

Molecular Approach to the Identification of Three Closely Related Slugs *Deroceras* Species (Gastropoda: Eupulmonata: Agriolimacidae)

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Some species of slugs belonging to the genus *Deroceras* are invasive and cause severe agricultural damage. Despite extensive knowledge about their invasiveness, data on molecular differentiation of these morphologically similar species are lacking. Here we present a molecular approach to the identification of three closely related species of the genus *Deroceras* - *D. agreste* (L., 1758), *D. reticulatum* (O. F. Müller, 1774) and *D. turcicum* (Simroth, 1894) (Gastropoda: Eupulmonata: Agriolimacidae) based on sequences of multiple molecular markers: cytochrome *c* oxidase subunit I (*COI*), cytochrome *b* (*cyt-b*), internal transcribed spacer 2 (ITS-2) and 28S ribosomal RNA (28S rRNA). We also provide detailed photomicrographs of the penis and penial gland of the three species as it is the latter that holds the most important phenotypic characters for distinguishing between these taxa. Since identification of the studied species, based solely on morphology is considered challenging, contributing a means of molecular differentiation will aid further ecological and biodiversity surveys of these important pests.

Key words: Taxonomy, *Deroceras*, Barcoding, Gastropods, Slugs.

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BACKGROUND

Species identification is central to estimation of biodiversity. In the past, traditional taxonomy was based mainly on the morphology of the investigated organism. Since the recognition

of cryptic diversity (*i.e.*, complexes of morphologically similar species), which was outcome consequence of the application of molecular techniques to the problem of species identification, morphology-based delineation became insufficient to describe biological diversity (Dayrat 2005). The modern methods of estimating species richness such as DNA-barcoding, metabarcoding and DNA-based taxonomy are now in standard use including examples in meiofauna, ants, decapod crustaceans, lizards and millipedes (Fontaneto et al. 2015; Troncoso-Palacios et al. 2018; Hosoishi and Ogata 2019; Shih et al. 2019; Nguyen et al. 2019). However, all of these methods suffer from limited correspondence between the libraries of DNA sequences and the correctly identified species.

Among terrestrial gastropods, identification of many species of slug is challenging, as some taxa can only be distinguished by minute morphological differences in their reproductive organs, and the anatomical sections, necessary for proper identification, are often problematic for non-specialists.

Deroceras Rafinesque, 1820 is a type genus within the family Agriolimacidae and comprises more than 120 species (Wiktor 2000). Some of the *Deroceras* species are invasive and drastically affect biodiversity and agriculture through reduction of native species richness and crop infestation (Barker 2002). Among *Deroceras* species, *D. reticulatum* (O. F. Müller, 1774) is a major agricultural pest and is known to cause considerable damage to crops of winter rape, winter wheat and many others (Kozłowski 2010). The majority of studies are concerned with the invasiveness and significance of *D. reticulatum* in agriculture (Ferguson et al. 1988; Birkett et al. 2004; Berman et al. 2011), whereas the molecular data supporting its pattern of distribution are lacking. Additionally, sequences of two closely related and morphologically very similar species, *Deroceras turcicum* (Simroth, 1894) (*locus typicus* – Lake Ohrid in Macedonia) and *Deroceras agreste* (L., 1758) (*terra typica* - Sweden) are unknown or uncertain, respectively, and what is more these species have overlapping distribution ranges. The *COI* sequences of *D. agreste* are known from a single study about gastropods of Britain and Ireland (Rowson et al. 2014). In that study, the authors found individuals with the genital morphology characteristic of *D. reticulatum* but mtDNA sequences characteristic of *D. agreste*. These results were explained as an example of a one-way introgression of *D. reticulatum* genes into a *D. agreste* population (Rowson et al. 2014). Notably, the population of *D. agreste* used in the study was not from the species' *terra typica*, thus the validity of the proposed conclusions remain open to question.

Deroceras reticulatum (O. F. Müller, 1774) is widely distributed synanthropic species occurring in Europe (Wiktor 2004). The origin of this species is unknown, but it likely comes from Central Europe (Kozłowski 2010). It was described from Frideriksdal in Denmark (*locus typicus*) in 1774. Recently, it was introduced to North and South America, Tasmania, New Zealand and Central

Asia (e.g. Crowley and Pain 1977; Hausdorf 2002; Tulli et al. 2009; Welter-Schultes 2012). Notably, the majority of these records have not been verified genetically. For instance, Hausdorf (2002) studied terrestrial gastropods (both slugs and snails) in Colombia and determined slug specimens anatomically. Crowley and Pain (1977) published a study about molluscs of Saint Helena in the last century and their identifications were based on classical taxonomic methods. Tulli et al. (2009) conducted research on predation on *D. reticulatum* by the carabid *Scarites athracinus* and collected slugs in Argentina. In this study there is no information about slug identification, thus it seems probable that species identification was based on morphological characters and/or external appearance. Finally, a book by Welter-Schultes (2012) summarizes data about the occurrence and distribution of particular species in Europe. *D. reticulatum* is regarded by the DAISIE organization (Delivering Alien Invasive Species Inventories for Europe) as one of the most invasive species in Europe. It mostly occurs in cultivated areas, open habitats, meadows, gardens, and cemeteries - in all places where human activity is observed (Wiktor 2004; Welter-Schultes 2012). *D. reticulatum* has a creamy or light coffee cream body pigmentation, rarely it is blackish and spotted, in which case the dark spots behind the mantle form a reticulate pattern. However, Wiktor (1989) noted considerable variability in the distribution and pigmentation of spots and stated that the most common *D. reticulatum* individuals are spotted, even though juveniles may be uniform in color. All this makes species identification based on the external appearance extremely difficult.

