Taxonomic, ecological and morphological diversity of Ponto-Caspian 1 gammaridean amphipods: a review 2 3 Denis Copilas-Ciocianu<sup>1\*</sup> (ORCID: 0000-0002-6374-2365), Dmitry Sidorov<sup>2</sup> (ORCID: 0000-0003-2635-9129) 4 <sup>1</sup>Laboratory of Evolutionary Ecology of Hydrobionts, Nature Research Centre, Vilnius, Lithuania; <sup>2</sup>Federal Scientific Centre 5 6 of the East Asia Terrestrial Biodiversity, Far Eastern Branch of the Russian Academy of Sciences, Vladivostok, Russian 7 Federation 8 \*Corresponding author: denis.copilas-ciocianu@gamtc.lt 9 10 11 Abstract Thanks to its dynamic geological history the Ponto-Caspian region harbors a unique and 12 unusually adaptable fauna, notorious for its invasive species. Gammarid amphipods attained 13 considerable diversity, becoming the world's second most speciose ancient-lake amphipod radiation. 14 Nonetheless, apart from a few invasive species, this group remains poorly studied. Herein, we review 15 and quantify the taxonomic, morphological and ecological diversity, as well as phylogenetic context 16 of Ponto-Caspian gammarids within the adaptive radiation framework. Published molecular 17 phylogenies indicate that this radiation has a monophyletic mid-Miocene Paratethyan origin, and is 18 nested within the morphologically-conserved Atlanto-Mediterranean genus Echinogammarus. We 19 find extensive disparity in body shape, size, ornamentation and appendage length, along a broad 20 ecological gradient from mountain springs to depths exceeding 500 m, on virtually all substrate types 21 (including symbiosis). We propose four putative ecomorphs that appear convergent with distantly 22 related oceanic and Baikal Lake taxa. Thus, the identified patterns support the adaptive radiation 23 model, although extensive further research is needed. A checklist and provisional key to all known 24 endemic species are provided to facilitate taxonomic research. Ponto-Caspian gammarids could be a 25 potentially powerful model for studying adaptive radiations and invasive species evolution. 26 27 Keywords: ancient lake, crustacean, evolution, radiation, systematics 28 29 30 Acknowledgements This study was financed by the Lithuanian Research Council (contract no. 09.3.3-LMT-K-712-19-0149). We thank Nadezhda Berezina for providing important literature. 31 32 33 Introduction 34 Ancient lakes are evolutionary cradles, harboring a rich endemic fauna that fascinated biologists for 35 centuries (M. E. Cristescu et al. 2010; Martens 1997). Their confined nature coupled with large size 36 and relative stability over geological time scales promoted lineage accumulation, diversification and 37 ecological specialization. Many of these lineages probably arose through adaptive radiation, an 38 evolutionary process wherein species rapidly evolve from a common ancestor and diversify to occupy 39 various ecological niches (Schluter 2000). Classical examples of adaptive radiations in ancient lakes 40 are cichlid species flocks in African Rift Valley lakes (Salzburger et al. 2014), or the gammarid 41 amphipods inhabiting Lake Baikal (Naumenko et al. 2017). 42 43 Situated in the Ponto-Caspian region, the Caspian Sea is the world's largest ancient lake (M. E. 44 Cristescu et al. 2010). The Azov, Aral and Black seas are also part of this system. These water bodies 45 are remnants of the once widespread epicontinental Paratethys Sea, which stretched from the foothills of the Alps to the Himalayas (Popov et al. 2004). The Paratethys had a turbulent geological 46 47 history with numerous regression-transgression phases causing drastic salinity fluctuations and 48 repeated episodes of isolation and reconnection with the world ocean (Audzijonyte et al. 2015; Palcu 49 et al. 2019; Popov et al. 2004; Rögl 1999). The uplift of the Caucasus range during the late Miocene

50 triggered the formation and separation of the Black and Caspian seas. During the last two million

51 years these two basins experienced recurrent phases of mutual isolation and reconnection

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54 It is thought that this tumultuous geological past drove the evolution of the unusually euryhaline 55 fauna that inhabits the region today (Reid and Orlova 2002). This plasticity has enabled many Ponto-56 Caspian species to spread across the Northern Hemisphere and become invasive due to human 57 interference (Adrian-Kalchhauser et al. 2020; Cuthbert et al. 2020; Vanderploeg et al. 2002). 58 Nevertheless, many Ponto-Caspian endemics face severe conservation challenges due to climate 59 change, invasive species and multifarious anthropogenic disturbances (Dumont 1995; Gogaladze et 60 al. 2020; Lattuada et al. 2019; Prange et al. 2020). The Ponto-Caspian region is a hot-spot of 61 endemicity and biodiversity with hundreds of species from various animal phyla, but particularly rich 62 in crustaceans (Birstein et al. 1968; Chertoprud et al. 2018; M. E. A. Cristescu and Hebert 2005; 63 Mordukhai-Boltovskoi 1979; Naseka and Bogutskaya 2009; Wesselingh et al. 2019). 64 65 Amphipod crustaceans radiated multiple times in the world's temperate ancient lakes. Several 66 radiations occurred in Lake Titicaca (Hyalellidae) (Adamowicz et al. 2018; Jurado-Rivera et al. 2020), 67 two in Lake Baikal (Gammaridae) (Macdonald et al. 2005; Naumenko et al. 2017), probably two in the 68 Ponto-Caspian basin (Gammaroidea, Corophiidae) (M. E. A. Cristescu and Hebert 2005; Hou et al. 69 2014), and apparently one radiation in other lakes such as Ohrid (Gammaridae) (Wysocka et al. 2013, 70 2014), and Fuxian Hu (Anisogammaridae) (Sket and Fišer 2009). Other lakes throughout Asia also 71 harbor endemic species, although their monophyly has yet to be proven. These are Lake Issyk-Kul in

Kyrgyzstan (Gammaridae) (Karaman and Pinkster 1977) and Lake Teletskoye (Gammaridae) in Russia
(Martynov 1930). In most of these lakes amphipods display a bewildering diversity in form and
ecology, with remarkable convergence in body armature among evolutionary and geographically
distant groups (Martens 1997).

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77 The endemic amphipod fauna of the Ponto-Caspian basin is one of the world's most diverse, second 78 only to Lake Baikal (Barnard and Barnard 1983; Väinölä et al. 2008). Among all endemic Ponto-79 Caspian organisms, amphipods seem to be the most species-rich and successful group, attaining 80 significant ecological and morphological disparity, akin to an adaptive radiation (Derzhavin 1948; 81 Pjatakova and Tarasov 1996; Sars 1895). However, despite these appealing features for evolutionary 82 and ecological studies, Ponto-Caspian amphipods are obscure and poorly known, even ignored in 83 some relatively recent reviews (Martens and Schön 1999). Most attention has been focused on the 84 invasive species that are spreading throughout European freshwaters (e.g. Cristescu et al., 2004; 85 Grabowski et al., 2007; Arbačiauskas et al., 2013; Rewicz et al., 2015), while the non-invasive ones 86 were largely ignored in the last two decades. The taxonomy of the group is rather chaotic due to old 87 and incomplete species descriptions, which led to fuzzy generic diagnoses and lack of a formal 88 system. Even online databases such as World Amphipoda Database 89 (http://www.marinespecies.org/amphipoda/) are incomplete (Horton et al. 2020). Furthermore, a 90 significant part of the literature predates the digital era and is published in Russian, thus not readily 91 available for the international community. As such, to date, there is no comprehensive overview of

92 the Ponto-Caspian amphipod diversity in terms of taxa, ecology and morphology. Some attempts

93 have been made in the past, but these either focused on taxonomy or ecology and never considered

the amphipods from all of the Ponto-Caspian basins (Birstein and Romanova 1968; Mordukhai Boltovskoi 1964, 1979; Pjatakova and Tarasov 1996).

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97 In this study we aim to provide a first comprehensive overview of endemic Ponto-Caspian

98 gammaridean amphipods (taxonomy, morphology and ecology) by examining all of the original

99 species descriptions and relevant literature. Furthermore, by integrating the results of this study with

100 previous phylogenetic research, we strived to identify to which extent the current knowledge on

101 Ponto-Caspian amphipods satisfies the adaptive radiation model (Schluter 2000; Simões et al. 2016).

<sup>52 (</sup>Krijgsman et al. 2019).

Specifically, we looked for evidence pointing to: I) monophyly of endemic Ponto-Caspian gammarids,
 II) an increase in their diversification rates, and III) ecomorphological divergence.

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105 This overview is intended to serve as a foundation and to encourage future eco-evolutionary and 106 taxonomic studies on Ponto-Caspian amphipods. To this end, we also provide a complete checklist 107 and a provisional key to all known endemic species in the hopes of reviving taxonomic interest and to 108 stabilize the systematics of the group.

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## 110 **Taxonomic diversity**

111 Our study focuses on the Ponto-Caspian amphipod taxa that belong to the superfamily 112 Gammaroidea. Specifically, we included the endemic genera of the family Gammaridae, as well as 113 the fully endemic families Behningiellidae, Caspicolidae, Iphigenellidae, and Pontogammaridae. 114 These taxa form the bulk of the endemic diversity and are most likely a monophyletic group (Hou et 115 al. 2014; Sket and Hou 2018), which is a necessary prerequisite for the adaptive radiation model 116 (Schluter 2000). The remaining Ponto-Caspian endemic amphipods such as Chelicorophium (9 spp., 117 Corophiidae), Gammaracanthus (1 sp., Gammaracanthidae), Niphargus (1 sp.), Onisimus (2 spp., 118 Uristidae), and Monoporeia (1 sp.) were excluded since they are unrelated to the focal gammarids 119 (Copilaș-Ciocianu et al. 2020; Lowry and Myers 2017; Väinölä et al. 2001). However, we include the 120 monotypic family Caspicolidae because it is very likely a highly derived gammarid (Derzhavin 1944). 121 Although this family is currently included in the infraorder Talitridira by Lowry & Myers (2013), we 122 consider this placement erroneous due to a character coding mistake (see Discussion for further 123 details).

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125 We compiled a checklist of all known Ponto-Caspian gammaroids by reviewing all of the original 126 species descriptions, including re-descriptions. It is presented in Table 1 along with species 127 systematics, native distribution and short taxonomic remarks where necessary. A total of 82 valid 128 extant species are known, belonging to 34 genera and five families: Behningiellidae (3 genera, 4 129 spp.), Caspicolidae (monotypic), Gammaridae (18 genera, 39 spp.), Iphigenellidae (monotypic) and 130 Pontogammaridae (11 genera, 37 spp.) (Fig. 1a). Five species are doubtful since they may be junior 131 synonyms and further study is needed (Table 1). The most diverse genus is Pontogammarus (8 spp.), 132 followed by Dikerogammarus and Obesogammarus (7 spp. each), Stenogammarus (6 spp.), 133 Chaetogammarus and Amathillina (5 spp. each). Eighteen genera (53%) are monotypic (Fig. 1a). The 134 extinct fossil genera Andrussovia (3 spp.) and Praeqmelina (2 spp.) are currently placed in the 135 Pontogammaridae (Table 1).

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The trend of species description through time reveals little taxonomic activity from the 18<sup>th</sup> to late 19<sup>th</sup> centuries, a sudden increase with Sars' monographs in the late 19<sup>th</sup> century, followed by a more or less steady increase towards the present day with peaks of activity in the middle 20<sup>th</sup> century by Russian and Romanian authors (Fig. 1b). A noticeable stagnation can be observed in the last two decades. By far the most prolific author was G. O. Sars (37 spp.), followed by A. N. Derzhavin (9 spp.) and S. Cărăuşu (8 spp.)(Fig. 1b inset).

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A provisional key to all known endemic families, genera and species (including non-Gammaroidea) is
 provided in the Appendix. We emphasize that some taxa are poorly known and have an uncertain
 generic placement.

