1	Phylogeny of the species-rich Pilea Lindl. (Urticaceae) supports its revised delimitation and
2	infrageneric classification, including the resurrection of Achudemia Blume
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21 ABSTRACT

Pilea Lindl., with 933 published names is the largest genus within the Urticaceae. Pilea was 22 23 last monographed in 1869 and whilst the monophyly of the genus has been proposed by previous authors, this has been based on incomplete taxon sampling and the failure to resolve 24 the position of key taxa. We aimed to generate a robust phylogeny for Pilea and allied genera 25 26 that could provide a framework for testing the monophyly of *Pilea*, revising its delimitation and for answering broader scientific questions about this species-rich genus. To do so, we 27 28 sought to sample taxa representative of previous infrageneric classifications and with 29 anomalous inflorescences or flower configurations and to use the resulting phylogeny to 30 evaluate the delimitation of *Pilea* and to establish an infrageneric classification. In addition, 31 we included a representative of the Polynesian genus Haroldiella which, morphologically, is 32 very similar to Pilea. Using Sanger sequence data from two plastid and one nuclear regions 33 we constructed a phylogeny using Bayesian Inference, Maximum Likelihood and Maximum Parsiomony approaches. We used our phylogeny to evaluate the informativness of 19 34 35 morphological traits and applied both to delimit a monophyletic genus and infrageneric 36 sections. Our results recovered *Pilea* as paraphyletic with respect to *Lecanthus*, a 37 consequence of the recovery of a monophyletic clade comprising sections Achudemia and Smithiella, neither of which had been adequately sampled in previous studies. We also 38 39 recovered *Pilea* as polyphyletic with respect to *Haroldiella*. We identified isomery between 40 male and female flowers, flower part number and male sepal arrangement as being 41 phylogenetically informative traits that can be used to delimit two genera, Achudemia, 42 including section Smithiella, recovered as sister to Lecanthus, and Pilea, including 43 Haroldiella, recovered as sister to both. On the basis of our evaluation of both morphological 44 traits and phylogenetic relationships we propose a new infrageneric classification for the 45 genus comprising seven sections, five of which we describe for the first time, § Trimeris Y.G.Wei & A.K.Monro, § Lecanthoides C.J.Chen, § Angulata L.F.Fu & Y.G.Wei, § 46 47 Tetrameris C.J.Chen, § Verrucosa L.F.Fu & Y.G.Wei, § Plataniflora L.F.Fu & Y.G.Wei and § 48 Leiocarpa L.F.Fu & Y.G.Wei. We also identify a trend of decreasing merism and fruit size, 49 and increasing species-richness as *Pilea* diverges. In addition, we recover strong geographical 50 structure within our phylogeny, sufficient to propose that *Pilea* originated in the IndoMalaya

51 biogeographic domain.

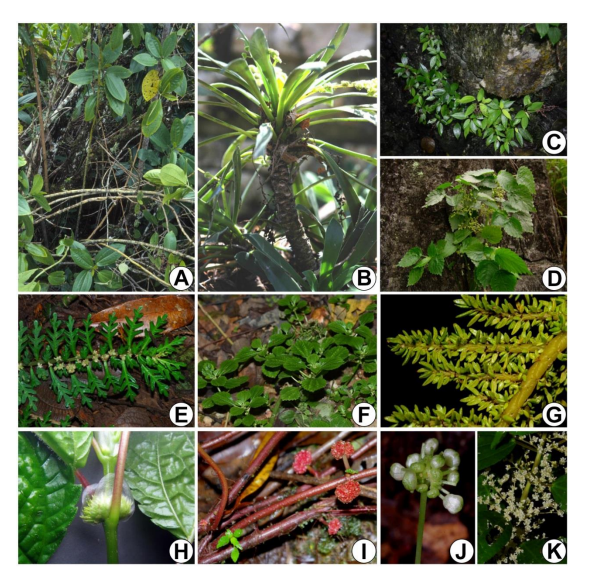
Key words: Lecanthus, Haroldiella, Aboriella, phylogenetics, taxonomy, merism

56 **1. Introduction**

Pilea Lindl., with 933 published names (IPNI, 2020), 604 accepted names (WCVP, 2020) 57 58 and likely 715 species worldwide (Monro, 2004) is the largest genus within the Urticaceae and has a pantropical and subtropical distribution. *Pilea* is characterized by succulent herbs, 59 shrubs and epiphytes whose flowers are wind pollinated, opening explosively, and seed which 60 61 is mechanically dispersed over short distances through the reflexing of the staminodes. It is most species-rich in forested rocky habitats, especially on limestone or ultramafic rocks at 62 63 elevations between of 500 and 2,000 masl, in the Greater Antilles, Central America and the Andes. Members of the genus may be distinguished from other genera in the family by the 64 65 combination of opposite (rarely alternate) leaves, intrapetiolar stipules, an absence of stinging 66 hairs, male inflorescences not fused to form a receptacle-like structure, and free female sepals (Fig. 1 & Fig. 2). As is the case for many species-rich genera, Pilea has not been 67 68 monographed since the 19thC (Weddell, 1869) at which time the genus comprised ca 150 spp... 69 Instead, its taxonomy has been revised piecemeal through flora treatments (Monro, 2006).

Pilea belongs to the Elatostemaea tribe (Gaudichaud, 1830) which, including Sarcopilea 70 71 Urb., has been recovered as monophyletic and sister to Lecanthus Wedd. (Monro, 2006; 72 Jestrow et al., 2012; Wu et al., 2013; Tseng et al. 2019). The Elatostemeae comprises mainly succulent, shade-loving, wind-pollinated species which, as is the case for each tribe in the 73 74 family, show a great variation in female inflorescence arrangement, ranging from open 75 panicles to spikes and fused receptacle-like structures. Despite several molecular studies, 76 doubts remain over the status of Achudemia Blume. Different accessions of A. japonica, 77 having been recovered within, or sister to, Pilea (Monro, 2006) and currently it is included 78 within Pilea (Friis, 1989; Chen and Monro, 2003). In addition, neither the Polynesian 79 endemic, Haroldiella J.Florence, whose morphological circumscription is congruent with 80 *Pilea*, or section *Smithiella*, characterised by strongly asymmetrical spicate inflorescences, 81 were sampled in previous studies, suggesting that the monophyly of the genus remains 82 untested.

