



Two new combinations of *Middletonia* (Gesneriaceae) in China

Qi-Yang Li¹, Xin-Xiang Bai¹, Song-Tao He¹¹ Forestry College, Guizhou University, CN-5500252 Guiyang, ChinaCorresponding author: Song-Tao He (sthe@gzu.edu.cn)

Abstract

In 2016, systematic studies of the tribe Loxocarpinae A.DC. supported the establishment of the new genus *Middletonia* C. Puglisi. This study was based on inference and parsimony analyses of a phylogenetic tree derived from the nuclear ITS and plastid *trnL-trnF* regions, constructing the most recent phylogeny that includes the genera *Middletonia* and *Paraboea*. The results showed that the current generic delimitation within the tribe differs from the clades delineated by phylogenetic analysis, and both morphological and molecular evidence support treating the Hainan endemics *Paraboea changjiangensis* Xing & Z.X.Li and *P. hainanensis* (Chun) Burtt as species of *Middletonia*. Therefore, this paper re-delimits the genera *Middletonia* and *Paraboea*, aiming to establish a more natural classification, and proposes the new combinations *Middletonia changjiangensis* (F.W.Xing & Z.X.Li) X.X.Bai, **comb. nov.**, and *Middletonia hainanensis* (Chun) X.X.Bai, **comb. nov.**, providing a valuable framework for the development and comparative studies of the molecular systematics of the family Gesneriaceae.

Key words: Loxocarpinae, *Middletonia changjiangensis*, *Middletonia hainanensis*, morphology, *Paraboea*, phylogeny

Introduction

Middletonia C. Puglisi belongs to the subtribe Loxocarpinae A.DC. (1845) of Gesneriaceae. In 2016, Puglisi et al. used molecular data from 68 species of five genera within the subtribe Loxocarpinae to construct a phylogenetic tree. They taxonomically treated *Middletonia evrardii*, *M. monticola*, and *M. multiflora*, which were originally classified in *Paraboea*, and established the new genus *Middletonia*. They also mentioned that this genus shares similarities in floral morphology with *Paraboea*, which requires further investigation (Puglisi et al. 2016). In the latest taxonomic revision framework, *Middletonia* comprises five species, with their fruit types being twisted or erect (Puglisi and Middleton 2017; GRC 2025).

Paraboea (C.B. Clarke) Ridl., established in 1905, also belongs to the subtribe Loxocarpinae of Gesneriaceae (Ridley 1905; Xu et al. 2008). It is the most species-rich genus within this subtribe and has fruit types that are twisted or erect (Weber et al. 2013; GRC 2025). Although *Paraboea* has undergone multiple systematic revisions (Wang et al. 1990, 1998; Li and Wang 2005; Xu et al. 2008; Wei et al. 2010), many taxonomic problems remain unresolved. Thereafter, Guo (2016) conducted a comprehensive study on the genus *Paraboea* in China using



Academic editor: Eberhard Fischer

Received: 4 March 2025

Accepted: 9 July 2025

Published: 5 August 2025

Citation: Li Q-Y, Bai X-X, He S-T (2025) Two new combinations of *Middletonia* (Gesneriaceae) in China. *PhytoKeys* 261: 33–43. <https://doi.org/10.3897/phytokeys.261.151963>

Copyright: © Qi-Yang Li et al.

This is an open access article distributed under terms of the Creative Commons Attribution License (Attribution 4.0 International – CC BY 4.0).

molecular systematics combined with morphological traits. The results revealed that *Paraboea hainanensis* (Burt 1984) and *Paraboea changjiangensis* (Xing and Li 1993) are significantly distinct from the morphological characteristics of *Paraboea*, with oblique tubular corollas and distinct upper and lower lips. It was proposed that these two species be treated as a new genus, “Huàn Yōng Jù Tái Shǔ” (Guo 2016).

At present, the taxonomic statuses of *Paraboea hainanensis* and *Paraboea changjiangensis* remain unclear due to limitations in molecular sampling and uncertainty regarding their resource backgrounds. Xu et al. (2017) published three new species of *Paraboea*, using *Middletonia* as an outgroup to construct a phylogenetic tree, but *P. changjiangensis* and *P. hainanensis* were not included in that analysis. Subsequently, Wang et al. (2022) used the chloroplast genomes of 12 *Paraboea* species to construct a phylogenetic tree to verify the monophyly of *Paraboea*, which also did not include *P. changjiangensis* and *P. hainanensis*. However, in the report of the new species *P. zunyiensis*, *P. hainanensis* was found to cluster separately, which does not support the monophyly of *Paraboea* (Deng et al. 2023).

In order to further resolve the complex evolutionary events and developmental relationships, we employed the internal transcribed spacer (ITS) region of nuclear ribosomal DNA (nrDNA) and the *trnL-trnF* intergenic spacer region of chloroplast DNA (cpDNA) to test whether the current classification is consistent with the phylogenetic structure of this group. We aimed to identify robust phylogenetic entities suitable for redefining generic boundaries, address intergeneric and intrageneric issues, and complement the taxonomic treatment with morphological characteristics.

