1	Original Article
2	Target Enrichment Data Uncovers Rapid Radiation, Whole Genome
3	Duplication, and Extensive Hybridization in Slipper Orchid Genus
4	Cypripedium L.
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Background and Aims: *Cypripedium* is the most widespread and morphologically
 diverse genus of slipper orchids. Despite several published phylogenies based on Sanger
 sequencing data, the topology and monophyly of its infrageneric taxa remained uncertain.
 Here, we aimed to reconstruct a robust section-level phylogeny of *Cypripedium* and
 explore its evolutionary history using target capture data for the first time.

- Methods: We used the orchid-specific bait set "Orchidaceae963" to reconstruct the phylogeny of *Cypripedium* based on 614 nuclear loci, covering 11 out of 13 sections.
 Subsequently, we investigated tree discordance, estimated divergence times and ancestral ranges, searched for anomaly zones, polytomies, and diversification rate shifts, and identified gene duplication and hybridization events.
- Key Results: All sections were recovered as monophyletic, contrary to the subsections 11 ٠ within sect. Cypripedium. Although the two subclades within this section did not 12 correspond to its two subsections, they matched the geographic distribution of their 13 14 species. Additionally, we discovered high levels of discordance in the short backbone branches of the genus and within sect. Cypripedium, which can be attributed to gene 15 duplication and hybridization events, a potential whole genome duplication, and 16 incomplete lineage sorting caused by rapid radiation. Our biogeographic analysis 17 18 suggested a Neotropical origin of the genus during the Early Miocene (~20 Ma). The rapid radiations at the backbone likely occurred in Southeast Asia around the Middle 19 20 Miocene Climatic Transition (~15-13 Ma), followed by several independent dispersals 21 back to the New World. Moreover, the Pliocene-Quaternary glacial cycles may have contributed to further speciation and reticulate evolution, giving rise to a hybrid swarm 22 23 within sect. Cypripedium.

1	•	Conclusions: Our study provided novel insights into the evolutionary h	nistory of
2		Cypripedium based on high-throughput molecular data, shedding light on the dy	namics of
3		its distribution and diversity patterns from its origin to the present.	
4	K	y words: Cypripedium, slipper orchids, phylogenomics, target enrichment,	historical

- 5 biogeography, anomaly zone, rapid radiation, whole genome duplication, reticulate evolution,
- 6 hybridization.

1 INTRODUCTION

2 The family Orchidaceae comprises the most species-rich family of vascular plants, with c. 28,000 species in five subfamilies and ~750 genera (Chase et al., 2015; Christenhusz et al., 2017). Their 3 4 great diversity has fascinated and puzzled scientists for centuries, including the father of evolutionary theory, Charles Darwin, who once wrote, "I never was more interested in any 5 subject in my life, than in this of Orchids" (Darwin Correspondence Project, n.d.). 6 7 Unfortunately, today, their diversity is highly threatened mainly due to habitat destruction and 8 unsustainable harvesting (DL Roberts and Dixon, 2008), prompting their protection by local and 9 national laws, as well as the Convention on International Trade in Endangered Species (aka 10 CITES; Appendices I, II, and III, 2023). In efforts to describe their diversity and facilitate informed conservation measures, a variety of molecular data, including high-throughput genomic 11 and transcriptomic data, has been used to reveal the relationships between orchid subfamilies in 12 recent decades (Cameron et al., 1999; Freudenstein et al., 2004; Givnish et al., 2015; Kim et al., 13 2020; Pérez-Escobar et al., 2021; Serna-Sánchez et al., 2021). However, phylogenetic support at 14 lower taxonomic ranks in Orchidaceae has been low primarily due to the limited genetic variation 15 in the commonly used markers (i.e., *rbcL*, *matK*, ITS, chloroplast intergenic spacers). The genus 16 Cypripedium L. is one such orchid taxon whose internal phylogenetic relationships have vet to be 17 18 resolved.

Cypripedium is a genus of temperate perennial herbs in the subfamily of lady's slipper orchids,
Cypripedioideae, and it currently consists of approximately 50 accepted species (Frosch and
Cribb, 2012; SC Chen *et al.*, 2013; Christenhusz *et al.*, 2017; POWO, 2023). Although *Cypripedium* has about half the species number of the largest slipper orchid genus, *Paphiopedilum* Pfitzer, it is the most morphologically diverse (Fig. 1) and widespread
(Supplementary Data Fig. S1) of all five cypripedioid genera. Its distribution is mainly

circumboreal, but its range extends from the Arctic Circle to Central America (~14°-70° North; J 1 Li et al., 2011; Frosch and Cribb, 2012). Eastern Asia, especially temperate China, constitutes the 2 genus' main center of diversity, harboring approximately 70% of all Cypripedium species (J Li et 3 4 al., 2011). They occur in various habitats and altitudes, from forests to wetlands and grasslands and from sea level to 4,900 m in the Himalayas (Frosch and Cribb, 2012). 5

(a) Irapeana





Arietinum







Obtusipetala

Trigonopedia



Californica



Flabellinervia











Sinopedilum





Figure 1(a): Pictures of the *Cypripedium* taxa per section included in the final phylogeny. (A) C. irapeanum, (**B**) *C. plectrochilum*, (**C**) *C*. subtropicum (= C. singchii; Frosch & Cribb, 2012), (**D**) C. wardii, (E) C. margaritaceum, (**F**) C. fargesii, (G) C. lichiangense, (**H**) C. lentiginosum, (I) C. sichuanense, (**J**) C. passerinum, (K) C. flavum, (L) C. reginae, (M) C. reginae var. alba, (N) C. californicum, (**O**) *C. acaule*, (**P**) C. yatabeanum, (**Q**) $C. \times alaskanum$, (**R**) C. guttatum, (S) C. japonicum, (**T**) C. formosanum, (**U**) *C. bardolphianum*, (**V**) *C. micranthum.* Credits: (A) by M. Béhar;

(B), (K), (P), (Q), (T), and (V) by J.-B. Chazalon; (C), (D), (H), (J), and (O) by W. Frosch; (R) and (S) by L. Chen; (E)-(G), (I), (N), and (U) by S. Urban; (L) by B. Isaac; (M) by M. Sunouchi (see Acknowledgements for more details).

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(b) Cypripedium



Credits: (A), (L), (Q), and (Y) by V. Steindl; (D), (E), (G), (J), (S)-(U), (X), (Z), and (AA) by J.-B. Chazalon; (P) by L. Chen; (W) by Orchi; (B), (F), (H), (I), (K), (M), (O), and (R) by S. Urban; (C), (N), (V), and (BB) by W. Frosch (see **Acknowledgements** for more details).

Like other slipper orchids, flowers of *Cypripedium* species have a profoundly inflated, slipper-1 2 shaped lip (i.e., labellum) that gives them their distinctive morphology. The lip traps pollinators 3 that enter through the upward-facing opening thanks to its incurved, glabrous, slippery margins, 4 with the only escape routes passing through its basal orifices under the two anthers at each side of the column (Cribb, 1997; Frosch and Cribb, 2012). Unlike other slipper orchids, Cypripedium 5 6 species are traditionally recognized by their (usually) plicate leaves and unilocular ovaries with 7 parietal placentation (Cox et al., 1997; Cribb, 1997). Although the reliability of these distinctive 8 characters has been questioned (Atwood, 1984), phylogenetic studies consistently support the 9 monophyly of the genus (Fatihah et al., 2011; J Li et al., 2011; Guo et al., 2012; H Liu et al., 10 2021a; Szlachetko et al., 2021; J-Y Zhang et al., 2022). On the other hand, its infrageneric 11 classification has constantly changed during the last two centuries.

Following *Cypripedium*'s description, the great interest in Cypripedioideae led to numerous 12 taxonomic revisions in the subfamily with often incongruent results (Linnaeus, 1753; Rafinesque, 13 14 1836; Lindley, 1840; Reichenbach, 1854; Pfitzer, 1888, 1894; Rolfe, 1896; Atwood, 1984; Cox et 15 al., 1997; Eccarius, 2009; Perner, 2008; Supplementary Data Table S1). To name a few, Linnaeus (1753) initially recognized only one species of Cypripedium (i.e., C. calceolus L.) and a few 16 17 varieties currently holding a species status. Lindley (1840) described 22 species within the genus 18 classified in a number of subgeneric groups. The classifications of Pfitzer (1903) taxonomically expanded *Cypripedium* with 28 species and numerous subgeneric categories, including four 19 20 sections. In Cribb's (1997) taxonomic treatment, the number of species increased to 45 and the 21 sections to 11, while Eccarius (2009) divided *Cypripedium* into two subgenera, 13 sections, and 37 species, lowering the rank of multiple species to subspecies or varieties. 22

Recent molecular phylogenies based on nrDNA ITS and five cpDNA markers by J Li *et al.*(2011) indicated that, among the non-monotypic groups, eight sections are monophyletic

[Arietinum C. Morren, Bifolia (Lindl.) S. C. Chen, Cypripedium, Flabellinervia (Pfitzer) 1 2 Hennessy ex P. J. Cribb, Obtusipetala (Pfitzer) P. J. Cribb, Sinopedilum Perner, Subtropica S. C. 3 Chen & K. Y. Lang, and Trigonopedia Franch.] while two sections [Irapeana P. J. Cribb and 4 Retinervia (Pfitzer) S. C. Chen] and the two subsections of sect. Cypripedium [Cypripedium and Macrantha (Kraenzl) P. J. Cribb] are non-monophyletic, following the classification by Cribb 5 6 (1997) and Perner (2008). These results prompted further infrageneric treatments by Frosch and 7 Cribb (2012) and SC Chen et al. (2013), producing the two currently used classification systems 8 of Cypripedium. Although based on the same phylogenies by J Li et al. (2011), Frosch and Cribb 9 (2012) proposed 13 sections with 48 species, whereas SC Chen et al. (2013) increased these 10 numbers to 15 and 51, respectively, adding two new monotypic sections: Palangshanensia S. C. Chen & Z. J. Liu and Wardiana S. C. Chen & Z. J. Liu (Supplementary Data Table S1). 11 After the publication of J Li et al. (2011), several studies included molecular phylogenies with 12 *Cypripedium* species, four of which specifically focused on the relationships of the infrageneric 13 taxa of Cypripedium (Fatihah et al., 2011; Guo et al., 2012; H Liu et al., 2021a; Szlachetko et al., 14 15 2021; J-Y Zhang et al., 2022). These studies used a multilocus approach with up to eight Sangersequenced nuclear and chloroplast DNA markers in different combinations and with different 16 17 phylogenetic reconstruction methodologies (i.e., Parsimony, Maximum Likelihood, and Bayesian 18 Inference). The topologies and the monophyly of some subgeneric taxa were congruent among the produced phylogenies (e.g., sect. Irapeana being sister to the rest; monophyly of sect. 19 20 Arietinum, Bifolia, Cypripedium, Flabellinervia, Obtusipetala, Sinopedilum, and Trigonopedia). 21 However, the topology and monophyletic status of other taxa (e.g., the monophyly of the two subsections within sect. *Cypripedium*) and the topology at the backbone of the phylogeny remain 22

23 uncertain.

24 The unresolved phylogeny of the genus *Cypripedium* not only prevents the accurate evaluation of

1 the relationships between the currently established subgeneric groups but also our understanding 2 of their evolutionary history. A well-resolved and robust phylogeny is fundamental for addressing 3 further questions regarding their divergence time, diversification rate shifts, ancestral spatial 4 distribution patterns, and hybridization events. Furthermore, it will provide a solid foundation for the efficient management of their conservation, especially as their continuous human-driven 5 6 population decline is predicted to exacerbate due to climate change (Nicolè *et al.*, 2005; Izawa *et* 7 al., 2007; Minasiewicz et al., 2018; Kolanowska and Jakubska-Busse, 2020; H Liu et al., 2021b; 8 Chandra et al., 2023; Yamashita et al., 2023).