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- 151

152 Table 1. Checklist, taxonomy and native distribution of extant and fossil Ponto-Caspian gammaroid

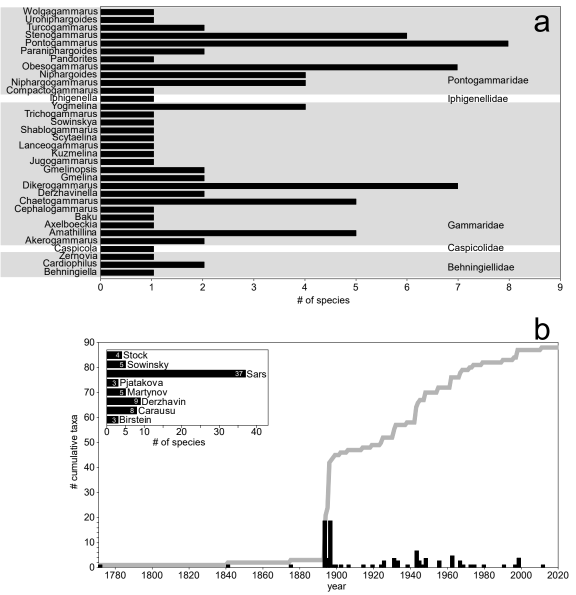
### amphipods.

Family	Species	Authority	Distribution basin	Status	Taxonomic remarks
Behningiellidae	Behningiella brachypus	Derzhavin, 1948	Caspian	valid	
Behningiellidae	Cardiophilus baeri	G.O. Sars, 1896	Caspian	valid	
Behningiellidae	Cardiophilus marisnigrae	Miloslavskaja, 1931	Ponto-Azov	valid	Original spelling: <i>C. maris- nigirs</i> Emended by Barnard (1958) according to ICZN article 32.5.2.3 Synonyms: <i>Cardiophilus</i> <i>miloslavskajae</i> Cărăuşu, 1955
Behningiellidae	Zernovia volgensis	Derzhavin, 1948	Caspian	valid	
Caspicolidae	Caspicola knipovitschi	Derzhavin, 1944	Caspian	valid	
Gammaridae	Akerogammarus contiguus	(Pjatakova, 1962)	Caspian	valid	
Gammaridae	Akerogammarus knipowitschi	Derzhavin & Pjatakova, 1967	Caspian	valid	
Gammaridae	Amathillina affinis	G.O. Sars, 1894	Caspian	valid	
Gammaridae	Amathillina cristata	G.O. Sars, 1894	Caspian, Ponto-Azov	valid	
Gammaridae	Amathillina maximowiczi	G.O. Sars, 1896	Caspian	valid	
Gammaridae	Amathillina pusilla	G.O. Sars, 1896	Caspian	valid	
Gammaridae	Amathillina spinosa	G.O. Sars, 1896	Caspian	valid	
Gammaridae	Axelboeckia spinosa	(G.O. Sars, 1894)	Caspian	valid	
Gammaridae	Baku paradoxus	(Derzhavin in Derzhavin & Pjatakova, 1967)	Caspian	valid	
Gammaridae	Cephalogammarus macrocephalus	(G.O. Sars, 1896)	Caspian	valid	Possibly a derived Dikerogammarus
Gammaridae	Chaetogammarus hyrcanus	Pjatakova, 1962	Caspian	valid	
Gammaridae	Chaetogammarus ischnus	(Stebbing, 1899)	Caspian, Ponto-Azov	valid	Nom. nov. for <i>Gammarus</i> <i>tennelus</i> Sars 1896; Synonyms: <i>Gammarus</i> <i>sowinskyi</i> (Behning, 1914)
Gammaridae	Chaetogammarus pauxillus	(G.O. Sars, 1896)	Caspian	valid	
Gammaridae	Chaetogammarus placidus	(G.O. Sars, 1896)	Caspian, Ponto-Azov	valid	
Gammaridae	Chaetogammarus warpachowskyi	(G.O. Sars, 1894)	Caspian, Ponto-Azov	valid	Reassessment of generic status needed. Close morphological affinities with <i>Gmelina</i> group.
Gammaridae	Derzhavinella cava	Stock, Mirzajani, Vonk, Naderi, Kiabi, 1998	Caspian	valid	
Gammaridae	Derzhavinella macrochelata	Birstein, 1938	Caspian	valid	
Gammaridae	Dikerogammarus aralychensis	(Birstein, 1932)	Caucasus	doubtful	Probably junior synonym of Pontogammarus setosus (Schäferna, 1914). Further research needed.
Gammaridae	Dikerogammarus bispinosus	Martynov, 1925	Caspian, Ponto-Azov	valid	
Gammaridae	Dikerogammarus caspius	(Pallas, 1771)	Caspian	valid	
Gammaridae	Dikerogammarus fluviatilis	Martynov, 1919	Caspian, Ponto-Azov	doubtful	Initially described by Martynov 1919 as a morph o D. haemobaphes. Further

					research needed.
Gammaridae	Dikerogammarus gruberi	Mateus & Mateus, 1990	Ponto-Azov	valid	
Gammaridae	Dikerogammarus haemobaphes	(Eichwald, 1841)	Caspian, Ponto-Azov	valid	Synonym: <i>Dikerogammarus balatonicus</i> Pony, 1958
Gammaridae	Dikerogammarus istanbulensis	Özbek and Özkan, 2011	Ponto-Azov	valid	
Gammaridae	Dikerogammarus oskari	Birstein, 1945	Caspian	valid	Nom. nov. for <i>Gammarus</i> grimmi Sars 1896
Gammaridae	Dikerogammarus villosus	(Sowinsky, 1894)	Caspian, Ponto-Azov	valid	
Gammaridae	Gmelina aestuarica	Cărăușu, 1943	Ponto-Azov	valid	
Gammaridae	Gmelina costata	G.O. Sars, 1894	Caspian, Ponto-Azov	valid	
Gammaridae	Gmelinopsis aurita	G.O. Sars, 1896	Caspian	valid	
Gammaridae	Gmelinopsis tuberculata	G.O. Sars, 1896	Caspian, Ponto-Azov	valid	
Gammaridae	Kuzmelina kusnezowi	(Sowinsky, 1894)	Caspian, Ponto-Azov	valid	
Gammaridae	Jugogammarus kusceri	(S. Karaman, 1931)	Balkan	valid	
Gammaridae	Lanceogammarus andrussowi	(G.O. Sars, 1896)	Caspian, Ponto-Azov	valid	
Gammaridae	Scytaelina simplex	Stock, Mirzajani, Vonk, Naderi, Kiabi, 1998	Caspian	valid	
Gammaridae	Shablogammarus shablensis	(Cărăușu, 1943)	Caspian, Ponto-Azov	valid	Original spelling: Gammarus chablensis Emended by Cărăuşu et al. (1955)
Gammaridae	Sowinskya macrocera	Derzhavin, 1948	Caspian	Valid	
Gammaridae	Trichogammarus trichiatus	(Martynov, 1932)	Ponto-Azov	Valid	We do not agree that Chaetogammarus tennelus major Cărăuşu, 1943 is a junior synonym. Further study is needed.
Gammaridae	Yogmelina brachyura	(Derzhavin & Pjatakova, 1962)	Caspian	valid	
Gammaridae	Yogmelina laeviuscula	(G.O. Sars, 1896)	Caspian	valid	
Gammaridae	Yogmelina limana	G.S. Karaman & Barnard, 1979	Ponto-Azov	valid	
Gammaridae	Yogmelina ovata	(Martynov, 1924)	Ponto-Azov	doubtful	Probably junior synonym of Chaetogammarus warpachowskyi. Further study is needed.
Gammaridae	Yogmelina pusilla	(G.O. Sars, 1896)	Caspian, ?Ponto-Azov	valid	
Iphigenellidae	Iphigenella acanthopoda	G.O. Sars, 1896	Caspian, Ponto-Azov	valid	
Pontogammaridae	Compactogammarus compactus	(G.O. Sars, 1895)	Caspian, Ponto-Azov	valid	
Pontogammaridae	Niphargogammarus aequimanus	(G.O. Sars, 1895)	Caspian	valid	
Pontogammaridae	Niphargogammarus borodini	(G.O. Sars, 1897)	Caspian	valid	
Pontogammaridae	Niphargogammarus intermedius	(Cărăușu, 1943)	Ponto-Azov	valid	
Pontogammaridae	Niphargogammarus quadrimanus	(G.O. Sars, 1895)	Caspian	valid	
Pontogammaridae	Niphargoides boltovskoyi	Derzhavin & Pjatakova, 1968	Caspian	valid	
Pontogammaridae	Niphargoides caspius	G.O. Sars, 1894	Caspian	valid	
Pontogammaridae	Niphargoides	G.O. Sars, 1894	Caspian,	valid	1

	corpulentus		Ponto-Azov		
Pontogammaridae	Niphargoides grimmi	G.O. Sars, 1896	Caspian	valid	
Pontogammaridae	Obesogammarus acuminatus	Stock, Mirzajani, Vonk, Naderi, Kiabi, 1998	Caspian	valid	
Pontogammaridae	Obesogammarus boeoticus	(Schellenberg, 1944)	Balkan	valid	Poorly described, generic status needs revision
Pontogammaridae	Obesogammarus crassus	(G.O. Sars, 1894)	Caspian, Ponto-Azov	valid	Synonyms: Obesogammarus crassus f. mediodanubialis S. Karaman, 1953
Pontogammaridae	Obesogammarus obesus	(G.O. Sars, 1894)	Caspian, Ponto-Azov	valid	
Pontogammaridae	Obesogammarus olvianus	(Sowinsky, 1902)	Caspian, Ponto-Azov	valid	
Pontogammaridae	Obesogammarus platycheir	(G.O. Sars, 1896)	Caspian	valid	
Pontogammaridae	Obesogammarus subnudus	(G.O. Sars, 1896)	Caspian, Ponto-Azov	valid	Provisionally placed in Obesogammarus. Generic status needs revision.
Pontogammaridae	Pandorites podoceroides	(G.O. Sars, 1896)	Caspian	valid	
Pontogammaridae	Paraniphargoides derzhavini	Pjatakova, 1962	Caspian	valid	
Pontogammaridae	Paraniphargoides motasi	(Cărăușu, 1943)	Caspian, Ponto-Azov	valid	
Pontogammaridae	Pontogammarus abbreviatus	(G.O. Sars, 1894)	Caspian, Ponto-Azov	valid	
Pontogammaridae	Pontogammarus aestuarius	(G.O. Sars, 1894)	Caspian, Ponto-Azov	valid	
Pontogammaridae	Pontogammarus borceae	Carausu, 1943	Caspian, Ponto-Azov	valid	
Pontogammaridae	Pontogammarus palmatus	(Martynov, 1925)	Caspian	doubtful	Poor description without illustrations.
Pontogammaridae	Pontogammarus robustoides	(G.O. Sars, 1894)	Caspian, Ponto-Azov	valid	
Pontogammaridae	Pontogammarus setosus	(Schäferna, 1914)	Caucasus	valid	
Pontogammaridae	Pontogammarus (=Euxinia) maeoticus	(Sowinsky, 1894)	Caspian, Ponto-Azov	valid	Removed to <i>Pontogammarus</i> since <i>Euxinia</i> Tucolesco, 1933 is preoccupied by the platyhelminth genus <i>Euxinia</i> Graff, 1911. According to ICZN articles 23.1 and 52.4 Synonyms: <i>Euxinia fagei</i> Tucolesco, 1933
Pontogammaridae	Pontogammarus (=Euxinia) sarsi	(Sowinsky, 1898)	Caspian, Ponto-Azov	valid	Removed to <i>Pontogammarus</i> since <i>Euxinia</i> Tucolesco, 1933 is preoccupied by the platyhelminth genus <i>Euxinia</i> Graff, 1911. According to ICZN articles 23.1 and 52.4
Pontogammaridae	Pontogammarus (=Euxinia) weidemanni	(G.O. Sars, 1896)	Caspian, Ponto-Azov	valid	Removed to <i>Pontogammarus</i> since <i>Euxinia</i> Tucolesco, 1933 is preoccupied by the platyhelminth genus <i>Euxinia</i> Graff, 1911. According to ICZN articles 23.1 and 52.4
Pontogammaridae	Stenogammarus compressus	(G.O. Sars, 1894)	Caspian, Ponto-Azov	valid	
Pontogammaridae	Stenogammarus compressosimilis	Cărăușu 1955	Caspian, Ponto-Azov	valid	Original spelling: S. compresso-similis Emended according to ICZN article 32.5.2.3 Synonyms: Stenogammarus