The second species examined in our study, *D. agreste*, was originally described from Sweden. This species likely occurs throughout Europe, but its distribution is not precisely known (Wiktor 2000; Welter-Schultes 2012). Currently there is no data relating to where *D. agreste* has been introduced and where it is indigenous. These slugs prefer open habitats, especially gardens, meadows, and less often farmland. It can be described as synanthropic and can be found together with, and in similar habitats to, *D. reticulatum* and *D. sturanyi* (Wiktor 2000). Its body is yellowish white to light greyish yellow, transparent, without spots (Welter-Schultes 2012).

Our third species, *D. turcicum*, occurs in the Balkan region (Wiktor 1989). It was originally described from Lake Ohrid in Macedonia in 1894. *D. turcicum* has expanded its distribution, and currently it is present in Austria, Hungary, Slovenia, Croatia, Bosnia and Herzegovina, Serbia, Montenegro and is probably present in Albania, Greece, as well as Turkey (Wiktor 2000). Reise and Hutchinson (2001) reported the presence of *D. turcicum* in the Czech Republic and Slovakia, whereas Reise et al. (2005) recorded new localities of this species in Poland. *D. turcicum* inhabits mostly mixed or beech woodland, parks, and gardens. Its body is dirty-cream or light cream, with dark spots covering the mantle; overall it is extremely similar in appearance to *D. reticulatum* (Welter-Schultes 2012). Most reports about the occurrence of these slugs are based on observations and/or species identification based on traditional taxonomic methods such as anatomical sections. In

the majority of reports, species identification was not supported by any molecular analysis, e.g. by comparisons of *COI* sequences (Szybiak 2004; Reise et al. 2005; Sulikowska-Drozd 2007; Hutchinson and Reise 2015). Thus, cited records should be treated with caution as the species under consideration are morphologically very similar.

In this study, we molecularly characterise and compare three closely related *Deroceras* species (*D. agreste*, *D. reticulatum*, *D. turcicum*) providing for each of them DNA sequences of four molecular markers (cytochrome *c* oxidase subunit I (*COI*), cytochrome *b* – (*cyt-b*), internal transcribed spacer 2 (ITS-2), 28S ribosomal RNA (28S rRNA)). We also provide comparative genetic analyses of p-distances to show differences between the studied species. Moreover, we visualise phylogenetic relationships between *Deroceras* species based on available *COI* sequences for the named species deposited in the GenBank database as well as the sequences obtained in this study. Finally, we also provide a set of photomicrographs that detail the morphology of the penis and penial gland in the studied species. Our contribution will be helpful to other researchers using modern approaches for the identification of the difficult to distinguish *Deroceras* species. Our findings may also have significance in biological conservation as these slugs have been shown to have considerable economic importance in agriculture.

MATERIALS AND METHODS

Material collection and species identification

All specimens of *Deroceras agreste*, *D. turcicum* and *D. reticulatum* were obtained from museum collection deposited in the Museum of Natural History, Wrocław University, Poland (Table 1). Gastropods were preserved in 70–75% ethyl alcohol (Wiktor and Jurkowska 2007). All specimens were identified and labelled by Professor Andrzej Wiktor (Wrocław University, Poland) and further verified by us based on anatomical characteristics under Nikon SMZ1500 stereomicroscope (Table 2). From each slug, a small piece of tissue, preserved in 99.8% ethanol, was taken for DNA extraction. The photos of penial glands were taken using a digital camera (Nikon DS-Fil-U2) associated with the stereomicroscope and later assembled into figures in Corel Photo-Paint X6, ver. 16.4.1.1281. Schematic drawings of the genitalia of each species were made using Inkscape software based on Wiktor's (2000) monograph.