9 It is widely recognized that the use of multiple genes can improve the accuracy of phylogenetic 10 reconstruction, and single- or low-copy nuclear genes are increasingly used due to their rapid 11 evolutionary rates and biparental inheritance (Guo et al., 2012; N Zhang et al., 2012; Z Li et al., 2017). A target enrichment approach would allow the sequencing of hundreds of low-copy 12 markers via high-throughput sequencing methods and, therefore, more robust estimates of 13 relationships with greater support. Moreover, the use of the recently designed orchid-specific 14 15 baits "Orchidaceae963" by Eserman et al. (2021) could provide sufficient information to resolve recent and rapid radiations in deep and shallow phylogenetic scales, allowing for the 16 17 characterization of species-level relationships and the resolution of long-debated polytomies 18 within Orchidaceae.

In this study, we used a target enrichment approach using the "Orchidaceae963" baits to reconstruct a well-supported phylogeny of the genus *Cypripedium* at the section level. Based on our results, we evaluated the two most recently published classification systems of the genus by Frosch and Cribb (2012) and SC Chen *et al.* (2013) and the congruence of the recovered relationships with published phylogenies based on Sanger data. Additionally, we aimed to gain new insights into the evolution of *Cypripedium* by answering the following questions: (1) Does the current classification stand up to phylogenetic reconstructions based on genomic data? (2) Which biological processes explain higher levels of gene tree discordance in some parts of the phylogeny? (3) When and where did *Cypripedium* originate and diversify and how did this diversification relate to geographic expansion of the lineages and paleoclimate? (4) Is the hybrid status of some taxa supported by molecular data? To answer these questions, we explored tree discordance, estimated divergence times and ancestral ranges, searched for anomaly zones and diversification rate shifts, and identified gene duplication and hybridization events.

8

9 MATERIALS AND METHODS

10 Taxon Sampling

We sampled leaf tissue from 53 specimens representing 36 species, eight varieties, and three natural hybrids of the genus *Cypripedium* (following the taxonomy of Frosch and Cribb, 2012; Supplementary Data Table S2). Fifty of the sampled individuals came from the Botanical Collection at Oberhof, Eurasburg, associated with the Botanical Garden Munich-Nymphenburg (BGM), and three from the Botanische Staatssammlung München herbarium (BSM-SNSB, herbarium acronym M). The material from the living collection was stored in silica-gel to be dried immediately after collection.

The sequence data from the above-collected tissue samples was combined with publicly available orchid genomes and transcriptomes. These represented species from all slipper orchid genera (incl. ten *Cypripedium* species, three of which are new to our sampling) as well as orchids from three outgroup subfamilies (i.e., Apostasioideae, Epidendroideae, and Vanilloideae). As a result, our final dataset included *Cypripedium* species from all sections except sect. *Enantiopedilum* 1 Pfitzer, consisting of *C. fasciculatum* and *C. palangshanense*, and sect. *Retinervia*, consisting of

- 2 *C. elegans* and *C. debile* (following Frosch and Cribb, 2012; Supplementary Data Table S3).
- 3

4 Library Preparation, Target Enrichment, and Sequencing

We isolated total genomic DNA from silica-dried or herbarium leaf tissue using the NucleoSpin 5 6 Plant II kit: Genomic DNA from plants (Macherey-Nagel, Düren, Germany) following a 7 modified version of the manufacturer's manual (Supplementary Data Table S4). Next, we quantified the DNA concentration with a Qubit 4 fluorometer using a Broad Range (BR) or High 8 9 Sensitivity (HS) assay kit (Thermo Fisher Scientific, Waltham, Massachusetts, USA). DNA was 10 sheared to an average fragment size of 350 bp with a Covaris M220 Focused-ultrasonicator 11 (Covaris, Woburn, Massachusetts, USA). We assessed the DNA fragment size distribution using 12 a High Sensitivity DNA ScreenTape on a 4150 TapeStation System (Agilent Technologies, Santa Clara, California, USA). 13

We prepared dual indexed libraries according to the instruction manual using the NEBNext Ultra 14 15 II DNA Library Prep Kit for Illumina and the NEBNext Multiplex Oligos for Illumina (Dual 16 Index Primers Set 1, New England Biolabs, Ipswich, Massachusetts, USA) and following the 17 recommended conditions of bead-based size selection according to distribution of DNA 18 fragments per sample. Next, we amplified the adaptor-ligated libraries with eight PCR cycles and measured DNA concentration using the Qubit. The average fragment size of the libraries was 19 20 assessed with the TapeStation. Prior to hybridization, the libraries were pooled in equal concentrations to include 250 ng of each library, with a maximum of 15 libraries per pooled 21 22 library.

For the hybridization enrichment reaction, we combined the pooled libraries with the custom orchid-specific bait set "Orchidaceae963" (Daicel Arbor Biosciences myBaits Target Capture

Kit, Ann Arbor, MI, USA) and incubated at 60 °C (hybridization temperature, TH) for 16 hours 1 2 overnight, following the Standard Protocol and the Blockers Mix setup designed for plants 3 (myBaits Hybridization Capture for Targeted NGS, User Manual v. 5.02). The bead-based 4 cleanup of the bait-target hybrids was performed at a wash temperature (TW) of 60 °C, and the hybridized libraries were subsequently amplified for 14 PCR cycles at 60 °C (annealing 5 6 temperature, TA). Then, we purified the amplification reaction following the PCR clean-up 7 protocol of the NucleoSpin Gel and PCR Clean-up kit (Macherey-Nagel, Düren, Germany). Finally, we checked the concentration and fragment size distribution of the libraries as before, 8 9 using the Qubit and the TapeStation. The enriched pooled libraries were sequenced on an 10 Illumina NovaSeq 6000 sequencing system (SP flow cell) at the Core Facility Genomics (CF-11 GEN) of the Helmholtz Zentrum München, Germany (Deutsches Forschungszentrum für 12 Gesundheit und Umwelt, GmbH).

13

14 Read Processing and Assembly

15 We created a set of references from orchid genomes and transcriptomes available on the Sequence Read Archive (SRA) of NCBI to improve gene extractions (Supplementary Data Table 16 17 S4; Sayers et al., 2022). Original target exon sequences from the Orchidaceae963 bait set 18 (https://github.com/laeserman/Orchidaceae963/blob/main/Orchidaceae963-targets.fa) were concatenated into 'genes' and used to identify the corresponding complete CDS from the 19 20 Phalaenopsis equestris (Schauer) Rchb. f. genome using BLAST. When no hits were produced, 21 we used the genome of *Dendrobium catenatum* Lindl. instead. In the end, 950 out of the original 963 genes were extracted. Then, we used the raw transcriptome assembly from 18 orchids 22 mentioned in the Taxon Sampling section, including Vanilla shenzhenica Z. J. Liu & S. C. Chen, 23 24 and 17 species of slipper orchids to extend the genome references. RNAseq data processing and transcriptome assembly followed Morales-Briones *et al.* (2021). We used CAPTUS v.0.9.90
(Ortiz *et al.*, 2023) to extract the corresponding loci from the 18 transcriptomes and the genomes
of *Apostasia shenzhenica* Z. J. Liu & L. J. Chen, *D. catenatum*, *P. equestris*, and *Vanilla planifolia* Andrews. The extracted loci were used as the extended reference dataset for loci
extraction in our own generated target enrichment data of *Cypripedium*.

We checked the quality of the raw reads using FastOC v0.11.9 (Andrews, 2010) and MultiOC 6 7 v1.14 (Ewels et al., 2016). Then, we used CAPTUS to trim the sequencing adaptors and low-8 quality bases, assemble the reads, and extract the nuclear loci based on the reference dataset. To 9 decrease the retention of contigs resulting from potential erroneous reads, we set the minimum 10 contig depth to eight for the assembly step and the minimum percentages of identity and coverage 11 to 75 and 50, respectively, for the extraction step. Similarly, we assembled and extracted the nuclear loci from the genomes and transcriptomes used as references to combine them with our 12 data for further analysis. We then extracted the coding sequences from the combined dataset 13 14 using CAPTUS, with the removal of paralogs disabled.

15

16 Orthology Inference

17 To infer orthologs for the phylogenetic reconstruction, we followed a modified version of the 18 methods described in Morales-Briones al. (2022;et https://bitbucket.org/dfmoralesb/target enrichment orthology). 19 First, we re-aligned the 20 untrimmed and unfiltered output alignments from CAPTUS using MACSE v2.07 (Ranwez et al., 21 2018) with default parameters. Next, we replaced "!" with gaps at the frameshifts and used Phyx (JW Brown et al., 2017) to remove aligned columns with more than 90% missing data. We 22 inferred maximum likelihood (ML) homolog gene trees using IQ-TREE 2 v.2.0.7 using extended 23 model selection (Kalyaanamoorthy et al., 2017) and no clade support (Minh et al., 2020). Then, 24

we masked mono- and paraphyletic tips that belong to the same taxon, keeping the tips with the 1 2 most unambiguous characters in the trimmed loci alignments for each taxon as described in 3 (Yang and Smith, 2014). Spurious tips with unusually long branches were removed by reducing 4 the tree diameter with TreeShrink v.1.3.9 (Mai and Mirarab, 2018). We ran TreeShrink twice with the quantile set to 0.01, using the output of the first run for the second run to avoid over-5 6 trimming. We wrote FASTA files from the output homolog trees and followed the same steps as 7 for the output alignments from CAPTUS, aligning them using MACSE with the default parameters, replacing the "!" with gaps at frameshifts, and removing aligned columns with >90% 8 9 missing data with Phyx. To infer the final homolog gene trees, we used IQ-TREE 2, extended 10 model selection, and assessed the clade support with 1,000 ultrafast bootstrap (BS) replicates.

11 Orthology inference was carried out with the tree-based "monophyletic outgroup" (MO) method described in Yang and Smith (2014). The MO method searches for clusters with monophyletic 12 ingroups rooted at the outgroups in the homolog trees, discarding those with duplicated taxa in 13 the outgroups. Subsequently, it infers the orthologs from root to tip, keeping the ortholog subtree 14 15 with the most taxa. To infer the orthologs, we set all Cypripedioideae members as ingroup, and the remaining taxa (i.e., A. shenzhenica, D. catenatum, Phalaenopsis equestris, Vanilla 16 17 *planifolia*, V. *shenzhenica*) as outgroups. In an initial MO run, we kept only ortholog groups with 18 at least 20 ingroup taxa. Paphiopedilum malipoense S. C. Chen & Z. H. Tsi, was causing the outgroups of most loci trees to be "non-monophyletic", due to low quality of the transcriptome or 19 20 a mislabeling of the raw data sample submission; therefore, it was removed, and the MO ortholog 21 inference was repeated. Following the second MO run, we removed the samples present in $\leq 35\%$ of the ortholog trees. Finally, we repeated the orthology inference, keeping only ortholog groups 22 23 with at least 35 ingroup taxa (~50% of retained samples), retaining a dataset of 74 samples and 792 ortholog trees (326–789 ortholog trees per sample; Supplementary Data Table S5). 24

1

2 Phylogenetic Reconstruction

We used concatenation and coalescent-based methods to reconstruct the phylogeny of 3 4 *Cypripedium.* First, a concatenated alignment was produced using the clean ortholog alignments (following the Phyx step), retaining 614 orthologs with at least 500 characters and 50 taxa. We 5 estimated a ML tree of the concatenated matrix with IO-TREE 2. We searched for the best 6 7 partition scheme using ModelFinder implemented within IQ-TREE (Kalyaanamoorthy et al., 8 2017) and 1,000 ultrafast BS replicates to assess clade support. Regarding the coalescent-based 9 approach, we first inferred ML trees from the same 614 individual orthologs used for the 10 concatenation-based phylogeny. Individual ortholog ML trees were inferred as previously 11 described for the final homolog trees. Then we used the quartet-based species-tree inference method ASTRAL v1.15.2.4 (wASTRAL-unweighted), which is statistically consistent under the 12 multispecies coalescent (MSC) model and thus useful for handling incomplete lineage sorting (C 13 14 Zhang et al., 2018; C Zhang and Mirarab, 2022). We used the 614 individual ML ortholog trees, 15 default ASTRAL parameters, and branch support was assessed using local posterior probabilities 16 (LPP; Sayyari and Mirarab, 2016). Due to the similarity in the topologies recovered between the 17 concatenation and coalescent-based approaches, all subsequent analyses were carried out using 18 the ASTRAL species tree unless stated otherwise.

