What happened to the genus *Gammarus* (Crustacea: Amphipoda) 
in Central Asia

Kaj se je zgodilo s postranicami *Gammarus* (Crustacea: Amphipoda) 
v centralni Aziji

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Abstract: We examined the phylogenetic relationships in the unusually structured fauna of the amphipod genus *Gammarus* in Central Asia, trying to explain its causes. Mitochondrial DNA sequence data were used in MP and ML and Bayesian analyses to generate a phylogenetic hypothesis. All of the recently collected *Gammarus* spp. specimens belong to the *G. lacustris* aggregate, except scarce marginal populations of *G. balcanicus* and *G. komareki* aggregates. Although molecularly closely related, these *G. lacustris* agg. populations are morphologically and ecologically equally diverse as are the numerous European *Gammarus* spp. Only few populations are morphologically close to the morphotype of *G. lacustris*. There is a more than 2,000 km wide gap between the eastern and western populations-species of *G. balcanicus* agg. Since *Gammarus lacustris* can be spread by birds, and its Asian relatives are molecularly significantly diverse, we suppose that it invaded Central Asia several times and from several sources. It most probably differentiated to diverse morphotypes after reaching Asia. The area of the cryophilic and less nomadic *G. balcanicus* could have been reached and conquered only by waterways. Its area was probably continuous in the past and divided by aridification during the Neogene. The strong historical aridifications probably extirpated the probably higher *Gammarus* diversity in Central Asia, including *G. balcanicus*. This opened the area and its habitats for the reiterated invasions of the particularly euryoecious *G. lacustris s.l.*, followed by its morphological and ecological diversification.

Keywords: Amphipoda, biogeography, diversification, Palearctic

**Introduction**

*History of discovery and problems in taxonomy*

Amphipods of Central Asia are insufficiently studied; distributional data are fragmentary and descriptions are mostly poor, often not mutually comparable. Only a few of the previously described *Gammarus* taxa have recently been confirmed with data that are more detailed. In addition, administrative and orthography changes in the near history added to the problems of the distributional data. Just as an example: the Martynov’s (1935) ‘Eastern Bukhara’ is not part of the recent town Buhara, Uzbekistan, it is the eastern part of the extended Bukhara Emirate; its eastern part mainly included the central and western parts of the modern Tajikistan (Chibilev 1993).

Another problem concerns the numerous findings of ‘*Gammarus pulex*’ in Central Asia and adjacent regions. Its alleged first mention was by G.O. Sars (1901) from territories near Astana in Kazakhstan and eastern Mongolia, and then again by Sars (1903) from Lake Teleckoe, Altai. Much later, Martynov (1930b) reported four Amphipoda species for Teleckoe ozero, but *G. pulex* was not mentioned among them. However, material identified by Sars as *G. pulex* was described as new species, *G. teletzkensis* (see Martynov, 1930b). Surprisingly, *G. pulex* was listed by Martynov (1930b) from another Siberian lake and from the Biya river. Furthermore, Chevreux (1908), studying collections of prof. D.D. Pedashenko in 1906-1908, reported *G. pulex* from a number of points located in the ‘Lake Issyk-Koule, stream Gorge of Karakol, and Lake Chatyr-Koule’ of Kirghizia. Afterwards, Martynov (1930a), based partly on the same collections of Pedashenko, described *Gammarus ocellatus* Martynov 1930, and *Gammarus bergi* Martynov, 1930 from Lake Ysyk Köl (Issyk Kul). Once more later, *G. ocellatus* was indicated by Martynov and Behning (1948) from the high mountain lakes of the Hissar Range in Tajikistan. According to Martynov and Behning, 1948, *G. ocellatus* is a widely distributed species in the mountainous regions of Central Asia, occurring from Tajikistan to Tien Shan in the northeast and to the Himalayas in the southeast (Ueno 1934). The question of the existence of *G. pulex* in Asia gets significantly complicated with the impossibility to accurately identify a number of previous findings due to their superficial descriptions. In Turkmenistan, *G. pulex* was reported by Birstein (1945) from rivers Šerlok and Firjuzinka (central Kopetdag); this is the only mention of *G. pulex*, where some morphological details show similarity with that European species.

The great majority of localities, treated by Martynov, are actually situated in southern Kazakhstan or very close to its border in Uzbekistan and Kyrgyzstan. The recently collected samples from Central Asia, which we could identify molecularly, are from Kyrgyzstan, E Uzbekistan, Tajikistan, and from Turkmenistan. They all virtually only contain members of the *G. lacustris* aggregate. They inhabit springs, streams, lakes, and caves, habitats,
which in Europe would mainly be inhabited by *G. pulex*, *G. fossarum*, and *G. balcanicus* aggregates. Based on the available data and by analyzing the geographical distribution (see the map and species list, Supplement table S2), we made an attempt to conduct a comparative morphological analysis only for two new species described in another paper (Sidorov et al. 2018). Here, we are trying to establish, what happened with the rest of the *Gammarus* phylogenetic clades/aggregates, which had evidently inhabited the region in the past and only left behind some scarce relics.

**Material and methods**

**Study area**

Our study area was wider Central Asia, with exclusion of the Caspian in the West, and inclusion of Krasnojarsk in the East and North and of Islamabad in the south, but excluding the Tibetan plateau.

The areas sampled were as follows. In Turkmenistan, the surroundings of Aşgabat (Ashgabat; where some localities were not accessible) and Köytendag area in the extreme NE. In Kyrgyzstan the lake Ysyk Köl (Issyk Kul) with surroundings...
and the SW of the country. In Tajikistan, the NW third of the country (sampled by C. Fišer) and the South-East (sampled by D. Palatov). In other countries, some scattered localities by different collectors. All localities and collectors are listed in the supplementary table (Supplement table S1). Geographical maps with locations of the sampling sites were constructed using an open source software Generic Mapping Tools, GMT 4.5.14. (Fig. 1).

Gammarids were sampled in springs and streams with a hand net. In lakes, small dredge and baited falls made of plastic bottles and left on deeper bottoms overnight were also used. Samples were fixed and stored in 96% ethanol with a small amount of glycerol added; such animals are appropriate for DNA analyses and still not too rigid for morphological studies. Single specimens were partly dissected and some muscle tissues or small body parts taken for DNA analysis. The rest of the bodies, including all taxonomically relevant parts, were retained as voucher specimens for subsequent morphological studies.