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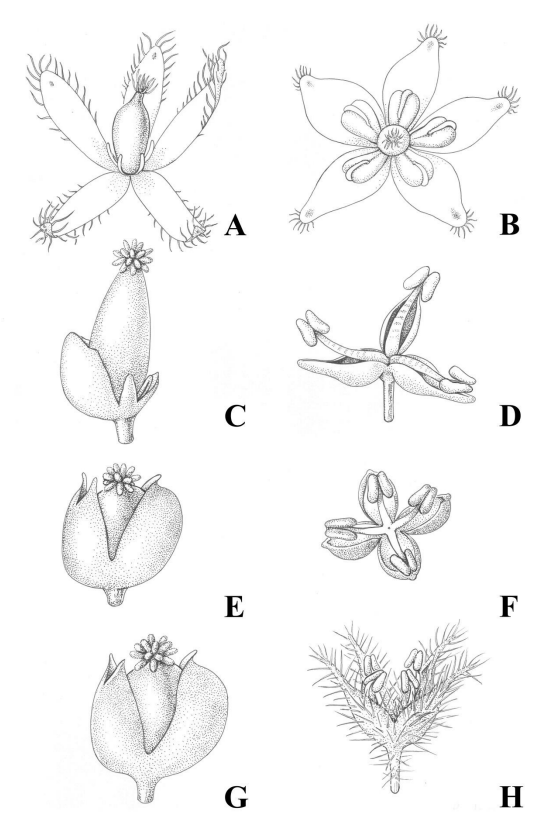


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87 Fig. 1. Morphological diversity of Pilea and Achudemia. A, P. longicaulis (shrub); B, P. 88 fairchildiana (shrub, with alternate, spirally arranged leaves); C, A. boniana (herb, epipetric); D, P. paniculigera (herb, epipetric); E, P. matama (unequal opposite leaves, epiphytic with capitate 89 90 female inflorescences); F, P. peploides (herb, clumped); G. P. sp aff. microphylla; H, P. rivularis 91 (female inflorescence enclosed by stipules); I, P. aff. pittieri (herb, male capitate inflorescence 92 arising from stolons); J, P. angustifolia (herb, male capitate inflorescence); P. notata (male cyme 93 inflorescence). A-B, E, G-J were photographed by Alexandre K. Monro; D was photographed by 94 Yi-Gang Wei; C-D, F, K were photographed by Long-Fei Fu.

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Fig. 2. Illustration of male and female flowers of *Pilea* and *Achudemia*. A, *A. javanica* (female
flower); B, *A. javanica* (male flower); C, *P. tripartite* (female flower); D, *P. tripartite* (male
flower); E, *P. plataniflora* (female flower); F, *P. plataniflora* (male flower); G, *P. microphylla*(female flower); H, *P. microphylla* (male flower). Illustration by Margaret Tebbs.

104 Achudemia, currently treated as a section of *Pilea*, comprises four species of herb that 105 grow in deep forest shade, stream sides, gorges and caves in Indomalaya. Achudemia was 106 established by Blume (1856: 57) to account for a *Pilea*-like collection from Java (Indonesia) 107 which had bisexual (hermaphrodite) five-parted flowers. It appears that Blume described the 108 flowers as bisexual in error as neither the holotype, type illustration, or any other collections 109 have been observed to have bisexual flowers.

Pilea section *Smithiella*, comprises a single species of herb from Indomalaya, also growing in deep shade (Chen, 1995; Chen and Monro, 2003). *Smithiella* was generated by Dunn (1920) to account for *Pilea*-like material from the Eastern Himalayas characterised by strongly asymmetrical spicate inflorescences of five-parted flowers. Dunn had been unaware of an earlier homonym with priority, *Smithiella* H. Perag. & Perag. and in 1981, Bennet (1981) created a replacement name, *Aboriella*.

116 Haroldiella comprises two species from Austral Polynesia growing on rocky outcrops or in rain forest. Haroldiella was described by Florence (1997) based on plants with alternate, 117 spirally arranged pinnately nerved leaves. As with all Pilea from French Polynesia (Florence, 118 119 1997), they also share the trait of two-sepalate female flowers, a condition very rare 120 elsewhere in the genus. With the recovery of Sarcopilea domingensis Urb., a taxon with spirally arranged, pinnately nerved alternate leaves, within a monophyletic Pilea (Monro, 121 122 2006, Jestrow et al., 2012), the characters used to delimit *Haroldiella* no longer support its separation as a distinct genus. 123

Both *Achudemia* and *Aboriella* differ from other *Pilea* species in having male and female flowers with five sepals (Fig. 2). Where free, the same number of perianth parts in male and female flowers is uncommon within the Urticaceae and within *Pilea*, it is a condition restricted to a basal, relatively species-poor clade (Monro, 2006) comprising Afrotropical, Indomalayan and neotropical species.

Previous phylogenetic studies have consistently recovered *Lecanthus* as the genus most closely related to *Pilea* (Monro, 2006; Wu et al., 2013, 2018), from which it be distinguished by its male inflorescences being fused to form a concave receptacle-like structure reminiscent of *Elatostema* J.R.Forst. & G.Forst. It also has an equal number of male and female perianth parts, either four or five (Chen and Monro, 2003). Given the above, and with the limited sampling of *Pilea* species with anomalous inflorescence arrangements (spicate, receptacle-like) or flower-part number (five), together with the ambiguous position of *Achudemia*, generic delimitation is potentially unstable. Furthermore, the two main infrageneric classifications of *Pilea* (Weddell, 1856; Chen, 1982) have been demonstrated to be largely para- or polyphyletic.

For the above reasons, we aimed to generate a robust phylogeny for *Pilea* and allied genera that could provide a framework for revising the delimitation of the genus and the answering of broader scientific questions about this species-rich and poorly studied genus. To do so we sought to increase taxon sampling for the genus, encompassing all previous sections, and the full range of morphological variation and geographical occurrence, using an evaluation of the informativeness of morphological traits support the establishment of an infrageneric classification.

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147 **2. Materials and methods**

148 2.1. Taxon sampling

149 We included 137 accessions representing 125 taxa (Table 1). These included 18 outgroup taxa 150 from the Cannabaceae, Moraceae and representatives of all Urticaceae tribes except for Cecropieae (see Table 1) and 107 ingroup taxa (Lecanthus + Pilea). Within the Elatostemeae, 151 152 all genera were sampled except for the monotypic *Metapilea* which is likely extinct (Wu et al., 2018), known only from the type and could not be sampled. This encompassed the following 153 154 taxa, Elatostema (3 spp), Elatostematoides (1 sp), Lecanthus (2 spp), Pilea (105 spp), Procris 155 (1 sp) and *Polychroa* (1 sp). We focussed on the ITS nuclear region and the *rbcL* and *trnL-F* 156 plastid regions. We combined sequences generated by previous studies (Monro, 2006; Jestrow 157 et al., 2012; Wu et al., 2013, 2018; Kim et al., 2015) 141 sequences of which were generated 158 by ourselves, excluding those accessions where we felt that the identifications were 159 ambiguous, or where sequence data for only a single region could be obtained. An exception 160 was made for the single sequence of *Haroldiella* that we were able to obtain. Our sampling of 161 Pilea included representatives of all infrageneric sections proposed by Chen (1982, Chen and Monro, 2003). We did not structure our sampling to include representatives of Weddell's 162 163 sections as these were all demonstrated to be para- and polyphyletic by Monro (2006).