19

20 Gene Tree Discordance Estimation

We quantified the conflict among gene trees on each node of the inferred species tree by estimating the number of conflicting and concordant bipartitions with Phyparts (Smith *et al.*, 2015). To do this, we used the individual ML ortholog trees and set a threshold of at least 50% BS support for a node to be considered informative. We plot the Phyparts result using the "missing and uninformative" script (i.e., "phypartspiecharts_missing_uninformative.py;
<u>https://bitbucket.org/dfmoralesb/target_enrichment_orthology</u>) with Python v3.10.10 to add pie
charts at the nodes while taking into consideration missing data (i.e., when input trees do not
have the same number of tips).

We also used Quartet Sampling (QS; Pease et al., 2018) to differentiate between lack of support 5 and conflicting nodes on the species tree. QS estimates branch support and conflict by sampling 6 7 quartets from the species tree and the corresponding concatenated alignment and calculating the 8 proportion of the three possible topologies at each node. As a result, it simultaneously evaluates 9 the consistency of information (Quartet Concordance, QC), the presence of secondary 10 evolutionary histories (Quartet Differential, QD), and the amount of information (Quartet Informativeness, QI) of internal nodes. We ran with 1,000 QS replicates with RAxML-NG 11 (Kozlov et al., 2019) as ML inference tool. The results were plotted using R by color-coding the 12 values of QC on each node and annotating them with the rest of the estimated values 13 (https://bitbucket.org/yanglab/conflict-analysis/src/master/). 14

15

16 Anomaly Zone Test

The anomaly zone, characterized by the presence of a set of short internal branches in the species tree, occurs when gene tree topologies that are discordant with the species tree topology are observed more frequently than those that are concordant (Linkem *et al.*, 2016). It arises from consecutive rapid diversification events leading to incomplete lineage sorting (ILS).

We estimated the boundaries of the anomaly zone for the internal nodes of our species tree following the calculations in Linkem *et al.* (2016) (<u>https://github.com/cwlinkem/anomaly_zone</u>) to investigate whether the high amount of gene tree discordance observed in numerous short branches of the tree could be explained by ILS. The calculations are based on equation 4 of Degnan and Rosenberg (2006), which defines the boundaries of the anomaly zone, α(x). In this
 equation, x is the length of an internal branch in the species tree, and its descendant internal
 branch has a length y (in coalescent units). If y is < α(x), then the internode pair is considered to
 be in the anomaly zone.

5

6 *Polytomy Test*

Additionally, due to the presence of short branches with low support, we tested whether we could 7 reject the null hypothesis that any branch in the species tree has a length equal to 0, or in other 8 9 words, is a polytomy. We used the polytomy test (-t 10) option in ASTRAL version 5.7.8 with default parameters (Sayyari and Mirarab, 2018). The ASTRAL polytomy test relies on the 10 11 distribution of the quartet frequencies of gene trees around each branch of the species tree to test 12 this hypothesis, annotating the branches of the output tree with the resulting p-values. Under the 13 null hypothesis, the three unrooted quartet topologies defined around the branch are expected to 14 have equal frequencies. Although failure to reject the null hypothesis may indicate a real (i.e., hard) polytomy, it might also be caused by lack of power or signal (i.e., soft polytomy). 15

16

17 Mapping Whole Genome Duplications

18 We mapped gene duplication events on our species tree based on the subclade orthogroup tree 19 topology method described in Yang al. (2018; et 20 https://bitbucket.org/blackrim/clustering/src/master/). This method extracts the rooted 21 orthogroups from each homolog tree. Then, it detects gene duplication events when the 22 orthogroup subclades share two or more ingroup taxa and maps the percentage of duplicated genes to the corresponding branch of the species tree. Alternatively, the duplications are mapped 23 24 on the most recent common ancestor branch if the gene tree has missing taxa or if its topology is incongruent with the species tree. To avoid the overestimation of the duplication percentages due
to nested duplications, each branch of the species tree is restricted to one duplication event for
each extracted clade (Yang *et al.*, 2015).

4 Similarly to Yang *et al.* (2018), we tested two filters to map the gene duplications: a bootstrap and a local topology filter. The bootstrap filter requires orthogroups to have an average bootstrap 5 percentage of \geq 50% (Z Li *et al.*, 2015), while the local topology filter requires the sister clade of 6 7 the gene duplication branch in the orthogroup to include a subset of the taxa in the corresponding sister clade in the species tree (Cannon et al., 2015). We plotted the percentages of gene 8 9 duplications per number of branches in R (R Core Team, 2023). A WGD will produce a large 10 burst (i.e., an outlier percentage, usually $\geq 20\%$) of shared duplications across taxa and loci 11 (Yang *et al.*, 2018).

12

13 *Testing Hybridization Events*

Our dataset included three taxa described as hybrids by both Frosch and Cribb (2012) and SC 14 15 Chen et al. (2013); namely, $C. \times alaskanum$ P. M. Br. (C. guttatum Sw. $\times C.$ yatabeanum 16 Makino), C. × columbianum Sheviak [C. montanum Douglas ex Lind]. × C. parviflorum Salisb. 17 var. pubescens (Willd.) O. W. Knight] and C. \times ventricosum Sw. (C. calceolus \times C. macranthos 18 Sw.). We tested whether our data supports their status as hybrids of their putative parent taxa using explicit phylogenetic networks in PhyloNet (Wen et al., 2018). Phylonet allows for 19 20 horizontal edges that visualize the genetic inheritance through gene flow, mapping the inheritance 21 probabilities (γ) for each parent hybrid edge to estimate the percentage of loci a hybrid inherited from each parent. 22

To reconstruct the phylogenetic networks, we first rooted the final ortholog trees and extractedthe three subclades containing each hybrid, along with the putative parents and other taxa sharing

their MRCA. We reduced computational load by removing duplicated taxa, leaving a single representative for each monophyletic taxon. In the case of paraphyletic taxa, one representative taxon was left from each conspecific monophyletic subclade or from a group of consecutively diverging conspecific varieties. The taxa present in most orthologs were favored to maximize the final number of loci used for Phylonet. Similarly, *Phalaenopsis equestris* was chosen as an outgroup taxon because it had the highest amount of retained loci. Gene trees missing any of these selected taxa were excluded from the analysis.

8 Since calculating the likelihood of a phylogenetic network is computationally intensive in 9 PhyloNet, we inferred the phylogenetic networks based on a maximum pseudo-likelihood (MPL) 10 measure via the InferNetwork_MPL command (Yu and Nakhleh, 2015). We set the number of 11 maximum reticulation events to one and the number of optimal output networks to ten for all three tests. The option "po" was specified to optimize the branch lengths and inheritance 12 probabilities under full likelihood for the inferred C. \times alaskanum networks. This optimization 13 14 was only performed for the C. \times alaskanum networks, which contain only four taxa, as it gets 15 more time-consuming with an increasing number of taxa.

Furthermore, since our sampling included other taxa that have been proposed—but not widely accepted—as hybrids (e.g., *C. froschii* Perner, also known as *C.* × *froschii* Perner) and since hybridization is considered to be pervasive within *Cypripedium* in nature (Klier *et al.*, 1991; S-J Hu *et al.*, 2011; Frosch and Cribb, 2012; Szlachetko *et al.*, 2017; Pupulin and Díaz-Morales, 2018), we ran further tests setting the number of maximum reticulation events from two to ten for the networks containing *C.* × *columbianum* and *C.* × *ventricosum*.

The phylogenetic networks with the highest total log probability were visualized in Dendroscope
v3.8.8 (Huson and Scornavacca, 2012) and the inheritance probabilities were mapped with

PhyloNetworks' v0.16.2 (Solís-Lemus *et al.*, 2017) companion package PhyloPlots v1.0.0
 (<u>https://github.com/cecileane/PhyloPlots.jl</u>) in Julia v1.9.2 (Bezanson *et al.*, 2012).

3

4 Divergence Time Estimation

We used a Bayesian Inference approach for divergence time estimation. To decrease 5 6 computational resources, we reduced the volume of the datasets to a subset of genes providing 7 the most useful information relevant to time calibration via a "gene shopping" method. 8 Specifically, we used the SortaDate package developed by Smith *et al.* (2018) to filter the 20 best 9 ortholog genes based on the (a) least topological conflict with a focal species tree (i.e., bipartition 10 calculation), (b) clock-likeness (i.e., root-to-tip variance statistic calculation), and (c) discernible information content (i.e., total tree length), sorting the genes in the respective order of these 11 12 properties (i.e., a, b, c).

We concatenated the resulting subset of genes and defined the positions of the 20 loci as data 13 14 blocks to find the best partitioning schemes and models of nucleotide evolution with PartitionFinder 2 (Lanfear et al., 2017) to inform our site model selection for the molecular 15 calibration. The branch lengths were estimated independently for each subset (i.e., unlinked), the 16 corrected Akaike's Information Criterion (AICc) was used to select the best-fit nucleotide 17 18 substitution models among those available in BEAST 2, while the "greedy" algorithm was used to search for a good partitioning scheme. The results suggested 18 partitioning sets, with 19 20 GTR+I+G4+X as the best-fit model for 17 sets and GTR+G4+X as the best-fit model for the 21 remaining one. For this reason and to reduce computational time, we decided to carry out molecular dating with the nucleotide substitution model GTR+I+G4+X without partitioning the 22 23 selected loci.

In detail, the analysis was performed using an unpartitioned scheme in BEAST v2.7.4 (Bouckaert 1 2 et al., 2019) under a relaxed uncorrelated lognormal clock model (Optimized Relaxed Clock, 3 ORC; Mean clock rate: 1.0), a GTR+I+G4+X site model and using a random starting tree for 200 4 million generations, sampling every 10,000 generations. Since no available fossil of slipper 5 orchids can be used for calibration and our sampling of orchids apart from Cypripedioideae is 6 scarce, we used a secondary calibration point based on the age estimates by Givnish *et al.* (2015). 7 The study above reconstructed a broad-scale phylogeny with species representing all orchid 8 subfamilies with 75 plastid genes and calibrated against 17 angiosperm fossils using BEAST v. 9 1.8.0 (Drummond *et al.*, 2012). Based on their estimates, we set a uniform distribution for the 10 crown age of Orchidaceae with the lower and upper bounds equal to 79.7 Ma and 99.5 Ma, 11 respectively, and used a Birth-Death tree model (Gernhard, 2008). The rest of the priors were not modified from their default values. Four identical runs with distinct seed numbers were 12 performed simultaneously to determine whether they converged on the same stationary 13 14 distribution. A fifth run was performed, sampling from the prior to examine whether the results 15 were significantly skewed by the prior assumptions or informed by our data.

Convergence of the Markov Chain Monte Carlo (MCMC) chains was checked with Tracer v1.7.2 (Rambaut *et al.*, 2018) by checking that the Effective Sample Size (ESS) of the combined runs was >200 for all trace statistics, and that the trace plots of the individual runs converged on the same posterior distribution. The tree files from the four independent runs were combined after removing 10% as burn-in using LogCombiner v1.8.2, and the maximum clade credibility chronogram was reconstructed using TreeAnnotator v1.8.2 with (maximum clade median node height and 95% highest posterior density (HPD) intervals.

23

1 Detection of Diversification Rate Shifts

We investigated if diversification rates changed throughout the evolutionary history of *Cypripedium* and whether there were significant rate shifts. To achieve this, we used BAMM v2.5 (Rabosky *et al.*, 2013), a program developed to model the dynamics of speciation and extinction on phylogenetic trees. It considers time-dependent (e.g., a lineage's age) and diversitydependent (e.g., the number of lineages in a clade) effects to quantify diversification rates using a reversible-jump MCMC approach.

8 Regarding the input tree used for this analysis, we modified the time-calibrated maximum clade 9 credibility tree obtained from the divergence time estimation analysis by removing all non-10 Cypripedium species, as well as taxon duplicates, hybrids, and varieties to avoid inflating the diversification rates. To account for non-random taxon sampling between the included 11 Cypripedium sections, section-specific sampling fractions were calculated based on the 12 classification by Frosch and Cribb (2012). The expected number of shifts was set to one, 13 14 following the recommendation for small trees with less than 500 tips. The priors on the initial 15 lambda, the lambda shift parameter, and the time mode for the speciation rate were calculated with the R package BAMMtools v2.1.10 (lambdaInitPrior = muInitPrior = 1.35, lambdaShiftPrior 16 = 0.05; Rabosky *et al.*, 2014), the segment length (segLength) was set to 0.1, and the rest of the 17 18 parameters were left as default. We ran four MCMC chains for 50 million generations and sampled every 1,000 generations. 19

Subsequently, we used BAMMtools to check whether the MCMC runs converged (ESS >200) and discarded the first 25% of samples as burn-in. Then we estimated the prior and posterior distributions, plotted the speciation rate through time and the set of distinct shift configurations that account for 95% of the probability of the data (i.e., 95% credible set, threshold = 5), checking which of these configurations had the maximum a posteriori (MAP) probability (aka
 best shift configuration).