All samples were molecularly identified up to the level of phylogenetic clades, defined as aggregates. Note that our aggregates (abbreviations: aggs, agg) (Hou and Sket 2016) are not identical with ‘groups’ of Karaman and Pinkster (1977a, 1987), who explicitly define them just as morphologically (rather than phylogenetically) defined groups. Species identification was only taken as reliable if we could molecularly define morphologically appropriate samples from the type locality. If not, identification was marked as ‘cf.’; such identification only shows a certain degree of similarity (i.e., morphotypes), allowed by the identification using available keys; it does not mean any degree of phylogenetic relatedness, let alone the taxonomical identity. In addition, a terrifying degree of homoplasy was noted in the morphological characters of Amphipoda (Hou and Sket 2011; Moškrič 2016).

**Molecular methods**

Total genomic DNA was extracted from specimens using the Tiangen Genomic DNA kit. Mitochondrial cytochrome oxidase subunit I (COI) and nuclear 28S rRNA genes were amplified using primers as in Hou et al. (2007). Chromatograms were proofed and edited using Sequencher 4.2 DEMO (Gene Codes Corporation, Inc). Sequences were aligned using Clustal X (Thompson et al. 1997) and translated with the invertebrate mitochondrial genetic code to check for stop codons to detect pseudogenes in MacClade 4.06 (Maddison and Maddison 2000).

The best-fit substitution model of HKY+I+Γ was selected for COI and GTR+I+Γ for 28S using jModelTest (Posada, 2008). The two fragments were analyzed separately to emphasize possible mitonuclear differences. The phylogeny was reconstructed under maximum parsimony (MP), maximum likelihood (ML), and Bayesian approach. MP analyses were performed using PAUP* 4.0b10 (Swofford 2002). All phylogenetically uninformative characters were excluded from the analysis, and gaps were treated as missing data. Heuristic searches were conducted using tree bisection reconnection branch swapping, with a limit of one million rearrangements for each replicate. Bootstrap support indices were generated based on 1,000 bootstrap replicates with ten random-addition sequences. ML analysis was conducted using RAxML 8.2.9 (Stamatakis 2014), starting with 1,000 rapid bootstrap replications followed by a thorough tree search. Bayesian analyses were conducted using MrBayes 3.2.6 (Ronquist et al. 2012), with 8,000,000 generations sampling every 100 generations. A majority consensus tree was constructed to estimate the Bayesian posterior probabilities, after 25% burn-in.

Molecular operational taxonomic units (OTUs or MOTUs) are defined as monophyletic clades with distinguishable morphological or geographic properties.

**Results**

Except for some representatives of the *G. balcanicus* agg and the only member of *G. komareki* agg, all of the recently collected *Gammarus* samples from Central Asia are members of the *G. lacustris* agg (all sensu Hou and Sket 2016; Figure 1). However, for most of the previously described taxa, samples for molecular analyses have not been obtained. Unfortunately, the same also goes for
the endemic genus and species *Issykogammarus hamatus* Chevreux, 1908. None of our samples appeared to be morphologically identical to the lacustrine species *G. ocellatus* Martynov, 1930 or *G. inberbus* Karaman & Pinkster, 1977.

The phylogenetic analyses of COI and 28S markers produced very similar results. The COI dataset included a larger taxa assemblage, which allowed rough survey on the entire *G. lacustris* aggregate, while the 28S dataset was more useful for exploring the relationships between Central Asian species. The phylogenetic relationships within the *G. lacustris* agg. were largely unresolved. However, OTUs clustering for the more variable COI (Brown et al. 1979) and for 28S was similar.

In its proximal part, the phylogram of the *G. lacustris* agg. is split into a high number of long branches. Members of these long branches are also morphologically diverse, in part already described as species living throughout western Asia and the Mediterranean (Kyrgyzstan, Turkmenistan, Iran, Lebanon, Italy, Morocco).

Terminally, the tree is split into two clades, called here *G. lacustrisA* and *G. lacustrisE*. This is a morphologically homogeneous part of the tree, most of its MOTUs have been mainly recognized by different authors and eventually published as ‘*G. lacustris*’. The clade *G. lacustrisE* is the European clade. The *G. lacustrisA* contains taxa from E Asia, from Himalayas (new data by ZH), some from Central Asia, but also from N America (USA, Canada), and few from Europe (Norway, Ukraine). The morphologically distinct *G. bergi* from Ysyk Köl make this clade morphologically more diverse, although the majority of members have been mentioned in literature as *G. lacustris*.

The widely dispersed *G. cf. subaequalis*-Garlyk and the lacustrine *G. bergi* are hardly distinguishable in the 28S tree, while they are broadly separated in the COI tree. In addition to that, *G. bergi* is morphologically distinct, while *G. cf. subaequalis*-Garlyk belongs to the *G. lacustris* morphotype (Figs. 2a, b).
Figure 2: Phylogenetic tree of the aggregate *Gammarus lacustris* derived from maximum likelihood analysis of 28S (a) and COI (b) sequences. Bootstrap values (>70) are shown above branches. OTUs in both distal branches of the tree are strongly grouped and condensed. Marked subclades are from Central Asia.

Slika 2: Filogenetsko drevo agregata *Gammarus lacustris* po analizi maximum likelihood markerjev 28S (a) in COI (b). Vrednosti bootstrap (če >70) so nad vejami. OTUji v obeh končnih vejah so močno združeni. Označene podveje so v Centralni Aziji.
List of taxa and their distribution

The diversity of the Central Asian *Gammarrus* can be mainly illustrated by the analysis of the phylogram of the *G. lacustris* aggregate (Fig. 2a and b).

