	HQ247035
Dicella julianii (J. F. Macbr.) W. R. Anderson	NW	Burnham 1903 (MICH)	HQ246815	HQ247261	HQ247480	HQ247036
Dicella macroptera Adr. Juss.†	NW	Fonseca et al. 2257 (MICH)	HQ246816			HQ247037
Dicella macroptera Adr. Juss.†	NW	Fonseca et al. 2404 (MICH)	HQ246817	HQ247262		HQ247038
Dicella macroptera Adr. Juss.†	NW	Pereira 2047 (MICH)		HQ247263		HQ247039
Dicella nucifera Chodat	NW	Anderson 13607 (MICH)	AF351048	AF344541	AJ235802	AF500553
Digoniopterys microphylla Arenes	OW NW	Du Puy et al. MB08 (P) Simman 82, 10, 22, 20 (MICH)	HQ246818	HQ24/264	HQ24/481	HQ247040
Dinemagonum gayanum Adr. Juss.	IN W NW	Dillon & Tailliar 5103 (MICH)	AF351064	AE344542	AF344408	AE500570
Dinlonterus cabrerana (Cuatrec) B. Gates [†]	NW	Burnham 1774 (MICH)	AF351009	HO247266	HO247482	AF500570
Diplopterys eubreratia (edudee.) B. Gues	NW	Anderson 13638 (MICH)	AF351037	AF344530	AF344460	HO247042
C. Davis†						(
Diplopterys lutea (Griseb.) W. R. Anderson & C. Davis†	NW	FTG 60-85 (FTG)	HQ246819	HQ247267	HQ247483	HQ247043
Diplopterys nutans (Nied.) W. R. Anderson & C. Davis	NW	Matezki 175 (MICH)	HQ246820	HQ247268	HQ247484	HQ247044
Diplopterys pubipetala (Adr. Juss.) W. R. Anderson &	NW	Salino 3472 (MICH)	HQ246821	HQ247269	HQ247485	HQ247045
C. Davis						
Diplopterys valvata (W. R. Anderson & B. Gates) W. R.	NW	Lombardi 2203 (MICH)	HQ246822	HQ247270	HQ247486	HQ247046
Anderson & C. Davis	N T N 7		11004(000	110047071		110047047
Diplopterys virguitosa (Adr. Juss.) W. K. Anderson & C.	IN W	Fonseca et al. 2800 (MICH)	HQ240823	HQ247271		HQ247047
Echinopterys ealandulosa (Adr. Juss.) Small ⁺	NW	Van Devender 98-178 (MICH)	AE351047	AE3//5//3	AE3///70	AE500552
Ectopopterys sociartoi W R Anderson ⁺	NW	Calleias 11806 (MICH)	AF351064	HO247272	AF344471	AF500552
Excentradenia propingua (W. R. Anderson) W. R.	NW	Lindeman et al. 51 (MICH)	AF351062			AF350995
Anderson†						
Flabellaria paniculata Cav.†	OW	Congdon 414 (K)	AF351083		AF344472	
Flabellariopsis acuminata (Engl.) Wilczek†	OW	Luke & Robertson 2683 (MO)	HQ246826	HQ247274	AF344473	HQ247050
Gallardoa fischeri Hicken†	NW	Anderson 13580 (MICH)	AF351035	AF344544	AF344474	AF500543
Galphimia brasiliensis (L.) Adr. Juss.	NW	Alencar 3 (MICH)	HQ246827	HQ247275	HQ247487	HQ247051
Galphimia glandulosa Cav.	NW	Anderson & Anderson 5849 (MICH)	HQ246828			HQ247052
Galphimia glauca Cav.†	NW	Ventura & López 8481 (MICH)	A E251015	A E2 4 4 5 4 5	A E2 4 4 4 7 5	HQ247053
Gaipnimia gracilis Barti. ⁺	IN W	FIG /9-233 (FIG)	AF351015	AF344545	AF544475	AF500530

				Plastid		Nuclear
Taxon	Distr. ^a	Voucher	ndhF	matK	rbcL	РНҮС
Galphimia mexiae C. E. Anderson	NW	Anderson & Anderson 6122 (MICH)	HO246829			HO247054
Galphimia mirandae C. E. Anderson	NW	Cabrera s.n. (ENCB)	HO246830	HO247276		HO247055
Galphimia multicaulis Adr. Juss.	NW	Díaz B. 4760 (MICH)	HQ246831		HO247488	HQ247056
Galphimia platyphylla Chodat	NW	Schinini & Marmori 30112 (MICH)	HQ246832			HQ247057
Galphimia speciosa C. E. Anderson	NW	Anderson 12991 (MICH)	HQ246833	HQ247277	HQ247489	HQ247058
Gaudichaudia albida Schlechtend. & Cham.†	NW	Anderson 13309A (MICH)	AF351034	AF344546	AF344476	AF500542
Gaudichaudia cynanchoides H. B. K.	NW	Koch 82260 (MICH)	HQ246834		HQ247490	HQ247059
Gaudichaudia hexandra (Nied.) Chodat	NW	Haber 3782 (MICH)	HQ246835	HQ247278		HQ247060
Gaudichaudia krusei W. R. Anderson	NW	Jessup 4064 (MICH)	HQ246836	HQ247279		HQ247061
Gaudichaudia mcvaughii W. R. Anderson†	NW	Anderson 12699C (MICH)	AF351032	AF344547	AF344477	AF500540
Gaudichaudia sp.	NW	Anderson 13291B, C (MICH)	HQ246837	HQ247280	HQ247491	HQ247062
Glandonia macrocarpa Griseb.†	NW	Anderson 13776 (MICH)	AF351072	AF344548	AF344478	AF500573
Heladena multiflora (Hook & Arn.) Nied.†	NW	Anderson 13599 (MICH)	AF351044	AF344549	AF344479	AF500549
Henleophytum echinatum (Griseb.) Small†	NW	Curtiss 688 (NY)	HQ246838	HQ247281	HQ247492	
Heteropterys aureosericea Cuatrec.	NW	Burnham 3344 (MICH)	AY499082	HQ247282	HQ247493	AY499056
Heteropterys bahiensis Nied.	NW	Amorim 3664 (CEPEC)	AY499083	HQ247283	HQ247494	AY499057
Heteropterys bicolor Adr. Juss.	NW	Amorim 2710 (MICH)	AY499084	HQ247284	HQ247495	AY499058
Heteropterys brachiata (L.) DC.†	NW	FTG PRC-105 (MICH)	HQ246839	HQ247285	HQ247496	HQ247063
Heteropterys byrsonimifolia Adr. Juss.	NW	Amorim 3663 (CEPEC)	AY499085	HQ247286	HQ247497	AY499059
Heteropterys capixaba Amorim	NW	Amorim 3317 (MICH)	AY499086	HQ247287	HQ247498	AY499060
Heteropterys catingarum Adr. Juss.	NW	Sousa 455 (CEPEC)	AY499087	HQ247288	HQ247499	AY499061
Heteropterys chrysophylla (Lam.) DC.	NW	Amorim 3097 (CEPEC)	AY499088	HQ247289	HQ247500	AY499062
Heteropterys ciliata Nied. [†]	NW	Anderson 13778 (MICH)	AF351049	AF344550	AF344480	AF500554
Heteropterys conformis W. R. Anderson	NW	Amorim 3583 (MICH)	HQ246840	HQ247290	HQ247501	HQ247064
Heteropterys cordifolia Adr. Juss.	NW	Amorim 308/ (MICH)	HQ246841	HQ247291	HQ247502	HQ247065
Heteropterys dumetorum (Griseb.) Nied. ⁷	NW	<i>Tuppo s.n.</i> (CEPEC 96908)	HQ246842	HQ247292	HQ247503	HQ247066
Heteropterys glabra Hook. & Arn.	IN W	Anderson 13390 (MICH)	HQ246843	HQ247293	HQ247504	HQ247067
Heteropterys imperata Amorim	IN W	Amorim 5128 (CEPEC)	HQ240844	HQ247294	HQ247505	HQ247068
Heleroplerys leona (Cav.) Exeli	NW &	Davis 99-3 (A)	AF551050	ПQ247293	ПQ247300	AF300333
Heteropterys leschengultiang Adr. Juss	NW	Amorim 3352 (MICH)	HO246845	HO247296	HO247507	HO247069
Heteropterys leschenauthan Adr. Juss	NW	Amorim 3375 (MICH)	AY499089	HQ247290	HQ247508	AY499063
Heteropterys macrostachya Adr. Juss	NW	Anderson 13788 (MICH)	HO246846	HQ247298	11Q21/200	HO247070
Heteropterys megantera Adr. Juss	NW	Amorim 3667 (CEPEC)	AY499090	HQ247299	HO247509	AY499064
Heteropterys nitida (Lam.) DC	NW	Amorim 3327 (CEPEC)	AY499091	HQ247300	HQ247510	AY499065
Heteropterys nordesting Amorim	NW	Amorim 3229 (CEPEC)	HO246847	HQ247301	HQ247511	HO247071
Heteropterys ovata (Nied.) W. R. Anderson & C. Davis†	NW	Beck 15105 (MICH)	HQ246848	11221/201	HO247512	1122.0000
Heteroptervs pauciflora (Adr. Juss.) Adr. Juss.	NW	Amorim 3623 (CEPEC)	AY499092	HO247302	HO247513	AY499066
Heteropterys pteropetala Adr. Juss. [†]	NW	Anderson 13637 (MICH)	HQ246849	HQ247303	HQ247514	HQ247072
Heteropterys racemosa Adr. Juss. [†]	NW	Rabelo et al. 3685 (MICH)	HQ246850	HQ247304	HO247515	
Heteropterys rhopalifolia Adr. Juss. ⁺	NW	Amorim 2879 (MICH)	AY499093	HQ247305	HQ247516	AY499067
Heteropterys rufula Adr. Juss.†	NW	Amorim 3666 (CEPEC)	HQ246851		HQ247517	HQ247073
Heteropterys sanctorum W. R. Anderson	NW	Amorim 3653 (MICH)	AY499094	HQ247306	HQ247518	AY499068
Heteropterys sericea (Cav.) Adr. Juss.	NW	Amorim 3342 (MICH)	AY499095	HQ247307	HQ247519	AY499069
Heteropterys sincorensis W. R. Anderson	NW	Queiroz 7145 (CEPEC)	AY499096	HQ247308	HQ247520	AY499070
Heteropterys sp.	NW	Amorim 3671 (CEPEC)	HQ246852		HQ247521	HQ247074
Heteropterys steyermarkii W. R. Anderson	NW	Maguire & Politi 27524 (MICH)	AY499097	HQ247309	HQ247522	AY499071
Heteropterys ternstroemiifolia Adr. Juss.	NW	Amorim 3619 (MICH)	AY499098	HQ247310	HQ247523	AY499072
Heteropterys trichanthera Adr. Juss.†	NW	Amorim 3019 (CEPEC)	AY499099	HQ247311	HQ247524	AY499073
Hiptage benghalensis (L.) Kurz†	OW	Averyanov et al. VH 1381 (P)	HQ246853	HQ247312		HQ247075
Hiptage benghalensis (L.) Kurz†	OW	Herbst 9221 (MICH)	AF351057	HQ247313	HQ247525	AF500561
Hiptage candicans Hook. f.	OW	Johnson 2059A (MICH)	HQ246854	HQ247314	HQ247526	HQ247076
Hiptage detergens Craib	OW	Middleton et al. 2095 (MICH)	HQ246855	HQ247315	HQ247527	HQ247077
<i>Hiptage</i> sp.	OW	Middleton 2496 (A)	HQ246856	HQ247316	HQ247528	HQ247079
<i>Hiptage</i> sp.	OW	Middleton 2556 (A)	1 22 24 0 40			HQ247078
Hiraea fagifolia (DC.) Adr. Juss.	NW	Anderson 13593 (MICH)	AF351060	AF344552	AF344483	AF500564
Hiraea smilacina Standl.†	NW	Anderson 13794 (MICH)	AF351061	110045015	AF344484	HQ247080
Hiraea sp.	NW	Anderson 13813 (MICH)	HQ246857	HQ24/31/	HQ247529	HQ247081
Hiraea wiedeana Adr. Juss.	NW	Anderson 13022 (MICH)	HQ246858	HQ24/318	HQ247530	HQ247082
Janusia anisandra (Adr. Juss.) Griseb.	IN W	Anderson 13094 (MICH)	AF351028	AF344555	AF344483	AF500556
Janusia christianeae W. R. Anderson	NW	Silva et al. 1304 (MICH)	HQ246859	HQ247319		HQ247083
Janusia guaranitica (A. StHil.) Adr. Juss.	IN W	Cristobal & Schinini 2498 (MICH)	HQ246860	HQ247321	110247521	HQ247084
janusia guaraniiica (A. StHil.) Adr. JUSS.7	IN W	мирочиская & Uristobal 40/30 (MICH)	TQ240801	пQ24/322	пQ247531	пү247085
Ianusia hexandra (Vell.) W. R. Anderson	NW	Thomas et al 11423a (MICH)	H02/6862	HO2/7320	HO247532	HO247086
Ianusia ianusioides (Adr. Juse.) W. R. Anderson ⁺	NW	McDaniel et al 2600 (UIS)	112270002	HO247323	112271332	11227/080
Janusia janusioides (Adr. Juss.) W. R. Anderson ⁺	NW	Anderson 13759 (MICH)	HO246863	11227/323	HO247533	HO247087
Janusia linearifolia (A. StHil.) Adr. Juss †	NW	Krapovickas & Cristóbal 46451	AY499102	AF344572	AF344505	AF500535
		(MICH)	.,,			2 2 00000

