1 Plastid phylogenomics clarifies broad-level relationships in

2 Bulbophyllum (Orchidaceae) and provides insights into range evolution

3 of Australasian section Adelopetalum

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- 18

19 Abstract

20 The hyper diverse orchid genus *Bulbophyllum* is the second largest genus of flowering plants and

- 21 exhibits a pantropical distribution with a center of diversity in tropical Asia. The only Bulbophyllum
- 22 section with a center of diversity in Australasia is sect. *Adelopetalum*. However, phylogenetic
- 23 placement, interspecific relationships, and spatio-temporal evolution of the section have remained
- 24 largely unclear. To infer broad-level relationships within Bulbophyllum and interspecific
- 25 relationships within sect. *Adelopetalum*, a genome skimming dataset was generated for 89 samples,
- 26 yielding 70 plastid coding regions and the nuclear ribosomal DNA cistron. For 18 additional samples,
- 27 Sanger data from two plastid loci (*mat*K, *ycf*1) and nuclear ITS were added using a supermatrix
- 28 approach. The study provided new insights into broad-level relationships in *Bulbophyllum*, including
- 29 phylogenetic evidence for the non-monophyly of sections *Beccariana, Brachyantha, Brachypus,*
- 30 Cirrhopetaloides, Cirrhopetalum, Desmosanthes, Minutissima, Oxysepala, Polymeres and
- 31 Sestochilos. Section Adelopetalum and sect. Minutissima s.s. formed a highly supported clade that
- 32 was resolved in sister group position to the remainder of the genus. Divergence time estimations
- based on a relaxed molecular clock model placed the origin of *Bulbophyllum* in the early Oligocene
- 34 (ca. 33.2 Ma) and of sect. *Adelopetalum* in the late Oligocene (ca. 23.6 Ma). Ancestral range
- 35 estimations based on a BAYAREALIKE model identified the Australian continent as ancestral area 36 of sect. *Adelopetalum*. The section underwent crown diversification during the mid-Miocene to the
- 37 late Pleistocene, predominantly in continental Australia. At least two independent long-distance
- dispersal events were inferred eastwards from the Australian continent to New Zealand, and New
- 39 Caledonia from the early Pliocene onwards, likely mediated by the predominantly westerly winds of

40 the southern hemisphere. Retraction and fragmentation of eastern Australian rainforests from the

41 early Miocene onwards are discussed as likely drivers of lineage divergence within sect.

42 Adelopetalum, facilitating allopatric speciation.

43 1 Introduction

44 The hyper diverse orchid genus *Bulbophyllum* Thouars (Epidendroideae) is the second largest genus

45 of flowering plants with more than 2,100 species and exhibits exceptional morphological and

46 ecological diversity (Frodin, 2004; Pridgeon et al., 2014, WCSP 2022). Species of this predominantly

47 epiphytic genus occur in a wide range of tropical and subtropical habitats, from montane rainforests

48 to dry deciduous forests, savannah woodlands, and rocky fields with shrubby vegetation (Pridgeon et

49 al., 2014). *Bulbophyllum* is distributed pantropically, occupying all botanical continents defined by

50 Brummit (2001) except for Antarctic and Eurasia. The genus is most diverse on the botanical 51 continent of tropical Asia (1562 species), also occurring on the botanical continents of Africa (305),

52 temperate Asia (152), Southern America (88), the Pacific (49), Australasia (Australia and New

52 Temperate Asia (152), Southern America (88), the Factice (49), Australiasia (Australia and New 53 Zealand; 35), and Northern America (7) (WCSP, 2022). Centres of diversity are found in tropical

Asia in the floristic regions of Malesia (667) and Papuasia (656) and in the Afrotropics in the western

55 Indian Ocean region, on the islands of Madagascar, and the Mascarenes (218) (WCSP, 2022).

56 The high number of species and complex patterns of morphological variation has presented

57 significant challenges for resolving relationships in *Bulbophyllum* and this is reflected in substantial

58 taxonomic revisions that have been proposed. Traditionally, the subtribe Bulbophyllinae Schltr. (tribe

59 Dendrobieae Endl.) included the large genus *Bulbophyllum* along with smaller genera, such as

60 Cirrhopetalum Lindl., Drymoda Lindl., Pedilochilus Schltr., Sunipia Buch.-Ham. ex Sm., and Trias

61 Lindl. (Dressler, 1993; Garay et al., 1994; Szlachetko and Margonska, 2001). Recent revisions treat

62 all genera within the subtribe Bulbophyllinae in a more broadly defined *Bulbophyllum* and recognise

97 sections within the genus (Pridgeon et al., 2014; Vermeulen et al., 2014). Molecular phylogenetic
 studies have largely focused on species from specific geographic regions such as Madagascar and the

- 65 Mascarenes (Fischer et al., 2007; Gamisch et al., 2015), the Neotropics (Smidt et al., 2013, 2011),
- and Peninsular Malaysia (Hosseini et al., 2012) or on taxonomic groups such as the *Cirrhopetalum*

67 alliance (Hu et al., 2020), and few have taken a global perspective (e.g., Gamisch and Comes, 2019).

68 These studies revealed a strong biogeographic pattern within the genus with four main clades that

69 include species largely confined or endemic within one broader geographical area: 1) continental

Africa, 2) Madagascar and the Mascarene Islands, 3) Southern America, or 4) Asia (Fischer et al.,

- 71 2007; Gamisch et al., 2015; Gamisch and Comes, 2019; Smidt et al., 2011). The Southern American
- clade, the Madagascan clade, and the continental African clade together form a highly supported

73 lineage (Fischer et al., 2007; Gamisch et al., 2015; Gamisch and Comes, 2019; Smidt et al., 2013;

2011), in sister group position to the Asian clade (Fischer et al., 2007; Gamisch et al., 2015; Gamisch

and Comes, 2019). Previous molecular phylogenetic studies have mainly elucidated relationships

76 within Madagascan, continental African and Neotropical sections and within the *Cirrhopetalum*

alliance (Fischer et al., 2007; Gamisch et al., 2015; Smidt et al., 2011; Hu et al., 2020). However,

78 evolutionary relationships within the Asian clade, which also includes taxa from the Australasian and

79 Pacific regions, are still poorly understood, and the monophyly of sections within this clade has

80 remained largely untested within a phylogenetic framework.