In detail, taxa, attributed to *Gammarrus* or *Rivulogammarus* with revised geographical data and with some comments where necessary, are listed in the supplementary data (Supplement table S2). To avoid the accumulation of weakly supported nomenclatural combinations, we are only directly citing Martynov’s taxa, since any serious revision would demand previous molecular analyses. This means that we also preserved the original nomenclature with *Rivulogammarus* S. Karaman 1931 (one of the authors - (DS) - does not agree with such solution). *Rivulogammarus* is anyway an objective junior synonym of *Gammarrus* (Stock, 1969). Note that virtually all of Martynov’s taxa (and subtaxa) are based on single populations. Molecularly studied samples are marked here in text with ‘(DNA)’. Numbers in brackets after species names are designations of OTUs in the phylogram as well as in the supplement table S1.

*Gammarrus lacustris* aggregate

The majority of Central Asian taxa and its genetic lineages are morphologically within the morphological diversity of the *G. lacustris* aggregate. Species of *G. tianshan*, *Gammarrus* sp5, *G. parvioculatus*, *G. troglomorphus*, *G. decorosus*, *Gammarrus* sp2, and *Gammarrus* sp1 are distinctly branched from close to the base of the aggregate’s phylogram. Being morphologically and molecularly so diverse, they might be the most ancient Central Asian *Gammarrus* species with a long evolutionary history in the earlier biodiversity of the region. The species *G. bergi* and *G. cf. subaequalis* are nested in the clade of *G. lacustris* with short branches, which could be explained by recent diversification of the clade *lacustris* in Central Asia. Since *G. bergi* and *G. cf. subaequalis*—Garlyk inhabit special niches, namely a brackish lake and a streaming river, this may explain morphological differences from *G. lacustris*.

(1) Close to the phylogram base, a loose clade of long-branched OTUs splits, including *G. tianshan* Zhao, Meng & Hou (432; syn. *G. montanus* Hou, Li & Platvoet, 2004), which is our only taxon from the lake Teleckoe (257) and other populations from Central Asia, and also a population from Ukraine (420) and the type population of *G. varsoviensis* from Poland. Martynov (1930b, 1935) described *Gammarrus teletzkensis* Martynov, 1930, *G. ocellatus* subsp. *angulatus* Martynov, 1930, *Gammarrus korbuensis* forma *reducta* Martynov, 1930 from the lake Teleckoe ozero (Altyn Köl), Altaj, Russia. We could not join our *Gammarrus* sp. (DNA) OTU 257 G.teleck. to any of the nominal taxa described from that lake.

(2) In the basal part of the tree, there is also a long-branched clade from Turkmenistan and Xinjiang. These are *G. decorosus* Meng, Hou & Li, 2003 (693; Xinjiang, China), *G. troglomorphus* Sidorov, Hou et Sket (513), and *G. parvioculatus* Sidorov, Hou et Sket (515), the latter ones from the extreme East of Turkmenistan. *G. troglomorphus* Sidorov, Hou & Sket, 2018 (DNA) Turkmenistan E, Lebap Province, Garlyk, is one of the most extremely troglomorphic *Gammarrus* species at all, eyeless and with long pereopods. It seems to be the only troglomorphic species and the only one with strongly carinate urosomites within the aggregate. It was found accompanied by a troglobiotic fish *Triplophysa starostini* (Parin, 1983) (syn. *Noemacheilus s.*, *Paracobitis s.*, *Troglocobitis s.*) in the collapse doline (karst sink) Suw Oyuk.

*Gammarrus parvioculatus* Sidorov, Hou & Sket, 2018 (DNA) from Turkmenistan E, Lebap Province, Köytendag District, Khodzhapil (= Hojeypil) was found in two very different springs near Köytén. Although evidently an eutroglophile, it is a comparatively short-legged species with variable pigmentation and only slightly reduced eyes.

Thus, *G. troglomorphus* and *G. parvioculatus* are very morphologically different, not only at the expense of troglomorphy. On the other hand, they are molecularly sister species in both COI and 28S analyses. These molecular relations are weakly supported (with only 45%) in COI. In the 28S tree, the grouping of *G. troglomorphus* and *G. parvioculatus* is supported with a value of only 78. In the COI tree, their further sister
southern sides of the lake (Bokonbaevo and Cholpon Ata). The species stands out of the *G. lacustris* agg. for its long and dense setosity of antennae I-II. It is roughly reminiscent to *G. komareki*, but four specimens analyzed molecularly clearly speak for its position in the *G. lacustris* agg. and away from the *G. komareki* agg. Moreover, in the tree it is surrounded by animals of the *lacustris* morphotype (i.e., morphologically identified as *G. lacustris*). An individual from deeper water (-30 m) has less setose (but still *komareki*-like) antennae. Terminal clades *G. lacustris*E and *G. lacustris*A loosely correspond to both allozyme races of Vainio and Väinölä (2003).

The other important subclade here has been designated as *G. cf. subaequalis*-Garlyk (Sidorov et al. 2018), morphologically related to the following Martynov’s taxa. *Rivulogammarus subaequalis* Martynov, 1935, with 4 subordinate taxa (and/or synonyms; Martynov’s subordinate/subspecific taxa are: subspecies, natio, morpha, forma) is from S Kazakhstan and Tajikistan; *Rivulogammarus turanus* Martynov, 1935 with 6 subordinate taxa, which was found in Uzbekistan and S Kazakhstan. As *Gammarus* (*Rivulogammarus*) *syrriacus subaequalis*, the former was supposed to be (Birštejn, 1948) ‘very common, widely spread in Tajikistan’ (springs at fortress Hisor (= Gissar, Hissar, r. Hanakinka, kishlak near st. Hanaka, kišlak Hodžambio).

