Supplementary Materials

Supplementary Materials and Methods

Sampling

During 2014 and 2017, we conducted a joint Sino-Kazh field investigation on the herpetofauna in eastern Kazakhstan and the adjacent Junggar Depression in China. Thirty specimens (15 males, 15 females) from seven sites were collected from eastern Kazakhstan and northwestern Altay region, Xinjiang, China (Supplementary Figure S1 and Table S1), which were initially identified as *E. multiocellata*. The lizards were euthanized with an overdose of sodium pentobarbital via intraperitoneal injection, and liver or tail tissues were extracted and preserved in 95% ethanol following the animal use protocols approved by the Chengdu Institute of Biology (CIB), Chinese Academy of Sciences (CAS). All voucher specimens are preserved in CIB, CAS.

Molecular data collection

Genomic DNA was extracted from liver or tail tissues following the modified high-salt protocol proposed by Aljanabi & Martinez (1997). Fragments of the mitochondrial COI gene (709 bp) were amplified via polymerase chain reaction (PCR) using primers designed according to Ward et al. (2005) with minor modifications (E COIF 5'-TCAACCAACCACAAAGACATTGGCAC-3' and E COIR 5'-TAGACTTCTGGGTGGCCAAAGAATCA-3'). The PCR reaction mixture contained 12.5 µL of 2 × EasyTaq SuperMix (Tsingke Biol-Tech, Chengdu, China), 0.2 μ M of each primer, and 1–2 μ L of genomic DNA for a total volume of 25 μ L. The amplification protocols included an initial denaturation at 94 °C for 4 min, followed by 35 cycles of 94 °C for 30 s, 52 °C for 30 s and elongation at 72 °C for 50 s, and a final extension at 72 °C for 10 min. All PCR products were commercially purified and sequenced for double strands with the primers used for amplification. All novel deposited GenBank under the accession numbers sequences were in MW172523-MW172552.

Nineteen individuals from China and 11 from Kazakhstan were sequenced for *COI*. All novel sequences were translated to amino acids with MEGA v7.0.26 (Kumar et al., 2016), and no stop codons were detected. In total, 156 sequences – which included those from Orlova et al. (2017) (available in GenBank) – were aligned using Clustal X v2.0 (Larkin et al., 2007) with default settings. Identical sequences were collapsed into a single haplotype using DnaSP v5.0 (Librado & Rozas, 2009).

DNA barcoding diversity analysis

The dataset containing all sequences was uploaded into the BOLD system under the project '*Eremias multiocellata-przewalskii* complex barcoding' (Barcode of Life Data System website, www.barcodinglife.com; Ratnasingham & Hebert, 2007). We measured the genetic distance with the Kimura 2-parameter (K2P; Kimura, 1980) model, which has been widely used in barcoding studies. The intraspecific and intra-genus pairwise distances were calculated with the 'Distance Summary' analysis tool in BOLD. The maximum intraspecific genetic distance and corresponding minimum distance to nearest neighbor at the species level were calculated and compared using the 'Barcoding Gap Analysis' tool in BOLD.

Phylogenetic analysis

The neighbor-joining (NJ) (Saitou & Nei, 1987) tree was constructed in MEGA with a gamma distribution rate variation among sites (shape parameter = 4) and pairwise deletion option for the treatment of missing data. Bootstrap support for clades was evaluated by nonparametric bootstrap analysis (Felsenstein, 1985) with 1 000 replicates. Bayesian inference (BI) was conducted with MrBayes v3.2.2 (Ronquist et al., 2012). The best-fit models of nucleotide substitution for each partition scheme were selected using PartitionFinder v2.1.1 (Lanfear et al., 2017). Three codon partitions and their corresponding substitution model for the *COI* gene sequences were proposed: first codon, K80+G+I; second codon, F81; third codon, GTR+G. Two parallel runs of one cold and three heated Monte Carlo Markov chains (MCMCs) were performed with sampling every 200 generations and were automatically stopped

after reaching a mean standard deviation between split frequencies of less than 0.01. Parameters and topologies were estimated from 13 405 steps after discarding the first 30%. Convergence of the runs was assessed by effective sample size (ESS \geq 200) with Tracer v1.7.1 (Rambaut et al., 2018). Partitioned ML (Maximum Likelihood) analyses were conducted in RAxML v8.2.4 (Stamatakis, 2014) with the GTR+G model for all subsets. A complete random starting tree (option -d) was initiated with 100 independent search replicates where the tree with the best likelihood was chosen. Bootstrap support proportion (BSP) for the clades was obtained with a sufficient number of bootstrap replicates (450), which was automatically determined by setting the default cutoff threshold (option -# autoMRE).

Species delimitation

We first employed two distance-based methods dedicated to barcoding datasets. The Barcode Index Number (BIN) system clusters (Ratnasingham & Hebert, 2013) was conducted with the 'BIN Discordance' option in BOLD, which was employed to assign individuals to operational taxonomic units (OTUs) that represent presumable species. The BIN system uses the refined single linkage algorithm (RESL) for clustering process, and hence is independent of any prior taxonomic assignment. Similarly, Automatic Barcode Gap Discovery (ABGD; Puillandre et al., 2012) does not need a priori species hypotheses, and clusters sequences into candidate species by detecting a 'barcode gap' that demarcates intra- and interspecific distance. Analysis implemented ABGD web was in the server (https://bioinfo.mnhn.fr/abi/public/abgd/abgdweb.html) with the K2P model and other parameters as default.

Two coalescent methods were further employed to delimit species: the generalized mixed Yule coalescent (GMYC) model implemented in the GMYC web server (http://species.h-its.org/gmyc/), and Bayesian species identification under the multispecies coalescent (MSC) model with the program BPP v3.3 (Yang & Rannala, 2010). GMYC was proposed to delimit species using the reconstructed gene tree for a single locus (Fujisawa & Barraclough, 2013). We reconstructed ultrametric gene trees

in BEAST v1.8.2 (Drummond et al., 2012) with three codon partitions. The substitution models were calculated in PartitionFinder; the first codon used K80+G+I, the second used HKY, and the third used TRN+G. To reduce compounding errors from time calibration uncertainties or molecular substitution rates, we left the prior 'clock.rate' to default 1.0. Posterior samplings were drawn every 1 000 steps over 15 million MCMC steps with the uncorrelated lognormal relaxed clock model and Yule speciation tree prior. Two independent runs were performed. Stationarity of the Markov chains for each run was checked using Tracer for $ESS \ge 200$. Finally, the tree files were combined using LogCombiner v1.8.2, and a maximum clade credibility (MCC) tree with median node heights was generated with TreeAnnotator after discarding the first 25% of posterior sampling as burn-in. The MCC tree was used as an input in the single-threshold GMYC analysis implemented in the GMYC server. Bayesian species identification under the MSC model was considered to have better performance in species assignment than other single-locus delimiting methods for DNA barcoding analyses (Yang & Rannala, 2017). We ran the A11 model with the presumption of relatively large ancestral population sizes ($\theta \sim G$ (1, 200)) and deep divergence ($\tau_0 \sim G$ (1.5, 10)); this single set of priors was based on our unpublished data (Liu, 2019). To explore the potential cryptic species within our study system, we designated the significantly supported lineages within E. stummeri (two reciprocal monophyletic lineages) and Eremias sp. 3 (two reciprocal monophyletic lineages and a singleton) as different unique populations in our prior population assignments in the imap file. The reversible-jump Markov chain Monte Carlo (rjMCMC) algorithm 1 with parameters combination $(\alpha, m) = (1, 0.5)$ and species model prior = 1 were used to repeat the BPP runs twice with different starting seeds to assess the consistency between the different runs. Each run was conducted with rjMCMC analysis for 500 000 generations (sampling interval of five), with the first 20 000 samples discarded as burn-in.

In addition, species delimitation with the multi-rate Poisson tree processes model (mPTP) has been shown to have reliable performance (Blair & Bryson, 2017), as it explicitly accounts for differences in sampling intensity and/or effective population

sizes of species (Kapli et al., 2017). We implemented mPTP analysis in the web server (<u>https://mptp.h-its.org</u>) using the MCC tree constructed in the previous BEAST analysis.

All species delineation analyses described above were conducted with haplotypes (except BIN, which used all sequences) and without outgroups.