3

4 Ancestral Range Estimation

We used the R package BioGeoBEARS (Matzke, 2013) to infer the biogeographic history of 5 6 *Cypripedium.* BioGeoBEARS reconstructs the ancestral geographic distributions on phylogenies 7 while testing for the best-fit model of range evolution. It replicates the basic assumptions of three widely used models in historical biogeography: DEC (Dispersal-Extinction-Cladogenesis; 8 9 Ree and Smith, 2008), DIVA (Dispersal-Vicariance Analysis; Ronquist, 1997) and BayArea 10 (Bayesian Inference of Historical Biogeography for Discrete Areas; Landis et al., 2013), 11 implementing them in a Maximum Likelihood framework to allow for direct comparison. 12 Together, these models allow for a broad range of processes, such as vicariance, sympatric speciation, range expansion, and contraction. They can also be combined with a founder-event 13 ("jump") speciation model specified with the parameter "j" (Matzke, 2014). 14

15 We conducted different BioGeoBEARS runs using DEC, DEC+J, DIVALIKE, DIVALIKE+J, 16 BAYAREALIKE, and BAYAREALIKE+J to find the best-fitting model. Again, we used a 17 modified version of the time-calibrated maximum clade credibility tree obtained from the 18 divergence time estimation analysis for the input tree. However, this time, hybrids and duplicated species were removed, but when multiple varieties were present, a single specimen per each 19 20 accepted variety, according to Frosch and Cribb (2012), was kept due to distinct distributions. Cypripedium amesianum Schltr. and the ambiguous C. macranthos var. alba were also kept since 21 22 they were not monophyletic with their presumably synonymous taxa [i.e., C. yunnanense Franch. and C. macranthos var. albiflorum Makino (now synonym of C. macranthos Sw. var. 23 *macranthos*), respectively], and therefore considered distinct taxonomic units for this analysis. 24

We also removed non-slipper orchid taxa because of scarce sampling. The taxon distributions 1 2 were based on Eccarius (2009), Frosch and Cribb (2012), SC Chen et al. (2013), and Walid et al. (2019). The distributions of C. amesianum and C. macranthos var. alba were considered the 3 4 same as their synonyms. A distance matrix was also used in the analysis to adjust the dispersal probabilities. The matrix included the distances between the closest points at the perimeters of 5 every area pair combination in kilometers, measured in Google Maps. When two areas were 6 7 adjacent, we set the distance between them to 1 km as recommended by the guidelines. To reduce 8 computational time, we decided on nine total areas and allowed up to five areas to be combined 9 in an ancestral range.

10 The areas were divided based on the current distribution of the taxa included in the analysis, their proximity, and their distinct floristic and topoclimatic characteristics (e.g., climate, precipitation, 11 elevation). South America (area A) was specified as a large distinct area since only a few 12 outgroup slipper orchids are restricted to its Northern part [i.e., Selenipedium aequinoctiale Garay 13 14 and Phragmipedium lindlevanum (R. H. Schomb. ex Lindl.); POWO, 2023]. We separated 15 Central America and Mexico (area B, containing section Irapeana) from South America at the Isthmus of Panama and from North America at the deserts to its north (i.e., Baja, Mojave, 16 Sonoran, and Chihuahuan deserts). North America was divided into three areas: the Northern area 17 18 E (colder to polar, humid climate), the Eastern area D (lower altitudes, higher humidity and precipitation than the Western area), and the Western area C (higher altitudes, lower humidity 19 20 and precipitation than Eastern area; Kottek et al., 2006; Schmidt, 2018; Xiao et al., 2020). All 21 three areas match the distributions of different Cypripedium species, with only five species found in area E, while the Great Plains in the middle of North America seemingly create a distribution 22 23 boundary for multiple *Cypripedium* species (Supplementary Data Fig. S1). The Mediterranean and Scandinavian regions were grouped with Western and Central Europe (area F) because only 24

C. calceolus occurs in all three areas (Eccarius, 2009; Frosch and Cribb, 2012; SC Chen et al., 1 2 2013; Walid et al., 2019). We split area F from Eastern Europe and Russia (area G) at the boundaries of the Sarmatic and Pontic-South Siberian floristic provinces according to Schroeder 3 4 (1998), as they have a more continental climate and match the limit of C. guttatum's distribution in the European continent (Pfadenhauer and Klötzli, 2020; Supplementary Data Fig. S1). Area G 5 6 was separated from the two Asiatic areas (namely, the Southeast Asian area I and the Northeast 7 Asian area H) due to the higher number of unique *Cypripedium* species occurring there, as well as their different climates and floristic provinces (Kottek et al., 2006; Fridley, 2008). The 8 9 Southeast and Northeast Asian areas are split around the Qinling Mountains-Huaihe River Line 10 (aka Qinling-Huaihe line), a natural topographic boundary that separates North temperate from 11 South tropical China (Y Hu et al., 2020), which also seems to create a boundary for the distribution of several *Cypripedium* species. To reduce the state space and thus the computational 12 time of the analysis, Taiwan was included in the same area as Southeast Asia (area I), and Japan 13 14 in the same area as Northeast Asia and Eastern Russia (area H) due to their proximity.

In addition to these runs, a separate analysis with only two specified areas—the Old and the New World—was performed for comparison. In this case, the same six models were tested, setting the maximum number of areas to two, but a distance matrix was not included. Finally, the best-fit models of both analyses were plotted on the maximum clade credibility tree.

19

20 RESULTS

21 Assembly and Orthology Inference

The number of total extracted loci per species (with ≥75% and ≥50% of the target length and
identity, respectively) ranged from 489 (*C. parviflorum* Salisb. var. *pubescens* (Willd.) O. W.

Knight, herbarium sample Nr. 69) to 906 (C. irapeanum La Llave & Lex., Oberhof sample Nr. 1 2 14) out of the 950 loci from the extended reference dataset. We recovered ~804 loci on average, 3 which is notably higher than the corresponding proportion of loci reported for slipper orchids in 4 the original publication by Eserman et al. [2021; i.e., 430 loci for Paphiopedilum exul (Ridl.) Rolfe and 533 loci for Phragmipedium longifolium (Warsz. & Rchb. f.) Rolfe]. Paralogs were 5 found in all samples, from 57 (C. irapeanum, Oberhof sample Nr. 14) to 1,845 (C. micranthum 6 7 Franch., Oberhof sample Nr. 22) with ~771 paralogs per sample on average. The orthology 8 inference resulted in 792 MO orthologs available for \geq 35 ingroup taxa. Further filtering for loci 9 with at least 50 out of the 74 retained taxa and at least 500 sites per locus reduced the number of 10 orthologs to 614. Unfortunately, most herbarium samples were removed as they generally yielded 11 fewer loci with shorter length coverage than the living specimens. In total, 63 Cypripedium 12 specimens representing 52 different taxa and 11 sections (Frosch and Cribb, 2012) were included in the final dataset (Fig. 1). 13

14

15 Inferred Species Phylogeny and Discordance

Both concatenation and coalescent-based phylogenetic reconstruction analyses recovered the 16 genus Cypripedium as monophyletic with maximum support (i.e., BS = 100, LPP =1; Fig. 2; 17 18 Supplementary Data Fig. S2). Additionally, both employed gene tree concordance analysis approaches for the ASTRAL species tree showed high concordance for the MRCA of 19 20 Cypripedium, with Phyparts identifying 521 informative concordant genes out of 554 and QS 21 giving full support (i.e., 1.0/-/1.0), indicating that all sampled quartet replicates support the node (Supplementary Data Fig. S3 and S4). The phylogenetic relationships among the slipper orchid 22 genera were congruent between the ASTRAL and IQ-TREE trees, with Cypripedium being the 23 sister to the rest. Within its sister clade, the plicate-leaved genus Selenipedium Rchb. f. was 24

recovered as sister to the clade of the conduplicate-leaved genera *Mexipedium* V. A. Albert & M.
 W. Chase, *Phragmipedium* Rolfe, and *Paphiopedilum*, with the two New World genera
 Mexipedium and *Phragmipedium* more closely related to each other than to the Old World
 Paphiopedilum.

All 11 sections were monophyletic within *Cypripedium*. On the other hand, the subsections 5 Cypripedium and Macrantha within sect. Cypripedium were non-monophyletic based on the 6 7 classification of Frosch and Cribb (2012). However, when considering the classification of SC 8 Chen et al. (2013), subsect. Macrantha was monophyletic whereas subsect. Cypripedium was 9 paraphyletic, and these results were consistent between the two phylogenies. Although the 10 species grouped in the two largest sister clades within sect. Cypripedium (clades I and II) did not 11 correspond to the species compositions of its two subsections, they matched the distribution of the species within them, with clade I only found in the New World and clade II in the Old World 12 (Supplementary Data Fig. S1). 13

14 Maximum support was recovered for most branches in both inferred phylogenies, except for 15 some branches along the backbone and within the sections Trigonopedia, Sinopedilum, Bifolia, and Cypripedium. Among the topologies with maximum support, high concordance, and 16 congruence between the two phylogenies was the placement of the Mesoamerican sect. Irapeana 17 18 being sister to the rest (LPP = 1; BS = 100; Phyparts: 521/554; QS: 1/-/1), followed by sect. Arietinum (LPP = 1; BS =100; Phyparts: 454/569; QS: 1/-/1). Additionally, sect. Subtropica, 19 20 following the classification by Frosch and Cribb (2012), was recovered as monophyletic (LPP =21 1; BS = 100; Phyparts: 431/550; QS: 1/-/1), including both sect. Wardiana and sect. Subtropica as described by SC Chen et al. (2013). The two fly-pollinated sections Sinopedilum and 22 23 Trigonopedia were also supported as most closely related to each other, which was congruent between the gene trees and between the ASTRAL and IQ-TREE trees (LPP = 1; BS = 100;
 Phyparts: 489/571; OS: 1/-/1).

Nevertheless, the remaining inter-sectional relationships showed decreased gene tree 3 4 concordance based on Phyparts and QS. For instance, although both inferred phylogenies supported the sister relationship between sect. Subtropica and the clade containing the 5 sections *Californica* Z. J. Liu, P. J. Cribb & Y. B. Luo and *Flabellinervia* (LPP = 1; BS = 100), 6 7 both gene tree concordance analyses indicated elevated levels of discordance for these nodes 8 (Phyparts: 68/531; OS: 0.15/0.77/0.99). The gene tree disagreement regarding the rest of the 9 section-level relationships is largely attributed to the five short-length backbone (Fig. 3a; 10 branches 3 to 7; Supplementary Data Fig. S5), with the proportion of informative concordant genes estimated by Phyparts falling between 2.4–22%, and QS indicating either weak or counter 11 support for these branches. This is likely to be the cause of the mismatched topology of sect. 12 Acaulia (Lindl.) C. Morren between the ASTRAL and the IQ-TREE trees, as it was recovered as 13 14 sister to the (Obtusipetala, (Bifolia, Cypripedium)) clade in the former and sister to the 15 (Sinopedilum, Trigonopedia) clade in the latter.

Gene tree heterogeneity was higher for nodes within sections than between sections, except for 16 the sect. Cypripedium (LPP = 1; BS = 100; Phyparts: 455/568; QS: 1/-/1). Although this section 17 18 was highly supported as monophyletic with high concordance, the number of informative concordant genes was lower than the number of discordant genes for almost all nodes within it. 19 20 Only two nodes within this clade showed substantial concordance; namely, the (C. henryi Rolfe, 21 C. segawai Masam.) clade (LPP = 1; BS =100; Phyparts: 321/383; QS: 1/-/1) and the node corresponding to the MRCA of clade II within the section (LPP = 1; BS = 100; Phyparts: 314/501; 22 23 QS: 1/–/1).