Thus, *Gammarus* cf. *subaequalis*-Garlyk (DNA) seems to be (a part of) the most important taxon of the Central Asia. The Martynov’s *G. subaequalis* and *G. turanus* are morphologically mutually very close and may be phylogenetically identical with the molecularly studied OTU 516, described nowadays (Sidorov et al. 2018) as *G. cf. subaequalis*-Garlyk. The population from Garlyk (= Karlyuk), Köyten area in NE Turkmenistan resembles *G. turanus* particularly in the shape of pereopod V base, but it is very different in telson shape and setation. In telson and in uropod III dimensions and setation, it is closer to *G. subaequalis*. Our taxon is very different from both in urosomal spines. Molecularly very close to 516 (Karabulak, Garlyk) are OTUs TAD3 (Zeravshan Valley), 261 (lake Alaudin), 369 (Ysyk Köl, -10 m), 423 (lake Iskandar Köl), 465 (lake Karakul,
Pamir). Their phylogenetic relation (affinity) is also highly supported (90 BP). Considering the morphology and distribution, we suppose that these populations are really conspecific with one or both of the mentioned Martynov’s species. Seven molecularly closely related populations are distributed within the area between the easternmost Turkmenistan, Tajikistan, and up to the eastern Kyrgyzstan with ca. 1,200 km distance between localities 516 and 369 (i.e., between Garlyk and Ysyk-Köl), while the Martynov’s G. subaequalis and G. turanus are ca. 250 km away from this area to the NW. The phylogenetic relation between this taxon and G. troglomorphus, its very close geographical neighbor, is very weak.

If our supposition is correct, Gammarus subaequalis Martynov, 1935 is one of the most widespread species of the region, as indirectly already suggested by Birštejn (1948). Morphologically it is a lacustris-like species with a series of features occurring in phylogenetically diverse species within the genus Gammarus. The greatest morphological similarity outside Central Asia is with G. crinicaudatus Stock et al., 1998 from the southern Zagros region of Iran.

Gammarus balcanicus aggregate

Members of the G. balcanicus aggregate are morphologically comparatively easily recognizable. The reliability of such identification is, however, low, but in our case it does not contradict the few molecularly supported data. Characteristic of this aggregate is the poor setation of pereopods III–IV and of uropod III (Karaman and Pinkster 1987). In the latter, the spines along the exopodite outer margin are associated with only few and short finer setae.

Molecularly proven members in Central Asia are Gammarus takesensis Hou, Li & Platvoet, 2004 (DNA) from China, Xinjiang, Takes, and Gammarus taisiensis (Hou, 2002, thesis not printed) (DNA) from China, Xinjiang, Yumin. The only morphologically recognized members of this aggregate are Gammarus brevipodus Hou, Li & Platvoet, 2004 from China, Xinjiang, Xinyuan, Gammarus angustatus Martynov, 1930a with two subordinate taxa from Russia, river Ob drainage (incl. Novosibirsk), Gammarus pellucidus Gurjanova, 1930 from Russia, Krasnojarsk, Gammarus (Rivulogammarus) balcanicus Birštejn, 1950 from Russia, river Listvjanka, Jurginskij rajon, Kemerovska oblast’ and Rivulogammarus spinulatus Martynov, 1935 from Kazakhstan, E of Almaty.

Thus, we can confirm the presence of the eastern assemblage of the agg. balcanicus along the E Kazakhstan–Xinjiang (China) border and in the adjacent areas of Russia. No member of agg. balcanicus could be detected in Iran. The closest western agg. balcanicus members are probably (not DNA-confirmed) in Asia Minor – G. orientalis (S. Karaman 1934) (Karaman 2017; in Central Turkey), G. burduri S. & G. Karaman, 1959 (SW Turkey), and in Azerbaijan – G. alarodius Deržavin, 1938 (in Nakhchivan, river Araks), G. talyschensis Deržavin, 1939 (S Azerbaijan).

Gammarus komareki aggregate

The Gammarus komareki aggregate is represented by Gammarus turcomanicus Birštejn, 1945 (DNA) (syn. G. (Rivulogammarus) balcanicus turcomanicus Birštejn 1945) at the edge of the area, in Turkmenistan, rivers Aşgabatka, Kişinka, Kara-su, Zolotoj ključ, Çulinka, Sakiz-jab around Aşgabat (Birštejn 1945). Molecular analysis (new data) shows the appurtenance of the spring population of Çulinka to agg. komareki. Although described as G. balcanicus ssp., its morphology is not in disagreement with such solution. This is the easternmost known population of the G. komareki agg. Further members of this aggregate are present in Iran (Stock et al. 1998, Zamanpoore et al. 2011), in Caucasus (Birstein 1933, Behning 1940, Behning and Popowa 1947), in E Europe, and probably in Asia Minor (Alanyali et al. 2011).

Discussion

Ruling of the G. lacustris aggregate in Central Asia

Our gammarid samplings clearly show that - beside the Sarothrogammarus group in high mountains - we could only find members of G. lacustris in Central Asia. However, the core area is sparsely surrounded by members of the
G. balcanicus aggregate and by G. komareki in the West. We were also not able to phylogenetically classify the Martynov’s (1935) taxa, originating mainly in southern Kazakhstan. Members of this aggregate even inhabit aberrant habitats, such as a brackish mountain lake and subterranean waters, the latter highly morphologically modified. The G. lacustris relatives here are morphologically very diverse, not as unified as the populations in Central and Northern Europe, which could be identified as a unique G. lacustris ‘morphospecies’ (Karaman and Pinkster 1977, Alther et al 2016).

Some additional taxa had been described, for which we find no close phylogenetic relations to our molecularly studied samples. These are G. ocellatus Martynov, 1930a with three subordinate taxa from Ysyk Köl and Teleckoe ozero, indicated later by Martynov and Behning, 1948 for a wider high mountain area in Central Asia. Farther R. truncatus Martynov, 1935 with 2 subordinate taxa from Uzbekistan and Tajikistan; Gammarus gracilis Martynov, 1935 from Kazakhstan, eastern Karatau; Gammarus angusticoxalis Martynov, 1935 from Kyrgyzstan; Rivulogammarus brevicornis Martynov, 1935 from Kazakhstan S, Karatau, springs at rivulets Kichkin, Ishfan-Mazar, Martynov (1935): ‘A separate species, but close to R. turanus’.


Rivulogammarus hirsutus Martynov, 1935 (syn.: Rivulogammarus brachyurus Birstein, 1935, Gammarus birsteini Karaman et Pinkster, 1977) with 3 subordinate taxa from S Kazakhstan is morphologically very different and taxonomically intriguing. Antennae resemble G. komareki or G. bergi; according to Martynov (1935), it is close to G. turanus; the uropod III endopodite is shortened to 50% of exopodite or less. The uniquely long dorsal setae on urosomites, in Martynov’s morph ‘hirsutissimus’ even on pleonites, make its belonging to agg. lacustris or agg. komareki unlikely, but not impossible.