Morphological analyses

Morphological measurement was performed on all 30 specimens collected in this study, following the traits analyzed in Szczerbak (1974), Dujsebayeva et al. (2009), and Orlova et al. (2017). Morphological analyses were conducted with 11 metric and 12 meristic traits (Supplementary Table S5). Of these traits, four meristic traits (Supplementary Table S5) were counted on both the left and right sides of the body. Metric measurements were conducted using a digital caliper and recorded to two decimal places.

We first employed analysis of variance (ANOVA) to test for sexual dimorphism in both metric and meristic datasets collected in this study. All metric traits (except Dist.P.fm) and one meristic trait (Sq.c.cd) exhibited significant sexual dimorphism (p< 0.05). As a result, the comparative demonstration of these sex-dependent characters (values given as ranges and means ± SE) between the individuals from this study and the holotype and paratypes of *E. dzungarica* from Orlova et al. (2017) were given separately for males and females. The values of the other age- and sex-independent metric and meristic characters were calculated for all specimens combined (including males, females and juveniles in this study and Orlova et al. (2017), respectively). Moreover, to test whether the specimens in this study differed significantly from *E. dzungarica* in morphological traits with a sample size greater than 11, we performed a two independent samples *t*-test. All morphometric analyses were performed with SPSS 19 (SPSS Inc., Chicago, USA).

REFERENCES

- Aljanabi SM, Martinez I. 1997. Universal and rapid salt-extraction of high quality genomic DNA for PCR-based techniques. *Nucleic Acids Research*, 25(22): 4692–4693.
- Blair C, Bryson Jr RW. 2017. Cryptic diversity and discordance in single-locus species delimitation methods within horned lizards (Phrynosomatidae: *Phrynosoma*). *Molecular Ecology Resources*, **17**(6): 1168–1182.
- Drummond AJ, Suchard MA, Xie D, Rambaut A. 2012. Bayesian phylogenetics with BEAUti and the BEAST 1.7. *Molecular Biology and Evolution*, **29**(8): 1969–1973.
- Dujsebayeva TN, Chirikova MA, Belyalov OV. 2009. New finds of the racerunner of *Eremias multiocellata* complex in Kazakhstan. *Russian Journal of Herpetology*, 16(1): 51–56.
- Felsenstein J. 1985. Confidence limits on phylogenies: an approach using the bootstrap. *Evolution*, **39**(4): 783–791.
- Fujisawa T, Barraclough TG. 2013. Delimiting species using single-locus data and the Generalized Mixed Yule Coalescent approach: a revised method and evaluation on simulated data sets. *Systematic Biology*, **62**(5): 707–724.
- Kapli P, Lutteropp S, Zhang J, Kobert K, Pavlidis P, Stamatakis A, et al. 2017.
 Multi-rate Poisson tree processes for single-locus species delimitation under maximum likelihood and Markov chain Monte Carlo. *Bioinformatics*, 33(11): 1630–1638.
- Kimura M. 1980. A simple method for estimating evolutionary rates of base substitutions through comparative studies of nucleotide sequences. *Journal of Molecular Evolution*, **16**(2): 111–120.
- Kumar S, Stecher G, Tamura K. 2016. MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets. *Molecular Biology and Evolution*, **33**(7): 1870–1874.
- Lanfear R, Frandsen PB, Wright AM, Senfeld T, Calcott B. 2017. Partitionfinder 2:

new methods for selecting partitioned models of evolution for molecular and morphological phylogenetic analyses. *Molecular Biology and Evolution*, **34**(3): 772–773.

- Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, et al. 2007. Clustal W and Clustal X version 2.0. *Bioinformatics*, 23(21): 2947–2948.
- Librado P, Rozas J. 2009. DnaSP v5: a software for comprehensive analysis of DNA polymorphism data. *Bioinformatics*, **25**(11): 1451–1452.
- Liu J. 2019. Molecular Phylogeny and Phylogeography of *Eremias velox* Complex and *Eremias multiocellata* Complex. Ph.D. dissertation, Sichuan University, Chengdu. (in Chinese)
- Orlova VF, Poyarkov Jr NA, Chirikova MA, Nazarov RA, Munkhbaatar M, Munkhbayar K, et al. 2017. MtDNA differentiation and taxonomy of Central Asian racerunners of *Eremias multiocellata-E. przewalskii* species complex (Squamata, Lacertidae). *Zootaxa*, **4282**(1): 1–42.
- Puillandre N, Lambert A, Brouillet S, Achaz G. 2012. ABGD, Automatic Barcode Gap Discovery for primary species delimitation. *Molecular Ecology*, 21(8): 1864–1877.
- Rambaut A, Drummond AJ, Xie D, Baele G, Suchard MA. 2018. Posterior summarization in Bayesian phylogenetics using Tracer 1.7. *Systematic Biology*, 67(5): 901–904.
- Ratnasingham S, Hebert PDN. 2007. BOLD: The Barcode of Life Data System (http://www.barcodinglife.org). *Molecular Ecology Notes*, 7(3): 355–364.
- Ratnasingham S, Hebert PDN. 2013. A DNA-based registry for all animal species: the Barcode Index Number (BIN) system. *PLoS One*, **8**(7): e66213.
- Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Höhna S, et al. 2012.
 MrBayes 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space. *Systematic Biology*, 61(3): 539–542.
- Saitou N, Nei M. 1987. The neighbor-joining method: a new method for reconstructing phylogenetic trees. *Molecular Biology and Evolution*, **4**(4):

406-425.

- Stamatakis A. 2014. RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. *Bioinformatics*, **30**(9): 1312–1313.
- Szczerbak NN. 1974. Yashchurki Palearktiki (*Eremias* Lizards of the Palearctic). Kiev: Naukova Dumka Press. (in Russian)
- Ward RD, Zemlak TS, Innes BH, Last PR, Hebert PDN. 2005. DNA barcoding Australia's fish species. *Philosophical Transactions of the Royal Society B: Biological Sciences*, **360**(1462): 1847–1857.
- Yang ZH, Rannala B. 2010. Bayesian species delimitation using multilocus sequence data. Proceedings of the National Academy of Sciences of the United States of America, 107(20): 9264–9269.
- Yang ZH, Rannala B. 2017. Bayesian species identification under the multispecies coalescent provides significant improvements to DNA barcoding analyses. *Molecular Ecology*, 26(11): 3028–3036.





Supplementary Figure S1 Frequency distribution of pairwise Kimura 2-parameter (K2P) distances (%) at intra- and interspecific taxonomic level in *Eremias multiocellata-przewalskii* species complex

Figure S2



Max Intra-Specific Distance (%)

Supplementary Figure S2 Scatterplot of maximum intraspecific variation and minimum genetic distance to the nearest-neighbor species for each species in *Eremias multiocellata-przewalskii* species complex



Supplementary Figure S3 50% majority-rule consensus tree of *E. multiocellata-przewalskii* species complex resulting from partitioned Bayesian analysis based on barcoding *COI* haplotypes

Numbers beside nodes indicate Bayesian posterior probabilities (BPP). Colored symbols correspond to Figure 1 and those in Orlova et al. (2017), except light green branches and light green circle, which represent samples collected in this study.



Supplementary Figure S4 Maximum-likelihood (ML) tree of *E*.

multiocellata-przewalskii species complex resulting from partitioned ML analysis based on barcoding *COI* haplotypes

Numbers beside nodes indicate bootstrap support proportion (BSP). Colored symbols correspond to Figure 1 and those in Orlova et al. (2017), except light green branches and light green circle, which represent samples collected in this study.

Figure S5



Supplementary Figure S5 Automatic Barcode Gap Discovery (ABGD) species delimitation

A: Frequency histogram of Kimura 2-parameter (K2P) pairwise distances for all barcoding mitochondrial *COI* haplotypes. B: Partitions under different prior intraspecific divergences.

