

Review

Evaluating taxonomic inflation: towards evidence-based species delimitation in Eurasian vipers (Serpentes: Viperinae)

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Abstract. The designation of taxonomic units has important implications for the understanding and conservation of biodiversity. Eurasian vipers are a monophyletic group of viperid snakes (Serpentes, Viperinae), currently comprising four genera (*Daboia*, *Macrovipera*, *Montivipera* and *Vipera*) and up to 40 species. Taxonomic units have been described using a wide variety of methods and criteria, and consequently, considerable controversy still surrounds the validity of some currently listed species. In order to promote a consensus- and evidence-based taxonomy of Eurasian vipers, we analysed published mitochondrial and nuclear DNA sequences for this group to reconstruct phylogenetic relationships among currently recognized viper species. We also compiled information on external morphology to assess their morphological distinctiveness. Phylogenetic inference based on mtDNA sequences shows contrasting levels of divergence across genera and species and identifies several instances of non-monophyly in described species. Nuclear DNA sequences show extremely low levels of genetic variation, with a widespread pattern of allele sharing among distant species, and even among genera. Revision of morphological data shows that most species designations rely on scalation traits that overlap extensively among species of the same genus. Based on our combined assessment, we recognize 15 taxa as valid species, three taxa which likely represent species complexes, 17 taxa of doubtful validity as species, and five taxa for which species status is maintained but further research is highly recommended to assess taxonomic arrangements. We stress the need to implement integrative taxonomic approaches for the recognition of evidence-based taxonomic units in Eurasian vipers.

Keywords: integrative taxonomy, morphology, mt-DNA, nuclear DNA, phylogeny, Viperidae.

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Introduction

The designation of taxonomic units has important implications for the way we study, describe and understand biodiversity, as well as for how we mobilize efforts and allocate resources to develop conservation strategies. Over the years, different criteria and tools have been used to define species, leading to a succession of species concepts that resulted in extended controversy within the research community (Mayden, 1997; de Queiroz, 2007). Nowadays, a species is often defined as a separately evolving metapopulation lineage that possesses relevant characteristics that allow assessing its distinctiveness from others (i.e., the unified species concept; de Queiroz, 2007, and its precursor, the evolutionary species concept; Simpson, 1961; Wiley, 1978; Frost and Hillis, 1990). This definition is linked to the integrative taxonomy framework, which is based on the combination of different lines of evidence (e.g., genetic, morphological, ecological) and methodologies (e.g., phylogenetic inference, ordination methods, ecological modelling) to objectively identify taxa (Dayrat, 2005) that – in an ideal case – would represent independently evolving species. Names of

species are therefore intended to identify biologically cohesive populations with recent common ancestry rather than to recognize unusual patterns of distribution or morphology (Kaiser et al., 2013).

Eurasian vipers are a monophyletic group within the subfamily Viperinae (Serpentes, Viperidae), whose members are distributed primarily in the Palaearctic region, i.e., non-tropical Eurasia and North Africa (Phelps, 2010). This group is phylogenetically sister to a clade of Middle Eastern vipers, constituted by the genera *Eristicophis* and *Pseudocerastes* (see Phelps, 2010; Zheng and Wiens, 2016), which are not considered in this work. At the time of writing, the most recent and comprehensive list of reptiles (i.e., The Reptile Database; Uetz, Freed and Hošek, 2019) lists four genera and 40 species within Eurasian vipers (table 1): *Daboia*, with 4 species; *Macrovipera*, with 3 species; *Montivipera*, with 8 species; and *Vipera*, with 25 species. However, Eurasian vipers have a long taxonomic history, and different authors have used a wide variety of methods and criteria to define taxonomic units, as reflected in previous species lists (e.g., Mallow, Ludwig and Nilson, 2003; Phelps, 2010).

At the genus level, the history of Eurasian vipers is relatively simple. Through most of the 20th century, all species considered here were included in the single genus *Vipera* (Boulenger, 1896, 1913; Schwarz, 1936; Klemmer, 1963; Minton, Dowling and Russel, 1968). The maverick German herpetologist Albert Franz Theodor Reuss described numerous genera within the Eurasian vipers (reviewed by Krecsák, 2007), but these gained little traction with subsequent authors, except where the names had priority for subsequently validated clades. Obst (1983) was the first author to challenge the monogeneric classification of Eurasian vipers, by separating the larger species into the genus *Daboia*, together with *Pseudocerastes*. This split however, was not adopted by most subsequent researchers. Herrmann, Joger and Nilson

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Table 1. Currently listed species of Eurasian vipers according to Uetz, Freed and Hošek (2019), depicting the criteria used for species designation, our recommended status, in some cases provisional, including justification, suggestions for further work and IUCN red list category (also for included taxa). Percentages of divergence from the closest sister species or clade are given for a small fragment of *cyt b* (196 bp; supplementary table S3). “DVAS” means Doubtful Validity As Species, may represent geographic variation, a subspecies or diverged population, hence, currently we recommend to decline its species status; “LSC” means Likely Species Complex (i.e., group of closely related taxa, possibly more than one species, but delimitations are not clear yet), hence, currently maintain species status; and “pending” indicates that, despite some incongruences with the divergence threshold delimitation, single-species recognition is currently maintained, but further research is required to assess taxonomic status. *Cyt b* divergence categories: Low = 2% or less; Moderate = 3 or 4%; High = 5% or more. LC = Least Concern; NT = Near Threatened; VU = Vulnerable; EN = Endangered; CR = Critically Endangered.

Species	Criteria	Status	Reason	Further work	IUCN
<i>Daboia mauritanica</i> Gray, 1849	morphological	species	high genetic divergence (15%)		NT (<i>D. deserti</i> NT)
<i>Daboia palaestinae</i> (Werner, 1938)	morphological	species	high genetic divergence (15%)		LC
<i>Daboia russelii</i> (Shaw and Nodder, 1797)	morphological	species	high genetic divergence (9%)		
<i>Daboia siamensis</i> (Smith, 1917)	genetic + morphological	species	high genetic divergence (9%)		LC
<i>Macrovipera lebetina</i> (Linnaeus, 1758)	morphological	species	high divergence from <i>M. razii</i> (9%)		
<i>Macrovipera razii</i> Orta et al., 2018	genetic + morphological	species	high genetic divergence from the remaining <i>Macrovipera</i> (9%)	subspecies should be examined for genetic divergence and extent of gene flow	
<i>Macrovipera schweizeri</i> (Werner, 1935)	morphological	DVAS	low divergence from <i>M. lebetina</i> , incl. Cyprus (2%) and shared haplotype with <i>M. lebetina</i> from the Turkish Mediterranean coast (Stümpel and Joger, 2009, Stümpel, 2012), despite 600 km distance between its range in Milos to nearest mainland population (Manavgat, Turkey)	extent of gene flow with <i>M. lebetina</i> should be examined relationship to <i>M. lebetina</i> should be clearly addressed, low genetic difference suggests a subspecific status for <i>M. schweizeri</i>	EN
<i>Montivipera albizona</i> (Nilson, Andrén and Flärdh, 1990)	morphological	DVAS	low genetic divergence from <i>M. bulgardaghica</i> (2%), subspecific status was suggested (Stümpel et al., 2016)	evaluate extent of gene flow with <i>M. wagneri</i> and <i>M. bulgardaghica</i>	EN
<i>Montivipera bommuelleri</i> (Werner, 1898)	morphological	species	high genetic divergence from <i>M. xanthina</i> (5-6%) and <i>M. raddei</i> (9%) complexes	low divergence within <i>M. bommuelleri</i> -group (2-3%), incl. <i>M. albizona</i> (see above), <i>M. bulgardaghica</i> and <i>M. wagneri</i> , extent of gene flow among these units should be evaluated	EN

Table 1. (Continued.)

Species	Criteria	Status	Reason	Further work	IUCN
<i>Montivipera bulgardaghica</i> (Nilson and Andrén, 1985)	morphological	DVAS	low to moderate genetic divergence from <i>M. albizona</i> (2%) and <i>M. bommuelleri</i> (4%); <i>M. albizona</i> should be included as ssp. (Stümpel et al., 2016)	evaluate extent of gene flow with <i>M. albizona</i> and <i>M. bommuelleri</i>	LC
<i>Montivipera kuhrangica</i> Rajabizadeh, Nilson and Kami, 2011	morphological	DVAS	included within <i>M. raddei</i> ; low genetic divergence (2%)		
<i>Montivipera latifii</i> (Mertens et al., 1967)	morphological	DVAS	included within <i>M. raddei</i> ; low genetic divergence (2%)		EN
<i>Montivipera raddei</i> (Boettger, 1890)	morphological	species	high genetic divergence to <i>M. xanthina</i> and <i>M. bommuelleri</i> (9%)	<i>M. kuhrangica</i> and <i>M. latifii</i> might be included, extent of gene flow with other units should be evaluated	NT (<i>M. albicornuta</i> VU)
<i>Montivipera wagneri</i> (Nilson and Andrén, 1984)	morphological	DVAS	sister to <i>M. albizona</i> - <i>M. bulgardaghica</i> ; low to moderate divergence within <i>M. bommuelleri</i> -group (2-3%)	evaluate extent of gene flow with other <i>M. bommuelleri</i> lineages	CR
<i>Montivipera xanthina</i> (Gray, 1849)	morphological	LSC	including 4 divergent lineages (Stümpel et al., 2016), with paraphyletic position and low branch support; high genetic divergence to <i>M. bommuelleri</i> (6%) and <i>M. raddei</i> (9%)	Taurus populations recently described as a new subspecies based on morphology of one live and three road-killed specimens is insufficient (Afsar et al., 2019); extent of geographic variation, limits, and gene flow between Taurus, Greek, Aegean and Lycian populations must be evaluated	LC
<i>Vipera altaica</i> Tuniyev, Nilson and Andrén, 2010	morphological	DVAS	included within <i>V. renardi</i> , low genetic divergence (1%)		
<i>Vipera ammodytes</i> (Linnaeus, 1758)	morphological	LSC	high genetic divergence to the remaining <i>Vipera</i> (11-5%); includes several divergent lineages (4-6%), two of them corresponding to <i>V. transcaucasiana</i>	evaluate extent of gene flow among lineages	LC
<i>Vipera anatolica</i> Eiselt and Baran, 1970	morphological	species	high genetic divergence to <i>V. ursinii</i> and <i>V. walseri</i> (6-8%)		CR

Table 1. (Continued.)

Species	Criteria	Status	Reason	Further work	IUCN
<i>Vipera aspis</i> (Linnaeus, 1758)	morphological	species	high genetic divergence to <i>V. latastai-monticola</i> (11%); several species recognized by Zuffi (2002) were subsequently rejected by genetic studies (Ursenbacher et al., 2006; Barbanera et al., 2009). West and East lineages show moderate to high genetic divergence (4–5%) nested within <i>V. berus</i> , showing low genetic divergence (2%), although apparent geographic isolation; subspecific status should be more appropriate	evaluate extent of gene flow among lineages	LC
<i>Vipera barani</i> Böhme and Joger, 1983	morphological + geographic isolation	DVAS	moderate to high genetic divergence (5%) and high genetic divergence (5%) and geographic isolation to <i>V. sevoanei</i>	evaluate extent of gene flow between western and eastern lineages	NT
<i>Vipera berus</i> (Linnaeus, 1758)	morphological	species	high genetic divergence to <i>V. sevoanei</i>	<i>V. barani</i> must be included, as well as other subspecies and clades already recognized such as <i>V. berus nikolskii</i> or <i>V. berus bosniensis</i>	LC
<i>Vipera dorevskii</i> Vedmederja et al., 1986	morphological	pending	high genetic divergence to <i>V. walser</i> (6%); sister to <i>V. kaznakovi</i> from Georgia, with moderate divergence (4%), and moderate to high genetic divergence to <i>V. sakoi</i> (4–5%)	relationships and gene flow to <i>V. kaznakovi</i> and <i>V. sakoi</i> must be clarified using integrative methods	CR
<i>Vipera dimiki</i> Nikolsky, 1913	morphological	DVAS	sister to <i>V. kaznakovi</i> from Russia, showing low genetic divergence (<1%)	test for gene flow with <i>V. kaznakovi</i>	VU
<i>Vipera erivanensis</i> (Reuss, 1933)	morphological	DVAS	low to moderate genetic divergence to <i>V. renardi</i> (2–4%); it includes <i>V. ebneri</i> , which was previously synonymized and shows low genetic divergence to <i>V. erivanensis</i> (1%); geographic isolation doubtful (Rajabizadeh et al., 2011, Tuniyev et al., 2018); private haplotypes in some nuDNA genes; a subspecific status must be appropriate.	nuclear inferences should be increased	VU (<i>V. ebneri</i> VU)
<i>Vipera graeca</i> (Nilsson and Andrén, 1988)	genetic + morphological	pending	moderate to high genetic divergence to <i>V. renardi</i> , <i>V. ursinii</i> and <i>V. kaznakovi</i> from Russia (3–5%); private haplotypes in one nuDNA gene and apparent geographic isolation in relation to closely related species	search for new populations over larger areas and test for gene flow with <i>V. ursinii</i>	EN

Table 1. (Continued.)