Birštejn (1945) considered G. hirsutus Martynov, 1935 a junior synonym of G. brachyurus Birštejn, 1935; since the name G. brachyurus appeared to have been preoccupied (Karaman and Pinkster 1977), the name hirsutus may nevertheless be used. The Karaman and Pinkster’s (1977) G. birsteini has been illustrated according to specimens from Turkey; since it lacks those characteristic urosomal setae, it is evidently a species per se.

Issykogammarus hamatus

A specific problem is Issykogammarus hamatus Chevreux, 1908. It is an aberrant taxon of an uncertain position. It was obtained from different depths of the lake Ysyk Köl, Kyrgyzstan NE. It was described as a separate genus (Chevreux 1908), later attributed to principally Bajkalian acanthogammarids as a ‘Baikalian Escapee’ (Barnard and Barnard 1983). But, if we only overlook the recast of the pereopod coxae, the species fits easily into the diversity of the aggregate lacustris or even very close to the morphotype lacustris. The evolutionary mutability of the mentioned coxal armament in amphipods can be best documented by the marine genus Epimeria (Rauschert and Arntz 2015) or by the general body shape variability in Bajkalian gammarids, nested phylogenetically within Gammarus (Hou et al. 2016). Unfortunately, a molecular analysis was impossible and the old-fashioned presentation of the morphology, conducted even mainly on females, does not allow for a serious reevaluation of the species’ taxonomy and phylogeny.

Gammarus pulex and its aggregate

Gammarus pulex and its aggregate in Central Asia remain a mystery. The only seemingly reliable data on G. pulex are those by Birštejn, 1945 from the rivers Sherlok and Firjuzinka near Asgabat (Ashgabat) (central Kopetdag, neighbor to G. turcomanicus). Explicitly mentioned are curled posterior setae on pereopod III (not so in pereopod IV) and widened and densely setose flagellar articles of antenna II; we could only hardly expect these striking characters of G. pulex pulex (Linnaeus) in G. lacustris relatives. Nevertheless, both species are very similar and the Asian members of the aggregate lacustris are explicitly diverse.
Martynov (1930a, 1935) is persistently mentioning ‘G. pulex’ for lentic(!) waters in Central Asia without giving any further data. One could suppose that he was misled by the similar G. lacustris, which he surprisingly did not mention at all. According to Karaman (1984), ‘almost not any of these (cited here) populations belong to the G. pulex’ - two pages of citations are given, more than 60 populations for the entire Asia between the Caspian and the Pacific shores, with exception of its Indian (Southeast Asia, Indonesia) region.

Ecology and ethology

Gammarus lacustris has been characterized as an euryhaline, eurythermal, and euryoxibiotic species-ubiquist (Bazikalova 1941, Birstein and Taliev 1946, Bekman 1954, Pirozhnikov 1932, Økland and Økland 1985). In the vast territory of its rather well documented geographic distribution (see also Straškraba 1966), the morphotype ‘Gammarus lacustris’ inhabits various water bodies: limnic and running waters, cold and warm, fresh and salty, rich in oxygen and hypoxic. In the Trans-Urals and in the Middle Urals, it inhabits soft fresh water with very low salt content (dry residue 60-77 mg/l) and brackish water bodies, for example in the Chelyabinsk region of the lakes Chebarkul and Cancul with dry residue 8,83 g/l (Dexbach 1952). In the Baikal region, it inhabits low-mineralized water bodies with a content of dry residue from 25 mg/l to 152 mg/l (Bekman 1954). In the lakes of the West Siberian lowland, it lives in water bodies whose salinity reaches up to 10% (Balabanova 1941, 1949).

A number of authors (Buschkiel 1931; Haempe 1908) pay attention to the concentration of calcium ions in the water. Therefore, Wundsch (1922) managed to show the mosaic distribution of Gammarus in the rivers of the right bank of the Rhine, where he stated the absence of Gammarus if the content of calcium ions was less than 9 mg/l. Experimentally, Bekman (1954) found out that the normal ontogenetic development of the Baikal Gammarus was disturbed in waters containing less than 7 mg/l of calcium.

In Central Asia, members of the G. lacustris agg. inhabit all types of perennial aquatic habitats. They have mostly been found in springs, while rivers were sampled occasionally. Since temperatures of ground waters (and springs) normally approach mean yearly temperatures of the area, we can expect the temperatures of springs to range from ca. 18 °C (at Aşgabat, at 300 m a.s.l.) in the foothills to 7.5 °C in higher areas (around Ysyk Köl at 1,650 m a.s.l.). The terrain is mainly magmatic, dictating low calcium content in water; there are smaller karst i.e., carbonate areas (particularly in SE Kyrgyzstan and NE Turkmenistan), where we may expect hard waters. An even higher level of mineralization is reached in the lake Ysyk Köl itself, where water is brackish, with 6 g/l chloride/sulfate/sodium/magnesium-based salinity (Aladin and Plotnikov 1993, Baetov 2005).

Martynov (1935) mentioned stagnant waters as a common dwelling place for G. pulex, but we only found some normal G. lacustris agg. members in some smaller lakes. Unfortunately, Martynov explicitly states only ponds close to Bishkek, the area where we found G. cf. lacustris. Several gammarids have been described from the lake Teleckoe, as well as from Ysyk Köl. Some species may be present in the depths of some tens of meters. Issykogammarus hamatus and Gammarus bergi are definitely endemic, while we cannot confirm this for G. iniberbus or G. ocellatus. I. hamatus exhibits characteristics of the ‘lacustrine morphology’ (Hou et al. 2016), while other species from this lake are morphologically trivial.

G. troglomorphus is an evidently highly specialized troglobiont. G. parvioculatus is present in both, springs and underground; its only troglobomorphies are slightly smaller eyes and integumental depigmentation in some individuals. Some other troglobiotic amphipods are also Tadzocrangonyx spp. (Sarothrogammarus group of Gammaridae) and Bogidiella spp. (Bogidiellidae; found in Turkmenistan).