164 Sequence data were obtained for the four species of P. sect. Achudemia, the monotypic P. sect. 165 Smithiella, ten from P. sect. Tetrameris (approximately 2/3 of the species), 85 from P. sect. Urticella (approximately 1/4 of the species), seven from P. sect. Pilea (approximately 2/3 of 166 the species), two from P. sect. Dimeris (1/2 of the species) and one from P. sect. Lecanthoides 167 (1/2 of the species). Four species of Moraceae (Fatoua villosa Nakai, Morus alba L., Sorocea 168 169 affinis Hemsl., Trophis racemosa (L.) Urb.) and two species of Cannabaceae (Cannabis sabiva L., Humulus lupulus L.) were chosen as outgroups based on the previous analyses 170 171 (Zhang et al., 2011; Kim et al., 2015). Species names, the accession numbers of sequences 172 downloaded from GenBank, and newly generated sequences used in this study are listed in 173 Supplementary Text 1.

174

175 Table 1. Statistics for the molecular datasets used in this study.

	Number of sequences (ingroup/outgroup)	Aligned length (bp)	Length variation (bp)	Variable characters (bp)	Parsimony-infor mative characters (bp)	Model selected (AIC)
ITS	119/18	716	472-588	490	398	GTR+I+G
trnL-trnF	117/18	1059	402-1059	532	356	GTR+I+G
rbcL	61/18	637	629-637	137	82	GTR+I+G
Combined plastid	119/18	1696	1038-1696	669	438	GTR+I+G
Combined all	119/18	2412	1572-2231	1159	836	GTR+I+G

176

177 2.2. DNA isolation, PCR amplification and sequencing

178 Genomic DNA was extracted from fresh or dried materials using a modified CTAB protocol 179 (Chen et al., 2014). The nrITS region was amplified using primers ITS 4 and ITS 5 (White et al., 1990) and rbcL using primers 1F and 724R (Fay et al., 1997). The trnL-F spacer was 180 181 amplified using primers e and f (Taberlet et al., 1991) for most accessions while for few 182 problematic cases, we employed primers c and d to separate *trnL-F* into two overlap regions then concatenated sequences (Taberlet et al., 1991). The PCR amplification were set at 94 $\,^{\circ}$ C 183 for 5 min, 30 cycles of 94 °C for 30 s, 55 °C for 30 s, and 72 °C for 45 s and a final extension 184 185 at 72 °C for 10 min. The PCR products were checked on 1% agarose gels before being 186 purified using a QiaQuick gel extraction kit (Qiagen, Inc., Valencia, California, USA) and directly sequenced in both directions using the amplification primers on an ABI 3730 187 188 automated sequencer (Applied Biosystems, Forster City, California, U.S.A.).

189 2.3. Phylogenetic analyses

190 Raw sequences were edited and assembled using the software Lasergene Navigator (DNAStar, 191 Madison, Wisconsin, USA) with subsequent manual adjustments. The output DNA sequences were then aligned using MAFFT version 7.0 (Katoh and Standley, 2013) with default settings, 192 followed by manual adjustment. The three datasets (nrITS, rbcL, and trnL-F spacer) were 193 194 aligned independently. Alignments were adjusted manually in MEGA 5.1 (Tamura et al., 2011). Phylogenies were reconstructed based on the nrITS dataset, the combined plastid 195 196 datasets (*rbcL*, and *trnL-F* spacer), and all three datasets combined (nrITS, *rbcL*, and *trnL-F* 197 spacer), respectively. All of these reconstructions were analysed using Bayesian inference (BI), maximum likelihood (ML), and maximum parsimony (MP) methods. A visual 198 comparison of the two best tree topologies generated by ML analyses of cpDNA and nrITS 199 200 datasets were performed to compare topological incongruence. A conflict in tree topologies of 201 each tree was considered significant when incongruent topologies both received bootstrap 202 values $\ge 80\%$ (Monro, 2006; Tseng et al., 2019).

Best-fit DNA substitution models were selected using the Akaike Information Criterion (AIC) in Modeltest v 2.7 (Posada and Crandall, 1998) for each data partition. The substitution model of the sequences was set to GTR+G+I for each single dataset based on Modeltest. BI analyses were based on a Markov chain algorithm implemented in MRBAYES 3.2.6 (Huelsenbeck and Ronquist, 2001).

ML analyses with 1000 bootstrap resampling (MLBS) were conducted using the online version of RAxML-HPC2 v8.2.9 (Stamatakis et al., 2008) available at the CIPRES Science Gateway version 3.3 (http://www.phylo.org/index.php/portal/) (Miller et al., 2010) with the gamma model of rate heterogeneity.

MP analyses were performed using PAUP* v4.0b10 (Swofford, 2002), in which all characters were unordered and equally weighted, and gaps were treated as missing data. Heuristic searches of MP were conducted with 100 random addition replicates with tree tree-bisection-reconnection (TBR) branch swapping and MulTrees in effect. Branch supports were assessed using 1000 bootstrap replicates (maximum parsimony bootstrap; MPBS) with the sample settings the same as those for heuristic searches.

218 2.4. Estimates of support

In Bayesian analyses, posterior probabilities (PP) below 0.9 were considered as providing no support, between 0.9 and 0.94 as providing weak support, between 0.95 and 0.99 as providing moderate support, and 1.0 as providing strong support (Tseng et al., 2019).

In bootstrap analyses of the ML (BSML) and MP (BSMP) analyses, values below 70% were considered as providing no support, between 70-79% as providing weak support, between 80-89% as providing moderate support, and 90-100% as providing strong support (Tseng et al., 2019).

226 2.5. Morphological trait evolution

Based on existing phylogenetic studies we performed ancestral state reconstructions (ASR) in order to evaluate the phylogenetic informativness of selected morphological traits and so apply these to the delimitation of *Pilea* and allied genera, and the establishment of an infrageneric classification of *Pilea*. Our aim was to establish a classification that was both phylogenetically congruent and morphologically diagnosable.

Nineteen morphological traits were coded for analysis (see Supplementary Text 2). Traits were selected on the basis that they had been used in previous classifications and revisions of *Pilea* (Weddell, 1856; Chen, 1982; Monro, 2006, 2015). Traits were scored based on the examination of herbarium specimens and description in the literatures (Chen, 1982; Friis, 1989; Monro, 1999, 2015; Chen and Monro, 2003; Monro et al., 2012; Fu et al., 2017a; Yang et al., 2018).

238 Likely transitions between trait states through evolution were reconstructed using ML 239 methods in Mesquite v.3.51 (Maddison and Maddison, 2015). We sampled the last 1000 trees 240 from the post burn-in set of the Bayesian analysis using combined dataset and an equal rate 241 model (Mk1) was selected for all traits. To account for phylogenetic uncertainty, we used 242 'Trace character over trees'. All reconstructions were integrated over the 1000 trees from the 243 post burn-in set and summarized on one of these trees that most matched our hypothesized 244 topology. The results were summarized as a percentage of changes of trait states using the 245 option of 'Average frequencies across trees'.

