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- What should we call the Levant mole? Unravelling the systematics and demography of
- 2 Talpa levantis Thomas, 1906 sensu lato (Mammalia: Talpidae).

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Abstract

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Turkey hosts five of the eleven species of *Talpa* described to date, Anatolia in particular 19 appearing to be an important centre of diversity for this genus. Of these taxa, the Levant mole, 20 Talpa levantis Thomas, 1906 has been suggested to consist of two genetically divergent 21 sublineages, which may represent separate species. Here we use a combination of 22 23 mitochondrial and nuclear DNA sequences from specimens of T. levantis s.lat. collected 24 across the species geographical range to explore the systematics and demographic history of Levant moles. Both mitochondrial and nuclear markers confirm the existence of distinct 25 eastern and western sublineages, which apparently diverged from each other in the early 26 Pleistocene. Given the degree of cytochrome-b divergence between these (7.28%), we 27 28 consider them to represent independent, cryptic species. By including topotypic specimens of 29 T. levantis s. str. in our study we are able to show that this name applies to the western 30 sublineage, distributed across most of the Anatolian Black Sea coastal region, from the vicinity of Trabzon in the east, westwards to Marmara. The earliest name available for the 31 32 eastern taxon, found in Transcaucasia and adjacent parts of northeastern Anatolia, is T. transcaucasica Dahl, 1944. Cytochrome-b haplotype diversity in T. levantis is relatively high, 33 demographic analyses suggesting that the species may have survived in multiple, separate, 34 refugial areas during the Pleistocene. Our work brings the total number of named mole 35 species recognized in Turkey to six, emphasising the importance of this region as a global 36 centre of mole diversification. 37

Key words: *Talpa*; Cytochrome *b*; *BRCA2*; Phylogeny; Anatolia

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Introduction

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Talpa Linnaeus, 1758 is a strictly subterranean genus of moles, distributed throughout the 41 western Palearctic region, from the Iberian Peninsula to China and Siberia (Hutterer, 2005), 42 which has been recovered as monophyletic in a number of phylogenetic studies (e.g., 43 Colangelo et al., 2010; Bannikova et al., 2015). Bannikova et al. (2015) dated the most recent 44 common ancestor of Talpa to 5.49-7.64 Myr, based on four concatenated nuclear genes, a 45 time window in keeping with that suggested by Colangelo et al. (2010) from studies of 46 mitochondrial cytochrome b (cyt-b) sequences, and corresponding to the Late Miocene (latest 47 Tortonian and Messinian stages). During this time several dramatic changes to terrestrial 48 environments and ecosystems occurred, large areas of continents experiencing drying, 49 enhanced seasonality, and a consequent restructuring of terrestrial plant and animal 50 51 communities (Herbert et al., 2016). The most recent version of Mammal Species of the World (Hutterer, 2005) recognized nine valid species in the genus, divided into a western group 52 53 including the common mole *T. europaea* Linnaeus, 1758, the blind mole *T. caeca* Savi, 1822, the Roman mole *T. romana* Thomas, 1902, the Levant mole *T. levantis* Thomas, 1906, the 54 55 Iberian blind mole T. occidentalis Cabrera, 1907 and the Balkan mole T. stankovici Martino and Martino, 1931 and an eastern group comprised of the Siberian mole *T. altaica* Nikolasky, 56 1883, the Père David's mole T. davidiana Milne-Edwards, 1884 and the Caucasian mole T. 57 caucasica Satunin, 1908. Based on genetic data, Bannikova et al. (2015) recognized three 58 additional species: T. talyschensis Vereschagin, 1945; T. ognevi Stroganov, 1948 and Talpa 59 ex gr. levantis. More recently two new mole species, T. aquitania Nicolas, Martínez-Vargas 60 and Hugot, 2017 (Nicolas et al., 2017a) from southern France and northern Spain and T. 61 martinorum Kryštufek, Nedyalkov, Astrin and Hutterer, 2018 from the south-western Black 62 Sea coast (Thrace), were described, initially on the basis of genetic data(Nicolas et al. (2017b; 63 Kryštufek et al. 2018). While most species of *Talpa* are narrowly endemic with predominantly 64 non-overlapping ranges, one species, T. europaea, is widespread and relatively eurytopic 65 across much of Europe, with a range extending from the Ebro River in Spain to the Ob and 66 Irtish Rivers in Russia (Mitchell-Jones et al., 1999; Hutterer, 2005; Loy et al., 2005; Wilson 67 68 and Reeder, 2005; Nicolas et al., 2017b). 69 70 The descriptions of the majority of the above *Talpa* species are based primarily on 71 morphometry (Corti et al., 1985; Corti and Loy, 1987; Loy et al., 1993; Kryštufek, 1994; 72 Rohlf et al., 1996; Loy and Capanna, 1998; Kryštufek and Benda, 2002; Kryštufek et al.,