Materials and methods

Morphological comparisons

In 2023, during a botanical survey in Guizhou Province and Hainan Province, China, the species *Paraboea hainanensis* and *Paraboea changjiangensis* were found. Subsequently, some living specimens were introduced and cultivated at Guizhou University for further research. We obtained information on *P. hainanensis*, *P. changjiangensis*, and their related species from the Internet, including descriptions in original literature (Chen 1974; Xing and Li 1993) and relevant literature (Brown 1839; Xu et al. 2008), as well as geographic distribution data from the Global Biodiversity Information Facility (GBIF, <https://www.gbif.org>). In addition, we consulted digital plant specimens collected by E, IBK, PE, NY, GH, and IBSC to examine the type specimens and high-resolution images, as well as other specimens. Finally, we conducted a preliminary taxonomic treatment of the genus by combining existing morphological data and molecular systematic evidence.

Genomic DNA extraction, PCR amplification, and sequencing

Species leaf samples collected from the place of origin were quickly dried with silica gel for DNA extraction (Chase and Hills 1991). The nuclear ribosomal internal transcribed spacer (ITS) and chloroplast DNA sequences (*trnL-trnF*) of these samples were amplified by polymerase chain reaction (PCR) using the primers described in Taberlet et al. (1991) and White et al. (1990). All DNA samples were sent to Sangon Biotech Co. Ltd. (Shanghai, China) for sequencing and splicing.

Phylogenetic analysis

In the phylogenetic analysis, the ingroup consisted of 33 species from six genera of the Loxocarpinae subtribe, including 26 species of the genus *Paraboea*. Two species from the genus *Petrocodon* in the Didymocarpinae subtribe, *Petrocodon ainsliifolius* and *P. viridescens*, were selected as outgroups (Suppl. material 1).

All sequences were compared using MAFFT v.7.5.1.1 (<https://mafft.cbrc.jp/alignment/server/>) (Kato and Standley 2013), conserved regions were selected using Gblock, and the substitution saturation index (Iss) of the data matrix was evaluated using DAMBE v5.3.19 (Xia 2013), which showed the $Iss < Iss.c$, $P = 0.0000$ of the data matrix < 0.05 , which is not saturated, can be used to construct phylogenetic trees. Multi-gene syndication was performed using Model Finder software, and polygenes were performed in PAUP*4.0 b10 (Swofford and Sullivan 2003). Thereafter, in PhyloSuite v.1.2.3 (Kalyaanamoorthy et al. 2017; Zhang et al. 2020; Xiang et al. 2023), for ML analysis and BI inference. The *trnL-trnF* and ITS sequences were concatenated in series in the Concatenate Sequence module of PhyloSuite v.1.2.3 (Zhang et al. 2020), and the optimal base substitution models for four partitions of the combined dataset were determined in the PartitionFinder2 module using the corrected Akaike Information Criterion (AICc). The best-fitting evolutionary models were GTR+G (*trnL-trnF*) and GTR+I+G (ITS). The ML method employed 1,000 bootstrap replicates to assess the reliability of each node in the phylogenetic tree. The BI rule uses the best substitution model for different segments and independently estimates the Bayesian posterior values of each segment's parameters, with a random tree as the starting tree, and the initial setting runs for 100,000,000 generations, with one tree reserved for every 10,000 generations. The first 25% of the trees are discarded as burn-in, and the remaining trees are used to generate consensus trees and calculate Bayesian posterior probabilities. Finally, use the iTOL v4 version of the online tool (<https://itol.embl.de>) to beautify the phylogenetic tree.

Results

Morphological comparisons

Based on field surveys and specimen examinations, morphological comparisons were conducted between *Paraboea changjiangensis* and *Paraboea hainanensis* with the type species of *Middletonia* and *Paraboea*, respectively (Table 1). Like *Paraboea*, *Middletonia* presents a matted, interwoven indumentum of long and fine hairs on the abaxial side of the leaf, a flat-faced corolla, and a capsular fruit (Puglisi and Middleton 2017). Combining morphological descriptions from specimens and original literature, the two species can be well distinguished from *Paraboea* based on four morphological characters: corolla, filament, anther, and style morphology, as well as ovary indumentum (Table 1, Fig. 1).

In addition, we compared the collected *Paraboea hainanensis* with the digital specimens (isotypes E00265039, NY00074065, GH00025112) from PPBC and GBIF, as well as the morphological descriptions in relevant literature (Fig. 2, Xu et al. 2008). Sessile leaves are one of the main characteristics of *Paraboea hainanensis*, which are present in all three digital specimens. Moreover, the felt-like indumentum on the lower leaf surface is thick, tightly appressed, and reddish-brown,

Table 1. Comparison among *Middletonia multiflora*, *Paraboea changjiangensis*, *P. hainanensis*, and *P. sinensis*.