81

82 The study of hyper diverse groups such as *Bulbophyllum* requires a robust phylogenetic framework to

83 assess monophyly of its infrageneric taxa. High throughput sequencing approaches facilitate the

84 establishment of such a framework phylogeny to clarify broader evolutionary relationships and to

assess the monophyly of infrageneric taxa and their trait evolution (Hassemer et al., 2019; van

86 Kleinwee et al., 2022, Nargar et al. 2022). However, phylogenomic studies which provide insights

87 into broad-level evolutionary relationships within the Asian clade of *Bulbophyllum* are still lacking.

- 88 This hampers progress in understanding diversification of its evolutionary lineages in time and space
- 89 and trait evolution within this highly diverse genus.

90 Within Bulbophyllum, section Adelopetalum has a unique distribution, being the only section with a 91 centre of diversity in Australia (Brummit, 2001; Pridgeon et al., 2014, Vermeulen, 1993), and thus 92 presents an interesting study case for range evolution within Bulbophyllum. The section comprises 93 twelve tropical to temperate epi-lithophytic species. Nine species occur along Australia's east coast 94 in the montane forest communities of the Great Dividing Range, with one species (B. argyropus) also 95 found on Australian islands (Lord Howe Island, and Norfolk Island). Two species are endemic to the 96 montane forests of New Caledonia (B. corythium, B. lingulatum) and one to the lowland coastal 97 forests of New Zealand (B. tuberculatum). The section was circumscribed based on morphological 98 affinities recognised among ten species from Australia and New Caledonia previously assigned to 99 Bulbophyllum sections Desmosanthes, Racemosae and Sestochilus (Dockrill, 1969; 1992; 100 Vermeulen, 1993). Subsequent treatments recognised two additional species within the section, B. 101 weinthalii and B. exiguum (Jones and Clements, 2002; Clements and Jones, 2006). Section 102 Adelopetalum is characterised by plants having thin creeping rhizomes adpressed to the host, 103 anchored by filamentous roots with small pseudobulbs that are crowded to widely spaced, and a 104 small single flat leaf arising from the apex of the pseudobulb. The inflorescence is single to few-

105 flowered, with small white, cream or yellow flowers, sometimes with red or purple patterns. The

- 106 petals are smaller than the sepals but similar in shape, with the bases of the lateral sepals fused to the
- 107 column foot. The fleshly three-lobed labellum is firmly hinged to the apex of the column foot.

108 Previous cladistic analysis of sect. *Adelopetalum* based on morphological characters resolved two

- 109 main clades within the section, differentiated by the size and shape of the lower margin of the
- 110 stelidia: the filiform column appendages typical for most *Bulbophyllum* (Vermeulen, 1993). Previous
- 111 molecular phylogenetic studies based on the nuclear ribosomal ITS region (ITS1 + 5.8S + ITS2)
- included two to three representatives of the section (Gamisch et al., 2015; Gamisch and Comes,
- 2019), placing these in an early diverging position within the Asian clade. However, phylogenetic
 placement of the section within *Bulbophyllum* was not strongly supported (PP<90, BS 97) and thus
- requires further study. Further, phylogenetic relationships within sect. *Adelopetalum* and its ancestral
- requires further study. Further, phylogenetic relationships within sect. *Adelopetatum* and its ancest range evolution are poorly understood and have not vet been investigated using a molecular
- range evolution are poorly understood and have not yet been investigated using a molecular
- 117 phylogenetic approach.
- 118 The aims of this study were to 1) build a phylogenomic framework for *Bulbophyllum* with focus on
- 119 the Asian clade 2) assess the monophyly and phylogenetic placement of sect. *Adelopetalum* within

120 Bulbophyllum; 3) to infer interspecific relationships within sect. Adelopetalum 4) and to reconstruct

121 the range evolution of sect. *Adelopetalum*.

122 **2** Materials and methods

123 **2.1 Sampling**

124 In total, 136 orchid samples representing 114 species were included in the study. From the Asian,

- Australasian and Pacific regions a broad sampling was included representing 41 sections, i.e. 60% of
- sections recognised from these regions in the most recent treatment of the group (Pridgeon et al.,
- 127 2014). From the Australasian region, all *Bulbophyllum* species were sampled (Australia: 30, New

- 128 Zealand: 2). For Bulbophyllum sect. Adelopetalum, 28 samples were included, representing all 12
- 129 species recognised for the section. The morphologically closely related sect. Minutissima was
- 130 included with nine samples representing five species comprising all four Australasian and Pacific
- 131 species and two (of ca. 19) tropical Asian species. Sampling of representative species for the
- 132 Afrotropical and Neotropical clades of *Bulbophyllum* was informed by previous phylogenetic studies
- 133 (Fischer et al., 2007; Smidt et al., 2011; Gamisch and Comes, 2019). Species names follow the
- 134 accepted taxonomy based on the World Checklist of Selected Plant Families (WCSP, 2022) and
- 135 sectional taxonomy IOSPE (2022). Exceptions were made for *B. exiguum* which was placed in
- 136 section Adelopetalum and B. wolfei which was placed in section Polymeres based on Jones and 137
- Clements (2002) and Clements and Jones (2006). The outgroup comprised representatives of subtribe
- 138 Dendrobineae which is sister to Bulbophyllinae, and tribes Malaxideae, Arethuseae, Nervilleae, and 139
- Neottieae based on previous molecular phylogenetic studies (Givnish et al., 2015; Górniak et al., 140 2010, Serna-Sánchez et al. 2021). Details of the plant material studied, voucher information and the
- 141 number of loci included for each sample are provided in Supplementary Material S1 and a complete
- 142 list of loci analysed is provided in Supplementary Material S2.