Table 1. Collection data on *Deroceras* specimens used in this study (N – number of sequenced individuals from each locality)

Species	N	Locality	Collector	Collection date	Fig. 1.	GenBank Accession numbers			
						<i>COI</i>	<i>cyt-b</i>	ITS-2	28S rRNA
<i>D. reticulatum</i>	3	Aragón, Pirineos Mts, Diazas near Torla (N of Huesca), Spain	A. Wiktor	07.07.1987	A	MN934414	MN931235	MT361823	MN930516
	3	Leon province, Villaturel and Villasabariego (NE of Leon), Spain	A. Wiktor	19.06.1987	B, C				
<i>D. agreste</i>	2	Västergötland, Sweden	H. W. Walden	30.09.1964	D	MN934413	MN931234	MT361821	MN930515
	2	Bohuslän, Sweden	H. W. Walden	10.10.1965	E				
	2	Cantabria, Spain	A. Wiktor	12.11.1988	F				
<i>D. turcicum</i>	2	near Maçka in Trabzon, Turkey	A. Riedel	07.12.1985	G	MN934415	MN931236	MT361822	MN930517
	2	Zafanos, Trabzon, Turkey	A. Riedel	01.12.1985	H				
	2	near Maçka in Trabzon, Turkey	A. Wiktor	29.04.2001	I				

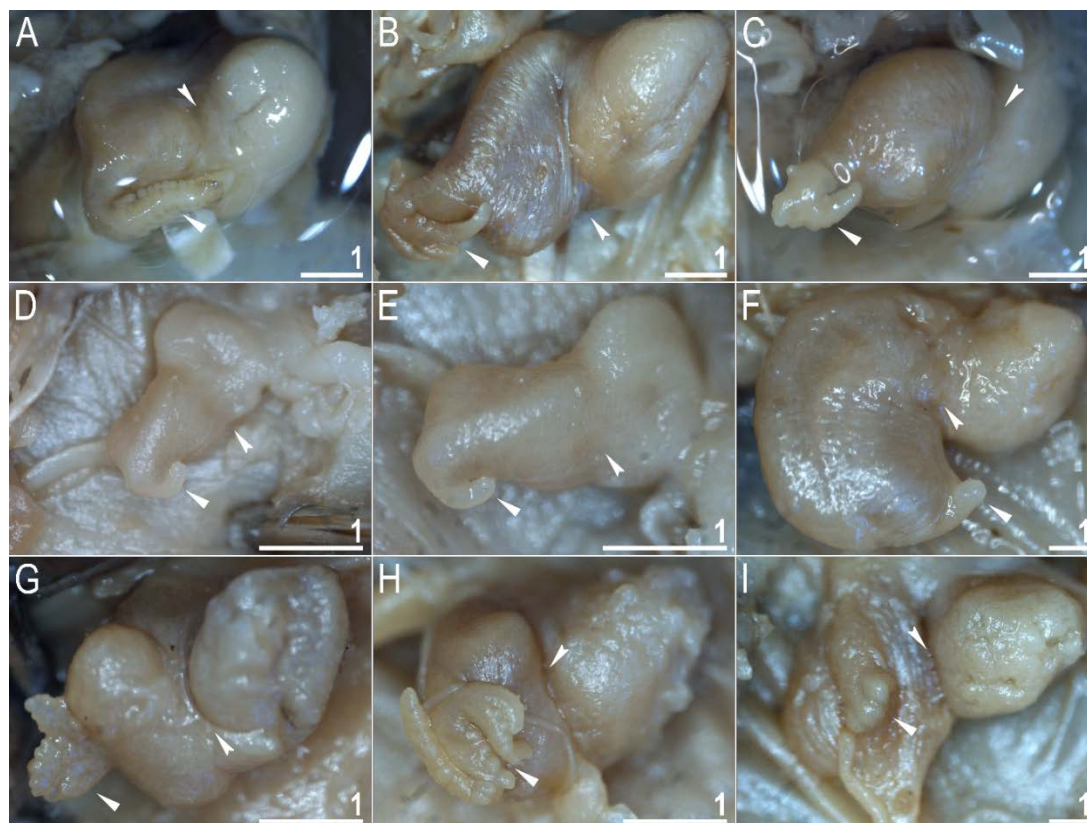


Fig. 1. Genitalia of *D. reticulatum* (A: Aragón, Pirineos Mts, Diazas near Torla (N of Huesca), Spain; B-C: Leon province, Villaturel and Villasabariego (NE of Leon), Spain), *D. agreste* (D:

Västergötland, Sweden; E: Bohuslän, Sweden; F: Cantabria, Spain) and *D. turcicum* (G: near Maçka in Trabzon, Turkey; H: Zafanos, Trabzon, Turkey; I: near Maçka in Trabzon, Turkey) detailing penis and penial gland morphology. Penial gland and constriction in the penis are indicated by flat and indented arrowheads respectively. Scale bars in mm.

Table 2. Morphological differences in reproductive system between *D. reticulatum*, *D. agreste* and *D. turcicum* according to Welter-Schultes (2012)

	<i>D. reticulatum</i>	<i>D. agreste</i>	<i>D. turcicum</i>
penis	fleshy and with a silky sheen, in the shape of an irregular sac, divided into 2 parts by a deep lateral constriction	oval in juveniles, with lateral constriction in adults	strong swelling on the anterior part of the penis
penial gland	variable shape; usually a few branches or a single long branch	finger-shaped, never branched, smooth, without glandular papillae	variable or reduced
stimulator	large, conical and narrow	cone-shaped, with narrow base	conical
rectal caecum	large	caecum on rectum well-developed, clearly longer than wider	well-developed
vas deferens	opens into the penis wall facing the external body side	opens laterally close to penial gland	opens at the base of the penial gland

