

Research Article

Phalaenopsis zhanhuoensis (Orchidaceae, Vandeae), a new species from Yunnan, China

Shiyu Qin^{1,2,3}, Hanchen Wang^{2,3,4}, Yajun Wang^{2,3,4}, Chongbo Ma^{2,3}, Zan Li^{2,3}, Boyun Yang¹, Xiaohua Jin^{2,3}

1 School of Life Sciences, Nanchang University, Xuefudadao 999, Shajing, Nanchang, Jiangxi, 330031, China

2 State Key Laboratory of Plant Diversity and Prominent Crops, Institute of Botany, Chinese Academy of Sciences, Nanxincun 20, Xiangshan, Beijing, 100093, China

3 China National Botanical Garden, Beijing, 100093, China

4 University of Chinese Academy of Sciences, Beijing, 100093, China

Corresponding authors: Boyun Yang (yangboyun@163.com); Xiaohua Jin (xiaohuajin@ibcas.ac.cn)

Abstract

A new species of Orchidaceae, *Phalaenopsis zhanhuoensis*, from Xichou County, Yunnan, China, is described and illustrated. The novelty is close to *P. taenialis*, *P. wilsonii*, and *P. stobartiana*, but differs from them by having a distinct, fleshy anterior callus with a deeply lobed apex at the base of the labellum and lateral lobes of labellum reflexed and facing outward.

Key words: China, new species, Phalaenopsis zhanhuoensis, Xichou County, Yunnan

Introduction

The moth orchid genus, *Phalaenopsis* Blume, comprises approximately 80 recognized species (https://powo.science.kew.org/) and is extraordinarily prominent in the field of horticulture. *Phalaenopsis* is distributed in India, Southeast to East Asia, and Australia, with most of the diversity in Indonesia and the Philippines (Pridgeon et al. 2014). Recent molecular results based on ITS nrDNA and plastid regions (*trnL* intron, *trnL-F* spacer, and *atpB-rbcL* spacer) indicated that the number of pollinia was not a good morphological character to distinguish *Phalaenopsis* from its alliance, such as *Doritis* Lindl., *Kingidium* P.F.Hunt., and *Nothodoritis* Z.H. Tsi, and proposed broadening *Phalaenopsis* to include its alliance (Christenson 2001; Padolina et al. 2005; Tsai et al. 2005; Yukawa et al. 2005; Tsai et al. 2010; Pridgeon et al. 2014). *Phalaenopsis* s.I. is characterized by roots more or less depressed and verrucose, stem short, leaves usually elliptic and fleshy, lip three lobed, column usually with column foot, pollinia two or four.

Based on molecular data and morphological characters, such as the presence or absence of column foot and the number of pollinia, *Phalaenopsis* was subdivided into four subgenera, subgen. *Parishianae* (Sweet) Christenson (26 spp.), subgen. *Phalaenopsis* Blume (45 spp.), subgen. *Hygrochilus* (Pfitzer) Kocyan & Schuiteman (5 spp.) and subgen. *Ornithochilus* (Lindl.) Kocyan & Schuiteman (4 spp.) (Kocyan and Schuiteman 2014; Pridgeon et al. 2014; Higgins and Alrich 2015).



Academic editor: M. Simo-Droissart Received: 6 September 2023 Accepted: 31 December 2023 Published: 22 January 2024

Citation: Qin S, Wang H, Wang Y, Ma C, Li Z, Yang B, Jin X (2024) *Phalaenopsis zhanhuoensis* (Orchidaceae, Vandeae), a new species from Yunnan, China. PhytoKeys 237: 153–160. https://doi. org/10.3897/phytokeys.237.112270

Copyright: © Shiyu Qin 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). The subgenus *Parishianae* is mainly distributed in India, Southeast to East Asia (Pridgeon et al. 2014), with species morphologically characterized by small plant size, few-flowered inflorescence, small scarious floral bracts, and biseriate callus (Pridgeon et al. 2014). There are 25 *Phalaenopsis* species in four subgenera in China (Zhou et al. 2021; Ma et al. 2022), of which 14 species belong to the subgenus *Parishianae* (Chen and Wood 2009; Zhou et al. 2021). During our fieldwork in Yunnan Province, China, in April 2023, a new species of *Phalaenopsis* belonging to subgenus *Parishianae* (Sweet) Christenson was found in evergreen broad-leaved forests and is described here.