Taxon	Voucher No.	Sex	Population	Haplotype	Locality	Longitude	Latitude	GenBank No.	Reference
				No.					
E. dzungarica	KZL91	Ŷ	56	Hap 10	near the bank of Irtysh river, Bastaushy, Kokpekti	83.42	48.82	MW172548	This study
					District, Kazakhstan				
E. dzungarica	KZL96	Ŷ	56	Hap 10	near the bank of Irtysh river, Bastaushy, Kokpekti	83.42	48.82	MW172549	This study
					District, Kazakhstan				
E. dzungarica	KZL97	8	56	Hap 10	near the bank of Irtysh river, Bastaushy, Kokpekti	83.42	48.82	MW172550	This study
					District, Kazakhstan				
E. dzungarica	KZL98	8	56	Hap 15	near the bank of Irtysh river, Bastaushy, Kokpekti	83.42	48.82	MW172551	This study
					District, Kazakhstan				
E. dzungarica	KZL99	8	56	Hap 10	near the bank of Irtysh river, Bastaushy, Kokpekti	83.42	48.82	MW172552	This study
					District, Kazakhstan				
E. dzungarica	KZL100	Ŷ	56	Hap 10	near the bank of Irtysh river, Bastaushy, Kokpekti	83.42	48.82	MW172542	This study
					District, Kazakhstan				
E. dzungarica	KZL111	Ŷ	57	Hap 11	near the bank of Irtysh river, Zaisan desert,	85.08	47.95	MW172543	This study
					Kazakhstan				
E. dzungarica	KZL113	Ŷ	57	Hap 12	near the bank of Irtysh river, Zaisan desert,	85.08	47.95	MW172544	This study
					Kazakhstan				
E. dzungarica	KZL114	Ŷ	57	Hap 3	near the bank of Irtysh river, Zaisan desert,	85.08	47.95	MW172545	This study
					Kazakhstan				
E. dzungarica	KZL115	Ŷ	57	Hap 13	near the bank of Irtysh river, Zaisan desert,	85.08	47.95	MW172546	This study
					Kazakhstan				
E. dzungarica	KZL116	ð	57	Hap 14	near the bank of Irtysh river, Zaisan desert,	85.08	47.95	MW172547	This study
					Kazakhstan				

Supplementary Table S1 List of analyzed specimens, along with their sex, geographic origin and GenBank accession number

E. dzungarica	Guo8275	Ŷ	58	Hap 3	Western Habahe County, Xinjiang, China	85.69	48.01	MW172538	This study
E. dzungarica	Guo8276	ð	58	Hap 5	Western Habahe County, Xinjiang, China	85.69	48.01	MW172539	This study
E. dzungarica	Guo8278	Ŷ	58	Hap 5	Western Habahe County, Xinjiang, China	85.69	48.01	MW172540	This study
E. dzungarica	Guo8279	ð	58	Hap 9	Western Habahe County, Xinjiang, China	85.69	48.01	MW172541	This study
E. dzungarica	Guo3034	ð	59	Hap 3	Zaisan desert, Jeminay County, Xinjiang, China	85.60	47.71	MW172526	This study
E. dzungarica	Guo3035	ð	59	Hap 3	Zaisan desert, Jeminay County, Xinjiang, China	85.60	47.71	MW172527	This study
E. dzungarica	Guo3036	ð	59	Hap 4	Zaisan desert, Jeminay County, Xinjiang, China	85.60	47.71	MW172528	This study
E. dzungarica	Guo3023	ð	60	Hap 1	Southeast Jeminay County, Xinjiang, China	86.15	47.28	MW172523	This study
E. dzungarica	Guo3024	Ŷ	60	Hap 1	Southeast Jeminay County, Xinjiang, China	86.15	47.28	MW172524	This study
E. dzungarica	Guo3026	ð	60	Hap 2	Southeast Jeminay County, Xinjiang, China	86.15	47.28	MW172525	This study
E. dzungarica	Guo8240	ð	61	Hap 8	Southeast Habahe County, Xinjiang, China	86.58	47.94	MW172534	This study
E. dzungarica	Guo8241	ð	61	Hap 5	Southeast Habahe County, Xinjiang, China	86.58	47.94	MW172535	This study
E. dzungarica	Guo8242	Ŷ	61	Hap 5	Southeast Habahe County, Xinjiang, China	86.58	47.94	MW172536	This study
E. dzungarica	Guo8243	Ŷ	61	Hap 5	Southeast Habahe County, Xinjiang, China	86.58	47.94	MW172537	This study
E. dzungarica	Guo3096	Ŷ	62	Hap 5	Baishashan desert, Habahe County, Xinjiang,	86.90	48.03	MW172529	This study
					China				
E. dzungarica	Guo3097	8	62	Hap 6	Baishashan desert, Habahe County, Xinjiang,	86.90	48.03	MW172530	This study
					China				
E. dzungarica	Guo3098	Ŷ	62	Hap 7	Baishashan desert, Habahe County, Xinjiang,	86.90	48.03	MW172531	This study
					China				
E. dzungarica	Guo3099	ð	62	Hap 5	Baishashan desert, Habahe County, Xinjiang,	86.90	48.03	MW172532	This study
					China				

E. dzungarica	Guo3100	Ŷ	62	Hap 5	Baishashan desert, Habahe County, Xinjiang,	86.90	48.03	MW172533	This study
					China				
E. szczerbaki	ZMMU R-14342-1	\	1	Hap 46	Naryn Prov., Naryn Distr., N from Naryn,	75.98	41.48	KY366618	Orlova et al. (2017)
					Kyrgyzstan				
E. szczerbaki	ZMMU R-14342-2	١	1	Hap 47	Naryn Prov., Naryn Distr., N from Naryn,	75.98	41.48	KY366619	Orlova et al. (2017)
					Kyrgyzstan				
E. stummeri	ZMMU R-14341-2	\	2	Hap 44	Naryn Prov., Kochkor Distr., S from Kochkor,	75.66	42.08	KY366616	Orlova et al. (2017)
					Kyrgyzstan				
E. stummeri	ZMMU R-14341-3	\	2	Hap 45	Naryn Prov., Kochkor Distr., S from Kochkor,	75.66	42.08	KY366617	Orlova et al. (2017)
					Kyrgyzstan				
E. stummeri	ZMMU R-14339-1	\	3	Hap 41	Naryn Prov., Kochkor Distr., Kochkor, Kyrgyzstan	75.75	42.22	KY366615	Orlova et al. (2017)
E. stummeri	ZMMU R-12556-1	١	4	Hap 37	Issyk-Kul Prov., env. of Balykchy, road to Akolen,	76.17	42.35	KY366611	Orlova et al. (2017)
					Kyrgyzstan				
E. stummeri	ZMMU R-12556-2	\	4	Hap 37	Issyk-Kul Prov., env. of Balykchy, road to Akolen,	76.17	42.35	KY366612	Orlova et al. (2017)
					Kyrgyzstan				
E. stummeri	ZMMU R-12557-1	\	4	Hap 43	Issyk-Kul Prov., env. of Balykchy, road to Akolen,	76.17	42.35	KY366613	Orlova et al. (2017)
					Kyrgyzstan				
E. stummeri	ZMMU R-12557-2	\	4	Hap 43	Issyk-Kul Prov., env. of Balykchy, road to Akolen,	76.17	42.35	KY366614	Orlova et al. (2017)
					Kyrgyzstan				
E. stummeri	ZMMU R-12427-1	\	5	Hap 42	Issyk-Kul Prov., NW bank of issyk-Kul lake,	76.41	42.58	KY366609	Orlova et al. (2017)
					20-25 km N from Toru Aygyr, Kungei-Alatau				
					Mts., Kyrgyzstan				
E. stummeri	ZMMU R-12427-2	\	5	Hap 42	Issyk-Kul Prov., NW bank of issyk-Kul lake,	76.41	42.58	KY366610	Orlova et al. (2017)
					20-25 km N from Toru Aygyr, Kungei-Alatau				
					Mts., Kyrgyzstan				
E. stummeri	ZMMU R-14338-1	١	6	Hap 41	Issyk-Kul Prov., 100 km SW from Karakol,	77.18	42.16	KY366607	Orlova et al. (2017)