DNA extraction, amplification and sequencing

DNA was extracted from the collected tissue using Sherlock AX kit (A&A Biotechnology, Poland) according to the manufacturer's protocol. This kit is intended for material with a trace of DNA content. To obtain DNA sequences of the *COI*, *ITS-2*, *28S rRNA*, *cyt-b* genes PCR reactions were run using the primers listed in table 3. For *COI*, *ITS-2* and *28S rRNA* specific primers for *Deroceras* species were designed *de novo* based on sequences of *Deroceras* (*D. laeve*, *D. reticulatum*, *D. invadens*), *Arion* (*Arion* sp., *A. silvaticus*, *A. vulgaris*, *A. distinctus*, *A. rufus*) and *Limax* (*L. maximus*, *L. flavus*, *L. cinereoniger*) species available in GenBank, respectively (for exact accession numbers see Table S1). A PCR cocktail and profiles/conditions for specific markers are given in supplementary materials (Tables S2 and S3). To check DNA quality 3 µl sample of PCR product was run on a 1.5% agarose gel for 30 min at 100 V. PCR products were cleaned by using NucleoSpin Gel and PCR Clean-up (Macherey-Nagel, Germany). A sequencing reaction was performed in 10 µl reaction mixture, consisting of 2 µl of PCR product, 0.15 µl of primer, 1 µl of sequencing buffer (BrilliantDye Terminator Sequencing Kit, Nimagen, The Netherlands), 5.85 µl of

ddH₂O and 1 µl of Terminator (BrilliantDye Terminator Sequencing Kit, Nimagen, The Netherlands). The sequencing program consisted of four steps: 1 min initial denaturation at 96°C, followed by 10 s denaturation at 96°C, 5 s annealing at 55°C, 4 min elongation at 60°C for 25 cycles. Sequencing products were cleaned up by using ExTerminator (A&A Biotechnology, Poland) and sequenced in both directions in Genomed company (Warsaw, Poland). Obtained sequences of single haplotypes per marker and species were deposited in GenBank.

Table 3. Primers used for amplification of the four DNA fragments sequenced in the study

Fragment	Primer name	Primer sequence (5'-3')	References
<i>COI</i>	COI_Der_Ff	TATATATAATTTTTGGGGTTTGATGTGG	This study
	COI_Der_Rr	CAAAAAGATGTTGATATAAAAATAGG	
CytB	151-F	TGTGGRGCNACYGTWATYACTAA	Merritt, 1998
	270-R	AANAGGAARTAYCAYTCNGGYTG	
ITS-2	ITS2_Der_Ff	GTCGGCTAGTCWAAAGCAATCG	This study
	ITS2_Der_Rr	CCGCTTCACTCGCCGTTACT	
28S rRNA	28S_Der_Ff	GCTAAATACTTGCACGAGTCCG	This study
	28S_Der_Rr	ACGGTTGCCAGTCTCTCC	

Data analysis

All obtained *COI* sequences were blasted by NCBI BLAST (Altschul et al. 1990) to verify species identification and to find similar homologous sequences. In order to visualize phylogenetic relationships between studied species, selected *COI* sequences (the longest and with published status) of six named *Deroceras* species, *D. golcheri* (Accession numbers: JN248291-293), *D. laeve* (MG421043, MG423214, KX959499-501, HM584699), *D. panormitanum* (JN248304-313), *D. reticulatum* (MG421618, MG421373, MG421157, MG421125, MG421099, MF545181, MF545161, MF545125, MF545107, FJ481179), *D. agreste* (KF894312, KF894346-247, KF894375), *D. invadens* (JN248295-297, JN248301-303, JN248314-315, KX959490), were downloaded from GenBank (Eskelson et al. 2011; Reise et al. 2011; Rowson et al. 2014; Araiza-Gómez et al. 2017) and together with sequences obtained in this study for *D. reticulatum*, *D. agreste* and *D. turcicum*, were aligned in BioEdit 5.0.0 (Hall 1999) with the multiple alignment function of ClustalW (Thompson et al. 1994). Aligned sequences were trimmed and translated into protein sequences in MEGA v. 7 to check against pseudogenes. Uncorrected pairwise distances between species were calculated in MEGA v. 7 (Kumar et al. 2016). Uncorrected genetic distances for *COI* were calculated based on all sequences obtained for *D. agreste*, *D. reticulatum* and *D.*