Species	Criteria	Status	Reason	Further work	IUCN
<i>Vipera kaznakovi</i> Nikolsky, 1909	morphological	pending	it includes two very divergent paraphyletic units, with high genetic divergence (10%); one in Georgia (type locality), sister to <i>V. darevskii</i> ; another in Russia sister to <i>V. orlovi</i> (admixed population of <i>V. kaznakovi</i> x <i>V. renardi</i>) and <i>V. dimniki</i> . RAD-seq data indicates low differentiation among these mtDNA lineages (Oleksandr Zinenko, unpublished).	relationships between both mitochondrial lineages of <i>V. kaznakovi</i> must be clarified, also to understand the relationship of Georgian populations to <i>V. darevskii</i> , Russian populations to <i>V. dimniki</i>	EN
<i>Vipera latastei</i> Boscá, 1878	morphological	LSC	it includes two divergent clades (7-8%), geographically isolated, one in Iberia and another in North Africa (<i>V. monticola</i> included), and further divergent lineages within each clade	populations from Iberia and North Africa must be recognized as distinct; extent of gene flow among further levels of structure within each main clade should be evaluated	VU
<i>Vipera lotievi</i> Nilsson et al., 1995	morphological	DVAS	paraphyletic, included within <i>V. renardi</i> with low genetic divergence (1-2%)		NT
<i>Vipera magnifica</i> Tuniyev and Ostrovskikh, 2001	morphological	DVAS	admixed population (<i>V. kaznakovi</i> x <i>V. renardi</i> ; Zinenko et al., 2016)		EN
<i>Vipera monticola</i> Saint Girons, 1953	morphological	pending	highly divergent lineage (6%) included within North African <i>V. latastei</i> clade	part of the North African <i>V. latastei</i> clade; extent of gene flow among lineages within this clade should be evaluated	NT
<i>Vipera olguni</i> (Tuniyev et al., 2012)	morphological	DVAS	nested within <i>V. darevskii</i> , with low genetic divergence (<1%), already suggested as a subspecies by Tuniyev et al. (2018)		
<i>Vipera orlovi</i> Tuniyev and Ostrovskikh, 2001	morphological	DVAS	admixed population (<i>V. kaznakovi</i> x <i>V. renardi</i> ; Zinenko et al., 2016)		CR
<i>Vipera pontica</i> Billing et al., 1990	morphological	DVAS	only two specimens known, represents a hybrid between sympatric <i>V. kaznakovi</i> and <i>V. transcaucasiana</i> (Baran et al., 2001; Oleksandr Zinenko, unpublished; Mebert et al., 2015a)		EN

Table 1. (Continued.)

Species	Criteria	Status	Reason	Further work	IUCN
<i>Vipera renardi</i> (Christoph, 1861)	morphological	species	moderate to high genetic divergence to <i>V. ursinii</i> and <i>V. kaznakovi</i> from Russia (4-6%)	<i>V. altaica</i> , <i>V. ebnieri</i> , <i>V. lotievi</i> and <i>V. eriwanensis</i> must be included	
<i>Vipera sakoii</i> (Tuniyev et al., 2018)	genetic + morphological	pending	moderate to high genetic divergence to <i>V. darevskii</i> (4-5%); however, only one independent sample was tested and geographic isolation from <i>V. darevskii</i> is uncertain	search for new populations over larger areas and test for gene flow with <i>V. darevskii</i>	
<i>Vipera seoanei</i> Lataste, 1879	morphological	species	high divergence (5%) and geographic isolation from <i>V. berus</i>		LC
<i>Vipera shenakhensis</i> (Tuniyev et al., 2013)	morphological + genetic	DVAS	low divergence to <i>V. eriwanensis</i> (less than 1%)		
<i>Vipera transcaucasiana</i> Boulenger, 1913	morphological	DVAS	two lineages included within <i>V. ammodytes</i> , with moderate to high genetic divergence (3-5%); recommended with subspecific status in Ursenbacher et al. (2008)	extent of gene flow must be evaluated to bridge large geographic gaps	NT
<i>Vipera ursinii</i> (Bonaparte, 1835)	morphological	species	moderate to high divergence to <i>V. renardi</i> and <i>V. kaznakovi</i> from Russia (4-6%)		VU
<i>Vipera walser</i> Ghielmi et al., 2016	genetic	species	high genetic divergence to <i>V. darevskii</i> (6%), geographically isolated, private haplotypes in two mtDNA genes (Ghielmi et al., 2016)		

(1992), using immunological distances, recognised *Macrovipera* for the *lebetina* group (including the taxa *mauritanica* and *deserti*) and restricted *Daboia* to the species *russelii*. More recently, Nilson et al. (1999) described *Montivipera* as a new subgenus for the *xanthina* group. This was subsequently raised to full genus level by Joger (2005). Lenk et al. (2001), using mitochondrial DNA sequences, assigned the species *mauritanica*, *deserti* and *palaestinae* to *Daboia*, leading to the current generic arrangement of the group. The recent use of subgenus *Pelias* (Merrem, 1820) as a full genus for the *berus* and *ursinii* groups (e.g., Avcı et al., 2010; Tuniyev et al., 2012, 2013, 2018a, b), on the other hand, has remained a minority opinion in the literature. Certain changes proposed outside the peer-reviewed scientific literature are not considered here for reasons given in Kaiser et al. (2013).

One third of the currently recognized species were described in the 18th and 19th centuries by recognized taxonomists and zoologists of the time (e.g., Carl Linnaeus described *Coleber lebetinus*, *C. berus*, *C. aspis* and *C. amodytes* (now in *Macrovipera* and *Vipera*) in 1758; John Edward Gray described *Clotho mauritanica* and *Daboia xanthina* (now in *Daboia* and *Montivipera*, respectively) in 1849; Eduard Boscá described *Vipera latastei* in 1878; table 1). During the 20th century, eighteen of the currently recognized species were described; ten of them were described before or during the 1970s, and eight after that period. All species descriptions were based on morphological traits (i.e., scale counts, biometric measures, colour patterns), with the application of statistical analyses of these morphological and other phenotypic traits gradually becoming incorporated during more recent times (e.g., Herrmann, Joger and Nilson, 1992; Nilson and Andrén, 2001).

Nine species were named since 2001 (Uetz, Freed and Hošek, 2019; table 1). Three of them

were described using integrative taxonomy approaches, first addressing phylogenetic relationships and later characterizing phenotypic variability (i.e., *Vipera walser* Ghielmi et al., 2016; *Vipera sakoi* Tuniyev et al., 2018; *Macrovipera razii* Oraie et al., 2018). However, the descriptions for the other six species were done in a traditional way, i.e., by solely recording or analysing morphological traits, without the support of molecular data or phylogenetic evidence (*Vipera magnifica* Tuniyev and Ostrovskikh, 2001; *Vipera orlovi* Tuniyev and Ostrovskikh, 2001; *Vipera altaica* Tuniyev, Nilson and Andrén, 2010; *Montivipera kuhrangica* Rajabzadeh, Nilson and Kami, 2011; *Vipera olguni* Tuniyev et al., 2012; *Vipera shemakhensis* Tuniyev et al., 2013).

Recent phylogenetic and phylogeographic studies have transformed the taxonomic panorama in Eurasian vipers considerably, validating some taxa as species (e.g., *Vipera graeca*, Mizsei et al., 2017), rejecting or synonymising others (e.g., *Vipera altaica* with *V. renardi*, Zinenko et al., 2015, *Montivipera albicornuta* with *M. raddei* and *M. albizona* with *M. bulgardaghica*, Stümpel et al., 2016; *Daboia deserti* with *D. mauritanica*, Martínez-Freiría et al., 2017a; *Vipera magnifica* and *V. orlovi* were identified as admixed populations, Zinenko et al., 2016), or modifying previously designated taxonomic units (e.g., assigning species to four genera; Garrigues et al., 2005). However, despite this multitude of studies, there is still considerable uncertainty regarding the validity of some species, which is hampering the development of optimized conservation strategy for the whole group.

Here, we apply an integrative approach to review the taxonomy of the Eurasian vipers, by bringing together and analysing existing molecular and morphological data. We compiled and analysed published and new mitochondrial and nuclear DNA sequences for this group to reconstruct phylogenetic relationships among currently recognized species. We also compiled information on external morphology, as well as

on the criteria used for species delimitation, in order to assess the morphological distinctiveness of currently recognized species. Our objectives are: 1) to provide an updated taxonomy of Eurasian vipers; 2) to evaluate the validity of some of the newest species designations under the unified species concept and recommend an appropriate status reflecting our current data base; 3) to identify remaining knowledge gaps and the research required to achieve a robust, stable and evidence-based taxonomic framework for this group of vipers.

Material and methods

Taxonomic inference

Our taxonomic evaluation is built upon an integrative and evolutionary framework, based on the unified species concept (de Queiroz, 2007), under which species are defined as separately evolving lineages and their biological properties (e.g., monophyly, reproductive isolation, differentiated ecological niches, morphological distinctiveness) as “operational criteria” that ultimately provide evidence for their separation through a variety of methods. We employ the integrative approach of Padial et al. (2010) by using phylogenetic analysis of mitochondrial (mtDNA) sequences, the most widely available and standardised marker, to identify divergent lineages, that are then tested, to the extent that the available data allow, with additional data, in particular single-copy nuclear DNA (nuDNA) markers and morphological data. We set a sequence divergence percentage to propose a threshold for provisional taxonomic categorization (see phylogenetic inference section for more details). Above this threshold, evolutionary lineages may simply confirm established species, or if not described as such, they may be considered as candidate species and should be targeted in future studies to evaluate their taxonomic status. If currently recognized species are composed of several divergent lineages, we classify them as a Likely Species Complex (LSC). Below the *cyt b* threshold, currently recognized species are categorized as Doubtful Valid as Species (i.e., DVAS), until there is enough evidence to indicate species integrity. In addition, we used the “pending” category to indicate that single-species recognition is currently maintained, despite incongruences with the divergence threshold delimitation, but further research is recommended to assess taxonomic arrangements.

Phylogenetic inferences

We searched on GenBank for mtDNA and nuDNA gene sequences representing all the relevant lineages within Eurasian vipers (see supplementary table S1 and S2). Selection of sequences was based on published phylogenetic

and phylogeographic studies (e.g., Ursenbacher et al., 2006, 2008a, b; Velo-Antón et al., 2012; Zinenko et al., 2015; Stümpel et al., 2016; Mizsei et al., 2017; Freitas et al., 2018; Martínez-Freiría et al., 2020) to ensure that the selected sequences represent the genetic structure and units reported in those studies. Mitochondrial DNA sequences were available for a total of nine gene fragments, as well as the whole mitochondrial DNA genome for *D. siamensis* (mislabelled as *D. russelii* in GenBank). For phylogenetic analysis, we selected DNA fragments from a subset of seven mtDNA markers with a higher representation across species: CR (control region), COI (Cytochrome c oxidase subunit I), *cyt b* (cytochrome b), ND2 (NADH dehydrogenase subunit 2), ND4 (NADH dehydrogenase subunit 4), ND5 (NADH dehydrogenase subunit 5) and 16S (mitochondrial gene coding for 16S rRNA).

Additionally, we also provide unpublished DNA sequences generated for other studies to complement available data from GenBank. Sequences were concatenated when they originate from the same lineage or geographic locality, using SequenceMatrix software (Vaidya, Lohman and Meier, 2011). Since Eurasian viper species generally contain geographically cohesive, parapatric mitochondrial lineages, it is unlikely that they generate chimeras when concatenating sequences from multiple individuals from the same geographic locality. The final dataset included 97 units (representing several lineages for 39 of the 40 recognized species) with sequences ranging from 654 to 4621 base pairs (bp). Details of sequences used in mtDNA analyses are available in supplementary table S1. Sequences for nuDNA were available for 17 gene fragments, from which we selected six protein-coding genes, BDNF (Brain Derived Neurotrophic Factor), CMOS (Oocyte maturation factor Mos), MC1R (Melanocortin 1 Receptor), NT3 (Neurotrophin-3), PRLR (Prolactin Receptor), RAG1 (Recombination Activating protein 1) and one intron, B-fib (Beta-fibrinogen intron 7), as they allowed the most comprehensive taxonomic coverage (24 species, 61.5% of the currently recognized total). This set includes the most relevant nuclear genes used to support the designation of some of the most recently recognized species (e.g., RAG1, Ghielmi et al., 2016; NT3, Mizsei et al., 2017). Details on nuDNA sequences used are provided in supplementary table S2. Sequences were manually aligned and edited using Geneious v 4.8.5 (Kearse et al., 2012). For the nuclear genes, haplotype phases were produced by a coalescent-based Bayesian reconstruction implemented in PHASE (Stephens, Smith and Donnelly, 2001) available in DNAsp (Librado and Rozas, 2009).

Phylogenetic relationships and time of divergence between species were inferred using a Bayesian Inference (BI) method implemented in BEAST v 1.7.5 (Drummond et al., 2012) on the concatenated mtDNA dataset. An exhaustive search with PartitionFinder 1.1.1 (Lanfear et al., 2012) was conducted to select appropriate partitioning schemes and evolutionary models based on the Bayesian Information Criterion (BIC). The GTR + G + I model applied to all mtDNA fragments combined in a single partition was determined as the best-fit model and partitioning scheme.

Substitution rates were estimated under a strict molecular clock (Drummond et al., 2006) that assumes uniform rates

across branches. A Yule model, most suitable for species-level phylogenies, was implemented as tree prior. Since the fossil record of Eurasian vipers is fairly incomplete and does not provide reliable and verified calibration dates (see Stümpel et al., 2016), our molecular dating strategy relied on secondary calibrations, including the splits of *Vipera-Daboia* and *Macrovipera-Montivipera*, dated at 26 Mya (Zheng and Wiens, 2016 – but see Šmíd and Tolley, 2019). We used a lognormal prior with a mean of 26.3 Mya and a standard deviation of 0.07 to constrain node ages. Three independent runs of 100 million generations were performed, sampling trees and parameter estimates every 10 000 generations with 10% of the trees discarded as burn-in. Convergence was verified by looking at the effective sample sizes of all parameters (ESS > 300) using Tracer v1.7 (Rambaut et al., 2018). Trees obtained from multiple independent runs were then combined using LogCombiner v 1.7.5 (Drummond et al., 2012) and summary trees were generated with TreeAnnotator v1.7.1 (Drummond et al., 2012).