246 2.6. Delimitation of infrageneric groupings

Given the number of species in *Pilea*, an infrageneric classification can be a practical way toease identification, as well as providing a framework for answering broader evolutionary

questions. With these aims our classification needed to reflect both phylogenetic relationships and be morphologically diagnosable. We decided to base our classification of *Pilea* on sections rather than subgenera as the distinction between the two is unclear (Brizicky, 1969) and in this way we maintain the terminology adopted by Weddell (1856, denoted by the symbol '§') and Chen (1982). We have also aimed to establish sections in accordance with the International Code of Nomenclature for algae, fungi and plants (Turland et al., 2018).

255

256 **3. Results**

257 3.1. Phylogenetic reconstruction

Characteristics and statistics of the datasets used in this study are summarized in Table 1. The 258 comparison of trees for cpDNA (trnL-F, rbcL) and nrITS revealed an incongruence between 259 260 the outgroup taxa (Figs. S1 & S2). Because this incongruence did not affect the topology of 261 the ingroup taxa and the phylogeny of the combined dataset showed better resolved trees with higher support values, than individual trees, we used the combined dataset for subsequent 262 analyses, including that of transitions between morphological trait states. The ingroup taxa 263 264 were recovered as monophyletic (Fig. 3) with strong support

265 (PP1.0/BSML100%/BSMP100%).

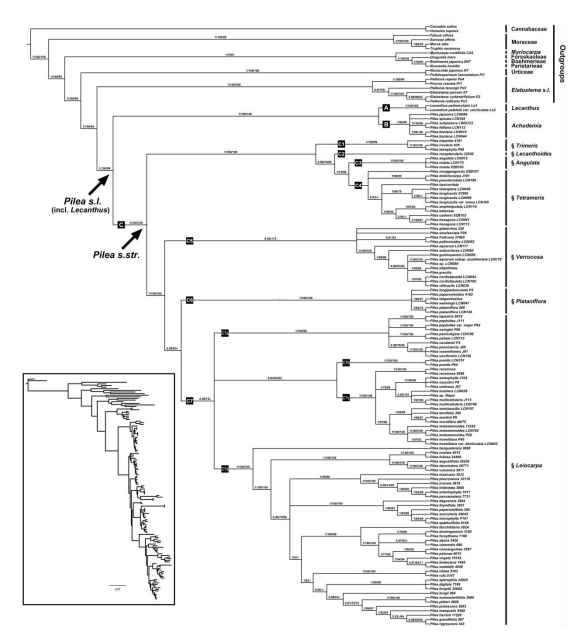
266 3.2. Phylogenetic relationships of Pilea

267 *Pilea*, including Achudemia and Haroldiella was recovered as paraphyletic with respect to Lecanthus (Fig. 3 & Fig. S3). Two strongly supported clades attributable to Pilea sensu latu 268 269 were recovered. The first (Fig. 3, Clade B, labelled as Achudemia), was recovered sister to 270 Lecanthus (Clade A) and included all accessions from P. sect. Achudemia and P. sect. Smithiella (P. subpubera, P. boniana, P. hilliana, P. japonica, P. spicata) with strong support 271 272 (PP1.0/BSML100%/BSMP100%). The second clade (Fig. 3, Clade C) comprised all other 273 accessions of *Pilea* with strong support (PP1.0/BSML100%/BSMP100%) and, in the analysis 274 of the ITS sequence data (Fig. S3), Haroldiella, with strong support (1/100/100). Within 275 Clade C, seven subclades were recovered with strong to weak support (C1 (1/100/99), C2 276 (0.96/74/92), C3 (1/100/100), C4 (-/-/), C5 (0.94/-/74), C6 (1/100/100) and C7 (0.99/72/-)) 277 (Fig. 3). Haroldiella was recovered within clade C7d (Fig. S3, 1/100/98). Clade C4 and C5 were no and weakly supported by all methods used to analyse the data, albeit they comprise 278

279 groups united by the morphological trait states of four-parted female flower and ornamented

achenes, respectively. Clade C7 was recovered with strong to weak support but comprised

- 281 four strongly supported subclades (C7a (1/100/99), C7b (1/100/100), C7c (1/100/100), C7d
- 282 (1/100/100)).
- 283



284

Fig. 3. Phylogenetic tree of *Pilea* generated from Bayesian Inference (BI) of combined dataset
(nrITS, *trnL-F* spacer and *rbcL*). Numbers on the branches indicate the posterior probability (≥0.8)
of BI and bootstrap values (≥60%) of the maximum likelihood (ML) and the maximum parsimony
(MP) analyses.

290 *3.3. Geographical structure*

Clade A (Figs. 4-5 & Fig. S23) comprises taxa with an Asia distribution, although *Lecanthus*also includes species (not sampled) from Africa. Clade B (Figs. 4-5 & Fig. S23) comprises
taxa from East and Southeast Asia.

Clade C (Figs. 4-5 & Fig. S23) comprises species with a pantropical distribution except 294 295 for Australia and New Zealand. Within clade C, subclade C1 comprises taxa from Africa, Asia and Latin America; subclades C2, C3, C4, C5 and C6 comprise taxa from East Asia; and 296 297 subclade C7 taxa from the pantropics. Haroldiella, restricted to Polynesia was recovered 298 within a polytomy within clade C7d meaning that it is more closely related to neotropical species, than to African or Asian ones. Within clade C7, three subclades show a strong 299 geographical association. Subclade C7d, which harbours the greatest number of species, is 300 301 strongly associated with the neotropics, the Greater Antilles, Andes and Central America in 302 particular. Subclades C7b and C7c comprise predominantly palearctic taxa.

303 *3.4. Morphological trait evolution*

304 The 19 morphological traits were mapped onto the Bayesian Inference (BI) tree based on the 305 Maximum Likelihood analyses (Fig. 4 & Figs. S4-S22). This recovered five-sepalate male and female flowers, and imbricate male flower sepals as synapomorphies for Clade A + B 306 (Achudemia), with a reversal in L. petelotii var. corniculata which has four-sepalate male and 307 308 female flowers. Achene bearing a crescent-shaped protuberance and unbranched male inflorescences were recovered as autoapomorphies for Clade A (Lecanthus). Five-sepalate 309 310 female and male flowers were recovered as a plesiomorphy, and branched male inflorescences 311 as a synapomorphy, for Clade B (*Pilea* section Achudemia + P. section Smithiella). The 312 presence or absence of a crescent-shaped protuberance on the achene and branching, or not, 313 of the male inflorescence enable clades A and B to be readily distinguished from each other.

Four plesiomorphies were recovered for Clade C (all remaining *Pilea* species), stipules \leq 10 mm, male flowers 3-sepalate, achene > 0.8 mm, achene ornamented. These were manifested in the basal subclades, transforming to other states through the tree. All other traits were recovered as homoplastic for the clade.

