

Deception in taxonomy – the example of *Truncatellina rothi* (Hesse, 1916) (Pupilloidea: Truncatellinidae)

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In this study, *Truncatellina cylindrica* (A. Férussac, 1807), *Truncatellina rothi* (Hesse, 1916), and *Truncatellina haasi* Venmans, 1957 were investigated, and a lectotype for *Isthmia rothi* was selected. So far, the three species have only been defined by their shell morphology. Intensified collection efforts throughout the past 100 years increased the number of specimens available, and it has become increasingly questionable whether these three species represent different taxa. In this study 1070 specimens were photographed, and measurements of several shell traits were taken. A multivariate ratio analysis revealed that the morphological trait variation of *T. rothi* (and probably *T. haasi* as well) is mainly covered by the variability of *T. cylindrica*. There is no morphological evidence that the investigated specimens represent different taxa. A genetic study is needed to reveal possible cryptic clades.

Key words: Dwarf snails, Greece, morphometrics, shape PCA, taxonomic revision, *Truncatellina*, Turkey.

INTRODUCTION

The shell of a land snail protects it from enemies or from drought and is influenced by environmental conditions as well as by the genetics that determine the bauplan. As a con-

sequence, many land snails have similar shells. This similarity often makes it difficult to distinguish species or even genera. Adult land snails often develop a strong lip when they stop growing, and this distinguishes adults from juveniles (Goodfriend, 1986). In general, only adult specimens should be used for species identification. Nonetheless, serious difficulties for a reliable determination may remain. Differentiation criteria are often presented too optimistically (or uncritically) and cannot be applied correctly in practice.

In this study we test specimens collected by Peter Subai, who used the traditional shell-morphological approach to reliably distinguish the toothless species *Truncatellina cylindrica* (A. Férussac, 1807), *T. rothi* (Hesse, 1916) and *T. haasi* Venmans, 1957, whether the traditional criteria lead to an unambiguous determination of the species, or not.

Truncatellina rothi is described from the vicinity of Athens, and since then, it is recorded in large numbers from Greece, Cyprus (Vardinoyannis et al., 2012), and the western part of Turkey. Reinhardt (1916: 164), distinguished the new species from *T. cylindrica* by the following shell character states: 1) a “blunt” shell, 2) the peculiar shell form, where the upper whorls are wider if compared to the lower, 3) six strongly convex whorls, 4) a deep suture and 5) the ribbing pattern, where rib interspaces are wider than rib thickness. Upon receipt of a specimen from the Sea of Galilee, which superficially matched his new species, Reinhardt stated that *T. rothi* is common in the Eastern Mediterranean region. However, this is erroneous and based on a misidentification with a species, which today is known as *T. haasi*, a species known from the Levante region and Turkey (Pavliček et al., 2008; Heller, 2009; Neubert et al., 2015).

The authorship of *T. rothi* has so far been attributed to Reinhardt (1916). Much to our surprise we noticed that *T. rothi*



Figs 1-8. Species and specimens of *Truncatellina* as identified by authors. **1.** *T. cylindrica*, SMF 273803/3a, Istanbul, Büyük Ada, sh = 1.8 mm. **2.** Syntype *Pupa minutissima* var. *obscura*, ZMZ 514580, Greece, Janina, sh = 2.2 mm. **3.** Lectotype *Isthmia rothi*, SMF 4082, Greece, Athens, sh = 1.8 mm. **4.** *T. cylindrica*, NMBE 543359, Romania, Dambovicioara, Arges, sh = 1.79 mm. **5.** *T. cylindrica*, NMBE 543395, Greece, Kefalari Mountain, Korinthia, sh = 1.83 mm. **6.** *T. rothi*, NMBE 543991, Greece, Karpenissi, sh = 1.72 mm. **7.** *T. haasi*, NMBE 545643, Turkey, Silifke, cave Narlıkuyu Mağarisi, sh = 1.55 mm. **8.** *T. haasi*, NMBE 545642, Turkey, Side, Roman theatre, sh = 1.47 mm. All photographs $\times 25$.

was already made available a few months earlier by Hesse in a paper on “Rumelia” in the same journal as Reinhardt (Hesse 1916: 121). It is based on the same specimens Reinhardt used for his description, and which once was collected by Roth. In parallel, Hesse included a specimen from Palestine (later described as *T. haasi*) from Tabgah, Lake Tiberias. We here-with select SMF 4082, “bei Athen”, coll. Reinhardt ex Roth as lectotype for *Isthmia rothi* to fix the use of this name.