Gammarids are not provided with ontogenetic stages that were particularly suited for lateral dispersal. However, appropriate behavior may enable the adult to disperse laterally, outside the water. Segerstråle (1954) has shown that the ethology of G. lacustris enables it to take the advantage of transportation by birds (ornithochory). The author could experimentally establish that this is not a general ability of gammarids, since G. pulex was behaving differently. Therefore, we also
do not know whether this is only an ethological adaptation or is it also morphologically dependent (supported). This capability may also be either inherent in the whole aggregate or limited to the lacustris morphotype. Only the remotely related Crangonyx pseudogracilis Bousfield, 1958 was later a subject for a similar experiment (Rachlewski et al. 2013). In South Dakota, Hyalella azteca (Saussure) was found in duck’s plumage (Rosine 1956). In Canada, the same amphipod was found in the fur of muskrat and beaver, while numerous G. lacustris were found in the fur of beaver (Peck 1975). We have no information about the respective behavior of other gammarids. We do not know which morphological characters contribute to the ability to hang on birds and we do not know the ethological differences and similarities between all of the numerous members of the G. lacustris aggregate. But, the exceedingly large distribution area of the G. lacustris agg. (and even of the morphotype ‘G. lacustris’) shows that this group is extraordinarily capable of dispersal. It either dispersed in morphologically diverse types, or it dispersed as the morphotype lacustris, while its populations morphologically diversified only after reaching their recent dwelling areas.

The supposed historical biogeography of Gammarus in Central Asia

Gammarus in Eurasia (Palearctic). The Eurasian continent is inhabited by phylogenetically and geographically very distinct gammarid groups (Hou et al. 2011, Hou and Sket 2016). The Asian continental freshwaters east of Ural and north of 55 °N are virtually devoid of Gammarus and mostly inhabited by Pallasea and Synurella, which regularly significantly dominates in tundra and forest-steppe in the northern biotopes. The few known populations (Deržavin 1923, Kurenkov and Mednikov 1959, Labay 1998) are either not attributable to an aggregate, or belonging to G. lacustris agg., with the exception of G. koreanus Ueno, 1940 in Sahalin, probably belonging to the Oriental aggregate. It is also necessary to take into account the fact that G. lacustris was purposely introduced almost everywhere in the lakes in the Central Russian Upland, Siberia and even in Kamčatka during the Soviet times. These doings have not always been properly documented. The eastern portion of China and areas NE from it (Japan and Korea, Russian Far East) are inhabited by the Oriental aggregate of Gammarus; Lake Bajkal has its own endemic fauna, nominally of diverse genera (and even families, e.g. Tahteev 2000), which are in fact phylogenetically members of Gammarus and probably sister to the aggregates balcanicus and lacustris (Hou and Sket 2016). In the West, the European aggregates G. pulex, G. fossarum, and G. roeselii do not cross the eastern borders of Europe. Two SE European aggregates are extending towards Central Asia. The G. komareki agg. is richly represented in Iran and with G. turcomanicus Birštejn, 1945 (syn. G. balcanicus turcomanicus) in S Turkmenistan at Aşgabat (Ashgabat). The repeatedly mentioned G. pulex (e.g., Martynov 1935) remains a total mystery regarding Asia; molecularly, the morphologically similar aggregates G. pulex, G. fossarum, and G. roeselii could have only been verified for the European territory, with inclusion of Georgia (Hou et al. 2014, Supplement table S1).

The G. balcanicus aggregate occupies the E and SE Europe with some probable, but not molecularly proven representatives between Europe and the Caspian (see above). Molecularly proven members are only present in Ukraine and further west (Hou et al. 2014). In SE Europe, G. balcanicus agg. populations, cited mainly simply as ‘G. balcanicus’, are evenly distributed, although populations of three other aggregates are also present there. Karaman and Pinkster (1987) mention more than 1,000 samples (of the supposed unified species!) in their area, Poland to SW USSR.

However, an additional detached area exists with a small number of reliable agg. balcanicus members (Hou and Sket 2016) along the border between Xinjiang and Kazakhstan, and (not molecularly proven) in the adjacent areas of southern Sibir’ (Siberia, Russia). This results in more than 2,000 km wide gap between both parts of the entire agg. balcanicus distribution. This gap includes the heart of Central Asia. In recent years, we sampled ca. 40 springs and clear lakes in Central Asia, typical habitats of ‘Gammarus balcanicus’, but not one sample contained balcanicus-like animals. It was molecularly proven that they all belong to the G. lacustris agg. Even if some undiscovered populations of the G. balcanicus agg. do exist
closely positioned UTM quadrates, the agg. where agg. pulex areas of central and northern parts of Europe, the situation is similar in the Alpine and Scandinavian ca. 40 quadrates, G. elvirae in ca. 45 quadrates of 10x10 km (Ruffo 2003), agg. p.p.).

Echinogammarus inhabited by the related man 1993). However, all these areas are richly are even ‘very similar to (Kara-

G. lacustris’ representatives are limited (with a small exception) to the continental North of the country, while Sicilia (Sicily) and the French Corse (Corsica) are devoid of a freshwater Gammarus. Some populations of ‘G. italicus’ from the central Italian lakes (later separated as G. elvirae Iannilli & Ruffo, 2002) are even ‘very similar to G. lacustris’ (Karaman 1993). However, all these areas are richly inhabited by the related Homoeogammarus (syn. Echinogammarus p.p.).

Within the peninsular part of Italy (Stoch and Ruffo 2003), agg. balcanicus is present in two closely positioned UTM quadrates, the agg. lacustris in ca. 45 quadrates of 10x10 km (G. elvirae ca. 40 quadrates, G. lacustris 5 quadrates). The situation is similar in the Alpine and Scandinavian areas of central and northern parts of Europe, where agg. pulex is somehow limited, while the others are close to absent, and G. lacustris agg. (by G. cf. lacustris) is mainly common in lakes (Fryer 1953; Segerstråle 1954, Alther et al. 2016). On the contrary, in Slovenia, which lies at the SE edge of the Alps, we registered G. fossarum agg. from 120 localities (but hundreds of noted remain unregistered), G. roeselii agg. from 13 in the East, G. balcanicus agg. from 6 scattered localities in the West, and G. lacustris agg. only in one lake in pre-Alps (our own data).