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2018; Selçuk et al., 2018; Sansalone et al., 2019) and dental traits (Nicolas et al., 2017a;
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       Kryštufek et al., 2018). However, the highly conservative morphology of the genus, resulting
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       from the functional constraints associated with fossoriality, have led to inconsistent
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       morphological recognition of taxa that have confused taxonomic assignments in the past
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       (Kryštufek and Vohralík, 2001; Bannikova et al., 2015; Kryštufek et al., 2018). Cytogenetic
       studies have also shown that the karyotypes of Talpa species are quite stable, with a diploid
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       chromosome number (2n) of 34, and 62-64 autosomal arms (NFa) (Meylan, 1966; Dzuev et
       al., 1972; Todorovic´ et al., 1972; Capanna, 1981; Jimenez et al., 1984; Gornung et al., 2008;
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       Sözen et al., 2012) throughout the genus, with the exception of T. caeca (2n = 36, NFa = 64)
       and T. caucasica (2n = 38, NFa = 62). Additionally it has been shown that Talpa species with
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       almost identical karyotypes (T. europaea and T. romana) differ only slightly in localisation of
       5S rRNA genes (Gornung et al., 2008), further suggesting taxa are not easily distinguished on
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       cytogenetic features.
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       In contrast to this morphological and karyotypic homogeneity, molecular genetic techniques
       have been highly successful in Talpa taxonomy - highlighting the existence of
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       morphologically cryptic, but genetically divergent lineages, which appear to constitute
       species. Bannikova et al. (2015) recently separated three such well-defined lineages in the
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       Caucasus and Anatolia: T. talyschensis Vereschagin, 1945, T. ognevi Stroganov, 1948, and
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       Talpa levantis sensu lato (s.l.). The first of these taxa was formerly considered a junior
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       synonym of T. levantis; the second a junior synonym of T. caucasica (Hutterer, 2005). These
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       findings highlight the fact that lineage diversity within the genus Talpa likely remains under-
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       estimated, particularly in areas known to habour high biodiversity, and which likely served as
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       refugia during Pleistocene glaciations. Turkey is likely to be one such area, since it not only
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       hosts five out of the eleven currently known Talpa species (T. caucasica, T. davidiana, T.
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       europaea, T. levantis, and T. martinorum - Kryštufek and Vohralík, 2009; Kryštufek et al.,
       2018; Selçuk et al., 2018) but it is also the region in which a number of other recent small
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       mammal species/lineages originated (Gündüz et al., 2007). More generally, it is an important
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       biodiversity hotspot in the West Palaearctic due to its complex topography, climatic
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conditions, and tectonic history (Myers et al., 2000). Talpa are strictly subterranean animals,

with limited dispersal capability (Steinberg and Patton, 2000) and a high degree of

territoriality (Ognev, 1928; Stein, 1950; Godfrey, 1957; Loy et al., 1994), and their

diversification is likely to have been strongly impacted by such factors.

The Levant mole was first described from the vicinity of Trabzon (Maçka, Altındere, northeastern Turkey) as a subspecies of the European T. caeca Savi, 1822 – Talpa caeca levantis Thomas, 1906. After being considered as subspecies of T. caeca for a period of time (Spitzenberger and Steiner, 1962; Osborn, 1964; Grulich, 1972), it was elevated to species rank by Spitzenberger (in Felten et al., 1973) based on external measurements and craniometrical variables, especially the relationships between condylobasal length and rostral breadth. Subsequent studies showed that the two species are clearly distinguished on diploid karyotypes (2n = 36 in T. caeca vs. 2n = 34 in T. levantis) and skull morphology (Zima and Král, 1984; Kryštufek, 1994; Selçuk and Kefelioğlu, 2017). Bannikova et al. (2015) showed that these two taxa are not particularly closely related, estimating that they diverged as early as the Late-Middle Pliocene, approximately 2.68 Myr, based on four concatenated nuclear genes. The range of the T. levantis s.l. extends west-east across the Anatolian Black Sea region, from Southeastern Bulgaria and Turkish Thrace as far as the southwestern coast of the Caspian Sea (Osborn, 1964; Doğramaci, 1989; Sokolov and Tembotov, 1989; Vohralík, 1991, Kryštufek, 2001; Popov and Miltchev, 2001; Bannikova et al., 2015). As such, it is the most common and widespread mole in Turkey, and inhabits various habitats from sea level to ca. 2000 m, in areas with rainfall ranging from 1000 to 2500 mm/year (Doğramaci, 1989; Kryštufek, 2001; Popov and Miltchev, 2001; this study). Bannikova et al. (2015) found relatively large cyt-b divergences (7%) between two sublineages within Anatolian and Caucasian T. levantis s.l., suggesting that this taxon, as currently defined, constitutes a pair of cryptic species. Of these sublineages, one occupies the majority of the Turkish range of T. levantis s. 1., throughout the Black Sea coast, westwards to the Marmara region in Anatolia, whilst the other is found in Transcaucasia, and the adjacent parts of northeastern Anatolia. Bannikova et al., (2015) recognized that these two sublineages likely constitute separate species, but refrained from naming them since they lacked genetic data for moles from or close to the type locality of T. levantis, and so could not determine whether this name applied to the eastern or western sublineage.

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Here we use a combination of nuclear and mitochondrial DNA sequence data to revisit the systematics and phylogeography of *T. levantis* s. l., including material from the type locality, in an attempt to stabilize the taxonomy of these moles and better understand their evolutionary history and population expansion.