Due to intensified collection efforts in the past century, more specimens of the taxa *T. cylindrica*, *T. rothi* and *T. haasi* are now available for comparison (Figs 1-8). Recognising consistent differences between the three species became increasingly difficult the more specimens were inspected. Therefore, we hypothesise that the morphological trait variation of *T. rothi* and *T. haasi* is covered by the variability of *T. cylindrica*. This hypothesis is investigated by using morphometric methods. The specimens used in this study pre-

dominantly originate from the collection Subai, which is housed in the NMBE. Thus, they were identified by a single collector (except *T. haasi*, identified by Bank & Neubert), who utilized the aforementioned conchological traits to separate the species. Thus, any bias due to a probably differing view or concept could largely be excluded.

The distribution of *T. cylindrica*, *T. rothi*, and *T. haasi* is shown in Fig. 9. A distinct line of demarcation in the occurrence of *T. cylindrica* and *T. rothi* is observed in Greece with only one exception in Kefalari. In the south of Greece, all specimens were automatically identified as *T. rothi*. It is striking that in each locality, all specimens recorded were identified as either *T. cylindrica* or as *T. rothi*. Syntopy of different species is indirectly reported by Holyoak et al. (2012) for other species of *Truncatellina*. Thus, the “allopatric” situation for these two taxa in Greece is quite remarkable and probably caused by a “human” bias.



Fig. 9. Distribution of investigated *T. cylindrica* (violet dots), *T. rothi* (green dots), and *T. haasi* (brown dots) populations in Greece and Turkey as present (with a few exceptions) in the collection of Peter Subai (now NMBE).

MATERIAL AND METHODS

Abbreviations used for collections

NMBE = Natural History Museum Bern, Switzerland; SMF = Research Institute Senckenberg, Frankfurt am Main, Germany; ZMZ = Zoological Museum Zürich, Switzerland

Specimens investigated

The specimens from the Subai collection were collected in Greece and Turkey between 1975 and 2008. The total sample size comprised 1070 specimens. To investigate the trait variation in a large population of *T. cylindrica*, 100 randomly selected specimens from a single population in Romania (out of > 900) were used. We investigated 26 lots with 430 specimens identified as *T. cylindrica* and 33 lots with 520 specimens identified as *T. rothi* from Greece and Turkey. When possible, 20 randomly chosen adult specimens from each lot were photographed and measured. In less specious lots, at least 10 individuals were processed, whereas in some smaller lots, all specimens were assessed. Additionally, 18 specimens of *T. haasi* from Turkey and the Levante were included in the study. The lectotype of *Isthmia rothi* and a syntype of *Pupa minutissima* var. *obscura* Mousson,

1859 (ZMZ 514580, Janina, coll. Mousson ex Schläfli) were included. A detailed list of all the lots and their localities is presented in Table 1. The map in Figure 9 was produced with QGIS (2016, v2.18.12) using the Natural Earth data set.

Measurements

Each specimen was placed on a piece of clay in standardized vertical orientation with the aperture to the right and imaged with a Leica DFC425 microscope camera. Using an image processing program (IMS Client v15Q4, Imagic, Switzerland), the shell width (sw) and shell height (sh), the aperture width (aw) and aperture height (ah), the maximum diagonal diameter of the penultimate whorl (dpw), whorl width of the third whorl (ww), and the suture of the third whorl (s) were measured. All measurements were taken in µm. Whorl number (n) was counted from the aperture towards the apex (Fig. 10). In order to compare different specimens, the number of ribs per 500 µm (R_{500}) on the penultimate teleoconch whorl was counted. Measurements were chosen according to Reinhardt’s differences (1916: 164) between *T. rothi* and *T. cylindrica*. We measured shell width (sw) and whorl width of the third whorl (ww) to account for Reinhardt’s criteria 1) and 2). The number of whorls (n)