Characters	<i>M. multiflora</i>	<i>P. changjiangensis</i>	<i>P. hainanensis</i>	<i>P. sinensis</i>
Leaf blade shape	oblong or ovate, 7–10 × 4–6 cm, apex rounded-obtuse, base cuneate	elliptic or oblong, 2–7 × 1–3 cm, apex obtuse, base cuneate	narrowly oblanceolate to oblanceolate, 8–18 × 3–6 cm, apex rounded, base gradually attenuate	oblong, oblanceolate or lanceolate, 5.5–25 × 2.4–9 cm, apex acute, base cuneate or broadly cuneate.
Petiole	3–5 cm long, with a matted indumentum	1–2.5 cm long, with a matted indumentum	leaves sessile	3–6 cm long, tomentose brown
Corolla	almost flat-faced	almost flat-faced	almost flat-faced	obliquely campanulate
Anthers	erect	erect	erect	borne at a right angle
Style shape	linear	linear	linear	upper part swollen and saccate, lower part curved and attenuate
Ovary indumentum	grayish-white waxy powder	farinose glandular	some minute pubescence	glabrous

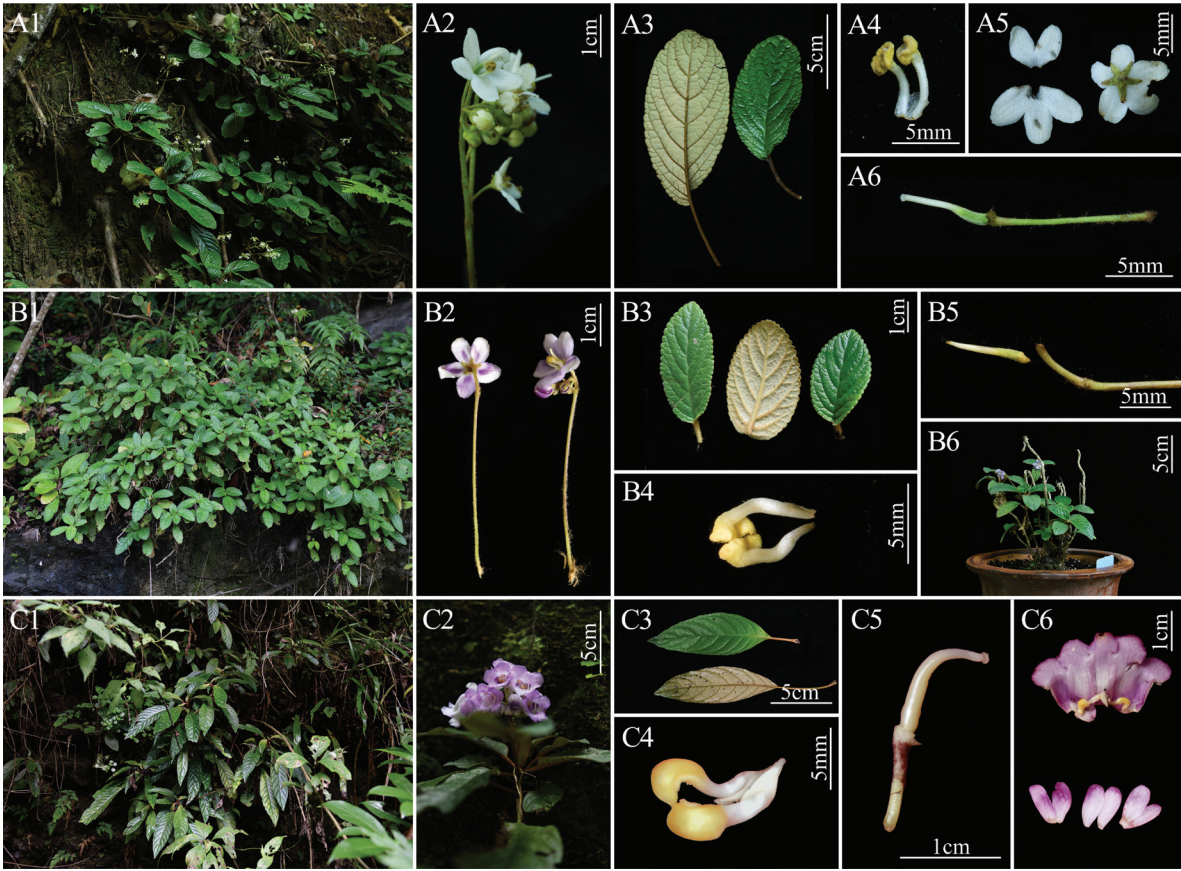


Figure 1. The morphologically and molecularly related species *Middletonia multiflora*, *Paraboea changjiangensis*, and *P. sinensis*. A1–6. *M. multiflora*; A1. Habit; A2. Cymes; A3. Petiole and abaxial leaf surfaces; A4. Filaments and anthers; A5. Top view of the expanded corolla tube and calyx; A6. Pistil; B1–6. *P. changjiangensis* B1. Habit; B2. Cymes; B3. Petiole and abaxial leaf surfaces; B4. Filaments and anthers; B5. Calyx; B6. Plants; C1–6. *P. sinensis*; C1. Habit; C2. Cymes; C3. Petiole and abaxial leaf surfaces; C4. Filaments and anthers; C5. Pistil; C6. Top view of opened corolla showing the interior surface of the corolla tube, stamens, and staminodes (photographed by Xin-Xiang Bai).

which is morphologically identical to that of the digital specimens. Furthermore, we carefully compared the leaf shape and size and observed that the leaves of the collected specimens are mostly narrow obovate, rarely narrow elliptic, 2.5–4 times as long as wide, with minute serrations on the margin, which is highly consistent with the images in PPBC and the records in Xu et al. (2008) (Fig. 2).