				Kaji-Say env Kyrgyzstan				
ZMMU R-14338-2	\	6	Hap 41	Issyk-Kul Prov., 100 km SW from Karakol,	77.18	42.16	KY366608	Orlova et al. (2017)
				Kaji-Say env., Kyrgyzstan				
ZMMU R-14335-2	\	7	Hap 39	Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of	78.36	42.46	KY366604	Orlova et al. (2017)
				Karakol, Kyrgyzstan				
ZMMU R-14335-3	\	7	Hap 40	Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of	78.36	42.46	KY366605	Orlova et al. (2017)
				Karakol, Kyrgyzstan				
ZMMU R-14335-4	\	7	Hap 40	Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of	78.36	42.46	KY366606	Orlova et al. (2017)
				Karakol, Kyrgyzstan				
ZMMU R-12551-1a	\	8	Hap 34	Almaty Prov., Rayimbek Distr., Ketmen Mts.	79.32	42.97	KY366589	Orlova et al. (2017)
				Foothills, 7-8 km E from Kegen, Kazakhstan				
ZMMU R-12551-2	\	8	Hap 35	Almaty Prov., Rayimbek Distr., Ketmen Mts.	79.32	42.97	KY366590	Orlova et al. (2017)
				Foothills, 7-8 km E from Kegen, Kazakhstan				
ZMMU R-12551-2a	\	8	Hap 34	Almaty Prov., Rayimbek Distr., Ketmen Mts.	79.32	42.97	KY366591	Orlova et al. (2017)
				Foothills, 7-8 km E from Kegen				
ZMMU R-12551-3	\	8	Hap 36	Almaty Prov., Rayimbek Distr., Ketmen Mts.	79.32	42.97	KY366592	Orlova et al. (2017)
				Foothills, 7-8 km E from Kegen, Kazakhstan				
ZMMU R-12551-4	\	8	Hap 34	Almaty Prov., Rayimbek Distr., Ketmen Mts.	79.32	42.97	KY366593	Orlova et al. (2017)
				Foothills, 7-8 km E from Kegen, Kazakhstan				
ZMMU R-12551-5	\	8	Hap 34	Almaty Prov., Rayimbek Distr., Ketmen Mts.	79.32	42.97	KY366594	Orlova et al. (2017)
			•	Foothills, 7-8 km E from Kegen, Kazakhsan				
ZMMU R-12551-5a	\	8	Hap 34	Almaty Prov., Rayimbek Distr., Ketmen Mts.	79.32	42.97	KY366595	Orlova et al. (2017)
			1	Foothills, 7-8 km E from Kegen, Kazakhstan				× ,
ZMMU R-12551-6	\	8	Hap 34	Almaty Prov. Ravimbek Distr., Ketmen Mts.	79.32	42.97	KY366596	Orlova et al. (2017)
		-	r • ·	Foothills 7-8 km E from Kegen Kazakhsan				
ZMM∐ R_12551_7	\	8	Han 34	Almaty Prov. Ravimbek Distr. Ketmen Mts	79.32	42.97	KV366597	Orlova et al. (2017)
	ZMMU R-14338-2 ZMMU R-14335-2 ZMMU R-14335-3 ZMMU R-14335-4 ZMMU R-14335-4 ZMMU R-12551-1a ZMMU R-12551-2a ZMMU R-12551-3 ZMMU R-12551-4 ZMMU R-12551-5a ZMMU R-12551-5a ZMMU R-12551-6 ZMMU R-12551-7	ZMMU R-14338-2 \ ZMMU R-14335-2 \ ZMMU R-14335-3 \ ZMMU R-14335-3 \ ZMMU R-14335-4 \ ZMMU R-12551-1a \ ZMMU R-12551-2a \ ZMMU R-12551-3a \ ZMMU R-12551-5a \ ZMMU R-12551-5a \ ZMMU R-12551-6a \ ZMMU R-12551-7a \	ZMMU R-14338-2 \ 6 ZMMU R-14335-2 \ 7 ZMMU R-14335-3 \ 7 ZMMU R-14335-4 \ 7 ZMMU R-12551-1a \ 8 ZMMU R-12551-2a \ 8 ZMMU R-12551-3a \ 8 ZMMU R-12551-5a \ 8 ZMMU R-12551-6a \ 8 ZMMU R-12551-75 \ 8	ZMMU R-14338-2 \ 6 Hap 41 ZMMU R-14335-2 \ 7 Hap 39 ZMMU R-14335-3 \ 7 Hap 40 ZMMU R-14335-4 \ 7 Hap 40 ZMMU R-14335-4 \ 7 Hap 40 ZMMU R-12551-1a \ 8 Hap 34 ZMMU R-12551-2a \ 8 Hap 34 ZMMU R-12551-3a \ 8 Hap 34 ZMMU R-12551-5a \ 8 Hap 34 ZMMU R-12551-6 \ 8 Hap 34	ZMMU R-14338-2\6Hap 41Issyk-Kul Prov., 100 km SW from Karakol, Kaji-Say env., KyrgyzstanZMMU R-14335-2\7Hap 39Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of Karakol, KyrgyzstanZMMU R-14335-3\7Hap 40Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of Karakol, KyrgyzstanZMMU R-14335-3\7Hap 40Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of Karakol, KyrgyzstanZMMU R-14335-4\7Hap 40Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of Karakol, KyrgyzstanZMMU R-12551-1a\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts. Foothills, 7-8 km E from Kegen, KazakhstanZMMU R-12551-2\8Hap 35Almaty Prov., Rayimbek Distr., Ketmen Mts. Foothills, 7-8 km E from KegenZMMU R-12551-3\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts. Foothills, 7-8 km E from Kegen, KazakhstanZMMU R-12551-5\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts. Foothills, 7-8 km E from Kegen, KazakhstanZMMU R-12551-5\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts. Foothills, 7-8 km E from Kegen, KazakhstanZMMU R-12551-6\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts. Foothills, 7-8 km E from Kegen, KazakhstanZMMU R-12551-6\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts. Foothills, 7-8 km E from Kegen, KazakhstanZMMU R-12551-6\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts. Foothills, 7-8 km E from Kegen, Kaz	ZMMU R-14338-2\6Hap 41Issyk-Kul Prov., 100 km SW from Karakol, Kaji-Say env., Kyrgyzstan77.18ZMMU R-14335-2\7Hap 39Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of Karakol, Kyrgyzstan78.36ZMMU R-14335-3\7Hap 40Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of Karakol, Kyrgyzstan78.36ZMMU R-14335-4\7Hap 40Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of Karakol, Kyrgyzstan78.36ZMMU R-12551-1a\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts.79.32ZMMU R-12551-2\8Hap 35Almaty Prov., Rayimbek Distr., Ketmen Mts.79.32ZMMU R-12551-3\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts.79.32ZMMU R-12551-3\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts.79.32ZMMU R-12551-5\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts.79.32ZMMU R-12551-5\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts.79.32Foothills, 7-8 km E from Kegen, Kazakhstan79.32Foothills, 7-8 km E from Kegen, Kazakhstan79.32ZMMU R-12551-5\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts.79.32Foothills, 7-8 km E from Kegen, Kazakhstan79.32Foothills, 7-8 km E from Kegen, Kazakhstan79.32ZMMU R-12551-5\8Hap 34Almaty Prov., Rayimbek Distr., Ketmen Mts.79.32Foothills, 7-8 km E from Kegen, Kazakhstan<	ZMMU R:14338-2 \ 6 Hap 41 Issyk-Kul Prov., 100 km SW from Karakol, Kaji-Say env., Kyrgyzstan 71.18 42.16 ZMMU R:14335-2 \ 7 Hap 39 Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of Karakol, Kyrgyzstan 78.36 42.46 ZMMU R:14335-3 \ 7 Hap 40 Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of Karakol, Kyrgyzstan 78.36 42.46 ZMMU R:14335-4 \ 7 Hap 40 Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of Karakol, Kyrgyzstan 78.36 42.46 ZMMU R:12551-13 \ 7 Hap 40 Issyk-Kul Prov., Rayimbek Distr, Ketmen Mts. 79.32 42.97 ZMMU R:12551-2 \ 8 Hap 35 Almaty Prov., Rayimbek Distr, Ketmen Mts. 79.32 42.97 ZMMU R:12551-3 \ 8 Hap 36 Almaty Prov., Rayimbek Distr, Ketmen Mts. 79.32 42.97 ZMMU R:12551-4 \ 8 Hap 36 Almaty Prov., Rayimbek Distr, Ketmen Mts. 79.32 42.97 CMMU R:12551-3 \ 8 Hap 36 Almaty Prov., Rayimbek Distr, Ketmen Mts. 79.32 42.97 CMMU R:12551-4 \ 8 <td< td=""><td>ZMMU R-14338-2 \ 6 Hap 41 Issyk-Kul Prov., 100 km SW from Karakol, Kaji-Say env., Kyrgyzstan 7.18 42.16 KY36608 ZMMU R-14335-2 \ 7 Hap 39 Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of 78.36 42.46 KY366004 Karakol, Kyrgyzstan . 7 Hap 40 Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of 78.36 42.46 KY366005 ZMMU R-14335-4 \ 7 Hap 40 Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of 78.36 42.46 KY366006 ZMMU R-12351-10 \ 7 Hap 40 Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of 78.36 42.46 KY366006 ZMMU R-12551-2 \ 8 Hap 34 Almaty Prov., Rayimbek Distr., Ketmen Mts. 79.32 42.97 KY366590 ZMMU R-12551-2 \ 8 Hap 36 Almaty Prov., Rayimbek Distr., Ketmen Mts. 79.32 42.97 KY366591 ZMMU R-12551-3 \ 8 Hap 36 Almaty Prov., Rayimbek Distr., Ketmen Mts. 79.32 42.97 KY366592 ZMMU R-12551-5 \ 8 Hap 36 Almaty Prov., Rayimbek Distr., Ketmen</td></td<>	ZMMU R-14338-2 \ 6 Hap 41 Issyk-Kul Prov., 100 km SW from Karakol, Kaji-Say env., Kyrgyzstan 7.18 42.16 KY36608 ZMMU R-14335-2 \ 7 Hap 39 Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of 78.36 42.46 KY366004 Karakol, Kyrgyzstan . 7 Hap 40 Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of 78.36 42.46 KY366005 ZMMU R-14335-4 \ 7 Hap 40 Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of 78.36 42.46 KY366006 ZMMU R-12351-10 \ 7 Hap 40 Issyk-Kul Prov., E bank of issyk-Kul lake, env. Of 78.36 42.46 KY366006 ZMMU R-12551-2 \ 8 Hap 34 Almaty Prov., Rayimbek Distr., Ketmen Mts. 79.32 42.97 KY366590 ZMMU R-12551-2 \ 8 Hap 36 Almaty Prov., Rayimbek Distr., Ketmen Mts. 79.32 42.97 KY366591 ZMMU R-12551-3 \ 8 Hap 36 Almaty Prov., Rayimbek Distr., Ketmen Mts. 79.32 42.97 KY366592 ZMMU R-12551-5 \ 8 Hap 36 Almaty Prov., Rayimbek Distr., Ketmen