Species	NMBE.no.	Country	Location	Latitude	Longitude	Altitude [m]
<i>cylindrica</i>	543359	Romania	Dambovicioara, gorge 200-300 m N / Arges	45.454	25.219	970
<i>cylindrica</i>	543386	Greece	Nikissiani, south edge	40.947	24.144	350
<i>cylindrica</i>	543387	Greece	Akrovouni, 17.8 km for Pangéo-summit	40.915	24.147	1280
<i>cylindrica</i>	543320	Greece	Akrovouni, 11.6 km for Pangéo-summit	40.917	24.191	900
<i>cylindrica</i>	543424	Greece	Megali- and Mikra-Prespaseen, confluence	40.811	21.071	855
<i>cylindrica</i>	543417	Greece	Loutra Arides, entry to valley	40.972	21.912	450
<i>cylindrica</i>	543420	Greece	Prossotsani, 6 km N	41.223	23.969	340
<i>cylindrica</i>	543389	Greece	Kato Vrontou, 3.8 km, marble quarry	41.262	23.788	750
<i>cylindrica</i>	543451	Greece	Loutra Arides, entry to valley	40.972	21.912	450
<i>cylindrica</i>	543310	Greece	Stavroupoli, west edge next to river Nestos	41.203	24.706	120
<i>cylindrica</i>	543419	Greece	Aetia, 2.2 km E for Grevena, river breakthrough	40.074	21.202	1005
<i>cylindrica</i>	543302	Greece	Prossotsani, 6 km N	41.223	23.969	340
<i>cylindrica</i>	543395	Greece	Kefalari, 2 km E. mountain / Korinthia	37.915	22.532	730
<i>cylindrica</i>	543332	Greece	Gérmas, valley w	40.453	21.415	850
<i>cylindrica</i>	543383	Greece	Stavroupoli, west edge next to river Nestos	41.203	24.706	120
<i>cylindrica</i>	543468	Greece	Monastir Panagia Mavriotissa, 50-100 m w	40.505	21.279	640
<i>cylindrica</i>	543382	Greece	Zarkadia, 2.5 km NNE	41.029	24.642	450
<i>cylindrica</i>	543384	Greece	Profitis Ilias summit region / Ossa-mountains / Thessalia	39.795	22.671	1550
<i>cylindrica</i>	549113	Turkey	Bilecik, gorge underneath the mosque of Sey Edebalı	40.144	29.99	400
<i>cylindrica</i>	549112	Turkey	Nusaybin, bridge over Çağcag Dere	37.09	41.215	470
<i>cylindrica</i>	549111	Turkey	right shore of Kizilirmak, 13 km SE Duragan / Sinop	41.340	35.140	180
<i>cylindrica</i>	549110	Turkey	Yeniçağa Gölü	40.782	32.034	988
<i>cylindrica</i>	543459	Turkey	Manavgat, bridge 13 km N of the cascade / Antalya	36.588	32.079	no data
<i>cylindrica</i>	547695	Turkey	Mekece	40.452	30.048	100
<i>cylindrica</i>	543390	Turkey	Road Akseki-Seydisehir, 1.5 km for Bozkir / Konya	37.220	31.936	1720
<i>cylindrica</i>	547694	Turkey	Gülümbe, 6 km before Bilecik	40.199	29.967	500
<i>cylindrica</i>	543558	Turkey	Dere (= Dereköy), west border / Bozkir / Konya	37.176	32.167	1260
<i>rothi</i>	544029	Greece	Klisura gorge, east edge, Ag. Eleusis / Etoloakarnania	38.501	21.376	350
<i>rothi</i>	543991	Greece	Karpenissi, 7.5 km WNW / Evritania	38.925	21.745	1250
<i>rothi</i>	544060	Greece	Klidonia, Turkish bridge, north end of the Vikos gorge / Ipiros	39.968	20.663	425
<i>rothi</i>	543999	Greece	Sandoméri, mountain Skollis / Ahaia	37.990	21.578	550
<i>rothi</i>	544073	Greece	Githio, 14 km for Areopoli / Lakonia	36.702	22.471	100
<i>rothi</i>	543995	Greece	Agios Pétros, 5.5 km SSE / Parnon mountains / Arkadia	37.311	22.578	970
<i>rothi</i>	544027	Greece	Agionori, 2 km for Limnes / Argolida	37.744	22.874	700
