# Incomplete lineage sorting and hybridization underlie tree discordance in *Petunia* and related genera (Petunieae, Solanaceae)

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11 Abstract

12 Despite the overarching history of species divergence, phylogenetic studies often reveal distinct

13 topologies across regions of the genome. The sources of these gene tree discordances are

14 variable, but incomplete lineage sorting (ILS) and hybridization are among those with the most

15 biological importance. *Petunia* serves as a classic system for studying hybridization in the wild.

16 While field studies suggest that hybridization is frequent, the extent of reticulation within

17 *Petunia* and its closely related genera has never been examined from a phylogenetic perspective.

18 In this study, we used transcriptomic data from 11 *Petunia*, 16 *Calibrachoa*, and 10 *Fabiana* 

19 species to illuminate the relationships between these species and investigate whether

20 hybridization played a significant role in the diversification of the clade. We inferred that gene

tree discordance within genera is linked to hybridization events along with high levels of ILS due

to their rapid diversification. Moreover, network analyses estimated deeper hybridization events

between *Petunia* and *Calibrachoa*, genera that have different chromosome numbers. Although

these genera cannot hybridize at the present time, ancestral hybridization could have played a

role in their parallel radiations, as they share the same habitat and life history.

## 2627 Introduction

28 Over the last two decades, systematic biology has experienced significant advances, managing to

29 work with vast volumes of data and constructing robust phylogenies to elucidate species'

30 relationships and evolutionary histories. Nevertheless, using different methods and datasets (i.e.,

31 genetic markers and sampling schemes) often results in conflicting tree topologies. These

32 discrepancies may stem from errors in model specifications, data processing, or evolutionary

processes such as incomplete lineage sorting (ILS) and hybridization (Galtier and Daubin, 2008).
 Coalescent-based methods are commonly employed to mitigate conflicts in trees caused by ILS,

such as anomaly zones, where the topology of 'anomalous gene trees' with short branch lengths

differs from the species tree topology (Degnan and Rosenberg, 2006). However, these methods

are unreliable in situations involving gene flow among lineages (Solís-Lemus et al., 2016).

38 Despite the advances in phylogenomic methods that account for ILS and gene flow (Hibbins and

Hahn, 2022), detecting and distinguishing between such events remains a complex task that

40 heavily depends on the extent to which they occur (Kong and Kubatko, 2021).

41 Botanists already recognize that plant evolution likely follows a web-like pattern due to

42 the numerous examples of plant hybridization (Stull et al., 2023). However, the potential

43 outcomes of such events are highly variable (Abbott et al., 2016; Soltis and Soltis, 2009).

Hybridization can facilitate speciation through novel trait combinations or polyploidization
(Abbott et al., 2013), lead to extinction through genetic swamping (Todesco et al., 2016), or

46 introgress adaptive alleles (Suarez-Gonzalez et al., 2018). Regardless of the outcomes,

47 hybridization is a frequent evolutionary phenomenon at both shallow (Nevado et al., 2018) and

48 deep timescales (Rothfels et al., 2015) with extensive impacts on plant diversification and

49 evolution (Goulet et al., 2017; Whitney et al., 2010).

50 The *Petunia-Calibrachoa-Fabiana* Solanaceae clade presents a unique opportunity to investigate the evolutionary dynamics of plant diversification in southern South America. 51 Notably, Petunia has become an important taxon to understand the role of hybridization in the 52 region due to the multiple hybrid zones documented (e.g., Binaghi et al., 2023; Caballero-53 Villalobos et al., 2021; Giudicelli et al., 2019). Lineages in the Petunia-Calibrachoa-Fabiana 54 clade have experienced rapid diversification over the past ca. 20 million years (Zuntini et al., 55 2024), with approximately 65 species exhibiting a diverse range of pollination syndromes and 56 inhabiting distinct biomes, from rainforests to savannahs and deserts (e.g., Alaria et al., 2022, 57 Mäder and Freitas, 2019; Reck-Kortmann et al., 2014). While Petunia and Calibrachoa are very 58 similar in morphology and ecological conditions, Fabiana stands out due to its drastic 59 differences from related genera: its species are xerophytic, have small flowers, and display 60 reduced or even absent leaves (Alaria et al., 2022). The three genera are classified within the 61 well-established tribe Petunieae, forming a strongly supported clade (Särkinen et al., 2013; 62 63 Wheeler et al., 2022). However, the internal relationships within this clade remain controversial. Phylogenetic studies using distinct genetic markers, sampling schemes, and analytical 64 approaches have produced conflicting topologies (Fig. 1). Some place Petunia as a sister to the 65 remaining genera (Alaria et al., 2022; Olmstead et al., 2008; Wheeler et al., 2022), whereas 66 others place it as a sister genus to Calibrachoa (Särkinen et al., 2013) or to Fabiana (Mäder and 67 Freitas, 2019; Reck-Kortmann et al., 2015). These incongruences might be attributed to the 68 69 substantial levels of ILS due to their rapid diversification (Särkinen et al., 2013; Wheeler et al., 2022), or potentially from instances of ancient hybridization. 70 The likelihood of hybridization depends on how effective reproductive barriers are at 71