Phylogeography of Gammarus taxa in Köytendag

Very instructive, predicting the next surprises in the Gammarus fauna of Central Asia, are the relations among the eastern Lebap (Köytên-Garlyk (=Karlyuk)), extreme East of Turkmenistan) species. Two cave-related species (G. trogloomorphus, G. parvioculatus) are morphologically the most dissimilar species of Central Asia, and among the most divergent species pairs within Gammarus at all, which is only to some extent the result of troglomorphy. As exposed above, both cave-related species are extremely morphologically different, and molecularly loosely, but firmly related. Their localities are ca. 40 km apart, while their probably (with low support again) closest relative is G. decorosus Meng, Hou & Li, 2003 from ca. 2,000 km away in Urumqi, Xinjiang, which is not inclined to inhabit subterranean waters. All of the mentioned species represent relatively long branches in the proximal part of the aggregate phylogram, with contrast to the basal G. tianshan and others.

The third species from the Köytên–Garlyk area is clearly epigean, geographically the closest neighbor of G. troglomorphus at Garlyk, but evidently broadly dispersed. We tentatively named it G. cf. subaequalis-Garlyk (Sidorov et al. 2018). See above for some more details. It is a member of a small clade of 7 samples (our OTUs 261, 423, 465, 516, 369, 359, S135) in the tree from the 1,000 km long stripe between Garlyk (=Karlyuk) in Turkmenistan, Tajikistan, and Ysyk Köl area in Kyrgyzstan. Only future field sampling can enable us to tell whether these series of populations is conspecific with the named Martynov’s taxa. The samples of G. cf. subaequalis-Garlyk are molecularly close or nearly identical, and as a group very close to the branchlets of the terminal part of the tree, its clade lacustrisA, including numerous samples from Eurasia and North America. The endemic G. bergi Martynov from the lake Ysyk-Köl is also included. Only ca. 300 km to the NW from the central part of this area are Martynov’s (1935) localities of G. subaequalis and G. turanus, which are morphologically similar and might be
both, mutually conspecific and conspecific with the mentioned group.

The climatic past possibly influencing the distribution of gammarids in Central Asia

Central Asia has been the area of the highest aridity because of the rain shadow caused by Tibetan Plateau and some other mountain ranges (Caves et al. 2014). This aridity was increasing during the Cenozoic. The precipitation should have ‘more than halved’ over the Neogene. It is likely that this aridity extirpated the majority of the previously evidently existing agg. balcanicus populations; this opened the territory for the invasion of the vagile and euryoecious G. lacustris agg., which may have happened several times during short climatically less adverse periods.

The Central Asian taxa are very diversely positioned in the phylogram; some have branched off close to the tree root, while some are members or sisters of the most distal branches. One has to bear in mind that the distribution area of the entire G. lacustris agg. covers the entire area of the genus Gammarus and even crosses the other gammarid species (Hou et al. 2011, Hou and Sket 2016, Väinölä et al. 2017). To a great degree, it occurs even in the shape, morphotype of G. lacustris, and is therefore often morphologically identified and published as ‘Gammarus lacustris’.

Such distribution can probably be explained by the ability of G. lacustris to be transported by birds, as shown by Segerstråle 1954, hand in hand with the above-mentioned ecological universality of ‘G. lacustris’. Waterfowl from the North (here Jamal and Tajmyr,) migrate in a wide front, but birds often fly along the coast of the seas and along large rivers (here Irtyš) and Central Asian lakes (Thompson et al. 2001, Šnitnikov 1980, Kydryaliev 1990). Therefore, the populations of amphipods can be steadily replenished in their basins. The erratic spreading of this group is also supported by the fact that molecularly very close populations occur in very distant localities, including the sisterhood between the populations from North America and Ysyk Köl.

The morphological diversity of the G. lacustris agg. members shows its evolutionary plasticity and a possibility that its populations (and future species) could have distributed at different times, but every one probably still in the ‘original’ lacustris morphotype, i.e. conquistador’s appearance. Thus, numerous subsequent colonizing groups formed lacustris-morphotype populations and diversified only after colonization of new territories, in this case in Central Asia. Such diversification might be a consequence of the enforcement between at least some lake (e.g., in Ysyk Köl) populations. No case of species sympatry within the agg. lacustris has been recognised outside Central Asia.

Precipitation quantity had been fluctuating in the near past (Wolff et al. 2017), and it was probably so during the whole Cenozoic period as well. Therefore, the G. balcanicus agg. could have spread its area from the West into Central Asia and beyond either before the onset of aridification, or also during a longer or shorter wetter period within its course.

The only additional gammarids (i.e., non-Gammarus) in Central Asia are members of the eastern (Asian) clade of the Sarothrogammarus group of genera (Hou and Sket 2016, Barnard and Barnard 1983), including Tadzocrangonyx spp. They are ecologically specific, living mainly in springs and to some extent in the subterranean, between 2,000 and 3,000 m a.s.l., and hardly ever below 1,000 m a.s.l. They are limited to the SE part of Central Asia: Uzbekistan - Tajikistan - N Afghanistan – N Pakistan. It is possible that different ecological regime in high elevations prevented extirpation of the Sarothrogammarus gammarids.

Conclusions

(1) Recent samplings of gammarids in the core area of Central Asia, approximately between E Turkmenistan - N to E Kirgyzstan – Tajikistan, which we were able to study molecularly, delivered only members of the G. lacustris aggregate. The area is sparsely surrounded by members of the G. balcanicus aggregate. Members of the related Sarothrogammarus group of genera are only present in springs and mainly in higher elevations.

(2) The G. lacustris aggregate appeared to consist of species with high phenotypic and evolutionary plasticity, resulting in a morphologically diverse and geographically widely distributed
taxon. Its distribution area is broader than that of the rest of the genus *Gammarus*. In addition, within the genus’ area, it is present where other species of *Gammarus* are absent – including in Central Asia. (3) While the *G. balcanicus* relatives and other Palearctic groups are evidently only able to disperse along streams, the *G. lacustris* relatives are able to disperse laterally outside water, by ornitochory. We do not know how general this ability is within the aggregate.

