Research paper

New insights into the evolutionary history of *Megacodon*: Evidence from a newly discovered species

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A B S T R A C T

*Megacodon* is an ideal genus to study speciation and ecological adaptation in the Sino-Himalayan region. The genus contains two species distributed at different elevations and in two separate areas. However, studies of this genus have long been impeded by a lack of fieldwork on one of its species, *Megacodon venosus*. In this study, we collected specimens of two *Megacodon* species and found an extraordinary new species of *Megacodon* in Lushui county of north-west Yunnan province, which we have since named *Megacodon lushuiensis*. We propose new species based on both morphological and molecular evidence. The finding of this new species emphasized the importance of ecological divergence in the divergence of *Megacodon stylophorus* and its parapatric low-elevation *Megacodon* species. To identify genetic determinants that underlie adaptations to different elevations, we characterized transcriptomes of the new species *M. lushuiensis*, which is distributed at low elevations, and *M. stylophorus*, which is distributed at high elevations. Comparative transcriptome analysis identified 8926 orthogroups containing single-copy genes, and 370 orthogroups containing significantly positively selected genes. The set of positively selected genes was enriched into 25 Gene Ontology terms, including “response to water deprivation”, “response to osmotic stress”, and “cellular response to external stimulus”. Our results provide new insights into how ecological adaptation and speciation occurred in *Megacodon* and highlight the role of heterogeneous habitats in the speciation of plants in the Sino-Himalayan region.

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1. Introduction

The Sino–Himalayan region is a temperate biodiversity hotspot with a high level of species endemism (Myers et al., 2000). Much of this diversity and endemism is attributed to the heterogeneous topography of the region and the diverse climates it supports (Shrestha et al., 2018). The formation of plant diversity in this region is synchronous with the uplift of the Qinghai–Tibet Plateau and has been intensified by climatic oscillations, especially the glacial–interglacial cycle during the Quaternary (Muellner-Riehl et al., 2019; Sun et al., 2017; Wen et al., 2014; Wu, 1988). The mechanisms of plant speciation and diversification in this region are complicated and various (Muellner-Riehl et al., 2019; Wen et al., 2014). On the one hand, heterogeneous topography can create geographical barriers, which promote allopatric species divergence (e.g., Luo et al., 2017), and recent climatic oscillations can give rise to habitat fragmentation, which also interrupts the gene flow (e.g., Ma et al., 2017). On the other hand, divergent selection pressures across elevational gradients or heterogeneous habitats can also cause adaptive divergence in sympathy or parapatry (Funk et al., 2016), and even lead to adaptive radiation. This type of ecological divergence has been shown to be an essential speciation process for plants in the Sino-Himalayan region (e.g., Zhao et al., 2016).

*Megacodon* (Hemsrl.) H. Smith is a small genus nested within a polytomy that contains *Swertia* L. and other genera in the *Swertii*-nae (Chassot et al., 2001; Favre et al., 2016). However, *Megacodon* is morphologically distinct from other *Swertiniae* genera by having large campanulate flowers (Ho and Pringle, 1995). *Megacodon* contains two species: *Megacodon stylophorus* (C. B. Clark) H. Smith and *Megacodon venosus* (Hemsrl.) H. Smith (Ho and Pringle, 1995).
*M. stylophorus* is found at high elevations between 3000 and 4400 m a.s.l. in western Sichuan, north-western Yunnan, southern Tibet of China, also in Bhutan, Nepal, and north-east India. *M. venosus* is endemic to China and is mainly found in Sichuan, western Hubei, and northern Chongqing at low to middle elevations between 600 and 2400 m a.s.l (Ho and Pringle, 1995). In contrast, embryology (Xue and Li, 2005), and breeding system (Meng et al., 2016; Zhang et al., 2019), *Swertia* species have extraordinarily disjunctive distributions. The new species of *Megacodon* in Lushui county of north-west Yunnan at 4400 m a.s.l in western Sichuan, north-western Yunnan, southern Lushui and *M. stylophorus* from Daxueshan Mountain were sampled by liquid nitrogen and stored at –80 °C for RNA-seq. Each sample was obtained from one individual.