<i>rothi</i>	544078	Greece	Githio, 8.5 km for Areopoli / Lakonia	36.740	22.494	150
<i>rothi</i>	544072	Greece	Kalamata, 9.4 km, Artemissia gorge / Messinia	37.084	22.158	310
<i>rothi</i>	543993	Greece	Alivéri, west border / isle Évía	38.402	24.014	350
<i>rothi</i>	544041	Greece	Agios Ioannis, 1 km for Astros / Arkadia	37.354	22.640	750
<i>rothi</i>	544077	Greece	Astakos, mountain Veloutsá, SE-slope / Etoloakarnania	38.537	21.069	185
<i>rothi</i>	544037	Greece	Gliki, road to Koukoulii / Ipiros	39.332	20.603	160
<i>rothi</i>	543990	Greece	Paralia Sarandi, 3 km for Prodromos / Viotia	38.257	22.876	220
<i>rothi</i>	544020	Greece	Petralona, cave of Petralona / Makedonia	40.372	23.169	335
<i>rothi</i>	544031	Greece	Amfiklia, SE-border, small chapel, entry to the gorge / Fthiotida	38.631	22.596	500
<i>rothi</i>	544030	Greece	Gorge Klisura, W-edge, Ag. Eleusis / Etoloakarnania	38.501	21.368	290
<i>rothi</i>	544023	Greece	Argos, old castle, ruins / Argolida	37.639	22.715	250
<i>rothi</i>	544045	Greece	Kastritsa, mountain / Ipiros	39.639	20.915	500
<i>rothi</i>	544058	Greece	Langadia, 2 km for Arhea Olympia / Arkadia	37.682	22.015	850
<i>rothi</i>	544076	Greece	Petrona, W-edge / Etoloakarnania	38.926	21.407	500
<i>rothi</i>	544075	Greece	Triklino, 3.9 km for Alevrada / Etoloakarnania	38.954	21.466	no data
<i>rothi</i>	544004	Greece	Delfi, 2 km for Amfissa / Fokida	38.484	22.472	400
<i>rothi</i>	544070	Greece	Branch-off to Loutra Elefthero, 4.5 km W / Makedonia	40.721	24.060	30
<i>rothi</i>	544044	Greece	Vouliagméni, vis-à-vis of the isle Fléves / Atiki	37.809	23.784	20
<i>rothi</i>	544025	Greece	Theopetra, Kuveldi, 6 km SSE from Kalambaka / Thessalia	39.680	21.681	260
<i>rothi</i>	544063	Greece	Sarakina, 1 km W / Thessalia	39.661	21.628	180
<i>rothi</i>	544002	Turkey	Beskonak, bridge over Köprülü Canyon / Antalya	37.192	31.187	200
<i>rothi</i>	509508	Turkey	between Göynük and Çavusdere, 5 km W Çavusdere / Bolu	40.523	31.099	670
<i>rothi</i>	544003	Turkey	Kuru Cay, ca. 1 km E + 200 m N of the road / Konya	37.208	32.065	1420
<i>rothi</i>	544046	Turkey	Yarpuz, 14 km for Seydisehir / Konya	37.229	31.916	1610
<i>rothi</i>	544001	Turkey	Camlik (= Dalayman), 2 km SW. cave Maslialti-ıni / Konya	37.344	31.622	1395
<i>rothi</i>	544000	Turkey	Road Beysehir-Akseki, 9 km for Yesildag / Konya	37.557	31.518	1200
<i>haasi</i>	549116	Turkey	Çesme, 10 rkm S Mordogan / Izmir	38.456	26.550	50
<i>haasi</i>	549115	Turkey	Anamur to Silifke, 4.6 km E Aydıncık	36.152	33.372	40
<i>haasi</i>	549114	Turkey	Uzuncaburç 12 km SW	36.584	33.952	1070
<i>haasi</i>	548745	Turkey	Ruins of Ephesus	37.941	27.342	10
<i>haasi</i>	545643	Turkey	Narlıkuyu Magarisi, 20 km NE Silifke / Silifke	36.449	34.102	140
<i>haasi</i>	545642	Turkey	Side, Ruins	36.768	31.391	10
<i>haasi</i>	517756	Jordan	Zobya	32.433	35.767	900
<i>haasi</i>	508204	Israel	limestone rock, upper Galil / Nahf	32.935	35.935	no data
<i>haasi</i>	508203	Jordan	Cave / Wadi Shwaib	31.970	35.723	no data