143 2.2 DNA extraction, amplification, and sequencing

- 144 Total genomic DNA was extracted from ca. 10 to 20 mg silica-dried leaf material. Extractions were
- 145 carried out with commercial extraction kits (Qiagen DNeasy plant kit, Venlo, Netherlands;
- 146 ChargeSwitch gDNA plant kit, Invitrogen, Carlsbad, USA) following the manufacturer's protocols or
- 147 using the CTAB method (Doyle and Doyle, 1990), with modifications as described in Weising et al.
- 148 (2005). Sequence data was generated using both Sanger sequencing (46 samples) for the nuclear
- 149 ribosomal ITS region (ITS1, 5.8s, ITS2) and two plastid genes (matK, ycf1) and shotgun high-
- throughput sequencing (89 samples) to extract 70 plastid coding sequences (CDS) and the nuclear 150 151
- ribosomal DNA cistron (Supplementary Material S2) for subsequent analyses. Libraries for high-152 throughput sequencing were constructed from 50 to 100 ng total DNA using the TruSeq Nano DNA
- 153 LT library preparation kit (Illumina, San Diego, USA) for an insert size of 350 base pairs (bp) and
- 154 paired-end reads following the manufacturer's protocol. Libraries were multiplexed 96 times and
- 155 DNA sequencing with 125 bp paired-end reads was carried out on an Illumina HiSeq 2500 platform
- 156 at the Australian Genomic Research Facility, Melbourne (Australia).
- 157 For Sanger sequencing, amplifications for ITS were carried out with primers 17F and 26SER (Sun et
- 158 al., 1994), for matK with the primers 19F and, 1326R (Cuénoud et al., 2002), and for vcfl with
- 159 primers 3720F, intR, intF, 5500R (Neubig et al., 2008). PCR reaction protocols are provided in
- 160 Supplementary Material S3. Sequencing reactions were carried out using the amplification primers
- and sequencing was conducted on an AB3730x1 96-capillary sequencer (Australian Genome 161
- 162 Research Facility, Brisbane, Australia).

163 2.3 Assembly and alignment

- Sequences were assembled and edited in Geneious R10 (Kearse et al., 2012). Illumina sequences 164
- 165 were assembled to a reference set of plastid CDS extracted from *Dendrobium catenatum* (GenBank
- 166 accession numbers KJ862886) and for vcf68 from Anoectochilus roxburghii (KP776980). To build a
- 167 reference for the nuclear ribosomal ITS-ETS region, Illumina reads of *B. boonjee* (CNS G07175)
- 168 were first mapped to the ITS-ETS region of Corallorhiza trifida (JVF2676a). To extend the region
- 169 assembled, the B. boonjee Illumina reads were then mapped to the B. boonjee consensus sequence
- 170 generated in the initial step, yielding a *B. boonjee* reference of the nuclear ribosomal DNA cistron
- 171 (5'ETS, 18s, ITS1, 5.8s, ITS2, 28s, 3'ETS). Illumina sequences for all other samples were assembled
- 172 against the B. boonjee nuclear ribosomal DNA cistron reference. Assemblies were carried out with

- the highest quality threshold and a minimum coverage of ten reads. The quality of the assemblies was
- 174 checked and edited manually where required. Sequences were deposited in GenBank and ENA. For
- 175 Sanger sequences, bidirectional reads were assembled in Geneious and edited manually. Additional
- sequences were sourced from DRYAD (https://doi.org/10.5061/dryad.n9r58) for *Coelogyne flaccida*
- 177 (Givnish et al., 2015). DNA sequences were aligned using MAFFT v.7.222 (Katoh et al., 2005, 2002)
- 178 with the default settings, visually inspected and then concatenated into a nuclear and plastid
- supermatrix, respectively. The nuclear supermatrix included 136 accessions, partitioned into coding
 and non-coding regions (alignment length: 6,341bp, number of parsimony informative sites: 995
- and non-coding regions (alignment length: 6,341bp, number of parsimony informative sites: 99
 (16%)); and the plastid supermatrix included 130 accessions, and 70 plastid coding regions,
- partitioned by gene and codon position (alignment length: 61,553bp, number of parsimony
- informative sites: 5,789 (9%)). A plastid dataset comprised of high throughput sequencing data only
- (excluding Sanger sequences) was produced including 90 accessions, and 70 plastid coding regions
- (alignment length: 61,553bp, number of parsimony informative sites: 5682 (9%) and analysed
- 186 separately.
- 187 For divergence time estimations, the plastid supermatrix was reduced to one representative per
- 188 species (indicated by an asterisk in Supplementary Material S1), comprising 111 accessions
- 189 (alignment length: 60,984 bp, number of parsimony informative sites: 5,755 (9%)).

190 2.4 Phylogenetic analysis

- 191 Phylogenetic relationships were inferred using maximum likelihood (ML) in IQ-TREE v. 1.6.12
- 192 (Nguyen et al., 2015). The best-fit partition scheme and nucleotide substitution model for each
- 193 partition was determined with IQ-TREE's ModelFinder (Kalyaanamoorthy et al. 2017) based on the
- 194 Akaike information criterion (AIC) (Akaike, 1974). Nodal support was assessed based on 1000
- 195 replicates of ultrafast bootstrap approximation with clades receiving >95 ultrafast bootstrap support
- 196 (UFBS) considered as well supported (Minh et al., 2013; Hoang et al., 2018).