2.2. DNA extraction, PCR amplification, and sequencing

The Plant Genomic DNA Kit (Tiangen Biotech, Beijing, China) was used to extract total genomic DNA following the manufacturer’s protocol. The nuclear ribosomal internal transcribed spacer (nITS) and two chloroplast markers, *atpB-rbcL* and *trnl-trnf* were amplified and sequenced following the protocols of previous studies (Favre et al., 2016; Matuszak et al., 2016). Primers used in this study for *atpB-rbcL*, *trnl-trnf* and the nITS region were the same as those used in previous studies (see Hoot et al., 1995; Taberlet et al., 1991; Grudinski et al., 2014 for the nITS region). The 25 μL volume polymerase chain reactions (PCRs) contained 1.0 μL of genomic DNA (20 ng/μL), 0.3 μL of each primer (10 μM), 12.5 μL Ex Taq Version 2.0 (Takara, Dalian, China), and 10.9 μL of double-distilled H₂O. PCRs were performed in a C1000 Touch™ Thermal Cycler (Bio-Rad, Hercules, CA, USA), with initiation at 95 °C for 2 min followed by 35 cycles of denaturation at 95 °C for 1 min, primer annealing at 53 – 56 °C for 1 min, primer extension at 72 °C for 1 min and a final extension step at 72 °C for 10 min. Alignments and subsequent manual adjustment were performed in BioEdit 7.2.5 (Hall, 1999) by using the multiple alignment software ClustalW (Thompson et al., 1994).

2.3. Phylogenetic analyses

Maximum likelihood (ML) and Bayesian inference (BI) approaches were used for phylogenetic tree reconstruction with combined sequences from nITS, *atpB-rbcL*, and *trnl-trnf*. Three *Potalia* Aubl. and three *Lisianthius* P. Browne species were set as outgroups (Table S2). In addition, 11 *Gentiana* L. species and 29 *Swertia* species were included as closely related genera (Table S2). ML analyses were performed on the CIPRES Science Gateway (www.phylo.org; Miller et al., 2010) through RAxML-HPC2 v.8.2.8 (Stamatakis, 2014) with 1000 bootstrap replicates. The best-fitting nucleotide substitution model for BI analyses was GTR + G, selected using the Akaike information criterion (AIC) in jModeltest 2.1.7 (Darriba et al., 2012). BI analyses were also performed on CIPRES using MrBayes v.3.2.6 (Ronquist et al., 2012) on XSEDE with the following settings: *ngen = 100,000, samplefreq = 1000, printfreq = 1000 and sumt burnin = 2500.*

2.4. Climatic data analysis

Our fieldwork identified 15 precise locations where *Megacodon* is distributed. We used the ‘raster’ package to extract 19 bioclimatic variables for these sites from WorldClim (http://www.worldclim.org; Hijmans et al., 2005). Principal component analysis (PCA) of these environmental factors was performed by the dudi.pca function of the package ade4® in R 3.5.1 (R Core Team, 2018). We conducted paired two-tailed t-tests in Microsoft® Excel® 2016 (Microsoft Company, USA) to characterize factors that diverged between *M. stylophorus* and the other two *Megacodon* species (including the suspected new species from Lushui).

2.5. Transcriptome sequencing and functional annotation

TRlzol reagent (Sigma–Aldrich) was used to extract total RNA of samples following the manufacturer’s instructions. A total amount of 3 μg of RNA per sample was used to construct a cDNA library with a fragment length range of 250–300 bp. The cDNA libraries were
created with NEBNext® Ultra™ RNA Library Prep Kit for Illumina® (NEB, USA), following the manufacturer’s recommendations. Illumina Hiseq 2500 was used for sequencing the library preparations, and paired-end reads were generated. Library preparation and Illumina sequencing were performed at Novogene Bioinformatics Technology Co., Ltd., Beijing, China (www.novogene.cn). Clean data was obtained by trimming read adapter sequences, and filtering low-quality reads from raw data. De novo assembly of each individual was accomplished using the Trinity program (Raychowdhury et al., 2011) with min_kmer_cov set to 2, and all other parameters set to default.