Haplotype networks for the *cyt b* gene (the most widely used marker across studies) were reconstructed with TCS v 1.21 (Clement, Posada and Crandall, 2000), with a 90% parsimony connection limit; and the graphical output was visualized in TCSBU (dos Santos et al., 2015). The initial alignment of 1141 bp was trimmed down to 196 bp with no missing data across 96 units. The short length of the single sequence available for *V. shemakhensis* precluded its inclusion in the haplotype network. However, it was directly compared to the remaining sequences (180 bp of overlap). Uncorrected p-distances between taxa were estimated based on the same *cyt b* fragment using MEGA ver. 5 (Tamura et al., 2011). We propose the use of this short standardized *cyt b* fragment as a candidate DNA barcode for the delimitation of evolutionary units in Eurasian vipers (see Hebert and Grégory, 2005). Furthermore, we recommend a value of 5% *cyt b* sequence divergence as a provisional threshold, below which an untested species-level designation appears inappropriate or premature, unless other lines of evidence would validate species classification. The *cyt b*-threshold results from two facts: 1) a 5% threshold provides a good delimitation of currently recognized species within Eurasian vipers, also recognizing deep evolutionary lineages within them; 2) *cyt b* divergence levels are consistently higher than 5% between closely related species that co-exist with restricted or no hybridization, e.g. sympatric species of the genus *Vipera* differ by more than 10% (Tarroso et al., 2014; Mebert et al., 2015b), sympatric watersnakes *Nerodia fasciata* and *N. sipedon* differ by 9% (Mebert, 2008), *Natrix helvetica* and *N. natrix* by 6.9% (Kindler et al., 2017) and 9% difference between *Montivipera wagneri* and *M. raddei* with no signs of mixing along a sharp contact line (Mebert et al., 2016; Stümpel et al., 2016).

Haplotype networks were drawn for each of the seven nuclear genes following the same procedure. Sequences with high proportion of missing data (>30% of the total length) were excluded from the dataset. For B-fib and RAG1, datasets were divided in two sets each based on sequence length (B-fib-1, B-fib-2 and RAG1-1, RAG1-2) and analysed independently to avoid excluding shorter sequences from the haplotype networks.

Morphological characterization

From the published literature, we compiled a list of the criteria used to identify each of the currently recognized species. In addition, we gathered information on the variability of 14 external morphological traits in each species, including one biometric, 11 pholidotic and two dorsal colour pattern characters. Information from the literature was collated with data from specimens measured by the authors in the field or in museum collections. This allowed us to establish the range of variation of different morphological traits, in some cases for each sex separately (see Results). In addition, whenever possible, and in order to more accurately represent morphological variation in some pholidotic traits, modal or mean values were retrieved for specific groups or subspecies within each species. Furthermore, we included verbal descriptions of colouration, which provide an idea of variation in visually striking qualitative traits.

Results

Phylogenetic inferences from mtDNA

We obtained a well-supported (most major nodes with posterior probability > 0.95) mtDNA phylogenetic reconstruction for the 97 taxa (fig. 1, supplementary fig. S1). Three main clades are recognized, corresponding to genera *Daboia*, *Vipera* and *Macrovipera* + *Montivipera*. Diversification times varied widely, starting with *Daboia* (mean time to most recent common ancestor, TMRCA = 20.97 Mya), followed by *Vipera* (TMRCA = 16.65 Mya), and later the split between *Macrovipera* and *Montivipera* (TMRCA = 12.86 Mya). Diversification within *Macrovipera* and *Montivipera* started later, around 8 Mya.

Vipera is the most diverse genus and includes three subclades: (1) *Pelias*, comprising one lineage with *V. berus* (including *V. barani* nested in it) and *V. seoanei*, and another lineage with *V. renardi*, *V. ursinii*, *V. kaznakovi*, *V. graeca*, *V. sakoi*, *V. darevskii*, *V. walser* and *V. anatolica* (as well as other species nested inside *V. renardi*, *V. kaznakovi* and *V. darevskii*); (2) *Vipera* 1, including *V. aspis* and the *V. latastei-monticola* complex; and (3) *Vipera* 2, comprising the *Vipera ammodytes-transcaucasiana* complex. Diversification within each subclade is <10 Mya, with *Vipera* 1 being the oldest

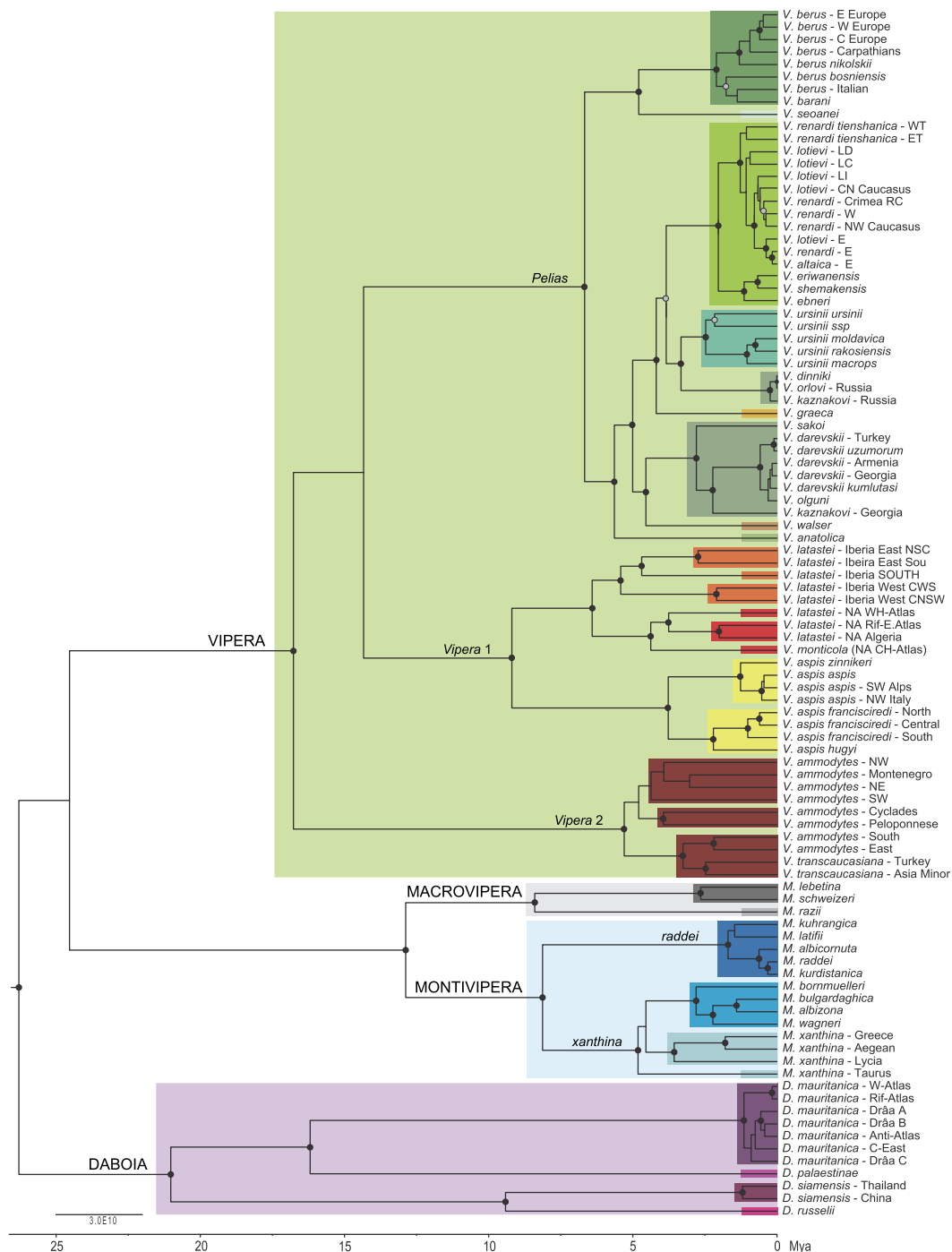


Figure 1. Dated Bayesian phylogenetic tree, obtained from the concatenated dataset of seven mitochondrial gene fragments, showing relationships for Eurasian vipers. The four genera (in uppercase) and major subclades (in italics) are highlighted. Lineages are grouped and distinctively coloured considering a divergence equal or higher than 5% for a small fragment (196 bp) of *cyt b* (supplementary table S3). Black and light-grey dots represent posterior probabilities higher than 0.95 and between 0.9-0.95, respectively. Names of taxa and lineages are given accordingly to publications from where sequences were retrieved. See supplementary table S1 for details.

(TMRCA = 9.11 Mya), followed by *Pelias* (TMRCA = 6.6 Mya) and *Vipera* 2 (TMRCA = 5.27 Mya).

The second most diverse clade is *Montivipera*, composed of two subclades: (1) *raddei*, which diversified ca. 1.68 Mya, and (2) *xanthina*, which diversified in three lineages ca. 4.78 Mya. *Macrovipera* is the least diverse clade and includes two subclades, *M. razii* and *M. lebetina* + *M. schweizeri*; the latter diverged about 2.62 Mya. *Daboia* includes three subclades, corresponding to *D. russelii* + *D. siamensis*, *D. palaestinae* and *D. mauritanica*. The latter two are sister species (TMRCA = 16.17 Mya). Divergence of *D. russelii* and *D. siamensis* is estimated about 9.3 Mya.

The range of divergence times between species and species complexes (i.e., group of closely related species) is highly variable (fig. 1), e.g., 9.1 Mya (*V. aspis* from *V. latastei-monticola*), 5.54 Mya (*V. anatolica* from the remaining *kaznakovi-ursinii*), 4.74 Mya (*V. seoanei* from *V. berus-barani*), 3.79 Mya (*V. ursinii* from *V. renardi* complex), 2.02 (*V. eriwanensis* from *V. renardi*) or 1.38 Mya (*V. barani* from *V. berus*). The TMRCA between species is found to be very recent for the pairs *V. altaica* with *V. renardi* – E, *V. orlovi* with *V. kaznakovi* – Russia, and *V. olguni* with *V. darevskii kumlutasi* (fig. 1). Several species were recovered as paraphyletic (e.g., *M. xanthina* and *V. lotievi*) or polyphyletic (e.g., *V. kaznakovi*).

Haplotype networks based on the 196 bp *cyt b* fragment recover similar genetic relationships to the mtDNA phylogenetic tree (supplementary fig. S2). The 82 identified haplotypes and matrices of uncorrected genetic distances based on these *cyt b* fragments are provided in supplementary material (supplementary tables S1 and S3, respectively). The single sequence available for *V. shemakhensis* differed in one position from the sequences of *V. ebneri* and *V. eriwanensis*.

Phylogenetic inferences from *nuDNA*

Haplotype networks constructed for the seven nuclear genes show a pattern of wide haplotype sharing and few mutational steps among species; however, some species present distinct, well differentiated haplotypes (fig. 2; supplementary table S4). Haplotype networks for BDNF, CMOS, MC1R and RAG1 (both sets) show very low levels of variability. Unique haplotypes were found for some species such as *V. berus* and *V. eriwanensis* in BDNF, *M. lebetina*, *M. raddei* and *D. siamensis* in CMOS, *V. aspis* in MC1R, or *V. ammodytes*, *V. berus* and *V. walser* in RAG1. However, haplotype sharing was found between *V. ursinii*, *V. renardi* and *V. graeca* in BDNF, between *V. berus*, *V. eriwanensis* and *V. aspis* in CMOS, and between *Montivipera* species in MC1R and RAG1. Similarly, some phylogenetically distant species (according to mtDNA and previous multilocus phylogenies), such as *M. lebetina* and *M. raddei* or *D. siamensis* and *V. ursinii* have very similar haplotypes (one or two mutational steps different) in the BDNF network; the same occurs for *M. lebetina* and *Montivipera* spp. in the set 2 of RAG1.

Haplotype networks for B-fib, PRLR and NT3 present higher variability than the other selected markers (fig. 2). The B-fib shows unique, well-separated haplotypes (more than three mutational steps) for *D. mauritanica*, *D. siamensis*, *M. lebetina* and *V. aspis*. However, B-fib-1 haplotypes are very similar in *V. seoanei* and *V. aspis*, and identical in North African and Iberian populations of *V. latastei*, as well as in *V. monticola*. The PRLR dataset displays unique, well-differentiated haplotypes in most taxa: *V. berus*, *V. aspis*, *V. renardi*, *V. seoanei*, *D. mauritanica*, *D. siamensis*, *M. lebetina*, some samples of *V. ursinii* and some samples of *V. latastei* from Iberia. However, it shows haplotype sharing between *V. ursinii* and *V. graeca*, between *V. latastei* from Iberia and North Africa

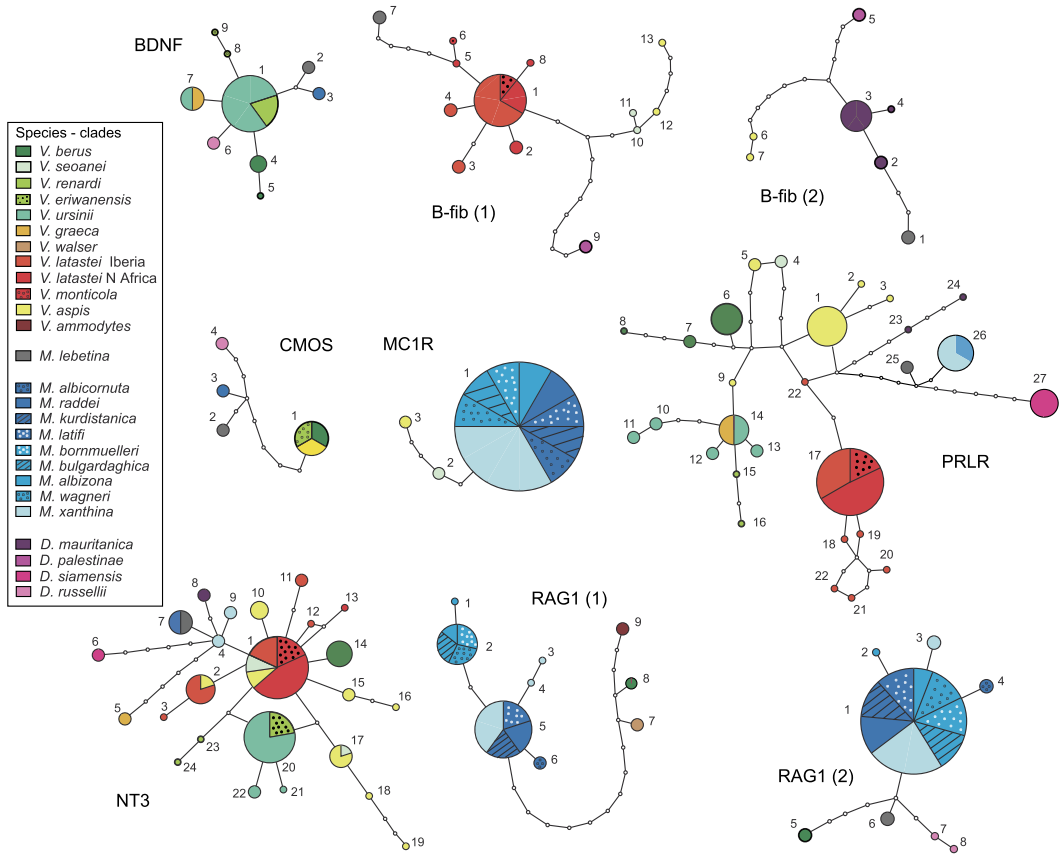


Figure 2. Haplotype networks for seven nDNA genes. B-fib and RAG1 genes were divided in two sets each (see text and supplementary tables S2-S3 for details).