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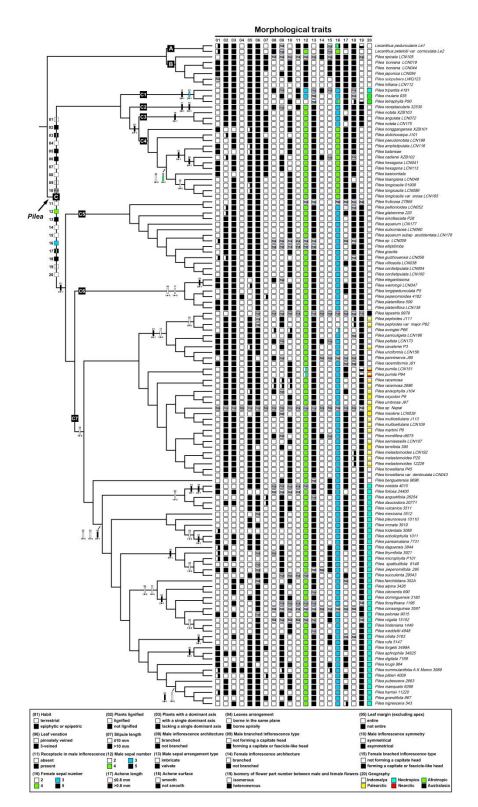


Fig. 4. Reconstruction of the evolution of twenty morphological traits in *Pilea* based on our Bayesian Inference analysis of the combined dataset. The trait states at the *Pilea* node indicate the ancestral states of the genus. Transitions are indicated as filled boxes on the branches. Traits are shown above boxes and state transitions below. Descriptions of traits and their states are provided in the legend. The three clades (A–C) and seven subclades (C1-C7) correspond to those in Fig. 2. Reconstructions for each trait can be seen in Supplementary Figures S4 to S23.

Within subclades of Clade C, however, several plesiomorphic and synapomorphic trait 327 states were recovered. For clade C1, we recovered male flowers 3-sepalate (with a reduction 328 329 to two-sepalate for *Pilea tetraphylla*) as a synapomorphy and male sepals valvate as a plesiomorphy. For C2, we recovered branches of the male inflorescence fused to form a 330 receptacle-like structure as a synapomorphy. For C3, we recovered stipules > 10 mm as a 331 332 synapomorphy. For C5 we recovered achene surface ornamented as a plesiomorphy. For C6 we recovered achene surface ornamented and leaf margin entire as plesiomorphies. For C7 we 333 334 recovered achene surface smooth as a plesiomorphy. For C4 we recovered no synapomorphies or plesiomorphies. It could, however, be morphologically diagnosed based on unique 335 336 combinations of morphological traits (male and female flowers 4-sepalate), as could several 337 other monophyletic groupings within clade C.

338

339 **4. Discussion**

340 *4.1. Resurrection and expansion of Achudemia*

Previous studies (Monro, 2006; Jestrow et al., 2012; Wu et al., 2013, 2015, 2018) did not 341 342 attempt to establish formal infrageneric classifications because of limited morphological trait 343 and taxon sampling, especially of the more basal taxa. Monro (2006) did, however, suggest a preliminary and informal classification of *Pilea* into six 'units' based on geography and 344 345 isomery of flower part number between male and female flowers, cystolith distribution on the leaves and the presence of multicellular hairs. In this study we sought to propose a formal 346 347 classification based increased taxon, morphological and geographical sampling. Our approach 348 resolved the positions of all members of section Achudemia, including Pilea subpubera, 349 synonym of Achudemia javanica, genus type for Achudemia, and Pilea (Achudemia) japonica, 350 ambiguously recovered within and outside of *Pilea* by Monro (2006). In doing so recovered a 351 paraphyletic *Pilea*. The paraphyly can be resolved through the exclusion of 5-sepalate male 352 flowered taxa, formally assigned to Chen's (Chen, 1982; Chen and Monro, 2003) sections 353 Achudemia and Smithiella, into a resurrected and expanded genus, Achudemia, which can be 354 distinguished from other Pilea species and Lecanthus, based on flower, male inflorescence and achene morphology. 355

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Haroldiella was recovered within Pilea. This result was not surprising given that the

justification for treating the taxa within a genus distinct to *Pilea* (Florence, 1997) was based
on the presence of alternate, spirally arranged leaves, a trait state expressed elsewhere within *Pilea* (e.g. *P. peperomiodes*, *P. fairchildiana*).

Whilst almost all of the morphological trait states assessed were found to be homoplastic, our expanded sampling did enable us to recover seven morphologically diagnosable clades within *Pilea* and to use these to establish seven sections, partly congruent with those proposed by Chen (1982) on morphology alone.

364 4.2. Pilea originated in IndoMalaya

Mapped onto our phylogeny, geographical occurrence suggests IndoMalaya as the centre of 365 origin for Pilea (Fig. 4 & Fig. S23), Lecanthus, Achudemia and the basal clades of Pilea (C2 366 367 to C6) predominantly comprising Indomalayan species. This confirms the findings of Monro (2006) and Wu et al. (2018). Given the relationships between geographical areas suggested by 368 369 our results, the most plausible scenario for the dispersal of *Pilea* is of two independent events. The first, early in the divergence of the genus (Fig. 4 & Fig. S23, clade C1) from IndoMalaya 370 to Africa and the northern Neotropics which has resulted in a lineage with a modest number 371 372 of species. The second, later in the divergence of the genus, involving dispersal to the Palearctic (Fig. 4 & Fig. S23, clade C7) and later from the Palearctic to the Neotropics (clade 373 C7) resulting in species radiations in the Andes and Greater Antilles. According to the dated 374 375 Urticaceae phylogeny of Wu et al. (2018, Fig. 1), this first dispersal event would have 376 occurred ca 35 MYA (late Eocene) and the second ca 25 MYA (late Oligocene). A plausible 377 mechanism and route for second dispersal to the Neotropics would be the Bering land bridge 378 at some point between the late Cretaceous and late Neogene (Wen et al., 2016). Whilst long 379 distance dispersal, invoked by Wu et al. (2018) may have played a role in the first.

380 4.3. Diversification accompanied by reductions in merism, achene size and ornamentation
381 and an increase in species number

Chen (1982) proposed a reduction in female flower part number as an evolutionary trend in *Pilea* and our results support this, five-parted flowers occurring amongst the basal clades of our phylogeny (*Lecanthus, Achudemia*), followed by three or four-parted flowers (clades C1-C4) and three or two-parted flowers (clades C5-C7). This includes a clade comprising seven South Pacific species (sampled here as *Haroldiella*) characterised by two-parted female 387 flowers (Florence, 1997). Our analysis of morphological traits recovered three-parted female flowers as plesiomorphy for the node comprising clades C5-C7 (Fig. S19). The trend in 388 389 reduced female part number was matched by a reduction in male flower part number from an ancestral number of five to a derived condition of four, or three in the case of clade C1 (Fig. 390 S15) a transition from imbricate to valvate sepals (Fig. S16). The transition from imbricate to 391 392 valvate arrangement may suggest a transition from non-explosive to explosive anthesis whereby imbricate sepals open in a controlled fashion whilst valvate sepals are torn apart by 393 394 the flexing filaments. The trend of reduced female and male flower part number parallels a 395 decrease in achene size and ornamentation, Lecanthus, Achudemia and clades C1-C4 having achenes > 0.8 mm and ornamented as the ancestral state, whilst clades C5-C7 have achenes \leq 396 397 0.8 mm and clade C7 has smooth surfaces as their ancestral states. It also parallels a change 398 from imbricate to valvate sepals in the male flowers, suggesting a transition from gradual to 399 explosive flower opening (Pedersoli et al., 2019).