197 **2.5 Divergence time estimation**

198 Divergence times were estimated based on the plastid dataset in Beast2 v. 2.4.8 (Bouckaert et al., 199 2014) applying the best fit partition scheme and substitution model as determined by IQ-TREE's 200 ModelFinder. We tested two molecular clock models: 1) strict clock (Zuckerkandl and Pauling, 201 1965) and 2) relaxed lognormal clock (Drummond et al., 2006) and two models of speciation and 202 extinction: 1) Yule and 2) birth-death (Yule, 1925; Gernhard et al., 2008). Three secondary 203 calibration points were used applying priors with a normal distribution and mean ages and 95% 204 higher posterior density (HDP) intervals based on the results of a family-wide molecular clock 205 analysis by Chomicki et al. (2015). The root age was set to 55.02 Ma (HDP: 42.0-68.0). The next 206 secondary calibration point was applied to the last common ancestor of Dendrobineae, Malaxideae, 207 and Arethuseae and was set to 47.77 Ma (HDP: 36.4-59.1). Monophyly was constrained for this 208 node consistent with relationships reconstructed in previous phylogenetic analyses (Chomicki et al. 209 2015; Givnish et al., 2015). The last secondary calibration was set at the stem node of Dendrobieae 210 and Malaxideae with 38.68 Ma (HDP: 30.8-46.6). An additional calibration based on the fossil 211 Dendrobium winikaphyllum (Conran et al., 2009) was applied to the stem node of the Australasian 212 Dendrobium clade (D. macropus, D. cunninghamii, and D. muricatum), using a uniform distribution 213 with an infinite maximum age and the minimum age constrained to 20.4 Ma, based on the minimum 214 age of the strata containing the fossil (Mildenhall et al. 2014). Ten independent Beast analyses were 215 run for 30 million MCMC generations, with trees sampled every $3x10^4$ generations. To assess 216 convergence of independent runs and determine burn-in fractions, log files were assessed in Tracer 217 v.1.7.1 (Rambaut and Drummond, 2007). Log and trees files from independent runs were combined

218 in LogCombiner (from the Beast package) with a cumulative burn-in fraction of 10%-31% and the

219 sampling frequency set to generate at least 10,000 tree and log files (Drummond and Bouckaert,

- 220 2015). The combined log file was assessed in Tracer to ensure the effective sample size of all
- 221 parameters was above 200. An additional five independent Beast runs were conducted for the final
- 222 analysis using a relaxed log normal clock with birth death speciation to achieve an effective sample 223 size above 200 for the ucldmean parameter. A maximum clade credibility tree was generated in
- 224
- TreeAnnotator (Beast package) with median node heights. To compare clock and speciation models, 225 the Akaiki information criterion by MCMC app from the BEAST 2 package v 2.6.2 was used to
- 226 measure the AICM for the combined MCMC runs generated in the BEAST analysis for each model
- 227 (Supplementary Material S4).

228 2.6 Ancestral range analysis

229 Species distributions were extracted from WCSP (2022). Biogeographic areas were largely

- 230 delineated based on botanical continents defined by Brummit (2001). The subcontinental regions of
- 231 Papuasia, Australia and New Zealand were recognised to allow a more fine-scaled resolution of range
- 232 evolution in section Adelopetalum (Brummit 2001). The following seven biogeographic areas were
- 233 coded: a, Africa; b, temperate Asia; c, tropical Asia; d, Papuasia; e, Australia; f, New Zealand, and g,
- 234 Pacific. Ancestral ranges were estimated in RASP v. 4.0 (Yu et al., 2015) with the BioGeoBEARS 235 package (Matzke, 2013) based on the maximum clade credibility tree obtained from the Beast
- 236 analysis of the plastid supermatrix, pruned of the outgroups to Dendrobieae. Three models of range
- 237 evolution were tested: the dispersal-extinction cladogenesis model (DEC) (Ree and Smith, 2008), a
- 238 ML version of Ronquist's parsimony dispersal-vicariance (DIVA; Ronquist, 1997), termed
- 239 DIVALIKE (Matzke, 2013), and a simplified likelihood interpretation of the Bayesian "BayArea"
- 240 program (Landis et al., 2013) known as BAYAREALIKE (Matzke, 2013). No constraints were
- 241 applied to dispersal direction and the maximum number of ranges was set to five based on the
- 242 maximum number of observed areas in extant species. Likelihood values were compared and the
- 243 model of best fit determined by AIC score (Akaike, 1974) was used to infer the marginal
- 244 probabilities of alternative ancestral ranges at each node in the phylogeny (Supplementary Material 245 S5).
- 246 3 **Results**
- 247 3.1 **Phylogenetic relationships**

248 3.1.1 Phylogenetic relationships – Plastid data

249 The ML phylogeny inferred from the 70 loci plastid supermatrix provided strong support for the

- 250 monophyly of *Bulbophyllum* and its sister group relationship to *Dendrobium* (Fig. 1). Section
- 251 Adelopetalum and Minutissima s.s. formed a highly supported clade, here termed the
- 252 Adelopetalum/Minutissima clade, which was resolved in sister group position to the remainder of the
- 253 genus (ultrafast bootstrap support/UFBS 98) (Fig. 1, Fig. 2). Within the Adelopetalum/Minutissima
- 254 clade, all Adelopetalum species plus B. pygmaeum (sect. Minutissima) formed a highly supported
- 255 lineage (UFBS 100), here termed the Adelopetalum clade. Within the Adelopetalum clade several
- 256 highly supported groups were resolved: 1) the argyropus clade consisting of *B. argyropus*, *B.*
- 257 corvthium and B. tuberculatum (UFBS 100), reconstructed in a highly supported sister group
- 258 relationship to *B. weinthalii* (UFBS 100); 2) the bracteatum clade, including *B. boonjee*, *B.*
- 259 bracteatum, and B. elisae (UFBS 99); and 3) the newportii clade comprised of B. exiguum, B.
- 260 lageniforme, B. lilianae, B. lingulatum, and B. newportii (UFBS 100). Relationships among B.
- 261 *pygmaeum*, the argyropus clade + *B. weinthalii*, bracteatum and newportii clades received weak

support. Sister to the Adelopetalum clade was the highly supported Minutissimum clade comprised

263 of three species of sect. *Minutissima* (B. globuliforme, B. keekee, B. minutissimum), including the

type species of the section (UFBS 100) (Fig. 1). Section *Minutissima* was identified as polyphyletic,

with sect. *Minutissima* species placed within the Adelopetalum clade (*B. pygmaeum*) and the Asian

clade (*B. mucronatum, B. moniliforme*). Within the Asian clade, sections *Beccariana, Brachyantha*,

267 Brachypus, Cirrhopetaloides, Cirrhopetalum, Desmosanthes, Oxysepala, Polymeres and Sestochilos

268 were identified as polyphyletic or paraphyletic. Phylogenetic relationships described here based on

the plastid supermatrix (Fig.1, Fig. 2) are supported by reconstructions based on the 70 gene plastid

270 dataset (Supplementary Material S6).

