A new species of Ranunculus (Ranunculaceae) from Western Pamir-Alay, Uzbekistan

Natalia V. Shchegoleva¹, Elena V. Nikitina², Inom J. Juramurodov³,⁴, Andrei A. Zverev⁵, Orzimat T. Turginov³, Anvarbek M. Jabborov³, Ziyoviddin Yusupov⁶,⁷, Davron B. Dekhkonov⁶, Tao Deng⁷, Hang Sun⁷

¹ Department of Botany, Institute of Biology, Tomsk State University, 36 Lenin Ave., Tomsk 634050, Russia
² Laboratory of Cadastre and Monitoring of Rare Plant Species, Institute of Botany of the Academy of Sciences of the Republic of Uzbekistan, 32 Durmon Yuli St., Tashkent, 100125, Uzbekistan
³ Laboratory Flora of Uzbekistan, Institute of Botany of the Academy of Sciences of the Republic of Uzbekistan, 32 Durmon Yuli St., Tashkent, 100125, Uzbekistan
⁴ University of Chinese Academy of Sciences, Beijing, China
⁵ Central Siberian Botanical Garden, Siberian Branch, Russian Academy of Sciences, 630090 Novosibirsk, Russia
⁶ Laboratory of Molecular Phylogeny and Biogeography, Institute of Botany of the Academy of Sciences of the Republic of Uzbekistan, 32 Durmon Yuli St., Tashkent, 100125, Uzbekistan
⁷ CAS Key Laboratory for Plant Diversity and Biogeography of East Asia, Kunming Institute of Botany, Chinese Academy of Sciences, Kunming 650201, Yunnan, China

Corresponding authors: Natalia V. Shchegoleva (schegoleva@outlook.com), Hang Sun (sunhang@mail.kib.ac.cn)

Abstract

New data on the phylogeny of four rare and endemic species of Ranunculus L. (sect. Ranunculastrum DC.) of western Pamir-Alai, one of which is new to science, have been obtained. Ranunculus tojibaevii sp. nov., from the Baysuntau Mountains, Western Hissar Range of Uzbekistan, is described. The new species is closely related to R. botschantzevii Ovcz., R. convexiusculus Kovalevsk. and R. alpigenus Kom., but differs in the blade of the radical leaves, which is rounded-reniform, segments 3–5-dissected, each 2–5-partite with elongated, rounded apical lobes. A phylogenetic analysis, using both the nuclear ribosomal internal transcribed spacer (ITS) and cpDNA (matK, rbcL, trnL-trnF), was informative in placing R. tojibaevii in context with its most closely-related species. Discussion on the geographic distribution, updated identification key, a detailed description, insights about its habitat and illustrations are provided.

Keywords

Endemic, Hissar Range, Pamir-Alay, phylogenetic analysis, Ranunculales, Ranunculastrum
Introduction

*Ranunculus* L., the largest genus in the Ranunculaceae Juss., includes ca. 600 genetically diverse species (Tamura 1995). The genus is distinguished by its high ecological-geographical diversity over a wide zonal spectrum ranging from the Arctic tundra through varied forests, steppes, deserts to exclusively aquatic habitats and high-altitude communities on nearly all continents (Paun et al. 2005). However, the main centres of speciation of *Ranunculus* are often in large mountain systems, where the formation of species is not only directly related to isolation, but also significantly depends on specific conditions of the highlands (Emadzade et al. 2015; Fernández Prieto et al. 2017; Shchegoleva 2018; Shchegoleva et al. 2020; Zverev et al. 2020).

More than 90 species of *Ranunculus* are distributed in Central Asia (Kovalevskaya 1972). Differentiation in the genus and the formation of locally endemic species are closely related to the history of the Tian Shan and Pamir-Alay Mountain formations. Here, more than half of the members of the genus are autochthonous representatives of the mountainous Central Asian flora, which arose in the process of regional adaptive diversification from ancient Mediterranean predecessors (Ovchinnikov 1971; Kamelin 1973). In the dry Central Asian seasonal climate, the features of these species are observed only in the short-term hydrothermal period of active vegetative growth.