72 preventing gene flow, and plants typically rely on a combination of barriers to achieve complete

reproductive isolation (Baack et al., 2015; Christie et al., 2022). In *Petunia*, gene flow is
 primarily prevented by prezygotic barriers, including geographic and floral isolation, with

primarry prevented by prezygotic barriers, including geographic and horal isolation, with
 postzygotic barriers playing a negligible role (Dell'Olivo et al., 2011). These barriers have been

result for the stabilished this genus as a model in plant hybridization

and pollination studies (Binaghi et al., 2023; Gübitz et al., 2009; Rodrigues et al., 2018;

78 Turchetto et al., 2019). However, a comprehensive investigation into *Calibrachoa* and *Fabiana*,

as well as the possibility of hybridization causing tree discordance among genera and congeneric

species has yet to be examined under a solid phylogenetic framework.



**Figure 1**. Distribution map of *Petunia* (blue outline), *Calibrachoa* (pink shaded area), and *Fabiana* (green shaded area) in South America. Map dots represent sampled localities, colored by genus, except for the species collected at greenhouses. Representatives of each genus are shown in circles: *Petunia altiplana*, *Calibrachoa eglandulata*, and *Fabiana bryoides* (photos: Lucas C. Wheeler). Phylogenetic relationships of the genera are presented on the right. Olmstead et al., (2008) used two plastid markers; Särkinen et al., (2013) used two nuclear and five plastid markers; Reck-Kortmann et al., (2015) used one nuclear and two plastid markers; Mäder et al., (2019) used eight nuclear and eight plastid markers; and Alaria et al., (2022) used one nuclear and three plastid markers.

- 81 Even though *Petunia* and *Calibrachoa* are similar in flower morphology, ecology, and 82 geographic distribution (Fig. 1), they have been split into two different genera due to their 83 chromosome numbers: *Petunia* has seven chromosome pairs (2n = 14), whereas *Calibrachoa* has
- nine (2n = 18) (Stehmann et al., 2009; Wijsman and De Jong, 1985). The persistence in nature of
- hybrids between species that have different chromosome numbers is unlikely as it leads to
- 86 meiotic mispairing—unless it involves polyploidization (Alix et al., 2017; Hegarty and Hiscock,

2008). To date, polyploidization has never been observed in *Petunia* or *Calibrachoa*. Hence, the 87

- occurrence of hybrids between Petunia and Calibrachoa in the wild seems unlikely, even though 88
- some species occur in sympatry. While intergeneric hybrids known as "Petchoa" have been 89
- 90 developed and are available commercially, these hybrids are sterile, and their creation requires
- significant human intervention (Shaw, 2007). In contrast, while Calibrachoa and Fabiana share 91
- the same chromosome count, which theoretically would allow successful meiosis in the hybrid, 92 their disjunct geographical distribution and distinct life histories serve as strong present-day
- 93
- 94 barriers that prevent gene flow.

In this study, we used floral transcriptome data from *Petunia*, *Calibrachoa*, and *Fabiana* 95 species to investigate the sources of discordance among phylogenetic trees. Specifically, we 96 aimed to evaluate the influence of ILS and reticulate evolution on the diversification of these 97 genera. We hypothesized that hybridization occurs frequently within genera, both in recent times 98 and throughout their evolutionary history, contributing to the observed phylogenetic discordance 99 within genera. Moreover, we tested whether intergeneric hybridization could have played a role 100 in the diversification of the clade. We predicted that intergeneric hybridization is unlikely due to 101

- robust reproductive barriers, including chromosome number differences and geographic 102
- 103 isolation.