					Plastid		Nuclear
Jonus molfermaner (Vell) W. R. Andersont NW Auderson J 2660 (MICH) AP34202 AP34458 AP34458 AP34458 AP34458 AP34458 AP34458 AP34458 AP34458 AP344587 AP344598 AP344587 AP344598	Taxon	Distr. ^a	Voucher	ndhF	matK	rbcL	PHYC
Januaria madhermanan (Vell, W. E. Anderson* NW Ferracci et al. Löfö (MICH) HQ24666 HQ247733 HQ24708 Abelitan rozen (Ma), Charte. NW Mori 2.2833 (MICH) HQ24866 HQ247733 HQ24709 Abelitan uncan (Na), Charte. NW Mori 2.2833 (MICH) HQ24666 HQ247734 HQ24709 Latiocorpus sp. ¹ NW Anderson* 1578 (MICH) HQ24666 HQ247334 HQ24737 Latiocorpus sp. ¹ NW Anderson* 1583 (MICH) HQ24666 HQ247334 HQ24737 Laphondrin lanomedit W. R. Anderson* NW Anderson* 1584 (MICH) HQ24666 HQ247334 HQ24733 Laphondrin lanomedit W. R. Anderson* NW Anderson* 1574 (MICH) HQ246676 HQ247334 HQ24733 Laphondrin lanomadit Diska NW Anderson* 1574 (MICH) HQ246677 HQ247334 HQ24734 Laphondring lanomadit Diska NW Anderson* 1574 (MICH) HQ24673 HQ247334 HQ24734 Laphondring lanomadit Diska NW Anderson* 1574 (MICH) HQ246677 HQ24733 HQ247344 HQ24734	Janusia mediterranea (Vell.) W. R. Anderson [†]	NW	Anderson 13686 (MICH)	AF351029	AF344556	AF344488	AF500537
Jahdim rippinia	Janusia mediterranea (Vell.) W. R. Anderson ⁺	NW	Ferrucci et al. 1636 (MICH)	HQ246864	HQ247324		HQ247088
Jubeling arear (Mig), Nied-1 NW Mori 2353 (MCH) H235100 AF34498 H9247000 Jubeling uniformi (Ne), Joansee, NW Modernmi 1776 (MCH) HQ 247081 HQ 247091 Jubeling uniformi (Ne), Joansee, NW Anderson 1776 (MCH) HQ 247081 HQ 247081 Jubeling uniformi (Ne), Joansee, NW Anderson 1787 (MCH) HQ 247081 HQ 247081 Lophanthern baromelit (Ne, Anderson 1 NW Anderson 1774 (MCH) HQ 247081 HQ 247039 Lophanthern baromelit (Ne, Anderson 1 NW Anderson 1774 (MCH) HQ 247081 HQ 247039 Lophanthern baromelit (Ne, Anderson 1 NW Anderson 1774 (MCH) HQ 246801 HQ 24733 HQ 24733 Lophanthern baromelit (Ne, Anderson 1 NW Anderson 1774 (MCH) HQ 246871 HQ 24733 HQ 24734 Lophanthern baromelit (Ne, Anderson 1 NW Anderson 1774 (MCH) HQ 246873 HQ 24733 HQ 24754 Lophanthern baromelit (Ne, Anderson 1 NW Anderson 17815 (MCH) HQ 246873 HQ 24733 HQ 24754 Lophanthern baromelit (Ne, Anderson 1 NW And	Jubelina riparia Adr. Juss.	NW	Mori & Souza 17320 (MICH)	HQ246865	HQ247325	HQ247534	HQ247089
Jabelien aufkanne (Nied.) Catatece. NW Burnham (757) (MICH) HQ24666 HQ247232 HQ247033 HQ247031 Lasiocarpuis sp. 1 NW Anderson 13729 (MICH) HQ246067 HQ21732 HQ21733 HQ217031 Lasiocarpuis sp. 1 NW Anderson 13729 (MICH) HQ246067 HQ21732 HQ21733 HQ247031 Laphanthera fammelii W, R. Anderson? NW Solar (LA) (MICH) HQ246069 HQ247733 HQ247031 Laphanthera factorsen Dacke ⁺ NW Anderson 17574 (MICH) HQ246069 HQ247733 HQ247031 Laphanthera factorsen Dacke ⁺ NW Anderson 17574 (MICH) HQ246069 HQ247733 HQ247031 Laphanthera factorsen Dacke ⁺ NW Anderson 17574 (MICH) HQ246069 HQ247733 HQ247734 HQ247081 Laphanthera factorsen Dacke ⁺ NW Imame 17574 (MICH) HQ246761 HQ24733 HQ24734 HQ247081 Laphanthera factorsen Dacke ⁺ NW Imame 17574 (MICH) HQ246761 HQ24733 HQ24734 HQ24734 HQ24734 HQ247341 HQ24734 HQ247334	Jubelina rosea (Miq.) Nied.†	NW	Mori 23855 (MICH)	AF351079	AF344557	AF344489	HQ247090
Jubelmon vilborni W. R. Andersoni T. N.W. Anderson 1525 (MICH) HQ 449010 HQ 47123 HQ 21735	Jubelina uleana (Nied.) Cuatrec.	NW	Burnham 1573 (MICH)	HQ246866	HQ247326	HQ247535	HQ247091
Labacapta Sp. NW Auderson 1583 (NILE) HQ24103 HQ24103 HQ24103 HQ24103 Lephonkerse hommeli W. R. Anderson † NW Solute (DM (CH) HQ24103 HQ24132 HQ24133 HQ24133 HQ24133 HQ24133 HQ24133 HQ24133 HQ24133 HQ24133 HQ24132 HQ24133 HQ24134 HQ24134 HQ24134 HQ24133 HQ24133 HQ24134 HQ241344	Jubelina wilburii W. R. Anderson†	NW	Anderson 13789 (MICH)	AY499100	HQ247327	HQ247536	AY499074
Laphandrein Australity Austra	Lasiocarpus sp. [†]	NW	Anderson 13828 (MICH)	HQ246867	HQ24/328	HQ247537	HQ247092
	Lastocarpus sp. 1	NW	Schatz 1034 (MICH)	HO246868	AF344338 HO247320	HO247538	AF300307
Image: Construction Displane NW Anderson 17371 (MICH) AF51000 AF541590 AF541590 AF541590 AF541590 AF541590 AF541590 AF541590 AF541590 AF541591 AF541590	Lophanthera hammelii W.R. Anderson [†]	NW	Hammel 17748 (MICH)	11Q240000	11Q24752)	11Q247550	HO247093
Lophanthera tonglófia (H. B. K.) Grissb.† NW Zummerman 27 (MICH) HQ246870 HQ247339 HQ24739 HQ24739 Lophanthera tonglófia arbitranda NR Anderson 17574 (MICH) AF31078 HQ247311 HQ247394 HQ247096 Lophanters tonglófia arbitranda NR Madessai IS754 (MICH) AF31078 AF34550 AF344492 AF30078 Madagasikaria andersoni IC. Davis† NW Anderson 13815 (MICH) HQ24671 HQ247333 HQ247334 HQ247342 HQ247333 HQ247342 HQ247394 HQ247342 HQ247333 HQ247342 HQ247342 HQ247354 HQ247333 HQ247342 HQ247354 HQ247352 HQ247354 HQ247354 HQ247354 HQ247354 HQ247356 HQ247341	Lophanthera lactescens Ducket	NW	Anderson 13781 (MICH)	AF351009	AF344559	AF344491	AF500524
Lophonters prindual Ducke NW Luma & Lima & Lima 31/85 (MICH) HQ247331 HQ247331 HQ247340 HQ247037 Lophonterys finandi W. R. Anderson NW Morawer, & Walhnifer 11-27888 MCH) HQ246731 HQ247331 HQ247332 HQ247332 HQ247332 HQ247096 Malpighia andi/ora (Cuntre-) Curtee, † NW Anderson 1357 (MICH) HQ246871 HQ247331 HQ247331 HQ247331 HQ247331 HQ247341 HQ247096 Malpighia andre Spreng, NW Garcria & Finemici 747 (MICH) HQ246871 HQ247331 HQ247341 HQ247331 HQ247341 HQ247331 HQ247341 HQ247341<	Lophanthera longifolia (H. B. K.) Griseb. [†]	NW	Zimmermann 27 (MICH)	HQ246869	HQ247330	HQ247539	HQ247094
Lophopterys, fioribanda W. R. Anderson E. Davis† NW Anderson 12574 (MICH) AP531576 AF534550 AF544540 AF534550 Madagasikaria andersoni C. Davis† NW Madagasikaria andersoni T. Davis† NW Andersoni 13815 (MICH) HQ246711 HQ247333 HQ247534 HQ247524 HQ247534 HQ247544 HQ24754	Lophanthera pendula Ducke	NW	Lima & Lima 3185 (MICH)	HQ246870	HQ247331	HQ247540	HQ247095