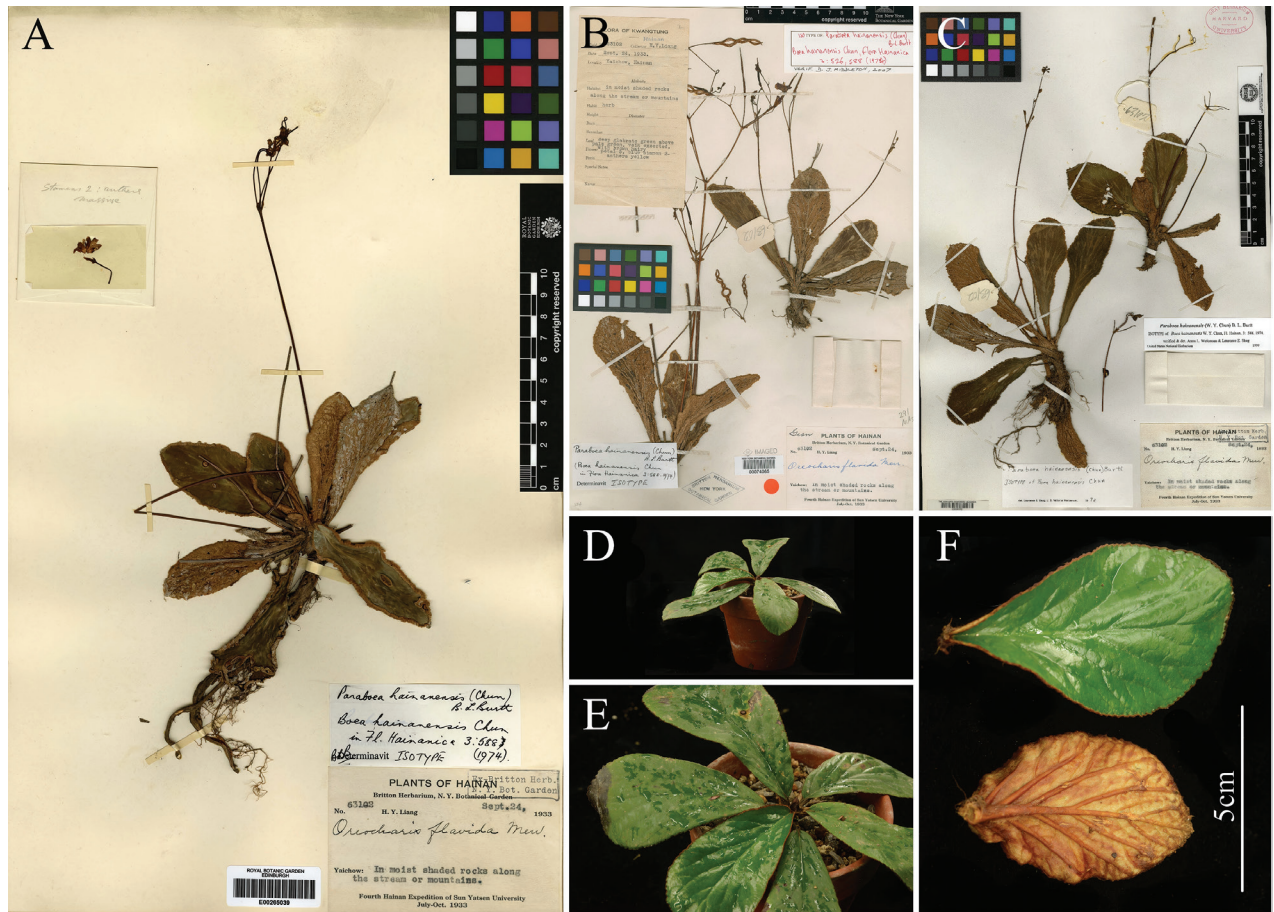


Figure 2. Morphological comparisons of *Paraboea hainanensis*. A. E00067459; B. NY00074065; C. GH00025112; D. Front view of the plant; E. Top view of the plant; F. Leaves (Images D–F were photographed by Qi-Yang Li).

Plastome genome features

Through combined parsimony and Bayesian inference analyses of ITS and *trnL-F* (Fig. 3), the results showed that in Clade 1, *Paraboea changjiangensis* was nested within *Middletonia* with high support (posterior probability [PP] = 1.00, bootstrap [BS] = 98%); *Paraboea hainanensis* also formed a strongly supported sister group relationship with *Middletonia* (posterior probability [PP] = 0.97, bootstrap [BS] = 75%). This indicates that classifying these two species in *Paraboea* is incorrect. Additionally, Clade 2 included all species except *Paraboea hainanensis* and *P. changjiangensis*, with high support (posterior probability [PP] = 1.00, bootstrap [BS] = 97%). Therefore, the taxonomic positions of *Paraboea hainanensis* and *P. changjiangensis* are inappropriate, and both should be classified within *Middletonia* (Fig. 3).