Material and methods

140	Specimen collection
141	Levant moles were collected from thirteen sites in northern Anatolia between 2007 and 2018
142	(Fig. 1), table 1 summarizes collection details. Standard voucher specimens (skins, skulls and
143	various tissues in ethanol) are deposited in the Department of Biology, Faculty of Sciences
144	and Arts, Ondokuz Mayıs University (OMUS), Samsun, Turkey for long term storage.
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146	Molecular analyses
147	DNA Extraction
148	Total genomic DNA was extracted using Qiagen DNeasy tissue kits (QIAGEN Inc.) from tail
149	tips, kidneys or liver preserved in 95% ethanol. Extracted DNA was suspended in nuclease-
150	free water and DNA concentration quantified using a Nano-Drop spectrophotometer
151	(NanoDrop Technologies, Wilmington, DE), adjusted to 25–50 ng/ μ L, and stored at -20 °C
152	until used for PCR.
153	PCR amplification and sequencing of the cyt-b gene
154	A 1754 bp fragment encompassing a small part of tRNA-Glu (18 bp), the whole of the cyt-b
155	gene (1140 bp), tRNA-Thr, tRNA-Pro (277 bp) and the 5' end (the hypervariable region I) of
156	the D-loop (554 bp, excluding indels), spanning positions 14 163 to 15 913 of the <i>T. europaea</i>
157	mitochondrial genome (GenBank Y19192, Mouchaty et al., 2000) was amplified with primers
158	L14162 5'-GACATGAAAAATCATCGTTG-3' (modified from L14727-SP in Jaarola and
159	Searle, 2002) and H15917 5'-CCTGAAGTAAGAACCAGATG-3' (modified from H16498 in
160	Meyer et al., 1990). PCR amplifications were carried out in an S1000 thermal cycler
161	(BIORAD) using Platinium Taq DNA polymerase (INVITROGEN). The PCR protocol
162	consisted of an initial 2 min denaturation step at 95 $^{\circ}$ C, 35 cycles of denaturation at 94 $^{\circ}$ C for
163	40~s, annealing at 56 °C for 45 s and extension at 72 °C for 1.5 min, and a final 7-min
164	extension step at 72 °C. PCR products were purified using QIAquick kits (QIAGEN).
165	Negative controls were included in all PCRs to check for contamination. After amplification,
166	an aliquot was taken from each PCR reaction mix, and the amplified DNA fragment
167	quantified in agarose gel by comparison with known quantities of phage λ DNA.
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169	The complete cyt-b gene was sequenced in both directions using one of the amplification
170	primers (L14162) plus three internal primers (H15351 5'-
171	TCTCCATTGCTGGTTTACAAGAC-3', modified from H15915 in Irwin et al., 1991 and two

- newly designed Levant mole specific primers, L14711 5'-GGTAGACAAAGCCACACTCAC
 -3' and H14935 5'-GAATGTAGTTGTCTGGATCTCC-3'). The position of the 3' end
 oligonucleotide of each primer is given relative to the published sequence of the common
- mole mtDNA (Mouchaty et al., 2000). Cycle sequencing reactions were carried out using
- 176 BigDye Terminator cycle sequencing kits (Applied Biosystems). Amplifications and
- sequencing reactions were performed in an S1000 thermal cycler. Sequencing products were
- purified using DyeEx 2.0 Spin Kits (QIAGEN) and run in an ABI 3100 automated DNA
- sequencer (Applied Biosystems).
- 180 Nuclear mitochondrial pseudogenes (numts) detection
- 181 It is crucial to avoid confusion of true mitochondrial sequences with copies in the nucleus
- 182 (pseudogenes Mirol et al., 2000; Bensasson et al., 2001; Dubey et al., 2009). To screen for
- pseudogenes, three different PCRs, producing overlapping fragments, were performed on five
- randomly selected individuals from distinct localities. First, partial tRNA-Glu (42 bp), the
- entire cyt-b (1140 bp) and partial tRNA-Thr (26 bp) were amplified with primers L14138 5'-
- 186 CCCACATGGAATTTAACCATGAC-3' (modified from Cb-M1 in Kurose et al., 2000) and
- H15351; second, partial tRNA-Glu (42 bp), the entire cyt-b (1140 bp), complete tRNA-Thr
- 188 (69 bp), complete tRNA-Pro (70 bp) and the 5' end of the D-loop (554 bp) were amplified
- with primers L14138 and H15933; and third, part of tRNA-Glu (18 bp), the entire cyt-b gene
- 190 (1140 bp), complete tRNA-Thr (69 bp), complete tRNA-Pro (70 bp) and the 5' end of the D-
- loop (554 bp) were amplified with primers L14162 and H15933. These three PCR products
- were then sequenced and aligned. Overlapping sequences were examined for any ambiguous
- bases, stop codons or open reading frame shifts, which might have indicated the presence of
- 194 nuclear copies.
- 195 *PCR amplification and sequencing of breast cancer type 2 susceptibility protein (BRCA2)*
- 196 gene
- Amplification of a 927 bp portion of exon 11 of *BRCA2* was performed using the primer pair
- 198 F1140a and R2050 (Bannikova et al., 2015). PCR conditions consisted of initial denaturing at
- 199 94 °C for 2 min; 30 cycles of 94 °C for 45 s, 64 °C for 45 s, and 72 °C for 1 min; and a final
- extension at 72 °C for 6 min. PCR primers were used for sequencing in both directions.
- 201 Sequence inspection and alignment