turcicum together with those used for phylogenetic tree calculation (alignment length: 402 bp). For this molecular marker, genetic distances were calculated in two ways, firstly between all sequences in the dataset (see Table S4), and secondly, between species defined as groups (Table 4). In the case of *cyt-b*, ITS-2 and 28S rRNA, only the newly obtained sequences for *D. agreste*, *D. reticulatum* and *D. turcicum* were used for genetic distances calculations (lengths of alignments: 299 bp, 280 bp, 352 bp, respectively) (Table 5). The concatenated dataset comprises 79 sequences in total (*COI* sequences of six *Deroceras* species from GenBank (42 sequences), *COI*, *cyt-b*, ITS-2 and 28S rRNA sequences for studied species obtained in this study (36 sequences) and one sequence of *Limax maximus* (Accession Number: KM612139) as outgroup). The obtained alignment of the concatenated sequences (1619 bp) were divided into eight data blocks constituting the ITS-2, 28S rRNA as well as *COI* and cytochrome b that were separated into three codon positions using PartitionFinder v. 2.1.1 under the Bayesian Information Criterion (BIC) (Lanfear et al. 2016). PartitionFinder retained five out of eight predefined partitions with the best fit models for them being (1) F81 for 28S rRNA, (2) TIM+G for the 2nd codon position in *COI* and 3rd codon position in cytochrome b, (3) TRN+G for the 3rd codon position in *COI* and 1st codon position in cytochrome b, (4) TRN+I for the 1st codon position in *COI* and 2nd codon position in cytochrome b, (5) JC for ITS-2. For the maximum-likelihood (ML) analysis in RAxML, models GTR, GTR+I, GTR+G and GTR+I+G were also tested (Stamatakis 2014). The best fit model for all partitions in this analysis was GTR+G.

Table 4. Genetic distances (%) between *Deroceras* species calculated from *COI* sequences

	1	2	3	4	5	6	7
1 <i>D. agreste</i>							
2 <i>D. golcheri</i>	17.1						
3 <i>D. invadens</i>	14.1	9.8					
4 <i>D. laeve</i>	14.4	10.3	7.8				
5 <i>D. panormitanum</i>	15.7	6.0	7.4	8.5			
6 <i>D. reticulatum</i>	5.6	15.1	14.7	13.3	13.6		
7 <i>D. turcicum</i>	9.8	16.6	14.2	14.4	15.2	8.6	

Table 5. Genetic distances (%) between three *Deroceras* species calculated from cytochrome b, ITS-2, 28S rRNA sequences

		1	2	3
cytochrome b	1	<i>D. agreste</i>	-	
	2	<i>D. reticulatum</i>	6.8	-

	3	<i>D. turcicum</i>	10.8	12.1	-
ITS-2	1	<i>D. agreste</i>	-		
	2	<i>D. reticulatum</i>	7.6	-	
	3	<i>D. turcicum</i>	8.0	3.5	-
28S rRNA	1	<i>D. agreste</i>	-		
	2	<i>D. reticulatum</i>	0.6	-	
	3	<i>D. turcicum</i>	1.2	1.7	-

ML topology was constructed using RAxML v. 8.0.19 (Stamatakis 2014). The tree branches were supported by bootstrap analysis with 1,000 replicates. Bootstrap support values $\geq 70\%$ were regarded as significant statistical support. Bayesian inference (BI) marginal posterior probabilities were calculated in MrBayes v. 3.2 (Huelsenbeck and Ronquist 2001, Huelsenbeck et al. 2001) with 1 cold and 3 heated Markov chains for 10 million generations and trees were sampled every 1000 generations. In the BI consensus tree, clades recovered with posterior probability (PP) between 0.95 and 1 were considered well supported, those with PP between 0.90 and 0.94 were considered moderately supported and those with lower PP were considered unsupported. Obtained trees were visualized in FigTree v.1.4.3, available at: <http://tree.bio.ed.ac.uk/software/figtree>.

RESULTS

All specimens previously identified by prof. Andrzej Wiktor were confirmed as *D. reticulatum*, *D. agreste* and *D. turcicum* based on morphology of their genitalia, especially penis and penial gland appearance (Figs 1A–I and 2A–C). *D. reticulatum* has a pronounced laterally constricted penis with both parts inscribed in an oval and with a large penial gland on the posterior end of the penis (Fig. 1A–C). The penial gland can vary in shape (Fig. 1A–C). It can be represented by a single process with both lateral sides covered by easily identifiable papillae-like structures (Fig. 1A) or made up of a few branches covered by glandular papillae and merged into a common short trunk (Fig. 1B). However, these papillae-like structures or glandular papillae on the lateral sides of the penial gland can sometimes be only faintly visible (Fig. 1C). *D. agreste* is characterized by sack-shaped penis, which can be laterally narrowed in older specimens. The penial gland does not show variability in its shape, it is visible as a small conical nodule or unbranched finger with a smooth surface without any papillae-like structures on the lateral sides (Fig. 1D–F). *D. turcicum* has a laterally constricted penis with the anterior part heavily bloated and a narrower posterior part. The penial gland can be a complex, multifurcated structure or may be significantly reduced (Fig. 1G–I).

More specifically, penial glands can consist of multiple knots, with the lateral sides covered by convexity/constrictions that can resemble papilla-like structures (but usually bigger than in *D. reticulatum*) (Fig. 1G–H), or it can be reduced and comprise only a few, usually short nodular processes set directly on the penis or on a short common stem (Fig. 1I).