					<i>kereuschi</i> Derzhavin & Pjatakova, 1962 according to ICZN article 23.3.7
Pontogammaridae	Stenogammarus deminutus	(Stebbing, 1906)	Caspian, Ponto-Azov	valid	Nom. nov. for <i>Gammarus</i> minutus G.O. Sars 1894
Pontogammaridae	Stenogammarus macrurus	(G.O. Sars, 1894)	Caspian, Ponto-Azov	valid	
Pontogammaridae	Stenogammarus micrurus	Derzhavin & Pjatakova, 1996	Caspian	valid	
Pontogammaridae	Stenogammarus similis	(G.O. Sars, 1894)	Caspian, Ponto-Azov	valid	
Pontogammaridae	Turcogammarus aralensis	(Uljanin, 1875)	Aral, Caspian, Ponto-Azov	valid	
Pontogammaridae	Turcogammarus spandli	(Karaman, 1931)	Balkan	valid	
Pontogammaridae	Turcogammarus turcarum	(Stock, 1974)	Caucasus	doubtful	Probably junior synonym of <i>Pontogammarus setosus</i> (Schäferna, 1914). Further research needed.
Pontogammaridae	Uroniphargoides spinicaudatus	(Cărăușu, 1943)	Caspian, Ponto-Azov	valid	
Pontogammaridae	Wolgagammarus dzjubani	(Mordukhai- Boltovskoi & Ljakhov, 1972)	Caspian	valid	
Fossil taxa					
Pontogammaridae	Andrussovia bogacevi	Derzhavin, 1927†	Caspian	valid	
Pontogammaridae	Andrussovia sokolovi	Derzhavin, 1927†	Caspian	valid	
Pontogammaridae	Andrussovia vassoievitschi	Derzhavin, 1941†	Caspian	valid	
Pontogammaridae	Praegmelina andrussovi	Derzhavin, 1927†	Caspian	valid	
Pontogammaridae	Praegmelina archangelskii	Derzhavin, 1927†	Caspian	valid	
Hellenidae	Hellenis saltatorius	Petunnikoff, 1914†	Caspian	valid	Uncertain affiliation with extant Ponto-Caspian taxa



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Fig. 1 a) Species richness within genera and families. Only valid and extant species were considered. b) Trends
in species descriptions through time. The thick gray line indicates the cumulative number of species while black
bars indicate the number of species described in that respective year. The inset graph depicts the number of
species described by the most prominent authors

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# 163 Morphology

164 To explore morphological diversity we extracted data only from those original species descriptions or 165 re-descriptions that provided good quality habitus illustrations (73% of all species) (Cărăuşu 1943; 166 Cărăuşu et al. 1955; Derzhavin 1944, 1948; Sars 1894a, 1894b, 1895, 1896). This was necessary 167 because we used the ratios of various body parts and appendages to total body length. In total, we 168 calculated ratios for 53 traits reflecting as much as possible the overall body shape and functional 169 morphology (see Supplementary information Tables S1-S2 and Fig. S1) (Fišer et al. 2009). The ratios 170 were measured using the Digimizer software (https://www.digimizer.com/). Whenever possible, 171 both sexes were included. We acknowledge that these illustration obtained ratios do not provide the 172 most exhaustive nor precise morphological detail. However, given that these data have a broad

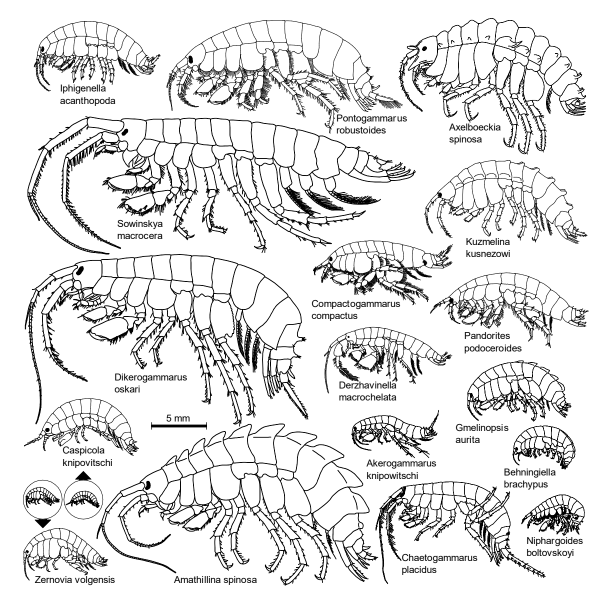
taxonomic coverage, we consider this analysis as a crucial preliminary step in quantifying andunderstanding the morphological diversity of Ponto-Caspian amphipods.

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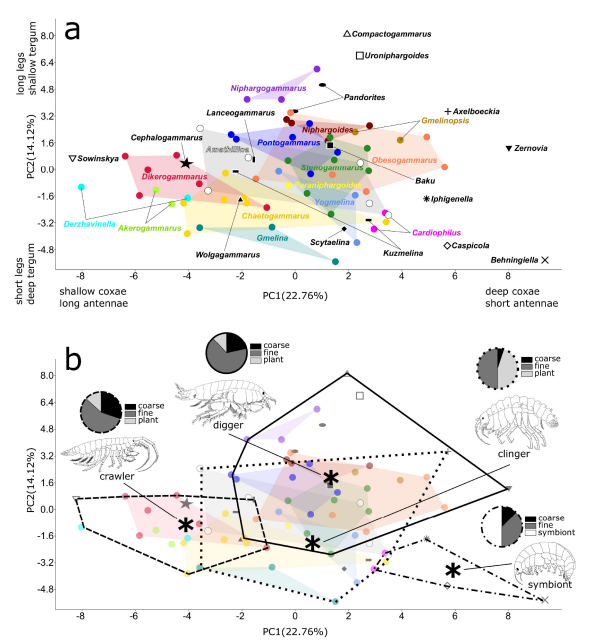
Apart from ratios, we also extracted body size information from the literature and included it in the
analysis as well. The 53 ratio + body-size dataset was subjected to a Principal Component Analysis
(PCA) based on a correlation matrix to visualize morphological gradients and similarity among
genera. Analysis was performed using Statistica 8.0 (StatSoft, Inc.,Tulsa, OK, USA).

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181 We find substantial diversity in body shape and size. The habitus of representative species is presented in Fig. 2. Body size varies by almost an order of magnitude (3.5 to 27 mm) (Figs. 2, 6). The 182 183 PCA plot indicates significant morphological disparity (Fig. 3). The first four PCA axes explained 22.76, 184 14.12, 10.17 and 9.28% (56.34%) of the total variation. The first principal component separated 185 species along a gradient from stout bodies with deep coxae and short antennae to slender bodies 186 with shallow coxae and long antennae (Fig. 3a). The second principal component distinguished a gradient along which species were separated by the length of walking appendages and the depth of 187 188 the tergum (Fig. 3a). The loadings of traits on the PCA axes are presented in Supplementary 189 information Table 3.



192 193 Fig. 2 Habitus and morphological diversity of Ponto-Caspian amphipods. Caspicola knipovitschi and Zernovia 194 volgensis are shown to scale in circles and enlarged outside the circles. All images are redrawn after the original 195 196 197 There is significant variation with respect to body armature as well. Although most species are 198 generally smooth, there are diverse patterns of ornamentation with either a medial keel that extends throughout different body regions (e.g. Amathillina, Gmelina and Gmelinopsis) to double dorso-199 200 lateral cuspidation (Kuzmeling), to lateral spines and dorsal protuberances (Axelboeckia) (Fig. 2). 201 202 Most genera seem to be relatively well defined in morphospace. However, Amathillina and 203 Obesogammarus overlap broadly with other genera (Fig. 3a). The monotypic genera (shown with 204 black and white symbols in Fig. 3a) are generally distinct from the more speciose ones, often lying 205 towards the extreme ends of the morphological gradients. 206 207 208 209



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Fig. 3 a) PCA scatterplot depicting the morphological gradients along the first two axes. Genera represented by
at least three data points are shown with a uniquely colored convex hull and dots. Monotypic genera are
depicted with various black symbols and shapes. b) The same PCA as in a) but with convex hulls delineating
putative ecomorphs. Asterisks indicate morph centroid. For each morph a representative species is shown. The
pie-charts indicate the proportion of species occurring on various substrates within each ecomorph

- 216
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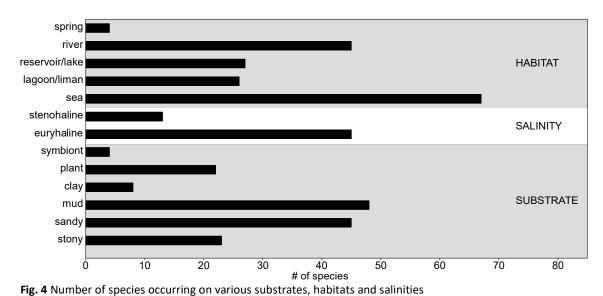
### 218 Ecology

To provide a synopsis of ecology we reviewed all the original species descriptions and the relevant literature (Birstein and Romanova 1968; Pjatakova and Tarasov 1996). We gathered data regarding

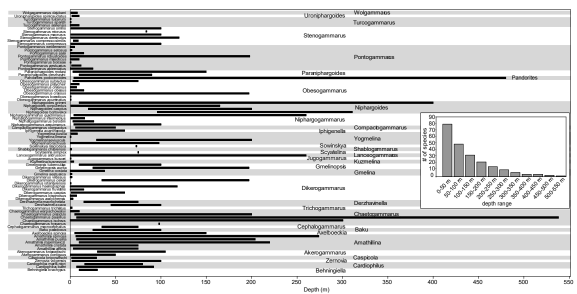
- 221 depth (minimum and maximum), habitat (sea, lagoon, lake/reservoir, river and spring), salinity
- 222 (steno- and/or euryhaline) and substrate type (stone, sand, mud, clay, plant and symbiotic
- 223 relationships).

### 

Our review highlights important ecological diversity within the Ponto-Caspian radiation. With respect to habitat, most species live in the sea (67 spp.) and lower courses of rivers (45 spp.), followed by brackish lagoons (26 spp.) and freshwater lakes or reservoirs (27 spp.). Only four species occur exclusively in springs and streams (Table 2, Fig. 4). With respect to salinity, it appears that most species are euryhaline, tolerating both fresh as well as brackish waters. However, salinity preference is not known for many species. With respect to substrate, the great majority of species occur on sandy and muddy substrates, followed by stones and plants. Four species seem to be associated with other organisms such as bivalve mollusks and crayfish (Table 2, Fig. 4). All of the ecological data is summarized in Table 2.



The depth gradient is broad, ranging from the wet sand of the supra-littoral to more than 500 m
depth (Table 2, Fig. 5). Individual species also seem to be quite plastic and can be found from shallow
depths (less than 50 m) to more than 200 m. The genera *Amathillina, Chaetogammarus, Niphargoides* and *Pandorites* have the broadest depth ranges. Species diversity is the highest in the
first 50 m (79 species), then rapidly decreases to below 10 species in the 250-550 m interval (Fig. 5).
The only species known to occur at depths greater than 500 m is *Chaetogammarus pauxillus*.



262

Fig. 5 Depth ranges structured by taxonomic composition. The inset graph depicts the number of species occurring in 50 m depth intervals

Table 2. Ecological diversity of Ponto-Caspian gammaroid amphipods.