#### 104

#### 105 **Material and Methods**

- 106 Taxa sampling and transcriptome data processing
- We used the raw RNA-seq data from Wheeler et al., (2022, 2023), focusing on the Petunia-107
- Calibrachoa-Fabiana clade and incorporating six outgroup species. In total, we employed 107 108
- 109 individuals, encompassing 11 Petunia species, 16 Calibrachoa species, 10 Fabiana species, and
- six outgroups (Table S1). Here, we expanded the Wheeler et al., (2022) dataset by including 110
- additional individuals for most sampled species derived from Wheeler et al., (2023), resulting in 111
- 112 three individuals per species collected at the same time and location (hereafter referred to as
- replicates). We corrected the raw RNA-seq reads using Rcorrector (Song and Florea, 2015) and 113
- removed adapters using Trimmomatic (Bolger et al., 2014). Subsequently, we mapped the reads 114 against the 3,672 protein-coding genes from conspecific transcriptomes, which were assembled 115
- for replicate 1 by Wheeler et al., (2022) using BWA (Li and Durbin, 2010). Consensus fasta 116
- sequences were generated through samtools 1.16 (Li et al., 2009) by calling the most frequent 117
- base (-m simple) and then aligned with MACSE 2.06 (Ranwez et al., 2018). 118
- 119
- 120 *Phylogenetic analyses and evaluation of tree discordance*
- We employed three distinct approaches to elucidate the phylogenetic relationships among species 121
- within the Petunia-Calibrachoa-Fabiana clade. Firstly, we estimated the maximum likelihood 122
- (ML) gene trees using the GTR+ $\Gamma$  model along with 1,000 bootstrap replicates in RAxML 123
- (Stamatakis, 2014) and estimated the species tree—both with and without assigning individuals 124
- to species—using ASTRAL III 5.7.8 (Rabiee et al., 2019; Zhang et al., 2018). Secondly, we 125
- constructed a supermatrix by concatenating the fasta alignments with the SuperMatrix function 126
- of the evobiR R-package (Jonika et al., 2023). This supermatrix was then used to generate a 127
- maximum likelihood species tree using IQTree 1.6.12 (Nguyen et al., 2015) setting the GTR+ $\Gamma$ 128
- model to each partition with 1,000 bootstrap replicates. Lastly, we estimated a species tree using 129
- SVDQuartets, a coalescent method originally designed for SNP data but also effective with 130
- multi-locus alignments (Chifman and Kubatko, 2014), implemented in PAUP\* 4a (Swofford, 131
- 2003), which infers relationships among quartets and subsequently summarizes these 132

- relationships into a species tree. We set the outgroups, assigned individuals to respective species, and assessed all quartets (evalq=all) using 200 multi-locus bootstrap replicates.
- We used phyparts (Smith et al., 2015) to evaluate the number of concordant and conflicting bipartitions among gene trees in comparison to the inferred ASTRAL species tree
- setting support level of at least 50% for the corresponding node (-*s* 0.5). Due to computational
- 138 limitations and the observed clustering of conspecific individuals (see Results), we pruned
- replicates 2 and 3 for all species using Newick utilities (Junier and Zdobnov, 2010). Since
- 140 phyparts requires rooted trees as input, we set *Bouchetia erecta* as the root, which led to a dataset
- 141 of 3,471 gene trees where the outgroup was present. For the ML phylogenetic tree, we evaluated
- 142 genealogical concordance with gene concordance factor (gCF) and site concordance factor (sCF)
- 143 with 100 randomly sampled quartets (*-scf*), where gCF measures how often a specific branch in 144 the species tree is supported by "decisive" gene trees, while sCF measures the percentage of sites
- the species tree is supported by "decisive" gene trees, while sCF measures the percethat support a branch in the tree (Minh et al., 2020).
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#### 147 *Detection of hybridization*

- 148 We used HyDe (Blischak et al., 2018) to search for hybridization signals. HyDe relies on
- 149 phylogenetic invariants to estimate admixture ( $\gamma$ ), where a  $\gamma$  value of 0.5 signifies an equal
- 150 genetic contribution from each parental species, and values approaching 0 or 1 indicate a greater
- 151 genetic contribution of one of the parental species. We used a concatenated matrix of alignments
- and trimmed sites with trimAl (Capella-Gutiérrez et al., 2009) with options -gt 0.5 (fraction of
- 153 gaps allowed) and *-cons* 60 (minimum percentage of positions to keep from the original
- alignment), which yielded 5,209,834 sites. We assigned individuals to species and set the six
- outgroup species as outgroups, which resulted in an evaluation of 23,310 triplets. As a second
- approach, we employed QuIBL (Edelman et al., 2019) which relies on branch lengths of gene
- trees to assess whether hybridization provides a more plausible explanation for the divergence
- patterns compared to ILS alone. Because QuIBL requires that all taxa be present in every gene
- tree, we created a dataset with no missing loci for all ingroup species and *B. erecta*. All trees
  were rooted in *B. erecta* and pruned to contain only one individual of each species with Newick
- 161 Utilities, which resulted in a final dataset of 826 gene trees.
- 162
- 163 *Reticulate evolution and network reconstruction*
- 164 Considering the possibility of a non-bifurcating evolutionary history of the *Petunia-Calibrachoa*-
- 165 *Fabiana*, we inferred phylogenetic networks that account for both ILS and gene flow among
- 166 taxa. Due to computational limitations, we constructed a reduced dataset comprising 18 ingroup
- taxa and a single outgroup species (Table S1), not allowing for missing loci, which resulted in a
- 168 dataset of 1,215 loci. We estimated gene trees with RAxML and the species tree with ASTRAL 169 as described in the previous section.
- 170 As our first approach, we estimated a phylogenetic network with the maximum
- pseudolikelihood method SNaQ implemented in the Julia package PhyloNetworks 0.16.2 (Solís-
- 172 Lemus et al., 2017; Solís-Lemus and Ané, 2016). We searched for up to five hybridization events
- 173 (h = 5) and used the ASTRAL phylogeny as the starting tree. For the following steps, we used
- the network from the previous estimation as the starting network. The best number of
- hybridization events was selected based on where we could detect a steep log-pseudolikelihood
   improvement. After selecting the best number of hybridization events, we ran 100 bootstrap
- replicates using the 1 000 bootstrap ML gaps trees inferred by DAVML for each of the 1 215 losi
- replicates using the 1,000 bootstrap ML gene trees inferred by RAxML for each of the 1,215 loci,
- 178 employing default settings.