Functions of all assembled unigenes were annotated based on the following databases: Nr (NCBI non-redundant protein sequences); Nt (NCBI non-redundant nucleotide sequences); Pfam (Protein family); KOG/COG (Clusters of Orthologous Groups of proteins); Swiss-Prot (A manually annotated and reviewed protein sequence database); KO (KEGG Ortholog database); and GO (Gene Ontology).

2.6. Homologous gene identified and positive selection analysis

We used OrthoFinder software v. 2.2.7 (Emms and Kelly, 2018, 2015) to identify homologous gene clusters (orthogroups). OrthoFinder was run with multiple sequences alignment (MSA) for gene tree inference and RAxML for tree inference method (-T ramxl -M msa). Orthogroups of single-copy genes (1:1 orthologs) were obtained for substitution rates estimation. Each orthogroup was aligned by Prank (Löytynoja and Goldman, 2005) at the condon level (with options “-F-codon”). To estimate a general evolutionary pattern of selective pressure for the two species, codeml program from PAML 4.9 (Yang, 2007) was used to calculate the \( d_N, d_S \) and \( d_N/d_S \) with the free ratio model (model = 2, NSsites = 0) and ML pairwise alignment (runmode = -2). To reduce false-positive findings in subsequent evolutionary analyses, genes with length less than 150 bp, \( d_S < 0.001 \) (to avoid a very large \( d_N/d_S \) ratio) and \( d_S > 0.1 \) (to avoid suspicious paralogs) were discarded (Bustamante et al., 2005; Sun et al., 2018; Zhang et al., 2013).

Positive selection genes (\( d_N/d_S > 1 \)) were used as the test set for the GO enrichment analysis, which was performed on web-based KOBAS software (Xie et al., 2011) by Fisher’s exact test, while whole single-copy genes were selected as the background set. Significant enriched GO terms (\( P < 0.05 \)) were selected and further analyzed using REVIGO (http://revigo.irb.hr; Supek et al., 2011) to investigate in-depth GO similarities.

3. Results

3.1. Morphological comparisons and molecular variation

Comparative analysis of morphological traits indicates that the Megacodon species from Lushui is closely related to M. venosus (Fig. 2 and Table 1). The Megacodon species from Lushui is similar to M. venosus by its narrow leaf blades shape, calyx shape and the existence of glands at the base of calyx, but differs from M. venosus by its taller and thinner plant, lower stem leaf blades elliptic-spathulate to elliptic-obovate (vs. elliptic-lanceolate), middle to upper stem leaf blades veins 3–5 (vs. 5–7), 11–15-flowered thyrses (vs. 7–11 flowered), and dense glands at the 1/3 base of the calyx.

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**Fig. 1.** Geographic distributions of *Megacodon* samples. The red pentagram represents the putative new species (ML) in Lushui county, north-west Yunnan. The red dot represents the *M. venosus* (MV) population from northern Chongqing.
Two individuals from Lushui and two individuals from *M. venosus* differed by a total of eleven nucleotide substitutions and six insertions/deletions in the alignment dataset (including ITS, *atpB-rbcL, trnL-trnF*, Table S3).

### 3.2. Phylogenetic relationships of Megacodon

The data set of phylogenetic analyses comprised 62 accessions with 2383 characters. Because the topologies of phylogenetic trees evolved from the putative new species *Megacodon lushuiensis* (A-E) and *Megacodon venosus* (F-J). A and F. Habitat and habit. B and G. Upper stem leaf blades. C and H. Middle stem leaves. D and I. Flower. E and J. Calyx. (A-D photographed by Jun-chu Peng; E-J photographed by Yi Yang).