	Min	Max	Substrate												
Species	depth	depth	sto1	san²	mud	cla <sup>3</sup>	pla⁴	symbiont	Salinity	sea	lag⁵	res <sup>6</sup>	riv <sup>7</sup>	spr <sup>8</sup>	Ecomorph
Behningiella brachypus	10	30	x	x					?	x					symbiont
Cardiophilus baeri	7	92			?	?		Cardium	?	х					symbiont
Cardiophilus maris- nigri	16	80		x		x		Mytilus	?	x					symbiont
Zernovia volgensis	2	100							?	х			х		digger
Caspicola knipovitschi	1.5	29		x				possibly molluscs	euryh.	x			x		symbiont
Akerogammarus contiguus	0.2	50	x	x	x				?	x					crawler
Akerogammarus knipowitschi	30	105			x				?	x					crawler
Amathillina affinis	3.5	75					х		euryh.	х					clinger
Amathillina cristata	0.5	75					х		euryh.	х	х	x	х		clinger
Amathillina maximowiczi	10	220					?		?	x	x				clinger
Amathillina pusilla	7	204					х		?	х					clinger
Amathillina spinosa	6.4	274		x	x		?		euryh.	х					clinger
Axelboeckia spinosa	5	150			?		х		euryh.	х			x		clinger
Baku paradoxus	25	100			x				?	x					digger
Cephalogammarus macrocephalus	35	75			x				?	x					crawler
Chaetogammarus hyrcanus		98			x				?	x					crawler
Chaetogammarus ischnus	0	300	x	x	x		x		euryh.	x	x	x	x		crawler
Chaetogammarus pauxillus	7	538		x					euryh.	x					crawler

		1	1	1		1		1			1		1	1	1
Chaetogammarus placidus	5	200			x				euryh.	х			x		crawler
Chaetogammarus warpachowskyi	0.5	10		x	x		x		euryh.	x	x	x	x		clinger
Derzhavinella cava	45	100							?	х					crawler
Derzhavinella macrochelata	15	75			x				?	x					crawler
Dikerogammarus aralychensis	0.2	3		x	x				stenoh.			x	x		digger
Dikerogammarus bispinosus	0	7	x	x					euryh.	x		x	x		crawler
Dikerogammarus caspius	1.2	60	x						euryh.	x		x	x		crawler
Dikerogammarus fluviatilis	0	15	x	x	x		x		euryh.				x		crawler
Dikerogammarus gruberi									?			x			?
Dikerogammarus haemobaphes	0	118	x	x	x				euryh.	x	x	x	x		crawler
Dikerogammarus istanbulensis	0.2	1			x				?			x			crawler
Dikerogammarus oskari	35	197			x				?	x					crawler
Dikerogammarus villosus	0	2	x				x		euryh.	x	x	x	x		crawler
Gmelina aestuarica	0	2		x	x		x		euryh.		х		x		clinger
Gmelina costata	0	38	x	x			х		euryh.	х			х		clinger
Gmelinopsis aurita	25	197							?	х					digger
Gmelinopsis tuberculata	10	100	x						euryh.	x	x		x		digger
Kuzmelina kusnezowi	1	4			x				euryh.	x	x		x		clinger
Jugogammarus kusceri	0	1	x	x					stenoh.					x	clinger
Lanceogammarus andrussowi	0.5	260		x			x		euryh.	x	x	x			crawler
Scytaelina simplex		75							?	х					clinger
Shablogammarus shablensis	0	1.5			x		x		euryh.		x	x	x		crawler
Sowinskya macrocera		73			x				?	x					crawler
Trichogammarus trichiatus	0	2	x	x					euryh.		x	x	x		crawler
Yogmelina brachyura	0.3	98		x	x		x		?	x					clinger
Yogmelina laeviuscula	29	200			x				?	x					clinger
Yogmelina limana		1		x	x		x		euryh.		х		?		clinger
Yogmelina pusilla	1	8		x	x		x		euryh.	x	x		?		clinger
Iphigenella acanthopoda	1	60						possibly Astacidae	euryh.	x	x	x	x		symbion
Compactogammarus compactus	6	50		x	x				euryh.	x	x		x		digger
Niphargogammarus aequimanus	4	100							stenoh.	x					digger
Niphargogammarus borodini	3	26							?	x					digger
Niphargogammarus intermedius	1	16		x	х	x			euryh.				x		digger

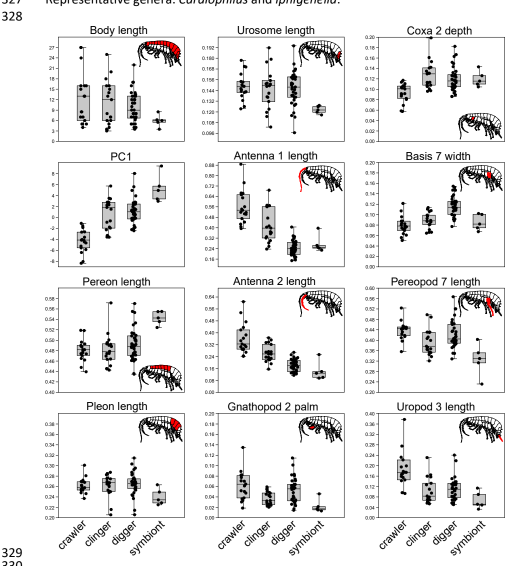
Niphargogammarus quadrimanus	4	260			x			stenoh.	x					digger
Niphargoides boltovskoyi	96	311			x			?	x					digger
Niphargoides caspius	20	200			x			stenoh.	x					digger
Niphargoides corpulentus	1	165			x			euryh.	х			x		digger
Niphargoides grimmi	10	400			x			stenoh.	x					digger
Obesogammarus acuminatus	0.2	1					x	euryh.	x	x				digger
Obesogammarus boeoticus	0.2	1	x	x				stenoh.			x			digger
Obesogammarus crassus	0.2	197	x	x	x		x	euryh.	х	x	x	x		digger
Obesogammarus obesus	0.5	15	x	x			x	euryh.	x	x	x	x		digger
Obesogammarus olvianus	0.5	7			x			euryh.	х					digger
Obesogammarus platycheir	1	10		x				euryh.	х			x		digger
Obesogammarus subnudus	3	75		x				?	x			x		digger
Pandorites podoceroides	4	480		x	x			euryh.	х					digger
Paraniphargoides derzhavini	10	90		x	x			?	х					digger
Paraniphargoides motasi	3	150		x	x			?				x		digger
Pontogammarus abbreviatus	0.5	25		x	x	x		euryh.	х		x	x		digger
Pontogammarus aestuarius	1.5	12	x	x	x			euryh.	х	x		x		digger
Pontogammarus borceae	0	0.1		x				euryh.	х	x		x		digger
Pontogammarus maeoticus	0	10		x				euryh.	х	x	x	x		digger
Pontogammarus robustoides	0	198	x	x	x		x	euryh.	х	х	x	x		digger
Pontogammarus sarsi	0.5	15		x	x	x		euryh.		x	x	x		digger
Pontogammarus setosus	0	2	x	x	x		x	stenoh.				x	х	digger
Pontogammarus weidemanni	0	5		x				euryh.	х	x				digger
Stenogammarus compressus	1	100			x			euryh.	x		x	x		digger
Stenogammarus compresso-similis	3.5	9	x	x	x	x		euryh.	x			x		digger
Stenogammarus deminutus	1	120						euryh.	х		x	x		digger
Stenogammarus macrurus	1	100	x	x	x	x		euryh.	x		x	x		digger
Stenogammarus micrurus	84	84			x			?	x					digger
Stenogammarus similis	1	100	x	x	x	x	x	stenoh.	x		x			digger
Turcogammarus aralensis	0	10	x	x				euryh.	x		x	x		digger
Turcogammarus spandli	0	0.5	x					stenoh.				x	х	crawler