Table 1. Museum number, country of origin and geographical data of the records of *T. cylindrica*, *T. rothi* and *T. haasi* used for the study.

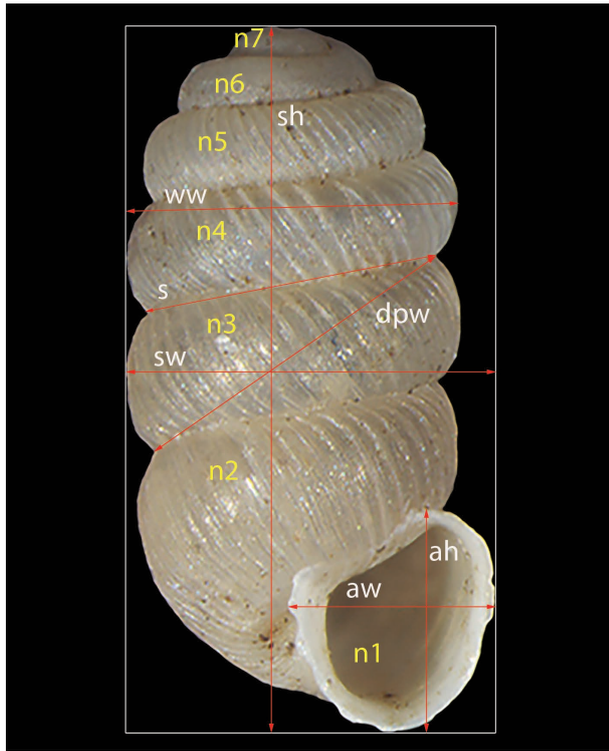


Fig. 10. Explanation of specimen measurements: shell height (sh), shell width (sw), aperture height (ah), aperture width (aw), maximum diagonal diameter of the penultimate whorl (dpw), whorl width of the third whorl (ww), and the suture of the third whorl (s). Whorl number (n) was counted from the aperture towards the apex.

were counted to meet Reinhardt's criteria 3). Criteria 4) is considered by the measurements of whorl width of the third whorl (ww) and suture of the third whorl (s) and criteria 5) is investigated by comparing the number of ribs per 500 μm (R_{500}).

Data analysis

A multivariate ratio analysis (MRA) was performed for shape analysis (R Development Core Team, 2016, v3.2.4). A slightly modified script from Baur & Leuenberger (2011, 2020) was used for the shape Principal Component Analysis (PCA) and the isosize calculation. Methods like PCA treat specimens as belonging to a single group, that is, specimens are not assigned to a particular group a priori. A PCA simply explores the most important pattern of variation in the entire dataset. Only later, when the data points are plotted on the first few axes, they are labelled by at least one factor variable, such as species or locality. But these have no influence on how principal components are calculated. In taxonomy, it is assumed that a PCA reveals the most interesting pattern of variation with respect to species separation. For instance, in sibling species, character variation related to

species membership is usually expected to be more significant than variation related to any intraspecific population. Hence, it is reasonable to assume that PCA do detect species differences (Baur & Leuenberger, 2011, László et al., 2013, Baur et al., 2014).

To calculate the reliability, 30 randomly picked specimens, each 15 specimens of *T. cylindrica* and 15 specimens of *T. rothi*, were photographed and measured three more times in a random order. MANOVA was calculated in R using the comment `manova()`. A pairwise Wilcoxon test was performed in R to calculate rib numbers in *T. cylindrica*, *T. rothi*, and *T. haasi*. The `lm()` function in R was used for investigating any effect of the geographic variables on shape.

RESULTS

Shape and size analysis

Data reliability turned out to be very high (data not shown). Thus, measurement error is low and differences among groups should be traceable.

In order to identify traits which differentiate *T. rothi*, *T. cylindrica*, and *T. haasi*, a shape PCA was performed to explore the variation in shape of these three taxa. The shape PCA used the values of shell height (sh), shell width (sw), aperture height (ah), aperture width (aw), whorl width of the third whorl (ww), suture of the third whorl (s) and maximum diagonal diameter of the penultimate whorl (dpw) for each specimen. Only shape PC1 showed some differences in mean shape of forms, while in shape PC2 they are completely overlapping (data not shown). Hence, we continue only with shape PC1 and discard the remaining axes of the shape PCA. Shape PC1 differences are also highly significant as revealed by MANOVA ($p < 0.001$) (data not shown).