Materials and methods

Morphological characters of the new species were observed, measured with a ruler (precision: 1 mm), and photographed based on living plants. Molecular phylogenetic analyses were conducted using one nuclear (nrITS) and four plastid markers (matK, trnL, trnL-F, and atpB-rbcL). Genomic DNA was extracted from the newly collected specimen of Phalaenopsis (silica dried) using the modified cetyltrimethylammonium bromide (CTAB) method (Li et al. 2013). Sequencing library was generated using Rapid Plus DNA Lib Prep Kit for Illumina and then delivered to Novogene Company (Beijing, China) for 150 bp paired-end sequencing on the Illumina HiSeq 2500 platform. Approximately 5 Gb of raw sequencing data were generated for the collected specimen. Plastid genome and ITS were assembled using GetOrganelle v.1.7.1 with Illumina sequencing reads as input and under default parameters (Jin et al. 2020), respectively. The assembled plastid genome was annotated using Geneious Prime v.2023.0.4 (https://www. geneious.com) and manually checked with P. lobbii (NC_059699) and P. stobartiana (NC_059917) as references. Four plastid markers (matK, trnL, trnL-F, and at*pB-rbcL*) were extracted from plastid genome using Geneious Prime v.2023.0.4.

Sixty-two species of *Phalaenopsis* were used for phylogenetic analyses. Two species, *Cleisostoma williamsonii* (Rchb. f.) Garay and *Pelatantheria rivesii* (Guillaumin)Tang & F. T.Wang, were used as outgroup based on previous results (Chase et al. 2015; Li et al. 2019; Ma et al. 2022). In total, 225 sequences from 64 Orchidaceae species were downloaded from NCBI (Suppl. material 1: table S1). The combined matrix thus includes 229 sequences for the five markers, belonging to 65 species. Sequence alignment, supermatrix generation, and substitution model selection were performed using PhyloSuite (Zhang et al. 2020). GTR+F+I+G4 was selected as the best model for *matK*, *atpB-rbcL*, *trnL-F*, and *trnL*, and GTR+F+G4 for ITS, respectively. Bayesian Inference of phylogeny was performed using Mr-Bayes v.3.2.7a on XSEDE in the CIPRES Science Gateway online web server (Miller et al. 2010). Two separate Markov Chain Monte Carlo (MCMC) analyses were performed 1,000,000 generations and sampling every 1000 generations. Maximum likelihood (ML) analyses were performed locally using IQTree2 (Minh et al. 2020). Support values for the clade were estimated using 1,000,000 bootstrap replicates.

Results

Phylogram of Maximum Likelihood based on nrITS and plastid DNA markers were used to illustrate the phylogenetic position of the new species. *Phalaenopsis* sp. nov. is nested within subgen. *Parishianae* and sister to *P. taenialis* with



Figure 1. Phylogram of Maximum Likelihood based on nrITS and plastid DNA markers (*matK*, *trnL*, *trnL-F* and *atpB-rbcL*). Numbers above branches indicate bootstrap percentages (BS) for ML.

high support (PP = 1, BSML = 92; Fig. 1, Suppl. material 2: fig. S1). *Phalaenopsis* sp. nov. and *P. taenialis* together formed a clade sister to the clade consisting of *P. stobartiana* and *P. wilsonii* with high support (PP = 0.994, BSML = 89; Fig. 1, Suppl. material 2: fig. S1).

The new species is morphologically close to *P. taenialis*, *P. wilsonii*, and *P. stobartiana* by sharing lip with two seriate of calli at base, lip more or less with spur, lateral lobes more or less erect. *Phalaenopsis* sp.nov., however, differs from its relatives by having a bifurcated, fleshy, yellow anterior callus, and lateral lobes flipping outward and center with large calli.