**Table 1**

|                | *M. lushuiensis* | *M. venosus* | *M. stylophorus* |
|----------------|-----------------|--------------|------------------|
| **Habit**      | Plants 130–240 cm tall | Plants 45–85 (–180) cm tall | Plants 30–60 (–100) cm tall |
| **Stems**      | Basal to middle stems erect, terete, 1.1–1.8 cm in diam at base; upper stems slim 3–9 mm in diam | Stems erect, terete, stout, 1–1.5 cm in diam | Stems erect, terete, stout, 1–1.5 cm in diam |
| **Lower stem leaf blades** | elliptic-lanceolate to elliptic-ovobovate | elliptic-lanceolate | elliptic to ovate-elliptic |
| **shape**      | 5–7              | 5–7          | 5–9              |
| **veins**      | 5–7              | 5–7          | 5–7              |
| **Middle stem leaf blades** | elliptic-lanceolate to lanceolate | elliptic-lanceolate | elliptic to ovate-elliptic |
| **shape**      | 15–45 × 5–14 cm  | 10–30 × 3–6 cm | 7–22 × 3–9 (–14) cm |
| **size**       | 3–5              | 5–7          | 7–9              |
| **Upper stem leaf blades** | linear-lanceolate to linear | linear-lanceolate | ova-lanceolate |
| **shape**      | 5–10 × 1.1–1.5 cm | 5–7 × 0.7–1.2 cm | 5–10 × 1.2–3 |
| **size**       | 3–5              | 5–7          | 7–9              |
| **Thyrses**    | 11–15-flowered   | 7–11-flowered | 2–8-flowered     |
| **Pedicel**    | 3–9 cm           | 2.5–6 cm     | 3–6 (–16) cm     |
| **Calyx**      | elliptic-lanceolate, apex acuminate | elliptic-lanceolate, apex acuminate | oblong-spatulate, apex obtuse |
| **lobes shape** | 1.7–3.6 cm       | 2.5–3 cm     | 2–2.6 cm         |
| **lobes length** | 4–6 mm           | 2–3 mm       | 6–8 mm           |
| **Nectary**    | Dense pustulose glands at the base ca. 1/3 | Sparse pustulose glands at the base ca. 1/2 | No glands |
| **Corolla size** | ca. 5–7 × 4–5 cm | 5–6 × 6–7.5 cm | 5–7 × 4–5 cm |
| **Corolla lobes shape** | oblong-spatulate | oblong-spatulate | oblong-spatulate |
| **Corolla lobes length** | 4–5.8 cm          | 4–5 cm       | 4–6 cm           |
| **Anthers**    | 8–11 mm          | 7–10 mm      | 1–1.2 cm         |
| **Style**      | ca. 1.7–2.0 cm   | 1.5–1.8 cm   | 1.5–1.8 cm       |
| **Phenology**  | Sep. to Nov.     | Sep. to Nov. | Jun. to Sep.     |
| **Distribution** | 1700 m, NW Yunnan | 600–2400 m, W Hubei, Sichuan, N Chongqing | 3000–4400 m, S Sichuan, SE Tibet, NW Yunnan [Bhutan, NE India, Nepal] |
from ML and BI analyses were mostly identical, only the ML tree is shown with the posterior probabilities (PP) from BI analysis (Fig. 3). Phylogenetic inference showed that Megacodon is most likely a monophyletic group (ML-BS = 100/BI-PP = 1.00 in Fig. 3). Twelve accessions of M. stylophorus from Tibet and Yunnan formed a well-supported clade (96/1.00). In comparison, two accessions of M. venosus from Chongqing and two accessions from Lushui formed the other clade of Megacodon. The population of M. venosus and the species from Lushui each formed a monophyletic clade in this branch.

3.3. Taxonomic treatment

According to both morphological and molecular evidence, as well as the extraordinary disjunctive distribution with M. venosus, we propose the Megacodon plants from Lushui as a new species here.

**M. lushuiensis J. C. Peng & H. Sun sp. nov.** (Fig. 2A–E, and Fig. 4).

**Holotype.** China, Yunnan Province: Laowo Xiang, Lushui County, on the hillside, 1700 m a.s.l, 25.85°N, 99.03°E, flowering, 21 September 2017, J. C. Peng & L. Sun PM001 (KUN).

**Paratype.** The same locality, flowering and fruiting, 7 November 2017, J. C. Peng, L. Sun & Z. Chen PM002 (KUN).

**Description.** Plant perennial, herbaceous ca. 130–240 cm tall. Basal to middle stems erect, terete, 1.1–1.8 cm in diam at base; upper stems slim, 3–9 mm in diam. Lower stem leaves elliptic-spathulate to elliptic-obovate, base cuneate and decurrent, veins 5–7 and arcuate; middle stem leaves elliptic-lanceolate to lanceolate, apex acuminate, 15–45 × 5–14 cm, veins 3–5 and arcuate; upper stem leaves linear-lanceolate to linear, 5–10 × 1.1–1.5 cm, veins 3–5 and arcuate. Thyrses 11–15-flowered. Pedicel 3–9 cm. Calyx campanulate, 1.7–3.6 cm, tube 4–6 mm; lobes elliptic-lanceolate, apex acuminate, dense pustulous glands at the base.