*Ranunculus tojibaevii* was first discovered in 2013 on the Baysuntau Highlands (Khodzha-Gurgur-ata Mountain) on the south-western spur of the Hissar Range (Pamir-Alay). The populations were detected again in 2019 while working on the *Flora of Uzbekistan Project* (Sennikov et al. 2016). It should be noted that the flora of the Western Pamir-Alay is characterised by a high taxonomic diversity (Kamelin 1973; Vasilchenko and Vasileva 1985; Tojibaev et al. 2016; Makhmudjanov et al. 2019; Yusupov et al. 2020).

The morphological features indicated that the unknown plants belonged to *R.* subg. *Ranunculus* sect. *Ranunculastrum* (Hörandl and Emadzade 2012; Baltisberger and Hörandl 2016). The main differences between sect. *Ranunculastrum* and other sections of *Ranunculus* are the presence of a triangular beak equal to or longer than the achene body, a receptacle completely glabrous, a taproot partly tuberous and a mostly elongate fruit (Hörandl and Emadzade 2012).

The unknown plants closely resembled *R. botschantzevii* Ovcz. (Ovchinnikov 1941) and *R. convexiusculus* Kovalevsk. (Kovalevskaya 1972), as well as *R. alpigenus* Kom. (Komarov 1896) in their main morphological features. The molecular results presented here also clearly substantiated its independent taxonomic status. In this study, we present a morphological description of these plants, which we named *Ranunculus tojibaevii* Schegol. & Turginov. Figures showing its features, a map of its distribution, taxonomy and an identification key separating it from the most closely-related species are also provided.
Materials and methods

Morphological observations and measurements of *R. tajibaevii* were made on two populations; in total, 34 individuals were compared. Voucher specimens have been deposited in the National Herbarium of Uzbekistan – TASH (Tashkent, Uzbekistan). Additionally, two closely-related species, *R. convexiusculus* and *R. botschantzevii*, from the same territory and high-altitude regions were examined. Herbarium specimens at TASH, TAD, LE, FRU, AA, MW, LE and TK were also compared. Studies of closely related species were performed during field expeditions to Uzbekistan, Tajikistan and Kyrgyzstan, from 2017–2021 (Shchegoleva et al. 2020). The distribution map was generated in ESRI ArcGIS 10 software using GPS coordinates (www.esri.com). The conservation status was established, based on IUCN Criteria (IUCN 2019).

Molecular methods

DNA extraction, amplification and sequencing

DNA isolation was performed using a Plant Genomic DNA Kit (TIENGEN Biotech, Beijing, China) according to the manufacturer’s protocol. CTAB extraction protocol with some modifications was used to extract genomic DNA from herbarium specimens of *R. alpigenus* (Doyle and Doyle 1987).

Selected nuclear DNA regions ITS1-ITS2 (for herbarium specimen *R. alpigenus*) and ITS1-ITS4; plastid DNA regions *matK*, *rbcL* and *trnL-F* were amplified on a thermal cycler (BioRad) using the 2X PCR Taq Plus MasterMix with dye (Applied Biological Materials Inc., Canada). Amplification of the DNA regions was carried out by using primers of the forward and reverse primer sets (TsingKe, China) (Table 1).

To obtain sequences of the genes of interest, PCR amplification was carried out according to the following parameters (except *R. alpigenus*): for ITS1-ITS4, initial denaturation for 3 min at 94 °C, followed by 35 amplification cycles: 30 s at 94 °C, 30 s at 50–54 °C, 1 min at 72 °C; elongation 7 min at 72 °C; for *matK*– an initial denaturation for 3 min 94 °C, followed by 35 amplification cycles: 30 s 94 °C, 1 min 51 °C, 1 min 72 °C; final extension 10 min at 72 °C; for *rbcL*, an initial denaturation for 4 min 95 °C, followed by 34 amplification cycles: 1 min 94 °C, 1 min 50 °C, 1 min 72 °C; final extension 10 min at 72 °C; for *trnL-F*, an initial denaturation for 3 min 94 °C, followed by 32 amplification cycles: 45 s 94 °C, 45 s 50 °C, 1 min 72 °C; final extension 8 min at 72 °C.