Key to Phalaenopsis zhanhuoensis sp. nov. and its relatives

- ear......**3a**

- 3a Lip mid-lobe obcordate with a central apical fleshy knob 2. P. wilsonii
- 3b Lip mid-lobe not obcordate, without a terminal notch......4a
- 4b Flowers rose-pink; mid-lobe with a conspicuous constriction**5a**
- 5a Lip mid-lobe flared below apex producing a 3-lobulate mid-lobe

Taxonomy

Phalaenopsis zhanhuoensis X.H.Jin & S.Y.Qin, sp. nov. urn:lsid:ipni.org:names:77334958-1 Figs 2, 3 吉氏蝴蝶兰

Type. CHINA. Yunnan, Wenshan Ctiy, Xichou County, alt. 1496 m, 11 Apr 2023, *Xiaohua Jin & Shiyu Qin 40050* (holotype, PE!).

Diagnosis. *Phalaenopsis zhanhuoensis* is similar to *P. wilsonii*, but differs from it by having a bifurcated yellow, fleshy anterior callus, lateral lobes with large calli and flipping outward (Table 1).

Description. Epiphytic plants. Roots fleshy, developing from the base or lower parts of the stem, elongated, flattened, densely verrucose and prostrate along



Figure 2. Flowers of *Phalaenopsis zhanhuoensis* X.H.Jin & S.Y.Qin, sp. nov. **A** front view of flower (1) rear view of flower (2) **B** column and lip; lateral view of column and lip, appendage and lateral lobes (1), front view of lateral lobes (2), lateral view of lateral lobes and mid-lobe (3) **C** petal, sepal and lip, lip (1), dorsal sepal (2), petal (3.4), lateral sepals (5.6). Photographed by Xiaohua Jin.

trunks. Stem very short, covered by tubular sheath at base. Leaves unseen. Inflorescence developing from the base of stem, suberect or arching, ca. 4.5 cm long, unbranched, with 3 laxly arranged flowers. Floral bracts ovate-triangular, 4–5 mm long. Flowers white with pale pink rib or white, 3–4 cm in diameter. Dorsal sepal broadly elliptic or spoon-shaped, ca. 2×1 cm, with semi-transparent veins abaxially; lateral sepals elliptic, acute at apex, slightly curved toward labellum, lilac spots at the apex in the dorsal, ca. $1.8-2.0 \times 0.9-1.1$ cm, obtuse and notched at base. Petals spathulate, ca. $1.8-2.0 \times 0.9-1.1$ cm, obtuse. Labellum three-lobed, clawed at the base, ca. 1-2 mm long; lateral lobes of labellum erect, purple, 0.5 cm long, flipping outward, adaxially center with a big callus; mid-lobe of labellum obcordate, ca. $1.3-1.5 \times 0.9-1.1$ cm, deep purple, with white stripes at the center, base with a yellow fleshy protuberant anterior callus; anterior callus deeply lobed at apex. Column subparallel to midlobe of labellum, lavender, ca. 0.6 cm long, with triangular wings; pollinarium yellow.



Figure 3. Habitat and plants of *Phalaenopsis zhanhuoensis* X.H.Jin & S.Y.Qin, sp. nov. **A** front view **B** roots **C** lateral view. Photographed by Xiaohua Jin.

	P. zhanhuoensis	P. wilsonii	P. stobartiana	P. taenialis					
Flower color	white with pale pink ribs	white with pale pink ribs or complete pale pink.	sepals and petals apple- green to dark olive-green.	petals pale pink, lip and anther cap rose-purple.					
Leaves	no leaves at anthesis.	leaves often deciduous in dry season.	leaves often deciduous during dry season, but present at anthesis.	leaves often deciduous at anthesis or during dry season					
Lateral sepals	lateral sepals elliptic, acute at apex, obtuse	lateral sepals obovate-elliptic, similar and equal to middle sepal.	lateral sepals slightly oblique, ovate-elliptic, subacute.	lateral sepals subelliptic, base adnate to column foot, apex obtuse.					
Lateral lobes of lip	lateral lobes flipping outward , adaxial center with a big callus	lateral lobes erect, adaxially with an incised-tipped keel.	lateral lobes erect, narrow, slightly constricted at middle	lateral lobes adaxially with a slightly thickened longitudinal ridge close to proximal margin					
callus	yellow, fleshy, bifurcated.	purple; anterior callus deeply lobed at apex; lobelets linear and long	purple; concave adaxially and distinctly convex abaxially on disk.	purple; ligulate, deeply bifid; lobelets linear and long, attached to front wall at base of mid-lobe.					