As a second approach, we estimated a network with the command "InferNetwork MPL" 179 180 in PhyloNet 3.8.2 (Than et al., 2008), also searching for up to five hybridization events and 10 runs for each search. To select the best-scored network, we used the "CalGTProb" function in 181 182 PhyloNet (Yu et al., 2012) to get network likelihoods. We compared the networks with model selection using the Akaike information criterion (AIC; Akaike 1973), the bias-corrected Akaike 183 information criterion (AICc; Sugiura 1978), and the Bayesian information criterion (BIC; 184 Schwarz 1978). We set the number of parameters to the number of estimated branch lengths and 185 hybridization probabilities, correcting for finite sample size with the number of gene trees used. 186

Considering the potential occurrence of intergeneric hybridization (see Results), we used 187 Twisst (Martin and Van Belleghem, 2017) on the reduced dataset of 19 species and 1,215 loci. 188 We categorized species according to their respective genera and designated *B. erecta* as the 189 outgroup, resulting in three potential topologies. We computed the topology weight and 190 determined the frequency of specific topologies within the gene tree set, that is, we counted the 191 number of trees supporting one of the three possible topologies. Subsequently, we conducted a 192 chi-square test to compare the occurrences of the two minor topologies (Owens et al., 2023; 193 Suvorov et al., 2022). Under the null hypothesis, i.e., without intergeneric hybridization, we

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195 expect the two minor topologies to occur with similar frequency (Baum, 2007). 196

#### Results 197

198 *Phylogenetic relationships and tree discordance within the Petunia-Calibrachoa-Fabiana clade* 

The gene count for each replicate ranged from 2,937 to 3,573 (Table S1), and the final 199

concatenated matrix consisted of 5,687.285 base pairs. The resulting phylogenetic trees 200 201 constructed using multiple methods consistently positioned Petunia as sister to Calibrachoa +

Fabiana (Olmstead et al., 2008; Alaria et al., 2022; Fig. 1), while revealing discordant 202

intrageneric topologies. Both the supertree (ASTRAL) and the supermatrix strategies (IQTree) 203

exhibited strong support for most branches (LPP = 0.95-1 in ASTRAL and bootstrap = 100 in 204

IOTree; Fig. 2). Nonetheless, the two methods estimated different relationships for multiple 205

branches within *Petunia* and within *Calibrachoa*, which might be expected given the high 206 207

proportion of conflict among gene trees apparent from the phyparts analysis (Fig. 2). We did, however, find that the replicates from a single species consistently group together in the 208

- ASTRAL analysis with robust support (Fig. S1), supporting assignment of individuals to species. 209
- As might be expected given the differences between the ASTRAL and supermatrix trees (Fig. 2), 210
- 211 SVDQuartets displayed high support for deeper nodes, but weaker support for shallower nodes
- within Calibrachoa and Petunia (Fig. S2), indicating extensive ILS and possibly intrageneric 212
- hybridization. IQTree Concordance Factor results indicated that the gCF values were notably low 213
- for shallow nodes, whereas sCF values offered greater support for these relationships than gCF, 214
- suggesting that genetic sites were more consistent in inferring evolutionary relationships at these 215
- shallower nodes than the genes themselves (Fig. 2). 216



Figure 2. Phylogenetic trees of the *Petunia-Calibrachoa-Fabiana* clade inferred from ASTRAL (A) and IQTree (B). The subgenera of *Calibrachoa* are annotated on their branches on ASTRAL

tree: Ca for *Calibrachoa*, and St for *Stimomphis*. All nodes are strongly supported (local

221 posterior probability=1 for ASTRAL/boostrap=100 for maximum likelihood tree), except when 222 otherwise noted by an asterisk (\*). Dashed lines represent species with differing positions in the

two trees. Pie charts on the ASTRAL tree depict gene support based on 3,471 gene trees:

corroborating (green), conflicting (orange), or uninformative (gray; < 50% bootstrap scores or

225 missing loci) relationships. The piecharts and numbers above the branches on the maximum

226 likelihood tree inferred from IQTree indicate gene and site Concordance Factors (gCF and sCF,

227 respectively). Outgroup species are not shown for simplicity.

228

#### 229 Reticulate evolution and species networks

230 The search for hybrids resulted in several significant hybrid triplets, both in QuIBL and HyDe

231 (Fig. 3). HyDe resulted in 3,352 significant triplets, even between intergeneric species (Table

S2). However, when we only considered events with  $0.2 < \gamma < 0.8$  (to detect more recent

hybridization events, where we can detect greater parental contribution from both species, and

discard spurious results with low contribution from either parent), these hybridization events

were constrained within genera (Fig. 3). QuIBL showed several minor topologies that could not

be explained by ILS alone, although the percentage of discordant loci explained by introgression

were lower than 10% in all cases (Fig. S3; Table S3).