Good quality sequences for all four molecular markers were obtained from all analyzed specimens. Only single haplotypes were recovered for each marker in each species. For the three analyzed species the following sequences were obtained:

- *D. agreste*: the *COI* sequence (GenBank: MN934413), 512 bp long, the cytochrome *b* sequence (GenBank: MN931234), 390 bp long, the ITS-2 sequence (GenBank: MT361821), 280 bp long, the 28S rRNA sequence (GenBank: MN930515), 400 bp long.
- *D. reticulatum*: the *COI* sequence (GenBank: MN934414), 651 bp long, the cytochrome *b* sequence (GenBank: MN931235), 299 bp long, the ITS-2 sequence (GenBank: MT361823), 352 bp long, the 28S rRNA sequence (GenBank: MN930516), 352 bp long.
- *D. turcicum*: the *COI* sequence (GenBank: MN934415), 420 bp long, the cytochrome *b* sequence (GenBank: MN931236), 299 bp long, the ITS-2 sequence (GenBank: MT361822), 350 bp long, the 28S rRNA sequence (GenBank: MN930517), 409 bp long.

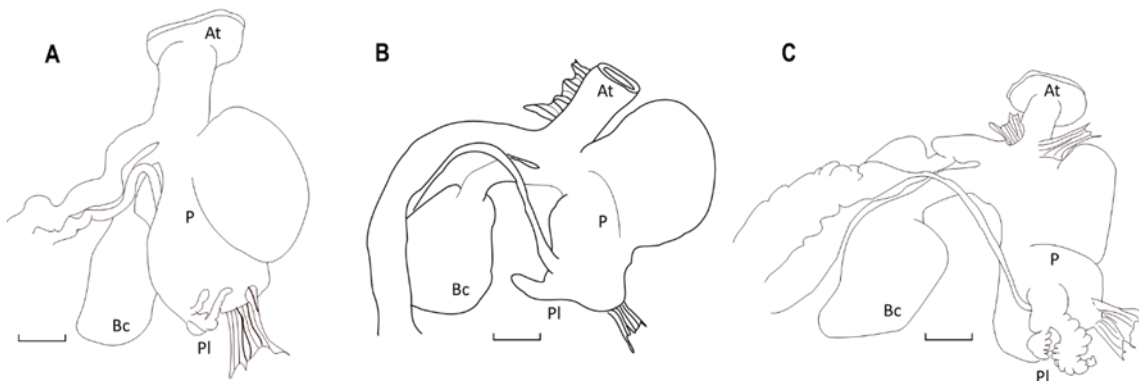


Fig. 2. Schematic drawings of genitalia of *D. reticulatum* (A), *D. agreste* (B) and *D. turcicum* (C) after Wiktor (2000). Bc, bursa copulatrix; Pl, penial gland; P, penis; At, atrium. Scale bars = 1 mm.

Regarding the *D. reticulatum* *COI* sequence, the verification in BLAST resulted in fitting to *D. reticulatum* (KY765589 (Ahn et al. 2017); Query cover: 100%; E-value: 0.00; Perc. ident: 98.92%). Best hit for the *D. agreste* sequence were *D. reticulatum* and *D. agreste* (LS974196, (Kropf, unpublished) and KF894312 (Rowson et al. 2014), respectively; for both species: Query cover: 100%; E-value: 0.00; Perc. ident: 99.41%). For the *D. turcicum* sequence the best hit was *D.*

reticulatum (MG421618 (Dewaard, unpublished); Query cover: 99%; E-value: 1e-162; Perc. ident: 91.87%).

Genetic distances calculated for *COI* sequences between *Deroceras* species, range from 5.6% (between *D. agreste* and *D. reticulatum*) to 17.1% (between *D. agreste* and *D. golcheri*) (12.01% on average) (Table 4). Regarding other molecular markers, genetic distances for cytochrome *b* range from 6.8% (between *D. agreste* and *D. reticulatum*) to 12.1% (between *D. reticulatum* and *D. turcicum*), for ITS-2 range from 3.5% (between *D. reticulatum* and *D. turcicum*) to 8.0% (between *D. agreste* and *D. turcicum*) and for 28S rRNA, genetic distances range from 0.6% (between *D. agreste* and *D. reticulatum*) to 1.7% (between *D. reticulatum* and *D. turcicum*) (Table 5).

Maximum-Likelihood (ML) and Bayesian (BI) phylogenetic trees resulted in similar tree topology (Fig. 3). Specimens belonging to particular species constituted seven separate clades, in most cases strongly supported in BI analysis and moderately or not supported by ML analysis (Fig. 3). The three species on which we focused in this study (*D. reticulatum*, *D. agreste* and *D. turcicum*) have been found to be distinct but closely related to one another and together form a clade with a sister relationship to the other analyzed species (Fig. 3). However, due to polytomy, specific relationships between these three species cannot be inferred. Individuals identified, based on anatomical studies conducted by Wiktor and by us (GenBank: MN934413, this study) and by Rowson et al. (2014) (GenBank: KF894312, KF894346-247, KF894375) as *D. agreste*, cluster together, thus tentatively providing an additional and independent confirmation of species affiliation (Fig. 3). The remaining four *Deroceras* species form another big clade which is a sister relationship to (*D. reticulatum* + *D. agreste* + *D. turcicum*). Here, similarly, the specific relationships between these species cannot be fully recovered due to the several weakly supported nodes that indicate polytomy (Fig. 3).