**Fig. 3.** Molecular phylogeny of Megacodon and four other genera (Swertia, Gentiana, Lisianthius, and Potalia) based on nrITS, atpB-rbcL, and trnL-trnF sequences. ML bootstrap support values (BS) and Bayesian posterior probabilities (PP) are indicated on nodes and separated by a slash. Values for nodes with less than 75 (BS) or 0.85 (PP) are not shown. **M. lushuiensis** are shown bold.
ca. 1/3. Corolla white to pale yellow, with green veins, ca. 5–7 × 4–5 cm, tube 8–10 mm; lobes oblong-spatulate, 4–5.8 cm, apex rounded. Filaments flat and linear, 2.2–2.8 cm, alternate with lobes; anthers oblong-quadrate, 8–11 mm; Capsule ovoid-ellipsoid, 1.7–2.0 cm.

**Phenology.** Flowering and fruiting from Sep. to Nov.

**Distribution and habitat.** *M. lushuiensis* was found in Laowo Xiang of Lushui County, north-west Yunnan, south-west China (Fig. 1), growing on hillsides (north-west slope) and under shrub at ca. 1700 m a.s.l.

**Conservation status.** *M. lushuiensis* is only known from one population in Lushui county (criterion B2a), with an area of occupancy estimated to be less than 500 km² (criterion B2b (i, ii)). The habitat of this population is near to cultivated field (less than 2 km) and suffering from invasive plants such as *Ageratina adenophora* (criterion B2b (iii)). The new species should be regarded as “Endangered” based on the IUCN (2012) criteria.

**Etymology.** The specific epithet refers to the type locality, Lushui County of Yunnan Province, in China.

### 3.4. Climatic niche

PCA of 19 bioclimatic variables indicated clear climate niche differentiation between 13 *M. stylophorus* sites, one *M. venosus* site, and one *M. lushuiensis* site (Fig. 5). The first two PCs identified two climatic groups and explained 59% and 26.2% of the total variation, respectively (Fig. 5). The climatic niche of *M. lushuiensis* is closer to the population of *M. venosus* from Chongqing, whereas 13 sites of *M. stylophorus* formed another group. Among these 19 bioclimatic variables, 12 bioclimatic variables (BIO1, BIO5, BIO6, BIO8–BIO12, BIO14, BIO15, BIO17, and BIO19) showed significant differences

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Fig. 4. *Megacodon lushuiensis* J. C. Peng & H. Sun sp. nov. (drawn by Wen Sha). A. Habit B. Flower C. Opened corolla lobes D. Partial calyx.
between *M. stylophorus* and the other two species (*P* < 0.01; Table S4).

### 3.5. Overview of RNA-seq analysis

We generated 55.2 and 55.3 million clean reads of RNA-seq data for *M. lushuiensis* and *M. stylophorus*, with a total size of 8.28 and 8.30 Gb, respectively (NCBI GenBank accession number: SRR11743766, SRR11743765 and Table 2). The median unigene length for *M. lushuiensis* after de novo assembly was 981 bp, with an N50 length of 2043 bp. The median unigene length for *M. stylophorus* was 1018 bp, with an N50 length of 2058 bp. A total of 24,086 unigenes from *M. lushuiensis* and 23,262 unigenes from *M. stylophorus* were annotated in at least one database with a significant match (e-value < 10^-5).