Taxonomic treatment

Based on the outcomes of phylogenetic and morphological research, both *Paraboea changjiangensis* and *P. hainanensis* are incorporated into *Middletonia*. The new combinations of *Middletonia* are provided below.

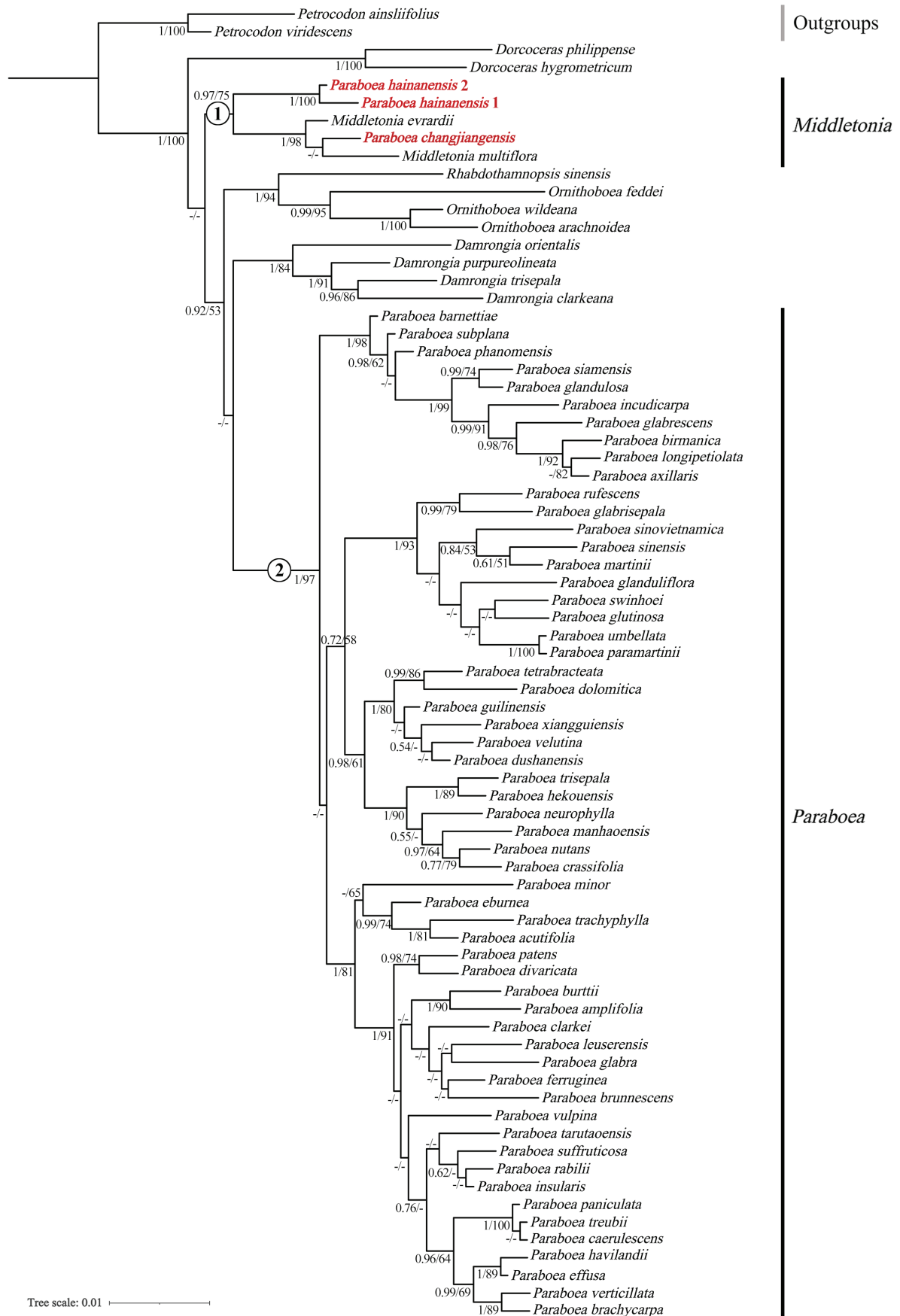


Figure 3. Bayesian tree from analysis of combined ITS and *trnL-F* data. The posterior probabilities (PP) of BI and bootstrap values (BS) of ML are listed at each node. A dash denotes branches with support rates below 50%.

***Middletonia changjiangensis* (Xing & Z.X.Li) X.X.Bai, comb. nov.**

[urn:lsid:ipni.org:names:77366493-1](https://nomenclature.ipni.org/names/77366493-1)

Fig. 4

≡ *Paraboea changjiangensis* Xing & Z.X.Li in *Acta Botanica Yunnanica* 15(2): 121–122, f. 1. 1993. Type: China, Hainan: Changjiang County, Wangxia, 600 m, 25 July 1989, Z.X.Li & F.W.Xing 5134 (holotype: IBSC!).