PCR amplification for *R. alpigenus* was performed for ITS1-ITS2, with the following programme: initial denaturation at 94 °C/5 min; 35 amplification cycles at 94 °C/30 s, at 54 °C/30 s, at 72 °C/ 45 s; elongation at 72 °C/7 min; for *rbcL*, 94 °C/4 min, 34 cycles: 94 °C/30 s, 54 °C/ 45 s, 72 °C/45 s; final extension at 72 °C/10 min.
Table 1. Primers used in this study.

| Primer name | Sequences (forward / reverse) | DNA fragment size, bp | Primer source |
|-------------|-------------------------------|-----------------------|---------------|
| ITS1-18S    | 5’-TCCGTAGGTGAACCTGCGG-3’ | ~ 700 bp              | White et al. (1990) |
| ITS4-26S    | 5’-TCCTCCGCTTATTGATATGC-3’ | ~ 650 bp              | White et al. (1990) |
| ITS1        | 5’-TCCGTAGGTGAACCTGCGG-3’   | ~ 650 bp              | White et al. (1990) |
| ITS2        | 5’-GCTGCGTTCTTCATCGATGC-3’ | ~ 900 bp              | Cuenoud et. al. (2002) |
| matK-390F   | 5’-TGATCTATTCATTCAATATTTC-3’| ~ 900 bp              | Cuenoud et. al. (2002) |
| matK-1326R  | 5’-TCATAGCAGAAAGTCGAAAGT-3’| ~ 900 bp              | Taberlet et al. (1991) |
| trnL-F_F    | 5’-CGAAATCGGTAGACGCTACG-3’  | ~ 900 bp              | Taberlet et al. (1991) |
| rbcLaF      | 5’-ATGTCACCACAAACAGAGAAGC-3’| ~ 600 bp              | Kress and Erickson (2007) |
| rbcLaR      | 5’-GTAATCAGTCCACCCRCG-3’    | ~ 600 bp              | Kress and Erickson (2007) |

To determine the taxonomic status and systematic position of *R. tojibaevii*, we sampled 24 species of *Ranunculus*. New nDNA (ITS) and cpDNA intergenic spacers (*matK*, *rbcL*, *trnL-trnF*) sequences for nine species were generated. We also used available sequences of 15 *Ranunculus* species from GenBank (www.ncbi.nlm.nih.gov/Genbank) (Table 2). We used *Ranunculus* subgenus *Auricomus* as the outgroup (Hörandl and Emadzade 2012; Almerekova et al. 2020).

Phylogenetic analyses

Sequence alignments were performed using ClustalW (Thompson et al. 2002) as implemented in MEGA X software (Kumar et al. 2018). The best partitioning scheme for the combined dataset contained two partitions: the ITS data; and the three plastid sequences data (*matK*, *rbcL*, *trnL-trnF*). Phylogenetic reconstruction was first conducted separately, based on the nuclear and the plastid data. Visual inspection determined that differences between the nuclear and the plastid trees were solely due to resolved/collapsed clades. No topological incongruence with a high support value (posterior probabilities and bootstrap percentages) was found. To further test whether the nuclear and plastid data could be combined for phylogenetic reconstruction, the incongruence length difference (ILD, Farris et al. 1995) test was conducted in PAUP* 4.0a169 (current) by using only the informative sites, heuristic search, tree-bisection-reconnection (TBR) branch-swapping algorithm, simple addition sequence and 1,000 replicates. The ILD test between the nuclear and the plastid data found $p = 0.322$, indicating insignificant support for incongruence between the two datasets. Therefore, the nuclear and the plastid sequences were combined into one dataset for phylogenetic analyses using SequenceMatrix software (Vaidya et al. 2011).
 Phylogenetic trees were reconstructed using Maximum Likelihood (ML), Maximum Parsimony (MP) and Bayesian Inference (BI). For ML, we employed raxmlGUI 2.0 (Edler et al. 2020), with 1,000 bootstrap replicates and, for BI, we used MrBayes v.3.1.2 (Huelsenbeck and Ronquist 2001) with 10,000,000 generations with random trees sampled every 1,000 generations. In the latter analysis, after discarding the first 25% trees as burn-in, a 50% majority-rule consensus tree was constructed from the remaining trees to estimate Posterior Probabilities (PP). For analyses, a model of nucleotide substitution was selected, based on the Akaike Information Criterion (AIC) using jModelTest2 on XSEDE (www.phylo.org). Phylogenetic analyses were also performed with the MP method using PAUP* 4.0a169. The MP bootstrap analysis was performed with heuristic search, TBR branch-swapping, 1,000 bootstrap replicates, random addition sequence with ten replicates, a maximum of 1,000 trees saved per round. Trees were visualised in FigTree v.1.4.0 (Rambaut 2012).