In Figure 11, isosize is plotted against shape PC1 to reveal a possible effect of allometry. This could be ruled out, as the three forms were not at all separated in isosize. Hence, size is no longer considered in the analysis.

The PCA ratio spectrum of shape PC1 (Fig. 12) showed that the differences among mean shape of the three forms was related to their main shell proportions (height to width). The variable shell height (shell.h) is at the upper end of the spectrum, while the five variables related with width, i.e. suture of the third whorl (suture.w), aperture width (aperture.w), shell width (shell.w), maximum diagonal diameter of the penultimate whorl (dp.w), and whorl width of the third whorl (whorl.w) are altogether at the lower end of the spectrum. The variable aperture height (aperture.h) lays in the middle of the spectrum and has thus no effect on shape PC1. One could thus say that generally the shape varied from the rather elongate shape of some *T. cylindrica* (and *T. rothi*) to the sturdier shape of some *T. haasi* (Fig. 11).

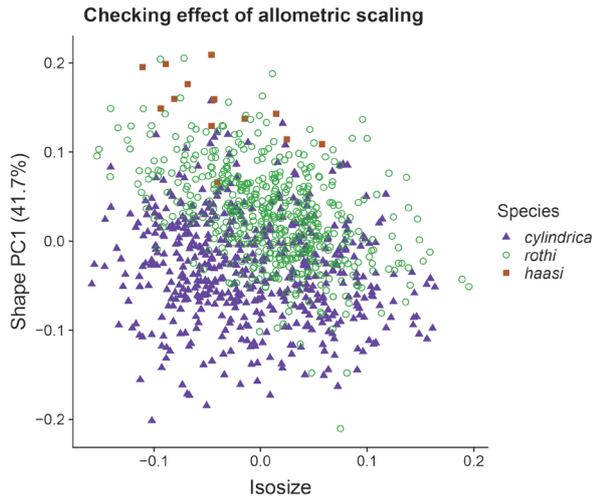
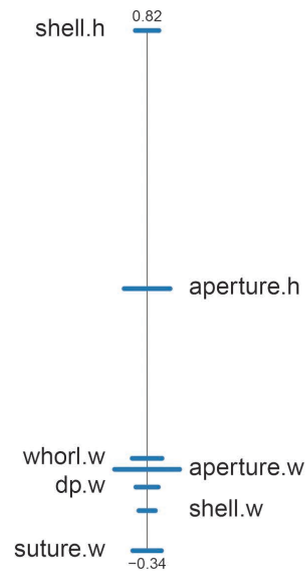


Fig. 11. Isize plotted against shape PC1. Violet rectangles represent *T. cylindrica*, green circles represent *T. rothi*, and brown squares represent *T. haasi*.

PCA Ratio Spectrum for shape PC1



bars = 68% confidence intervals based on 1000 bootstrap replicates

Fig. 12. The PCA ratio spectrum of shape PC1 with the shell measurements of shell height (shell.h), shell width (shell.w), aperture height (aperture.h), aperture width (aperture.w), maximum diagonal diameter of the penultimate whorl (dp.w), whorl width of the third whorl (whorl.w), and the suture of the third whorl (suture.w).

Frequency of whorl number in percentage within species

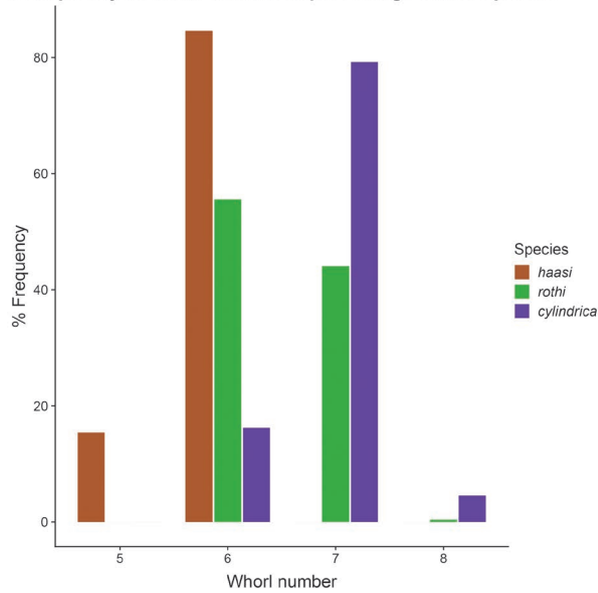


Fig. 13. Frequency of whorl number in percentage within the three species *T. cylindrica* (violet), *T. rothi* (green), and *T. haasi* (brown).