Turcogammarus turcarum	0	1	x	x				stenoh.				x	crawl
Uroniphargoides spinicaudatus	2	10		x	x			stenoh.	x		x		digge
Wolgagammarus dzjubani	1	8		x				stenoh.		x	x		digge
	ny, ²–sai	ndy, <sup>3</sup> –cla	ay, ⁴–p	lant, <sup>5</sup>	–lagoc	on, <sup>6</sup> –r	servoir/lake	, <sup>7</sup> –river, <sup>8</sup> –spring	5				
270													
271													
272 <b>Pro</b>	posed	ecomo	rphs										
273 By in	tegrating	g morpho	ology a	and sul	bstrate	e type	we aimed to	classify the spec	ies into	putati	ve		
274 econ	horphs. S	pecifical	ly, we	looked	d for c	ommo	n morpholog	gical characteristi	cs amor	ng taxa	a, whil	е	
275 takin	g into ac	count th	eir sin	nilarity	in PC/	A mor	hospace. W	e also took into a	ccount	previo	us inf	ormal	
276 grou	pings of a	genera (I	Barnar	d and	Barna	rd 198	3). Once the	se groups were id	dentified	l, thei	r subs	trate	
277 prefe	erence w	as estab	lished	by est	imatin	g the	roportion of	f species occurrin	ig on a p	articu	lar		
278 subs	trate. Th	e substra	ate cla	ssificat	tion wa	as sim	lified and di	vided into four g	roups: c	oarse			
279 (corr	espondir	ng to sto	nes an	d grav	el), fin	e (cor	esponding t	o sand, mud and	clay), pl	ant ar	nd		
280 symb	oiotic. W	e acknov	vledge	that t	his is a	some	what arbitra	ry approach. How	vever, n	nore			
281 soph	isticated	analyse	s could	l not b	e perf	ormed	given the so	arce data at han	d. Quan <sup>.</sup>	titativ	e data	Ì	
282 rega	rding ecc	ology (sul	bstrate	e or tro	ophic r	niche)	re only limit	ed to a few inva	sive spe	cies. L	ikewis	ie,	
283 morp	ohology i	s incomp	letely	know	n in ma	any sp	ecies (especi	ally mouthparts)	. We em	phasi	ze tha	t our	
284 goal	here was	s to prov	ide a f	irst ex	plorate	ory ste	o into under	standing the con	nection	betwe	en		
285 morp	hology a	and ecolo	ogy.										
286													
287 Wet	entative	ly define	d four	ecom	orphs:	clinge	s, crawlers,	diggers and syml	oionts. L	.oosel	, the	se	
		-			•	-		ies and informal			-		
								marids" + "Dikerd					
290 Barn	ard & Ba	rnard, 19	983), c	lingers	s with	Gamn	aridae or "G	melinids" ( <i>sensu</i>	Barnard	l & Ba	rnard,		
291 1983	), digger	s with Po	ontoga	mmar	idae o	r "Pon	ogammarid	s" + "Compactog	ammario	ds" ( <i>se</i>	ensu		
292 Barn	ard & Ba	rnard, 19	983), a	nd syr	nbiont	s with	Behningielli	dae, Caspicolidae	and Ipł	nigene	llidae	or	
								describe the mo					
	ogical pe	-											
295													
296 1) Cli	nger. Sto	out body	often	keeled	d and/	or orn	mented wit	h spines and tub	ercles, a	ntenn	ae are	ė	
297 slend	ler, shor	t to med	ium le	ngth, c	coxal p	lates	nedium to de	eep, gnathopods	weak, a	nd pe	reopo	ds	
298 short	t to medi	ium leng	th witl	n pairs	3-4 st	rongly	opposable t	o pairs 5-7 (Fig. 6	i). Clinge	ers are			
299 inter	mediate	in morp	hospa	ce betv	ween o	rawle	s and digger	s, although there	e is signi	ficant	overla	ар	
300 signi	ficantly v	vith the l	atter g	group	(Fig. 3l	o). Mc	st species ar	e associated with	plants	and fir	ne sub	strate	
	2h) Arou	und 19%	of all :	species	s belor	ng to t	is ecomorpl	h Tavonomic cor	npositio	n is gi	ven in	Table	
-	3D). AI UI			volhaa	ckia a	nd Cm		1. 10,011011110 001					
301 (Fig.	presenta		era: A	xeibbe	chiù ai	iu Gii	elina.						
301 (Fig.	-		era: A	xeibbe		iu Gii	elina.						
301 (Fig. 302 2. Re 303	presenta	ative gen						e are long and sle	nder, co	oxal pl	ates		
301 (Fig. 302 2. Re 303 304 2) Cr	presenta awler. Bo	ative gen ody is sle	nder a	and ge	nerally	, smoo	h, antennae					Fig.	
301 (Fig. 302 2. Re 303 304 2) Cr 305 shall	presenta awler. Bo ow, pere	ative gen ody is sle opods sl	nder a ender,	and ge , short	nerally to me	' smoo dium,	h, antennae gnathopods	e are long and sle	and uro	pods	long (	•	
301 (Fig. 302 2. Re 303 304 2) Cr 305 shall 306 6). It	presenta awler. Bo ow, pere is genera	ative gen ody is sle opods sl ally well-	nder a ender define	and ge , short ed in m	nerally to me iorpho	' smoo dium, space	th, antennae gnathopods naving little	e are long and sle generally strong,	and uro gers and	opods I digge	long ( ers (Fig	g. 3b).	
301       (Fig.         302       2. Re         303       304         304       2) Cr         305       shall         306       6). It         307       Spec	presenta awler. Bo ow, pere is genera ies are m	ative gen ody is sle opods sl ally well- nainly ass	nder a ender, define sociate	and ge , short ed in m ed with	nerally to me orpho fine a	smoo dium, space and co	th, antennae gnathopods naving little rse substrat	e are long and sle generally strong, overlap with clin	and uro gers and ind 26%	opods d digge of spe	long ( ers (Fig	g. 3b).	
301       (Fig.         302       2. Re         303       304         304       2) Cr         305       shall         306       6). It         307       Spec         308       to th	presenta awler. Bo ow, pere is genera ies are m	ative gen ody is sle opods sl ally well- nainly ass orph. Tax	nder a ender, define sociate	and ge , short ed in m ed with ic com	nerally to me orpho fine a positic	smoo dium, space and co on is g	th, antennae gnathopods naving little rse substrat	e are long and sle generally strong, overlap with clin es (Fig. 3b). Arou	and uro gers and ind 26%	opods d digge of spe	long ( ers (Fig	g. 3b).	
301       (Fig.         302       2. Re         303       304         304       2) Cr         305       shall         306       6). It         307       Spec         308       to th         309       Chae	presenta awler. Bo ow, pere is genera ies are m is ecomo	ative gen ody is sle opods sl ally well- nainly ass orph. Tax	nder a ender, define sociate	and ge , short ed in m ed with ic com	nerally to me orpho fine a positic	smoo dium, space and co on is g	ch, antennae gnathopods naving little rse substrat	e are long and sle generally strong, overlap with clin es (Fig. 3b). Arou	and uro gers and ind 26%	opods d digge of spe	long ( ers (Fig	g. 3b).	
301       (Fig.         302       2. Re         303       304         304       2) Cr         305       shall         306       6). It         307       Spec         308       to th         309       Chae         310	awler. Bo ow, pere is genera ies are m is ecomo togamm	ative gen ody is sle opods sl ally well- nainly ass orph. Tax parus and	nder a ender, define sociate onom I <i>Diker</i>	and gei , short ed in m ed with ic com rogami	nerally to me orpho fine a positio marus	dium, space and co on is g	ch, antennae gnathopods naving little rse substrat ven in Table	e are long and sle generally strong, overlap with clin es (Fig. 3b). Arou 2. Representativ	and uro gers and Ind 26% e genera	opods I digge of spe a:	long ( ers (Fig ecies b	g. 3b). pelong	
301       (Fig.         302       2. Re         303       304       2) Cr         305       shall         306       6). It         307       Spec         308       to th         309       Chae         310       311	presenta awler. Bo ow, pere is genera is are m is ecomo togamm gger. Sto	ative gen ody is sle opods sl ally well- nainly ass orph. Tax <i>parus</i> and out body	nder a ender, define sociate onom I <i>Diker</i> and al	and get , short ed in m ed with ic com ogami most e	nerally to me orpho fine a positio marus exclusi	dium, space and co on is g vely si	ch, antennae gnathopods naving little rse substrat ven in Table nooth, anter	e are long and sle generally strong, overlap with clin es (Fig. 3b). Arou 2. Representativ	and uro gers and ind 26% e genera nd thick	opods d digge of spe a: , with	long ( ers (Fig ecies b 1 <sup>st</sup> art	g. 3b). belong cicle of	
301       (Fig.         302       2. Re         303       304       2) Cr         305       shall         306       6). It         307       Spec         308       to th         309       Chae         310       311         312       ante	presenta awler. Bo ow, pere is genera is are m is ecomo togamm gger. Sto nna 1 oft	ative gen ody is sle opods sl ally well- nainly ass orph. Tax parus and out body cen swoll	nder a ender, define sociate onom <i>Diker</i> and al en, co	and get , short ed in m ed with ic com rogami most e xal pla	nerally to me orpho fine a positio marus exclusi tes de	v smoo dium, space ind co on is g vely si ep, gn	th, antennae gnathopods naving little rse substrat ven in Table nooth, anten athopods ge	e are long and sle generally strong, overlap with clin es (Fig. 3b). Arou 2. Representativ	and uro gers and ind 26% e genera nd thick ereopod	opods d digge of spe a: , with s med	long ( ers (Fi ecies b 1 <sup>st</sup> art ium to	g. 3b). belong cicle of b long,	

315 this ecomorph predominantly occur on fine substrates and are characterized by a fossorial behavior 316 (Fig. 3b). This appears to be the most common ecomorph since almost half of the Ponto-Caspian 317 species are classified as diggers (49%). Taxonomic composition is given in Table 2. Representative 318 genera: Pontogammarus and Niphargoides.

319

320 4) Symbiont. Very stout and generally minute bodies, with well-developed coxal plates and pereopod 321 bases, usually characterized by diminished mouthparts (palps of maxilla 2 and maxilliped), pleon, 322 urosome, antennae and pereopods (Fig. 6). The gnathopods can be very specialized (Caspicola and 323 Iphigenella), or rudimentary (Behningiella and Cardiophilus). This ecomorph is the most distinct in 324 morphospace, with hardly any overlap (Fig. 3b). Its species are known to live on or inside bivalve 325 mollusks (Cardiophilus and Caspicola), or commensals with crayfish (Iphigenella). This ecomorph is 326 the rarest and accounts for 6 % of all species. Taxonomic composition is given in Table 2. 327 Representative genera: Cardiophilus and Iphigenella.



329 330

331 Fig. 6 Boxplots comparing selected traits among the four proposed ecomorphs. PC1 refers to the first principal 332 component resulting from the PCA analysis. It mainly describes the gradient from slender bodies with long 333 antennae (negative values) to stout bodies with short antennae (positive values). All traits except body length 334 and PC1 values are presented relative to total body length

### 336 **Discussion**

Our study reviewed and quantified for the first time the rich taxonomic, ecological and morphological
diversity of Ponto-Caspian amphipods. Although we consider these findings preliminary, our synopsis
will serve as a foundation for future eco-evolutionary and systematic studies. Below we discuss the
evidence accrued so far that point towards a remarkable, yet unrecognized adaptive radiation.
Within each of the following sub-sections we also highlight the gaps in existing knowledge and

- 342 recommend further research.
- 343

### 344 **Ponto-Caspian gammarid amphipods – an adaptive radiation?**

The main prerequisites that define an adaptive radiation are: monophyly, species sympatry,
speciation rate increase, and ecomorphological divergence (Schluter 2000; Simões et al. 2016). With
respect to Ponto-Caspian amphipods the sympatry criterion is the most readily fulfilled since most of
the species co-occur in the Caspian Sea and Lower Volga (Table 1). Furthermore, most species seem
to be widespread in the Caspian Sea, occurring in all of its main areas (north, middle and southern)
(Pjatakova and Tarasov 1996). A significant number of species are also found in sympatry in the
Ponto-Azov region (Cărăuşu et al. 1955).

352

The monophyly condition is supported by recent molecular phylogenies which indicate that several morphologically disparate Ponto-Caspian genera form a well-supported clade (Copilaş-Ciocianu, Borko, et al. 2020; Hou et al. 2014; Hou and Sket 2016; Sket and Hou 2018). Although relatively few taxa have been sequenced so far, it is likely that the remaining species would fall within the same clade. The Ponto-Caspian amphipod radiation also satisfies the requirement of speciation rate increase since it experienced a higher diversification rate in comparison to its sister clades (Hou et al. 2014).

360

361 We consider that our current study fulfills, at least partially, the criterion of ecomorphological 362 divergence, which is perhaps the most relevant to the adaptive radiation model. We highlight 363 significant ecological and morphological disparity within the Ponto-Caspian amphipod radiation. 364 Along an order of magnitude body-size gradient, morphology ranges from minute (several 365 millimeters), stout-bodied symbiotic species with attenuated appendages, to large and slender 366 (several centimeters), stocky and setose, or heavily armored species. Likewise, ecological diversity is 367 also remarkable, with species being encountered along a >500 m depth gradient on virtually all types 368 of substrates and water bodies (mountain springs to deep sea). By integrating morphology and 369 ecology, we propose a provisional classification into four main ecomorphs: clingers, crawlers, diggers 370 and symbionts. Although this classification is only tentative, we consider it a necessary first step 371 towards understanding the evolution of Ponto-Caspian amphipods. We highlight that these 372 ecomorphs have potential analogues in distantly related marine or Lake Baikal taxa that occupy 373 similar habitats (see Morphological evolution section below), further strengthening the environment-374 phenotype association.

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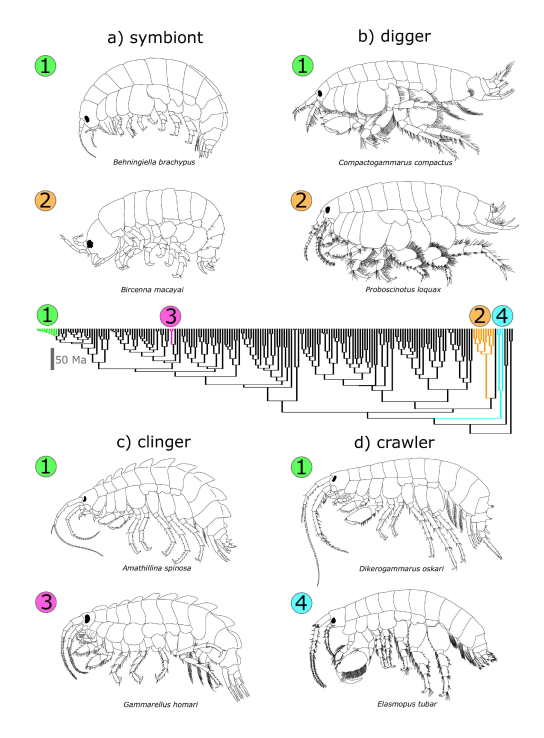
376 Overall, it appears that Ponto-Caspian amphipods fulfill, at least to some extent, the main 377 prerequisites of the adaptive radiation model. However, our findings provide only a first glimpse. 378 Extensive further research is needed to corroborate the patterns highlighted herein. Specifically, the 379 criteria of monophyly and speciation rate increase have to be tested on larger multilocus phylogenies 380 with a greater taxonomic coverage. The morphology-environment association needs to be refined 381 with newly collected field data. Specifically, fine-scale morphometry of functionally relevant traits 382 coupled with trophic niche (gut content DNA metabarcoding and stable isotopes) and ecology 383 (depth, substrate and salinity) in a phylogenetic context will provide a more comprehensive 384 ecomorphological understanding. Furthermore, it is important to test whether these ecomorphs 385 have a common ancestor or evolved several times independently (Trontelj et al. 2012). It is likely that 386 upon more detailed investigation they could be split into more specialized forms. Comparative

transcriptomics and genomics could provide important insight into adaptation and selection at the
 molecular level. A well-sampled time-calibrated molecular phylogeny could also prove invaluable for
 understanding the historical circumstances that promote the evolution of invasive species.