24

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1 the remaining discordant topologies, light grey denotes the proportion of gene trees with missing taxa, dark grey 2 denotes the proportion of uninformative gene trees. Tip symbols: filled circles "●" denote transcriptomes, unfilled 3 circles "O" denote genomes, unfilled stars "☆" denote specimens from herbarium M. Tips without symbols come 4 from living specimens of the Botanical Collection at Oberhof. Flowers of representative species from each section are displayed to the right. Species names corresponding to the flower pictures, from top to bottom: C. franchetii, C. 5 6 parviflorum var. pubescens, C. guttatum, C. flavum, C. acaule, C. lentiginosum, C. micranthum, C. japonicum, C. 7 californicum, C. subtropicum, C. plectrochilum, and C. irapeanum. See the legends of Figure 1 and the 8 Acknowledgements section for the credits of the flower pictures.

9

Regarding the monophyly at the species level, two species within sect. *Cypripedium* that included 10 numerous infraspecific taxa (i.e., varieties and one form), C. macranthos and C. parviflorum 11 Salisb., were consistently recovered as paraphyletic in both phylogenies, with C. franchetii 12 13 Wilson and C. yunnanense nested in the former, and C. kentuckiense C. F. Reed nested in the latter. Two pairs of synonyms (according to Frosch and Cribb, 2012) formed monophyletic 14 clades: that is, (C. parviflorum Salisb. var. makasin (Farw.) C. J. Sheviak, C. parviflorum Salisb. 15 16 var. parviflorum), and (C. subtropicum S. C. Chen & K. Y. Lang, C. singchii Z. J. Liu & L. J. Chen). However, C. amesianum was more closely related to C. froschii rather than its 17 synonymous species, C. yunnanense. Additionally, the ambiguous taxon C. macranthos var. alba, 18 which was presumed to be either C. macranthos var. albiflorum Makino (now a synonym of C. 19 20 macranthos var. macranthos) or C. macranthos var. album Mandl, was more closely related to the equally white-flowered C. macranthos var. rebunense (Kudo) Ohwi. Regarding the rest of the 21 species, only ASTRAL recovered them as monophyletic, grouping all conspecific specimens 22 retrieved from different sources (i.e., Botanical Collection at Oberhof, herbarium M, and SRA). 23 24 In contrast, IQ-TREE also produced non-monophyletic groupings of C. micranthum and C. bardolphianum W. W. Sm. & Farrer within sect. Sinopedilum, which matched their sequence 25 26 type (i.e., transcriptome or target enrichment data). Notably, the three included hybrids following

Frosch and Cribb (2012) were placed in the same clades as one or both of their putative parent
 taxa.

3

4 Anomaly Zone and Polytomy Test

5 The anomaly zone boundary estimations detected four pairs of nodes at the backbone of 6 *Cypripedium* that are in the anomaly zone [i.e., $y < \alpha(x)$], as well as the internode between the 7 MRCAs of the (Subtropica, (Californica, Flabellinervia)) clade and the (Californica, 8 Flabellinervia) clade (Fig. 3a; Supplementary Data Table S6 and Fig. S5). Additionally, several internode pairs within the two clades of sect. Cypripedium also fell into the anomaly zone 9 (Supplementary Data Table S6 and Fig. S5). This suggests that most of the gene tree 10 incongruence at these nodes could be explained by ILS. Interestingly, the nodes between the 11 12 MRCA of sect. Cypripedium and the MRCAs of its subclades I and II were not found in the anomaly zone despite elevated gene tree heterogeneity at these nodes, especially for the former 13 14 subclade, which could indicate that other factors are playing a role in this discordance, such as 15 hybridization.

The ASTRAL polytomy test failed to reject the null hypothesis that the branch length is 0 (i.e., pvalue > α ; $\alpha = 0.05$) for branches 5 and 6 (Fig. 3b). These branches were also found to be in the anomaly zone, while branch 5 also received a low LPP support (LPP = 0.44). This could explain the topology of sect. *Acaulia*, which varies between the ASTRAL and the IQ-TREE phylogenies, switching places within the potential polytomy. Additionally, branches within the subclades I and II of sect. *Cypripedium* found in the anomaly zone were also identified as soft polytomies (Supplementary Data Fig. S6).

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Figure 3: Results of the anomaly zone, polytomy, and gene duplication analyses annotated on the backbone of the
 collapsed ASTRAL species tree (branch numbers are shown in circles). (A) Anomaly zone test: Branches shown in

1 red are in the anomaly zone. (B) Polytomy test: Resulting p-values are annotated on the backbone branches. 2 Polytomies based on $\alpha = 0.05$ are shown in red. (C) Gene duplications: (Left) Percentages of duplicated genes are 3 annotated on the backbone nodes based on the two used filtering methods (i.e., min 50% bootstrap and local 4 topology). Branches with potential WGDs are shown in red (outlier duplication percentages > 20%). Known diploid 5 chromosome counts from species within each clade are shown to the right of the collapsed clades, with "?" marking 6 sections with unknown chromosome numbers [based on SC Chen et al. (2013) and Eccarius (2009)]. (Right) The 7 number of branches is plotted against the gene duplication percentages per filtering method. Outlier gene duplication 8 percentages correspond to branches 7 and 8 with potential WGDs. Key to collapsed clades: Cyp I = clade I of sect. 9 *Cypripedium* (see Fig. 2); *Cyp* II = clade II of sect. *Cypripedium* (see Fig. 2); *Bif* = *Bifolia*; *Obt* = *Obtusipetala*; *Aca* 10 = Acaulia; Tri = Trigonopedia; Sin = Sinopedilum; Fla = Flabellinervia; Cal = Californica; Sub = Subtropica. C. 11 *plectrochilum* and *C. irapeanum* represent sections *Arietinum* and *Irapeana*, respectively. OUT = Outgroups.

12

13 Gene Duplication and Whole Genome Duplication Events

We mapped gene duplications on the ASTRAL species tree using two filtering methods (i.e., 14 15 bootstrap and local topology filters; Fig. 3c; Supplementary Data Fig. S7). Most percentages of 16 duplicated genes reached a maximum of ~14%, from 0.12-14.3% using the bootstrap filter and 17 from 0.2–12.79% using the local topology filter. However, an outlier percentage of elevated gene duplications was identified by both methods (28.25% by the bootstrap filter and 26.67% by the 18 19 local topology filter), although it was mapped to two different nodes, namely, the MRCA of sect. 20 Cypripedium, or on the MRCA of the (Cypripedium, Bifolia) clade. The observed outliers could be caused by a WGD event, which could be further supported by the higher 2n chromosome 21 number count found in species from both sections compared to other sections (i.e., > 20 22 23 chromosomes).

24

25 Hybridization Networks

In our PhyloNet analysis, we tested for a maximum of one reticulation event for the phylogenetic network containing *C*. × *alaskanum*, with the "po" option enabled, one to ten events for the network with *C*. × *columbianum*, and one to nine events for the network with *C*. × *ventricosum* as the run for ten events has proven to be overly time-consuming. Each test produced ten optimal
output networks, and we plotted the total log probabilities of the most likely network from each
run for comparison (Supplementary Data Fig. S8).

4 To address our first question of whether the three known hybrid species are indeed supported as products of hybridization between their putative parent taxa by our molecular data, we plotted the 5 6 most likely phylogenetic networks that tested for one reticulation event (Fig. 4). Regarding the 7 test with C. \times alaskanum, the most likely network indicated one hybridization event (Fig. 4a; 8 total log probability ~ -550.38). However, our results suggested that C. guttatum is most probably 9 a hybrid between C. \times alaskanum ($\gamma = 0.84$) and an unsampled or extinct species that shares a 10 common ancestor with the (C. × alaskanum, C. yatabeanum) clade ($\gamma = 0.16$). The most likely hybridization network of C. \times columbianum on the other hand (total log probability = ~ -11 17545.08) supports its status as a hybrid between C. parviflorum var. pubescens and C. 12 montanum as described in Frosch and Cribb (2012), and SC Chen et al. (2013), with relatively 13 14 even inheritance probabilities (i.e., $\gamma = 0.48$ and 0.52, respectively; Fig. 4b). As for the network that includes $C. \times ventricosum$ (total log probability = ~ -88871.73), the results indicate that both 15 $C. \times ventricosum$ and C. calceolus are sister taxa that arose from a hybridization event between 16 C. macranthos var. rebunense ($\gamma = 40$) and C. shanxiense S. C. Chen ($\gamma = 60$; Fig. 4c). 17

To address our second question of whether other hybrids could be detected within the *Cypripedium* section, we plotted the most likely overall phylogenetic network from all runs that tested from one to nine or ten reticulation events for the networks including *C*. × *ventricosum* and *C*. × *columbianum*, respectively (Supplementary Data Fig. S8). The analyses that tested for up to nine hybridization events produced phylogenetic networks with the highest total log probabilities for both tests, identifying nine reticulation events for the network with *C*. × *ventricosum* (total log probability = ~ -88508.91) and seven reticulation events for the network with *C*. × *columbianum* (total log probability = ~ -17501.44; Supplementary Data Fig. S8d and S8e). These phylogenetic
networks indicate that in addition to *C*. × *columbianum*, *C*. × *ventricosum*, and *C. calceolus*,
which were already identified as hybrids in our previous tests, *C. froschii*, *C. calcicola* Schltr., *C. amesianum*, *C. shanxiense*, *C. macranthos* var. *macranthos*, *C. franchetii*, *C. candidum* Muehl.
ex Willd., *C. kentuckiense*, all sampled taxa of *C. parviflorum*, and some unsampled ancestral
species also constitute potential hybrids (Supplementary Data Fig. S8a and S8b).

7



Figure 4: Phylogenetic networks with the highest total log probabilities resulting from the PhyloNet analysis testing for one hybridization event for the extracted subclades with each of the three hybrids: (A) C. × *alaskanum*, (B) C. × *columbianum*, (C) C. × *ventricosum*. The branch lengths and inheritance probabilities of Network 1 were optimized under full likelihood (option "po"). The inheritance probabilities are shown for each parent hybrid edge (blue, solid = major hybrid edge; red, dotted = minor hybrid edge).

14

15 *Divergence Times*

The time-calibrated maximum clade credibility tree, which was produced with BEAST 2 using nuclear genes amounting to 59,202 sites, supported that the subfamily Cypripedioideae diverged from the rest of the slipper orchids close to the K-Pg boundary (64.84 Ma; 95% HPD 89.09–41.18 Ma) while genus *Cypripedium* split from the rest of the slipper orchids in the Oligocene (30.26 Ma; 95% HPD 46.05–17.24 Ma). The Mesoamerican section *Irapeana* was the
first to diverge within the genus, originating in the Early Miocene (20.27 Ma; 95% HPD 31.63–
11.57 Ma). Following this split, rapid diversification occurred in the middle Miocene (15.93–
13.74 Ma), giving rise to most *Cypripedium* sections or lineages from which the sections
diverged (15.93–9.29 Ma). Section *Cypripedium* bifurcated during the late Miocene (8.52 Ma;
95% HPD 13.29–4.78 Ma), producing its two subclades.

7

8 Diversification Rates

9 The BAMM analysis illustrated a pattern of an initial elevated net diversification rate during the 10 early stages of Cypripedium's evolution, which steadily declined through time (from 0.23 to 11 0.13) due to decreasing speciation rate (from 0.28 to 0.17; Fig. 5a). No significant rate shifts have been detected in the maximum a posteriori probability shift configuration (f = 0.9; Fig. 5b; 12 Supplementary Data Fig. S10), suggesting that a single macroevolutionary rate better explains the 13 14 diversification within *Cypripedium* over time. The second most frequently sampled configuration in the 95% credible set indicated one significant rate shift on the MRCA branch of the sister clade 15 of sect. Irapeana (f = 0.1), with speciation rate increasing sharply and slowing down towards the 16 17 tips. In both cases, the analysis showed that the internode pairs previously identified to be in the 18 anomaly zone indeed had higher diversification rates compared to the rate at later time points in the phylogeny, providing support to the hypothesis of ILS playing a role in their elevated levels 19 20 of discordance.