					Foothills, 7-8 km E from Kegen, Kazakhsan				
E. stummeri	ZMMU R-12551-8	١	8	Hap 34	Almaty Prov., Rayimbek Distr., Ketmen Mts.	79.32	42.97	KY366598	Orlova et al. (2017)
					Foothills, 7-8 km E from Kegen, Kazakhsan				
E. stummeri	ZMMU R-12551-10	١	8	Hap 34	Almaty Prov., Rayimbek Distr., Ketmen Mts.	79.32	42.97	KY366599	Orlova et al. (2017)
					Foothills, 7-8 km E from Kegen, Kazakhsan				
E. stummeri	ZMMU R-1 2495-1	\	8	Hap 37	Almaty Prov., Rayimbek Distr., Ketmen Mts.	79.32	42.97	KY366600	Orlova et al. (2017)
					Foothills, 7-8 km E from Kegen, Kazakhsan				
E. stummeri	ZMMU R-12552-1	١	9	Hap 38	Almaty Prov., Rayimbek Distr., central Tian Shan	80.08	42.92	KY366601	Orlova et al. (2017)
					Mts., 15 km S from Tuzkol Lake, Zhabyrtau Mt.,				
					Kazakhstan				
E. stummeri	ZMMU R-12552-2	\	9	Hap 38	Almaty Prov., Rayimbek Distr., central Tian Shan	80.08	42.92	KY366602	Orlova et al. (2017)
					Mts., 15 km S from Tuzkol Lake, Zhabyrtau Mt.,				
					Kazakhstan				
E. stummeri	ZMMU R-12552-4	١	9	Hap 38	Almaty Prov., Rayimbek Distr., central Tian Shan	80.08	42.92	KY366603	Orlova et al. (2017)
					Mts., 15 km S from Tuzkol Lake, Zhabyrtau Mt.,				
					Kazakhstan				
Eremias sp. 1	ZMMU R-14329-1	\	10	Hap 50	Xinjiang Prov., 60 km NE rrom Aksu, China	80.79	41.54	KY366626	Orlova et al. (2017)
Eremias sp. 1	ZMMU R-14330-1	\	11	Hap 51	Xinjiang Prov., 75 km NE from Aksu, China	80.83	41.74	KY366627	Orlova et al. (2017)
Eremias sp. 1	ZMMU R-14330-2	١	11	Hap 52	Xinjiang Prov., 75 km NE from Aksu, China	80.83	41.74	KY366628	Orlova et al. (2017)
Eremias sp. 1	ZMMU R-14330-3	١	11	Hap 53	Xinjiang Prov., 75 km NE from Aksu, China	80.83	41.74	KY366629	Orlova et al. (2017)
Eremias sp. 1	ZMMU R-14327-1	\	12	Hap 49	Xinjiang Prov., 35 km NE rrom Aksu, China	81.05	41.40	KY366625	Orlova et al. (2017)
Eremias sp. 1	ZMMU R-14328-1	\	13	Hap 54	Xinjiang Prov., 89 km NE from Aksu, China	81.21	41.56	KY366630	Orlova et al. (2017)
Eremias sp. 1	ZMMU R-14328-2	\	13	Hap 55	Xinjiang Prov., 89 km NE from Aksu, China	81.21	41.56	KY366631	Orlova et al. (2017)
Eremias sp. 1	ZMMU R-14328-3	١	13	Hap 56	Xinjiang Prov., 89 km NE from Aksu, China	81.21	41.56	KY366632	Orlova et al. (2017)

E. yarkandensis	ZMMU R-14344-1	\	14	Hap 48	Osh Prov., vicinity of Nura, Kyrgyzstan	73.87	39.65	KY366620	Orlova et al. (2017)
E. yarkandensis	ZMMU R-14344-2	١	14	Hap 48	Osh Prov., vicinity of Nura, Kyrgyzstan	73.87	39.65	KY366621	Orlova et al. (2017)
E. yarkandensis	ZMMU R-14344-3	١	14	Hap 48	Osh Prov., vicinity of Nura, Kyrgyzstan	73.87	39.65	KY366622	Orlova et al. (2017)
E. yarkandensis	ZMMU R-14344-4	١	14	Hap 48	Osh Prov., vicinity of Nura, Kyrgyzstan	73.87	39.65	KY366623	Orlova et al. (2017)
E. yarkandensis	ZMMU R-14344-5	١	14	Hap 48	Osh Prov., vicinity of Nura, Kyrgyzstan	73.87	39.65	KY366624	Orlova et al. (2017)
E. cf. buechneri	ZMMU R-8910-1a	١	15	Hap 28	Xinjiang Prov., Qarqan (Chemo) Distr., Altintag	86.05	37.51	KY366574	Orlova et al. (2017)
					Mt., Chinbulak, 60 km S from Turav, China				
E. cf. buechneri	ZMMU R-8910-1b	\	15	Hap 28	Xinjiang Prov., Qarqan (Chemo) Distr., Altintag	86.05	37.51	KY366575	Orlova et al. (2017)
					Mt., Chinbulak, 60 km S from Turav, China				
Eremias sp. 2	١	\	16	Hap 26	Qinhai Prov., Delingha env., China	97.35	37.35	KY366572	Orlova et al. (2017)
Eremias sp. 2	١	\	17	Hap 27	Qinhai Prov., Gonghe (Qiabugia), Chekhou, env.	100.75	36.20	KY366573	Orlova et al. (2017)
					Of Kyiking (Chin-Kon), China				
E. cf. multiocellata	١	١	18	Hap 65	Gansu Prov., env. of Shandan, China			KY366641	Orlova et al. (2017)
E. przewalskii	ZMMU R-13209	١	19	Hap 17	Inner Mongolia Prov., 30 km W from Bayanhot	101.27	39.12	KY366551	Orlova et al. (2017)
E. przewalskii	\	\	20	Hap 16	Gansu Prov., Minqin County, Wuwei, China	103.11	38.63	KM507330	Orlova et al. (2017)
E. przewalskii	ZMMU R-13214	١	21	Hap 18	Inner Mongolia Prov., Alashan desert, 150 km W	105.59	40.41	KY366552	Orlova et al. (2017)
					from Bayanmod, China				
E. przewalskii	ZMMU R-12046-1	\	22	Hap 19	Bayankhongor Aimaq, Zhinst, Mongolia	100.21	45.62	KY366553	Orlova et al. (2017)
E. przewalskii	ZMMU R-13038-1	\	23	Hap 19	Govi-Altai Aimaq, Biger valley, Mongolia	97.4	45.73	KY366554	Orlova et al. (2017)
E. przewalskii	ZMMU R-13038-3	\	23	Hap 19	Govi-Altai Aimaq, Biger valley, Mongolia	97.4	45.73	KY366555	Orlova et al. (2017)
E. przewalskii	ZMMU R-13038-4	١	23	Hap 19	Govi-Altai Aimaq, Biger valley, Mongolia	97.4	45.73	KY366556	Orlova et al. (2017)
E. przewalskii	ZMMU R-12842-1	١	24	Hap 19	Govi-Aitai Aimaq, 3 km W from Biger Sum,	97.15	45.72	KY366557	Orlova et al. (2017)
					Mongolia				