## 390

### 391 Morphological evolution

392 Recent molecular phylogenies revealed that the morphologically diverse Ponto-Caspian amphipod 393 radiation is nested within the genus Echinogammarus (Hou and Sket 2016; Sket and Hou 2018), 394 which is characterized by morphological conservatism (Pinkster 1993). This is in good agreement with 395 previous hypotheses that postulated a close relationship between these two groups (Barnard and 396 Barnard 1983). A similar pattern is also encountered in the two highly diverse Baikal amphipod 397 radiations which are classified into several families (Hou and Sket 2016; Lowry and Myers 2013), yet 398 they are both nested within the genus Gammarus (Hou et al. 2011, 2014; Macdonald et al. 2005; 399 Naumenko et al. 2017), notorious for its low morphological diversity, morphological crypsis (Copilas-400 Ciocianu and Petrusek 2015; Katouzian et al. 2016; Mamos et al. 2014) and generalist ecology 401 (MacNeil et al. 1997; Piscart et al. 2011). And yet again the same pattern appears in the distantly 402 related American genus Hyalella where morphologically conserved riverine species (Witt et al. 2006) 403 colonized the ancient Titicaca Lake multiple times, giving rise to a remarkable array of forms 404 (Adamowicz et al. 2018; González and Coleman 2002; Jurado-Rivera et al. 2020). These compelling 405 patterns indicate that species living in ephemeral, highly fluctuating and ecologically limited 406 environments (springs, streams, rivers and shallow lakes/ponds) are under stabilizing selection for 407 maintaining a generalist life-style and a conserved, non-specialized morphology (Wellborn and 408 Broughton 2008). On the other hand, species inhabiting stable ancient lakes with broad niche space 409 are probably under disruptive selective pressures which in turn promote specialization and ecological 410 speciation (Seehausen 2015; Wellborn and Langerhans 2015). Thus, it would seem that the ecological 411 transition from ephemeral habitats to long-lived ancient lakes promotes adaptive radiations in some 412 freshwater amphipod groups. These intriguing patterns are worth pursuing further and could shed 413 more light on the role of ecological opportunity in driving adaptive radiations.



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- 416

417 Fig. 7 Putative examples of ecomorphological convergence of Ponto-Caspian and distantly related oceanic taxa. 418 Ponto-Caspian species are shown with green. a) Symbiotic ecomorph adapted to piercing various organic 419 substrates (redrawn from Derzhavin (1948) and Loerz et al. (2010)), b) digger ecomorph adapted for digging 420 and burrowing in fine substrates (redrawn from Sars (1895) and Barnard (1967)), c) clinger ecomorph adapted 421 to cling on algal and vegetal substrates (redrawn from Sars (1896)), and d) crawler ecomorph adapted to a 422 generalist life-style, usually hiding in coarse stony substrates (redrawn from (Sars (1896) and Garcia-Madrigal 423 (2010)). The phylogenetic tree is a time-calibrated molecular phylogeny of Amphipoda modified after Copilaş-424 Ciocianu et al. (2020) 425

#### 426

427 We propose that the ecological and morphological diversity of Ponto-Caspian gammarids can be 428 distilled into four ecomorphs. Remarkably, all of them apparently have analogues in distantly related 429 lineages inhabiting oceanic waters or other ancient lakes (Figs. 7-8). The Ponto-Caspian symbiotic 430 ecomorph is the most specialized and morphologically distinct due to its reduced mouthparts, 431 antennae, pereopods and urosome, presumably due to a semi-parasitic life-style. We highlight a 432 striking resemblance between the Ponto-Caspian genus Behningiella and the oceanic algae-boring 433 genus Bircenna Chilton, 1884 (Fig. 7a). Both exhibit typical features for substrate boring such as a 434 large head with protruding mandibles adapted to cutting into tough material, and extremely short 435 antennae and pereopods due to living in narrow self-constructed tunnels (Mejaes et al. 2015). Within 436 the Baikal Lake Acanthogammaride radiation, the symbiotic ecomorph is probably represented by 437 the parasitic genus Pachyschesis (Naumenko et al. 2017; Takhteev 2019).

438

439 The fossorial ecomorph seems to be the most common among Ponto-Caspian amphipods. These 440 species are generally adapted for digging in fine substrates and have stout, strong bodies with very 441 short yet powerful and thick antennae, and broadened pereopods usually fringed with dense rows of 442 setae. This ecomorph is widely encountered among amphipods in general, albeit under slightly 443 different iterations (Bousfield 1970). Morphologically, most fossorial amphipods are classified within 444 the superfamily Haustorioidea (Lowry and Myers 2017). However, molecular phylogenies indicate 445 that the fossorial body-type evolved multiple times independently (Copilas-Ciocianu, Borko, et al. 446 2020; Hancock et al. 2020). A noticeable resemblance can be observed between the Ponto-Caspian 447 genus Compactogammarus and the hyaloidean Proboscinotus Barnard, 1967 (Fig. 7b). Additionally, in 448 Lake Baikal this ecomorph is possibly represented by the Micruropodidae radiation, comprising 449 fossorial species living on fine substrate (Naumenko et al. 2017; Takhteev 2019).

450

The clinger ecomorph characterizes species with elaborate body armature/ornamentation and 451 452 preference for living (plant) substrate. These species often have elongated and curved dactyls for 453 improved grasping of the substrate. Given the exposed nature of their life-style, the armature might 454 serve as protection against predators (Bollache et al. 2006; Copilas-Ciocianu, Borza, et al. 2020) or, in 455 combination with variegated coloration (as is often the case with armored taxa), may act as 456 camouflage by disrupting the body contour (d'Udekem d'Acoz and Verheye 2017). We point out the 457 high similarity among the Ponto-Caspian genus Amathilling and the oceanic algae-clinging 458 Gammarellus Herbst, 1793 (Fig. 7c). Although the Ponto-Caspian clingers are diverse in 459 ornamentation and armature, some striking resemblance can be observed with Baikal Lake taxa. For 460 example Amathillina and Eucarinogammarus (Baikal), Axelboeckia and Acanthogammarus (Baikal), 461 and Kuzmeling and Propachygammarus (Baikal) (Naumenko et al. 2017; Takhteev 2019). 462

463 The crawler ecomorph is the second-most encountered in Ponto-Caspian amphipods, characterizing 464 species living on coarse or fine substrate, often in shallow water. Typically, these taxa are strongly 465 sexually dimorphic, males possessing very large second gnathopods, relatively long antennae and 466 slender bodies with shallow coxal plates. Morphologically, this morph is probably the most 467 plesiomorphic, being widespread among the amphipod evolutionary tree, especially in some basal 468 branches (Copilas-Ciocianu, Borko, et al. 2020; Lowry and Myers 2017) as well as in the oldest known 469 fossils (Jarzembowski et al. 2020). As an example, we emphasize the similarity among the Ponto-470 Caspian genus Dikerogammarus and the widespread littoral genus Elasmopus Costa, 1853 (Fig. 471 7d). The Baikalian analogues of this ecomorph could be envisioned in Eulimnogammarus and 472 Corophiomorphus (Naumenko et al. 2017; Takhteev 2019).

473

474 Body armature is extremely diverse in amphipods, with similar phenotypes having evolved

independently multiple times (Copilaş-Ciocianu, Borko, et al. 2020; Lowry and Myers 2017;

- 476 Naumenko et al. 2017). We highlight a remarkably convergent evolution of body armature in some
- ancient lake radiations where strong lateral spines appear on the pereonites, the longest one being

478 located on the 4<sup>th</sup> or 5<sup>th</sup> segment (Fig. 8). In some cases the spine is an outgrowth of the tergum,

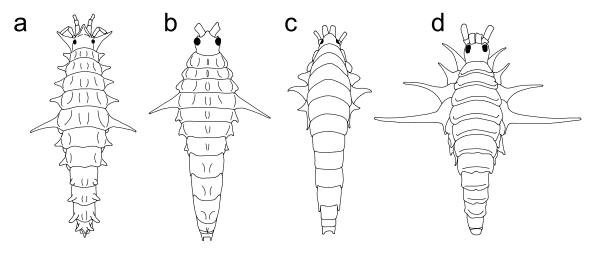
479 while in others an outgrowth of the coxal plate. These analogous convergent structures point

480 towards a strong selective pressure. Most likely these spines function as a mechanism for deterring

481 ingestion by predatory fish (Bollache et al. 2006; Copilaş-Ciocianu, Borza, et al. 2020), although the

482 exact mechanical interactions are unknown.

483



484 485

Fig. 8 Examples of evolutionary convergent patterns in body armature of species inhabiting different ancient
lakes. a) Axelboeckia spinosa (Caspian Sea, redrawn after Sars (1894b)), b) Acanthogammarus lappaceus (Lake
Baikal, redrawn after Daneliya et al. (2011)), c) Issykogammarus hamatus (Lake Issyk-Kul, redrawn after
Chevreux (1908)), and d) Hyalella armata (Lake Titicaca, redrawn after González & Coleman (2002))

490 491

### 492 Spatio-temporal origin

493 The phylogenetic position of Ponto-Caspian amphipods within the Atlanto-Mediterranean 494 Echinogammarus clade (sensu Hou et al., 2014; Sket & Hou, 2018) indicates that this radiation likely 495 has a Mediterranean origin. Specifically, its sister clade is represented by the genus 496 Dinarogammarus, which is endemic to freshwaters of the Western Balkans (Sket and Hou 2018). 497 Regarding the temporal time-frame, several recent studies proposed a Middle Miocene origin 498 (ca.12-14 Ma) (Copilas-Ciocianu, Borko, et al. 2020; Hou and Sket 2016), coeval with the final closure 499 of the Paratethys, which caused a switch from marine to brackish conditions and promoted the 500 evolution of endemic faunas after initial mass extinctions (Popov et al. 2004; Rögl 1999). This time 501 frame is also supported by Late Miocene (ca. 9-10 Ma) Caucasian fossil taxa (two genera and five 502 species) that have clear affinities with extant Ponto-Caspian genera Axelboeckia, Gmelina, Kuzmelina 503 and Yogmelina (Derzhavin 1927, 1941). Alternatively, an earlier study suggested an origin dating back 504 to the Eocene (30-40 Ma) (Hou et al. 2014). However, this analysis was based on biogeographical 505 calibration of the molecular clock rather than fossils, thus possibly resulting in biased inferences (Ho 506 et al. 2015). Furthermore, a Late Eocene origin does not correspond with an isolation of the 507 Paratethys realm from the world ocean (Popov et al. 2004). As such, we consider that a middle 508 Miocene origin is more plausible considering the data at hand.

509

510 A densely sampled, multilocus and time-calibrated phylogeny will be of critical importance in

511 understanding the historical biogeography and evolution of Ponto-Caspian gammarids. Furthermore,

such a phylogeny could complement geological studies regarding the palaeogeographic history of the

513 Paratethyan region, as seen with other freshwater gammarids (Copilaș-Ciocianu et al. 2019; Copilaș-

514 Ciocianu and Petrusek 2017; Hou et al. 2011; Mamos et al. 2016). It could provide additional time

515 constraints on some important palaeogeographic events such as the final Paratethys closure, the

isolation of the Pannonian, Pontic and Caspian basins, the emergence of the Caucasus, as well as therecurrent episodic connections of the Pontic and Caspian basins during the Plio-Pleistocene.

### 519 Taxonomic and systematic remarks

520 The Ponto-Caspian gammaroid amphipods as defined in this study are formally split into 5 families: 521 Behningiellidae, Caspicolidae, Gammaridae, Iphigenellidae and Pontogammaridae. However, 522 molecular research has revealed that Pontogammaridae is nested within Gammaridae, and also 523 harbors the gammarid genus Dikerogammarus (Copilaş-Ciocianu, Borko, et al. 2020; Hou et al. 2014). 524 Members of this family correspond to the digger ecomorph, which probably evolved on more than 525 one occasion. Moreover, the Ponto-Caspian "Gammaridae" form a paraphyletic grade at the base of 526 Pontogammaridae (Sket and Hou 2018). A taxonomically more inclusive morphological and molecular 527 study will clarify this issue, but most likely will not recover Pontogammaridae as monophyletic. 528 However, for the sake of stability we do not propose any taxonomic changes until this issue is firmly 529 resolved.