Lophopterys inpana W. R. Anderson NW Morawer, & Wallhöfer 11-27888 HQ246871 HQ247332 HQ247332 Malagissi andlerson (2. Davis † OW Davis 20-01 (A) AP436700 HQ247333 HQ24731 HQ247331 HQ24791 Malagissi andlegon (2. Surse, c. + NW Garcia & Finemic / Ar (MCH) HQ246871 HQ247331 HQ247341 HQ247104 Malpighia mericana Adi, Juss. NW Torres, C. 629 (MICH) HQ246871 HQ247341 HQ247104 Malpighia mericana Adi, Juss. NW Rougeton 1527 (MICH) HQ246871 HQ247341 HQ247104 Malpighia baronesang Spreng. NW Word et al. 3982 (MICH) HQ246881 HQ247341 HQ247104 Malpighia baronesang Spreng. NW Boarlett 17568 (MICH) HQ24688	Lophopterys floribunda W. R. Anderson & C. Davis†	NW	Anderson 13754 (MICH)	AF351078	AF344560	AF344492	AF500579
	Lophopterys inpana W. R. Anderson	NW	Morawetz & Wallnöfer 11-27888 (MICH)	HQ246871	HQ247332		HQ247096
Malpiphia abliftora (Cuatrec.) NW Anderson 18815 (MICH) HQ24782 HQ247334 HQ247342 HQ247097 Malpiphia coccigera L.7 NW <i>Cance & Phinestel 74</i> (MICH) HQ247335 HQ247333 HQ247333 HQ247334 HQ247099 Malpiphia merginata DC.7 NW <i>Malerson 1532</i> (MICH) HQ247357 HQ247334 HQ247334 HQ247344 HQ247334 HQ247344 HQ247354 HQ247344 HQ247357 HQ247357 HQ247357 HQ247357 HQ247357 HQ247354 HQ247344 HQ247344 HQ247344 HQ247344 HQ247344 HQ247344 HQ247344 HQ247344 HQ247344 HQ247105 Malpiphia mericana Adi. Juss. NW <i>Calcada et al. 18581</i> (MICH) HQ247344 HQ247344 HQ247105 HQ247344 HQ247344 HQ247104 HQ247344 HQ247104 HQ247104 HQ247104 HQ247104 HQ247104 HQ247104 HQ247104 HQ247344 HQ247344 HQ247104 HQ	Madagasikaria andersonii C. Davis†	OW	Davis 20-01 (A)	AF436790	HQ247333	HQ247541	AF436800
	Malpighia albiflora (Cuatrec.) Cuatrec.†	NW	Anderson 13815 (MICH)	HQ246872	HQ247334	HQ247542	HQ247097
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	Malpighia cnide Spreng.	NW	García & Pimentel 747 (MICH)	HQ246873	110017005	110017510	HQ247098
Mathylia Intergenia Chargenia De.1 NW Adderson 150:21 (MICH) AF2-44-39 AF2-44-39 AF2-44-39 AF2-44-39 AF2-44-34 IQ247136 Malpylia Jatoria Ker Gaul. NW FTG F.2.1 (J4 (FTG)) IIQ246875 IIQ247137 IIQ247138 IIQ247137 IIQ247138 IIQ247137 IIQ247138 IIQ247138 IIQ247137 IIQ247138 IIQ247137 IIQ247138 IIQ247137 IIQ247138 IIQ247137 IIQ247138 IIQ247138 IIQ247138 IIQ247138 IIQ247138 IIQ247138 IIQ247134	Malpighia coccigera L. [†]	NW	UMBG 20626 (MICH)	HQ2468/4	HQ24/335	HQ247543	HQ247099
$ \begin{array}{c} \text{Integration function} & \text{Integration} \\ \text{Molpinghia glatine} L: \\ \text{Molpinghia glatine} L: \\ \text{Molpinghia glatine} L: \\ \text{Molpinghia micrand} Mill: \\ \text{Molpinghia micrand} Calculate \\ \text{Molpinghia micrand} \\ \text{Molpinghia micrand} Calculate \\ \text{Molpinghia micrand} Calculate \\ \text{Molpinghia micrand} Calculate \\ \text{Molpinghia micrand} Calculate \\ \text{Molpinghia micrand} \\ \text{Molpinghia micrand micrand \\ \text{Molpinghia micrand}$	Malpighia emarginala DC.	IN W NW	Proctor 23111 (MICH)	AF551025 HO246875	HO247336	AF 344493	HO247100
	Malpighia glabra L *	NW	$FTG X_{-1}2_{-1}04$ (FTG)	HQ240875	HQ247330	HO247544	HQ247100
$ \begin{array}{c} Malpighia leticiana (W. R. Anderson) W. R. Anderson & NW \\ \hline Torres C. 629 (MICH) \\ Molpighia moneroana Cuatec. \\ Davis \\ Malpighia moneroana Cuatec. \\ NW \\ Wend et al. 3982 (MICH) \\ HQ246878 \\ HQ247341 \\ HQ247341 \\ HQ247341 \\ HQ247103 \\ HQ247104 \\ HQ246881 \\ HQ247341 \\ HQ247341 \\ HQ247104 \\ HQ246881 \\ HQ247341 \\ HQ247105 \\ HQ246881 \\ HQ247341 \\ HQ247141 \\ HQ246881 \\ HQ247341 \\ HQ247141 \\ HQ247105 \\ HQ247105 \\ HQ14688 \\ HQ247341 \\ HQ247141 \\ HQ247105 \\ HQ247105 \\ HQ14734 \\ HQ247141 \\ HQ247105 \\ HQ247105 \\ HQ247141 \\ HQ24734 \\ HQ24735 \\ HQ24734 \\ HQ24735 \\ HQ24734 \\ HQ24735 \\ HQ24734 \\ HQ24735 \\ HQ24735 \\ HQ24735 \\ HQ24734 \\ HQ24735 \\ HQ24735$	Malpighia incana Mill †	NW	Houghton 1152 (MICH)	HQ246877	HO247338	HO247545	HO247102
	Malpighia leticiana (W. R. Anderson) W. R. Anderson & C. Davis†	NW	Torres C. 629 (MICH)	HQ246878	HQ247339	HQ247546	HQ247103
	Malpighia mexicana Adr. Juss.	NW	Calzada et al. 18581 (MICH)	HQ246879	HQ247340		HQ247104
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	Malpighia romeroana Cuatrec.	NW	Wendt et al. 3982 (MICH)	HQ246880	HQ247341		HQ247105
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	Malpighia setosa Spreng.	NW	Bartlett 17568 (MICH)	HQ246881	HQ247342		HQ247106
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	Malpighia souzae Miranda	NW	Zembrano C. 1252 (MICH)	HQ246882	HQ247343		HQ247107
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	Malpighia stevensii W. R. Anderson [†]	NW	Davis 1019 (MICH)	AF351022	HQ247344	HQ247547	AF436798
Materson 13// (MICH)Ar53103Ar34305Ar34493Ar300359Mascagnia drinis W. Anderson C. DavisNWGerlach et al. (10)/01 (MICH)AF351025AF344505AF34499AF340804Mascagnia arenicola C. E. Anderson†NWChase 90165 (MICH)HQ247848HQ247344HQ247549HQ247110Mascagnia australis C. E. Anderson†NWAnderson 13592 (MICH)HQ246885HQ247344HQ247549HQ247111Mascagnia cordifolia (Arb. Juss.) Griseb.†NWAnderson 13592 (MICH)HQ246888HQ247350HQ247550HQ247111Mascagnia dissimilis C. V. Morton & MoldenkeNWNe af1409 (MICH)HQ246888HQ247351HQ247551HQ247111Mascagnia dissimilis C. V. Morton & MoldenkeNWMalnerson 13604 (MICH)HQ246889HQ247351HQ247554HQ247151Mascagnia polybotrya (Adr. Juss.) Nied.†NWAnderson & Anderson 46506 (MICH)HQ246891HQ247354HQ247554HQ247116Mascagnia polybotrya (Adr. Juss.) Nied.†NWAnderson 4604 (MICH)HQ246894HQ247355HQ247156HQ247116Mascagnia tomentosa C. E. AndersonNWChareson 13670 (MICH)HQ246894HQ247354HQ247354HQ247116Mascagnia tomentosa C. E. AndersonNWAnderson 13690 (MICH)HQ246894HQ247354HQ247354HQ247116Mascagnia tomentosa C. E. AndersonNWAnderson 13690 (MICH)HQ246894HQ247354HQ247354HQ247112Mascagnia tomentosa C. E. AndersonNWAnderson 13690 (MICH)HQ246894 <td>Malpighia urens L.</td> <td>NW</td> <td>Bartlett 17518 (MICH)</td> <td>HQ246883</td> <td>HQ247345</td> <td>4 52 4 4 405</td> <td>HQ247108</td>	Malpighia urens L.	NW	Bartlett 17518 (MICH)	HQ246883	HQ247345	4 52 4 4 405	HQ247108