as well as *V. monticola*, and between *M. xanthina* and *M. albizona*. The NT3 dataset includes samples of several *Vipera* species, *M. lebetina*, *M. raddei*, *M. xanthina*, *D. mauritanica* and *D. siamensis*. It shows haplotype sharing among species of the same genus (e.g., *V. aspis* – *V. latastei-monticola* – *V. seoanei*, *V. ursinii* – *V. renardi*), and even of distinct genera (*Macrovipera lebetina* – *Montivipera raddei*). Some species such as *D. mauritanica*, *M. xanthina*, *V. berus*, *V. renardi*, *V. graeca* or *D. siamensis* have distinct haplotypes, the two latter species being well separated from the rest.

Morphological characterization

We obtained data from 39 studies providing morphological descriptions or addressing the

variability of species, 13 assessing phylogenetic relationships among species, and six using phylogenetic inferences and phenotypic characterization to describe taxa (see supplementary references S1). One book (Phelps, 2010) was used to extract maximum body size for some species for which other published data were lacking. Information retrieved from publications, complemented with data collected by the authors from museum specimens, is shown in supplementary table S5.

The most commonly used criteria to diagnose species relied on head (24 species) and body (26 species) pholidosis, and dorsal colouration (27 species). In contrast, phylogenetic analyses based on molecular data were initially considered for species description in only five cases (table 1, supplementary table S5).

Data compilation showed that species within *Macrovipera* and *Daboia* are the largest in body size, followed by *Montivipera* and with *Vipera* being the smallest. Sexual dimorphism in body size is reported in most species (supplementary table S5).

Regarding pholidosis, some traits of head scalation (e.g., canthal, supralabial or infralabial scales) exhibit low variation, particularly within each genus (supplementary table S5). Modal or mean values of other head traits, however, show important variation within genera (e.g., apical and intercanthal + intrasupraocular scales within *Vipera*) and among them (e.g., loreal scales; fig. 3). Modal or mean values of body scalation exhibit variation among species (e.g., ventral and subcaudal scales; fig. 5) or genera (e.g., number of dorsal rows; fig. 4). Nevertheless, the ranges of variation for most pholidotic traits overlap extensively among species of the same genus and even among distinct genera (figs 3, 4). Sexual dimorphism in subcaudal scale counts is also mirrored in ventral scale variation in many species (fig. 4).

With respect to dorsal colouration, there is high variability at both the inter- and intraspecific levels, particularly in *Vipera* species, which display more dorsal marks and a higher number of distinct dorsal pattern types than species of the other genera (supplementary table S5).

Discussion

Eurasian vipers are a taxonomically challenging group due to the long-standing use of diverse species criteria and the prevalence of morphology-based classifications and species delimitations. Here, we inferred the molecular phylogeny of Eurasian vipers based on mitochondrial data, encompassing for the first time almost all the described species and evolutionary lineages within this group. We also gathered information on the morphological variability of currently recognized species and reviewed the

criteria used to identify them. By assessing phylogenetic and morphological variability, we provide recommendations and future research directions for robust species delimitation, which will aid the advancement towards a more informed and coherent taxonomy for this group of vipers.

Phylogenetic inference from mtDNA

Phylogenetic analyses based on seven mtDNA fragments produced a mostly resolved topology that strongly supports the monophyly of the four genera (fig. 1). Phylogenetic relationships and divergence dates estimated in this study mostly agree with those reported in previous works (e.g., Alencar et al., 2016; Stümpel et al., 2016; Zheng and Wiens, 2016), with the exception of divergence dates recently reported using alternative time calibration procedures (Šmíd and Tolley, 2019).

Overall, our inferences suggest distinct diversification dates and divergence levels for each genus. Diversification within *Daboia* is estimated in the early Miocene, followed by the diversification within *Vipera* in the middle Miocene; *Macrovipera* and *Montivipera* appear as the most recent genera, diverging from each other in the late Miocene (fig. 1). Despite its old origin, *Daboia* shows low diversity levels, with only four species recognized: two tropical Asian species, *D. russelii* and *D. siamensis*, and two Mediterranean species, *D. mauritanica* and *D. palaestinae*. These species, however, are extremely divergent, with genetic distances based on a small (196 bp), variable fragment of *cyt b* ranging from 9%, between the Asian relatives *D. russelii* – *D. siamensis*, to 21%, between *D. siamensis* and the Mediterranean *D. palaestinae* (supplementary table S3). The old diversification and wide distributional range of this genus, together with the spatial gaps separating species ranges, suggest that this group likely experienced major extinctions along its evolutionary history, with only some of the representative taxa currently persisting. This suggestion is supported by the occurrence of *Daboia*-like vipers

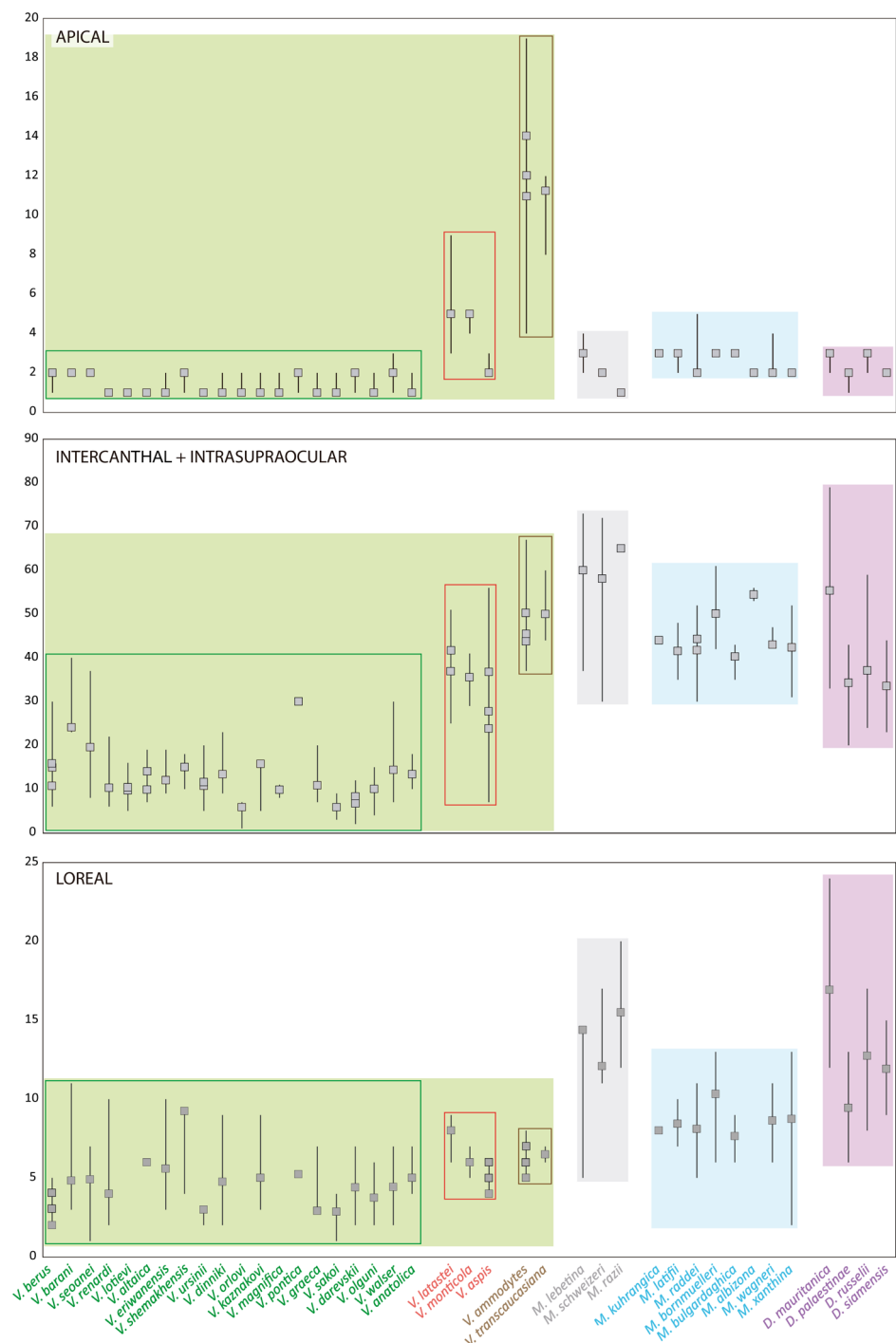


Figure 3. Variation across species and genera in three pholidotic head traits (number of apical, intercanthal + intrasupraocular, and loreal scales). For each species, variation range (minimum-maximum) is represented as a vertical line. Modal, in apical and loreal scales, and mean values, in intercanthal + intrasupraocular scales, are represented as grey squares. Both values were retrieved for specific groups or subspecies (supplementary table S5). Species are displayed and grouped according to mtDNA phylogenetic relationships in fig. 1.

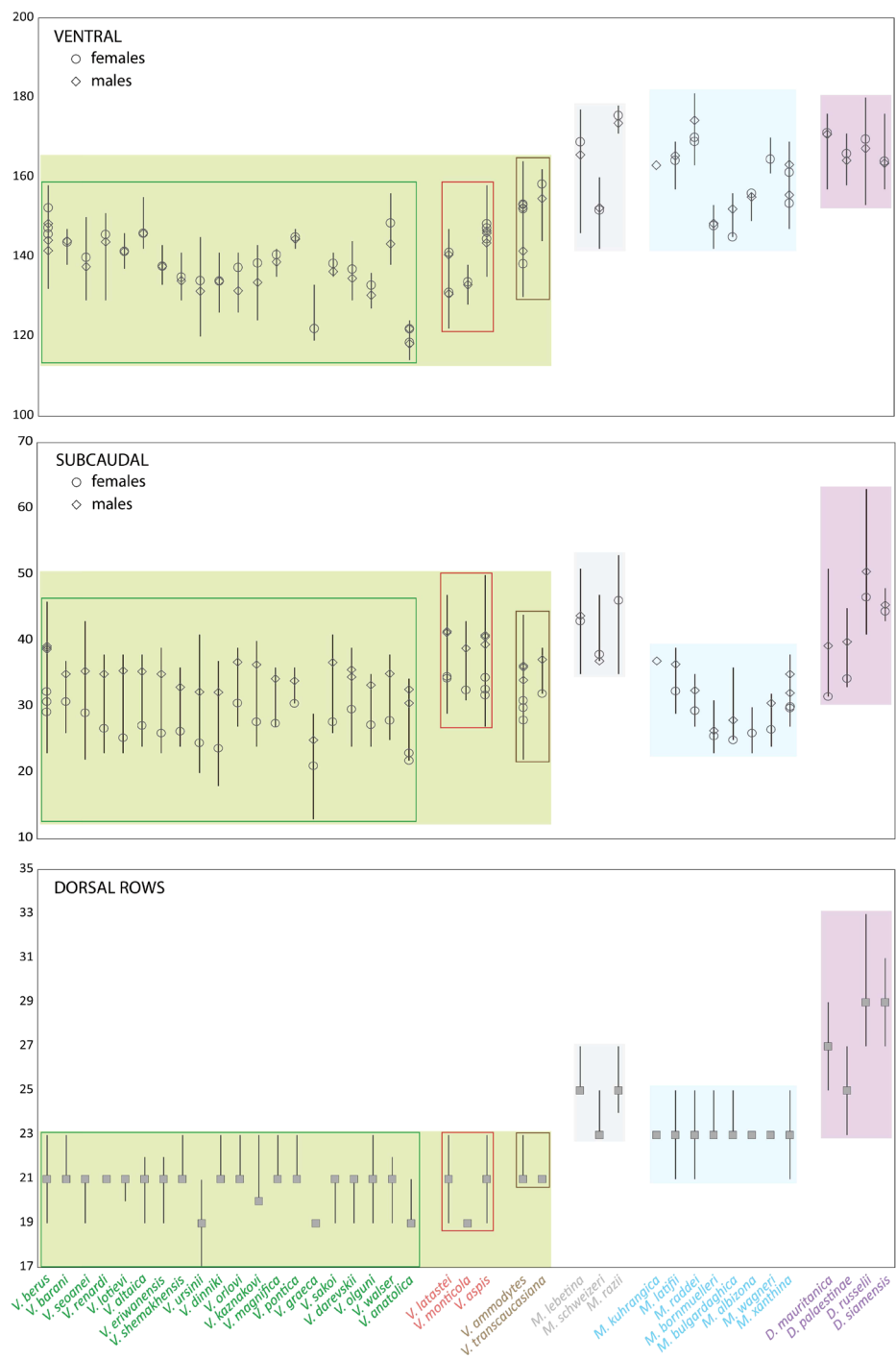


Figure 4. Variation across species and genera in three pholidotic body traits (number of ventral and subcaudal scales, and dorsal rows). For each species, variation range (minimum-maximum) is represented as a vertical line. Mean values are represented as circles (females) and rhomboids (males) in ventral and subcaudal scales, while modal values are represented as a grey square in dorsal rows. Both values were retrieved for specific groups or subspecies (supplementary table S5). Species are grouped according to mtDNA phylogenetic relationships in fig. 1.

in the fossil record from the Miocene in areas such as Western Europe, where this group is currently absent (Paleobiology Database, 2019).