21

22 Historical Biogeography

23 Our comparison of biogeographic models in BioGeoBEARS showed that allowing founder-event

speciation (parameter "j") provided the best fit to our data. Namely, the DEC+J model (d =


1

Figure 5: Results of BAMM analysis in *Cypripedium*. (A) Rate variations in (top) speciation, (middle) extinction, and (bottom) net diversification through time, based on all samples in the posterior distribution (density shading on confidence regions). (B) Maximum a posteriori probability shift configuration represented as a phylorate plot showing variations in speciation rates (cooler colors = slow, warmer colors = fast) along each branch of the dated *Cypripedium* phylogeny (posterior median node height estimates of the divergence times in Ma are shown on the nodes). The clades are annotated with the first three letters of the name of each section.

0.0201; e = 0; x = -0.2711; j = 0.0299; LnL = -119.27) had the lowest AIC (246.5) and AICc 1 2 (247.4) scores for the first test with nine defined areas, while the BAYAREALIKE+J model (d = 0.0026; e = 0; j = 0.0747; LnL = -25.97) had the lowest AIC (57.94) and AICc (58.44) scores for 3 4 the second test with the New and Old World areas. The relative probabilities of the ancestral geographical ranges are illustrated with pie charts in Supplementary Data Fig. S11 and S12 for 5 the first and second test, respectively, while Fig. 6 combines the single most probable ancestral 6 7 ranges from both analyses with the results of the latter shown only for nodes where they disagree. 8 Regarding the results of the first test, the DEC+J model indicated that the ancestors of the 9 Cypripedioideae and Cypripedium clades were more widespread, distributed across the Old and 10 the New World (ranges ABI and BI, respectively; Fig. 6) and that potential long-distance 11 dispersals took place when Cypripedium diverged from the rest of the slipper orchids in the Oligocene, as well as when the Mesoamerican sect. Irapeana split from the Southeastern ancestor 12 of its sister clade in the Miocene. On the other hand, the BAYAREALIKE+J model implemented 13 14 in the second test supported the New World being the ancestral range of both Cypripedioideae 15 and *Cypripedium*, with the ancestor of sect. *Irapeana*'s sister clade speciating after its dispersal to the Old World. In both cases, the models suggested that the sister clade of sect. Irapeana rapidly 16 diversified in the Old World in the Middle Miocene, specifically in Southeast Asia (i.e., area "I"), 17 18 where most *Cypripedium* species are still found today.

Many of the lineages produced during these rapid diversification events dispersed and speciated
in other Old World regions, such as in Northeast Asia and the nearby islands of Japan and
Taiwan (e.g., *C. japonicum* Thunb. and *C. formosanum* Hayata). There were also multiple
independent dispersals back to the New World between the Miocene and the Pliocene (e.g., sect. *Acaulia*, sect. *Californica*; and MRCA of *C. reginae* Walter and *C. passerinum* Richardson),
while the MRCA of sect. *Bifolia* spread both Eastwards and Westwards, acquiring a wide



Figure 6: Estimations of ancestral ranges with the highest likelihood, plotted on the dated maximum clade credibility
 tree of slipper orchids and their state probabilities at the nodes. Results from both BioGeoBEARS runs (i.e., run with
 nine areas and with only New and Old World areas) are shown together when they disagree (below and above the

1 branch, respectively; the corresponding nodes/corners are marked with yellow stars); otherwise, only the results of 2 the former test are shown. Distribution on the nodes and branch corners are right before and right after cladogenesis, 3 respectively (the latter is not shown if it shares the same distribution with the following node). Key to area codes: A 4 = South America; B = Central America and Mexico; C = Western North America; D = Eastern North America; E = 5 Northern North America; F = Western and Central Europe, the Mediterranean, and Scandinavia; G = Eastern Europe 6 and Eurasia; H = Eastern Russia and Northeast Asia; I = Southeast Asia; N = New World; O = Old World. Plio. = 7 Pliocene; Quat. = Quaternary. Cyp I = clade I of sect. Cypripedium (see Fig. 2); Cyp II = clade II of sect. 8 Cypripedium (see Fig. 2).

9

distribution in both the Old (i.e., Eastern Europe, Eurasia, Northeast Asia, Japan, East Russia,
Southeast Asia) and the New World (i.e., Alaska), with the two species evolving sympatrically in
the broader sense.

The clades I and II of sect. *Cypripedium* diverged with the long-distance dispersal of the MRCA 13 of clade II to the New World during the Late Miocene. Following this, the two clades diversified 14 15 allopatrically, with subclade II spreading throughout North America, evolving several closely related taxa within the C. parviflorum complex. Within subclade I, some lineages and species 16 expanded their distribution from Southeast Asia to the adjacent area of continental Northeast Asia 17 18 [e.g., C. henryi; MRCA of (C. shanxiense, C. calceolus); C. fasciolatum Franch.] as well as the neighboring island of Taiwan [e.g., C. segawai, C. macranthos var. taiwanianum (Masam.) 19 Maekwa]. C. shanxiense also spread to the island of Japan, while C. calceolus speciated in 20 Northeast Asia, establishing populations in large latitudinal ranges as the only known species in 21 22 Western Europe, Scandinavia, and the Mediterranean. The ancestor of the sister clade of (C. yunnanense, C. macranthos var. taiwanianum) switched its distribution to Northeast Asia, from 23 where it probably further dispersed to Japan, producing multiple endemic varieties [e.g., C. 24 macranthos var. rebunense, C. macranthos var. speciosum (Rolfe) Koidz., and C. macranthos 25 26 var. *hotei-atsumorianum* Sadovsky] while C. *macranthos* var. *macranthos* also expanded to East, Asiatic, and European Russia. 27

1

2 DISCUSSION

3 Monophyly and Topology of Established Infrageneric Taxa

In the present study, we reconstructed the first robust phylogeny of the genus Cypripedium based 4 on high-throughput target enrichment data of 614 nuclear loci using the "Orchidaceae963" bait 5 set (Eserman *et al.*, 2021). The inferred phylogenetic tree showed that *Cypripedium* is sister to 6 the clade of the other four slipper orchid genera, (Selenipedium, (Paphiopedilum, 7 8 (*Phragmipedium*, *Mexipedium*))), which agrees with the topologies recovered by the supra-9 generic phylogenies of Guo et al. (2012), Wong and Peakall (2022) and Pérez-Escobar et al. 10 (2023) based on Sanger sequences (cpDNA and nDNA markers), transcriptomic data, and a 11 combination of target enrichment (low-copy nuclear loci) and Sanger sequences (matK and ITS), 12 respectively. All sections were monophyletic based on the classification system proposed by 13 Frosch and Cribb (2012) and SC Chen et al. (2013). However, subsect. Macrantha, which included the clade of C. farreri W. W. Sm. and C. fasciolatum as proposed by SC Chen et al. 14 15 (2013), was nested within subsect. Cypripedium.

16 Although the incongruence between the taxonomy and the monophyly of these subsections may 17 be affected by the elevated gene tree discordance within sect. *Cypripedium*, our results match the 18 findings of Szlachetko et al. (2021), who also evaluated the monophyly of these subsections based on the same two classification systems. Indeed, most published Cypripedium phylogenies 19 supported that one or both subsections may be non-monophyletic and that the (C. farreri, C. 20 21 fasciolatum) clade is more closely related to subsect. Macrantha rather than subsect. Cypripedium (Fatihah et al., 2011; J Li et al., 2011; H Liu et al., 2021a), as suggested by SC 22 Chen et al. (2013). Moreover, the species composition of the two clades within sect. Cypripedium 23

in our phylogeny matched their distributions, with clades I and II only found in the Old World
and the New World, respectively. A similar trend was observed in the phylogenies by Fatihah *et al.* (2011), J Li *et al.* (2011), and Szlachetko *et al.* (2021), where the division between the two
groups of species seemed to be based on their distribution rather than the traditionally used
morphological characteristics (i.e., the floral coloration and the shape of the labellum and the
lateral petals).

7 Within these two subclades, we recovered the two morphologically diverse species, C. 8 *parviflorum* and *C. macranthos*, as paraphyletic. Other authors also found that the latter is 9 paraphyletic, with C. kentuckiense embedded in the clade, similar to our phylogeny (Fatihah et 10 al., 2011; J Li et al., 2011; H Liu et al., 2021a; Szlachetko et al., 2021). However, even though C. yunnanense and C. franchetii were shown to be closely related to C. macranthos, with the 11 latter recovered monophyletic (Fatihah et al., 2011; J Li et al., 2011; H Liu et al., 2021a; 12 Szlachetko et al., 2021), our results showed that they are nested within the C. macranthos group, 13 which could have resulted due to the inclusion of different C. macranthos varieties (i.e., C. 14 15 macranthos var. alba and all five accepted natural varieties described by Frosch and Cribb, 2012). Cypripedium parviflorum and C. macranthos are widespread and morphologically 16 variable species, traditionally distinguished mainly based on flower size and coloration (SC Chen 17 18 et al., 2013; Cribb, 1997). Due to high morphological variation within their infraspecific taxa, as well as the existence of intermediate forms, they have been historically difficult to classify. This 19 20 difficulty could be attributed to the recent divergence of these varieties and forms within each 21 species that could cause ILS, in addition to hybridization, gene duplication, and WGD events, all occurring within sect. Cypripedium, according to our analyses. 22

Besides sect. *Cypripedium*, several supported topologies with high gene tree concordance
recovered in our phylogeny, are consistent with those in other studies. For instance, sect.

Irapeana, the only neotropical section of Cypripedium, was recovered as the sister to the rest in 1 2 the majority of published molecular phylogenies (Cox et al., 1997; Fatihah et al., 2011; J Li et al., 2011; Guo et al., 2012; H Liu et al., 2021a; Szlachetko et al., 2021) as well as in the present 3 study. The placement of sect. Irapeana is further supported by morphological and 4 biogeographical data, as it is considered to share "ancestral" morphological features with the 5 6 plicate-leaved genus Selenipedium, also found in the Neotropics (Cox et al., 1997; Szlachetko et 7 al., 2021). Similarly to the results of Cox et al. (1997), Fatihah et al. (2011), J Li et al. (2011), 8 and Szlachetko et al. (2021), sections Arietinum and Irapeana form a grade that is sister to the 9 remaining Cypripedium included in our phylogeny. Furthermore, sect. Trigonopedia and sect. 10 Sinopedilum, the two Southeast Asian fly-pollinated sections, previously grouped as a single section (Cribb, 1997), form a clade in our phylogeny. This sister relationship is consistently well-11 supported by other studies (Fatihah et al., 2011; J Li et al., 2011; H Liu et al., 2021a; Szlachetko 12 et al., 2021), while the infra-sectional topology we recovered for *Trigonopedia* matches that of H 13 14 Liu et al. (2021a).

Our molecular data also provides support for the classification of sect. Subtropica following 15 Frosch and Cribb (2012), as well as sect. Subtropica and Wardiana following SC Chen et al. 16 (2013). Specifically, two species have been described to belong to sect. Subtropica in the former 17 18 monograph, namely, C. subtropicum and C. wardii, with C. singchii being a synonym of C. subtropicum. However, SC Chen et al. (2013) redefined C. singchii as a distinct species and 19 20 transferred C. wardii to its own monotypic section, Wardiana. We showed that all three species 21 share an MRCA based on our molecular data, with C. subtropicum and C. singchii being more closely related to each other than to C. wardii, matching the corresponding topology in J Li et al. 22 23 (2011) and Szlachetko *et al.* (2021)

Concerning the remaining intersectional phylogenetic relationships, we uncovered high gene tree 1 2 discordance at several backbone nodes. Low backbone branch support has been observed in all 3 molecular phylogenies focusing on the section-level relationships within Cypripedium (Cox et 4 al., 1997; Fatihah et al., 2011; J Li et al., 2011; Szlachetko et al., 2021), except H Liu et al. (2021a), where four plastid regions were used for phylogenetic reconstruction. Our results 5 6 suggest that gene tree heterogeneity could explain why the evolutionary relationships between the 7 majority of the sections within Cypripedium remain unresolved, with different sets of loci and 8 phylogenetic inference methods producing incongruent topologies, such as the changing 9 placement of sect. Acaulia between the concatenation- and coalescent-based phylogenies in our 10 study.