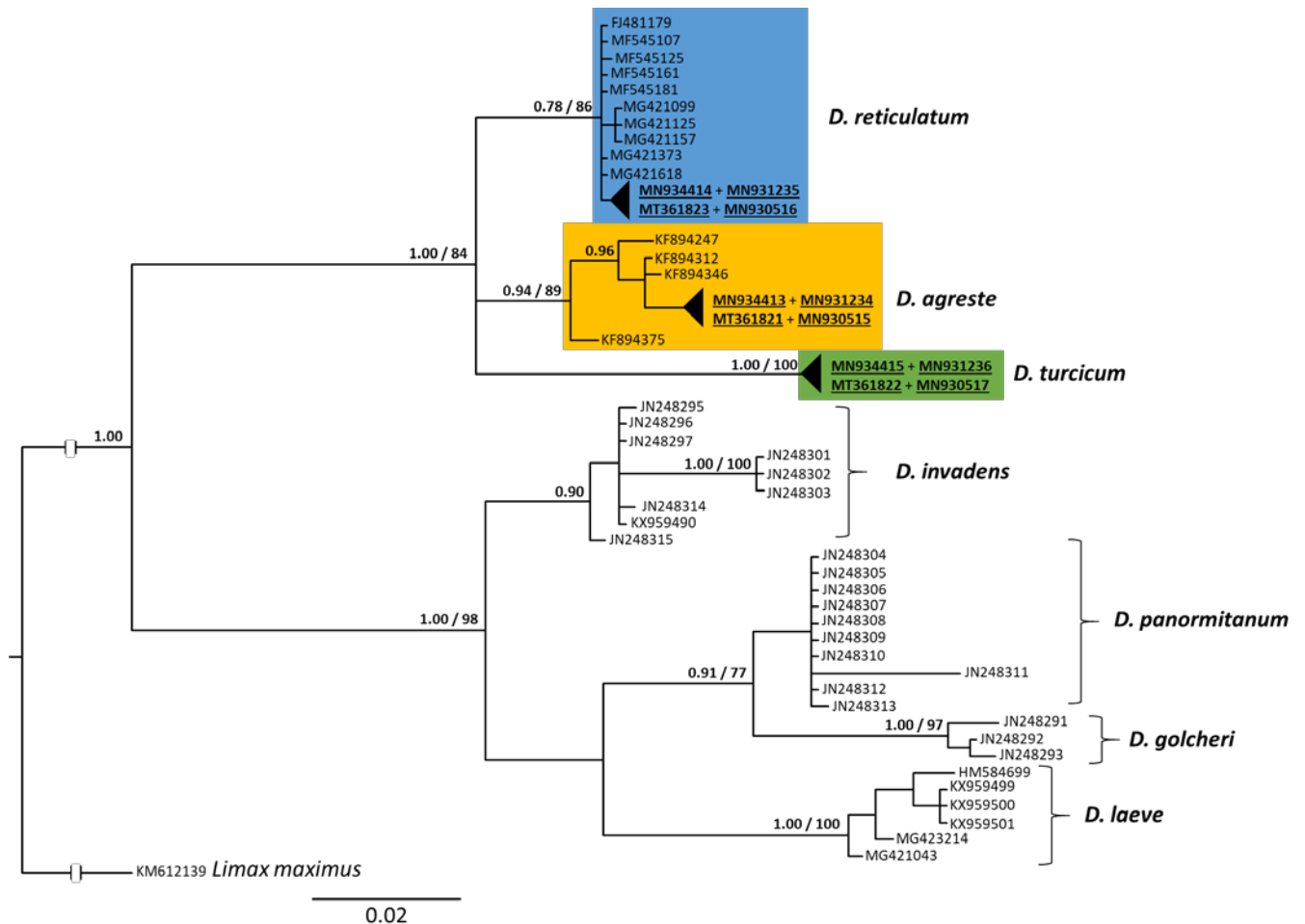


Fig. 3. A phylogenetic tree constructed from concatenated sequences of *COI*, *cyt-b*, *ITS-2* and 28S rRNA of *Deroceras* species. Only topology of Bayesian tree is shown. Bolded and underlined GenBank accession numbers indicate sequences obtained in this study. Numbers at nodes indicate Bayesian posterior probability and bootstrap values separated by “/” mark. Values less than 0.90 for BI and less than 70 for ML are not shown. The scale bar represents 0.02 substitutions per nucleotide position.