- All sequence traces were checked, aligned and ambiguous bases resolved by eye using
- Sequencher v4.5 (Gene Codes Corp.). Nucleotide and amino acid composition were analyzed
- using MacClade v4.08 (Maddison and Maddison, 2000).
- 205 Phylogenetic analyses
- Nucleotide composition was analyzed and the frequency of each haplotype estimated using
- 207 MacClade v4.08. To evaluate the degree of differentiation between sublineages of *T. levantis*
- s.l. compared to other species of the genus *Talpa*, and revisit the phylogenetic position of
- these, we also downloaded 78 cyt-b sequences from GenBank, from fourteen Talpa taxa (T.
- altaica, T. aquitania, T. caeca, T. caucasica, T. davidiana, T. europaea, T. levantis 'eastern'
- 211 T. levantis 'western' (sensu Bannikova et al., (2015)), T. martinorum, T. occidentalis, T.
- 212 ognevi, T. romana, T. stankovici and T. talyschensis see Appendix). Phylogenetic
- 213 relationships amongst cyt-b haplotypes were inferred with Maximum Parsimony (MP) and
- Maximum Likelihood (ML) algorithms implemented in PAUP v4.10b (Swofford, 2002) as
- well as Bayesian inference of phylogeny (BI) as implemented in MRBAYES v3.1.2 (Ronquist
- and Huelsenbeck, 2003). The Akaike information criterion (AIC) implemented in
- jMODELTEST v1.0 (Posada, 2008) was used to establish the most appropriate model of
- 218 DNA substitution for our data, and this then employed in ML and BI analyses. The parsimony
- analyses were replicated 10 times with the heuristic search approach using the TBR swapping
- algorithm, steepest descent option and 10 random repetitions. Strict and 50% majority rule
- consensus trees were constructed from equally parsimonious MP trees. Bootstrap analysis of
- the MP tree was conducted with 1000 replications using 10 random repetitions of each
- replication. ML analysis was conducted using the heuristic search, the 'as is' addition
- replicate. Branch support was assessed using 1000 non-parametric bootstrap replicates. BI
- analysis involved four Markov Chains of 20 million generations, with trees being sampled
- every 100 generations and a burn-in of 25%. The software tool TRACER v1.6 (Rambaut et
- al., 2014) was used to observe the parameters and to determine the number of trees needed to
- reach stationarity (burn-in). After discarding burn-in trees and evaluating convergence,
- remaining samples were retained for generating 50% majority rule consensus tree and
- 230 calculating posterior probabilities. Based on the existing supported phylogenetic hypothesis
- 231 (He et al., 2014), cyt-b sequences of four eastern Asian mole species of three genera (Mogera
- 232 robusta Nehring, 1891, Euroscaptor mizura (Günther, 1880), Euroscaptor longirostris
- 233 (Milne-Edwards, 1870) and *Parascaptor leucura* (Blyth, 1850)) were used as outgroups in
- phylogenetic analyses (see Appendix).

235 Relationships amongst cyt-b sequences of T. levantis s.l. were also investigated by 236 constructing a network using the median-joining (MJ) algorithm implemented in the software 237 NETWORK v4.6.1.2 (Bandelt et al., 1999; http://www.fluxus-engineering.com). We included 238 all newly sequenced specimens (24 individuals) and all specimens of *T. levantis* 'western' 239 (seven individuals) and *T. levantis* 'eastern' (five individuals) available in GenBank (Table 1). 240 241 As some of the BRCA2 sequences in GenBank were shorter than ours or contained some 242 unresolved positions, the BRCA2 alignment was limited to 656 bp to match newly obtained 243 haplotypes with published data, leading to some redundant sequences and some loss of 244 245 phylogenetic signal. Phylogenetic relationships between BRCA2 sequences were also 246 investigated using MP (1000 bootstrap replicates), ML (1000 bootstrap replicates) and BI approaches using the settings described above. The nucleotide substitution model selected 247 248 according to the AIC by iMODELTEST was employed in ML and BI analyses. We included all newly described haplotypes (Hap.1-9) and those of *Talpa* species available in GenBank 249 250 (Table 1, Appendix). Unfortunately, two recently described species (*T. aquitania* and *T.* martinorum) could not be included in the analyses since there are no BRCA2 sequences 251 available. The full dataset contained 31 Talpa haplotypes. Again, BRCA2 sequences of the 252 253 same four eastern Asian mole species were used as an outgroup in the phylogenetic analysis 254 (Appendix). 255 Molecular diversity 256 Genetic diversity estimates were calculated for cyt-b sublineage of interest (western). 257 Nucleotide diversity (π) and haplotype diversity (H_d) were calculated using ARLEQUIN v3.5 (Excoffier and Lischer, 2010). Another measure of nucleotide diversity, θw (computed from 258 the number of segregating sites; Watterson, 1975), was calculated using DNASP v5 (Librado 259 260 and Rozas, 2009). DNA net and mean divergences (Da and Dxy; Nei, 1987) between 261 sublineages were estimated under the Kimura 2-parameter (K2P; Kimura, 1980) model using MEGA X (Kumar et al., 2018). 262 263 Divergence time estimation and population demographics We obtained an approximate estimation of the divergence time between T. levantis 264 sublineages using the formula $T = Da/2\mu$, where 2μ is the divergence rate, using a substitution 265 rate for the cyt-b gene of 0.01407 changes per site per lineage per million years (Colangelo et 266

al., 2010). We performed a series of statistical tests to test the hypothesis of sudden population expansion for the two sublineages. First, mismatch distributions (Rogers and Harpending, 1992; Rogers, 1995) were calculated for each sublineage to examine historical changes in population size with ARLEQUIN. To compare observed data with those expected under a sudden expansion model we conducted goodness-of-fit tests based on the sum of squared deviations (SSD) and Harpending's raggedness index (*rg*) (Harpending, 1994; Schneider and Excoffier, 1999) using 10 000 parametric bootstrap replicates.