E. przewalskii	ZMMU R-12842-2	١	24	Hap 19	Govi-Aitai Aimaq, 3 km W from Biger Sum,	97.15	45.72	KY366558	Orlova et al. (2017)
					Mongolia				
E. przewalskii	ZMMU R-12888-1	\	25	Hap 20	Govi-Aitai Aimaq, Tsogt, Dzahuin-Govi, 10 km W	96.65	45.67	KY366559	Orlova et al. (2017)
					Bayan-Toroo, Mongolia				
E. przewalskii	ZMMU R-12888-2	١	25	Hap 20	Govi-Aitai Aimaq, Tsogt, Dzahuin-Govi, 10 km W	96.65	45.67	KY366560	Orlova et al. (2017)
					Bayan-Toroo, Mongolia				
E. przewalskii	ZMMU R-13060-1	\	26	Hap 19	Tuva Republic, Ovyurskiy Distr., Tere-Khollake,	93.21	50.68	KY366561	Orlova et al. (2017)
					Tsuger-Els sands, Russia				
E. przewalskii	ZMMU R-13060-2	\	26	Hap 19	Tuva Republic, Ovyurskiy Distr., Tere-Khollake,	93.21	50.68	KY366562	Orlova et al. (2017)
					Tsuger-Els sands, Russia				
E. przewalskii	ZMMU R-13060-3	\	26	Hap 19	Tuva Republic, Ovyurskiy Distr., Tere-Khollake,	93.21	50.68	KY366563	Orlova et al. (2017)
					Tsuger-Els sands, Russia				
E. przewalskii	ENS-T07-1	\	26	Hap 19	Tuva Republic, Ovyurskiy Distr., Tere-Khollake,	93.21	50.68	KY366564	Orlova et al. (2017)
					Tsuger-Els sands, Russia				
E. przewalskii	ENS-T07-2	\	26	Hap 19	Tuva Republic, Ovyurskiy Distr., Tere-Khollake,	93.21	50.68	KY366565	Orlova et al. (2017)
					Tsuger-Els sands, Russia				
E. cf. reticulata	ZMMU R-12855-1	١	27	Hap 22	Gobi-Altai Aimaq, 4 km NW from Altai Sum, env.	95.86	44.64	KY366567	Orlova et al. (2017)
					Danshig-Khuduk, Mongolia				
E. cf. reticulata	ZMMU R-12855-2	١	27	Hap 23	Gobi-Altai Aimaq, 4 km NW from Altai Sum, env.	95.86	44.64	KY366568	Orlova et al. (2017)
					Danshig-Khuduk, Mongolia				
E. cf. reticulata	ZMMU R-12856-1	١	27	Hap 23	Gobi-Altai Aimaq, 4 km NW from Altai Sum, env.	95.86	44.64	KY366569	Orlova et al. (2017)
					Danshig-Khuduk, Mongolia				
E. cf. reticulata	MNUE-Em-S	\	28	Hap 21	Bayankhongor Aimaq, Tsagan Bogdo Uul (Mt.)	98.82	42.88	KY366566	Orlova et al. (2017)
E. cf. przewalskii	ZMMU R-12858-1	١	29	Hap 24	ömnögovi Aimaq, 8-10 km S from Bayandalai	103.45	43.39	KY366570	Orlova et al. (2017)
*					Sum, Mongolia				
	ZMMI R-12859-1	\	29	Han 25	ömnögovi Aimaa 8-10 km S from Bayandalai	103 45	43 39	KV366571	Orlova et al. (2017)

					Sum, Mongolia				
Eremias sp. 3	١	١	30	Hap 64	Gansu Prov., env. of Lanzhou, Baiyin, China	104.21	36.46	KJ664798	Li and Song
									(unpublished data)
Eremias sp. 3	λ	١	31	Hap 63	Gansu Prov., Minqin County, China	103.11	38.63	NC025304	Tong et al. (2014)
Eremias sp. 3	ZMMU R-13208	١	32	Hap 62	Inner Mongolia Prov., 80 km SW Sayan Nur,	103.87	40.00	KY366640	Orlova et al. (2017)
					China				
Eremias sp. 3	ZMMU R-13207	١	33	Hap 60	Inner Mongolia Prov., 120 km W from Wuhai,	105.45	39.53	KY366638	Orlova et al. (2017)
					China				
Eremias sp. 3	ZMMUR-13206	١	34	Hap 61	Inner Mongolia Prov., Alashan desert, 40 km S	105.62	38.67	KY366639	Orlova et al. (2017)
					from Bayan-Hot, China				
Eremias sp. 3	ZMMU R-13205	١	35	Hap 59	Inner Mongolia Prov., W Ordos, 30 km SW from	108.50	40.02	KY366637	Orlova et al. (2017)
					Bayan-Us, China				
Eremias sp. 3	ZMMU R-13215	\	36	Hap 58	Inner Mongolia Prov., 50 km S from Baotou,	110.00	40.28	KY366636	Orlova et al. (2017)
					China				
Eremias sp. 3	ZMMU R-12610-1a	١	37	Hap 57	Dornogovi Aimaq, Sainshand Sum, vicinity of	110.10	44.89	KY366633	Orlova et al. (2017)
					Saishand, Mongolia				
Eremias sp. 3	ZMMU R-12610-1b	\	37	Hap 57	Dornogovi Aimaq, Sainshand Sum, vicinity of	110.10	44.89	KY366634	Orlova et al. (2017)
					Saishand, Mongolia				
Eremias sp. 3	ZMMU R-14523	\	38	Hap 57	Dornogovi Aimaq, Sainshand Sum, ca. 100 km NE	111.79	45.18	KY366635	Orlova et al. (2017)
					from örgön, Mongolia				
E. multiocellata	ZMMU R-12863-1	\	39	Hap 72	Umnugovi Aimaq, 102 km NW from	105.32	44.42	KY366651	Orlova et al. (2017)
					Dalandzadagat, Tsogt-Ovu, Mongolia				
E. multiocellata	ZMMU R-12863-2	/	39	Hap 73	Umnugovi Aimaq, 102 km NW from	105.32	44.42	KY366652	Orlova et al. (2017)
	7) 0 (U.D. 10072	,	40		Dalandzadagat, Tsogt-Ovu, Mongolia	106.54	46.50	1131266647	
E. multiocellata	ZMMU R-12952a	١	40	Hap 70	Dundgovi Aimaq, env. of Deren, Mongolia	106.54	46.59	KY366647	Orlova et al. (2017)