Figure 3. Detected hybridization events using QuIBL (top) and HyDe (bottom), using ASTRAL phylogeny as the reference species tree. The HyDe graph displays hybridization events with 0.2  $<\gamma < 0.8$ ; refer to Table S2 for complete results. QuIBL identified introgression events based on branch lengths in 826 gene trees with single individuals from each taxon; refer to Table S3 for complete results.

Although network inferences yielded different optimal numbers of reticulations (one in 244 SNaQ, two in PhyloNet; Fig. 4), both agreed on an ancient hybridization edge from Petunia to 245 Calibrachoa subgenus Stimomphis. However, the inheritance probabilities for this introgression 246 were low in both analyses (less than 1% in SNaQ and 3.4% in PhyloNet). The bootstrap analyses 247 for SNaQ showed high support for the species network nodes, but low support for the hybrid 248 249 edge. The placement of minor and major edges was not consistent, with low consistency for both the origin and the source of hybridization. The addition of more hybridization events in SNaQ 250 usually led to the impossibility of rooting the tree in the outgroup (supplemental material online), 251 which suggests incorrect placement of that hybridization edge. 252



Figure 4. Inferred species networks using SNaQ (left) and PhyloNet (right) based on a reduced 254 255 dataset comprising 18 ingroup species and 1,125 genes, rooted in *B. erecta*. The SNaO tree identified the optimal network with one hybridization event. All branches received 100% 256 bootstrap support from 100 replicates, except those signed with gray circles. The dashed line 257 represents the minor edge, displaying the inheritance probability of the best network. The 258 bootstrap values for minor (origin) and major (source) edges of alternative networks are colored 259 green and orange, respectively. The PhyloNet network indicated the optimal network with two 260 hybridization events. The dashed lines also indicate the minor edges with corresponding 261 inheritance probabilities. 262

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The model selection for PhyloNet revealed the network with two hybridization edges as the optimal network and highlighted that any species network is better fitting than the bifurcating species tree (Table 1). In addition to the intergeneric hybridization edge, PhyloNet also suggested a hybridization within *Fabiana* as a second hybridization event. In this case, it showed a high inheritance probability of 0.46 from *F. patagonica* to *F. australis*.

269

270 Table 1. Network likelihoods derived from the reduced dataset using PhyloNet. The number of

271 parameters (k) represents the number of estimated branch lengths and admixture probabilities.

Topology	Maximum number of reticulations	Number of inferred reticulations	Total log probability	lnL	k	AIC	ΔΑΙϹ	AICc	BIC
Astral	0	NA	-	-30833.49	37	61740.98	4676.06	61743.37	61781.11
Network 0	0	NA	-389607.21	-29310.04	37	58694.08	1629.16	58620.08	58734.21
Network 1	1	1	-388824.25	-28520.99	39	57119.98	55.06	57041.98	57162.28
Network 2	2	2	-388777.78	-28491.46	41	57064.92	0.00	56982.92	57109.39
Network 3	3	3	-388791.27	-28511.48	43	57108.96	44.04	57022.96	57155.60
Network 4	4	3	-388805.06	-28524.33	43	57138.66	73.74	57048.66	57181.30
Network 5	5	3	-388827.51	-28543.19	43	57180.38	115.46	57086.38	57219.02

272 The optimal network is in bold.

Results from Twisst revealed that, between the two minor topologies, the topology
positioning *Fabiana* as the sister group to *Petunia* and *Calibrachoa* exhibited a marginally
greater frequency (Fig. 5), although this difference was not statistically significant (chi-square =
1.15, *P* value = 0.28). Thus, the difference in the number of topologies of gene trees is primarily
attributed to ILS and not gene flow.



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Figure 5. Total count of topologies by Twisst for the subset of 1,215 gene trees. The *P* value
indicates the lack of significance for the chi-square test between the two minor topologies that
places *Calibrachoa* or *Fabiana* as the outgroup.

282

#### 283 Discussion

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285 Extensive ILS and ancient hybridization are the sources of tree discordance in the Petunia-

- 286 *Calibrachoa-Fabiana clade*
- 287 Historically, phylogenies based on Sanger sequences often yielded conflicting species trees
- among *Petunia*, *Calibrachoa*, and *Fabiana*. However, our results consistently placed *Petunia* as
- a sister group to *Calibrachoa* + *Fabiana*, mirroring previous findings by Olmstead et al., (2008),
- Alaria et al., (2022), and Wheeler et al., (2022). These results are surprising due to the
- 291 morphological and ecological similarities between *Petunia* and *Calibrachoa* and suggest that
- bee-pollinated herbs (such as most extant *Calibrachoa* and *Petunia* species) represent the
- ancestral state with the extreme xerophyte traits found in *Fabiana* (tiny flowers and reduced
- leaves) being derived features. Moreover, the more arid and temperate range of the clade likely
- represents a southward expansion from the shared distribution of *Petunia* and *Calibrachoa*, both
- of which is inferred to have originated in the lowland grasslands of southern Brazil, Uruguay,

and northeast Argentina (Reck-Kortmann et al., 2014; 2015; Mäder and Freitas, 2019).
Nonetheless, the relationships within each genus remained inconsistent, with high levels of
conflict among all reconstructed trees. Moreover, we found extensive gene flow within genera,
as supported by both QuIBL and HyDe (Fig. 3) analyses.