Distribution and habitat. Hainan Province: Changjiang County and Dongfang County. The species grows on calcareous formations at an elevation of 600 m.

Vernacular name. Chāng Jiāng Fěn Máo Jù Tái (Chinese pronunciation); 昌江粉毛茛苔 (Chinese name).

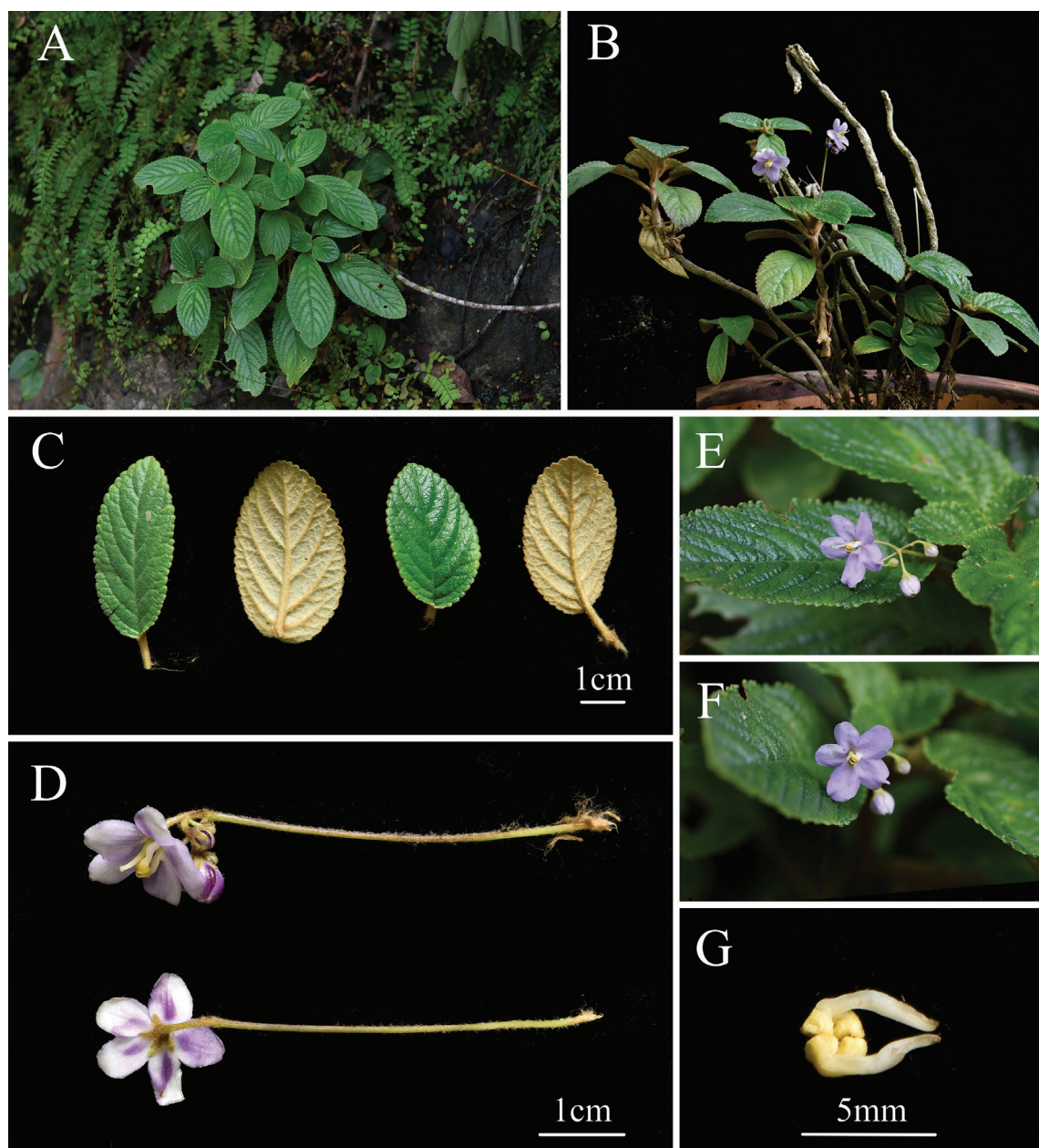


Figure 4. *Middletonia changjiangensis*. A. Habit; B. Plants; C. Petiole and abaxial leaf surfaces; D. Cymes; E–F. Plant with flowering; G. Filaments and anthers (photographed by Xin-Xiang Bai).

Representative specimens examined. CHINA. Hainan Province: • Changjiang County, 25 Jun 1975, *Guoyuan Fu* 264 (IBSC); • Changjiang County, 5 May 1988, *Zexian Li et al.* 4167 (IBSC).

***Middletonia hainanensis* (Chun) X.X.Bai, comb. nov.**

[urn:lsid:ipni.org:names:77366494-1](https://nomenclature.ipni.org/names/77366494-1)

Fig. 2D–F

≡ *Boea hainanensis* Chun in *Flora Hainanica* 3:526, 588, f. 903. 1974. Type: China, Hainan, Yaichow, on moist shaded rocks along streams in mountains, 18°30'N, 109°08'E, 24 September 1933, *H. Y. Liang* 63102 (isotype: E!, NY!, GH!).