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The mismatch distribution of pairwise genetic differences was also used to calculate expansion times (Rogers and Harpending, 1992). The change of effective female population size (N) since population expansion is used to estimate the time since expansion in generations. A population at equilibrium with N_0 changes to N_1 at τ units of mutational time. The modal value τ is determined from the distribution of pairwise genetic differences in the extant population. The parameters of the model are given by: $\theta_0 = 2N_0u$, $\theta_1 = 2N_1u$ and $\tau =$ 2ut. Here, t is the time since expansion in generations and u is the mutation rate of the entire DNA fragment (expressed as $u = 2\mu k$, where μ is the mutation rate per nucleotide per generation and k is the length of the sequence). Population expansion times were estimated assuming a mean clock rate of 1.407x10⁻⁸ substitution/site/year, as observed in the genus Talpa (Colangelo et al., 2010). Second, we conducted tests of neutral equilibrium assumptions using two widely used statistics, Tajima's D (Tajima, 1989) and Fu's F_S (Fu, 1997), as additional assessments of possible population expansion. Tajima's D statistic tests the null hypothesis that the average number of pairwise nucleotide differences and number of segregating sites in the sample are equal (Tajima, 1989). This method explicitly tests for selective neutrality and is based on an infinite-sites model assuming no recombination. Fu's Fs statistic is also a selective neutrality test based on the infinite-sites model, assumes no recombination and tests the probability of having no fewer haplotypes than the observed number in the sample (Fu, 1997). Negative values for these two statistics are most often attributed to positive selective sweeps, population size expansion or background selection. Used in combination, the tests can provide evidence for or against particular evolutionary mechanisms. As population size expansion leads to changes in the frequency distribution of haplotypes (i.e. an excess of haplotypes or an excess of singleton mutations), we also calculated Strobeck's S statistic (Strobeck, 1987) and R₂ (Ramos-Onsins and Rozas, 2002). Strobeck's S compares the observed number of haplotypes to that expected based on the

frequency distribution derived from the inferred mutation rate, whereas R_2 is based on the difference between the number of singleton mutations and the average number of nucleotide differences, where lower R_2 values are expected under population expansion. The R_2 test is very powerful in detecting population expansions in small sample sizes (Ramos-Onsins and Rozas, 2002). Significance values for Tajima's D, Fu's F_S and R_2 were obtained from 10 000 coalescent simulations conditioned on theta as implemented in DNASP. The cut-off level for statistical significance was 0.05. For Fu's F_S , significance at the 0.05 level was indicated when P values were < 0.02 (Excoffier and Lischer, 2010).

Results

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- 309 Sequence analysis and phylogenetic reconstructions
- 310 Mitochondrial DNA
- 311 The entire cyt-b gene (1140 bp) was sequenced from 24 Levant moles from 13 localities,
- revealing 15 unique haplotypes (Hap.1-15) with K2P-distance values 0.09-3.34% (overall
- mean K2P distance of 1.54%). Eleven of these haplotypes were found in one individual, four
- in more than one animal (Table 1). No internal stop codons or insertion/deletions were
- detected. Sequences with the same properties were also obtained using the alternative primer
- pairs for amplification and sequencing, making it reasonable to assume that no mitochondrial
- 317 nuclear mitochondrial insertions (numts) were sequenced. The full dataset, including novel
- and published *Talpa* sequences contained 425 variable sites, of which 381 were parsimony
- informative, with 293 synonymous and 32 non-synonymous changes across all taxa. The
- 320 substitution model supported was the General Time Reversible, with specified substitution
- 321 types (AC-1.0052, AG-31.6171, AT-0.7223, CG-0.1456, CT-18.9234, GT-1.0), proportion of
- invariable sites (0.5570), gamma shape parameter (1.1580) and nucleotide frequencies
- 323 (A-0.3572, C-0.3041, G-0.0941, T-0.2446).
- 325 The phylogenetic hypothesis for species of *Talpa* based on cyt-b gene sequences from ML is
- shown in Fig. 2. The different phylogenetic methods used (BI, ML and MP) all produced
- similar topologies. Our tree broadly agrees with previous phylogenetic studies of the genus
- 328 (Colangelo et al., 2010; Bannikova et al., 2015; Kryštufek et al., 2018), including the
- separation of Levant moles into an eastern and western sublineage. Specimens from the type
- locality of *T. levantis* (possessing Haplotype 1), fall into the western sublineage (see Fig. 2).
- The network profile of 26 haplotypes of *T. levantis* is shown in Fig. 3A. This haplotype