Table 1.	Morphological	comparison o	f Phalaeno	psis zhanhuc	<i>ensis</i> and	close taxa.

Etymology. The epithet *zhanhuoensis* was designated in honor of the Chinese botanist Zhanhuo Tsi.

Distribution and habitat. *Phalaenopsis zhanhuoensis* is currently known only from the type locality in Xichou, Yunnan, China. It is epiphytic on trunks and twigs at elevations 1400–1500 m in evergreen broad-leaved forests.

Phenology. Flowering in March and April.

Conservation status. *Phalaenopsis zhanhuoensis* grows in evergreen broadleaf forests in Xichou County Yunnan Province, China. One subpopulation of about 10 individuals was discovered during our fieldwork. The habitat has been severely fragmented due to the development of agriculture. During our survey in nearby forests, we did not find any additional subpopulation of the new species. According to IUCN criteria v15.1 (IUCN 2022), we putatively assessed this new species as Critically Endangered CR C2a(i).

Additional information

Conflict of interest

The authors have declared that no competing interests exist.

Ethical statement

No ethical statement was reported.

Funding

This research was supported by the Project of Orchid Biodiversity Survey of China, National Forestry and Grassland Administration (2019070714, 2020100707, 2022070702 to XJ) and the National Natural Science Foundation of China (32270214, 31870195 to XJ).

Author contributions

Conceptualization: BY, XJ. Funding acquisition: XJ. Investigation: SQ, XJ. Methodology: CM. Validation: ZL, YW. Writing – original draft: SQ. Writing – review and editing: HW.

Author ORCIDs

Shiyu Qin [®] https://orcid.org/0009-0008-2680-6789 Hanchen Wang [®] https://orcid.org/0000-0002-4690-8052 Yajun Wang [®] https://orcid.org/0009-0001-1035-3964 Chongbo Ma [®] https://orcid.org/0000-0001-5869-8649 Zan Li [®] https://orcid.org/0009-0000-3098-1418 Boyun Yang [®] https://orcid.org/0000-0003-2123-0027 Xiaohua Jin [®] https://orcid.org/0000-0002-9987-5602