Meristic variables

All three forms show a variable number of whorls (Fig. 13), but are largely overlapping. Whorl number is clearly related to shell height, as the overall correlation between whorl number and shell height is highly significant (Spearman's rank correlation 0.541, $p < 0.001$).

rib number per 500 μm

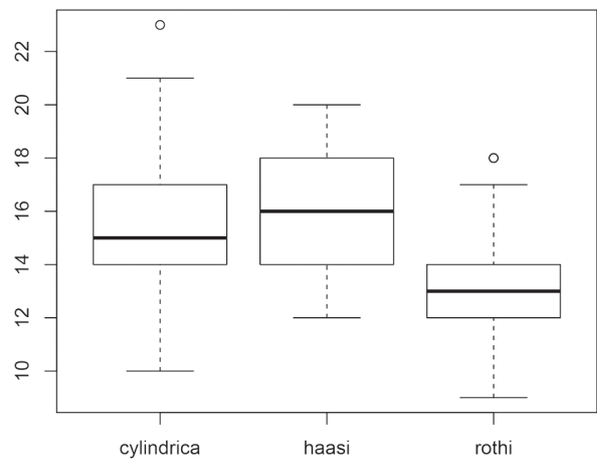


Fig. 14. Number of ribs per 500 μm of *T. cylindrica*, *T. rothi*, and *T. haasi*. A pairwise Wilcoxon test revealed a significant difference in the number of ribs per 500 μm ($p < 0.001$).

As it is the case for the other characters, the number of ribs per 500 μm are almost completely overlapping in range (Fig. 14). Nevertheless, mean number is significantly smaller in *T. rothi* compared to the other forms (pairwise Wilcoxon test with Bonferroni correction $p/3$, $p < 0.001$).

	<i>T. cylindrica</i>	<i>T. haasi</i>	<i>T. rothi</i>	percentage correct
<i>Truncatellina cylindrica</i>	395	3	132	74.5
<i>Truncatellina haasi</i>	0	10	3	76.9
<i>Truncatellina rothi</i>	90	33	397	76.3

Table 2. LDA cross validation of the shell measurements of *T. cylindrica*, *T. rothi* and *T. haasi*.

LDA cross validation of shell measurements

The above analyses clearly show that it is impossible to separate any of the three “species” based on their shell morphology. There were some significant differences in mean shape, but as the ranges were largely overlapping, it was impossible to classify most specimens. For demonstrating, how difficult the separation was, we calculated LDA cross validation of the shell measurements. Although this time specimens are assigned to forms beforehand, the success of the classification was very low (Table 2). Only 76 % of *T. rothi*, 75 % of *T. cylindrica* and 77 % of *T. haasi* were correctly classified. Altogether these are very poor findings, as normally it should be possible that at least 95 % of specimens are classified correctly with LDA cross validation (Baur et al., 2014).

Modelling shape

We could show that the three forms cannot be separated using shell measurements and meristic data. However, the shape PCA clearly revealed that there is significant variation in shape PC1 (data not shown). We therefore applied a linear model to find out whether shape depended from geographic location or altitude. For some terrestrial snail

species, a relation between altitude and shell traits has been suggested (e.g. Engelhard & Slik, 1994; Welter-Schultes, 2000). The model used in this study included latitude, longitude and altitude and allowed for their interaction. Only latitude was significant ($p < 0.01$) (Fig. 15).

DISCUSSION

The analysis of 1070 specimens shows that the variation of *T. rothi* is mainly covered by the variability of *T. cylindrica*. If a trait would have differentiated *T. rothi* from *T. cylindrica*, we would expect two distinct clusters of data points in the shape PCA with almost no overlap but a shift along shape PC1 or along shape PC2.