### 3.6. Orthogroups identified and positive selection analysis

In this study, a total of 20,679 and 20,471 unigenes were detected in *M. lushuiensis* and *M. stylophorus*, respectively, and 23,611 orthogroups were identified. Among these, 8926 orthogroups contained putative one-to-one single-copy genes between two species. To detect genes that are involved in ecological adaptation to high elevations, *d*<sub>K</sub>, *d*<sub>σ</sub>, and *d*<sub>N</sub>/*d*<sub>S</sub> ratio of single-copy genes were calculated by PAML. A peak of *d*<sub>S</sub> distribution between *M. stylophorus* and *M. lushuiensis* was observed at 0.0411 ± 0.0210 (Fig. 6A). The *d*<sub>N</sub>/*d*<sub>S</sub> of most genes was less than one (Fig. 6B), which indicates purifying selection. Only 370 single-copy genes were identified as positively selected genes (PSGs, *ω* > 1) between two species.

In enrichment analyses, the 370 positively selected genes were designated as the test sets, and 7749 single-copy genes (0.001 < *d*<sub>S</sub> < 0.1) were designated as the background set. Twenty-five GO categories were annotated to 49 pairs of positively selected genes, which were overrepresented (Fisher’s exact test, *p* < 0.05) in the test sets (Table 3) by GO enrichment analysis in the KOBAS website. Then based on GO similarities, five clusters of biological processes were shown in the REVIGO treemap (Fig. 6C). A majority of biological processes in treemap were found to be related to specific adaptation traits, such as response to water (11 PSGs), regulation of response to osmotic (3 PSGs), cellular response to nutrient levels (7 PSGs), and cellular response to external stimulus (7 PSGs) (Fig. 6C, Table 3 and Table S5).

### 4. Discussion

#### 4.1. New insights into the evolutionary history of Megacodon

In this study, we described a new species of *Megacodon*, *M. lushuiensis*, and determined the phylogenetic relationships among *Megacodon* species. The three existing *Megacodon* species formed a well-supported monophyletic clade (Fig. 3). All samples of *M. stylophorus* formed a monophyletic clade. Furthermore, in our phylogenetic tree, *M. lushuiensis* is closely related to *M. venosus*, indicating a close evolutionary relationship between these two species.

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**Table 2** Statistics of clean data and unigenes of RNA-seq.

| Species          | Statistics of clean data | Frequency distribution of unigene lengths |
|------------------|--------------------------|------------------------------------------|
|                  | Clean Reads             | Clean Bases (G) | Error (%) | Q20 (%) | Q30 (%) | GC Content (%) | Total number | min length (bp) | mean length (bp) | median length (bp) | max length (bp) | N50 (bp) | total nucleotides (bp) |
| *M. lushuiensis* | 55,224,736              | 8.28         | 0.03       | 96.3     | 90.2    | 43.2    | 35,573       | 301           | 1400             | 981               | 16,580             | 2043           | 49,784,866       |
| *M. stylophorus* | 55,301,198              | 8.30         | 0.03       | 96.7     | 91.0    | 43.5    | 33,202       | 301           | 1420             | 1018              | 15,568             | 2058           | 47,132,153       |

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**Fig. 5.** PCA analysis of bioclimatic factors of the putative new species (*M. lushuiensis* black square), *M. stylophorus* (red dots), and *M. venosus* (blue triangle). Each arrow represents a projection of a bioclimatic variable.
Fig. 6. The $d_{s}$ (A) and $d_{ls}/d_{s}$ ratio (B) distribution of orthologs between M. stylophorus and M. lushuiensis. REVIGO treemap summarizing GO biological process categories overrepresented regarding ecological divergence of M. lushuiensis and M. stylophorus (C). KOBAS was used to identify GO biological processes that were overrepresented at p-values (Fisher's exact test) $< 0.05$. Similar colors indicate semantic similarity related to a similar functional category. The size of each rectangle is proportional to the p-value (Fisher's exact test) for that category. TS: frequency in the test set; BS: frequency in the background set.

Table 3: Partial list of positively selected genes related to ecological adaptation.