network was consistent with the results of the phylogenetic analyses, providing an enhanced 332 333 visualization of intraspecific genetic variation, with many substitutions accumulated between the western and eastern sublineages of *T. levantis* s.l. The two sublineages were 334 connected via two median vectors, separated by 45 mutational steps. There were no shared 335 haplotypes between the two sublineages, suggesting complete lineage sorting. The majority 336 of the haplotypes in the western (Hap.5-15, KP717339, KP717340 and FN640572) 337 sublineage belonged to one group with short to long branches between the haplotypes, 338 connected via median vectors. Three smaller group of haplotypes formed more distant sub-339 340 clusters within the western sublineage (Hap.1-3; FN640571; KP717336 and KP717338), 341 suggesting additional subdivisions within this sublineage. One sub-cluster, comprising 342 haplotypes Hap.1-3 is restricted to north-east Anatolia, including the locality of topotypes of T. levantis (localities 5 and 6), another sub-cluster comprising haplotypes FN640571, 343 344 KP717336 and KP717338 is restricted to north-west Anatolia (localities 21-23), whilst the sub-cluster comprising the remaining 18 haplotypes is distributed across north-central 345 346 Anatolia (localities 7-20) (Fig. 1, Table 1). The eastern sublineage is represented by fewer sequences than the western, but it is nevertheless clear that the haplotype (FN640570) from 347 348 Çam geçidi, Ardahan (locality 4) in northeastern Anatolia is rather strongly differentiated from the four relatively closely related haplotypes (KP717334, KP717335, KP717337 and 349 FN640574) from Russia (Nalchik, locality 1) and Armenia (Fioletovo and Margahovit, 350 localities 2 and 3). 351 352 353 Nuclear DNA Sequences of a 927 bp portion of exon 11of BRCA2 were generated from 24 moles from 14 354 localities, revealing nine unique haplotypes (Hap.1-9), six of which were found in one 355 individual, three in more than one (Table 1). When combined with available (shorter – see 356 Materials and methods) published Talpa BRCA2 sequences, the dataset contained 31 Talpa 357 haplotypes of 656 bp long (see Appendix). This alignment contained 43 variable sites, 22 of 358 359 which were parsimony informative. jMODELTEST supported the Hasegawa-Kishino-Yano (HKY) substitution model for these data, maximum likelihood analysis resulting in an optimal 360 361 which was only partly resolved due to the relatively low number of polymorphic sites (Fig. 4). 362 BI and MP analyses both produced topologies similar to that of ML. As with the cyt-b 363 phylogeny, the BRCA2 data separated T. levantis s.l. into a geographically widespread western sublineage represented by nine haplotypes (Hap.1-9, including Hap.1 from the 364 topotypes of *T. levantis*, locality 5 in Fig. 1) from this study, plus KP717122 from Uzungüney 365

- Köyü, Zonguldak (locality 25 in Fig. 1), and a more restricted eastern sublineage represented
- here by one haplotype only, KP717115 from the Nalchik region in Russia (locality 1 in Fig.
- 368 1).
- 369 *Diversity, divergence time and demographic history*
- 370 The western sublineage of *T. levantis* contained 21 cyt-b haplotypes (15 new in the present
- 371 study, six additional haplotypes from GenBank), resulting in a haplotype diversity (H_d) of
- 372 0.996, nucleotide diversity (π) of 1.894% and a θw of 2.447%. Mean and net divergence
- estimates between cyt-b sequences in eastern and western sublineages of T. levantis were
- 374 7.28% and 5.75%, respectively. This net divergence estimate equates to a separation time of
- 375 ∼1.91 Mya, suggesting that the two sublineages diverged in the early Pleistocene. For the
- western sublineage of *T. levantis*, neutrality tests for cyt-*b* data revealed no significant deviation
- from neutrality and the expected equilibrium. Both Tajima's D (D = -0.9346, P = 0.1765) and
- Fu's Fs (Fs = -0.8062, P = 0.3982) yielded negative but nonsignificant values. Strobeck's S
- was low (0.816) and Ramos-Onsins and Rozas' R_2 test was not significant ($R_2 = 0.0894$, P =
- 380 0.1816). These results were corroborated by a mismatch distribution that showed a wave signal
- 381 (multimodal distribution) consistent with constant population size or demographic equilibrium
- 382 (Fig. 3B). Conversely, the goodness of fit test was consistent with a hypothesis of
- demographical expansion (SSD = 0.0239, P = 0.1241; rg = 0.0165, P = 0.1812). Demographic
- analyses were not possible for the eastern sublineage due to low sample size (n = 5).

Discussion

- 386 Our data confirm that *Talpa levantis* s.l. is divisible into distinct eastern and western
- sublineages, which apparently diverged from each other in the early Pleistocene. Given the
- degree of DNA sequence divergence observed between sublineages, and the fact that these
- remain robust and geographically coherent in light of our more extensive sampling, we agree
- with Bannikova et al. (2015) that these should be considered as separate species. Mean cyt-b
- K2P divergences observed between most mole species examined previously range from 8.6%
- 392 (T. europaea vs. T. occidentalis) to 15.6% (T. altaica vs. T. romana) (Colangelo et al., 2010;
- Feuda et al., 2015). However, slightly lower genetic divergence (7.7%) was observed between
- 394 T. occidentalis and its recently described relative T. aquitana (Nicolas et al., 2017a,b). The
- 395 7.28% cyt-*b* divergence between sublineages revealed here is very close to this value.
- The inclusion of *T. levantis* specimens from the type locality allows us to fix the
- 397 nomenclature of Levant moles for the first time. Topotypical *T. levantis* belong to the western
- sublineage, on both mitochondrial and nuclear data and it is therefore this taxon which is the