E. multiocellata	ZMMU R-12952b	\	40	Hap 70	Dundgovi Aimaq, env. of Deren, Mongolia	106.54	46.59	KY366648	Orlova et al. (2017)
E. multiocellata	ZMMU R-12608-1	١	41	Hap 71	Dundgovi Aimaq, env. of ölziit Sum, Mongolia	102.51	48.03	KY366649	Orlova et al. (2017)
E. multiocellata	ZMMU R-12608-2	١	41	Hap 71	Dundgovi Aimaq, env. of ölziit Sum, Mongolia	102.51	48.03	KY366650	Orlova et al. (2017)
E. multiocellata	ZMMU R-12045-1	١	42	Hap 74	Bayankhongor Aimaq, Zhinst, Mongolia	100.21	45.62	KY366653	Orlova et al. (2017)
E. multiocellata	ZMMU R-12045-2	١	42	Hap 74	Bayankhongor Aimaq, Zhinst, Mongolia	100.21	45.62	KY366654	Orlova et al. (2017)
E. multiocellata	ZMMU R-12841-1	١	43	Hap 74	Bayankhongor Aimaq, N bank of Bööntsagan	99.12	45.67	KY366655	Orlova et al. (2017)
					Nuur Lake, Mongolia				
E. multiocellata	ZMMU R-12841-2	\	43	Hap 74	Bayankhongor Aimaq, N bank of Bööntsagan	99.12	45.67	KY366656	Orlova et al. (2017)
					Nuur Lake, Mongolia				
E. multiocellata	ZMMU R-13039-1	\	44	Hap 74	Govi-Altai Aimaq, Delger, Mongolia	98.05	45.93	KY366657	Orlova et al. (2017)
E. multiocellata	ZMMU R-12843-1	١	45	Hap 75	Govi-Altai Aimaq, Shargyn-Govi, 2 km SW from	96.12	45.93	KY366658	Orlova et al. (2017)
					Khalium Sum, Mongolia				
E. multiocellata	ZMMU R-12843-2	١	45	Hap 75	Govi-Altai Aimaq, Shargyn-Govi, 2 km SW from	96.12	45.93	KY366659	Orlova et al. (2017)
					Khalium Sum, Mongolia				
E. multiocellata	ZMMU R-12081-1	١	46	Hap 76	Govi-Aitai Aimaq, Shargyn-Govi, 30 km S from	95.23	46.36	KY366660	Orlova et al. (2017)
					Sharga Sum, Mongolia				
E. multiocellata	ZMMU R-13058-1	١	47	Hap 66	Tuva Republic, Erzin Distr., Ubsu-Nw valley, 12	94.79	50.16	KY366642	Orlova et al. (2017)
					km SE from Yamalyg Mt. , Russia				
E. multiocellata	ZMMU R-13058-2	١	47	Hap 67	Tuva Republic, Erzin Distr., Ubsu-Nw valley, 12	94.79	50.16	KY366643	Orlova et al. (2017)
					km SE from Yamalyg Mt., Russia				
E. multiocellata	ZMMU R-13058-3	١	47	Hap 68	Tuva Republic, Erzin Distr., Ubsu-Nw valley, 12	94.79	50.16	KY366644	Orlova et al. (2017)
					km SE from Yamalyg Mt., Russia				
E. multiocellata	ZMMU R-13058-4	١	47	Hap 69	Tuva Republic, Erzin Distr., Ubsu-Nw valley, 12	94.79	50.16	KY366645	Orlova et al. (2017)
					km SE from Yamalyg Mt., Russia				

E. multiocellata	ZMMU R-13059-1	\	47	Hap 67	Tuva Republic, Erzin Distr., Ubsu-Nw valley, 12	94.79	50.16	KY366646	Orlova et al. (2017)
					km SE from Yamalyg Mt., Russia				
E. multiocellata	ZMMU R-12844-1	\	48	Hap 74	Govi-Aitai Aimaq, Junggar Govi, 4 km W from	93.58	45.59	KY366661	Orlova et al. (2017)
					Bidzh, Mongolia				
E. multiocellata	ZMMU R-12844-2	١	48	Hap 74	Govi-Aitai Aimaq, Junggar Govi, 4 km W from	93.58	45.59	KY366662	Orlova et al. (2017)
					Bidzh, Mongolia				
E. multiocellata	ZMMU R-12853-1	\	49	Hap 74	Govi-Aitai Aimaq, Junggar Govi, Bugat, Mongolia	93.53	45.22	KY366663	Orlova et al. (2017)
E. multiocellata	ZMMU R-12853-2	١	49	Hap 74	Govi-Aitai Aimaq, Junggar Govi, Bugat, Mongolia	93.53	45.22	KY366664	Orlova et al. (2017)
E. multiocellata	ZMMU R-12848-1	١	50	Hap 74	Khovd Aimaq, 58 km SE from Altai Sum, E	92.98	45.65	KY366665	Orlova et al. (2017)
					foothills of Sertengiyn-Khuvch-Ula, Mongolia				
E. multiocellata	ZMMU R-12849-1	١	50	Hap 77	Khovd Aimaq, 58 km SE from Altai Sum, E	92.98	45.65	KY366666	Orlova et al. (2017)
					foothills of Sertengiyn-Khuvch-Ula, Mongolia				
E. multiocellata	ZMMU R-12849-2	١	50	Hap 74	Khovd Aimaq, 58 km SE from Altai Sum, E	92.98	45.65	KY366667	Orlova et al. (2017)
					foothills of Sertengiyn-Khuvch-Ula, Mongolia				
E. multiocellata	ZMMU R-12847-1	\	51	Hap 75	Khovd Aimaq, I km SW from Altai Sum,	92.25	45.80	KY366668	Orlova et al. (2017)
					Mongolia				
E. multiocellata	ZMMU R-13132-1	١	52	Hap 74	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366669	Orlova et al. (2017)
					Mongolia				
E. multiocellata	ZMMU R-13132-2	١	52	Hap 74	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366670	Orlova et al. (2017)
					Mongolia				
E. multiocellata	ZMMU R-13132-3	\	52	Hap 74	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366671	Orlova et al. (2017)
					Mongolia				
E. multiocellata	ZMMU R-13132-4	\	52	Hap 74	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366672	Orlova et al. (2017)
					Mongolia				
E. multiocellata	ZMMU R-13132-5	\	52	Hap 74	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366673	Orlova et al. (2017)
					Mongolia				

E. dzungarica	ZMMU R-12845-2	\	52	Hap 31	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366580	Orlova et al. (2017)
					Mongolia				
E. dzungarica	ZMMU R-12845-4	\	52	Hap 31	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366581	Orlova et al. (2017)
					Mongolia				
E. dzungarica	ZMMU R-12845-5	\	52	Hap 32	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366582	Orlova et al. (2017)
					Mongolia				
E. dzungarica	ZMMU R-12845-6	\	52	Hap 31	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366583	Orlova et al. (2017)
					Mongolia				
E. dzungarica	ZMMU R-12845-7	\	52	Hap 31	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366584	Orlova et al. (2017)
					Mongolia				
E. dzungarica	ZMMU R-12845-8	\	52	Hap 31	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366585	Orlova et al. (2017)
					Mongolia				
E. dzungarica	ZMMU R-12845-9	\	52	Hap 31	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366586	Orlova et al. (2017)
					Mongolia				
E. dzungarica	ZMMU R-12845-10	\	52	Hap 31	Khovd Aimaq, 7 km SW from Uyench Sum,	93.61	46.93	KY366587	Orlova et al. (2017)
					Mongolia				
E. dzungarica	ZMMU R-12862-1	\	53	Hap 31	Khovd Aimaq, Bulgan Sum, Bayan-Mod, 11 km	92.91	47.06	KY366578	Orlova et al. (2017)
					W from Ikher-Toli, Mongolia				
E. dzungarica	ZMMU R-12862-3	\	53	Hap 31	Khovd Aimaq, Bulgan Sum, Bayan-Mod, 11 km	92.91	47.06	KY366579	Orlova et al. (2017)
					W from Ikher-Toli, Mongolia				
E. dzungarica	ZMMU R-12550-1	\	54	Hap 33	Khovd Aimaq, 24 km N from Uyench Sum,	92.05	46.27	KY366588	Orlova et al. (2017)
					Mongolia				
E. dzungarica	ZMMU R-11989-1	\	55	Hap 29	East-Kazakhstan Prov., Aigyrkum sands, 5-7 km	84.89	47.98	KY366576	Orlova et al. (2017)
					SW from Buran				
E. dzungarica	ZMMU R-11989-2	\	55	Hap 30	East-Kazakhstan Prov., Aigyrkum sands, 5-7 km	84.89	47.98	KY366577	Orlova et al. (2017)
					SW from Buran				

E. argus barbouri	ZMMU R-12605-1	\	\	\	\	\	\	KY366548	Orlova et al. (2017)
E. brenchleyi	/	١	١	١	Qianshan, Suzhou, Anhui Province, China	\	\	EF490071	Rui et al. (2009)
E. nikolskii	ZMMU R-11673-1	١	١	١	/		\	KY366550	Orlova et al. (2017)
E. vermiculata	ZMMU-R12047-1	١	\	١	/	\	\	KY366549	Orlova et al. (2017)

References

Orlova VF, Poyarkov Jr NA, Chirikova MA, Nazarov RA, Munkhbaatar M, Munkhbayar K, Terbish K. 2017. MtDNA differentiation and taxonomy of Central Asian racerunners of Eremias multiocellata-E. przewalskii

species complex (Squamata, Lacertidae). Zootaxa, 4282(1): 1-42.