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Malpighiodes bracteosa (Griseb.) W. R. Anderson [†]	IN W	Anderson 13/// (MICH)	AF351055	AF344563	AF344495	AF500559
$ \begin{array}{c} \mbox{mb} (Macagnia drenicola C. E. Anderson† NW Chase 90165 (MICH) H246885 H0247347 H0247548 H024714 $	Mascagnia argnicola C E Anderson [†]	NW	Chase 90160 (MICH)	AF351025	AE3//540	A F3///00	AE436804
Mascagnia australis C. E. Anderson [†] NW Anderson 13592 (MICH) HQ246886 HQ247348 HQ247549 HQ247111 Mascagnia brevifuia Griseb. [†] NW Nee 51409 (MICH) HQ246887 HQ247349 HQ247549 HQ247111 Mascagnia cordificia (Adr. Juss.) Griseb. [†] NW Nee f1409 (MICH) HQ246888 HQ247350 HQ247551 HQ247113 Mascagnia dissimilis C. V. Morton & Moldenke NW Walthright (ICH) HQ246889 HQ247353 HQ247554 HQ247154 Mascagnia litacina (S. Waston) Nied. NW Anderson & Anderson 4656 (MICH) HQ246891 HQ247353 HQ247554 HQ247155 Mascagnia tomentosa C. E. Anderson NW Anderson & Laskowski 4098 (MICH) HQ246891 HQ247355 HQ247157 HQ247156 HQ247111 Mascagnia tomentosa C. E. Anderson NW Webster 23307 (MICH) HQ246891 HQ247355 HQ247557 HQ247157 HQ247157 HQ247157 HQ247157 HQ247157 HQ247157 HQ247157 HQ247157 HQ247157 HQ24711 Mascagnia tomentosa C. E. Anderson 10000 HQ247111 HQ24758 HQ247157<	Mascagnia arenicola C. E. Anderson [†]	NW	Chase 90165 (MICH)	HO246885	HO247347	HO247548	HO247110
Mascagnia brevifolia Griseb.†NWNee \$1409 (MICH)HQ246887HQ247349HQ247550HQ247112Mascagnia contifolia (Adr. Juss.) Griseb.†NWAnderson 13626 (MICH)HQ246888HQ247351HQ247551HQ247111Mascagnia dissimilis C. V. Morton & MoldenkeNWMultinifer 114-291088 (MICH)HQ246888HQ247351HQ247551HQ247151Mascagnia divaricata (H. B. K.) Nied.†NWAnderson & Anderson 4656 (MICH)HQ246891HQ247351HQ247551HQ247151Mascagnia polybotrya (Adr. Juss.) Nied.†NWAnderson & Laskowski 4098 (MICH)HQ246891HQ247355HQ247155HQ2471117Mascagnia tomipiola Nied.NWAnderson & Laskowski 4098 (MICH)HQ246891HQ247355HQ24756HQ247119Mascagnia tomipiola Nied.NWMcebster 32307 (MICH)HQ246894HQ247355HQ247157HQ247157Mascagnia tomentosa C. E. AndersonNWCortes 728 (MICH)HQ246895HQ247357HQ24758HQ247121Mascagnia vaccinifola Nied.†NWAnderson 13797 (MICH)HQ246896HQ247358HQ247122Mevaughia bahiana W. R. Anderson†NWAnderson 13797 (MICH)HQ246896HQ247361HQ247576Mezia includens (Benth.) Cuatrec.†NWAnderson 13784 (MICH)HQ246898HQ247361HQ247561Microsteira ambovombensis Arènes†OWAnderson 13784 (MICH)HQ246899HQ247361HQ247561Microsteira ambovombensis Arènes†OWAnderson 13784 (MICH)HQ246891HQ247361HQ247561 <tr<< td=""><td>Mascagnia australis C. E. Anderson†</td><td>NW</td><td>Anderson 13592 (MICH)</td><td>HQ246886</td><td>HO247348</td><td>HO247549</td><td>HO247111</td></tr<<>	Mascagnia australis C. E. Anderson†	NW	Anderson 13592 (MICH)	HQ246886	HO247348	HO247549	HO247111
$\begin{array}{llllllllllllllllllllllllllllllllllll$	Mascagnia brevifolia Griseb. ⁺	NW	Nee 51409 (MICH)	HQ246887	HQ247349	HQ247550	HQ247112
$\begin{array}{llllllllllllllllllllllllllllllllllll$	Mascagnia cordifolia (Adr. Juss.) Griseb.†	NW	Anderson 13626 (MICH)	HQ246888	HQ247350	HQ247551	HQ247113
$\begin{array}{lll} Mascagnia divaricata (H, B, K,) Nied, ^{+} & NW & Anderson 13604 (MICH) & HQ246891 & HQ247352 & HQ247533 & HQ247155 \\ Mascagnia nolybotrya (Adr. Juss.) Nied, ^{+} & NW & Anderson & Laskowski 408 (MICH) & HQ246891 & HQ247354 & HQ247155 & HQ247117 \\ Mascagnia polybotrya (Adr. Juss.) Nied, ^{+} & NW & Anderson & Laskowski 408 (MICH) & HQ246892 & HQ247354 & HQ247555 & HQ247117 \\ Mascagnia tenuifolia Nied, ^{+} & NW & McYaugh 24004 (MICH) & HQ246892 & HQ247355 & HQ247757 & HQ247119 \\ Mascagnia tenuifolia Nied, ^{+} & NW & Webster 23307 (MICH) & HQ246895 & HQ247357 & HQ247557 & HQ247120 \\ Mascagnia tomentosa C. E. Anderson & NW & Cortes 728 (MICH) & HQ246895 & HQ247357 & HQ247558 & HQ247122 \\ Mascagnia vaccinifolia Nied, ^{+} & NW & Anderson 13797 (MICH) & HQ246896 & HQ247359 & HQ247558 & HQ247122 \\ Mcvaughia bahiana W. R. Anderson ^{+} & NW & Anderson 13797 (MICH) & HQ246894 & HQ247359 & HQ247155 \\ Mezia araujoi Nied, ^{+} & NW & Anderson 13784 (MICH) & AF351070 & AF344568 & AF344501 & AF500571 \\ Mezia araujoi Nied, ^{+} & NW & Anderson 13784 (MICH) & AF351071 & AF344569 & AF344502 & HQ247122 \\ Microsteira ambovombensis Arènes^{+} & OW & Davis 38-01 (A) & AF436791 & HQ247361 & HQ247558 & HQ247126 \\ Microsteira ambovombensis Arènes^{+} & OW & Bavis 38-01 (A) & AF436791 & HQ247361 & HQ247160 \\ Microsteira ambovombensis Arènes^{+} & OW & Randerson 13585 (MICH) & AF351080 & AF344570 & AF344503 & AF500580 \\ Niedenzuella acutifolia (Cav.) W. R. Anderson^{+} & NW & Anderson 13555 (MICH) & AF351080 & AF344570 & AF34570 & HQ247126 \\ Microsteira (Adr. Juss.) W. R. Anderson^{+} & NW & Anderson 13601 (MICH) & HQ246901 & HQ247365 & HQ247161 \\ Miedenzuella sericae (Adr. Juss.) W. R. Anderson^{+} & NW & Anderson 13604 (MICH) & HQ246901 & HQ247365 & HQ247164 \\ Niedenzuella sericae (Adr. Juss.) W. R. Anderson^{+} & NW & Anderson 13604 (MICH) & HQ246901 & HQ247365 & HQ247164 \\ Niedenzuella sericae (Adr. Juss.) W. R. Anderson^{+} & NW & Anderson 13604 (MICH) & HQ246904 & HQ247365 & HQ247126 \\ Niedenzuella sericae (Adr. Juss.$	Mascagnia dissimilis C. V. Morton & Moldenke	NW	Wallnöfer 114-291088 (MICH)	HQ246889	HQ247351	HQ247552	HQ247114