Data availability

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

References

- Chase MW, Cameron KM, Freudenstein JV, Pridgeon AM, Salazar G, Van den Berg C, Schuiteman A (2015) An updated classification of Orchidaceae. Botanical Journal of the Linnean Society 177(2): 151–174. https://doi.org/10.1111/boj.12234
- Chen XQ, Wood JJ (2009) *Phalaenopsis* Blume, Bijdr. 294. 1825. In: Wu CY, et al. (Eds) Flora of China, vol. 25. Beijing: Science Press and Missouri Botanical Garden Press, St. Louis, 478–483.
- Christenson EA (2001) Phalaenopsis: a monograph. Timber Press, Portland, 330 pp.
- Higgins W, Alrich P (2015) *Phalaenopsis*: the Genus in Pictures. International *Phalaenopsis* Alliance, Taiwan, 80–83.
- IUCN (2022) Guidelines for Using the IUCN Red List Categories and Criteria. Version 15.1. https://www.iucnredlist.org/resources/redlistguidelines [Accessed 4 Dec.2023]
- Jin JJ, Yu WB, Yang JB, Song Y, dePamphilis CW, Yi TS, Li DZ (2020) GetOrganelle: A fast and versatile toolkit for accurate de novo assembly of organelle genomes. Genome Biology 21(1): 241. https://doi.org/10.1186/s13059-020-02154-5
- Kocyan A, Schuiteman A (2014) New combinations in Aeridinae (Orchidaceae). Phytotaxa 161: 061–085. https://doi.org/10.11646/phytotaxa.161.1.3
- Li J, Wang S, Yu J, Ling W, Zhou S (2013) A Modified CTAB Protocol for Plant DNA Extraction. Zhiwu Xuebao 48(1): 72–78. https://doi.org/10.3724/SP.J.1259.2013.00072
- Li YX, Li ZH, Schuiteman A, Chase MW, Li JW, Huang WC, Hidayat A, Wu SS, Jin XH (2019) Phylogenomics of Orchidaceae based on plastid and mitochondrial genomes. Molecular Phylogenetics and Evolution 139: 106540. https://doi.org/10.1016/j. ympev.2019.106540
- Ma CB, Wang XL, Chen WS, Liu Z, Jin XH (2022) *Phalaenopsis medogensis* (Orchidaceae, Epidendroideae, Vandeae), a new species from Xizang, China. PhytoKeys 214: 39–46. https://doi.org/10.3897/phytokeys.214.93607
- Miller MA, Pfeiffer W, Schwartz T (2010) Creating the CIPRES Science Gateway for inference of large phylogenetic trees. In: Gateway Computing Environments Workshop. New Orleans, 1–8. https://doi.org/10.1109/GCE.2010.5676129
- Minh BQ, Schmidt HA, Chernomor O, Schrempf D, Woodhams MD, von Haeseler A, Lanfear R (2020) IQ-TREE 2: New Models and Efficient Methods for Phylogenetic Inference in the Genomic Era. Molecular Biology and Evolution 37(5): 1530–1534. https:// doi.org/10.1093/molbev/msaa015
- Padolina J, Linder C, Simpson B (2005) A Phylogeny of *Phalaenopsis* Using Multiple Chloroplast Markers. Selbyana 26: 23–27.

- Pridgeon AM, Cribb PJ, Chase MW, Rasmussen FN (2014) Genera orchidacearum: Epidendroideae, volume 6, Part 3. Oxford University Press, Oxford, 232–241.
- Tsai CC, Huang SC, Chou CH (2005) Molecular phylogeny of *Phalaenopsis* Blume (Orchidaceae) based on the internal transcribed spacer of the nuclear ribosomal DNA. Plant Systematics and Evolution 256(1–4): 1–16. https://doi.org/10.1007/ s00606-005-0356-y
- Tsai C, Chiang Y-C, Huang S, Chen C, Chou C (2010) Molecular phylogeny of *Phalaenopsis* Blume (Orchidaceae) on the basis of plastid and nuclear DNA. Plant Systematics and Evolution 288(1–2): 77–98. https://doi.org/10.1007/s00606-010-0314-1
- Yukawa T, Kita K, Handa T, Hidayat T, Ito M (2005) Molecular Phylogenetics of *Phalaenopsis* (Orchidaceae) and allied Genera: Re-evaluation of Generic Concepts. Acta Phytotaxonomica et Geobotanica 56: 141–161. https://doi.org/10.18942/apg.KJ00004623240
- Zhang D, Gao F, 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
- Zhou Z, Shi R, Zhang Y, Xing X, Jin X (2021) Orchid conservation in China from 2000 to 2020: Achievements and perspectives. Plant Diversity 43(5): 343–349. https://doi. org/10.1016/j.pld.2021.06.003

Supplementary material 1

GenBank accession numbers for sequences used for phylogenetic analyses

Authors: Shiyu Qin, Hanchen Wang, Yajun Wang, Chongbo Ma, Zan Li, Boyun Yang, Xiaohua Jin

Data type: docx

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.237.112270.suppl1

Supplementary material 2

Phylogram of Bayesian Inference (BI) based on nrDNA ITS and plastid sequences (*matK*, *trnL*, *trnL*-F, and *atpB-rbcL*)

Authors: Shiyu Qin, Hanchen Wang, Yajun Wang, Chongbo Ma, Zan Li, Boyun Yang, Xiaohua Jin

Data type: pdf

Explanation note: Numbers above branches indicate posterior probabilities (PP) for BI.

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.237.112270.suppl2