- 271 Our analyses showed that sect. *Adelopetalum* does not share a close relationship with other
- Australasian *Bulbophyllum* species, such as those in sect. *Brachypus (B. nematopodum)*, sect.
- 273 Brachystachyae (B. evasum), sect. Cirrhopetalum (B. longiflorum), sect. Ephippium (B.
- 274 gracillimum), sect. Monanthes (B. macphersonii), sect. Oxysepala (B. gadgarrense, B.
- 275 grandimesense, B. lamingtonense, B. lewisense, B. schillerianum, B. shepherdii, B. wadsworthii, B.
- 276 windsorense), sect. Polymeres (B. bowkettiae, B. johnsonii, B. radicans, B. wolfei), and sect.

277 Sestochilus (B. baileyi). Australian species from each of these sections were placed in nine different

- 278 positions within the Asian clade. Australian species from section *Polymeres* formed a highly
- supported clade (UFBS 100), while Australian species from sect. *Oxysepala* formed a moderately
- supported clade (UFBS 91) and together with the type species of section Oxysepala from Papuasia
- 281 (B. cladistinum) formed a close relationship with the Australian representative of section Monanthes
- 282 (B. macphersonii) (UFBS 100).



283

Figure 1.

284 Maximum likelihood phylogenetic reconstruction of *Bulbophyllum* based on the supermatrix of 70 285 plastid coding regions in inset a. with *Bulbophyllum* sections *Adelopetalum* and *Minutissima* s.s. in

285 plastid coding regions in inset a. with *Bulbophyllum* sections *Adelopetalum* and *Minutissima* s.s. in 286 detail. Ultrafast bootstrap values are given adjacent to nodes. Australian species are shown with an

asterisk.



288 289

Figure 2. Maximum likelihood phylogenetic reconstruction of Bulbophyllum based on the

290 supermatrix of 70 plastid coding regions in inset a. with Afrotropical and Asian Bulbophyllum clades 291 in detail. Ultrafast bootstrap values are given adjacent to nodes. Australian species are shown with an

292 asterisk.

293 3.1.2 Phylogenetic relationships – Nuclear data

294 The ML phylogeny based on the nuclear ribosomal DNA cistron was resolved with overall lower

- support compared to analyses based on 70 plastid loci supermatrix (Fig. 3, Fig. 4). Relationships
- among outgroup taxa were concordant with the plastid phylogeny and Bulbophyllum was resolved
- 297 with maximum support. Within Bulbophyllum, the Afrotropical (UFBS 97), Asian (UFBS 100) and
- Adelopetalum/Minutissima (UFBS 100) clades were resolved with high to maximum support,
- 299 however the relationships among them were poorly supported. Within the Adelopetalum/Minutissima
- 300 clade, the highly supported clades revealed in the plastid phylogeny were also reconstructed based on
- 301 the nuclear dataset (argyropus clade (UFBS 100), bracteatum clade (UFBS 93), minutissimum clade
- 302 (UFBS 86), and newportii clade (UFBS 83), however relationships among these remained poorly
- 303 supported. Similar to reconstructions based on the plastid phylogeny were the relationships among
- the argyropus, bracteatum and newportii clades, the poor support of *B. pygmaeum* and *B. weinthalii*, the polyphyly or paraphyly for sections *Beccariana*, *Brachyantha*, *Brachypus*, *Cirrhopetaloides*,
- 306 Cirrhopetalum, Desmosanthes, Minutissima, Oxysepala, Polymeres, and Sestochilos; and that
- 307 Australian species from sect. sect. *Brachypus*, sect. *Brachystachyae*, sect. *Cirrhopetalum*, , sect.
- 308 *Ephippium*, sect. *Monanthes*, sect. *Oxysepala*, sect. *Polymeres*, and sect. *Stenochilus* were placed in
- 309 nine clades across the Asian clade.

310



311 312

Figure 3. Maximum likelihood phylogenetic reconstruction of *Bulbophyllum* based on the nuclear

- ribosomal DNA cistron (5'ETS, 18s, ITS1, 5.8s, ITS2, 28s, 3'ETS) in inset a. with *Bulbophyllum*
- 314 sections Adelopetalum and Minutissima in detail. Ultrafast bootstrap values are given adjacent to
- 315 nodes. Australian species are shown with an asterisk.



316

- 317 Figure 4. Maximum likelihood phylogenetic reconstruction of *Bulbophyllum* based on nuclear
- 318 ribosomal DNA cistron (5'ETS, 18s, ITS1, 5.8s, ITS2, 28s, 3'ETS) in inset a. with Afrotropical and
- 319 Asian *Bulbophyllum* clades in detail. Ultrafast bootstrap values are given adjacent to nodes.
- 320 Australian species are shown with an asterisk.