**Table 2.** Accession numbers of samples used for phylogenetic analyses of *Ranunculus* (* newly-generated sequences).

| Species               | GenBank accession number |
|-----------------------|--------------------------|
|                       | ITS | matK | rbcL | trnL-F |
| *Ranunculus acris*    | AY680167 | AY954199 | HQ590232 | – |
| R. alpigenus*         | OM283824 |  | OM287560 | – |
| R. arvensis           | HQ650550 | HQ650551 | MK925091 | – |
| R. auricomus          | FM242803 | FM242739 | JN893758 | – |
| R. botchantzevii*     | MW540744 | MW748677 | MW748685 | MW748693 |
| R. convexicusculus*   | MW540743 | MW748676 | MW748684 | MW748692 |
| R. flavula            | AY680185 | AY954204 | MK526480 | – |
| R. glorerrimus        | KP687273 | JF509974 | MG247649 | – |
| R. inamoenus          | KP687279 | KP687302 | MG249011 | – |
| R. japonicus          | EU591982 | AY954200 | MH577741 | DQ410744 |
| R. leptorrhynchos*    | MW737444 | MW748673 | MW748681 | MW748689 |
| R. linealbatus*       | MW737445 | MW748674 | MW748682 | MW748690 |
| R. lingua             | AY680184 | AY954206 | JN92742 | – |
| R. maricatus          | DQ410718 | AY954191 | HM50296 | DQ410740 |
| R. paucidentatus*     | MW540747 | MW748679 | MW748687 | MW748695 |
| R. pygmaeus           | KP687287 | KP687310 | KC438650 | – |
| R. regelianus*        | MW737446 | MW748675 | MW748683 | MW748691 |
| R. repens             | MT271835 | HM561666 | MK925397 | EU382995 |
| R. seleratus          | MT271836 | GU257993 | AB517148 | DQ410746 |
| R. sulphurea          | JF509969 | JF509883 | KC438700 | – |
| R. talasicas*         | MW540748 | MW748680 | MW748688 | MW748696 |
| R. tajikiacii*        | MW540745 | MW748678 | MW748686 | MW748694 |
| R. tojibaevi*         | OM278385 | OM287558 | OM287559 | – |
| R. trichophyllus      | KC620483 | AY954133 | L08766 | – |
| R. turneri            | FM242817 | FM242741 | MG249550 | – |
Results and discussion

The phylogenetic tree, based on the nuclear and plastid sequences (Fig. 1), showed that *R. tojibaevii* is sister to *R. convexiusculus* and *R. botschantzevi* with high support values: PP = 1, MP = 94% and ML = 94%. *Ranunculus tojibaevii*, *R. convexiusculus* and *R. botschantzevi* formed a clade with well supported values (PP = 0.8, MP = 71% and ML = 64%).

The results of the phylogenetic analysis were similar to the results of Hörandl and Emadzade (2012) and Almerekova et al. (2020). Species of *R. sect. Ranunculastrum* are of particular interest. In our case, these native Asian species are mainly limited to the foothills and mountains of Central Asia (southern Kazakhstan, Uzbekistan, Kyrgyzstan, Tajikistan and Turkmenistan), as well as to the mountains of Afghanistan, Iran and Pakistan.

It is obvious that *R. sect. Ranunculastrum* in Central Asia is heterogeneous. The species forming sub-clusters in this section differ in their area of distribution, altitude confinement and time of origin, thereby confirming the neoendemic nature and origin of *R. tojibaevii*.

**Figure 1.** Bayesian tree based on combined nuclear (ITS) and plastid (*matK, rbcL, trnL-trnF*) sequence data showing phylogenetic position of *Ranunculus tojibaevii* Schegol. & Turginov in *R. sect. Ranunculastrum*. Bayesian Posterior Probability (PP) / Maximum Parsimony (MP) is given on each branch, respectively; Maximum Likelihood (ML) is below branches. The classification is according to Hörandl and Emadzade (2012). * denotes the *Ranunculus* species analysed in this study. The new species is highlighted in bold.
Taxonomy

*Ranunculus tojibaevii* Schegol. & Turginov, sp. nov.
urn:lsid:ipni.org:names:77296907-1

Figs 2, 3

**Diagnosis.** Similar to *R. botschantzevii*, *R. convexiusculus* and *R. alpigenus* morphologically, but differing in the rounded-reniform radical leaves dissected into 3–5 segments, each 2–5-partite into elongated lobes rounded at the apex (Fig. 4). *Ranunculus tojibaevii* differs from *R. alpigenus* in having fewer levels of leaf blade dissection. It differs from *R. convexiusculus* in the dissection of the radical leaf blades, larger flowers and having somewhat white, bristle-like hairs on the root collar and also from *R. botschantzevii* by the rounded apical lobes of the basal leaves and more xeromorphic habit.