400 Whilst we did not seek to test hypotheses about key innovations, we are able to use the results of our ASR analysis to propose hypotheses for future testing. Reduction in merism has 401 402 been associated with key innovations in floral evolution (De Craene, 2016; Kümpers et al., 403 2016; Simões et al., 2017). Our study supports this, a reduction in merism coinciding with an evolutionary radiation, basal five-parted flowers of *Lecanthus* and *Achudemia* being relatively 404 405 species-poor (3 spp, and 5 spp respectively). Pilea, in contrast, which has four-, three-, or 406 two-parted flowers comprises ca 715 spp (Monro, 2004). According to De Craene (2016) a 407 reduction in merism may be driven by an advantage in reducing flower size. Whilst we have 408 not directly measured flower size, achene size is an effective surrogate for the size of the 409 female flower and fruit, both of which are dominated by the single ovary, and later achene. As 410 discussed above, there has been a decrease in achene size with the diversification of *Pilea*, 411 and specifically the species-rich C7 clade. The decrease in achene size coincides with a loss 412 of achene surface ornamentation. Based on an assumption that achene ornamentation 413 indicates animal dispersal, our results suggest a shift in both pollen and fruit dispersal, pollen 414 dispersal becoming explosive and more kinetic and fruit dispersal less reliant on animals.

415 Whilst the above discussion may provide the basis for future research into a reduction in 416 merism and achene surface ornamentation as key innovations, the increase in species number 417 with which they are associated may be unrelated. Increasing species number, focussed in the 418 Greater Antilles and Andes could also be the result of increased reproductive isolation and 419 subsequent speciation through random drift in steeply dissected shaded habitats devoid of 420 strong air-currents. It could, therefore be, an example of an intrinsic driver of speciation rather 421 than a response to the colonisation of novel habitats or development of novel pollination 422 syndromes (Tilston Smith et al., 2014).

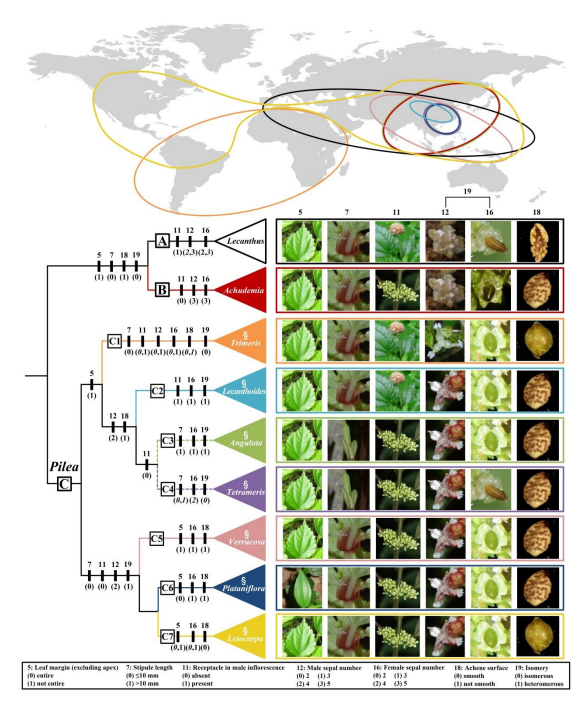
423 *4.4. Proposal for an infrageneric classification*

To date, there have been only two main infrageneric classifications both of which were based on morphological traits (Weddell, 1856, 1869; Chen, 1982; Chen and Monro, 2003). Whilst these classifications facilitate the identification of taxa, they are not good indicators of evolutionary relationships (Monro, 2006).

Chen (1982) proposed a classification for the Chinese taxa that focuses on the traits of female flower merism, leaf nervation, male inflorescence arrangement and male sepal arrangement. This was predated by Weddell's classifications (Weddell, 1856, 1869) based on geographical distribution, leaf incision and heteromorphy. With the exceptions of male sepal arrangement and male inflorescence becoming receptacle-like, the traits used in both classifications were recovered as homoplastic, confirming the results of previous studies (Monro, 2006; Wu et al., 2013, 2015).

We did, however, recover achene morphology as a plesiomorphy useful in the distinction of taxa in the species-rich clade C7 from clades C1-C6 (Fig. S21). Where traits were recovered as homoplastic, they could still be used in combination, and or, with the addition of geographical distribution to delimit monophyletic infrageneric groupings. For example, clade C6, which we propose as section (§) *Plataniflora*, can be delimited by the combination of entire leaf margins, four-parted valvate male flowers and ornamented achene surface. In this way we were able to delimit seven infrageneric groupings.

- The sections (§) that we propose for *Pilea* are, § *Trimeris*, § *Lecanthoides*, § *Angulata*, § *Tetrameris*, § *Verrucosa*, § *Plataniflora* and § *Leiocarpa* (Fig. 3 & Fig. 5).
- 444
- 445



447

Fig. 5. Proposed infrageneric classification of *Pilea* and their geographical distribution. Filled
boxes illustrate all states for each section. *Derived* states in italics, ancestral in regular font (based
on ancestral state reconstructions summarised in Fig. 3). Images illustrate the ancestral states
(replaced by common state when ancestral one is not available) for each infrageneric section.

452

453 **5. Conclusions**

454 We demonstrate that the species-rich genus *Pilea* is paraphyletic with respect to *Achudemia* 455 and polyphyletic with respect to *Haroldiella*. We identify sepal number, flower isomery, male 456 sepal arrangement and achene surface ornamentation as phylogenetically informative traits 457 useful in both the delimitation of generic and infrageneric groupings and the generation of 458 hypotheses about the evolution of the genus and the richness of its neotropical clade. 459 Translating these findings into taxonomic actions resulted in the resurrection of *Achudemia* 460 and the revised delimitation and infrageneric classification of *Pilea*. Our findings provide a 461 stable framework for future research aimed at answering broader questions in evolutionary 462 biology, such as whether intrinsic factors can drive species radiations.

463

464 6. Taxonomy

465 6.1. Revised delimitation of Achudemia and Pilea

466 Achudemia Blume, Mus. Bot. 2: 57, 1856. Genus type: *A. javanica* Blume, Mus. Bot. 2: 57, 1856.
467 1856.

Smithiella Dunn. Bull. Misc. Inform. 1920: 210, 1920. Genus type: *Smithiella myriantha*Dunn. nom. illeg., non *Smithiella* H. Perag. & Perag. = *Aboriella* Bennet, Indian Forester 107:
437, 1981. = *Dunniella* Rauschert, Taxon 31: 562, 1982.