Mascagnia lilacina (S. Watson) Nied.NWAnderson & Anderson & 4656 (MICH)HQ246891HQ247353HQ247154HQ247155HQ247155HQ247155HQ247155HQ247156HQ247117Mascagnia polybotrya (Adr. Juss.) Nied.†NWMcVaugh 24004 (MICH)HQ246892HQ247355HQ247355HQ247156HQ247118Mascagnia tomentosa C. E. AndersonNWWebster 23307 (MICH)HQ246894HQ247357HQ247575HQ247120Mascagnia tomentosa C. E. AndersonNWNeil 7103 (MICH)HQ246896HQ247358HQ247584HQ247121Mascagnia tomentosa C. E. AndersonNWAnderson 13690 (MICH)HQ246896HQ247358HQ247158HQ247121Mascagnia vacciniifolia Nied.†NWAnderson 13690 (MICH)AF351070AF344568AF344501AF500571Mezia includens (Benth.) Cuatrec.†NWAnderson 13772 (MICH)HQ246898HQ247360HQ247559HQ247123Microsteira ambovombensis Arènes†OWDavis 38-01 (A)AF436791HQ247361HQ2471561HQ247126Microsteira ambovombensis Arènes†OWRaderson 13585 (P)HQ246901HQ247363HQ247126Microsteira anbovombensis Arènes†OWRandrianaivo et al. 355 (P)HQ246901HQ247365HQ247163HQ247128Micensteira anbovombensis Arènes†NWAnderson 13585 (MICH)HQ246904HQ247365HQ247163HQ247127Micensteira anbovombensis Arènes†NWAnderson 13530 (MICH)HQ246904HQ247365HQ2471654HQ247125Micenstei	Mascagnia divaricata (H. B. K.) Nied.†	NW	Anderson 13604 (MICH)	HQ246890	HQ247352	HQ247553	HQ247115
Mascagnia polybotrya (Adr. Juss.) Nied.†NWAnderson & Laskowski 4098 (MICH)HQ246892HQ2471354HQ247155HQ247155HQ247155HQ247156HQ247118Mascagnia tomentosa C. E. AndersonNWWebster 23307 (MICH)HQ246894HQ247355HQ247157 <td< td=""><td>Mascagnia lilacina (S. Watson) Nied.</td><td>NW</td><td>Anderson & Anderson 4656 (MICH)</td><td>HQ246891</td><td>HQ247353</td><td>HQ247554</td><td>HQ247116</td></td<>	Mascagnia lilacina (S. Watson) Nied.	NW	Anderson & Anderson 4656 (MICH)	HQ246891	HQ247353	HQ247554	HQ247116
Mascagnia polybolrya (AGT, Juss.) Nied. †NWMcVaugn 24004 (MICH)HQ247035HQ247355HQ247135HQ247153HQ247135HQ247153HQ247154HQ247153HQ247154HQ247153HQ247154HQ247154HQ247154HQ247154HQ247154HQ247154HQ247155HQ247155HQ247155HQ247155HQ247156H	Mascagnia polybotrya (Adr. Juss.) Nied.†	NW	Anderson & Laskowski 4098 (MICH)	HQ246892	HQ247354	HQ247555	HQ247117
Mascagnia temutosia C. E. AndersonNWWebsiter 2307 (MICH)HQ246895HQ247350HQ247150HQ247157Mascagnia tomentosa C. E. AndersonNWNeill 7103 (MICH)HQ246895HQ247357HQ247557HQ247157Mascagnia vaccinitifolia Nied.†NWAnderson 13797 (MICH)HQ246895HQ247358HQ247158HQ247157Mcvaughia bahiana W. R. Anderson†NWAnderson 13690 (MICH)AF351070AF344568AF344501AF50571Mezia araujoi Nied.†NWAnderson 13672 (MICH)AF351051AF344569AF344502HQ247123Mezia includens (Benth.) Cuatrec.†NWAnderson 13784 (MICH)HQ246898HQ247360HQ247561HQ247124Microsteira ambovombensis Arènes†OWRabenantoandro et al. 750 (MO)HQ246909HQ247362HQ247160AF345083Microsteira abovombensis Arènes†OWRabenantoandro et al. 350 (MICH)HQ246901HQ247363HQ247126Microsteira abovombensis Arènes†OWRabenantoandro et al. 350 (MICH)HQ246901HQ247363HQ247126Microsteira abovombensis Arènes†OWRaberantoandro et al. 350 (MICH)HQ246901HQ247363HQ247126Microsteira abovombensis Arènes†OWRaberantoandro et al. 350 (MICH)HQ246901HQ247363HQ247120Microsteira abovombensis Arènes†NWAnderson 13574 (MICH)HQ246901HQ247363HQ247126Microsteira abovombensis Arènes†NWAnderson 13574 (MICH)HQ246903HQ247364HQ247125Micdenzuella acutifoli	Mascagnia polybotrya (Adr. Juss.) Nied.7	IN W	McVaugh 24004 (MICH)	HQ246893	HQ247355	HQ247556	HQ24/118
Mascagna iomeniosa C. E. AndersonNWCories 20 (MICH)HQ247057HQ247157HQ247157HQ247157Mascagnia tomeniosa C. E. AndersonNWAnderson 13797 (MICH)HQ246897HQ247358HQ247358HQ247121Mascagnia vacciniifolia Nied.†NWAnderson 13797 (MICH)HQ246897HQ247359HQ247558HQ247121Mezia araujoi Nied.†NWAnderson 13690 (MICH)AF351070AF344568AF344501AF500571Mezia includens (Benth.) Cuatrec.†NWAnderson 13784 (MICH)HQ247869HQ247360HQ247559HQ247124Microsteira ambovombensis Arènes†OWDavis 38-01 (A)AF436791HQ247361HQ247561HQ247126Microsteira ambovombensis Arènes†OWRabernatoandro et al. 750 (MO)HQ246899HQ247361HQ247561HQ247126Microsteira gp.OWRabernatoandro et al. 365 (P)HQ246900HQ247363HQ247121Mionandra camareoides Griseb.†NWAnderson 13585 (MICH)AF351080AF344570AF344503AF500580Niedenzuella acutifolia (Cav.) W. R. Anderson†NWAmorim 1574 (MICH)HQ246903HQ247364HQ247564HQ247128Niedenzuella sericea (Adr. Juss.) W. R. Anderson†NWAnderson 13604 (MICH)HQ246905HQ247366HQ247564HQ247128Niedenzuella sericea (Griseb.) W. R. Anderson†NWAnderson 13604 (MICH)HQ246904HQ247366HQ247564HQ247128Niedenzuella sericea (Adr. Juss.) W. R. Anderson†NWAnderson 13804 (MICH)HQ246905	Mascagnia temujolia Nicu. Mascagnia tementosa $C \in Anderson$	NW	Cortes 728 (MICH)	HQ240894	HQ247350	HO247557	HQ247119
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International detailing (Car.) W. R. Anderson†INWInternational formational formation formation formational formation formational formation formation formational formation formation formational formation formational formational formational formational formation formational formation formation formational formation formational formation formational formation formational formation formation formational formation formational formational formation formation formation formational formation format	Niedenzuella acutifolia (Cay.) W. R. Anderson ⁺	NW	Amorim 1574 (MICH)	HO246903	HO247365	HO247563	HO247129