**Type.** Uzbekistan. Hissar Ridge, Baysuntau, Khodzha-Gurgur-ata Mountains, vicinity of the Village Gumatag, 38°22.2888’N, 67°21.0834’E, 2482 m a.s.l., 30 May 2019, N. Shchegoleva, A. Jabborov, O. Turginov (holotype: TASH-003743; isotypes: TASH-003748, TASH-003749, TASH-003750, TK-002339).

**Description.** *Herbs* perennial. *Roots* dimorphic, some roots subulate, up to 0.5 mm thick, storage roots palmately-thickened, ca. 2.5 mm thick; root collar with milk-white bristle-like hairs. *Stems* 10–12(–15) cm tall, up to 2 mm diam., erect, branched, pubescent with white curly hairs, 1–3-flowered. *Leaves* dimorphic, radical leaves 2–3, 1.6–2 × 1.5–2.2 cm, blade rounded-reniform, 3–5-dissected, segments 2–5-partite, elongate, lobes apically rounded; cauline leaves 1–2, petiole short, slender, blade trisected, lobes 0.6–0.9 × 0.1–0.2 cm, oblong-lanceolate. *Flowers* 1.6–2.4 cm diam., sepals 0.4–0.6 mm long, ovate-concave, sparsely white pubescent; petals 0.9–1.2 cm long, well-developed, ovate, apex rounded. Infructescence globose-ovoid; receptacle oblongoid, glabrous; achenes 1–1.8 mm long, with white bristle-like hairs; beak hamate-curved.

**Specimen seen (paratype).** Uzbekistan. Pamir-Alay, South-western spurs of the Hissar Ridge, Baysuntau, vicinity of the Village Gumatag, amongst the stones, 4 June 2013, O. Turginov (TASH-003754).

**Phenology.** Flowering in May. Fruiting in May and June.

**Distribution.** *Ranunculus tojibaevii* is distributed in the Khodzha-Gurgur-ata of the Baysuntau Mountains area of Hissar Ridge (Fig. 5). The closely-related *R. convexiusculus* is endemic to Central Asia and *R. botschantzevii* is endemic to the Western Pamir-Alay.

**Habitat and plant associations.** *Ranunculus tojibaevii* is rupicolous on southern and western exposed limestone outcrops and in cracks and crevices of large boulders at 2,450–2,500 m. a.s.l. The region is alpine and rather xerophytic. The common taxonomic composition of phytocenos includes *Cerasus amygdaliflora* Nevski (Rosaceae), *Corydalis ledebouriana* Kar. & Kir. (Papaveraceae), *Cousinia regelii* C.Winkl. (Asteraceae), *Eremurus regelii* Vved. (Asphodelaceae), *Gagea gymnopoda* Vved.
Figure 2. *Ranunculus tojibaevii* Schegol. & Turginov (Shchegoleva, Jabborov, Turginov, holotype TASH-003743).
Figure 3. *Ranunculus tojibaevii* Schegol. & Turginov. Habitat (A flowering B fruiting) and flower (C). Scale bar: 1 cm.
Figure 4. Series of basal leaves in related species A *Ranunculus tojbaevii* (from the holotype) B *R. alpigenus* C *R. botschantzevii* D *R. convexiusculus*. Scale bar: 1 cm.
A new species of *Ranunculus* (Liliaceae), *Iris khassanovii* Tojibaev & Turginov, *I. parvula* (Vved.) T.Hall & Seisums, *I. stolonifera* Maxim. (all Iridaceae), *Rheum maximowiczii* Losinsk. (Polygonaceae), *Tulipa lanata* Regel (Liliaceae) and *Ziziphora pamiroalaica* Juz. (Lamiaceae).