With a more recent origin, *Macrovipera* also shows low diversity levels, comprising three described species (fig. 1): *M. schweizeri*, from the western Cyclades, *M. lebetina*, widely distributed from central Asia to the Middle East, and *M. razii* from southern and central Iran. High levels of polymorphism, especially in colouration, led to the description of distinct subspecies within *M. lebetina* (Nilson and Andrén, 1988; Stümpel and Joger, 2009). Molecular studies have suggested the validity of four subspecies, i.e., *lebetina*, *obtusa*, *turanica* and *cernovi* (Stümpel and Joger, 2009), but rejected the species status of *M. schweizeri*, which is suggested to be a subspecies of *M. lebetina* with a wider distribution than previously considered (Lenk et al., 2001; Stümpel and Joger, 2009). Our phylogenetic analyses recover *M. schweizeri* and *M. lebetina* as a single unit (2% of *cyt b* genetic distance; supplementary table S3). Shared mitochondrial haplotypes between *Macrovipera schweizeri* and some southern Turkish populations of *M. lebetina* further corroborate conspecificity (Stümpel and Joger, 2009). Oraie et al. (2018) found cytochrome *b* divergences between moderate and high (up to 4.4%) among Iranian populations of *M. lebetina obtusa* and *M.l. cernovi* and between these and specimens from Uzbekistan and Azerbaijan, suggesting that further diversity may exist within this species. Moreover, increased geographical sampling may uncover additional phylogenetic diversity, as has been the case with the description of *M. razii* from Iran, previously allocated to *M. lebetina* (Oraie et al., 2018).

The genus *Montivipera* consists of two well-supported species complexes, the *xanthina* and *raddei* clades. This genus was recently subjected to a comprehensive phylogenetic study based on a multilocus mitochondrial and nuclear dataset for all its constituent taxa (Stümpel et al., 2016). Here we confirm previous findings of a more recent origin of the *raddei*-complex

(ca. 1.68 Mya, 2% genetic divergence within the group) and an older divergence with higher levels of genetic diversity within the *xanthina*-complex, including the *bornmuelleri*-group (ca. 4.78 Mya, 6% genetic divergence within the entire complex). Major branches within the latter group are supported by high posterior probabilities, except for the monophyly of *M. xanthina*, for which low branch support was already shown in Stümpel et al. (2016).

Among all analysed genera, *Vipera* is the most diverse and well-studied group. Extensive phylogeographic and phylogenetic work has been conducted within this genus (e.g., Garrigues et al., 2005; Ursenbacher et al., 2006a, b, 2008; Velo-Antón et al., 2012; Zinenko et al., 2015, 2016; Martínez-Freiría et al., 2020). However, no previous study has included all known taxa (e.g., 12 taxa in Zheng and Wiens, 2016; 19 taxa in Šmíd and Tolley, 2019). Although traditionally grouped in two subgenera (Garrigues et al., 2005), *Vipera* forms three well-supported monophyletic groups (fig. 1): *Pelias*, *Vipera* 1 and *Vipera* 2 (see Nilson and Andrén, 1997 for a similar designation). Both *Vipera* 1 and *Vipera* 2 present deep phylogenetic structure and high divergence between and within taxa. *Vipera* 1 includes the western and Mediterranean *V. aspis* and *V. latastei-monticola*, and *Vipera* 2 is represented by *V. ammodytes-transcaucasiana* from the Balkans, Turkey and Asia Minor. *V. latastei-monticola* and *V. ammodytes-transcaucasiana* are species complexes that comprise highly divergent units (9% and 6% of *cyt b* genetic distance within each complex, respectively; supplementary table S3). Phylogeographic studies on these taxa show high levels of geographically structured genetic diversity and the oldest divergences among Eurasian vipers, with main diversification events likely occurring during late Miocene and early Pliocene (Ursenbacher et al., 2008; Velo-Antón et al., 2012; Freitas et al., 2018; Martínez-Freiría et al., 2020).

On the other hand, *Pelias* is the most diversified group in the phylogeny, with multiple described species. In general, genetic divergence within this group is shallower than that observed within the other *Vipera* subclades, with many taxa likely resulting from geographic splits during Pleistocene climatic oscillations (see Zinenko et al., 2015). Our phylogeny recovered two highly divergent subclades (TMRCA = 7 Mya, fig. 1; 9% genetic distance, supplementary table S3). One of the subclades includes *V. berus* (with *V. barani* nested within it) and *V. seoanei*. The other clade is highly diversified and includes three major groups: (1) *V. renardi* (with *V. shemakhensis*, *V. lotievi* and *V. altaica* nested within it) and *V. eriwanensis* (with *V. ebneri*) as sister group, together with *V. ursinii* and *V. kaznakovi* from Russia (*V. orlovi*, is an admixed population of *V. kaznakovi* and *V. renardi*, Zinenko et al., 2016; and *V. dinniki* is nested within *V. kaznakovi* from Russia), and *V. graeca*; (2) *V. sakoi*, *V. kaznakovi* from Georgia, *V. darevskii* (with *V. olguni* nested within) and *V. walseri*; and (3) *V. anatolica*, which appears as a separate lineage at the root of this clade.

Our results support most taxa and recognized complexes (e.g., *V. latastei-monticola*) as monophyletic lineages, with the exception of *V. kaznakovi* and *V. lotievi*. As already shown by Zinenko et al. (2015) and Ghielmi et al. (2016), *V. kaznakovi* from Russia (Greater Caucasus) and Georgia (Lesser Caucasus) appear in the phylogeny as two polyphyletic lineages separated by a considerable genetic distance (5%), and *V. lotievi* is also polyphyletic, with several lineages included within *V. renardi* (fig. 1, supplementary fig. S2). Multilocus RAD-sequencing data recovered low differentiation between both mitochondrial lineages within *V. kaznakovi* (Oleksandr Zinenko, unpublished data), a pattern that is further supported by low morphological differentiation and the existence of a continuous area of suitable habitats connecting these populations (Orlov and Tuniyev, 1990). Reasons for the discordance between mtDNA and nuDNA

are still unclear but a historical occurrence of introgressive hybridization with asymmetric mitochondrial DNA capture could explain this pattern (e.g., Barbanera et al., 2009). The polyphyletic status of *V. lotievi*, on the other hand, is thought to be the result of possible confusion in the identification of species due to morphological convergence or hybridization and introgression leading to admixture of traits (Zinenko et al., 2015, 2016).

Mito-nuclear discordance and low resolution of nuDNA

The nuclear data do not follow the same pattern observed for mtDNA. The haplotype networks constructed for each nuDNA marker show a pattern of widespread haplotype sharing among distant species, and even among genera, with extremely low levels of genetic variation (fig. 2), suggesting incomplete lineage sorting of ancestral polymorphism (Wan et al., 2004) or extremely low levels of sequence evolution. Nuclear genes were already known to provide insufficient resolution to infer the phylogenetic variability within Eurasian viper taxa (e.g., *V. latastei-monticola* and *V. aspis*, Velo-Antón et al., 2012; Freitas et al., 2018; Martínez-Freiria et al., 2020; *D. mauritanica*, Martínez-Freiria et al., 2017a; *Montivipera* taxa, Stümpel et al., 2016). Yet, relying on mtDNA as the only source for phylogenetic inference could be problematic (see Ballard and Whitlock, 2004), and thus, the phylogenetic patterns retrieved here should be interpreted with some caution. The existence of local hybridization between highly differentiated species (e.g., Tarroso et al., 2014; Guiller, Lourdais and Ursenbacher, 2017) and the growing evidence for extensive gene flow in more recently differentiated taxa (Zinenko et al., 2015, 2016) highlight the importance of nuDNA analyses. However, despite being commonly used in species delimitation studies, the sequenced nuDNA markers were too conserved to consistently differentiate between otherwise well supported species of Eurasian vipers. This applies even to introns and other

fast-evolving single copy nuclear genes (e.g., NT3, PRLR, Townsend et al., 2008). Consequently, other molecular approaches such as even faster evolving markers (e.g., Kindler and Fritz, 2018; Pöschel et al., 2018), phylogenomic methods (e.g., Blair et al., 2019; Heinicke et al., 2018), and increased sampling to evaluate current or past gene flow between most-proximate populations or contact zones of two or more closely related species (Mebert, 2008, 2015a, b, 2020; Hillis, 2019) should be favoured to infer evolutionary histories and resolve species limits among these taxa.

Taxonomic relevance of morphological traits

The value of morphological traits to define species in many groups has become questionable after the emergence of DNA-based methods in taxonomy. Morphological variation across populations often reflects local adaptation processes or phenotypic plasticity, rather than historical relationships (e.g., Kaliontzopoulou et al., 2011; Kaliontzopoulou, Carretero and Llorente, 2012; Alhajeri, Hunt and Stepan, 2015). Additionally, lack of morphological differentiation does not necessarily imply shared evolutionary histories, as is the case in cryptic species (e.g., Bickford et al., 2007; Kaliontzopoulou et al., 2011; Ghielmi et al., 2016) or taxa displaying convergent evolution (e.g., Harmon et al., 2005).

The taxonomy of Eurasian vipers has been traditionally based on differences in pholidotic (head and body scalation) and colouration (dorsal pattern and colour) traits. Our data compilation shows that the ranges of variation of most pholidotic traits overlap extensively among species of the same genus (figs 4, 5), while colouration traits exhibit high variability at both the inter- and intraspecific levels (supplementary table S5). This apparent low prevalence of diagnostic traits for taxonomic purposes must be taken with caution given the limitations of our data compilation. While some species descriptions relied on few specimens from a small,

not representative region and/or particular diagnostic traits (e.g., *M. albizona* Nilson, Andrén and Flärdh, 1990; *M. kuhrangica* Rajabizadeh, Nilson and Kami, 2011), robust studies using multivariate comparative analyses of morphological traits have shown morphological distinctiveness of some species (e.g., *V. graeca*, Nilson and Andrén, 2001; *M. razii*, Moradi, Rastegar-Pouyani and Rastegar-Pouyani, 2014), reinforcing the view that the combination of traits can in fact identify taxonomic units in some cases. On the other hand, the occurrence of morphologically-cryptic species, that have only been identified after molecular phylogenetic analyses (e.g., *V. walser* Ghielmi et al., 2016), suggests that other factors may be affecting the external morphological variability of Eurasian vipers.

Both pholidotic and colouration traits frequently display geographic variation associated with environmental gradients, reflecting adaptive processes (e.g., Shine, 2000; Sanders, Malhotra and Thorpe, 2004; Martínez-Freiría et al., 2009; Tomović, Crnobrnja-Isailović and Brito, 2010; Martínez-Freiría and Brito, 2013). The role of local adaptation in shaping intraspecific morphological differentiation has been highlighted in multiple studies on reptile species (e.g., Thorpe and Baez, 1993; Malhotra and Thorpe, 1997; Kaliontzopoulou, Pinho and Martínez-Freiría, 2018). In particular, traits related to fitness frequently present variation across different environmental and ecological conditions in order to meet the species-specific needs and enhance performance and fitness (Arnold, 1983; Kingsolver and Huey, 2003). In vipers, for instance, differences in dorsal pattern colouration can be an adaptive response to temperature gradients, enhancing thermoregulation capabilities, or to predation pressures, leading to aposematic signals or increased substrate-crypsis (Wüster et al., 2004; Valkonen et al., 2011; Santos et al., 2014; Dubey et al., 2015;

Martínez-Freiría et al., 2017). Increasing or decreasing scale numbers within species can influence water loss along environmental gradients (Malhotra and Thorpe, 1997; Sanders, Malhotra and Thorpe, 2004) or enhance locomotion over distinct substrates (Kelley, Arnold and Gladstone, 1997). These traits can also be plastic and depend on the thermal conditions experienced by the embryos during gestation. For example, both field and experimental studies indicate that thermal conditions at early embryonic stages influence the number of ventral scales, scale abnormalities, as well as dorsal colouration in vipers (Lourdais et al., 2004; Lorigoux et al., 2013). Additionally, reproductive programs in low-effective-size populations have shown that inbreeding can lead to subsequent decrease of scale counts in viper offspring (Üveges et al., 2012). Altogether, these studies suggest that morphological traits might exhibit high inter-population variability and little taxon-specific variation and thus, should no longer be used as the only source of data to inform taxonomic decisions in Eurasian vipers.