321 **3.2** Divergence time estimation

322 The divergence time analysis based on a relaxed log normal clock and birth death prior with

- 323 speciation and extinction, which was identified as the model of best fit based on the Akaike
- information criterion (Supplementary Material S4), is presented here (Fig. 5) with the Asian and
- Afrotropical clades collapsed and the complete chronogram provided in Supplementary Material S7. The divergence time analysis based on the plastid dataset was well resolved and highly supported
- 327 (Fig. 5, Supplementary Material S7). The divergence between *Bulbophyllum* and *Dendrobium* was
- estimated to have occurred during the early Oligocene, ca. 33.2 Ma (95% highest posterior
- 329 probability density, HPD: 27.7–39.0). The crown of *Bulbophyllum*, constituting the divergence of the
- Adelopetalum/Minutissima clade from the remainder of the genus, was dated to the late Oligocene,
- 331 ca. 24.9 Ma (HPD: 20.1–30.7). Divergence between the Asian clade and the Afrotropical clade was
- estimated to have taken place during the late Oligocene, ca. 24.3 Ma (HPD: 19.4–29.8) and
- 333 diversification within the Asian clade was estimated from the mid Miocene 21.4 Ma (HPD: 17.2–
- 334 26.4 Ma) and the Afrotropical clade from 16.0 Ma (HPD: 10.4–21.7). The crown age of the
- Adelopetalum/Minutissima clade was dated to the late Oligocene, ca. 23.6 Ma (HPD: 18.6–29.1),
- 336 with the split of the Minutissimum clade from the Adelopetalum clade. The crown age of the
- Adelopetalum clade was dated to the mid Miocene, ca. 15.3 Ma (HPD: 10.6–21.2). The stem
- 338 branches of major lineages within the Adelopetalum clade were estimated to have diversified during
- the mid-Miocene: the bracteatum clade was dated to ca. 14.5 Ma (HPD: 10.0–20.4); the lineage
- 340 giving rise to *B. weinthalii* to ca. 12.1Ma (6.9–17.6); the argyropus clade to ca. 12.1 Ma (6.9–17.6);
- and the newportii clade to ca. 14.9 Ma (HPD 10.3–20.7). Diversification among species within these
- 342 lineages took place from the mid-Miocene onwards with the most recent divergence identified during
- 343 the late Pleistocene among *B. argyropus*, *B. corythium*, and *B. tuberculatum*.



344

Figure 5. Maximum clade credibility chronogram for *Bulbophyllum* sect. *Adelopetalum* based on 70
plastid coding sequences, relaxed log normal clock and birth death prior. Divergence dates and 95%
highest posterior density values are indicated adjacent to nodes. Grey bars indicate 95% highest
posterior density. The asterisk denotes the node constrained with a fossil calibration point; the

349 diamond shape denotes nodes which were constrained by secondary calibration points.

350 3.3 Ancestral range analysis

351 Model testing of the three biogeographic models (DEC, DIVALIKE, BAYAREALIKE) using the

352 Akaike information criterion identified the BAYAREALIKE model as the model of best fit for the

- ancestral range estimation (Supplementary Material S5). Ancestral ranges estimated with the
- BAYAREALIKE model are presented here with the Asian and Afrotropical clades collapsed (Fig. 6).
- 355 The complete chronogram is provided in Supplementary Material S8, marginal probabilities for
- ancestral ranges at all nodes in Supplementary Material S9 and node IDs in Supplementary Material
- 357 S10.



358

Figure 6. Range evolution of *Bulbophyllum* sect. *Adelopetalum*. a) ancestral area reconstruction based on a the BAYAREALIKE model with species extant distributions shown within the grid and pie charts at internal nodes representing marginal probabilities for alternative ancestral areas; b) legend of color-coded geographic regions and shared ancestral areas; c) world map of color-coded geographic regions delineated in the biogeographic analysis; d) detail of Australasian color-coded geographic regions.

- 365 Australia was reconstructed as the most likely ancestral range for the MRCA of the Adelopetalum
- clade (range probability (RP) 82) and all nodes within this lineage (RP 72-99) except for the
- 367 argyropus clade (Fig. 6). Range shifts from Australia were inferred from the early Pliocene to the
- 368 Pacific region (New Caledonia) in the newportii clade, in the lineage giving rise to *B. lingulatum*.
- 369 Range shifts were also inferred from Australia to the Pacific region (New Caledonia) and New
- 370 Zealand either in the lineage giving rise to the MRCA of the argyropus clade or subsequently within
- this lineage. Three alternative ancestral ranges were reconstructed for the MRCA of the argyropus
- 372 clade: Australia (RP 36), or widespread distributions including Australia and New Zealand (RP 33)
- 373 or Australia and New Caledonia (RP 26). Two alternative ranges were also reconstructed for the

- 374 MRCA of *B. corythium* and *B. tuberculatum*: New Zealand (RP 41) and New Caledonia (RP 34).
- 375 Considering these alternative scenarios, range shifts within the argyropus clade were estimated to
- have occurred sometime between the mid Miocene and late Pliocene (12.1–0.5 Ma). The ancestral
- 377 range of MRCA of the Adelopetalum/Minutissima clade and Bulbophyllum remained unresolved in
- the ancestral range reconstruction. The most likely ancestral range for the MRCA of the
- 379 Adeloptalum/Minutissima clade was a widespread distribution across Australia and tropical Asia (RP
- 380 26), while alternative ranges reconstructed included a widespread range including Australia, tropical
- 381 Asia and Papuasia (RP 13) and Australia (RP 11). Two alternative ancestral ranges were
- 382 reconstructed for the MRCA of Bulbophyllum, both widespread distributions including Australia and
- 383 tropical Asia (RP 25.9) or Australia, tropical Asia and Papuasia (RP 21).