**Conservation status.** *Ranunculus tojibaevii* is a local, narrowly distributed endemic, represented by two or three individuals per m$^2$ within an area of < 500 m$^2$. It should therefore be assigned the status EN (Endangered), Criteria B1 ab(i, ii, iii)+ B2 ab(i, ii, iii), following the IUCN Standards and Petitions Committee (IUCN 2019).

**Notes.** *Ranunculus tojibaevii* differs from closely-related species by its habitat on well-heated limestone outcrops, as well as in cracks and crevices of large boulders at ca. 2,500 m a.s.l., which is atypical of related species. *Ranunculus convexiusculus* is on clayey-stony soil, less often on slopes of fine soil, at 2,000–2,600 m a.s.l. *Ranunculus botschantzevii* is hygrophilous in wet mountain meadows with melting snow and on slopes of fine clayey soil at 2,400–3,500 m a.s.l. *Ranunculus alpigenus* grows on slopes of fine soil of the alpine belt at 2,800–4,000 m a.s.l. All these species are endemic to the western Pamir-Alay. The vicariant species to *R. alpigenus* is *R. badachschanicus* Ovcz. & Koch. from the western Pamirs.

**Etymology.** *Ranunculus tojibaevii* is named after Komiljon Tojibaev, a leading botanist, professor and academician from Uzbekistan who actively promotes the botanical sciences in Central Asia.
Key to *Ranunculus tojibaevii* and similar species (Table 3)

1. Blades of basal leaves broadly ovate, dissected into pinnatipartite segments, with tripartite-oblong terminal lobules .............................................. *R. alpigenus*
   – Radical leaves 3–5-dissected or lobed-incised .............................................. 2

2. Blade of radical leaves 1/3 or nearly 1/2 unequally partite into broad incised-dentate lobes .................................................. *R. convexiusculus*
   – Blade of radical leaves 3–5-dissected ................................................................................................................................. 3

3. Blade of basal leaves triangular-reniform, 3–5-dissected almost to the base, wedge-shaped segments, unequally sharp-toothed .......... *R. botschantzevii*
   – Blade of radical leaves is round-reniform, 3–5-dissected, each section divided into 2–5 elongated lobules; apex of lobules rounded ............ *R. tojibaevii*

**Table 3.** Comparison of *R. tojibaevii*, *R. botschantzevii*, *R. convexiusculus* and *R. alpigenus*.

|                  | *R. tojibaevii* | *R. botschantzevii* | *R. convexiusculus* | *R. alpigenus* |
|------------------|-----------------|---------------------|---------------------|----------------|
| **Blade of radical leaves** | rounded-ovate, 3–5-dissected, segments further 2–5-dissected, lobules elongated, rounded at apex | triangular-reniform, 3–5-partite, segments narrowly cuneate, unequally and subcuteley dentate | reniform, dissected 1/3 to nearly 1/2 of its length, lobes broad incised-dentate | broadly ovate, dissected into pinnatipartite segments, segments triangite, terminal lobules oblong |
| **Sepals** | narrowly elliptic, concave, less than half as long as petals, with long reclinate hairs | elliptic, concave, some shorter than the petals, with sparse, long reclinate hairs | elliptic, concave, half as long as petals, with short reclinate spreading hairs | elliptic, concave, with scattered hairs |
| **Petals** | oblong-ovate, greenish-yellow, base cuneate, margin undulate | obovate, bright yellow, becoming dark when dry, base broadly cuneate, margin undulate | very broadly ovate, golden yellow, base cuneate, margin slightly undulate | oblong-ovate, yellow-green, base narrowly cuneate, marginundulate |
| **Achenes** | 1.0–1.8 mm long, asymmetrical ovate, slightly convex, with semi-appressed hairs | 2.2–2.5 mm long, oblong, slightly laterally compressed, with appressed hairs | 1.8–2.5 mm long, oblong, slightly convex, with appressed hairs | 1.5–2.0 mm long, asymmetrical obovate, laterally compressed, with scattered not appressed hairs |

**Acknowledgements**

This study was supported by the International Partnership Program of the Chinese Academy of Sciences (151853KYSB20180009), the framework of the Second Tibetan Plateau Scientific Expedition and Research (STEP) program (2019QZKK0502), the Strategic Priority Research Program of the Chinese Academy of Sciences (XDA20050203), the Key Projects of the Joint Fund of the National Natural Science Foundation of China (U1802232), the Youth Innovation Promotion Association of the Chinese Academy of Sciences (2019382), Yunnan Young & Elite Talents Project (YNWR-QNBJ-2019-033), the Chinese Academy of Sciences “Light of West China” Program, the state research project “Taxonomic revision of polymorphic plant families of the flora of Uzbekistan” (FZ-20200929321), the State Programs for 2021–2025
A new species of Ranunculus

years “Grid mapping of the flora of Uzbekistan” and “The creation of DNA bank and barcoding of endemic plants of Uzbekistan”, as well as in accordance with the current State Assignment of Central Siberian Botanical Garden at Siberian Branch of Russian Academy of Sciences (AAAA-A21-121011290026-9).