Taxonomic inflation and need for an integrative taxonomy

The taxonomy of Eurasian vipers has long been under intense debate, especially as it provides crucial underpinnings to the formulation of conservation management strategies and the allocation of economic resources for this purpose. Species have been described based on diverse criteria, and mostly using morphology and/or geographic isolation as the only source of inference. Not surprisingly, recent phylogenetic studies have often revealed major inconsistencies in relation to current taxonomic units (e.g., Ferchaud et al., 2012; Velo-Antón et al., 2012; Zinenko et al., 2015, 2016; Stümpel et al., 2016; Martínez-Freiría et al., 2017), also unveiling the existence of morphologically cryptic taxa (e.g., Ghielmi et al., 2016). Yet, many

recent species descriptions maintained the traditional methods of species delimitation, disregarding known limitations (e.g., *Vipera altaica* Tuniyev, Nilson and Andrén, 2010; *Montivipera kuhrangica* Rajabizadeh, Nilson and Kami, 2011; *Vipera olguni* Tuniyev et al., 2012; *Vipera shemakhensis* Tuniyev et al., 2013). In agreement with previous molecular studies, our phylogenetic reconstruction shows a clear mismatch between relevant evolutionary units and recognized species. This is particularly evident within *Vipera*, in which species complexes such as *V. latastei-monticola* or *V. ammodytes-transcaucasiana* include highly divergent lineages from the Miocene, while other species are much younger (from late Pleistocene), polyphyletic (e.g., *V. lotievi*) or are nested within others (e.g., *V. altaica*, *V. shemakhensis*). This puzzling scenario mainly results from the indiscriminate use of morphological traits and geographic isolation as exclusive criteria for species delimitation (see table 1). Polymorphism as a result of local adaptation and plasticity may lead to taxonomic inflation, whereas low morphological variability can hamper the identification of cryptic species, such as in the case of the *V. latastei-monticola* and *V. ammodytes-transcaucasiana* complexes. One example that illustrates well the problem of high morphological variation within species is the case of *V. lotievi*, for which morphological convergence across similar environments and confusion over species identification are highlighted as a possible explanation for its polyphyly (Zinenko et al., 2015). Similarly, the occurrence of ecotypes can lead to the designation of taxonomic units which are not concordant with evolutionary history (e.g., *V. aspis atra*, Ursenbacher et al., 2006b; *V. aspis montecristi*, Barbanera et al., 2009; Luiselli et al., 2015; *V. monticola*, Velo-Antón et al., 2012). Additionally, genetic introgression was also shown to be a confounding factor on species classification within the *Pelias* subgenus, leading to intermediate phenotypes in admixed populations (e.g., the description of *V. magnifica* and *V. orlovi*; Zinenko et al., 2016).

Due to these limitations, an increasing number of studies have applied molecular methods to inform taxonomic decisions. The amount of sequence divergence among groups has been extensively used as a criterion to delimit taxa (e.g., in small mammals, Bradley and Baker, 2001; in rat snakes, Hofmann et al., 2018). Although the use of a standard percentage of sequence divergence to separate species is debatable, in this work, we draw an arbitrary but conservative (in terms of recognizing most described species) threshold for the delimitation of evolutionary units in Eurasian vipers corresponding to an uncorrected genetic distance equal or higher to 5% for a 196 bp *cyt b* fragment (fig. 1; supplementary table S3). This strategy highlights the distinct levels of divergence for the currently recognized species, allowing the formulation of recommendations for specific status of already described species (table 1). Furthermore, our barcoding approach, using a short *cyt b* fragment that fully recovers all phylogenetic units (supplementary fig. S2), may be a useful tool to assess the distinctiveness of newly discovered populations that are suspected to represent new taxa (Hebert and Gregory, 2005).

Ultimately, the taxonomic status of candidate species should be best addressed in an integrative fashion, that is, by searching for concordant differences in genetic, morphological and ecological traits (Padial et al., 2010). Since mtDNA could be particularly misleading in vipers, both by inflating the number of taxa or by missing lineages that have lost their mtDNA due to introgression (e.g., Zinenko et al., 2016), we encourage the use of multilocus genetic data (e.g., UCE loci, Blair et al., 2019). Until now, among all Eurasian vipers, only three species have been described using integrative approaches (i.e., by addressing phylogenetic divergence and characterizing phenotypic variability; *Vipera walser* Ghielmi et al., 2016; *Macrovipera razii* Oraie et al., 2018; *Vipera sakoi* Tuniyev et al., 2018). In the absence of more integrative studies, which

are sometimes constrained by low financial support and by the scarcity or remoteness of populations (e.g., Göçmen et al., 2014a, b, 2017; Freitas et al., 2018), the systematic situation of the group will likely remain unresolved in the immediate future.

Species classification is not the sole controversial issue in Eurasian viper taxonomy; allocation below (i.e., subspecies) or above (i.e., genera) species level is even more problematic due to the lack of objective, operational concepts defining these ranks. While our phylogenetic inferences are directed towards assessing the validity of species as independently evolving lineages, some provisional consensus can be reached at higher hierarchical levels. For instance, if *Pelias* is used to refer to Euro-Siberian *Vipera* species (as proposed in several studies, see Tuniyev et al., 2009, 2012, 2013, 2018a, b; Avcı et al., 2010), the designation of an additional genus within the current *Vipera* and the separation of *Daboia* into two or three genera would be required to reflect equivalent phylogenetic distances (see fig. 1; supplementary table S3). However, in order to avoid ill-founded splitting procedures and even further confusion, we advise against taking such steps (see also Vences et al., 2013), and argue that the use of subgenera may be a better way of providing names for clades without disrupting the binomial nomenclature (Wallach, Wüster and Broadley, 2009).

Concluding remarks

In this work, we integrate currently available information on the phylogenetic and morphological variability of Eurasian vipers to advance into a more coherent and objective taxonomy for this group. Based on our integrative assessment, we provide recommendations on the specific status of 40 described species and propose some guidelines to clarify the taxonomic status of some of them (table 1). Species complexes such as *V. latastei-monticola* or *V. ammodytes-transcaucasiana* require further analyses on the

extent of gene flow among distinct lineages to delineate taxonomic units, while some of the currently recognized species, described using morphological data only (e.g., *V. altaica*, *V. magnifica*, *V. shemakhensis*), must be explicitly regarded as non-valid species due to low genetic differentiation in relation to other previously recognized species. Again, other taxa, such as eastern *M. xanthina* and *V. sakoi* require more extensive geographic sampling to arrive at robust conclusions. As discussed above, integrative taxonomic approaches bringing together independent evidence and using different methodological approaches, particularly incorporating genomic data instead of relying solely on mitochondrial data, will allow the robust delineation of coherent evolutionary units in a unified taxonomic framework.

Previous attempts to propose priorities for the conservation of vipers (i.e., Maritz et al., 2016) suffered from taxonomic inflation and lack of a geographic comprehensive sampling scheme (see table 1). It is striking that more than half of the Eurasian viper species listed as globally endangered in our assessment (nine of 16 species with categories CR, EN and VU; table 1) were classed as of doubtful validity as species. Advancing in a robust, evidence-based designation of taxonomic units is therefore essential for the future development of conservation strategies aimed to anticipate threats related to anthropogenic factors, while slowing down or even stopping the rapid decline of many populations of Eurasian vipers.

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References

- Afaser, M., Yakin, B.Y., Çiçek, K., Dinçer, A. (2019): A new subspecies of Ottoman viper, *Montivipera xanthina* (Gray, 1849), (Squamata: Viperidae) from Geyik Mountains, Mediterranean Turkey. *Ecologica Montenegrina* **22**: 214-225.
- Alencar, L.R., Quental, T.B., Graziotin, F.G., Alfaro, M.L., Martins, M., Venzon, M., Zaher, H. (2016): Diversification in vipers: phylogenetic relationships, time of divergence and shifts in speciation rates. *Mol. Phylogenetics Evol.* **105**: 50-62.
- Alhajer, B.H., Hunt, O.J., Steppan, S.J. (2015): Molecular systematics of gerbils and deomyines (Rodentia: Gerbillinae, Deomyiinae) and a test of desert adaptation in the tympanic bulla. *J. Zool. Syst.* **53**: 312-330.
- Arnold, S.J. (1983): Morphology, performance and fitness. *Am. Zool.* **23**: 347-361.
- Avcı, A., Ilgaz, Ç., Başkaya, Ş., Baran, I., Kumlutaş, Y. (2010): Contribution to the distribution and morphology of *Peliás darevskii* (Vedmederja, Orlov and Tuniyev 1986) (Reptilia: Squamata: Viperidae) in northeastern Anatolia. *Russ. J. Herpetol.* **17**: 1-7.
- Ballard, J.W., Whitlock, M.C. (2004): The incomplete natural history of mitochondria. *Mol. Ecol.* **13**: 729-744.
- Baran, I., Joger, U., Kutrup, B., Türkozan, O. (2001): On new specimens of *Vipera barani* Böhm and Joger, 1983, from northeastern Anatolia, and implications for the validity of *Vipera pontica* Billing, Nilsson and Sattler, 1990 (Reptilia, Viperidae). *Zool. Middle East* **23**: 47-53.

- Barbanera, F., Zuffi, M.A., Guerrini, M., Gentili, A., To-fanelli, S., Fasola, M., Dini, F. (2009): Molecular phylogeography of the asp viper *Vipera aspis* (Linnaeus, 1758) in Italy: evidence for introgressive hybridization and mitochondrial DNA capture. *Mol. Phylogenetics Evol.* **52**: 103-114.
- Bickford, D., Lohman, D.J., Sodhi, N.S., Rahayu, D.L., Meier, R., Winker, K., Das, I. (2007): Cryptic species as a window on diversity and conservation. *Trends Ecol. Evol.* **22**: 148-155.
- Billing, H., Nilson, G., Satler, U. (1990): *Vipera pontica* sp. n., a new viper species in the kaznakovi group (Reptilia, Viperidae) from northeastern Turkey and adjacent Transcaucasia. *Zoologica Scripta*. **19**: 227-231.
- Blair, C., Bryson Jr, R.W., Linkem, C.W., Lazcano, D., Klicka, J., McCormack, J.E. (2019): Cryptic diversity in the Mexican highlands: thousands of UCE loci help illuminate phylogenetic relationships, species limits and divergence times of montane rattlesnakes (Viperidae: Crotalus). *Mol Ecol. Resour.* **19**: 349-365.
- Boettger, O. (1890): Eine neue Viper aus Armenien. *Zool. Anz.* **13**: 62-64.
- Böhme, W., Joger, U. (1984): Eine neue Art des *Vipera berus*-Komplexes aus der Türkei. *Amphibia-Reptilia*. **4**: 265-271.
- Bonaparte, L. (1835): *Iconographia della Fauna Italica per le quattro classi degli animali vertebrati*. Tome. Rome.: 1832-1841.
- Boscá, E. (1878): Note sur une forme nouvelle ou peu connue de vipère. *Bull. Zool. Soc. France* **3**: 116-121.
- Boulenger, G.A. (1896): *Catalogue of the Snakes in the British Museum Natural History*, vol. III. British Museum (Natural History), London.
- Boulenger, G.A. (1913): On the geographical races of *Vipera ammodytes*. *Ann. Mag. Nat. Hist.* **11**: 283-287.
- Bradley, R.D., Baker, R.J. (2001): A test of the genetic species concept: cytochrome-b sequences and mammals. *J. Mammal.* **82**: 960-973.
- Christoph, H. (1861): *Pelias renardi* mihi. *Bull. Soc. Nat. Moscow* **34**: 599-606.
- Clement, M., Posada, D., Crandall, K.A. (2000): TCS: a computer program to estimate gene genealogies. *Mol. Ecol.* **9**: 1657-1659.
- Dayrat, B. (2005): Towards integrative taxonomy. *Biol J Linnean Soc.* **85**: 407-417.
- De Queiroz, K. (2007): Species concepts and species delimitation. *Syst. Biol.* **56**: 879-886.
- dos Santos, A.M., Cabezas, M.P., Tavares, A.I., Xavier, R., Branco, M. (2015): tcsBU: a tool to extend TCS network layout and visualization. *Bioinformatics* **32**: 627-628.
- Drummond, A.J., Ho, S.Y., Phillips, M.J., Rambaut, A. (2006): Relaxed phylogenetics and dating with confidence. *PLoS Biol.* **4**: 699.
- Drummond, A.J., Suchard, M.A., Xie, D., Rambaut, A. (2012): Bayesian phylogenetics with BEAUti and the BEAST 1.7. *Mol. Biol. Evol.* **29**: 1969-1973.
- Dubey, S., Zwahlen, V., Mebert, K., Monney, J.-C., Golay, P., Ott, T., Durand, T., Thiery, G., Kaiser, L., Geser, S., Ursenbacher, S. (2015): Diversifying selection and color-biased dispersal in the asp viper. *BMC Evol Biol.* **15**: 99.
- Eiselt, J., Baran, I. (1970): Ergebnisse zoologischer Sammelreisen in der Türkei: Viperidae. *Ann Nat Hist Mus Wien* **74**: 357-369.
- Ferchaud, A.-L., Ursenbacher, S., Luiselli, L., Jelić, D., Halpern, B., Major, A., Kotenko, T., Crnobrnja-Isailovic, J., Tomović, L., Ghira, I., Ioannidis, Y., Arnal, V., Montgelard, C. (2012): Phylogeography of the *Vipera ursinii* complex (Viperidae): mitochondrial markers revealed an east-west disjunction in the palearctic region. *J. Biogeogr.* **39**: 1836-1847.
- Freitas, I., Fahd, S., Velo-Antón, G., Martínez-Freiría, F. (2018): Chasing the phantom: biogeography and conservation of *Vipera latastei-monticola* in the Maghreb (North Africa). *Amphibia-Reptilia*. **39**: 145-161.
- Frost, D.R., Hillis, D.M. (1990): Species in concept and practice: herpetological applications. *Herpetologica*. **46**: 87-104.
- Garrigues, T., Dauga, C., Ferquel, E., Choumet, V., Failloux, A.B. (2005): Molecular phylogeny of *Vipera* Laurenti, 1768 and the related genera *Macrovipera* (Reuss, 1927) and *Daboia* (Gray, 1842), with comments about neurotoxic *Vipera aspis aspis* populations. *Mol. Phylogenetics Evol.* **35**: 35-47.
- Ghielmi, S., Menegon, M., Marsden, S.J., Laddaga, L., Ursenbacher, S. (2016): A new vertebrate for Europe: the discovery of a range-restricted relict viper in the western Italian Alps. *J. Zool. Syst.* **54**: 161-173.
- Göçmen, B., Mebert, K., İgci, N., Akman, B., Yıldız, M.Z., Oğuz, M.A., Altın, Ç. (2014a): New locality records for four rare species of vipers (Reptilia: Viperidae) in Turkey. *Zool. Middle East*. **60**: 306-313.
- Göçmen, B., Mebert, K., Karış, M., Oğuz, M.A., Ursenbacher, S. (2017): A new population and subspecies of the critically endangered Anatolian meadow viper *Vipera anatolica* Eiselt and Baran, 1970 in eastern Antalya province. *Amphibia-Reptilia*. **38**: 289-305.
- Göçmen, B., Mulder, J., Karış, M., Oğuz, M.A. (2014b): The poorly known Anatolian Meadow Viper, *Vipera anatolica*: new morphological and ecological data. *Herpetologica Romanica*. **8**: 1-10.
- Gray, J.E. (1849): *Catalogue of the specimens of snakes in the collection of the British Museum*. Edward Newman, London **15**: 1-125.
- Guiller, G., Lourdais, O., Ursenbacher, S. (2017): Hybridization between a euro-Siberian (*Vipera berus*) and a para-Mediterranean viper (*V. aspis*) at their contact zone in western France. *J Zool.* **302**: 138-147.
- Harmon, L.J., Kolbe, J.J., Cheverud, J.M., Losos, J.B. (2005): Convergence and the multidimensional niche. *Evolution*. **59**: 409-421.
- Hebert, P.D., Gregory, T.R. (2005): The promise of DNA barcoding for taxonomy. *Syst. Biol.* **54**: 852-859.
- Heinicke, M.P., Lemmon, A.R., Lemmon, E.M., McGrath, K., Hedges, S.B. (2018): Phylogenomic support for evolutionary relationships of new world direct-developing frogs (Anura: Terraranae). *Mol. Phylogenetics Evol.* **118**: 145-155.
- Herrmann, H.W., Joger, U., Nilson, G. (1992): Phylogeny and systematics of viperine snakes. III: resurrection of the genus *Macrovipera* (Reuss, 1927) as suggested by biochemical evidence. *Amphibia-Reptilia*. **13**: 375-392.