Our results corroborated the subdivision of Petunia into two main clades, the long corolla 301 tube and the short corolla tube clades (Reck-Kortmann et al., 2014). The former is characterized 302 by a wide range of flower colors and pollinators, represented here by P. axillaris subsp. parodii 303 (white, hawkmoth-pollinated), P. exserta (red, hummingbird-pollinated), and P. secreta (purple, 304 bee-pollinated). In contrast, the latter consists of species with purple flowers primarily pollinated 305 by bees, represented here by the remaining seven Petunia species. The long corolla tube clade is 306 noteworthy for its documented history of extensive hybridization (e.g., Caballero-Villalobos et 307 al., 2021; Giudicelli et al., 2019), whereas records of interspecific hybridization within the short 308 corolla tube clade are rare and, until now, limited to P. interior and P. inflata (Pezzi et al., 2022). 309

The short corolla tube clade displayed a higher number of discordant gene trees and 310 shorter branch lengths compared to the long corolla tube clade, indicative of rapid radiation. 311 These species are often rare and endemic (Souza et al., 2022), occurring primarily in southern 312 313 Brazil. Geographic isolation serves as the primary reproductive barrier among these species, and pollinators do not impose any reproductive barriers because they are shared among species 314 (Stehmann et al., 2009). Despite their geographic isolation due to microhabitat adaptation, many 315 316 of these species have overlapping distributions, and all are self-incompatible (Stehmann et al., 2009). This scenario presents an opportunity for interspecific gene flow. Whereas field 317 observations have documented only a few hybrids, the substantial level of polymorphism shared 318 319 between these species could be attributed to high levels of ILS (Lorenz-Lemke et al., 2010) or ongoing and recent hybridization events (Fig. 3). 320

Calibrachoa is classified into two subgenera: Calibrachoa and Stimomphis. The 321 322 subgenus Calibrachoa comprises just two species that exhibit significant differences in reproductive biology and habitat compared to species in the subgenus Stimomphis (Fregonezi et 323 al., 2013). Intriguingly, no instances of hybridization have been observed between Calibrachoa 324 and Stimomphis species, underscoring the presence of robust reproductive barriers between 325 subgenera. Stimomphis showed a similar evolutionary history to the one observed in the short 326 corolla *Petunia* clade: rapid radiation, high levels of conflicting gene trees, and extensive ILS. 327 The topology recovered here exhibits minimal congruence with prior studies (Fregonezi et al., 328 329 2012, 2013; Mäder and Freitas, 2019). Notably, the highland clade identified by Mäder and Freitas (2019), represented here by C. elegans, C. eglandulata, C. sendtneriana, and C. linoides, 330 did not emerge in any of our phylogenetic reconstructions. Such high levels of conflict among 331 phylogenetic methods are expected with extensive ILS and hybridization. Moreover, it is worth 332 noting that, except for C. parviflora, all these species are self-incompatible (Fregonezi et al., 333 2013), facilitating the potential for hybridization as these species cannot prevent heterospecific 334 335 gene flow through autonomous selfing (Brys et al., 2015).

It has been demonstrated that ML concatenation methods are often inadequate for accurately recovering species trees when extensive ILS is involved (Kubatko and Degnan, 2007; Mendes and Hahn, 2018). Therefore, ASTRAL is a more suitable alternative in scenarios such as the short corolla *Petunia* clade and *Calibrachoa* subgenus *Stimomphis*. These clades display extremely short branch lengths, suggesting they went through a rapid diversification process, not allowing for genes to coalesce. Species that underwent rapid radiation tend to fall in the "anomaly zone", where the most frequent gene trees do not align with the species tree (Degnan

and Rosenberg, 2006; Linkem et al., 2016) which could explain the high levels of tree 343 344 discordance observed here. To address this challenge, one potential strategy is to sample multiple individuals from the same species (Degnan and Rosenberg, 2006), as was done here. However, it 345 346 is important to note that our individuals were sampled on the same site (Table S1), and they might not fully represent the species' genetic diversity. Thus, they may fail to provide sufficient 347 resolution for phylogenetic inferences. However, when investigating clades harboring numerous 348 rare and endemic species, sampling from the same locality is often unavoidable, but it still 349 350 provides valuable biological insights.

Among the three genera, *Fabiana* exhibited a lower level of tree discordance and greater consistency among phylogenies. The previously available *Fabiana* phylogeny included only eight species but agreed on the close relationship between *F. imbricata* and *F. australis* (Alaria et al., 2022). Here, *F. viscosa* displayed varied phylogenetic placement and the highest level of introgression in QuIBL analyses, indicating significant gene flow, particularly with *F. australis* (Fig. 3). This implies that, in addition to ILS, hybridization plays a central role in causing tree discordance for these species. Intriguingly, these two species do not currently occur in sympatry.