References

Almerekova S, Shchegoleva N, Abugalieva S, Turuspekov Y (2020) The molecular taxonomy of three endemic Central Asian species of Ranunculus (Ranunculaceae). PLoS ONE 15(10): e0240121. https://doi.org/10.1371/journal.pone.0240121

Baltisberger M, Hörandl E (2016) Karyotype evolution supports the molecular phylogeny in the genus Ranunculus (Ranunculaceae). Perspectives in Plant Ecology, Evolution and Systematics 18: 1–14. https://doi.org/10.1016/j.ppees.2015.11.001

Cuénoud P, Savolainen V, Chatrou LW, Powell M, Grayer RJ, Chase MW (2002) Molecular phylogenetics of Caryophyllales based on nuclear 18S rDNA and plastid rbcL, atpB, and matK DNA sequences. American Journal of Botany 89(1): 132–144. https://doi.org/10.3732/ajb.89.1.132

Doyle JJ, Doyle JL (1987) A rapid DNA isolation procedure for small quantities of fresh leaf tissue. Phytochemical Bulletin 19: 11–15.

Edler D, Klein J, Antonelli A, Silvestro D (2020) raxmlGUI 2.0: A graphical interface and toolkit for phylogenetic analyses using RAxML. Methods in Ecology and Evolution 12(2): 373–377. https://doi.org/10.1111/2041-210X.13512

Emadzade K, Lebmann M, Hoffmann MH, Tkach N, Lone F, Hörandl E (2015) Phylogenetic relationships and evolution of high mountain buttercups (Ranunculus) in North America and Central Asia. Perspectives in Plant Ecology, Evolution and Systematics 17(2): 131–141. https://doi.org/10.1016/j.ppees.2015.02.001

Farris JS, Källersjö M, Kluge AG, Bult C (1995) Testing significance of incongruence. Cladistics 10(3): 315–319. https://doi.org/10.1111/j.1096-0031.1994.tb00181.x

Fernández Prieto JA, Sanna M, Pérez M, Pérez Haase A, Molero Mesa J, Cires E (2017) Evolutionary and biogeographic relationships of related Ranunculus taxa: Dispersal, vicariance and pseudovicariance as mechanisms of change. Plant Ecology & Diversity 10(5–6): 379–387. https://doi.org/10.1080/17550874.2018.1430187

Hörandl E, Emadzade K (2012) Evolutionary classification: A case study on the diverse plant genus Ranunculus L. (Ranunculaceae). Perspectives in Plant Ecology, Evolution and Systematics 14(4): 310–324. https://doi.org/10.1016/j.ppees.2012.04.001

Huelsenbeck JP, Ronquist F (2001) MRBAYES: Bayesian inference of phylogenetic trees. Bioinformatics (Oxford, England) 17(8): 754–755. https://doi.org/10.1093/bioinformatics/17.8.754

IUCN (2019) Guidelines for Using the IUCN Red List Categories and Criteria. Version 14. Prepared by the Standards and Petitions Committee.

Kamelin RV (1973) The florogenetical analysis of native flora of the mountain Middle Asia. Leningrad, 356 pp.
Komarov VL (1896) Materials to the flora of Turkestan mountains. Basin of the River Zeravshan. Travaux de la Société des Naturalistes de St. Pétersbourg. Section de Botanique 26: 31–162.

Kovalevskaya SS (1972) *Ranunculus* L. In: Bondarenko OM, Nabiev MM (Eds) Identification key to plants of Middle Asia. Tashkent 3: 204–225.