true T. levantis sensu Thomas, 1906. As discussed by Bannikova et al. (2015) this therefore 399 400 means that the oldest name available for the eastern clade is *T. transcaucasica* Dahl, 1944. Traditional delineation of species-level entities in morphologically constrained subterranean 401 402 mammals such as moles is intrinsically difficult (see Nevo, 1979), and there are, to date, no known diagnostic morphological characters on which T. levantis and T. transcaucasica can be 403 404 separated. Selçuk et al. (2018) examined skull and mandible morphometrics of selected Turkish mole species, including specimens of *T. levantis* s.l. from both eastern and western 405 406 areas; i.e. both T. levantis s. str. and T. transcaucasica as defined here. Their results reveal 407 relatively high morphological variability in *T. levantis* s.l. compared to the other taxa studied, 408 suggesting that morphometric separation between T. levantis and T. transcaucasica may be 409 possible. Future studies exploring this, together with additional nuclear markers, would clearly be useful to confirm our taxonomic hypothesis. There is no clear geographical barrier 410 411 currently separating T. levantis s. str. and T. transcaucasica, although the species appear to occur either side of the Anatolian Diagonal (sensu Davis, 1971), a biogeographical break 412 413 between central and eastern Anatolia which is observed in a wide variety of organisms (Çıplak, 2003; Gündüz et al., 2007; Mutun, 2010; Kapli et al., 2013; Stümpel et al., 2016; van 414 Riemsdijk et al., 2017). As discussed by Gür (2016), this seems to result largely from 415 differences in temperature seasonality, which apparently also existed during the Last Glacial 416 Maximum and possibly earlier. The identity of *T. levantis* s.l. reported from southeastern 417 Bulgaria and Turkish Thrace remains unclear, due to the absence of molecular data from these 418 regions. A number of small mammal species are known to span the Bosphorus (e.g. Dubey et 419 420 al., 2007, Helvacı et al., 2012) and it is possible that European populations are indeed conspecific with Anatolian ones. Alternatively, what has been considered as T. levantis in 421 422 Europe may belong to yet another distinct taxon, something which should be tested in the 423 future. It is clear, however, that Turkey is a very important centre for mole speciation and diversity, now being home to six distinct species, more than any other comparable 424 geographical region. 425 426 Compared with other mole species, T. levantis s. str. has relatively high haplotype and 427 nucleotide diversity ($H_d = 0.996$, $\pi = 1.894\%$). Based on cyt-b data, Canestrelli et al. (2010) 428 429 estimated haplotype and nucleotide diversities for the six haplogroups of T. romana to be 0.81-0.98 and 0.18-0.77%, respectively. Nicolas et al. (2017b) found almost equally high 430 levels of haplotype diversity but relatively low nucleotide diversity values for three mole 431 species: $H_d = 0.932$, $\pi = 0.371\%$ in *T. europaea*; $H_d = 0.908$, $\pi = 0.830\%$ in *T. aquitania* and 432

 $H_d = 0.892$, $\pi = 0.724\%$ in *T. occidentalis*. The high diversity values observed in *T. levantis* s. 433 str. suggest either the maintenance of relatively large populations throughout the species 434 evolutionary history, and/or its survival in a number of separate refugia during the Pleistocene 435 (Grill et al., 2009; Vega et al., 2010; Nicolas et al., 2017b). Our understanding of Pleistocene 436 palaeoenvironmental history in northern Anatolia remains incomplete, but it is clear that the 437 438 region hosts a large number of microendemics, in a variety of plant and animal taxa (e.g. Davis, 1971; Wielstra et al., 2010; Trizzino et al., 2013), which strongly suggests the 439 occurrence of multiple, long-term refugial areas in the region, as has been hypothesised 440 441 elsewhere (e.g. Lindell et al., 2006; Soltis et al., 2006; Byrne et al., 2008; Canestrelli and Nascetti, 2008; Ursenbacher et al., 2008; Gonçalves et al., 2009; Wang et al., 2009; Vega et 442 al., 2010; Bidegaray-Batista et al., 2016; van Riemsdijk et al., 2017; Wielstra et al., 2017). 443 Genetic substructuring and mutation rate heterogeneity may produce a multimodal pattern, 444 even in demographically unstable populations (Marjoram and Donnelly, 1994; Aris-Brosou 445 and Excoffier, 1996). The multimodality of the mismatch distribution of *T. levantis* s. str. 446 447 seems to result from the presence of different haplogroups, rather than demographical 448 stability. This interpretation is supported by the phylogenetic and network analyses of cyt-b 449 data, which suggest three distinct sub-clusters within this taxon (see Fig. 2 and 3A), perhaps 450 corresponding to populations which have persisted in separate refugial areas which retained suitable habitat during much of the species evolutionary history. If we assume a hypothesis 451 452 involving some degree of population growth for T. levantis s. str., demographic expansion for this taxon, based on our estimated values of τ, occurred approximately 296,000 years BP in 453 454 the late Pleistocene.