≡ *Paraboea hainanensis* (Chun) Burt in Notes from the Royal Botanic Garden, Edinb 41(3): 429. 1984.

Distribution and habitat. Hainan Province: Dongfang County. The species grows on moist, shaded rocks along the stream or mountains at an elevation of 800 m. Reported as occurring on acid soil.

Vernacular name. Hǎi Nán Fěi Máo Jù Tái (Chinese pronunciation); 海南粉毛茛苔 (Chinese name).

Representative specimens examined. CHINA. Hainan Province: • Ledong County, Aug 1985, *Xinqi Liu* 27407 (PE!); • Dongfang County, Aug 1985, *Shaoqing Chen* 11251 (IBSC!); • Changjiang County, 0–800 m elev., 25 Jun 1975, *Guoyuan Fu* 264 (IBSC!); • Changjiang County, 1400 m elev., 20 Aug 2004, *The Kadoorie Project Team of Hong Kong* 6632 (PE!); • Hainan, 26 Sep 1933, *Xiangri Liang* 63162 (IBK!); • Hainan, 24 Sep 1933, *H. Y. Liang* 63102 (E!, NY!).

Discussion

In this study, from the perspective of macroscopic morphology, *Paraboea changjiangensis* and *Paraboea hainanensis* are most closely related to *Middletonia*. Additionally, molecular phylogenetic analyses clarified the phylogenetic relationships between *Paraboea* and *Middletonia*, showing that *P. changjiangensis* is nested within *Middletonia* and *P. hainanensis* forms a strongly supported sister group with *Middletonia*. Future research should focus on resolving the relationships among clades. Additionally, in modern systematics, the integration of molecular and morphological data is indispensable. More molecular markers should be introduced, with priority given to conserved chloroplast region sequences, so as to provide strong support for the backbone topology of the phylogenetic tree. Therefore, we formally rename the two species as *Middletonia changjiangensis* and *Middletonia hainanensis*, contributing to a more comprehensive phylogenetic framework of Gesneriaceae.

Acknowledgments

We would like to thank Dr. Xu Xiao for providing the distribution site of *Middletonia multiflora* in Wangmo County, Guizhou, and Dr. Ming-Zhong Huang for providing the materials of *Paraboea changjiangensis*. We are also grateful to the above-mentioned herbaria for images/photos of available specimens.

Additional information

Conflict of interest

The authors have declared that no competing interests exist.

Ethical statement

No ethical statement was reported.

Use of AI

No use of AI was reported.

Funding

This study was supported by the National Natural Science Foundation of China (grant 32260782), the Special Fund for Innovation Capacity Construction of Guizhou Research Institution (Qiankehefuqi [2024]013), and the 2024 Guizhou Science and Technology Innovation Talent Team Construction Project: Wildlife Innovation Team of the Forestry College of Guizhou University (Qiankeherencai CXTD [2025]053).

Author contributions

Data curation: QYL,STH. Investigation: XXB, STH. Writing – original draft: QYL. Writing – review and editing: XXB,STH.

Author ORCIDs

Qi-Yang Li  <https://orcid.org/0009-0006-9336-7718>

Xin-Xiang Bai  <https://orcid.org/0000-0003-2449-6664>

Song-Tao He  <https://orcid.org/0009-0005-5223-7522>

Data availability

All of the data that support the findings of this study are available in the main text or Supplementary Information.

References

- Brown R (1839) On Cyrtandreae. From Horsfield's "Plantae Javanicae rariores" 120.
- Burt BL (1984) Studies in the Gesneriaceae of the Old World: XLVII. Revised generic concepts for *Boea* and its allies. Notes from the Royal Botanic Garden Edinburgh 41: 401–452.
- Chase MW, Hills H (1991) Silica gel: An ideal material for field preservation of leaf samples for DNA studies. Taxon 40(2): 215–220. <https://doi.org/10.2307/1222975>
- Chen HY (1974) Flora Hainanica. The BMJ 3(526): 588–903. <https://doi.org/10.1136/bmj.3.5930.588>
- Deng T, Wen F, Xie DJ, Wei RX, He L, Dou LQ, Qian ZM, Zhang RB (2023) *Paraboea zunyiensis* (Gesneriaceae), a new species from north Guizhou, China. PhytoKeys 235: 21. <https://doi.org/10.3897/phytokeys.235.111412>
- GRC (2025) Gesneriaceae Resource Centre, Royal Botanic Garden Edinburgh. <https://padme.rbge.org.uk/grc/> [Retrieved/Accessed: 21 January 2025]
- Guo J (2016) Phylogenetic study of the genus *Paraboea* in China. Doctoral dissertation, Guangxi Normal University.