530

518

531 The remaining families Behningiellidae, Caspicolidae and Iphigenellidae are poorly known and have 532 not yet been sequenced. Behningiellidae and Iphigenellidae have been classified into Gammaroidea 533 based on a morphological cladistic analysis (Lowry and Myers 2013). However, the monotypic 534 Caspicolidae is currently not recognized as part of Gammaroidea, but as a distinct superfamily 535 (Caspicoloidea) within the infraorder Talitrida (Lowry and Myers 2013). This classification is 536 erroneous because the authors mistakenly considered that the antenna I lacks an accessory flagellum 537 (a defining character state of the infraorder Talitrida). Derzhavin's (1944) original description clearly 538 indicates the presence of the accessory flagellum, although it is reduced and uniarticulate. Another 539 issue with assigning Caspicolidae to Talitrida is the presence of a well-developed mandibular palp, 540 whereas an absent/vestigial palp is another defining character state of the Talitrida (Lowry and 541 Myers 2013). Behningiellidae, Caspicolidae and Iphigenellidae belong to the symbiotic ecomorph and 542 represent highly specialized taxa which are difficult to classify using external morphology alone. It is 543 very likely that these small families are nothing but highly derived Ponto-Caspian gammarids, 544 possibly related to the various genera of the gmelinid facies (Gmelina, Kuzmelina and Yogmelina) 545 (Barnard and Barnard 1983; Bousfield 1977; Derzhavin 1944). Thus, the systematic position of these 546 families will be clarified only with additional morphological and molecular study.

547

We argue that most, if not all Ponto-Caspian amphipod species are in need of thorough, modern
revision using morphology, multilocus DNA sequences and ecology. Many species are only partially
illustrated and intraspecific variability has been studied in only a handful of taxa (Cărăuşu 1936;
Nahavandi et al. 2013). Moreover, cryptic lineages of potential specific status have been recently
discovered (Jażdżewska et al. 2020). As such, a first step towards a modern taxonomic revision could
be the generation of a well-sampled DNA barcode reference library.

554

555 Lastly, the Late-Miocene (Upper Sarmatian, ca. 9 Ma) fossil genera Andrussovia and Praegmelina 556 have long been considered ancestral to extant Ponto-Caspian genera such as Gmelina and 557 Amathillina, albeit without a formal analysis (Barnard and Barnard 1983; Derzhavin 1927, 1941). The 558 fossils were discovered in calcareous clay deposits at the foothills of the Caucasus near Grozny, 559 Solenaya balka (Chechnya, Russian Federation) and from Eldar Oyugu Ridge (Azerbaijan). We agree 560 that there are rather clear affinities with extant Ponto-Caspian species in general, mainly in the 561 combination of the following traits: shape of the basis of pereopod 7, ornamentation and armature, 562 short and thick antennae, and deep coxal plates. Some traits are considered plesiomorphic, such as 563 the lack of a postero-ventral lobe on the basis of pereopod 7, and the long endopod of uropod 3. A cladistic analysis is necessary to confidently assess evolutionary relationships with extant taxa. Until 564 565 then, these species should be conservatively treated as stem Ponto-Caspian amphipods (Copilaş-566 Ciocianu, Borko, et al. 2020).

568 Two more Miocene fossil taxa have been reported from the Caucasus that have less clear affinity 569 with extant Ponto-Caspian taxa. These are Gammarus praecyrius, Derzhavin, 1941 and Hellenis saltatorius, Petunnikoff, 1914. The former is indistinguishable from a typical Gammarus and it is thus 570 571 not considered a Ponto-Caspian taxon. The affinities of the latter taxon are less straightforward to 572 interpret due to its high degree of morphological specialization (very short antennae, large raptorial 573 gnathopods and unusually long pereopods). Such a combination of traits is not present in the extant 574 Ponto-Caspian fauna. Furthermore, Petunnikoff's illustrations are also not detailed enough to draw a 575 conclusion. At the moment we consider that it is possible that *H. saltatorius* could be related to 576 Ponto-Caspian amphipods but further detailed studies are needed.

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### 579 Conclusion

580 The Ponto-Caspian gammarid radiation fulfills, at least partially, the most important criteria of an 581 adaptive radiation: 1) apparent monophyly, 2) sympatric occurrence within a constrained area, 3) 582 accelerated diversification and 4) ecomorphological disparity. Nevertheless, these literature-based 583 results are only preliminary and a lot of in depth eco-evolutionary study is further needed. Moreover, 584 most species need a modern taxonomic revision within an evolutionary context. Nevertheless, we 585 consider that Ponto-Caspian amphipods could be an excellent future model for the study of adaptive 586 radiation, origin of invasive species, and could even help illuminate the region's dynamic

- 587 palaeogeographic history.
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881	Appendix
882	Key to endemic families, genera and species of Ponto-Caspian amphipods (including non-
883	Gammaroidea).
884	
885	Key to families (parentheses indicate non-Gammaroid genera that contain more than one
886	species; their species are keyed in the "Key to species" below)
887	1. Eyes absent Niphargidae
888	- Eyes present
889	2. Body dorso-ventrally compressed, coxal plates rudimentary, antenna 2 greatly developed
890	Corophiidae ( <i>Chelicorophium</i> )
891	- Body laterally compressed, coxal plates developed, antenna 2 normally developed
892	3. Head rostrum long and narrow, uropod 3 rami foliaceous of equal length Gammaracanthidae
893	- Head rostrum absent or rudimentary, uropod 3 rami slender, subequal 4
894	4. Telson uncleft Uristidae (Onisimus)
895	- Telson cleft
896	5. Pereopod 6 much longer than pereopods 5 and 6 Pontoporeiidae
897	- Pereopod as long as or slightly longer than pereopods 5 and 66
898	6. Gnathopods chelate Caspicolidae
899	- Gnathopods subchelate
900	7. Pereopods 5-7 prehensile Iphigenellidae
901	- Pereopods 5-7 not prehensile
902	8. Maxilliped and mandibular palp reduced, and/or meral articles of pereopod 5-7 with postero-distal
903	lobe
904 905	- Maxilliped and mandibular palp normal, meral articles of pereopod 5-7 without postero-distal lobe
906 907 908	<ul> <li>9. Antenna 1 peduncle article 1 slender, as long as or shorter than articles 2 and 3 combined</li> <li>Gammaridae</li> <li>- Antenna 1 peduncle article 1 swollen, longer than articles 2 and 3 combined Pontogammaridae</li> </ul>
909	Key te server (entre Commenciales)
910	Key to genera (only Gammaroidea)
911 912	1. Gnathopods chelate Caspicola (monotypic: Caspicola knipovitschi Derzhavin, 1945)
912 913	- Gnathopods subchelate
915 914	2. Head with lateral projections
914 915	- Head with lateral projections
916	3. Strong lateral-pointing spine on pereonite 5 Axelboeckia
917	(monotypic: Axelboeckia spinosa (G.O. Sars, 1894))
918	- No spines, only blunt knobs on pereonite 5
919	4. Body with two dorso-lateral rows of knobs
920	(monotypic: <i>Kuzmelina kusnezowi</i> (Sowinsky, 1894))
921	- Body with a central dorsal a keel on pereonal segments 6-7 and pleosome Gmelinopsis
922	5. Antenna 1 accessory flagellum uniarticulate
923	- Antenna 1 accessory flagellum two- or multiarticulated 10
924	6. Lateral head lobes form a characteristic "hood" Scytaelina
925	(monotypic: <i>Scytaelina simplex</i> Stock et al., 1998)
926	- Lateral head lobes normal
927	7. Bases of pereopods 5-6 lobed, uropod 3 reduced Behningiella
928	(monotypic: Behningiella brachypus Derzhavin, 1948)
929	- Bases of pereopods 5-6 not lobed, uropod 3 normal8
930	8. Body with an obvious central dorsal keel, more pronounced on the pleosome
931	- Body with a weak or absent central dorsal keel

932 933	9. Pereopod 7 basis with a large downward pointing postero-distal lobe Jugogammarus (monotypic: Jugogammarus kusceri (S. Karaman, 1931))
934 935	<ul> <li>Pereopod 7 basis with minute/without a downward pointing postero-distal lobe</li></ul>
936	- Gnathopod 1 equal or smaller than gnathopod 2
937	11. Uropod 3 reduced, exopod shorter than twice the peduncle length
938	(one species: <i>Baku paradoxus</i> (Derzhavin in Derzhavin & Pjatakova, 1967)
939	- Uropod 3 normal, exopod at least twice the peduncle length
940	12. Dactyli of pereopods 3-7 prehensile
941	(monotypic: <i>Iphigenella acanthopoda</i> G.O. Sars, 1896)
942	- Dactyli of pereopods 3-7 not prehensile Lanceogammarus
943	(monotypic: Lanceogammarus andrussowi (G.O. Sars, 1896))
944	13. Pereopod 7 basis not lobed
945	- Pereopod 7 basis lobed
946	14. Antenna 1 accessory flagellum bi-articulated Cardiophilus
947	- Antenna 1 accessory flagellum tri- or more articulated 15
948	15. Antenna 2 peduncle greatly expanded, article 3 with a downward projection Derzhavinella
949	- Antenna 2 peduncle normal, article 3 without a downward projection
950	16. Antenna 1 peduncle article 3 longer or equal with article 1
951	(monotypic: <i>Sowinskya macrocera</i> Derzhavin, 1948)
952	- Antenna 1 peduncle article 3 shorter than article 1
953	17. Uropod 3 setae curled, longer than spines Trichogammarus
954	(monotypic: Trichogammarus trichiatus (Martynov, 1932))
955	- Uropod 3 setae straight, shorter than spines
956	18. Pereopod 6 basis lobed
957	- Pereopod 6 basis not lobed
958	19. Pereopod 5-7 meral articles with a postero-distal lobe Zernovia
959	(monotypic: <i>Zernovia volgensis</i> Derzhavin, 1948)
960	- Pereopod 5-7 meral articles without a postero-distal lobe Shablogammarus
961	(monotypic: Shablogammarus shablensis (Cărăușu, 1943))
962	20. Antenna 1 flagellum shorter than peduncle 21
963	- Antenna 1 flagellum equal/longer than peduncle 25
964	21. Epimeron 3 with postero-ventral setal fan
965	- Epimeron 3 without postero-ventral setal fan
966	22. Uropod 3 endopod longer than half of exopod Uroniphargoides
967	(monotypic: Uroniphargoides spinicaudatus (Cărăușu, 1943))
968	- Uropod 3 endopod shorter than half of exopod 23
969	23. Ganthopod 2 propodus palm longer than posterior margin <i>Compactogammarus</i>
970	(monotypic: Compactogammarus compactus (G.O. Sars, 1895))
971	- Ganthopod 2 propodus palm shorter than posterior margin Niphargoides
972	24. Uropod 3 exopod 2 <sup>nd</sup> article well developed, bearing lateral setae <i>Niphargogammarus</i>
973	- Uropod 3 exopod 2 <sup>nd</sup> article minute/absent, bearing no lateral setae Paraniphargoides
974	25. Eyes round to ovoid, gnathopod 2 propodus large and triangular, armed with a palmar spine as
975	long as 1/3 of dactylus Pandorites
976	(monotypic: Pandorites podoceroides (G.O. Sars, 1896))
977 978	- Eyes reniform, gnathopod 2 propodus armed with a palmar spine(s) shorter than 1/3 of dactylus
979	26. Antenna 1 peduncular article 1 slender, width not exceeding 1/3 of length, pereopods 3-4 with
980	sparse setae shorter than the width of underlying segment
981	- Antenna 1 peduncular article 1 robust, width exceeding 1/3 of length, pereopods 3-4 with dense
982	setae as long as/longer than the width of underlying segment (except <i>O. subnudus</i> )
983	27. Body with a central dorsal a keel
984	- Body without a keel