- Hillis, D.M. (2019): Species delimitation in herpetology. *J. Herpetol.* **53**: 3-12.
- Hofmann, S., Mebert, K., Schulz, K.D., Helfenberger, N., Göçmen, B., Böhme, W. (2018): A new subspecies of *Zamenis hohenackeri* (Strauch, 1873) (Serpentes: Colubridae) based on morphological and molecular data. *Zootaxa*. **4471**: 137-153.
- Joger, U. (2005): *Montivipera* Nilson, Tuniyev, Andrén, Orlov, Joger und Herrmann, 1999. In: Handbuch der Reptilien und Amphibien Europas, Schlangen (Serpentes), p. 61-62. Joger, U., Stümpel, N., Eds, Aula-Verlag, Wiesbaden.
- Kaiser, H., Crother, B.I., Kelly, C.M., Luiselli, L., O'Shea, M., Ota, H., Passos, P., Schleip, W.D., Wüster, W. (2013): Best practices: in the 21st century, taxonomic decisions in herpetology are acceptable only when supported by a body of evidence and published via peer-review. *Herpetol. Rev.* **44**: 8-23.
- Kaliontzopoulou, A., Carretero, M.A., Llorente, G.A. (2012): Morphology of the *Podarcis* wall lizards (Squamata: *Lacertidae*) from the Iberian Peninsula and north Africa: patterns of variation in a putative cryptic species complex. *Zool. J. Linn. Soc.* **164**: 173-193.
- Kaliontzopoulou, A., Pinho, C., Harris, D.J., Carretero, M.A. (2011): When cryptic diversity blurs the picture: a cautionary tale from Iberian and north African *Podarcis* wall lizards. *Biol. J. Linn. Soc.* **103**: 779-800.
- Kaliontzopoulou, A., Pinho, C., Martínez-Freiría, F. (2018): Where does diversity come from? Linking geographical patterns of morphological, genetic, and environmental variation in wall lizards. *BMC Evol. Biol.* **18**: 124.
- Kearse, M., Moir, R., Wilson, A., Stones-Havas, S., Cheung, M., Sturrock, S., Buxton, S., Cooper, A., Markowitz, S., Duran, C., Thierer, T., Ashton, B., Meintjes, P., Drummond, A. (2012): Geneious basic: an integrated and extendable desktop software platform for the organization and analysis of sequence data. *Bioinformatics*. **28**: 1647-1649.
- Kelley, K.C., Arnold, S.J., Gladstone, J. (1997): The effects of substrate and vertebral number on locomotion in the garter snake *Thamnophis elegans*. *Funct. Ecol.* **11**: 189-198.
- Kindler, C., Fritz, U. (2018): Phylogeography and taxonomy of the barred grass snake (*Natrix helvetica*), with a discussion of the subspecies category in zoology. *Vertebr. Zool.* **68**: 269-281.
- Kindler, C., Chèvre, M., Ursenbacher, S., Böhme, W., Hille, A., Jablonski, D., Vamberger, M., Fritz, U. (2017): Hybridization patterns in two contact zones of grass snakes reveal a new central European snake species. *Sci. Rep.* **7**: 1-12.
- Kingsolver, J.G., Huey, R.B. (2003): Introduction: the evolution of morphology, performance and fitness. *Integr. Comp. Biol.* **43**: 361-366.
- Klemmer, K. (1963): Liste der rezenten Giftschlangen. In: Die Giftschlangen der Erde, Behringwerke, Marburg, p. 255-464. Elwert, N.G., Ed.
- Krecksák, L. (2007): An account of the generic and specific names, and type specimens of viperid taxa described by Albert Franz Theodor Reuss (Squamata: Viperidae). *Zootaxa*. **1514**: 1-36.
- Lanfear, R., Calcott, B., Ho, S.Y., Guindon, S. (2012): PartitionFinder: combined selection of partitioning schemes and substitution models for phylogenetic analyses. *Mol. Biol. Evol.* **29**: 1695-1701.
- Lataste, F. (1879): Diagnose d'une vipère nouvelle d'Espagne. *Bull. Soc. zool. Fr.* **4**: 132.
- Lenk, P., Kalyabina, S., Wink, M., Joger, U. (2001): Evolutionary relationships among the true vipers (Reptilia: *Viperidae*) inferred from mitochondrial DNA sequences. *Mol. Phylogenetics Evol.* **19**: 94-104.
- Librado, P., Rozas, J. (2009): DnaSP v5: a software for comprehensive analysis of DNA polymorphism data. *Bioinformatics*. **25**: 1451-1452.
- Linnaeus, C. (1758): *Systema naturae per regna tria naturae, secundum classes, ordines, genera, species, cum characteribus, differentiis, synonymis, locis*. Tomus I. Editio decima, reformata. Laurentii Salvii, Holmiae. 10th Edition: 824.
- Lorioux, S., Vaugoyeau, M., DeNardo, D.F., Clobert, J., Guillon, M., Lourdaïs, O. (2013): Stage dependence of phenotypical and phenological maternal effects: insight into squamate reptile reproductive strategies. *Am. Nat.* **182**: 223-233.
- Lourdaïs, O., Shine, R., Bonnet, X., Guillon, M., Naulleau, G. (2004): Climate affects embryonic development in a viviparous snake, *Vipera aspis*. *Oikos*. **104**: 551-560.
- Luiselli, L., Petrozzi, F., Mebert, K., Zuffi, M.A.L., Amori, G. (2015): Resource partitioning and dwarfism patterns between sympatric snakes in a micro-insular Mediterranean environment. *Ecol. Res.* **30**: 527-535.
- Malhotra, A., Thorpe, R.S. (1997): Microgeographic variation in scalation of *Anolis oculatus* (Dominica, West Indies): a multivariate analysis. *Herpetologica*. **53**: 49-62.
- Mallow, D., Ludwig, D., Nilson, G. (2003): *True Vipers: Natural History and Toxinology of Old World Vipers*. Krieger Publishing Company.
- Maritz, B., Penner, J., Martins, M., Crnobrnja-Isailović, J., Spear, S., Alencar, L.R., Sigala-Rodriguez, J., Messenger, K., Clark, R.W., Soorae, P., Luiselli, L. (2016): Identifying global priorities for the conservation of vipers. *Biol. Cons.* **204**: 94-102.
- Martínez-Freiría, F., Brito, J.C. (2013): Integrating classical and spatial multivariate analyses for assessing morphological variability in the endemic Iberian viper *Vipera seoanei*. *J. Zool. Syst. Evol.* **51**: 122-131.
- Martínez-Freiría, F., Crochet, P.A., Fahd, S., Geniez, P., Brito, J.C., Velo-Antón, G. (2017a): Integrative phylogeographical and ecological analysis reveals multiple Pleistocene refugia for Mediterranean *Daboia* vipers in north-west Africa. *Biol. J. Linn. Soc.* **122**: 366-384.
- Martínez-Freiría, F., Freitas, I., Zuffi, M., Golay, P., Ursenbacher, S., Velo-Antón, G. (2020): Climatic refugia boosted allopatric diversification in western Mediterranean vipers. *J. Biogeogr.* DOI:10.1111/jbi.13861.
- Martínez-Freiría, F., de Lanuza, G.P., Pimenta, A.A., Pinto, T., Santos, X. (2017b): Aposematism and crypsis are not enough to explain dorsal polymorphism in the Iberian adder. *Acta Oecol.* **85**: 165-173.

- Martínez-Freiría, F., Santos, X., Pleguezuelos, J.M., Lizana, M., Brito, J.C. (2009): Geographical patterns of morphological variation and environmental correlates in contact zones: a multi-scale approach using two Mediterranean vipers (Serpentes). *J. Zool. Syst. Evol.* **47**: 357-367.
- Mayden, R.L. (1997): A hierarchy of species concepts: the denouement in the saga of the species problem. In: *Species: the Units of Diversity*, p. 381-423. Claridge, M.F., Dawah, H.A., Wilson, M.R., Eds, Chapman & Hall.
- Mebert, K. (2008): Good species despite massive hybridization: genetic research on the contact zone between the watersnakes *Nerodia sipedon* and *N. fasciata* in the Carolinas, USA. *Mol. Ecol.* **17**: 1918-1929.
- Mebert, K., Göçmen, B., Igci, N., Oguz, M.A., Karis, M., Ursenbacher, S. (2015a): New records and search for contact zones among parapatric vipers in the genus *Vipera* (*barani*, *kaznakovi*, *darevskii*, *erivanensis*), *Montivipera* (*wagneri*, *raddei*), and *Macrovipera* (*lebetina*) in northeastern Anatolia. *Herpetol. Bull.* **133**: 13-22.
- Mebert, K., Göçmen, B., Igci, N., Karis, M., Oguz, M.A., Yildiz, M.Z., Teynié, A., Stümpel, M., Ursenbacher, S. (2020): Mountain vipers in central-eastern Turkey: huge range extensions for four taxa reshape decades of misleading perspectives. *Herpetol. Conserv. Bio.* **15**.
- Mebert, K., Jagar, T., Grželj, R., Cafuta, V., Luiselli, L., Ostanek, E., Golay, P., Dubey, S., Golay, J., Ursenbacher, S. (2015): The dynamics of coexistence: habitat sharing vs. segregation patterns among three sympatric montane vipers. *Biol. J. Linn. Soc.* **116**: 364-376. DOI:10.1111/bij.12582.
- Merrem, B. (1820): Versuch eines Systems der Amphibien I (Tentamen Systematis Amphibiorum). J. C. Kriegeri, Marburg.
- Mertens, R., Darevsky, I.S., Klemmer, K. (1967): *Vipera latifii*, eine neue Giftschlange aus dem Iran. *Senckenb. Biol.* **48**: 161-168.
- Minton, S.A., Jr., Dowling, H.G., Russell, F.E. (1968): *Poisonous Snakes of the World. A Manual for Use by U.S. Amphibious Forces*, 2nd Edition. Department of the Navy and Bureau of Medical Surgery, Washington.
- Mizsei, E., Jablonski, D., Roussos, S.A., Dimaki, M., Ioannidis, Y., Nilson, G., Nagy, Z.T. (2017): Nuclear markers support the mitochondrial phylogeny of *Vipera ursinii-renardi* complex (Squamata: Viperidae) and species status for the Greek meadow viper. *Zootaxa*. **4227**: 75-88.
- Moradi, N., Rastegar-Pouyani, N., Rastegar-Pouyani, E. (2014): Geographic variation in the morphology of *Macrovipera lebetina* (Linnaeus, 1758) (Ophidia: Viperidae) in Iran. *Acta Herpetol.* **9**: 187-202.
- Nikolsky, A.M. (1909): Eine neue Vipern-Art aus dem Kaukasus: *Vipera kaznakovi* sp. nov. *Tiflis Mitt. Kaukas. Mus.* **4**: 174.
- Nikolsky, A.M. (1913): *Herpetologia caucasica*. Tiflis. 272.
- Nilson, G., Andren, C. (1984): Systematics of the *Vipera xanthina* complex (Reptilia: Viperidae). An overlooked viper within the *xanthina* species-group in Iran. *Bonn. zool. Beitr.* **35**: 175-184.
- Nilson, G., Andren, C. (1985): Systematics of the *Vipera xanthina* complex (Reptilia: Viperidae). Taxonomic status of the Bulgar Daghi viper in south Turkey. *J. Herpetol.* **19**: 276-283.
- Nilson, G., Andrén, C. (1988): *Vipera lebetina transmediterranea*, a new subspecies of viper from north Africa, with remarks on the taxonomy of *Vipera lebetina* and *Vipera mauritanica*. *Bonn. zool. Beitr.* **39**: 371-379.
- Nilson, G., Andrén, C. (1997): Evolution, systematics and biogeography of Palaearctic vipers. In: *Venomous Snakes: Ecology, Evolution and Snakebite*, p. 31-42. Thorpe, R.S., Wüster, W., Malhorta, A., Eds, Oxford University Press.
- Nilson, G., Andrén, C. (2001): The meadow and steppe vipers of Europe and Asia – the *Vipera* (*Acridophaga*) *ursinii* complex. *Acta zool. Acad. Sci. Hung.* **47**: 87-267.
- Nilson, G., Andrén, C., Flärdh, B. (1990): *Vipera albizona*, a new mountain viper from central Turkey, with comments on isolating effects of the Anatolian “Diagonal”. *Amphibia-Reptilia*. **11**: 285-294.
- Nilson, G., Tuniyev, B., Andrén, C., Orlov, N., Joger, U., Herrmann, H.-W. (1999): Taxonomic position of the *Vipera xanthina* complex. *Kaupia* **8**: 99-102.
- Nilson, G., Tuniyev, B.S., Orlov, N., Hoggren, M., Andren, C. (1995): Systematics of the vipers of the Caucasus: polymorphism or sibling species? *Asiat. Herpetol. Res.* **6**: 1-16.
- Obst, F.J. (1983): Zur Kenntnis der Schlangengattung *Vipera*. *Zool. Abh. Staatl. Mus. Tierk. Dresden* **38**: 229-335.
- Oraie, H., Rastegar-Pouyani, E., Khosravani, A., Moradi, N., Akbari, A., Sehhatiasabet, M.E., Shafiei, S., Stümpel, N., Joger, U. (2018): Molecular and morphological analyses have revealed a new species of blunt-nosed viper of the genus *Macrovipera* in Iran. *Salamandra*. **54**: 233-248.
- Orlov, N.L., Tuniyev, B.S. (1990): Three species in the *Vipera kaznakovi* complex (Eurosiberian Group) in the Caucasus: their present distribution, possible genesis and phylogeny. *Asiat. Herpetol. Res.* **3**: 1-36.
- Padial, J.M., Miralles, A., De la Riva, I., Vences, M. (2010): The integrative future of taxonomy. *Front. Zool.* **78**: 16.
- Paleobiology Database (2019): The Paleo Biology Database Navigator. Search on March 2019. Available at: <https://paleobiodb.org/navigator/>.
- Phelps, T. (2010): Old world vipers: a natural history of the Azemiopinae and Viperinae. Ed. Chimaira.
- Pöschel, J., Heltai, B., Graciá, E., Quintana, M.F., Velo-Antón, G., Arribas, O., Valdeón, A., Wink, M., Fritz, U., Vamberger, M. (2018): Complex hybridization patterns in European pond turtles (*Emys orbicularis*) in the Pyrenean region. *Sci. Rep.* **8**: 15925.
- Rajabzadeh, M., Nilson, G., Kami, H.G. (2011): A new species of mountain viper (Ophidia: Viperidae) from the central Zagros Mountains, Iran. *Russ. J. Herpetol.* **18**: 235-240.
- Rambaut, A., Drummond, A.J., Xie, D., Baele, G., Suchard, M.A. (2018): Posterior summarization in Bayesian phylogenetics using Tracer 1.7. *Syst. Biol.* **67**: 901-904.