Figure 14 shows the distribution of the number of ribs per 500 µm. The reduction in number of ribs (Reinhardt, 1916: 164) was one important criterion for Reinhardt to separate the species *T. rothi* from *T. cylindrica*. Here it is shown that this difference exists in populations from Central to Southern Greece and Turkey. The difference between *T. cylindrica* and *T. rothi* is on average 1.8 ribs per 500 µm on the penultimate teleoconch whorl. However, Kemperman & Gittenberger (1988) showed that the ribbing pattern is influenced by humidity and temperature. Snails with many large ribs can retain more water between the shell surface and the substratum. This observation is supported by our data, and as ribbing pattern is obviously influenced by ecological factors, it cannot be considered as a reliable taxonomic trait discriminating between “species”.

Figure 15 shows that the population from Romania covers almost the entire range of variation of the populations of Greece and Turkey. How then would it be possible to separate those three forms? Clearly this is not possible, and from a morphological species concept, the three forms have to be considered as belonging to one and the same, geographically slightly variable species. It should be stressed that it cannot be excluded that certain populations might be separable by means of genetic data. But, again, based on morphometric data of 1070 specimens over a wide distribution range, there is no justification for the recognition of three separate taxa from a morphological point of view.

In their paper in 2012, Holyoak et al. discuss the highly variable species *T. callicratis* (Scacchi, 1833), which can have

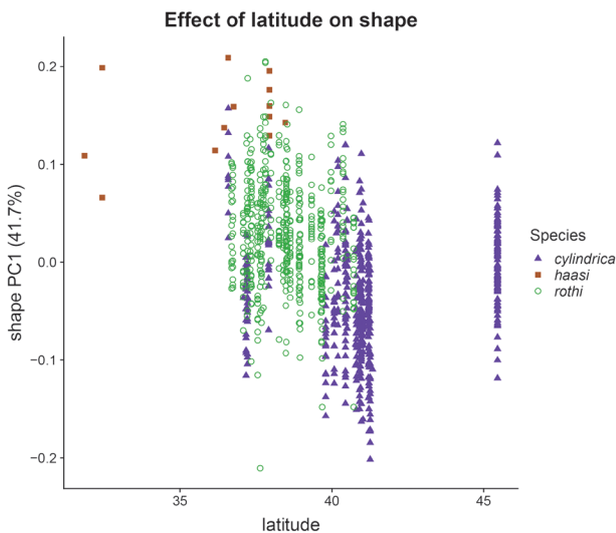


Fig. 15. Shape PC1 plotted against latitude for testing the effect of latitude on shape for *T. cylindrica* (violet rectangles), *T. rothi* (green circles), and *T. haasi* (brown squares).

up to three denticles in the aperture, and thus can be easily distinguished from our species. However, there are many populations of this taxon known with a reduced apertural armature: toothless specimens (or even populations) are well known. This raises the question on how to distinguish such toothless *T. callicratis* specimens from the toothless *T. cylindrica*. After measuring numerous specimens of both taxa from France and mainly the Iberian Peninsula and Northern Africa, Holyoak et al. conclude in their key that both species are mainly distinguished by the “flatness” of their shells, which is quite a subjective trait. This assumption ignores morphological traits that remain to be explored in Central Europe, the Central Mediterranean area and Eastern Europe, where both taxa are known to occur. In Switzerland, both species are recorded from numerous localities, many of them in sympatry, and some probably even syntopic. The question remains whether both species (*T. cylindrica* and *T. callicratis*) represent biological entities, or whether traditional taxonomy is misguided by a single species, which is highly variable in its shell morphology, including all variations regarding the presence or absence of teeth (or lamellae as demonstrated by Nekola et al. (2018) for many species in the Vertiginidae).

CONCLUSION

After the analysis of 1070 specimens, no evidence was found for any significant shell-morphological differentiation as used by Hesse and Reinhardt (1916) between the three nominal taxa *T. rothi*, *T. cylindrica*, and *T. haasi*. The identification of shells as one of the three species seems to follow – especially with respect to *T. cylindrica* and *T. rothi* – an anthropocentric approach rather than a reflection of actually differing morphological traits. However, since the sample size of *T. haasi* is rather small and the distribution range of this species is not covered very well by our samples, we provisionally consider it as a distinct species until more specimens are investigated. But for *T. rothi* and *T. cylindrica* the classical shell-morphological approach is inappropriate and these two taxa should be synonymised. This does not necessarily mean that there are no other toothless *Truncatellina* species existing in Greece or Turkey. If so, a genetic study is needed to reveal such cryptic species, and this is also needed to clarify the taxonomic concept of *T. cylindrica* and *T. callicratis*.

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