| Orthologs | Arabidopsis thaliana gene accession | Description                                                                 | $d_{ls}/d_{s}$ |
|-----------|-----------------------------------|-----------------------------------------------------------------------------|----------------|
| OG0007681 | AT1G32230                         | RCD1, WWE protein–protein interaction domain protein family                  | 1.99           |
| OG0010334 | AT3G17810                         | PYD1, pyrimidine 1                                                          | 1.31           |
| OG0002624 | AT2G39550                         | PGFL-I, Prenyltransferase family protein                                    | 1.05           |
| OG0009542 | AT1G17755                         | LEU1, Undecaprenyl pyrophosphate synthetase family protein                  | 1.10           |
| OG0009326 | AT3G808040                        | FRD3, MATE efflux family protein                                            | 1.12           |
| OG0008777 | AT3G33630                         | EX1, UvrB/UvrC domain protein                                               | 1.15           |
| OG0009854 | AT3G63060                         | EDL1, EID1-like 3                                                           | 1.15           |
| OG0003758 | AT1G06770                         | DRIP1, DREB2A-interacting protein                                           | 1.16           |
| OG0008137 | AT4G36020                         | CSIP1, cold shock domain protein                                            | 1.15           |
| OG0009134 | AT5G28770                         | BZIP28, BZIP transcription factor family protein                            | 1.75           |
| OG0004371 | AT3G62420                         | BZIP53, basic region/leucine zipper motif 53                               | 1.42           |
| OG0006371 | AT3G48170                         | ALDH10A9, aldehyde dehydrogenase family protein                            | 1.51           |
which is consistent with the morphological similarity between these two species. The phylogenetic relationship between *Megacodon* and four other lineages (*Potentilla, Lisianthus Gentiana*, and *Swertia*) is congruent with previous studies (Favre et al., 2016). Although sharing many morphological similarities, *M. lushuiensis* can be easily distinguished from *M. venosus* by its leaf blade, inflorescence, extrafloral nectaries, and more. Importantly, the extrafloral nectaries in the basal portion of the calyx were neglected in previous morphological descriptions of *M. venosus*. The extrafloral nectaries of *M. venosus* and *M. lushuiensis* not only secrete nectar but also cover the petal and make it sticky (Fig. 2D, E, I, and J). We speculate that in addition to attracting ants to participate in ant-plant mutualisms, the secretions from *M. venosus* and *M. lushuiensis* extrafloral nectaries may directly participate in the protection of the flowers (Marazzi et al., 2013).

The exact distribution of *M. lushuiensis* in north-west Yunnan indicates that the most recent common ancestor of *M. lushuiensis* and *M. venosus* may occupy a more extensive area, from west Yunnan to central China. In the past, *Megacodon* populations from the Nujiang River valley may have been connected to *M. venosus* populations from central China through the eastern edge of the Qinghai-Tibet Plateau region or the Yunnan-Guizhou Plateau. The current fragmented distribution of this species pair may be the result of a vicariance event closely related to the climatic changes of the Sino-Himalayan region and its low reproductive capacity. Because of its humid climate and large elevational gradient, the Nujiang River valley harbors many distinct species that are identical to or sister to species in central or south China, such as *Icthyoselminus macrantha* (Oliver) Lidén, *Vysilandra Franch.*, *Asteropyrum peltatum* (Franch.) Drumm. ex Hutch. The large elevational gradient and complex climate conditions in Nushan Mountain should provide more ecological opportunities for the maintenance of relict populations or species. Despite this, the fragmented distribution pattern of *M. venosus* and *M. lushuiensis* is still rare and may have important significance in biogeographical studies.

The occurrence of *M. lushuiensis* in north-west Yunnan also alters our understanding of the divergence of *M. stylophorus* and related species. Speciation of *M. venosus* and *M. stylophorus* was previously thought to have occurred through allopatric speciation, similar to that of *Pleuroserpurn hookeri* C. B. Clarke and *P. giralldii* Diels, which have a fragmented distribution between central China and the Qinghai-Tibet Plateau region (Bai et al., 2015). However, the fragmented distribution of *M. lushuiensis* and *M. stylophorus* occur at different elevations along the Nushan Mountain, suggesting that ecological differentiation may have played an important role in their divergence.