384 4 Discussion

385 4.1 Phylogenetic relationships

386 This study provided a broad plastid phylogenetic framework for Asian and Australasian sections of

- 387 Bulbophyllum and revealed a close relationship between sections Adelopetalum and Minutissima s.s.,
- that together form a highly supported early diverging lineage within the genus (Fig. 1, Fig. 2).
- 389 Relationships based on 70 plastid genes support a sister group relationship between the
- 390 *Adelopetalum/Minutissima* clade and the remainder of the genus (Asian + Afrotropical clades).
- 391 Within the Adelopetalum/Minutissima clade, analyses based on our 70 plastid loci supermatrix
- 392 showed a dichotomous split between the highly supported *Minutissima s.s.* and Adelopetalum clades.
- 393 Species were reconstructed in each of these clades according to their sectional placement except for
- 394 New Zealand endemic *B. pygmaeum* (sect. *Minutissima*), which was nested within the Adelopetalum
- 395 clade, rendering section *Adelopetalum* paraphyletic. Section *Minutissima* was identified as
- 396 polyphyletic, with the Australian (B. minutissimum (sect. type), B. globuliforme) and Pacific species
- 397 (B. keekee) placed in the Minutissima clade and New Zealand species (B. pygmaeum) in the
- 398 Adelopetalum clade while the Asian species, *B. mucronatum* and *B. moniliforme* were resolved
- 399 within the Asian clade. Section *Minutissima* has undergone numerous taxonomic changes with
- 400 treatments ranging from a narrower circumscription recognising species from Australia (Jones and
- Clements, 2001), to broader classifications including 23 species from Thailand, Indonesia, Australia,
 New Zealand, New Caledonia, and New Guinea (Pridgeon et al. 2014). Our phylogenetic analysis
- 403 based on plastid and nuclear markers did not reconstruct a close relationship between sect.
- 404 *Minutissima* species from the Australasian/Pacific region and Asian species *B. mucronatum* and *B.*
- 405 moniliforme. Rather, in our analyses the Australasian species fell within the
- 406 Adelopetalum/Minutissima clade while the Asian species were nested within the Asian clade. The
- 407 results support morphological studies differentiating sect. *Minutissima* species from Australasia and
- 408 Asia (Jones and Clements, 2001) and show minute pseudobulbs are a trait that has evolved more than
- 409 once independently in the genus.
- 410 Within section *Adelopetalum*, phylogenetic analyses supported current species concepts, except for
- 411 species within the argyropus clade, which exhibited shallow genetic differentiation (Fig. 1, Fig. 3).
- 412 The species of the argyropus clade share morphological affinities and previous taxonomic treatments
- 413 recognised up to three species within the group: *B. argyropus* (Australia's east coast and off shore
- 414 islands: Lord Howe Island, and Norfolk Island), *B. corythium* (New Caledonia), and *B. tuberculatum*
- 415 (New Zealand) (Clements and Jones, 2002; Halle, 1981; Vermeulen, 1993). Divergence dating
- analyses shows this group represents a relatively recent radiation reconstructing divergence among
- 417 species during the Pleistocene. Further studies are required to clarify species delimitation and
- 418 dispersal patterns utilising population-level sampling and genomic techniques suited to resolving

419 relationships among recently diverged lineages, such as reduced representation high-throughput

420 approaches like ddRAD, DArT or target sequence capture methods (Peterson et al., 2012; Sansaloni

- 421 et al., 2011; Weitemier et al., 2014; Folk et al., 2015; Bagley et al. 2020, Schmidt-Lebuhn et al.,
- 422 2022).
- 423 Previous cladistic analysis based on morphological traits in sect. Adelopetalum found two main
- 424 clades within the section, one comprising *B. lageniforme, B. lilianae, B. lingulatum,* and *B.*
- 425 *newportii*, and the other uniting *B. argyropus*, *B. bracteatum*, and *B. elisae* (Vermeulen 1993).
- 426 Phylogenetic relationships based on plastid and nuclear markers found strong to moderate support for
- 427 the first clade recovered in the cladistic analysis, corresponding to the newportii clade in the present
- 428 analyses (Fig 1, Fig. 3). The second group found in the cladistics analysis included three species
- 429 placed in phylogenetic analyses within either the bracteatum clade (<u>B. bracteatum, B. elisae</u>) or the 430 argvropus clade (<u>B. argvropus</u>). However, relationships among these two lineages remained unclear
- 430 argyropus clade (*B. argyropus*). However, relationships among these two lineages remained unclear
 431 due to low support. The sister group relationship between *B. bracteatum* and *B. argyropus* recovered
- 432 in the cladistic analysis was not supported in phylogenetic reconstructions based on molecular data,
- 433 suggesting character states shared by these species may be homoplasious. Further studies using
- 434 ancestral character reconstruction are required to test the phylogenetic utility of morphological traits
- 435 utilised in prior studies.
- 436 While plastid phylogenomics has clarified major clades and intraspecific relationships within sect.
- 437 Adelopetalum and broad-level relationships within Bulbophyllum, further studies are required. Non-
- 438 monophyletic sections identified in the present study (e.g., sections *Beccariana, Brachyantha*,
- 439 Brachypus, Cirrhopetaloides, Cirrhopetalum, Desmosanthes, Minutissima and Polymeres) (Fig. 2)
- 440 and in previous molecular phylogenetic studies (Fischer et al., 2007; Smidt et al., 2011; Pridgeon et
- 441 al. 2014; Hu 2020) highlight the need for further taxonomic revision within *Bulbophyllum*. Studies
- 442 are required with an expanded sampling of the diverse Asian and Pacific taxa to increase our
- 443 understanding of evolutionary relationships and assess sectional classification in more detail.
- Phylogenetic relationships reconstructed from the nuclear ribosomal DNA cistron were not strongly
 supported in line with previous molecular studies based in ITS (Gamisch et al., 2015; Gamisch and
- 446 Comes, 2019, Hu et al., 2020). Approaches yielding higher number of nuclear markers such as target
- sequence capture, provide an opportunity to improve the understanding of evolutionary relationships
- 448 in future studies. While assembling datasets with comprehensive species coverage within mega
- 449 diverse groups such as *Bulbophyllum* remains a challenge, the present study provides an example of
- 450 the use of a broad phylogenetic framework with targeted sampling within a section, to test the
- 451 monophyly and phylogenetic placement of groups of interest.