Herbs, perennial or annual. Stems without stinging hairs, not releasing watery latex when cut. Leaves opposite, frequently subequal at each node, the margins toothed; cystoliths fusiform; stipules borne in axils of the leaves, persistent or caducous. Inflorescences unisexual or bisexual, paniculate, capitate or racemose cymes; pedicels subtended by inconspicuous bracteoles. Male flowers 5-merous; sepals imbricate in bud, equal, each bearing a subapical appendage. Female flowers 5-merous; the sepals 5, equal or subequal, not dimorphic. Achenes compressed ovoid, ornamented.

Five species, restricted to the Palearctic and IndoMalayan biogeographic regions.
Associated with forested rocky habitats. *Achudemia japonica* is used as a medicine to treat
fever and as a diuretic (Chen and Monro, 2003)

Note:— Blume (1856) in his description of the genus refers to polygamous, hermaphrodite
flowers. We believe this to have been an editorial error as neither, material collected by
Blume at L, or the illustration which serves as type, include polygamous or hermaphrodite
flowers.

486 *New combinations and typifications*

- 487 Achudemia subpubera (Miq.) Y.G.Wei & A.K.Monro. comb. nov. = Pilea subpubera Miq.
- 488 Syst. Verz. Ind. Archip. 2: 102, 1854. Type: [Indonesia], Bandong Province, H. Zollinger 870

489 (holotype, U (U0226171*); isotypes P (P 02428341*), (P 02428342*))

- 490 Achudemia javanica Blume, Mus. Bot. 2: 57, 1856. TYPE: Mus. Bot. 2: 57, 1856, t. 20
- 491 (holotype). Epitype (selected here): [Indonesia] Java, C.L. Blume s.n. L (10039782)*
- 492 We have selected an epitype as the type material comprises an illustration
- 493 (<u>https://www.biodiversitylibrary.org/item/200679#page/274/mode/1up</u>) which is not adequate
- for making observations of anatomy of the leaf or stem. We selected material collected by
- Blume in Java, which may have served as the subject of the type illustration.
- 496
- Achudemia boniana (Gagnep.) L.F.Fu & Y.G.Wei. comb. nov. ≡ Pilea boniana Gagnap.,
 Bull. Soc. Bot. France 75: 71. 1928. Type: Indochina [Vietnam], [Hà Nam Province]
 Kien-kh é, Dong-ham rocks, *R.P. Bon* 2522 (holotype P (P06817992)*). Epitype (selected here): [Vietnam], Tonkin, Dong-Dang, on calcareous rocks, 12 Feb. 1886, B. *Balansa 581* (P
- 501 (P06817995)*).
- 502 *P. morseana* Hand.-Mazz., *Symb. Sin.* 7: 140. 1929. Type: China, Guangxi, Longzhou, *Morse*503 495 (holotype K (K000708579)*)
- *P. pentasepala* Hand.-Mazz., Symb. Sin. 7: 128. 1929. Type: China: Yunnan, mountains of
 Mengzi, 1800 m, *Henry 9771* (holotype K (K000708578)*)
- We have selected an epitype for *Achudemia boniana* as the holotype comprises leafless
 material and leaves include several traits useful for species delimitation in *Achudemia*.
- 508
- 509 Achudemia hilliana (Hand.-Mazz.) L.F.Fu & Y.G.Wei. comb. nov. ≡ Pilea hilliana
 510 Hand.-Mazz., Symb. Sin. 7: 129. 1929. Type: China, Yunnan, Möngdse [Mengzi], Henry
 511 10295 (lectotype (selected here) K (K000708583)*).
- 512
- 513 Achudemia myriantha (Dunn) L.F.Fu & Y.G.Wei. comb. nov. ≡ Smithiella myriantha Dunn,
- 514 Bull. Misc. Inform. Kew 1920: 211. 1920. Type: [India] Eastern Himalaya, Outer Abor Hills,
- sunless side of the Dihong Gorge below Rotung, 300 m. Jan 3 1912, *Burkill* 37636 (lectotype

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516	(selected here)	K	(K000708616)*).	Dilaa	myriantha	(Dunn)	C I Chan	nom	illag	non	D
510	(selected nere)	N	$(KUUU/U8010)^{*}$).	Рпеа	myriantna	(Dunn)	C.J.Chen	nom.	meg.,	non	Ρ.

517 myriantha Killip, Bull. Bot. Res., Harbin 2: 44. 1982. P. spicata C.J. Chen & A.K. Monro,

518 Novon 17: 26. 2007.

- 519
- 520 Pilea Lindl., nom. cons., Coll. Bot. ad t. 4. 1821. Genus type: P. muscosa Lindl. nom. illeg.

521 superfl. = *Parietaria microphylla* L. = *Pilea microphylla* (L.) Liebm.

522 Adicea Raf., nom. nud. First Cat. Gard. Transylv. Univ.: 13. 1824.

523 Adicea Raf. ex Britton & A. Br., Ill. Fl. N. U.S. 1: 533. 1896. nom. illeg. superfl.,

524 Adike Raf., New Fl. 1: 63. 1836. Genus type: A. pumila Raf.

525 *Chamaecnide* Nees & Mart. ex Miq., in C.F.P.von Martius & auct. suc. (eds.), Fl. Bras. 4:

526 203. 1853. Genus type: *C. microphylla* Nees ex Miq.

527 *Dubrueilia* Gaudich., Voy. Uranie: 495. 1830. Genus type: *D. peploides* Gaudich.

- 528 Haroldiella J.Florence, Fl. Polynésie Franç 1: 218. 1997. Genus type: *H. rapaensis*529 J.Florence.
- *Neopilea* Leandri, Ann. Mus. Colon. Marseille, s ér. 6, 7-8: 46. 1950. Genus type: N. *tsaratananensis* Leandri

532 *Sarcopilea* Urb., Symb. Antill. 7: 201. 1912. Genus type: *S. domingensis* Urb.

Herbs, rarely shrubs, occasionally epiphytic, perennial, rarely annual. Stems without 533 534 stinging hairs, not releasing watery latex when cut. Leaves opposite, frequently unequal at each node, the margins toothed or entire; cystoliths fusiform; stipules borne in axils of the 535 leaves, persistent, rarely caducous. Inflorescences unisexual, rarely bisexual, paniculate, 536 537 capitate or rarely fused cymes; pedicels subtended by inconspicuous bracteoles. Male flowers 4- or rarely 2- or 3-merous; sepals valvate, equal, each bearing a subapical appendage. Female 538 539 flowers 3-, or rarely 2- or 4-merous, unequal, dimorphic, the adaxial sepal of the larger sepal 540 frequently bearing a dorsal thickening. Achenes weakly to strongly compressed ovoid to 541 sub-ellipsoid, smooth or ornamented. Approx. 710 spp. Cosmopolitan, except for Australia 542 and New Zealand. A number of species cultivated as ornamentals.