DISCUSSION

In this study we have shown that the three morphologically similar species, *D. reticulatum*, *D. agreste*, *D. turcicum*, are genetically easily differentiable. The results from the BLAST search clearly underlined the problems with identification of these taxa, namely that *D. agreste* and *D. turcicum* are often wrongly designated as the more common *D. reticulatum*. The DNA sequences presented in our study are valuable as they came from specimens deposited in museum collection and had been correctly identified by Professor Andrzej Wiktor, a specialist in slug taxonomy, and additionally verified in this study. Additionally, sequences for *D. agreste* were obtained from the

specimens collected from *terra typica* in Sweden. Our phylogenetic results, presented above, additionally confirmed the molecular distinctives and monophyly of each studied species. As the number of traits associated with the external appearance of slugs are usually extremely limited, the morphology of the reproductive system are considered crucial for a precise identification of many slug species (*e.g.*, Reise and Hutchinson 2001). In accordance with Wiktor (2000), we identified penis and penial gland appearance to be the most useful character for at least a provisional, and not so complicated identification (even for non-specialist), due to the ease by which these structures can be located. In concordance with previous studies we also confirmed a considerable morphological variability in the penial glad in *D. reticulatum* (Wiktor 2000) and *D. turcicum* (Wiktor 2000; Reise and Hutchinson 2001), as well as almost no variation in *D. agreste* (Wiktor 2000). However, in dubious cases the application of genetic data and cross-verification with the specific DNA sequences provided in our study will be of great help to other researchers. Moreover, this may have increased significance when only juveniles are present in a given sampling area as anatomical studies of their genitalia will not be possible.

The mitochondrial *COI* and cytochrome b were the most variable molecular markers and thus most informative in terms of species distinction. The first of these is already known as the most useful molecular marker for species identification (Fontaneto et al. 2015), however as we show cytochrome b is also variable enough to distinguish *Deroceras* species. Moreover, cytochrome b was successfully used for species identification and delimitation of members of other gastropod families (*e.g.*, Merritt 1998; Groenenberg et al. 2016). Notably, amplification and/or sequencing of these two DNA fragments are often problematic because of their length, so obtaining good quality sequences is challenging but possible in most cases with more specific primers designed for shorter DNA fragments. Moreover, specimens deposited in museum collections are usually old, thus the DNA is usually degraded and highly fragmented. Work with such material often requires specific kits for DNA extraction that are intended for material with a trace of DNA content. Therefore, by presenting reliable sequences for *cyt-b* and *COI*, we increased the chances for correct species identifications in future research. Regarding nuclear markers, ITS-2 and 28S rRNA, it has already been shown that they are more conservative than the mentioned mitochondrial *cyt-b* and *COI* (Wade et al. 2006). This makes them especially suitable for resolving deeper phylogenetic nodes, especially those between genera, and of higher taxonomic levels (*e.g.*, Wade et al. 2006; Neiber and Hausdorf 2015). Similar to other studies, our results show that ITS-2 is more variable than 28S rRNA. As such, ITS-2 can be used as additional support for the identification by mitochondrial markers as well as *e.g.* in phylogeographic research (Schniebs et al. 2013; Zhou et al. 2017).

Our phylogenetic analysis revealed a similar tree topology for *Deroceras* species to that found in the recent study conducted by Rawson et al. (2014). The addition of new sequences for *D.*

turcicum enabled us to show a close relationship with the morphologically similar *D. reticulatum* and *D. agreste*. Even though the phylogenetic position of the genus *Deroceras* is known (Bouchet and Rocroi 2005), the specific and detailed relationships between its species have only been examined superficially (e.g., Koene and Schulenburg 2005). As mentioned above, molecular data regarding these slugs are extremely limited. Currently, the existing data enabled the testing of relationships between only seven named *Deroceras* taxa, from a genus comprising more than 120 species (Wiktor 2000). Importantly, many of them (including specimens of the type series) are deposited in museum collections, thereby providing an opportunity to obtain tissue for genetic studies from accurately identified specimens. This might be significant as morphological species determination is quite challenging and it may be difficult to obtain fresh material from the field because of the species status (e.g. endangered, endemic) or its rare occurrence. Thus, museum collections constitute a valuable source of reliable barcodes which might help enhance species identification and facilitate biodiversity estimations (Neiber and Hausdorf 2016; Neiber et al. 2017).

CONCLUSIONS

Our study presents a molecular approach to the identification of three closely related species of *Deroceras* slugs by providing, for each of them, definitive DNA sequences of four molecular markers. We designed *de novo* specific primers for three molecular markers (*COI*, *ITS-2*, *28S rRNA*), which successfully amplified DNA fragments, and which can be used on the other *Deroceras* species held within museum collections. The DNA sequences of *D. turcicum* are presented in our study for the first time, providing new and compelling evidence of its close relationship with the two morphologically similar taxa (*D. reticulatum* and *D. agreste*). The sequences provided in this study are reliable barcodes which may aid correct species identification even by non-specialists. This study underlines the important role of natural history museums and identifies their collections as priceless resources of global biodiversity. Finally, our results have application in studies concerning the distribution, conservation biology and introgression in *Deroceras* species.

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Authors' contributions: KSZ designed the study, analyzed the data and wrote the manuscript. DS designed primers and contributed to data analysis and manuscript writing.

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Supplementary Materials

Table S1. GenBank accession numbers of DNA sequences used to design specific primers for three molecular markers (*COI*, ITS-2, 28S rRNA). (download)

Table S2. Concentrations of chemicals for PCR reactions. (download)

Table S3. PCR profiles. (download)

Table S4. Genetic *p*-distances between available COI sequences of *Deroceras* species. (download)