- Kalyaanamoorthy S, Minh BQ, Wong TK, Haeseler A, Jermini LS (2017) ModelFinder: Fast model selection for accurate phylogenetic estimates. *Nature Methods* 14(6): 587–589. <https://doi.org/10.1038/nmeth.4285>
- Katoh K, Standley DM (2013) MAFFT multiple sequence alignment software version 7: Improvements in performance and usability. *Molecular Biology and Evolution* 30: 772–780. <https://doi.org/10.1093/molbev/mst010>
- Li ZY, Wang YZ (2005) Plants of Gesneriaceae in China. Henan Sciences & Technology Publishing House, Zhengzhou, 170–303.
- Puglisi C, Middleton DJ (2017) A revision of *Middletonia* (Gesneriaceae) in Thailand. *Thai Forest Bulletin (Botany)* 45(1): 35–41. <https://doi.org/10.20531/tfb.2017.45.1.07>
- Puglisi C, Yao TL, Milne R, Möller M, Middleton DJ (2016) Generic recircumscription in the Loxocarpinae (Gesneriaceae), as inferred by phylogenetic and morphological data. *Taxon* 65: 277–292. <https://doi.org/10.12705/652.5>
- Ridley HN (1905) The Gesneraceae of the Malay Peninsula. *Journal of the Straits Branch of the Royal Asiatic Society* 43: 1–92. <https://www.jstor.org/stable/41561022>
- Swofford DL, Sullivan J (2003) Phylogeny inference based on parsimony and other methods using PAUP*. *The phylogenetic handbook: a practical approach to DNA and protein phylogeny* 7: 160–206.
- Taberlet P, Gielly L, Pautou G, Bouvet J (1991) Universal primers for amplification of three non-coding regions of chloroplast DNA. *Plant Molecular Biology* 17(5): 1105–1109. <https://doi.org/10.1007/BF00037152>
- Wang WT, Pan KY, Zhang ZY, Li ZY, Tao DD, Yin WC (1990) Gesneriaceae. In: Wang WT (Ed.) *Flora Reipublicae Popularis Sinicae*. Vol. 69. Science Press, Beijing, 460–472.
- Wang WT, Pan KY, Li ZY, Weitzman AL, Skog LE (1998) Gesneriaceae. In: Wu ZY, Raven PH (Eds) *Flora of China* Vol. 18. Science Press, Beijing, and Missouri Botanical Garden Press, St. Louis, 362–370.
- Wang YF, Wen F, Hong X, Li Z, Mi Y, Zhao B (2022) Comparative chloroplast genome analyses of *Paraboea* (Gesneriaceae): Insights into adaptive evolution and phylogenetic analysis. *Frontiers in Plant Science* 13: 1019831. <https://doi.org/10.3389/fpls.2022.1019831>
- Weber A, Clark JL, Möller M (2013) A new formal classification of Gesneriaceae. *Selbyana* 31(2): 68–94. <http://www.jstor.org/stable/24894283>
- Wei YG, Wen F, Möller M, Monroe A, Zhang Q, Gao Q, Mou HF, Zhong SH, Cui C (2010) Gesneriaceae of South China. Guangxi Science & Technology Publishing House, Nanning, 606–645.
- White TJ, Bruns T, Lee S, Taylor J (1990) Amplification and direct sequencing of fungal ribosomal RNA genes for phylogenetics. *PCR Protocols: a guide to methods and applications* 18(1): 315–322. <https://doi.org/10.1016/B978-0-12-372180-8.50042-1>
- Xia XH (2013) DAMBE5: A comprehensive software package for data analysis in molecular biology and evolution. *Molecular Biology and Evolution* 30(7): 1720–1728. <https://doi.org/10.1093/molbev/mst064>
- Xiang CY, Gao FL, Jakovlić I, Lei HP, Hu Y, Zhang H, Zou H, Wang GT, Zhang D (2023) Using PhyloSuite for molecular phylogeny and tree-based analyses. *iMeta* 2(1): e87. <https://doi.org/10.1002/imt2.87>
- Xing XF, Li ZX (1993) A new species of *Paraboea* from Hainan Island. *Acta Botanica Yunnanica* 15(2): 121–122. <https://journal.kib.ac.cn/EN/Y1993/V15/I02/1>
- Xu ZR, Burtt BL, Skog LE, Middleton DJ (2008) A revision of *Paraboea* (Gesneriaceae). *Edinburgh Journal of Botany* 65: 161–347. <https://doi.org/10.1017/S0960428608005106>

- Xu W, Guo J, Pan B, Han M, Liu Y, Chung KF (2017) Three new species of *Paraboea* (Gesneriaceae) from limestone karsts of China based on morphological and molecular evidence. *Botanical Studies* 58: 1–14. <https://doi.org/10.12705/652.5>
- Zhang D, Gao FL, Jakovlic I, Zou H, Zhang J, Li WX, Wang GT (2020) PhyloSuite: An integrated and scalable desktop platform for streamlined molecular sequence data management and evolutionary phylogenetics studies. *Molecular Ecology Resources* 20(1): 348–355. <https://doi.org/10.1111/1755-0998.13096>

Supplementary material 1

Supplementary information

Authors: Qi-Yang Li, Xin-Xiang Bai, Song-Tao He

Data type: xlsx

Copyright notice: This dataset is made available under the Open Database License (<http://opendatacommons.org/licenses/odbl/1.0/>). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Link: <https://doi.org/10.3897/phytokeys.261.151963.suppl1>