543

544 New combinations

545 Pilea australis L.F.Fu & A.K.Monro, nom. nov. Replaced name: Haroldiella rapaensis

546	J.Florence,	Fl	Polyn ásie	Fran c	1.	220	(1997)	TYPE	French	Polynesia	Austral	Islands
540	J.I.IUICHCC.	I'l.	rownesie	r run c.	1.	220	(177/).	IIFL.	TICHCH	r urvnesia.	Ausuai	islanus.

- 547 Rapa, eastern flank of Nt. Perau, 610 m, 21 Jul. 1934, H. St. John, FR. Fosberg & J. Maireau
- 548 15643 (holotype B1SH).
- 549 Note:- Pilea australis was created as a replacement name because a homonym, Pilea
- 550 *rapensis*, has been published by Forest Brown (Brown, 1935).
- 551
- 552 Pilea sykesii (J.Florence) L.F.Fu & A.K.Monro. comb. nov. = Haroldiella sykesii
- J.Florence, Fl. Polynésie Franç 1: 221 (1997). TYPE: French Polynesia, Austral Islands,
- Raivavae, Anatonu, Falaise centrale, 140 m, 10 May 1992, J. Florence & W.R. Sykes 11336
- 555 (holotype P (P 00637067)*, isotype PAP)
- 556
- 557 6.2. Key to the sections of Pilea
- 1. Male and female flowers with the same merism, 3-parted or 4-parted 2
- 1. Male and female flowers with different merisms, 2-, 3- or 4-parted 3
- 560
- 561 2. Merism of 3, rarely 4 (*P. tetraphylla*), stipules ≤ 10 mm in length, tropical Africa,
- 562 neotropics. **§Trimeris**
- 563 2. Merism of 4, stipules > 10 mm in length, Indomalaya. **§ Tetrameris**

564

- 3. Male inflorescence an unbranched and fused receptacle-like capitulum, involucrate.
- 566 **§Lecanthoides**
- 567 3. Male inflorescence branched or unbranched, where unbranched capitulum globose or
- subglobose, not involucrate 4

- 570 4. Achenes ornamented 5
- 571 4. Achenes not ornamented or rarely so, where ornamented Indomalayan and either P.
- 572 *melastomatoides*, or *P. peploides*. **§Leiocarpa**
- 573
- 574 5. Stipules > 10 mm in length. **§Angulata**
- 575 5. Stipules $\leq 10 \text{ mm in length}$ 6

576 6. Leaf margins incised. § Verrucosa 577 578 6. Leaf margins entire. § Plataniflora 579 6.3. Infrageneric classification of Pilea 580 581 Pilea § Trimeris Y.G. Wei & A.K. Monro, sect. nov. — Section type: P. tripartita A.K. 582 Monro. 583 Herbs. Stipules < 10 mm in length. Leaf margin incised. Male inflorescence a capitate cyme, involucrate; male and female flowers with the same merism, three- rarely four- parted; 584 585 the achene > 0.8 mm in length, not ornamented. Ca three spp. Tropical Africa, Neotropics. 586 Pilea § Lecanthoides C.J.Chen, Bull. Bot. Res. 2(3): 118. 1982. Section type: P. 587 588 receptacularis C.J.Chen. Herbs. Stipules ≤ 10 mm in length. Leaf margin incised. Male inflorescence an 589 unbranched and fused capitulum, involucrate; male and female flowers with different merism, 590 591 male flowers four-parted, female flowers three-parted; the achene > 0.8 mm in length, 592 ornamented. Two spp. Indomalaya. 593 Pilea §Angulata L.F.Fu & Y.G.Wei, sect. nov. — Section type: P. angulata (Blume) Blume 594 595 Herbs. Stipules > 10 mm in length. Leaf margin incised. Male inflorescence a branched cyme; male and female flowers with different merism, male flowers four-parted, female 596 597 flowers three-parted; achene > or ≤ 0.8 mm in length, ornamented. Ca two spp. Indomalaya. 598 599 Pilea § Tetrameris C.J.Chen, Bull. Bot. Res. 2(3): 44. 1982. Section type: P. basicordata 600 W.T.Wang. Herbs. Stipules > 10 or rarely \leq 10 mm in length. Leaf margin incised. Male 601 inflorescence a branched cyme; male and female flowers with the same merism, flowers 602 603 four-parted; achene > 0.8 mm in length, ornamented. Ca 15 spp. Indomalaya. 604 Pilea § Verrucosa L.F.Fu & Y.G.Wei, sect. nov. Type: P. gracilis Hand.-Mazz. 605

606	Herbs. Stipules \leq 10 mm in length. Leaf margin incised. Male inflorescence a branched
607	cyme; male and female flowers with different merism, male flowers four- or occasionally
608	two-parted, female flowers three-parted, rarely two-parted; achene > or ≤ 0.8 mm in length,
609	ornamented. Ca 80 spp. Indomalaya.
610	
611	Pilea §Plataniflora L.F.Fu & Y.G.Wei, sect. nov. Type: P. plataniflora C.H.Wright.
612	Herbs. Stipules ≤ 10 mm in length. Leaf margins entire. Male inflorescence a branched
613	cyme; male and female flowers with different merism, male flowers four-parted, female
614	flowers three-parted; achene > or ≤ 0.8 mm in length, ornamented. Ca 34 spp. Indomalaya.
615	
616	Pilea §Leiocarpa L.F.Fu & Y.G.Wei, sect. nov. Type: P. micropylla (L.) Liebm.
617	Herbs. Stipules $>$ or \leq 10 mm in length. Leaf margins entire or incised. Male
618	inflorescence a branched or capitate cyme; male and female flowers with different merism,
619	male flowers four- or rarely two-parted, female flowers four- or rarely two-parted, achene >
620	or ≤ 0.8 mm in length, not ornamented. Ca 570 spp, Indomalaya, Neotropics, Australasia,
621	Palearctic, Nearctic.
622	
623	6.4. Excluded names
624	Metapilea W.T.Wang, Bull. Bot. Res., Harbin 36: 164. 2016. Genus type: M. jingxiensis
625	W.T.Wang.
626	We have some doubt over the position of this taxon based on the poor quality of the
627	material upon which the description and illustration are based (a single sheet, very few
628	immature flowers). One individual of this taxon was collected once in a relatively common
629	habitat in Guangxi Province, China, in 1973 (Wang s.n.) and it has not been collected since. It
630	is possible that the original material may be an immature collection of <i>Pilea</i> , to which it is

- vegetatively identical, or a distinct genus, which, based on the illustrations, could be allied to *Elatostema* or *Procris*. Due to the sampling policy of the herbarium where the type collection
 is stored it is not possible to sample this material for DNA and so its status and position
- 634 remains uncertain.

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- 645

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