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Figure legends

Figure 1. Sampling locations and habitat of *T. levantis* s.l. in Anatolia. A) Map of sampling locations (see Table 1). ● *T. levantis* s.l. 'western', this study; ○ *T. levantis* s.l. 'western'; previous studies and ▲ *T. levantis* s.l. 'eastern', previous studies. Western and Eastern sublineages follow Bannikova et al. (2015). B) Typical habitat of *T. levantis* s.l. at the type locality (Altındere, locality 5); 1650 m a.s.l., Trabzon, Anatolia. Photograph S. Demirtaş.

Figure 2. Results of BI, ML and MP analyses combined on a ML tree based on cyt-b sequences of Talpa and outgroup species. Numbers at nodes indicate posterior probabilities (BI) and bootstrap support values (ML and MP) which are reported only for the key nodes. Bayesian posterior probabilities (≥ 0.90) and bootstrap supports ($\geq 70\%$) are shown. The locality numbers for the haplotypes of 'western' and 'eastern' sublineages are given in parentheses just after the haplotype IDs according to the numbering on map on the Figure 1 and Table 1. For the geographical origins of the published sequences see Table 1 and Appendix.

Figure 3. A) Median-joining network constructed using cyt-*b* sequences of *Talpa levantis* s.l.

The size of each circle is proportional to the frequency of the particular haplotype in the sample.

Median vectors are indicated by blank circles. For the geographical origins of the published sequences see Appendix. B) Mismatch distribution of cyt-*b* sequences for the western sublineage of *Talpa levantis* s.l., showing observed and expected values (see text).

Figure 4. Results of BI, ML and MP analyses combined on a ML tree based on BRCA2 sequences of Talpa and outgroup species. Numbers at nodes indicate posterior probabilities (BI) and bootstrap support values (ML and MP). Bayesian posterior probabilities (≥ 0.90) and bootstrap supports ($\geq 70\%$) are shown. The locality numbers for the haplotypes of 'western' and 'eastern' sublineages are given in parentheses just after the haplotype IDs according to the numbering on map on the Figure 1 and Table 1. For the geographical origins of the published sequences see Table 1 and Appendix.

Locality	Locality	Latitude/longitude	n	GenBank		Reference
	code			Accession No/Haplotype ID		
	-			Cyt-b	BRCA2	
RUSSIA						
Nalchik	1	43°46' N, 43°66' E	2	KP717334	KP717115	Bannikova et al., 2015
				FN640574		Colangelo et al., 2010
ARMENIA						
Fioletovo	2	40°71' N, 44°71' E	1	KP717337		Bannikova et al., 2015
Margahovit	3	40°74' N, 44°70' E	1	KP717335		Bannikova et al., 2015
TURKEY						
Çam geçidi, ARDAHAN	4	41°20' N, 42°55' E	1	FN640570		Colangelo et al., 2010
Taşköprü, Altındere, TRABZON	5	40°64' N, 39°67' E	2	Hap.1	Hap.1	
Görele, GİRESUN	6	41°02' N, 39°02' E	2	Hap.2	Hap.1	
				Hap.3		
Batlama Deresi, GİRESUN	7	40°88' N, 38°34' E	1	Hap.4	Hap.2	
Ulubey, ORDU	8	40°85' N, 37°76' E	2	Hap.5	Hap.3	
				Hap.6	Hap.4	
Geyikçeli Köyü, Fatsa, ORDU	9	40°93' N, 37°37' E	1	Hap.7	Hap.5	
Kurupelit, SAMSUN	10	41°37' N, 36°19' E	3	Hap.8	Hap.6 (2)	

					Hap.7	
İncesu Köyü, SAMSUN	11	41°38′ N, 36°18′ E	1	Hap.9	Hap.8	
Erikli Köyü, SAMSUN	12	41°37' N, 36°05' E	1	Hap.10	Hap.6	
Karakavuk, SAMSUN	13	41°38' N, 36°11' E	1	Hap.10	Hap.6	
Kürtler, SAMSUN	14	41°48' N, 36°09' E	1	FN640572		Colangelo et al., 2010
Dereköy, Ondokuzmayıs, SAMSUN	15	41°45' N, 36°11' E	4	Hap.11	Hap.3 (2)	
				Hap.12		
				Hap.13 (2)	Hap.6 (2)	
Kızılırmak deltası, Bafra, SAMSUN	16	41°67' N, 35°98' E	4	Hap.12 (2)	Hap.6 (3)	
				Hap.13	Hap.1	
				Hap.14		
Uluağaç Köyü, Bafra, SAMSUN	17	41°26' N, 35°58' E	1	Hap.13	Hap.3	
Abalı Köyü, SİNOP	18	42°02' N, 34°98' E	1	Hap.15	Hap.9	
Sefercik, Filyos, ZONGULDAK	19	41°55' N, 32°03' E	2	KP717339		Bannikova et al., 2015
				KP717340		Bannikova et al., 2015
Çaycuma, ZONGULDAK	20	41°43' N, 32°15' E	1	KP717339		Bannikova et al., 2015
Uzungüney Koyu, ZONGULDAK	21	41°37' N, 31°70' E	1	KP717338	KP717122	Bannikova et al., 2015
Alapli, ZONGULDAK	22	41°18' N, 31°40' E	1	KP717336		Bannikova et al., 2015
Uludağ, BURSA	23	40°10′ N, 29°24′ E	1	FN640571		Colangelo et al., 2010