It is crucial to recognize that methods that rely on summary statistics of triplets or 358 359 quartets, such as HyDe, are highly sensitive to substitution rate variation across lineages and genes (Frankel and Ané, 2023), resulting in a high rate of false positives. In addition, such 360 methods can fail to identify hybridization involving ghost or unsampled taxa (Bjorner et al., 361 362 2022), and they often struggle to discern gene flow between sister species, where allele sharing could be interpreted as ancestral polymorphism (Mallet et al., 2016). Moreover, transcriptomic 363 data is inherently more prone to natural selection because it comprises coding regions (Liu et al., 364 365 2015). This can potentially influence rate variation between genes and the lengths of branches in phylogenetic trees (Edwards et al., 2016), ultimately impacting the power detection of 366 hybridization analyses (Frankel and Ané, 2023). Hybridization methods that rely on branch 367 368 length are also susceptible to the influence of rate variation. For instance, QuIBL has demonstrated a propensity for producing false positives when using shorter alignments compared 369 to alignments longer than 1,000 bp (Koppetsch et al., 2023). Notably, our QuIBL dataset 370 exhibited a median length of 494 bp, with 10% of the alignments being longer than 900 bp. The 371 Stimomphis species and Petunia short corolla clade diverged recently and are probably less prone 372 from high-rate variation across lineages. Thus, the intrageneric hybridization events detected by 373 both HyDe and QuIBL are likely authentic. However, we should interpret these results with 374 375 caution as there is still potential for the rate variation assumption to be violated.

376

377 Would it be possible for Petunia and Calibrachoa to hybridize?

Our network analyses suggested an introgression event from *Petunia* to *Calibrachoa* subgenus 378 Stimomphis, which, given recent dating estimates, would have occurred roughly between 8 mya 379 (Särkinen et al., 2013) and 10 mya (Lisa De-Silva et al., 2017). However, recent estimates 380 381 indicate the divergence of *Calibrachoa* and *Fabiana* around 20 mya (Zuntini et al., 2024), suggesting that interspecific hybridization would have to be at least this old. These two genera 382 differ in their chromosome number, with *Petunia* having a haploid chromosome number of seven 383 384 (Stehmann et al., 2009) and Calibrachoa having nine (Wijsman and De Jong, 1985), as in Fabiana (Acosta et al., 2006). Such differences in chromosome numbers typically impose a 385 strong postzygotic barrier against hybridization, either preventing it entirely or resulting in 386 387 hybrid sterility (Levin, 2002). Nevertheless, instances of hybridization and introgression between plant species with different ploidy numbers have been documented (Chapman and Abbott, 2010), 388

and both *Petunia* and *Calibrachoa* subgenus *Stimomphis* share similar geographic distribution,

- morphology, habitat, and potential group of pollinators (Stemann et al., 2009). One possible
- explanation for our result is that this change in chromosome number occurred in the ancestral
   lineage of *Petunia* after the admixture event, such as in the scenario posited by PhyloNet, where
- the admixture event is from the common ancestor of all *Petunia* species. However, SNaO
- 394 contradicts this hypothesis, as the introgression event is inferred to have occurred after a
- reduction in chromosome number in the common ancestor of *Petunia* (Fig. 4). Alternatively,
- 396 sufficient chromosomal homology may have enabled meiotic pairing during diversification.
- 397 Artificial crosses between *Calibrachoa* and *Petunia* demonstrated some success in embryo
- formation but failed in germination (Olschowski et al., 2013). However, Milicia et al., (2021)
- crossed *P. inflata* with *C. hybrida*, and despite a significantly lower percentage of viable pollen
   granules than intrageneric crosses, the hybrids produced 5% of viable pollen, highlighting plant
- 401 species' flexibility in chromosome rearrangement to allow successful meiosis. Thus,
- 402 hybridization between these genera may not be out of the question. Additionally, the low chance403 of current hybridization does not exclude the possibility of ancient hybridization.
- Despite the inference of intergeneric hybridization from SNaQ and PhyloNet, we did not 404 405 detect any support for such an event from our Twisst analysis. Instead, the discordance appears best explained by ILS as the two minor topologies are present in nearly equal frequencies (Fig. 406 5) We note that inheritance probabilities from SNaQ and PhyloNet were very low (1 to 3%, Fig. 407 4), and thus, this reticulation event, if it occurred, might be at the boundary of detection. The 408 absence --- or very low levels --- of gene flow between these two genera highlights how important 409 chromosome number difference was to prevent hybridization, which allowed Petunia and 410 Calibrachoa to undergo parallel radiation despite their many ecological similarities and 411 geographic overlaps. Regardless, hybridization between the two genera merits future 412 investigation when full genomes become available for these genera (Bombarely et al., 2010). 413 414 Introgression of even a small fraction of the genome could potentially carry a large phenotypic effect (Clarkson et al., 2014) and facilitate rapid radiations (Meier et al., 2017). However, 415 detecting such events with confidence is challenging as it involves identifying introgressed 416 genomic regions and linking them to adaptations (Taylor & Larson, 2019; Suarez-Gonzalez et 417 al., 2018). 418
- 418