4.2. Functions under positive selection between *M. lushuiensis* and *M. stylophorus*

To further investigate why *M. lushuiensis* and *M. stylophorus* occupy distinct ecological niches along an elevational gradient, we used comparative transcriptome analysis. The majority of genes identified by GO enrichment of *M. lushuiensis* and *M. stylophorus* function in response to water deprivation (GO:0009414) and other external stimuli (e.g., GO:0047484 and GO:0071436) (Fig. 6C). These findings suggest that genes that respond to the environment may have been triggered by natural selection during ecological adaptation (Table 3). However, unlike many other alpine plants, especially alpine scree plants (Chen et al., 2019; Guo et al., 2018; Jia et al., 2019; Zhang et al., 2019), *M. stylophorus* does not appear to have adapted to high radiation. This may be because *M. stylophorus* grows in the forest or scrub habitat. Analysis of climatic variables revealed that *M. stylophorus* favors the colder and higher climatic drought conditions of high elevations (Fig. 5 and Table S1). In contrast, field observations indicate that *M. lushuiensis* prefers to grow in drier soil conditions such as semi-humid hillside at low elevations (Fig. 2A). Habitat differentiation between these two species may have led some populations to experience water deficiency at different periods, which can influence plant growth and development. Our transcriptome analysis identified several genes (e.g., PGGT-1, ALDH110A9, and EDL3) under positive selection that is directly involved in water deprivation and hormone signaling pathways (Johnson et al., 2005; Koops et al., 2011; Missihoun et al., 2014). *DRIP1* is a negative regulator in drought tolerance-responsive gene expression that was also found to be under positive selection (Qin et al., 2008). Our study also found that the following genes were under positive selection: *LEW1*, which response to endoplasmic reticulum stress, drought, and dark-induced senescence by catalyzing dolichol biosynthesis (Zhang et al., 2008); *RCD1*, which is involved in the regulation of stomatal conductance and the flavonoid accumulation pathway (Ahflors et al., 2004; Suvorov et al., 2017); and *CSDP1*, which contains a cold shock domain and is involved in cold and salt stress responses (Yang and Karlson, 2013). The functions of these homologous genes emphasize that the genetic pathways that regulate efficient water use may have been important drivers of the divergence of *M. lushuiensis* and *M. stylophorus*.

Under a short growing season in the alpine region, *M. stylophorus* also require proper phenological responses. Our transcriptome analysis identified several genes that may play roles in these phenological responses. For example, in addition to its response to water deprivation, *CSDP1* can also regulate seed germination (Yang and Karlson, 2013). Furthermore, *bZIP53* is an enhancer for seed maturation gene transcription (Alonso et al., 2009), while *BZO2H3* (also known as bZIP62) is involved in the regulation of a circadian oscillator gene *PRR7* (Frank et al., 2018). The correct circadian clock guarantees that plants have photosynthetic advantages and can accurately anticipate environmental changes (Dodd et al., 2014, 2005). These homologous genes may strengthen the ability of *M. stylophorus* to respond to habitat-specific variability. Similarly, *EX1* and *FRD3*, which were identified in our analysis, may help *M. stylophorus* tolerate higher oxidative stress and iron deficiency that result from radical-initiating factors at high elevations (Dogra et al., 2019; Lei et al., 2014), especially at the end of the growing period (Sakata et al., 2006). Overall, adapting to extreme environments (such as climatic drought, oxidative stress, and low temperature) in Sino-Himalayan regions may play an important role in the speciation of *Megacodon*.

5. Conclusion

In this study, we used field observations in conjunction with morphological and molecular data to propose a new *Megacodon* species from the Nujiang River valley, *M. lushuiensis*. The discovery of this species calls attention to the importance of ecological divergence between *M. stylophorus* and low-elevation *Megacodon* species. We also presented the phylogenetic position of *M. venosus* for the first time and described a neglected trait in this species (sparse glands at 1/2 base of the calyx). Although population genetic and genomic studies are essential to understanding the demographic and divergence history of *Megacodon*, our analysis of the transcriptomes of *M. stylophorus* and *M. lushuiensis* provide valuable resources for the future genomic research and general insight into how ecological divergence occurred between these two *Megacodon* species. *M. venosus* and *M. lushuiensis* are both extremely rare in the
field; we, therefore, recommend that conservation efforts be carried out timely and properly for these two species.

Author contributions
HS, XGM and YHW conceived and designed experiments. JCP performed field investigations, analyzed the data, and wrote the manuscript. XGM and HS revised the manuscript.

Declaration of Competing Interest
We declare that we do not have any commercial or associative interest that represents a conflict of interest in connection with the work submitted.

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Appendix A. Supplementary data
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