Niedenzuella sericea (Adr. Juss.) W. R. Anderson†NWAnderson 13611 (MICH)HQ246904HQ247366HQ247564HQ247130Niedenzuella sericea (Adr. Juss.) W. R. Anderson†NWAnderson 13611 (MICH)HQ246905HQ247367HQ247565HQ247130Niedenzuella stannea (Griseb.) W. R. Anderson†NWAnderson 13804 (MICH)AF351056HQ247368AF344500AF500560Niedenzuella stannea (Griseb.) W. R. Anderson†NWPrance et al. 26362 (MICH)HQ246906HQ247369HQ247567HQ247132Peixotoa cordistipula Adr. Juss.†NWGerlach et al. 26/01 (MICH)HQ246907HQ247370HQ247567HQ247133Peixotoa glabra Adr. Juss.†NWAnderson 9197 (MICH)HQ246907HQ247370HQ247134AF500544Peixotoa paludosa Turcz.NWAnderson 9197 (MICH)HQ247372HQ247371HQ247134Peixotoa reticulata Griseb.NWFerrucci et al. 1678 (MICH)HQ246908HQ247373HQ247136Philgamia glabrifolia ArènesOWSchatz et al. 4098 (P)HQ246910HQ247374HQ247568HQ247137	Niedenzuella acutifolia (Cav.) W. R. Anderson†	NW	Amorim 3350 (MICH)	HQ246902	HQ247364	HO247562	HO247128
Niedenzuella sericea (Adr. Juss.) W. R. Anderson†NWHeringer et al. 1524 (MICH)HQ246905HQ247367HQ247565HQ247131Niedenzuella stannea (Griseb.) W. R. Anderson†NWAnderson 13804 (MICH)AF351056HQ247368AF344500AF500560Niedenzuella stannea (Griseb.) W. R. Anderson†NWPrance et al. 26362 (MICH)HQ246906HQ247369HQ247567HQ247132Peixotoa cordistipula Adr. Juss.†NWGerlach et al. 26/01 (MICH)HQ246907HQ247370HQ247567HQ247133Peixotoa glabra Adr. Juss.†NWAnderson 13636 (MICH)AF344501AF344504AF500564Peixotoa paludosa Turcz.NWAnderson 9197 (MICH)HQ247371HQ247371HQ247134Peixotoa parviflora Adr. Juss.NWHatschbach et al. 6718 (MICH)HQ247373HQ247133Peixotoa reticulata Griseb.NWFerrucci et al. 1678 (MICH)HQ247373HQ247368HQ247136Philgamia glabrifolia ArènesOWSchatz et al. 4098 (P)HQ246910HQ247374HQ247568HQ247137	Niedenzuella sericea (Adr. Juss.) W. R. Anderson ⁺	NW	Anderson 13611 (MICH)	HQ246904	HQ247366	HQ247564	HQ247130
Niedenzuella stannea (Griseb.) W. R. Anderson†NWAnderson 13804 (MICH)AF351056HQ247368AF344500AF500560Niedenzuella stannea (Griseb.) W. R. Anderson†NWPrance et al. 26362 (MICH)HQ246906HQ247369HQ247566HQ247132Peixotoa cordistipula Adr. Juss.†NWGerlach et al. 26/01 (MICH)HQ246907HQ247370HQ247567HQ247133Peixotoa glabra Adr. Juss.†NWAnderson 13636 (MICH)AF344571AF344504AF500564Peixotoa paludosa Turcz.NWAnderson 9197 (MICH)HQ247371HQ247371HQ247134Peixotoa parviflora Adr. Juss.NWHatschbach et al. 6718 (MICH)HQ246908HQ247372HQ247134Peixotoa parviflora Adr. Juss.NWFerrucci et al. 1678 (MICH)HQ246908HQ247373HQ247136Philgamia glabrifolia ArènesOWSchatz et al. 4098 (P)HQ246910HQ247374HQ247568HQ247137	Niedenzuella sericea (Adr. Juss.) W. R. Anderson†	NW	Heringer et al. 1524 (MICH)	HQ246905	HQ247367	HQ247565	HQ247131
Niedenzuella stannea (Griseb.) W. R. Anderson† NW Prance et al. 26362 (MICH) HQ247369 HQ247566 HQ247132 Peixotoa cordistipula Adr. Juss.† NW Gerlach et al. 26/01 (MICH) HQ246907 HQ247370 HQ247567 HQ247133 Peixotoa glabra Adr. Juss.† NW Anderson 13636 (MICH) AF351036 AF344571 AF344504 AF500544 Peixotoa paludosa Turcz. NW Anderson 9197 (MICH) HQ247371 HQ247132 Peixotoa parviflora Adr. Juss. NW Hatschbach et al. 67158 (MICH) HQ247372 HQ247135 Peixotoa parviflora Adr. Juss. NW Ferrucci et al. 1678 (MICH) HQ247373 HQ247135 Peixotoa palubrigolia Arènes OW Schatz et al. 4098 (P) HQ247374 HQ247568 HQ247137	Niedenzuella stannea (Griseb.) W. R. Anderson†	NW	Anderson 13804 (MICH)	AF351056	HQ247368	AF344500	AF500560
Peixotoa cordistipula Adr. Juss.† NW Gerlach et al. 26/01 (MICH) HQ246907 HQ247370 HQ247567 HQ247133 Peixotoa glabra Adr. Juss.† NW Anderson 13636 (MICH) AF351036 AF344571 AF344504 AF500544 Peixotoa paludosa Turcz. NW Anderson 9197 (MICH) HQ247371 HQ247134 Peixotoa parviflora Adr. Juss. NW Hatschbach et al. 67158 (MICH) HQ246908 HQ247372 HQ247135 Peixotoa reticulata Griseb. NW Ferrucci et al. 1678 (MICH) HQ246909 HQ247373 HQ247136 Philgamia glabrifolia Arènes OW Schatz et al. 4098 (P) HQ246910 HQ247374 HQ247137	Niedenzuella stannea (Griseb.) W. R. Anderson†	NW	Prance et al. 26362 (MICH)	HQ246906	HQ247369	HQ247566	HQ247132
Petxotoa glabra Adr. Juss. [†] NW Anderson 13636 (MICH) AF351036 AF344571 AF344504 AF500544 Peixotoa paludosa Turcz. NW Anderson 9197 (MICH) HQ247371 HQ247134 Peixotoa parviflora Adr. Juss. NW Hatschbach et al. 67158 (MICH) HQ246908 HQ247372 HQ247135 Peixotoa reticulata Griseb. NW Ferrucci et al. 1678 (MICH) HQ246909 HQ247373 HQ247136 Philgamia glabrifolia Arènes OW Schatz et al. 4098 (P) HQ246910 HQ247374 HQ247137	Peixotoa cordistipula Adr. Juss.†	NW	Gerlach et al. 26/01 (MICH)	HQ246907	HQ247370	HQ247567	HQ247133
reixotoa patuaosa Turcz.NWAnderson 919/ (MICH)HQ24/3/1HQ24/134Peixotoa parviflora Adr. Juss.NWHatschbach et al. 67158 (MICH)HQ246908HQ247372HQ247135Peixotoa reticulata Griseb.NWFerrucci et al. 1678 (MICH)HQ246909HQ247373HQ247136Philgamia glabrifolia ArènesOWSchatz et al. 4098 (P)HQ246910HQ247374HQ247168	Peixotoa glabra Adr. Juss.†	NW	Anderson 13636 (MICH)	AF351036	AF344571	AF344504	AF500544
Peixotoa parvijora Adi. Juss.INWHaischbach et al. 07156 (MICH)HQ240908HQ247572HQ247155Peixotoa reticulata Griseb.NWFerrucci et al. 1678 (MICH)HQ246909HQ247373HQ247136Philgamia glabrifolia ArènesOWSchatz et al. 4098 (P)HQ246910HQ247374HQ247168	reixoloa paluaosa 1urcz. Pairotoa parviflora Adr. Jucc	IN W NW/	Anuerson 9197 (MICH) Hatschhach at al. 67158 (MICH)	HO246000	HQ24/3/1		HQ24/134
Philgamia glabrifolia Arènes OW Schatz et al. 4098 (P) HQ247576 HQ247568 HQ247137	Peixotoa reticulata Griseh	NW	Ferrucci et al. 1678 (MICH)	HO246908	HO247373		HO247136
	Philgamia glabrifolia Arènes	OW	Schatz et al. 4098 (P)	HQ246910	HQ247374	HQ247568	HQ247137