Kress JW, Erickson LD (2007) A two-locus global DNA barcode for land plants: The coding rbcL gene complements the non-coding trnH-psbA spacer region. PLoS ONE 2(6): 1–10. https://doi.org/10.1371/journal.pone.0000508

Kumar S, Stecher G, Li M, Knyaz C, Tamura K (2018) MEGA X: Molecular evolutionary genetics analysis across computing platforms. Molecular Biology and Evolution 35(6): 1547–1549. https://doi.org/10.1093/molbev/msy096

Makhmudjanov D, Yusupov Z, Abdullaev D, Deng T, Tojibaev K, Sun H (2019) The complete chloroplast genome of *Eremurus robustus* (Asphodelaceae). Mitochondrial DNA. Part B, Resources 4(2): 3366–3367. https://doi.org/10.1080/23802359.2019.1674198

Ovchinnikov PN (1941) New species of the genus *Ranunculus* L. Botanical materials of the Herbarium of the Botanical Institute of the Uzbek branch of the USSR Academy of Sciences 3: 28–30.

Ovchinnikov PN (1971) Flora and vegetation of Varzob River basin. Nauka, Leningrad, 447 pp.

Paun O, Lehnebach C, Johansson JT, Lockhart P, Hörandl E (2005) Phylogenetic relationships and biogeography of *Ranunculus* and allied genera (Ranunculaceae) in the Mediterranean region and in the European alpine system. Taxon 54(4): 911–932. https://doi.org/10.2307/25065478

Rambaut A (2012) FigTree v.1. 4.0. University of Oxford, Oxford. http://tree.bio.ed.ac.uk/software/figtree

Sennikov AN, Tojibaev KS, Khassanov FO, Beshko NY (2016) The Flora of Uzbekistan Project. Phytotaxa 282(2): 107–118. https://doi.org/10.11646/phytotaxa.282.2.2

Shchegoleva NV (2018) New distribution data on Altai endemic *Ranunculus schischkinii* in floras of Russia and Kazakhstan. Vestnik Tomskogo gosudarstvennogo universiteta. Biologiya = Tomsk State University. Journal of Biology 41: 182–189. https://doi.org/10.17223/19988591/41/1

Shchegoleva NV, Turginov OT, Jabborov AM, Kodirov UH (2020) Ecological and geographical features of the Western Pamir-Alai endemic *Ranunculus botschantzevii* Ovcz. Vestnik Tomskogo gosudarstvennogo universiteta. Biologiya = Tomsk State University. Journal of Biology 49: 175–186. https://doi.org/10.17223/19988591/49/10

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

Tamura M (1995) Angiospermae. Ordnung Ranunculales. Fam. Ranunculaceae. II. Systematic Part. In: Hiepko P (Ed.) Natürliche Pflanzenfamilien, ed. 2, 17aIV. Duncker & Humblot, Berlin, 223–519.

Thompson JD, Gibson TJ, Higgins DG (2002) Multiple sequence alignment using ClustalW and ClustalX. Current Protocols in Bioinformatics 00(1): 2.3.1–2.3.22. https://doi.org/10.1002/0471250953.bi0203s00
A new species of Ranunculus

Tojibaev KSh, Beshko NYu, Popov VA (2016) Botanical-geographical regionalization of Uzbekistan. Botanicheskii Zhurnal 101(10): 1105–1132. https://doi.org/10.1134/S000681361610001X

Vaidya G, Lohman DJ, Meier R (2011) SequenceMatrix: Concatenation software for the fast assembly of multi-gene datasets with character set and codon information. Cladistics 27(2): 171–180. https://doi.org/10.1111/j.1096-0031.2010.00329.x

Vasilchenko IT, Vasileva LI (1985) Endemic and rare plants of Western Gissar. In: Rasteniya Srednei Azii. Leningrad, 42–121.

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, 315–322. https://doi.org/10.1016/B978-0-12-372180-8.50042-1

Yusupov Z, Deng T, Volis S, Khassanov F, Makhmudjanov D, Tojibaev K, Sun H (2020) Phylogenomics of Allium section Cepa (Amaryllidaceae) provides new insights on domestication of onion. Plant Diversity 43(2): 102–110. https://doi.org/10.1016/j.pld.2020.07.008

Zverev AA, Shchegoleva NV, Revushkin AS (2020) Determination of the ecological areas of rare high-mountain Ranunculus species by indicator values methods. Vestnik Tomskogo gosudarstvennogo universiteta. Biologiya = Tomsk State University. Journal of Biology 51: 212–222. https://doi.org/10.17223/19988591/51/12