11

12 Rapid Radiation, Whole Genome Duplication, and Hybridization Promoted Diversification

The concordance analyses indicated that most gene tree topologies disagree at certain nodes of 13 14 our phylogeny. High levels of discordance were particularly observed at nodes along the 15 backbone, the MRCA between sect. Subtropica and the (Californica, Flabellinervia) clade, and within the two subclades of sect. Cypripedium. Other authors (Fatihah et al., 2011; J Li et al., 16 2011; Szlachetko et al., 2021) have previously speculated that potential rapid radiation events 17 18 could interpret the low support at the branches along the backbone of their Cypripedium phylogenies, as well as the high morphological differentiation between the sections. Here, we 19 20 provided supporting evidence for this hypothesis based on our analyses of ~600 nuclear loci. 21 Specifically, our anomaly zone test showed that this incongruence largely owed to ILS caused by rapid radiation. This is further corroborated by the increased diversification rates at the 22 23 corresponding nodes and their chronologically close placement at the geological time scale.

1 The only intra-sectional nodes detected in the anomaly zone belonged to sect. *Cypripedium*, 2 indicating that rapid and recent diversification took place within its two subclades, leading to ILS 3 and producing multiple closely related species and infra-specific taxa with high morphological 4 variation (e.g., within the *C. macranthos* and *C. parviflorum* complexes). However, we showed 5 that other factors could have also contributed to the incongruence between the gene and the 6 species trees observed within this section and throughout the phylogeny. For instance, another 7 source of mixed gene tree signals could be reticulation.

8 The heterogeneous gene topologies at the MRCA nodes of the two subclades I and II of sect. 9 *Cypripedium* could be mainly explained by hybridization and WGD, as they have not been 10 shown to fall within the anomaly zone, contrary to the nodes within these subclades. As 11 suggested by Unruh et al. (2018), an increased taxon sampling in Cypripedium, which is the genus with the widest genome size range (4.1–43.1 pg/C) among all slipper orchids, revealed a 12 potential WGD event. Specifically, we located a WGD either at the MRCA branch of sect. 13 *Cypripedium*, or of the (*Cypripedium*, *Bifolia*) clade, and widespread gene duplications within the 14 15 phylogeny, using the subclade orthogroup tree topology method. These factors could contribute 16 to gene tree discordance (Górecki and Eulenstein, 2014) and diversification by producing 17 paralogous genes with the potential to develop novel functions. WGDs, in particular, are 18 considered one of the main driving forces of species-, and in some cases, niche-diversification, as polyploids may occupy new or wider ranges of niches in contrast to their diploid relatives 19 20 (Dodsworth et al., 2016; Ren et al., 2018). Thus, a WGD event could have promoted the higher 21 diversity observed within sect. *Cypripedium*, as well as the widespread distribution of both sect. Cypripedium and Bifolia, which contain species with the widest latitudinal (i.e., C. guttatum) and 22 23 longitudinal (i.e., C. calceolus) ranges in the genus, found in a variety of different altitudes (i.e., 50 to 4100 m and sea level to 2000 m, respectively; Frosch and Cribb, 2012; SC Chen *et al.*,
 2013).

The time interval between the WGD and the diversification of sect. Cypripedium and sect. Bifolia 3 4 could be interpreted by the 'WGD Radiation Lag-Time' model, where some lineages within a clade sharing an ancestral WGD expand millions of years following the event. This has been 5 6 proposed to occur not only due to the time needed for the subfunctionalization or 7 neofunctionalization but also for the right ecological conditions to uncover their adaptive 8 advantage (e.g., migration or climatic changes) or for re-diploidization to occur, as neopolyploids 9 are more likely to go extinct and less likely to speciate than diploids (Schranz et al., 2012; Tank 10 et al., 2015; Dodsworth et al., 2016; Robertson et al., 2017). On the other hand, this lag time 11 could also be a consequence of extinction events taking place between the divergence of these sections and their diversification, which we also observed in other sections of our phylogeny 12 (e.g., sect. Flabellinervia and sect. Obtusipetala). 13

Additionally, the increased number of chromosomes within sections *Cypripedium* (2n = 20-36)and *Bifolia* (2n = 20-30), compared to the usual diploid count in the genus (2n = 20), provides further evidence to support the existence of a potential WGD at either of the identified branches (Eccarius, 2009; SC Chen *et al.*, 2013). In particular, the species *C. macranthos* has the widest range of recorded chromosome counts (i.e., 20, 21, 30, 36) in the genus. These numbers nearly match different ploidy levels, from the usual 2n (20) to less than 4n (36).

The elevated chromosome count in these two sections may have also resulted from allopolyploidization, as hybrids are observed in both clades. All nodes leading to the hybrids identified through our phylogenetic network analyses had decreased concordance levels, including the MRCA node of the (*C. guttatum*, *C.* × *alaskanum*) clade. Although *C.* × *alaskanum* has been previously described as a natural hybrid between *C. yatabeanum* and *C. guttatum* by PM Brown (1995), it was difficult to assess its validity because no analysis accompanied the
 description (Cribb, 1997). We found that, most likely, *C. guttatum* is a hybrid between *C.* ×
 alaskanum and an unsampled species sharing an MRCA with the (*C.* × *alaskanum*, *C. yatabeanum*) clade.

As for the hybrids within sect. *Cypripedium*, testing for a maximum of one hybridization event yielded both expected and unexpected results. Sheviak (1992) proposed that $C. \times columbianum$ is a hybrid between *C. montanum* and *C. parviflorum* var. *pubescens* based on an extensive survey and analysis of wild and herbarium specimens, with hybrids having intermediate morphological characteristics between the presumed parent taxa. The phylogenetic network containing $C. \times$ *columbianum* agreed with Sheviak's taxon description, with both proposed parent taxa having almost equal inheritance probabilities.

Nonetheless, the results of a similar test on the dataset containing C. × ventricosum supported 12 that the taxon did not constitute a hybrid from a cross between C. calceolus and C. macranthos as 13 it was previously suggested based on the resemblance of its flower with an artificial hybrid 14 15 between the presumed parent taxa (Rolfe, 1904, 1910; Cribb, 1997). Instead, the results indicated that C. \times ventricosum and C. calceolus are products of a hybridization event between C. 16 shanxiense and C. macranthos. Another hybrid, C. × catherinae, which occurs in Far East Russia 17 18 (Siberia) and possibly in Korea and Northeast China, where the parent taxa's distributions overlap, was already described from the hybridization between these two taxa (Frosch and Cribb, 19 20 2012; SC Chen et al., 2013). Cypripedium calceolus and C. \times ventricosum can also be found sympatrically with the parent taxa identified in our analysis, namely, in Far Eastern Russia 21 22 (Siberia) and Northeastern China, while C. calceolus is also found in Japan and on Sakhalin Island (Cribb, 1997; Frosch and Cribb, 2012; SC Chen et al., 2013). Interestingly, a sister 23 relationship is consistently recovered between C. calceolus and C. shanxiense (Fatihah et al., 24

2011; J Li *et al.*, 2011; H Liu *et al.*, 2021*a*; Szlachetko *et al.*, 2021) or between *C. calceolus* and
 C. macranthos in the absence of *C. shanxiense* (Cox *et al.*, 1997). Additionally, *C. × ventricosum* and *C. macranthos* formed a clade closely related to the (*C. calceolus*, *C. shanxiense*) clade in the
 phylogeny of H Liu *et al.* (2021*a*).

In support of Rolfe's description, a statistical analysis of morphological and allozyme data from 5 $C. \times ventricosum$ corroborated its status as a hybrid between C. calceolus and C. macranthos 6 7 (Knyasev et al., 2000). However, the fact that C. calceolus and C. \times ventricosum may share the same parent taxa based on our results (i.e., C. shanxiense and C. macranthos) and that 8 9 introgressive hybridization has been previously reported between C. calceolus and C. \times 10 ventricosum (Knyasev et al., 2000) could elucidate the relationship between these species. 11 Furthermore, it has been shown that the genetic structure of C. shanxiense was similar to C. *calceolus* from the eastern part of the range where they are sympatric, with the authors arguing 12 that it may have been a result of introgressive hybridization, making the classification of these 13 taxa even more challenging (Filippov and Andronova, 2011). At the same time, a hybrid has also 14 been described between these two species (i.e., $C. \times microsaccos$ Kraenzl; Frosch and Cribb, 15 16 2012; SC Chen et al., 2013).

17 When looking at the most probable phylogenetic networks produced for our second test, the 18 results suggested that extensive hybridization events between sampled and unsampled taxa occurred within both subclades of sect. Cypripedium. Specifically, the patterns of reticulate 19 20 evolution in these networks indicated that the hybrids that persisted interbred with other taxa 21 (incl. other hybrids) and backcrossed with their parent taxa, producing a hybrid swarm. The mixed genetic and morphological signals created by these extensive and overlapping reticulation 22 23 events could have significantly contributed to the difficulty in specific delimitation and taxonomic classification within the sect. Cypripedium, with the same taxa receiving a species, 24

variety, or hybrid rank by different taxonomists. For example, a self-pollinating taxon that has 1 2 been reported as a synonym of C. yunnanense, C. amesianum (Frosch and Cribb, 2012; SC Chen 3 et al., 2013; P Cribb, Royal Botanical Garden, Kew, UK, 'pers. comm.'), was more closely 4 related to C. froschii and C. tibeticum King ex Rolfe in our phylogeny. Our results show that C. 5 amesianum may constitute a hybrid between unsampled parent taxa, one of which is closely 6 related to C. tibeticum and the other to C. yunnanense. Additionally, C. froschii, the nature of 7 which is still strongly debated (e.g., variety of C. tibeticum according to Eccarius, 2009; species 8 status according to Frosch and Cribb, 2012; C. tibeticum \times C. yunnanense hybrid according to SC 9 Chen et al., 2013) seems also to be a result of hybridization between the latter parent taxon of C. 10 amesianum, and a taxon within an unsampled sister clade of C. franchetii. However, an ancestor 11 of the latter parent taxon also experienced hybridization between an unsampled taxon and C. 12 macranthos.

Further studies are necessary to confirm the hybridization events that we identified in the evolutionary history of sections *Cypripedium* and *Bifolia*. Unfortunately, we could not test for reticulation events at a broader scope within *Cypripedium* as the phylogenetic network analysis becomes highly computationally intensive and time-consuming with increasing taxa and hybridization events. However, as hybridization is considered to be common in nature between sympatric *Cypripedium* species, we could assume that it might have also played a part in the discordance observed at other nodes of the genus' phylogeny.

20

21 Climatic Fluctuations Influenced Current Distribution and Diversity Patterns

Previous primary molecular calibrations placed the stem age of Cypripedioideae at 77 Ma
(Givnish *et al.*, 2016) or between ca. ~75–82 Ma (based on the youngest and oldest fossil ages,
respectively; Guo *et al.*, 2012). Our estimation places the divergence time of the slipper orchids

at a more recent time point around the K-Pg boundary (64.84 Ma), a time that was followed by rapid radiations of flowering plants at the genus and family level (e.g., Koenen *et al.*, 2021; Morales-Briones *et al.*, 2021), as they replaced some of the old lineages and filled empty niches left by the mass extinction (Berendse and Scheffer, 2009). Similarly to the fossil-assisted calibration of (Givnish *et al.*, 2016) and a secondary calibration based on the divergence time estimates above (H Liu *et al.*, 2021*a*), we dated the split of *Cypripedium* from its sister clade to ~30 Ma in the Oligocene.

8 During that time, according to our biogeographic analysis, the ancestors of the slipper orchids 9 either had a wider distribution across the New and the Old World or were initially limited to the 10 New World. Such uncertainty was also noticed in the genus-level Cypripedioideae phylogeny of 11 Guo et al. (2012), who identified either the New World or the Old World + the New World as the ancestral regions of slipper orchids using both S-DIVA and Lagrange methods. A biogeographic 12 analysis of a broader Orchidaceae phylogeny that included placeholders from all five orchid 13 14 subfamilies (representing 18/19 tribes and 40/43 subtribes), all genera of slipper orchids, and 96 15 outgroup angiosperms suggested that the ancestors of Cypripedioideae were in the Neotropics 16 with >75% probability. This result agreed with the analysis of H Liu et al. (2021a), whose 17 estimates pinpointed Central America as the origin of slipper orchids despite a different topology 18 recovered at the genus level. The more extensive taxon-sampling of outgroup taxa within non-*Cypripedium* slipper orchids, as well as other orchids and angiosperms in Givnish *et al.* (2016), 19 20 would have likely better informed the estimations of the ancestral region of the subfamily Cypripedioideae compared to our outgroup sampling, therefore, providing support to our second 21 22 biogeographic analysis indicating the New World as the MRCA.