Rui J, Wang Y, Nie L. 2009. The complete mitochondrial DNA genome of Eremias brenchleyi (Reptilia: Lacertidae) and its phylogeny position within squamata reptiles. Amphibia-Reptilia, 30: 25-35

Supplementary Table S2 Kimura 2-parameter (K2P) distance summary for sequence divergence between barcoding mitochondrial *COI* sequences at species and genus level

	Individuals	Taxa	Comparisons	Min%	Mean%	Max%	SE%
Within species	155	12	2018	0.00	1.16	6.18	0.00
Within genus	156	1	10072	0.77	9.09	13.19	0.00

Supplementary Table S3 Kimura 2-parameter (K2P) distance summary of maximum intraspecific variation and minimum genetic distance to the nearest-neighbor species calculated from barcoding mitochondrial *COI* sequences for each species in *Eremias multiocellata-przewalskii* species complex

Species	Max Intra-Sp (%)	Nearest Species	Distance to NN (%)
Eremias sp. 2	0.31	Eremias. cf. reticulata	0.77
Eremias. cf. reticulata	0.31	Eremias sp. 2	0.77
Eremias przewalskii	2.19	Eremias. cf. przewalskii	1.08
Eremias. cf. przewalskii	0.31	Eremias przewalskii	1.08
Eremias cf. buechneri	0	Eremias yarkandensis	1.48
Eremias yarkandensis	0	Eremias cf. buechneri	1.48
Eremias. dzungarica	2.99	Eremias cf. buechneri	1.87
Eremias sp. 1	2.14	Eremias yarkandensis	3.35
Eremias multiocellata	1.71	Eremias sp. 3	5.65
Eremias sp. 3	6.18	Eremias multiocellata	5.65
Eremias cf. multiocellata	N/A	Eremias multiocellata	6.85
Eremias stummeri	2.47	Eremias szczerbaki	7.58
Eremias szczerbaki	0.33	Eremias stummeri	7.58

Supplementary Table S4 Quantitative characteristics of specimens of *Eremias*

dzungarica from this study and Orlova et al. (2017)

Values are given as ranges and means \pm SE. Morphological characters detected with significant sexual dimorphism are given separately for females and males.

Tab	le S	54.1
-----	------	------

Metric Females of <i>E. dzungarica</i> in this			Females of <i>E. dzungarica</i> from Orlova et al.			
and study			(2017)			
meristic	Ν	Range	$Mean \pm SE$	Ν	Range	$Mean \pm SE$
traits						
SVL	15	46.55-64.46	57.79 ± 1.443	2	61.40-64.50	\
TL	7	63.65-76.19	69.96 ± 1.794	0	\setminus	\
Ga	15	21.48-33.07	28.72 ± 0.941	2	33.80-35.00	\
HL	15	11.59-15.40	13.64 ± 0.279	2	14.00-15.00	\
HW	15	6.99-9.89	8.66 ± 0.196	2	7.00-8.20	\
HH	15	5.47-8.07	6.96 ± 0.215	2	6.00-7.20	\
NL	15	15.98-20.96	18.40 ± 0.353	2	19.00-20.50	\
Pa	15	16.14-20.48	18.68 ± 0.346	2	18.00-19.00	\
Рр	15	23.95-29.66	26.98 ± 0.459	2	25.40-26.20	\
Dist.P.f						
m	15	3.52-5.42	4.73 ± 0.117	2	1.60-1.70	\
Sq.c.cd	15	24-27	25.13 ± 0.274	2	23-23	
Lpil.	15	10.74-14.15	12.34 ± 0.236	0	\	\

Table S4.2

Metric	Males of <i>E. dzungarica</i> in this study			Males of <i>E. dzungarica</i> from Orlova et al. (2017)			
and	N	Range	Mean \pm SE	N	Range	Mean \pm SE	
meristic							
traits							
SVL	15	56.14-75.72	63.44 ± 1.378	5	52.00-65.00	58.72 ± 2.506	
TL	11	72.74-94.82	80.79 ± 2.121	2	67.30-85.00	\	
Ga	15	25.23-35.61	31.08 ± 0.853	5	24.00-32.00	28.240 ± 1.574	
HL	15	15.10-18.98	16.45 ± 0.343	5	13.60-18.00	15.88 ± 0.779	
HW	15	8.82-12.20	10.50 ± 0.232	5	7.30-10.30	9.00 ± 0.640	
HH	15	7.16-10.72	8.84 ± 0.272	5	5.00-8.30	6.86 ± 0.616	
NL	15	19.53-26.14	21.93 ± 0.473	5	12.50-21.40	18.96 ± 1.688	
Pa	15	18.14-24.33	20.81 ± 0.414	5	17.50-20.00	19.20 ± 0.464	
Рр	15	26.64-35.41	31.16 ± 0.654	5	25.70-30.00	28.48 ± 0.813	
Dist.P.fm	15	3.52-6.34	4.68 ± 0.219	5	1.70-2.50	2.06 ± 0.129	
Sq.c.cd	15	25-29	26.67 ± 0.319	5	24-27	25.20 ± 0.583	
Lpil.	15	13.26-16.48	14.35 ± 0.270	0	\	\	

Meristic	All s	All specimens of E. dzungarica in this			All specimens of E. dzungarica from		
traits	study			Orlova et al. (2017)			
	Ν	Range	Mean \pm SE	N	Range	Mean \pm SE	
Lab.total.L	30	9-12	10.83 ± 0.145	11	9–11	10.36 ± 0.203	
Lab.total.R	30	9-12	10.60 ± 0.141	11	9–11	10.00 ± 0.234	
Infralab.L	30	6-7	6.63 ± 0.089	12	6-8	$\boldsymbol{6.83 \pm 0.167}$	
Infralab.R	30	6-8	6.77 ± 0.092	12	6v9	6.92 ± 0.260	
Sq.	30	44–55	47.67 ± 0.456	12	46-52	48.42 ± 0.609	
Ventr.	30	30-34	31.93 ± 0.185	12	30v32	30.92 ± 0.193	
G.	30	20-26	22.93 ± 0.335	12	21–26	22.83 ± 0.474	
Coll.	30	9–14	11.27 ± 0.267	11	7–13	11.00 ± 0.632	
Lam.subdig.	30	22-25	23.53 ± 0.224	12	19–24	21.58 ± 0.379	
P.fm.L	30	10-13	11.17 ± 0.167	12	9–15	12.17 ± 0.405	
P.fm.R	30	9–14	11.33 ± 0.188	12	9–14	12.00 ± 0.369	
Ventr.long.	30	14-18	16.30 ± 0.199	1	16	\	
supracil.L	30	3-8	5.43 ± 0.196	0	\	\	
supracil.R	30	4–7	5.47 ± 0.124	0	\setminus	\	
scal.f.p	30	7-12	9.70 ± 0.204	0	\	\	

Table S4.3

Supplementary Table S5 Morphological characters used for morphometric

Abbreviations	Description of morphological trait
SVL	Snout to vent length
TL	Tail length
Ga	Gleno-acetobular distance (from axilla to groin)
HL	Head length
HW	Head width
HH	Head height
Lpil.	Pileus Length (from rostrum to the posterior border of parietals)
NL	head length from snout tip to the anterior edge of collar (measured from ventral side)
Pa	Forelimb Length
Рр	Hindlimb Length
Dist.P.fm.	Distance between femoral pore rows
Lab.total.L	Number of supralabials on the left side
Lab.total.R	Number of supralabials on the right side
Infralab.L	Number of infralabials on the left side
Infralab.R	Number of infralabials on the right side
Sq.	Number of scales around middorsal
Ventr.	Number of transverse rows of ventral scales
G.	Number of gular scales along mid-line of throat
Coll.	Number of collar scales
Lam.subdig.	Number of subdigital lamellae on the 4th toe of hindlimb
P.fm.L	Number of femoral pores on the left side
P.fm.R	Number of femoral pores on the right side
Ventr.long.	Maximal number of longitudinal rows of ventral scales
Supracil.L	Number of supraciliary scales on the left side
Supracil.R	Number of supraciliary scales on the right side
Scal.f.p	Number of scales between femoral pore rows
Sq.c.cd.	Number of scales around the 9–10th tail ring

analysis and their corresponding abbreviations