				Plastid		Nuclear
Taxon	Distr. ^a	Voucher	ndhF	matK	rbcL	PHYC
Philgamia hibbertioides Baill. [†]	OW	Du Puy et al. M635 (P)	HQ246911	HQ247375	HO247569	HQ247138
Psychopterys dipholiphylla (Small) W. R. Anderson & S.	NW	Anderson 13816 (MICH)	AF351063	AF344565	AF344497	AF436795
Corso†		× , ,				
Pterandra arborea Ducke†	NW	Anderson 13766 (MICH)	AF351014	AF344573	AF344506	AF500529
Ptilochaeta bahiensis Turcz. [†]	NW	Anderson 13725 (MICH)	AF351068	AF344574	AF344507	AF500569
Ptilochaeta nudipes Griseb. ⁺	NW	Anderson 13588 (MICH)	AF351067	HQ247376	HQ247570	AF500568
Rhynchophora humbertii Arènes†	OW	Davis 18-01M (A)	HQ246912	HQ247377	HQ247571	HQ247139
Rhynchophora phillipsonii W. R. Anderson [†]	OW	Davis 23b-01 (A)	AF436793	HQ247378	HQ247572	AF436802
Ryssopterys intermedia Hochr.†	OW	Chase M241 (K)	AF351040	-	AF344509	AF500545
Ryssopterys sp.	OW	Motley & Cameron 2215 (NY)	HQ246913	HQ247379	HQ247573	HQ247140
Ryssopterys tiliifolia (Vent.) Adr. Juss. [†]	OW	<i>Chase M240</i> (K)	HQ246914	-	HQ247574	HQ247141
Spachea correae Cuatrec. & Croat†	NW	Hammel 17746 (MO)	AF351074	AF344575	AF344510	AF500575
Spachea elegans (G. Mey.) Adr. Juss. [†]	NW	Jansen-Jacobs et al. 3907 (MICH)	HQ246915	HQ247380	HQ247575	HQ247142
Sphedamnocarpus angolensis (Adr. Juss.) Oliv.	OW	Brummitt & Pope 19597 (MICH)	HQ246916	HQ247381	HQ247576	HQ247143
Sphedamnocarpus galphimiifolius (Adr. Juss.) Szyszyl.	OW	Steiner 2275 (MICH)	AF351041	AF344576	AF344511	HQ247144
Sphedamnocarpus poissonii Arènes	OW	Dorr 3959 (P)	HQ246917	HO247382	HQ247577	HO247145
Sphedamnocarpus pruriens (Adr. Juss.) Szyszyl.†	OW	Goldblatt s.n. (MO)		HO247383	HQ247578	HO247146
Sphedamnocarpus sp.	OW	Davis 03-01 (Å)	AY499104	HO247385	HQ247579	AF500546
Sphedamnocarpus sp.	OW	Davis 39-01 (A)	AY499105	HO247386	HO247580	AY499077
Sphedamnocarpus sp.	OW	Phillipson 2957 (P)	HO246918	HO247384	C	HO247147
Stigmaphyllon aberrans C. E. Anderson	NW	Rojas et al. 2347 (MICH)	HO246919	HO247387	HO247581	HO247148
Stigmaphyllon bogotense Triana & Planch.	NW	Rojas et al. 3058 (MICH)	HO246920	HO247388	HO247582	HO247149
Stigmaphyllon bogotense Triana & Planch.	NW	Wovtkowski 7035 (MICH)	HO246921	??		
Stigmaphyllon calcaratum N E Br	NW	Anderson 13595 (MICH)	HO246922	HO247389	HO247583	HO247150
Stigmaphyllon ciliatum (Lam.) Adr. Juss	NW	Anderson 13739 (MICH)	HO246923	HO247390	HO247584	HO247151
Stigmaphyllon finlayanum Adr. Juss	NW	Chase 90163 (MICH)	HQ246924	HQ247391	HQ247585	HO247152
Stigmaphyllon Jindenianum Adr. Juss	NW	Anderson 13796 (MICH)	112210721	HO247392	HQ247586	HO247153
Stigmaphyllon paralias Adr. Juss †	NW	Anderson 13693 (MICH)	AF351065	AF344577	AF344513	AF500566
Stigmaphyllon puberum (Rich) Adr. Juss †	NW	Anderson 13793 (MICH)	AF351038	HO247393	AF344514	AF436794
Stigmaphyllon sagraeanum Adr. Juss	NW	FTG 375A (FTG)	HO246925	HQ247395	HO247587	HO247154
Stigmaphyllon sagraeanum Adr. Juss	NW	$FTG X_{-1} - 6$ (FTG)	AF351077	HQ247394	AF344512	AF500578
Tetranterys ambigua (Adr. Juss.) Nied *	NW	Kral & Wanderley 75016 (MICH)	HO246926	11Q2+757+	HO247588	HO247155
Tetrapterys arcana C V Morton ⁺	NW	Daniel 9558 (MICH)	HQ240920	HO247396	11Q2+7500	HQ247156
Tetrapterys discolor (G Mey) DC *	NW	Chase 90158 (MICH)	AF351076	AF344578		AF500577
Tetrapterys autotiana Triana & Planch	NW	Anderson 13800 (MICH)	HO246928	HO247397	HO247589	HO247157
Tetrapterys megalantha W R Anderson	NW	Mori et al. 21584 (MICH)	HQ240920	HQ247398	HQ247500	HO247158
Tetrapterys micronhylla (Adr. Juss.) Nied *	NW	Anderson 13644 (MICH)	AF351052	AF344579	AF344515	AE500556
Tetranterys nhlomoides (Spreng) Nied *	NW	Anderson 13685 (MICH)	HO246930	HO247399	HO247591	HO247159
Tetrapterys salicifolia (Adr. Juss.) Nied	NW	Anderson 13616 (MICH)	HQ240930	11Q2+7577	HQ247591	HQ247160
Tetrapterys schiedeana Schlechtend & Cham +	NW	Anderson 13811 (MICH)	HQ240931	HO247400	HQ247592	HQ247161
Tetranterys tinifolia Triana & Planch *	NW	Anderson 13792 (MICH)	HQ240932	HQ247400	HQ247595	HQ247161
Thrvallis latifolia (Bartl.) Kuntze	NW	Lombardi 2213 (MICH)	HQ240933	HQ247401	11Q2+757+	HQ247162
Thryallis longifolia Mart +	NW	Anderson 13657 (MICH)	AF351046	AF344580	AF344516	AF500551
Triasnis glauconhylla Engl	OW	Goldblatt & Manning 10130 (MO)	HO246935	HO247403	HO247595	HO247164
Triaspis Stateophytia Engli Triaspis hypericoides Burch +	OW	Immelman 650 (PRF)	AF351021	AF344581	AF344517	AE500533
Triaspis hypericoides Burch *	OW	P Raal & G Rall 2429BB (M)	HO246936	HO247404	HO247596	HO247165
Triaspis macropteron Oliv	OW	Kuchar 23017 (MO)	HQ240930	HQ247404	HQ247590	HQ247166
Triaspis nelsonii Oliv	OW	Gleiss & Müller 13967 (M)	HQ240937	HQ247405	HO247598	HQ247167
Triaspis niedenzujana Engl	OW	Belsky 584 (MO)	HQ240930	11Q2+7+00	11Q2+7590	HQ247168
Tricomaria usillo Hook & Arn +	NW	Anderson 13581 (MICH)	AF351075	AF344582	AF344518	AE500576
Triontarus iamaicansis I +	NW	Hammel 17816 (MICH)	AF351075	AF344583	AF344519	AF500534
Trionterys paniculata (Mill.) Small [‡]	NW	Valor et al. 862 (MICH)	HO246940	HO247407	HO247500	HO2/7160
Tristellateia africana S. Moore	OW	$D_{avis} 00.25 (\Delta)$	AE3510/13	HQ247407	HQ247577	AE500548
Tristellateia madagascariansis Doir +	OW	McPharson 1/308 (MICH)	AF351043	AE344584	AE344520	AE500547
Tristellateia sp	OW	Bourgaois 16 (MO)	HO246041	HO247400	AI'544520	HO247170
Tristellataia sp.	OW	Davis 10 (110)	AV400104	HO2/7/12		AV400070
Tristellataia sp.	OW	Davis $17-01$ (A) Davis 20.01 (A)	AV400107	HO2/7/12	HO2/7601	AT 4990/0 AV/00070
Tristellateia sp.	OW	Lucile & Rakotozah, 2682 (D)	HO2/60/2	HO2/7/11	11Q24/001	HO2/7172
Tristellataia sp.	OW	Lacino & Rakolozaly 2002 (F)	HO246943	HO2/7/10		HQ24/1/2
тыснанеш эр.	0 11	14477 (MO)	11Q240942	11Q247410		11Q24/1/1
Verrucularia glaucophylla Adr. Juss.†	NW	Amorim 3662 (MICH)	AY499108	HQ247414	HQ247602	AY499080