#### 420 Conclusions

421 Here, we investigated the origins of tree discordance in the *Petunia-Calibrachoa-Fabiana* 

- 422 Solanaceae clade using a comprehensive genome-scale dataset encompassing multiple species.
- 423 Our results confirm *Petunia* as the sister genus to *Calibrachoa* + *Fabiana*. However, the
- 424 relationships among species within these genera remain unsolved. The discordance in tree
- 425 topologies within the short corolla tube *Petunia* clade and *Calibrachoa* subgenus *Stimomphis*
- 426 arises from a combination of ILS due to their rapid diversification and past and ongoing
- 427 hybridization events. Instances of high ILS and extensive hybridization are not uncommon in the
- evolutionary history of plants (e.g., Kleinkopf et al., 2019; Morales-Briones et al., 2021; McLay
- 429 et al., 2023), but pinpointing the specific taxa involved in the hybridization events is still a
- 430 daunting task, and one of the reasons why different methods often yield conflicting results (Gates
- et al., 2023). These introgression events likely contributed to the species' genetic diversity, aiding
  their adaptation during their radiation. Additionally, our network reconstructions indicated
- 432 then adaptation during then radiation. Additionary, our network reconstructions indicated
   433 potential intergeneric hybridization between *Calibrachoa* and *Petunia*, two genera characterized
- 434 by distinct chromosome numbers. Considering the weak hybridization signals observed in

- 435 network analyses, the lack of support from gene tree topology weights, and the known current
- 436 barriers due to differing base chromosome numbers, it leads us to believe that such a
- 437 hybridization event did not occur. However, both of our network analyses indicated intergeneric
- 438 gene flow, suggesting there is still a remote possibility that this could have occurred and may
- have been facilitated by strong selection despite the barrier imposed by differing chromosome
- base numbers between the two genera. Whole genome analyses could solve the intergeneric
- 441 hybridization puzzle and contribute to ascertaining which genomic regions may have been
- 442 involved in the *Petunia-Calibrachoa* introgression. Overall, our study sheds light on the complex
- evolutionary history of this charismatic South American clade, providing crucial insights into its
- 444 adaptation and diversification.
- 445

#### 446 Acknowledgments

- 447 This work used the RMACC Alpine supercomputer, funded by the University of Colorado
- 448 Boulder, the University of Colorado Anschutz, Colorado State University, and the National
- 449 Science Foundation (award 2201538). SDS was supported by NSF-DEB (award 1553114). PHP
- 450 was supported by a CAPES/PRINT fellowship.
- 451

#### 452 **Declaration of competing interest**

- 453 The authors declare that they have no competing interests.
- 454

## 455 Data availability

- 456 All scripts and processed data files are available at https://github.com/pedrohpezzi/Petunia-
- 457 Calibrachoa-Fabiana\_TreeDiscordance.git and https://figshare.com/s/c3f6e7305660e03031ec.
- 458 The raw RNA-seq data files are available in SRA under the BioProject accession number
- 459 PRJNA746328.
- 460

## 461 **CRediT author contributions**

- 462 **Pedro H. Pezzi:** Conceptualization, Methodology, Software, Formal analysis, Investigation,
- 463 Writing Original Draft, Visualization. Lucas C. Wheeler: Conceptualization, Data Curation,
- 464 Methodology, Software, Investigation, Writing Review and Editing. Loreta B. Freitas:
- 465 Conceptualization, Writing Review and Editing, Investigation, Supervision. Stacey D. Smith:
- 466 Conceptualization, Resources, Investigation, Writing Review and Editing, Supervision, Project
- 467 administration, Funding acquisition.

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#### 811 Supplemental figures



Figure S1. ASTRAL phylogenetic tree depicting individuals without species assignments.

814 Numerical values on branches indicate local posterior probabilities that are below 100. The scale

815 bar denotes branch length in coalescent units.



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 818 Figure S2. Phylogenetic tree depicting the relationship within the *Petunia-Calibrachoa-Fabiana*

819 clade and the outgroups. The tree was constructed using quartets of taxa and the coalescent

820 model with SVDQuartets. Branch numbers indicate nonparametric bootstrap values, derived

from 200 bootstrap replicates.

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Figure S3. Summary Graphs of QuIBL Results. The upper graph illustrates the proportion of
topologies not solely explained by Incomplete Lineage Sorting (ILS), while the lower graph
depicts the proportion of gene trees indicating introgression events.



Figure S4. Log-likelihood results from the SNaQ analysis. The steepest change in log-likelihood
signifies the optimal network with a single reticulation.

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832 Supplemental tables are available on GitHub.