452 **4.2** Spatio-temporal evolution of *Bulbophyllum* sect. *Adelopetalum*

- 453 Our divergence time analysis and ancestral range estimations showed that *Bulbophyllum* sect.
- 454 Adelopetalum represents an Australasian lineage that originated on the Australian continent during
- the late Oligocene to early Miocene (Fig. 5, Fig. 6). The Australian ancestral range is largely
- 456 conserved within the lineage, indicating diversification among species has predominantly taken place
- 457 on the Australian continent. The conservation of ancestral range observed within sect. *Adelopetalum*
- 458 is consistent with previous phylogenetic analyses of *Bulbophyllum* that have shown a strong
- 459 biogeographic signal among clades, being largely confined to biogeographic regions such as
- 460 Madagascar, continental Africa and South America (Fischer et al., 2007; Gamisch et al., 2015;
- 461 Gamisch and Comes, 2019; Smidt et al., 2011). The evolution of *Bulbophyllum* during the early
- 462 Oligocene occurred subsequently to the breakup of Gondwana (Matthews, et al. 2016; Zahirovic et
- al. 2016), implicating long-distance dispersal (LDD) in the evolution of biogeographical lineages

464 within the genus (Van den Berg, 2003, Smidt et al., 2011; Gamisch et al., 2015; Gamisch and Comes, 465 2019). Nevertheless, the conservation of ancestral ranges observed within the Adelopetalum lineage 466 in this study and strong biogeographic signal among clades identified in previous studies indicate 467 LDD with successful establishment and persistence has been relatively infrequent within 468 Bulbophyllum. Although the minute wind-dispersed seeds of orchids have a high dispersal potential, 469 successful establishment in a new area are limited by several factors, such as the presence of 470 mycorrhizal partners necessary for germination and development, a suitable host or substrate and 471 microclimatic conditions, and the availability of pollinators (Arditti and Ghani, 2000; Jersáková and 472 Malinová, 2007; McCormick et al., 2012). Our results are consistent with previous studies that have 473 identified in situ diversification as the dominant biogeographic process, despite evidence for LDD, 474 and provide further support for the hypothesis that the complex requirements for successful

475 establishment, rather than dispersal limitations, play an important role in constraining the geographic

476 distribution of orchids (Givnish et al., 2016; Perez-Escobar and Chomicki et al. 2017).

477

478 Our phylogenetic analysis further resolved interspecific relationships in sect. *Adelopetalum* (Fig. 1).

479 Divergence time estimation showed that divergence among species occurred mainly during the

480 Miocene and Pliocene (Fig. 5), during a period of extensive changes to the distribution of forest

481 vegetation on the Australian continent in response to drastic climatic changes. During the early

482 Miocene, Australian vegetation diversified in response to aridification of the Australian continent and

the abrupt shift to a cool dry climate during the mid-Miocene resulted in considerable fragmentation

484 of rainforest habitats (Martin, 2006, Byrne et al. 2011). *Bulbophyllum* sect. *Adelopetalum* comprises

485 epiphytic species that occur in mesic forest habitats and thus diversification and fragmentation of 486 these habitats were likely drivers of allopatric lineage divergence within this group. Sister group

487 relationships were identified between two species pairs with disjunct distributions in Australia's

488 northern wet tropical rainforests and south-eastern rainforests (*B. booniee/B. bracteatum* and *B.*

489 *newportii/B. exiguum*). These relationships support the hypothesis that the diversification and

490 fragmentation of forest habitats in Australia has been an important driver of lineage divergence in

491 Australia's mesic biome (Byrne et al., 2011, Simpson et al., 2018).

492 Whilst the ancestral range was predominantly conserved within the *Adelopetalum* lineage, range

493 expansion events were inferred from continental Australia across the Coral and Tasman Seas to New

494 Caledonia in the lineage giving rise to *B. lingulatum*, and to New Zealand and New Caledonia in the

495 argyropus clade (Fig. 6). New Caledonia and New Zealand each have a long history of isolation from

496 Australia that predates the evolution of *Bulbophyllum*, indicating colonisation of these islands by

497 Bulbophyllum species has been via LDD (Matthews et al., 2016). It remains unclear if LDD to New

498 Zealand and New Caledonia in the argyropus clade occurred from the early Miocene in the lineage

499 giving rise to the MRCA of the group or subsequently within this clade during the late Pleistocene,

500 thus the spatio temporal evolution of this lineage requires further study.

501 The pattern of eastward dispersal observed in range shifts from Australia, across the Coral and

502 Tasman Seas, is consistent with dispersal patterns inferred in other angiosperms, including

503 Abrotanella, Dendrobium, Dracophyllum, Hebe, Korthalsella, Leucopogon, Northofagus, Oreobolus,

504 Pterostylis, Rytidosperma (Chacón et al., 2006; Lavarack et al., 2000; Linder, 1999; Molvray et al.,

505 1999; Swenson et al., 2001; Puente-Lelièvre et al., 2013; Wagstaff et al., 2010, 2006, 2002, Nargar et

- al. 2022). The bias towards eastward dispersal observed within section Adelopetalum among other
- 507 plant groups may be facilitated by the predominant westerly winds occurring in the southern

hemisphere that initiated after the rifting of Australia and South America from Antarctica during theEocene (Sanmartín et al., 2007).

510 4.3 Conclusions

- 511 This study provided an important phylogenomic framework for the mega genus Bulbophyllum
- 512 facilitating studies into trait and range evolution within the genus. Several Asian sections were
- 513 resolved as paraphyletic warranting taxonomic revisions. Our plastid phylogenomic analyses
- 514 revealed an early-diverging lineage within *Bulbophyllum*, composed of sect. *Adelopetalum* and sect.
- 515 Minutissima s.s.. For Bulbophyllum sect. Adelopetalum, this study reconstructed an origin in the early
- 516 Oligocene and identified the Australian continent as ancestral range. Species diversification within
- 517 the section occurred predominantly on the Australian continent with fragmentation of mesic habitats
- 518 during the Miocene identified as likely drivers of allopatric lineage divergence. Multiple independent
- 519 long distance dispersal events were inferred from the Australian continent eastward to the islands of
- 520 New Zealand and New Caledonia.

521 5 Conflict of Interest

522 The authors declare that the research was conducted in the absence of any commercial or financial 523 relationships that could be construed as a potential conflict of interest.

524 6 Author Contributions

525 Conceptualization: LS, KN, MAC; Data curation: LS, MAC; Formal Analysis: LS, HKO; Funding

- 526 acquisition: LS, KN, DMC, MAC; Investigation and Methodology: LS; Supervision: KN, DCM,
- 527 MAC; Writing original draft: LS; Writing review and editing: KN, HKO, DCM, MAC. All
- 528 authors approved of the final version of the manuscript.

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