^a Distribution: NW = New World, OW = Old World.
* Outgroups
† Taxa scored for morphology

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APPENDIX 2. Morphological characters scored for 144 species representing all genera of Malpighiaceae, plus six outgroup taxa.

1. Habit—0: tree or woody shrub; 1: erect suffrutex; 2: trailing suffrutex; 3: vine; 4: herb

2. Stems rooting at nodes—0: no; 1: yes

3. Vegetative hairs-0: unicellular; 1: multicellular

4. Vegetative hairs—0: 2-branched; 1: basifixed; 2: stellate

5. Phyllotaxy-0: opposite or whorled; 1: alternate

6. Stipule position—0: stem between petioles or beside petiole; 1: petiole margin; 2: petiole inner face; 3: not present

7. Stipule connation-0: distinct; 1: connate, same leaf; 2: connate, opposite leaves; 3: connate, opposite and same leaves; 4: not present

8. Stipule persistence-0: long-persistent; 1: soon-deciduous; 2: not present

9. Stipules enclosing buds-0: no; 1: yes

10. Petiole glands-0: absent; 1: present

11. Lamina margin-0: without true teeth; 1: toothed

12. Lamina glands-0: absent; 1: abaxial surface; 2: margin or very slightly within; 3: adaxial surface

13. Inflorescence position-0: terminal or terminal and axillary; 1: axillary

14. Inflorescence ultimate units-0: pseudoracene or thyrse; 1: umbel of (2)4(6) flowers; 2: umbel of >6 flowers; 3: 1(2) flower(s); 4: verticil of 4 flowers; 5: spike; 6: consistently 2 flowers

15. Inflorescence decussate-0: mostly not; 1: all or mostly yes; 2: proximally decussate, distally not; 3: too tight to tell; 4: single flowers

16. Cincinni-0: 1-flowered; 1: 2-several-flowered

17. Bract and bracteole size-0: full-sized leaves; 1: much reduced; 2: apparently absent

18. Bracts and bracteoles scalelike-0: no; 1: yes; 2: absent

19. Bract persistence-0: persistent; 1: deciduous; 2: absent

20. Peduncle—0: 0–1 mm long; 1: >1 mm long

21. Pedicel-0: well developed, > bracteoles; 1: absent or very short, < bracteoles

22. Bracteole glands-0: eglandular; 1: 1 or both glandular; 2: bracteoles absent

23. Bracteole persistence-0: persistent; 1: deciduous; 2: absent

24. Bracteoles enclosing buds-0: no; 1: yes; 2: absent

25. Cleistogamous flowers-0: absent; 1: present

26. Breeding system—0: bisexual; 1: morphologically bisexual, functionally unisexual; 2: morphologically bisexual or male; 3: unisexual, dioecious

27. Sepal number-0: 5; 1: 2-4

28. Sepal aestivation-0: imbricate; 1: valvate

29. Sepal margin, glands-0: absent; 1: present

30. Calyx in anthesis—0: erect or appressed; 1: revolute; 2: reflexed

31. Calyx in fruit-0: hardly enlarged; 1: enlarged, papery 32. Calyx glands—0: none; 1: 10 on 5 sepals; 2: 8 on lateral 4 sepals; 3: 4

(fused) on lateral 4 sepals; 4: up to 5 glands; 5: 6 on lateral 4 sepals 33. Calyx gland attachment-0: completely on free sepals; 1: half or more

on free sepals; 2: mostly below free sepals; 3: not present

34. Calyx glands long-stalked—0: no; 1: yes; 2: not present

35. Petal number-0: 5; 1: 2-4

36. Petals in bud-0: exposed; 1: concealed by sepals

37. Petal color-0: yellow; 1: pink (+white); 2: white; 3: lilac

38. Petals clawed—0: yes; 1: no

39. Petal hairs-0: absent; 1: present, abaxial surface; 2: present, adaxial surface; 3: present, margin

40. Corolla symmetry-0: nearly radial; 1: bilateral, NW type; 2: bilateral, Acridocarpus type

41. Disc—0: absent; 1: present

42. Stamen number (in bisexual or male flowers)—0: >10; 1: 10; 2: 7–9; 3:

6 opposite sepals + posterior petal; 4: 5 opposite sepals; 5: 1-4

43. Stamens sterile (in bisexual or male flowers)-0: none; 1: staminodes opposite 5 sepals; 2: staminodes opposite anterior-lateral sepals; 3: posterior 3 staminodes; 4: filament opposite posterior petal without anther; 5: staminodes opposite posterior-lateral sepals; 6: staminodes opposite petals, posterior-lateral sepals; 7: staminodes opposite 3 anterior sepals; 8: staminodes opposite 5 petals

44. Filament sizes (fertile)-0: subequal; 1: longer opposite sepals; 2: longer opposite some or all petals; 3: longer opposite sepals, posterior-lateral petals; 4: anterior 3 shorter; 5: posterior 3 shorter; 6: shorter opposite posterior-lateral petals; 7: very long opposite anterior sepal; 8: stouter opposite posterior-lateral petals; 9: anterior 1 shorter

45. Filament fusion-0: distinct; 1: connate at base or higher

46. Anther hairs—0: absent; 1: present

47. Anther bristles, apex-0: absent; 1: present

48. Anther wings-0: absent; 1: present

49. Anther dehiscence-0: longitudinal slits; 1: apical or subapical pores

50. Connective extended, fleshy (fertile anthers)-0: no; 1: yes

51. Anther size (fertile)-0: subequal; 1: larger opposite sepals; 2: larger opposite petals; 3: larger opposite posterior-lateral petals; 4: posterior 3 smaller; 5: smaller opposite anterior-lateral petals; 6: anterior 3 smaller; 7: anterior 1 larger;

8: larger opposite 3 anterior sepals; 9: anterior 3 larger 52. Androecium symmetry-0: nearly radial; 1: strongly bilateral

53. Pollen symmetry-0: radial; 1: global

54. Pollen ectoapertures-0: present; 1: absent

55. Pollen endoapertures, number-0: 3; 1: 4-12

56. Pollen diameter (µm)—0: ≤22; 1: >22

57. Receptacle hairs between filaments and gynoecium-0: absent or nearly so; 1: present, abundant

58. Carpel number (chasmogamous flowers)-0: 3; 1: 2; 2: 5

59. Ovules per locule-0: 1 (or none); 1: 2 or more

60. Carpels sterile-0: none; 1: anterior; 2: 1 posterior; 3: anterior + 1 posterior

61. Carpel fusion in ovary-0: distinct or connate at base only; 1: connate whole ventral face or axis

62. Style number (chasmogamous flowers)-0: as many as carpels; 1: 1 on anterior carpel; 2: 2 on posterior carpels; 3: 2 on anterior carpel + 1 posterior carpel

63. Style fusion-0: distinct; 1: connate in styles; 2: connate in stigmas

64. Gynoecium symmetry-0: nearly radial; 1: strongly bilateral

65. Style thickness-0: subulate, slender; 1: uniform, thick; 2: uniform, slender

66. Stigma-0: terminal or nearly so, tiny; 1: internal, short, large; 2: internal, long-decurrent, large; 3: internal, tiny; 4: terminal, large, capitate or truncate; 5: terminal, large, reniform; 6: terminal, large, elongated

67. Style dorsal extension-0: none; 1: angle to hook; 2: foliole

68. Styles distally-0: entire; 1: bifid

69. Fruit texture-0: dry; 1: fleshy

70. Fruit dehiscence-0: schizocarpic, not releasing seed; 1: indehiscent; 2: loculicidally dehiscent, releasing seeds; 3: septicidally dehiscent, releasing

seeds

71. Fruit wall-0: smooth; 1: setiferous; 2: dorsal wing dominant; 3: lateral wing(s) dominant; 4: winglets, ruffles; 5: aculeate; 6: dorsal or dorsal+lateral crest(s)

72. Carpophore-0: absent; 1: present

73. Eumascagnioid disk-0: absent; 1: present

74. Aril-0: absent; 1: present

75. Endosperm-0: absent; 1: present