- Reuss, T. (1933): Fortsetzung und Schluss der Originalberichte. Nachr. Bl. Aquar. Terrar.-Ver.: 372-373.
- Saint Girons, H. (1953): Une vipère naine: *Vipera latastei montana*. Bull. Soc. zool. Fr. **78**: 24-28.
- Sanders, K.L., Malhotra, A., Thorpe, R.S. (2004): Ecological diversification in a group of Indomalayan pitvipers (*Trimeresurus*): convergence in taxonomically important traits has implication for species identification. J. Evol. Biol. **17**: 721-731.
- Santos, X., Vidal-García, M., Brito, J.C., Fahd, S., Llorente, G.A., Martínez-Freiría, F., Parellada, X., Pleguezuelos, J.M., Sillero, N. (2014): Phylogeographic and environmental correlates support the cryptic function of the zigzag pattern in a European viper. Evol. Ecol. **28**: 611-626.
- Schwarz, E. (1936): Untersuchungen über Systematik und Verbreitung der europäischen und mediterranen Ottern. In: Die europäischen und mediterranen Ottern und ihre Gifte, p. 159-362. Behringwerke-Mitteilungen.
- Shaw, G., Nodder, F.P., Eds (1997): The naturalist's miscellany. London, Nodder & Co. **8**: 255-300.
- Shine, R. (2000): Vertebral numbers in male and female snakes: the roles of natural, sexual and fecundity selection. J. Evol. Biol. **13**: 455-465.
- Simpson, G.G. (1961): Principles of Animal Taxonomy. Columbia University Press, USA.
- Šmíd, J., Tolley, K.A. (2019): Calibrating the tree of vipers under the fossilized birth-death model. Sci. Rep. **9**: 5510.
- Stephens, M., Smith, N.J., Donnelly, P. (2001): A new statistical method for haplotype reconstruction from population data. Am. J. Hum. Genet. **68**: 978-989.
- Stümpel, N. (2012): Phylogenie und Phylogeographie eurasischer Viperinae unter besonderer Berücksichtigung der orientalischen Vipern der Gattungen *Montivipera* und *Macrovipera*. Unpubl. PhD Thesis, TU Carolo-Wilhelmina zu Braunschweig, Germany.
- Stümpel, N., Joger, U. (2009): Recent advances in phylogeny and taxonomy of near and middle eastern vipers – an update. ZooKeys. **31**: 179-191.
- Stümpel, N., Rajabizadeh, M., Avci, A., Wüster, W., Joger, U. (2016): Phylogeny and diversification of mountain vipers (*Montivipera*, Nilson et al., 2001) triggered by multiple Plio-Pleistocene refugia and high-mountain topography in the Near and Middle East. Mol. Phylogenetics Evol. **101**: 336-351.
- Tamura, K., Peterson, D., Peterson, N., Stecher, G., Nei, M., Kumar, S. (2011): MEGA5: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. Mol. Biol. Evol. **28**: 2731-2739.
- Taroso, P., Pereira, R.J., Martínez-Freiría, F., Godinho, R., Brito, J.C. (2014): Hybridization at an ecotone: ecological and genetic barriers between three Iberian vipers. Mol. Ecol. **23**: 1108-1123.
- Thorpe, R.S., Baez, M. (1993): Geographic variation in scalation of the lizard *Gallotia stehlini* within the island of Gran Canaria. Biol. J. Linn. Soc. **48**: 75-87.
- Tomović, L., Crnobrnja-Isailović, J., Brito, J.C. (2010): The use of geostatistics and GIS for evolutionary history studies: the case of the nose-horned viper (*Vipera ammodytes*) in the Balkan Peninsula. Biol. J. Linn. Soc. **101**: 651-666.
- Townsend, T.M., Alegre, R.E., Kelley, S.T., Wiens, J.J., Reeder, T.W. (2008): Rapid development of multiple nuclear loci for phylogenetic analysis using genomic resources: an example from squamate reptiles. Mol. Phylogenetics Evol. **47**: 129-142.
- Tuniyev, B.S., Ostrovskikh, S.V. (2001): Two new species of vipers of “*kaznakovi*” complex (Ophidia, Viperinae) from the Western Caucasus. Russ. J. Herpetol. **8**: 117-126.
- Tuniyev, B.S., Avci, A., Ilgaz, Ç., Olgun, K., Petrova, T.V., Bodrov, S.Y., Geniez, P., Teynié, A. (2018a): On taxonomic status of shield-head vipers from Turkish Lesser Caucasus and East Anatolia. Proceedings of the Zoological Institute RAS. **322**: 3-44.
- Tuniyev, B.S., Iremashvili, G.N., Petrova, T.V., Kravchenko, M.V. (2018b): Rediscovery of the steppe viper in Georgia. Proceedings of the Zoological Institute RAS. **322**: 87-107.
- Tuniyev, B.S., Nilson, G., Andrén, C. (2010): A new species of viper (Reptilia, Viperidae) from the Altay and Saur Mountains, Kazakhstan. Russ. J. Herpetol. **17**: 110-120.
- Tuniyev, S.B., Avci, A., Tuniyev, B.S., Agasian, A.L., Agasian, L.A. (2012): Description of a new species of shield-head vipers – *Pelias olguni* sp. nov. from Basin of Upper Flow of the Kura River in Turkey. Russ. J. Herpetol. **19**: 314-332.
- Tuniyev, S.B., Orlov, N.L., Tuniyev, B.S., Kidov, F.F. (2013): On the taxonomical status of steppe viper from foothills of the south macroslope of the east Caucasus. Russ. J. Herpetol. **20**: 129-146.
- Uetz, P., Freed, P., Hošek, J., Eds (2019): The Reptile Database, <http://www.reptile-database.org>, accessed on March 2019.
- Ursenbacher, S., Carlsson, M., Helfer, V., Tegelström, H., Fumagalli, L. (2006a): Phylogeography and Pleistocene refugia of the adder (*Vipera berus*) as inferred from mitochondrial DNA sequence data. Mol. Ecol. **15**: 3425-3437.
- Ursenbacher, S., Conelli, A., Golay, P., Monney, J.-C., Zuffi, M.A.L., Thiery, G., Durand, T., Fumagalli, L. (2006b): Phylogeography of the asp viper (*Vipera aspis*) inferred from mitochondrial DNA sequence data: evidence for multiple Mediterranean refugial areas. Mol. Phylogenetics Evol. **38**: 546-552.
- Ursenbacher, S., Schweiger, S., Tomović, L., Crnobrnja-Isailović, J., Fumagalli, L., Mayer, W. (2008): Molecular phylogeography of the nose-horned viper (*Vipera ammodytes*, Linnaeus (1758)): evidence for high genetic diversity and multiple refugia in the Balkan Peninsula. Mol. Phylogenetics Evol. **46**: 1116-1128.
- Üveges, B., Halpern, B., Péchy, T., Posta, J., Komlósi, I. (2012): Characteristics and heritability analysis of head scales of the Hungarian meadow viper (*Vipera ursinii rakosiensis*, Méhely 1893). Amphibia-Reptilia. **33**: 393-400.
- Vaidya, G., Lohman, D.J., Meier, R. (2011): SequenceMatrix: concatenation software for the fast assembly of multi-gene datasets with character set and codon information. Cladistics. **27**: 171-180.

- Valkonen, J., Niskanen, M., Björklund, M., Mappes, J. (2011): Disruption or aposematism? Significance of dorsal zigzag pattern of European vipers. *Evol. Ecol.* **25**: 1047-1063.
- Vedmederjia, V.J., Orlov, N.L., Tuniyev, B.S. (1986): On taxonomy of three viper species of the *Vipera kaznakovi* complex. *Dokl. Akad. Nauk SSSR.* **157**: 55-61.
- Velo-Antón, G., Godinho, R., Harris, D.J., Santos, X., Martínez-Freiria, F., Fahd, S., Pleguezuelos, J.M., Brito, J.C. (2012): Deep evolutionary lineages in a western Mediterranean snake (*Vipera latastei* monticola group) and high genetic structuring in southern Iberian populations. *Mol. Phylogenetics Evol.* **65**: 965-973.
- Vences, M., Guayasamin, J.M., Miralles, A., de la Riva, I. (2013): To name or not to name: criteria to promote economy of change in Linnaean classification schemes. *Zootaxa.* **3636**: 201-244.
- Wallach, V., Wüster, W., Broadley, D.G. (2009): In praise of subgenera: taxonomic status of cobras of the genus *Naja* Laurenti (Serpentes: Elapidae). *Zootaxa.* **2236**: 26-36.
- Wan, Q.H., Wu, H., Fujihara, T., Fang, S.G. (2004): Which genetic marker for which conservation genetics issue?. *Electrophoresis* **25**: 2165-2176.
- Werner, F. (1898): Über einige neue Reptilien und einen neuen Frosch aus dem cilicischen Taurus. *Zool. Anz.* **21**: 217-223.
- Werner, F. (1935): Reptilien der Ägäischen Inseln. *Sitzungsber. Akad. Wiss. Wien.* **144**: 81-117.
- Werner, F. (1938): Eine verkannte Viper (*Vipera palaestinae* n. sp.). *Zool. Anz.* **122**: 313-318.
- Wiley, E.O. (1978): The evolutionary species concept reconsidered. *Systematic Zoology* **21**: 17-26.
- Wüster, W., Allum, C.S.E., Bjargardóttir, I.B., Bailey, K.L., Dawson, K.J., Guenioui, J., Lewis, J., Mcgurk, J., Moore, A.G., Niskanen, M., Pollard, C.P. (2004): Do aposematism and Batesian mimicry require bright colours? A test, using European viper markings. *Proc. Royal Soc. B.* **271**: 2495-2499.
- Zheng, Y., Wiens, J.J. (2016): Combining phylogenomic and supermatrix approaches, and a time-calibrated phylogeny for squamate reptiles (lizards and snakes) based on 52 genes and 4162 species. *Mol. Phylogenetics Evol.* **94**: 537-547.
- Zinenko, O., Sovic, M., Joger, U., Gibbs, H.L. (2016): Hybrid origin of European Vipers (*Vipera magnifica* and *Vipera orlovi*) from the Caucasus determined using genomic scale DNA markers. *BMC Evol Biol.* **16**: 76.
- Zinenko, O., Stümpel, N., Mazanaeva, L., Bakiev, A., Shiryayev, K., Pavlov, A., Kotenko, T., Kukushkin, O., Chikin, Y., Duisebayeva, T., Nilson, G. (2015): Mitochondrial phylogeny shows multiple independent ecological transitions and northern dispersion despite of Pleistocene glaciations in meadow and steppe vipers (*Vipera ursinii* and *Vipera renardi*). *Mol. Phylogenetics Evol.* **84**: 85-100.
- Zuffi, M. (2002): A critique of the systematic position of the asp viper subspecies *Vipera aspis aspis* (Linnaeus, 1758), *Vipera aspis atra* Meisner, 1820, *Vipera aspis francisciredi* Laurenti, 1768, *Vipera aspis hugyi* Schinz, 1833 and *Vipera aspis zinnikeri* Kramer, 1958. *Amphibia-Reptilia.* **23**: 191-213.

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