A similar dichotomy was observed in the results of our two biogeographic tests regarding the
most likely ancestral regions of the MRCA of *Cypripedium* following its divergence from the rest

of the Cypripedioideae (i.e., either Central America + Southeast Asia or the New World). Again, 1 2 the results of Guo et al. (2012) resembled ours at the corresponding node, which either 3 recognized both the Old World + the New World (S-DIVA) or the Old or New World (Lagrange) 4 as the ancestral regions of *Cypripedium*. This ambivalence was also reflected in the ancestral region estimations of Givnish et al. (2016), supporting that Neotropics + Eurasia was the most 5 probable distribution of the ancestor of Cypripedium, followed by the Neotropics. Contrarily, H 6 7 Liu et al. (2021a) found that the most probable ancestral area of slipper orchids was Central 8 America, while the second most probable was Central America + South Asia. Thus, despite 9 evidence favoring both scenarios, a wider ancestral distribution across the Old and the New 10 World received higher support in most studies. However, it is important to note that due to 11 incomplete taxon sampling (i.e., 11 out of 13 Cypripedium sections following Frosch & Cribb, 2012, and few outgroups) as well as the discordance in the backbone of the phylogeny, these 12 results should be interpreted with caution. 13

14 A wider ancestral distribution could have been facilitated by the Bering Land Bridge acting as a 15 dispersal corridor between the New and the Old World, as it was intermittently exposed 16 throughout most of the Cenozoic (Hopkins, 1959). Additionally, a significantly warmer climate 17 during the Early to Middle Miocene Climatic Optimum (MMCO; ca. 17–15 Ma), especially at 18 higher latitudes where mixed deciduous-evergreen forests dominated the vegetation type (Dutton and Barron, 1997; Herold et al., 2010; Steinthorsdottir et al., 2021), coupled with adaptations to 19 20 colder climates after diverging from the neotropical sister clade (H Liu *et al.*, 2021*a*), could have 21 allowed ancestors of the genus *Cypripedium* to spread northwards and disperse via the Bering Land Bridge or over the strait. The current distribution of sect. *Bifolia* at both sides of the Bering 22 23 Strait (i.e., Eastern Siberia and Alaska) supports the potential of *Cypripedium* to disperse via this corridor. 24

1 As global cooling took place following the MMCO (Frigola et al., 2018; Methner et al., 2020; 2 RM Brown et al., 2022), a climatic dispersal barrier may have formed between the Old and the 3 New World at higher latitudes, leading to a vicariance and allopatric diversification. The East 4 Asian monsoon system, whose development was associated with the global cooling and the uplift of the Tibetan Plateau in the Early Miocene, may have promoted Cypripedium's dispersal to 5 South Asia (H Lu and Guo, 2014; Hui et al., 2021; Ye et al., 2022). The Middle Miocene Climate 6 7 Transition (MMCT; 15-13 Ma), a period of climate transition between the MMCO and the 8 Middle Miocene Glaciation (MMG) associated with major Antarctic ice sheet expansion and 9 global cooling (Frigola et al., 2018), coincided with the rapid radiation of Cypripedium in 10 Southeast Asia during the Middle Miocene (~16–14 Ma), according to our results. Interestingly, our dating of the divergence time between sect. Irapeana and its sister clade (~21 Ma) coincides 11 with the stem ages of most seed plant clades originating in the Sino-Japanese floristic region (Y-12 S Chen et al., 2018), while the identified period of the rapid diversification events in the 13 mountains of China (~16-14 Ma) fits the observed trend of large-scale plant diversifications in 14 15 W. China during the Miocene (L-M Lu et al., 2018).

After dispersing to Southeast Asia, the time interval between the origin and the diversification of 16 17 this new clade (i.e., stem age: 20.27 Ma; crown age: 15.93 Ma) indicates that extinctions might 18 have taken place, which could explain why the ancestors of this lineage were limited to that region according to our analysis. The southeastern margins of the Tibetan Plateau may have 19 20 provided potential refugia to *Cypripedium* from the cooling climate of the MMCT due to their warmer and more humid conditions owing to the intensification of the Asian monsoons (Xing 21 22 and Ree, 2017; S Li et al., 2020; Zuo et al., 2022). Specifically, the isolation of the Cypripedium population in deep valleys between mountains, such as the Hengduan mountains, a biodiversity 23 and endemism hotspot harboring the most species of Cypripedium (H Liu et al., 2021a; J Liu et 24

al., 2022), could have promoted allopatric diversification at a highly dissected landscape from 1 2 which several sections or lineages leading to sections diverged, producing the observed pattern of 3 rapid radiations along the backbone of our phylogeny (Favre *et al.*, 2015; Xing and Ree, 2017; 4 Spicer et al., 2020). Additionally, the continuous uplift post-Middle Miocene and the orogeny of the mountains at the southeastern margins of the Tibetan Plateau may have provided a diverse 5 6 variety of habitats throughout its wide altitudinal range, promoting adaptive radiation (López-7 Pujol et al., 2011; Wang et al., 2012; Xing and Ree, 2017), while the relatively stable conditions 8 of the Hegduan mountains, which glaciated minimally, could have allowed time for speciation to 9 take place (Spicer et al., 2020; H Liu et al., 2021a).

10 As mentioned before, the cooling climate during the Middle to Late Miocene glacial events 11 (Hansen et al., 2013; Frigola et al., 2018) may have prevented the dispersal of many lineages back to the New World through the Bering Land Bridge, besides the most cold tolerant lineages 12 such as sect. *Bifolia* and sect. *Acaulia* (H Liu *et al.*, 2021*a*). In this case, "jump" dispersals across 13 14 the Pacific Ocean may have been a more likely route back to the New World for some lineages, as suggested by our biogeographic analysis (e.g., sect. Californica, and New World clades of 15 sections *Obtusipetala* and *Cypripedium*). Such long-distance dispersals could be attributed to 16 17 their small, dust-like seeds that lack endosperm, as their seedlings solely depend on their 18 mycorrhizal associates for nutrition until they become photosynthetically active (Arditti and Ghani, 2000). Due to the size of the orchid seeds, dispersal models indicated that they could 19 20 travel a distance of up to 2,000 km by wind (Arditti and Ghani, 2000). However, evidence of 21 endozoochory was also found for some orchid species (Suetsugu et al., 2015; Suetsugu, 2020; Y 22 Zhang et al., 2021), while dispersal via epizoochory (e.g., on bodies of mammals and birds) and water currents has been suggested, although it is still uncertain whether orchids seeds could 23 24 germinate after long exposure to salt water (Arditti and Ghani, 2000).

1 The glacial cycles of the Late Pliocene and the Ouaternary may have promoted the infra-sectional 2 diversification of *Cypripedium*. The isolation of populations in refugia, such as in the mountains at the margins of the Tibetan Plateau, the islands of Japan and Taiwan (Tang et al., 2018), as well 3 4 as numerous areas in North America, including the Californian coast and the Klamath Mountains 5 (DR Roberts and Hamann, 2015) would have allowed survival and allopatric diversification 6 throughout glacial periods. During the interglacial periods, population expansions resulting in 7 overlapping distributions could have led to hybridization and introgression, further increasing 8 genetic and morphological diversity (e.g., in the C. macranthos/C. tibeticum complex; P Cribb, 9 Royal Botanical Garden, Kew, UK, 'pers. comm.'). The hybrid swarms and the high number of 10 taxa endemic to Japan and Taiwan within sect. Cypripedium underpin this hypothesis.

11

12 *Conclusion*

The phylogenetic relationships within the genus *Cypripedium* have remained generally 13 14 unresolved despite numerous phylogenies based on different molecular markers and phylogenetic 15 reconstruction methods, largely due to the low support and conflicting topologies at nodes 16 between and within sections. Our study provided a new perspective on the causes of this 17 phenomenon, using high-throughput sequence data from over 600 nuclear loci to gain more insight into the evolution of this genus. Although all established sections were supported as 18 monophyletic, we uncovered potential rapid diversification events that led to incomplete lineage 19 20 sorting at the backbone of the tree. A similar pattern was also observed within the two subclades 21 of the largest section, *Cypripedium*, which did not correspond to its two subsections following 22 Frosch and Cribb (2012) and SC Chen et al. (2013) but matched their respective geographic 23 distributions. Additionally, a whole genome duplication and multiple hybridization events were linked to the sections *Cypripedium* and *Bifolia*. All these sources of gene tree discordance likely
 produced mixed phylogenetic signals, preventing the accurate resolution of *Cypripedium*'s
 phylogeny.

4 The events above were associated with climatic transitions that led to major ancestral distribution changes. After the slipper orchids likely originated in the Neotropics around the K-Pg boundary, 5 6 the milder climate of the Early Miocene could have allowed them to disperse to the Old World 7 via the Bering Land Bridge. The Middle Miocene Climatic Transition coincided with the rapid 8 radiations following *Cypripedium*'s dispersal to Southeast Asia, possibly due to isolation in local 9 refugia. Several independent dispersals back to the New World via the Bering Land Bridge or 10 across the Pacific Ocean likely took place, while the glacial-interglacial cycles probably played a 11 role in the further speciation and reticulate evolution observed in *Cypripedium*.

12

13 SUPPLEMENTARY DATA

14 Table S1: Selected examples of taxonomic revisions for the infrageneric classification of Cypripedium L. Table S2: Cypripedium specimens sampled from the Botanical Collection at 15 Oberhof associated with the BGM and the herbarium M. Table S3: List of publicly available 16 17 orchid genomes and transcriptomes used in this study. Table S4: Modifications to the Macherey-18 Nagel NucleoSpin Plant II kit: Genomic DNA from plant (Macherey-Nagel – 07/2014, Rev.09) protocol. Table S5: Final ortholog occupancy statistics per specimen after MO orthology 19 inference. Table S6: The $\alpha(x)$ calculations of the anomaly zone test with the corresponding 20 branch numbers and their branch lengths. Figure S1: Distribution of *Cypripedium* species per 21 section. Figure S2: The concatenation-based phylogeny of the genus Cypripedium, inferred with 22 23 614 nuclear loci using IQ-TREE 2. Figure S3: The output phylogeny of Cypripedium from the 24 Phyparts analysis. Figure S4: The output Cypripedium phylogeny from the Quartet Sampling

analysis. Fig. S5: The ASTRAL phylogeny of *Cypripedium* annotated with the corresponding 1 2 branch numbers referred to in Supplementary Table S6 for the results of the anomaly zone test 3 calculations and in Figure 3. Figure S6: Results of the ASTRAL polytomy test on the ASTRAL 4 phylogeny of *Cypripedium*. Figure S7: Results of the gene duplication test based on the subclade 5 orthogroup tree topology method implemented with the ASTRAL phylogeny of Cypripedium, 6 using the (A) bootstrap and the (B) local topology filters. Fig. S8: Supplementary figure for the 7 PhyloNet analysis. Figure S9: The maximum clade credibility tree of *Cypripedium* resulting from the molecular analysis with BEAST 2. Figure S10: Supplementary figures from the BAMM 8 9 analysis of the *Cypripedium* phylogeny. Figure S11: The relative proportions of the ancestral 10 ranges estimated by BioGeoBEARS (nine area test, DEC+J model) and plotted as pie charts on 11 the corresponding nodes (i.e., right before cladogenesis) or branch corners (i.e., right after cladogenesis) on the dated maximum clade credibility tree of slipper orchids. Figure S12: The 12 relative proportions of the ancestral ranges estimated by BioGeoBEARS (two area test, 13 14 BAYAREALIKE+J model) and plotted as pie charts on the corresponding nodes (i.e., right before cladogenesis) or branch corners (i.e., right after cladogenesis) on the dated maximum 15 clade credibility tree of slipper orchids. 16

17

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20

21 DATA AVAILABILITY

Target enrichment data generated for this study can be found in the NCBI BioProject XXXXX
(please refer to Table XXX for SRA accession numbers). Analyses files are available from the
Dryad repository https://doi.org/10.5061/dryad.XXXX

1

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18

19 CONFLICT OF INTEREST

20 The authors declare that they have no competing interests.

21

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