985	28. Head swollen and enlarged	
986 987	(monotypic: <i>Cephalogammarus macrocephalu</i> s) - Head normal	
988	29. Urosomites 1-2 with columnar tubercles	
989	- Urosomites 1-2 without columnar tubercles	-
990	30. Uropod 3 exopod 2 <sup>nd</sup> article at least 1/5 the length of 1 <sup>st</sup> article	
991	- Uropod 3 exopod 2 <sup>nd</sup> article much shorter than 1/5 the length of 1 <sup>st</sup> article	
992	31. Uropod 3 exopod outer margin armed with few simple setae	
993	- Uropod 3 exopod outer margin armed with many plumose setae	-
994	(monotypic: <i>Wolgagammarus dzjubani</i> (Mordukhai-Boltovskoi	
995	32. Setae on posterior margin of carpal articles of pereopods 3-4 arranged in a contin	-
996		
997	- Setae on posterior margin of carpal articles of pereopods 3-4 arranged in clusters	
998	33. Urosomites 1-2 with noticeable dorsal knobs 7	
999	- Urosomites 1-2 flat or humped	-
000	·	5
001		
.002	Key to species	
.003	Doubtful species are indicated with an asterisk (*).	
.004		
.005	Key to species of Akerogammarus	
006	1. Propodi of gnathopods 1 and 2 similar in size, telson with short apical setae	A. contiguous
.007	- Propodus of gnathopod 2 larger than gnathopod 1, telson with long apical setae	A. knipowitschi
800		
009	Key to species of Amathillina	
010	1. Body keel starts from the first pereonite	2
011	- Body keel starts from the sixth pereonite or later	
012	2. Pereonites 1-2 with well-developed dorsal spines, urosomite 1 humped	
013	- Pereonites 1-2 with barely visible crest, urosomite 1 not humped	
.014	3. Body keel present only on pleosome	A. maximowiczi
.015	- Body keel present on pereonites 6-7 and pleosome	
.016	4. Antenna 1 accessory flagellum bi-articulated, last pleonal projection rounded	•
017	- Antenna 1 accessory flagellum tri-articulated, last pleonal projection triangular	A. affinis
018		
.019	Key to species of <i>Cardiophilus</i>	
.020	1. Uropod 3 exopod less than twice the length of peduncle, 2 <sup>nd</sup> article present	
.021	- Uropod 3 exopod twice the length of peduncle, 2 <sup>nd</sup> article absent	C. maris-nigri
.022		
023	Key to species of Chaetogammarus	
.024	1. Antenna 1 accessory flagellum bi-articulated	
.025	- Antenna 1 accessory flagellum at least three articles	
.026	2. Pleosome covered with small spines	
.027	- Pleosome bare	
028	3. Eyes very elongated and constricted in the middle	•
029	- Eyes regular (reniform)	
.030	4. Antenna 1 and 2 set with dense setae longer than the underlying segment	
031	- Antenna 1 and 2 set with sparse setae shorter/equal with the underlying segment	C. pauxillus
.032		
.033	Key to species of Chelicorophium	
.034	1. Antenna 2 peduncular article 4 distal tooth simple	
.035	- Antenna 2 peduncular article 4 distal tooth with an additional simple or bidentate to	oth 2

1036	2. Antenna 2 peduncular article 4 distal tooth with an additional simple tooth
1030	- Antenna 2 peduncular article 4 distal tooth with an additional bidentate tooth
1037	3. Pereopods 3-4 meral articles stout, length is less than twice the width
1039	- Percopods 3-4 meral articles slender, length is twice the width
1040	4. Antenna 2 peduncular article 5 with a proximal and distal tooth
1040	- Antenna 2 peduncular article 5 with either a proximal or distal tooth
1042	5. Antenna 2 peduncular article 5 with a small proximal tooth, distal tooth missing
1043	- Antenna 2 peduncular article 5 with a strong distal tooth, proximal tooth missing <i>C. spinulosum</i>
1043	6. Antenna 2 peduncular article 5 with a proximal and distal tooth
1045	- Antenna 2 peduncular article 5 with a proximal tooth only
1046	7. Antenna 2 peduncular article 5 proximal tooth situated in the distal half of the article, inner side of
1047	uropod 3 peduncle without spines
1048	- Antenna 2 peduncular article 5 proximal tooth situated in the proximal half of the article, inner side
1049	of uropod 3 peduncle with spines
1050	8. Antenna 1 flagellum as long as peduncle
1050	- Antenna 1 flagellum half as long as peduncle
1051	
1053	Key to species of <i>Derzhavinella</i>
1055	1. Anterior margin of pereopod 7 with long setae
1055	- Anterior margin of percopod 7 with short setae
1056	
1057	Key to species of <i>Dikerogammarus</i>
1058	1. Pleosome segments keeled
1059	- Pleosome segments flat
1060	2. Urosomal tubercles low
1061	- Urosomal tubercles columnar, well developed
1062	3. Antenna 1 accessory flagellum bi-articulated
1063	- Antenna 1 accessory flagellum 4 or more articles
1064	4. Propodi of gnathopods 1 & 2 with mid-palmar spine
1065	- Propodi of gnathopods 1 & 2 without mid-palmar spine
1066	5. Medial surface of pereopod 7 basis with setae
1067	- Medial surface of percopod 7 basis without setae
1068	6. Antenna 2 peduncular segments with numerous clusters of setae longer than the underlying
1069	segment
1070	- Antenna 2 peduncular segments with few clusters of setae shorter than the underlying segment
1071	D. istanbulensis
1072	7. Propodi of gnathopods 1 & 2 with setae as long as propodus width
1073	- Propodi of gnathopods 1 & 2 with setae much shorter than propodus width
1074	8. Uropod 3 exopod with spines on inner and outer margins
1075	- Uropod 3 exopod without spines
1076	
1077	Key to species of <i>Gmelina</i>
1078	1. Pleonal humps high and triangular <i>G. costata</i>
1079	- Pleonal humps low and rounded G. aestuarica
1080	
1081	Key to species of <i>Gmelinopsis</i>
1082	1. Head lateral projections blunt, tubercle-like <i>G. tuberculata</i>
1083	- Head lateral projections pointed, spear-like G. aurita
1084	
1085	Key to species of Niphargogammarus
1086	1. Antenna 1 main flagellum as long as the first peduncular article 2
1087	- Antenna 1 main flagellum shorter than the first peduncular article

3 9 0	<ul> <li>2. Body size ca. 10 mm, gnathopod 2 propodus larger than gnathopod 1, telson lobes diverging and armed with 1 apical spine</li></ul>			
L	armed apically with 2 spines			
2	3. Body size ca. 13 mm, urosomite 1 bare, uropod 3 exopod external margin armed with many setae 			
1 5 5	- Body size ca. 7 mm, urosomite 1 armed with small setae, uropod 3 exopod external margin armed with few distal setae <i>N. intermedius</i>			
7	Key to species of <i>Niphargoides</i>			
3	1. Posterior margin of pereopod 6 basis armed with a few short setae proximally			
9	- Posterior margin of pereopod 6 basis armed with long setae along its entire length			
)	2. Ventral margin of coxal plate 4 armed with setae shorter than 1/2 of its length N. corpulentus			
L	- Ventral margin of coxal plate 4 armed with setae longer than 1/2 of its length			
2	3. Uropod 3 exopod bares a well-developed setal fan N. caspius			
	- Uropod 3 exopod lacks setal fan, armed with strong spines N. boltovskoyi			
	Key to species of Obesogammarus			
	1. Posterior margin of pereopods 3-4 with few sparse setae shorter than the underlying segment 			
	- Posterior margin of pereopods 3-4 with numerous setae as long as/longer than the underlying segment			
	2. Uropod 3 exopod less than twice peduncle length			
	- Uropod 3 exopod ites than twice peduncle length			
	3. Posterior margin of basis of percopods 5-7 with short setae			
	- Posterior margin of basis of percopods 5-7 with long setae			
	4. Coxae 1-2 tapering towards distal end, propodus of gnathopod 2 triangular			
	- Coxae 1-2 not tapering, propodus of gnathopod 2 roughly rectangular			
	5. Medial surface of pereopod 7 basis without setae			
	- Medial surface of percopod 7 basis armed with clusters of setae			
	6. Urosomites with dorsal elevations			
	- Urosomites without dorsal elevations			
	Key to species of <i>Onisimus</i>			
	1. Antenna 1 peduncle article 1 longer than head, accessory flagellum 6-segmented, postero-ventral			
	corner of 3 <sup>rd</sup> epimere almost straight			
	2. Antenna 1 peduncle article 1 shorter than head, accessory flagellum 4-segmented, postero-ventral			
	corner of 3 <sup>rd</sup> epimere sharp			
	corner or s reprinere sharp			
	Key to species of <i>Paraniphargoides</i>			
	1. Uropod 3 exopod without terminal article, plumose setae absent			
	- Uropod 3 exopod with minute terminal article, plumose setae absent			
	סיסטיט ש באסטיט אונוי ווווועני נכווווומו מרנוסי, טעוווטשי שכומב ערפשרוג אווווומי אונויא אוווייניט איניי אוווייניט איניי			
	Key to species of <i>Pontogammarus</i>			
	1. Uropod 3 endopod reaches half the length of the exopod 1 <sup>st</sup> article			
	- Uropod 3 endopod shorter than half the length of the exopod 1 <sup>st</sup> article			
	2. Dactylus nail of pereopods 5-7 hook-like			
	- Dactylus nail of pereopods 5-7 straight			
	3. Mandibular palp very large, plumose D-setae present, last article of antenna 2 bares continuous			
	setal fan P. maeoticus			
	- Mandibular palp normal, D-setae absent, last article of antenna 2 bares several setal clusters			

1140	4. Urosome with dorsal elevations	5		
1141	- Urosome flat	7		
1142	5. Dorsal elevations tall, pillar-like	P. setosus		
1143	- Dorsal elevations low, hump like	6		
1144	6. Urosomite 1 armed with a crown of spines	P. robustoides		
1145	- Urosomite 1 armed with setae only	P. aestuarius		
1146	7. Uropod 3 exopod 2 <sup>nd</sup> article as long as broad, uropod 2 exopod devoid of spines	P. borceae		
1147 1148	- Uropod 3 exopod 2 <sup>nd</sup> article longer than broad, uropod 2 exopod with 1 spine	P. abbreviatus		
1149	Key to species of Stenogammarus			
1150	1. Uropod 3 reduced, 1 <sup>st</sup> exopod article as long as peduncle	S. micrurus		
1151	- Uropod 3 not reduced, 1 <sup>st</sup> exopod article longer than peduncle			
1152	2. Uropod 3 exopod inner margin bare/with 1 long seta			
1153	- Uropod 3 exopod inner margin with multiple long setae			
1154	3. Pereopods 6-7 basis medial surface with clusters of setae, basis 6 with long setae on posterior			
1155	margin, basis 7 with long setae on anterior margin			
1156	- Pereopods 6-7 basis medial surface bare, basis 6 with short setae on posterior margin, basis 7 with			
1157	short setae on anterior margin			
1158	4. Uropod 3 exopod 2 <sup>nd</sup> article shorter than half of 1 <sup>st</sup> article			
1159	- Uropod 3 exopod 2 <sup>nd</sup> article as long/longer than half of 1 <sup>st</sup> article			
1160	5. Uropod 3 endopod longer than 2 <sup>nd</sup> exopod article and is 1/2 of 1 <sup>st</sup> article			
1161	- Uropod 3 endopod shorter than 2 <sup>nd</sup> exopod article and is 1/3 1 <sup>st</sup> article	S. macrurus		
1162				
1163	Key to species of <i>Turcogammarus</i>			
1164	1. Pleosome segments keeled	T. spandli		
1165	- Pleosome segments flat			
1166	2. Urosomal tubercles low	T. aralensis		
1167	- Urosomal tubercles tall and columnar	T. turcarum*		
1168				
1169	Key to species of <i>Yogmelina</i>			
1170	1. Uropod 3 reduced, 1 <sup>st</sup> article of exopod as long as peduncle	Y. brachyura		
1171	- Uropod 3 not reduced, 1 <sup>st</sup> article of exopod longer than peduncle	2		
1172	2. Basis of pereopod 7 truncated, abruptly tapering towards the distal edge			
1173	- Basis of pereopod 7 not truncated, gradually tapering towards the distal edge			
1174	3. Coxal plate 1 slightly bent forwards, fringed with short setae	Y. laeviuscula		
1175	- Coxal plate 1 strongly bent forwards fringed with long setae			
1176	4. Urosomites without setae, epimeral plates 2-3 with short distal setae	Y. pusilla		
1177	- Urosomites with setae, epimeral plates 2-3 with long distal setae	Y. limana		
1178				
1179				
1180				
1181				