The *Gammarus* assortment in C Asia consists of species from all parts of the phylogram. They evidently immigrated in part in different times from far abroad, but some speciation must have happened within the area, to a smaller extent probably even within the lake Ysyk Köl (Issyk Kul) alone.

No members of the generally European clade *G. lacustris* were found in Central Asia, but Central Asian species are in close relationship with other species from Europe, Asia, and North Africa – and even N America. (4) The absence of other *Gammarus* groups in this wide arid area and its paleoclimatic history indicates that the – probably originally richer - amphipod fauna in the lower regions could have been extirpated in particularly arid periods. The *G. lacustris* relatives could survive to some extent for their euryoecity and be restored by a steady import and immigration. The *G. balcanicus* populations could only survive sheltered in springs in marginal positions of the area.

So, a great gap in the Gammaridae diversity appeared in Central Asia (5) The morphological diversity of the *G. lacustris* agg. in Central Asia and also in general nearly equals the diversity of the European *Gammarus* spp.

The generally very poor subterranean fauna in Central Asia is represented by the troglobiotic and highly troglomorphic *G. troglomorphus*, and by the very stocky and still occulated eu-troglophile *G. parvioculatus*.

**Povzetek**

Ob raziskovanju podzemeljske biotske pes-rosti v Köytendag, Turkmenistan, v okviru misije britanskega Royal Society for the Protection of Birds, smo opazili nenavadno sestavo favne postranice rodu *Gammarus*. To nas je napeljalo na podrobnejšo raziskavo postranice v Centralni Aziji.


Naša raziskava zajema v Turkmenistanu Aşgabat z okolico in Köytendag na skrajnem sverovzhodu; v Kirgiziji jezero Ysyk Köl (Issyk Kul) z okolico ter jugozahod dežele; v Tadžikistanu kraje na severozahodu in jugovzhodu. Vse lokalitete in zbiralci so našteti v tabeli. V največjih meri smo vzorčili izvire in jezero, le malo pa reke. Živali smo molekulsko identificirali do agregatov, ‘skupin vrst’, ki se sploh ne ujemajo z izključno morfološko določenimi skupinami Karamana in Pinksterja. Za določitev vrst se zanesemo le na vzorce iz topotipskih lokalitet, v kolikor so tudi morfološko primerni. Pri vseh drugih veljajo imena le pogojno in so označena s ‘cf.’.

Za molekulsko analizo smo vzelih majhne dele živali, preostanek takšnih osebkov smo spravili za poznejše natančneji morfološko in taksonomsko obdelavo. DNK smo izolirali s pomočjo Qiagen kita in pomnožili mitohondrijski COI ter jedrni 28S gen s primeri, ki so opisani v Hou et al. (2007).

Razen maloštevilnih pripadnikov agregata balcanicus in vzorca iz agregata komareki smo vse vzorce molekulsko razkrili kot pripadnike agregata lacustris. Filogenetske analize s COI in 28S so dala zelo podobne rezultate. Žal nismo imeli na voljo aberantnega endema Issykogamarus hamatus Chevreux.


Starejši avtorji ga omenjajo pogosto, večinoma brez opisov, zdaj pa ga nismo našli in te vrste v Aziji verjetno ni. Od drugih rodv v Srednji Aziji živi rodovna skupina Sarothrogammarus; tukajsojna podskupina je vezana na višje lege in jo široko ločuje od mediteransko-atlantske podskupine rodov in vrst. V Centralni Aziji smo našli gamaride v vseh tipih voda. Tako v mehkih, kot v trdih in v somornici, v potokih, bistrih mlakah, jezerih in izvirih, od ca. 300 do nad 2,000 m n.m. V geološki preteklosti je Srednja Azija utrpela aridifikacije, ki so nedvomno povzročile siromašenje voda ter izumiranje manj evriekih gamaridov. Zlasti v vlažnejših obdobjih je bilo tako olajšano poseljevanje vod v izjemen G. lacustris.

Za razliko od srednje in severne Evrope, kjer je agregat lacustris zelo vezan na jezera, se tukaj pojavnja v vseh tipih voda. Gamaridi nimajo tega prikaznjen, posebej prirejenih za razširjanje, pač pa so za G. lacustris ugotovili možnost razširjanja s ptiči. Domnevamo, da so se na ta način razširili v Srednjo Azijo in po njej. Na ta način domneviva vrsta G. lacustris (torej morfotip tega imena) zaseda območje, ki presega siceršnji areal rodu Gammarus in celo družine Gammaridae.

Rod Gammarus v Srednji Aziji zastopa skoraj izključno agregat G. lacustris, le po robu semkaj sega še agregata balcanicus in komareki. Domnevamo, da so takšni poselitvi prispevala izumiranja gamaridov v izjemno sušnih obdo-
bjih. Tem je sledilo poseljevanje z evrieko vrsto *G. lacustris*, ki se morda edina razširja tudi s ptiči, in se je diverzificirala po naselivti v Srednji Aziji. Zdaj tako na območju najdemo med drugim tudi ekstremno troglomorfnost vrsto tega agregata in zelo aberantno jezersko vrsto.

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List of supplements (web edition)

**Supplement S1:** List of molecularly analyzed samples of the *Gammarus lacustris* aggregate, adequate to the tree in Fig. 2. Top samples (from the tree top to first Tajikistan samples) are only few representatives of the morphotype *lacustris* from Europe, the rest are all available samples of the aggregate. Highlighted samples are from Central Asia s.l..

**Supplement S2:** List of *Gammarus* spp., described from Central Asia, but not studied molecularly. Nomenclature and taxonomic subdivision are taken from original papers; the recent geographic, administrative, and orthographic changes have been considered as much as possible. For practical reasons, the original naming of taxa has been retained.

**Supplement S3:** Distribution map of *Gammarus* spp. in Central Asia. Red asterisks with names - important cities. Reversed triangles - molecularly not defined *Gammarus* spp., letters indicate abbreviated taxonomic names (see Suppl. S2). Rings - molecularly defined taxa of agg. *lacustris*, numerals are OTU designations (as in the second column in Suppl. S1); pale ribbons connect localities of *G. cf. subaequalis*-Garlyk, hatched ribbon connection with Martynov’s *G. subaequalis* and *G. turanus*. Squares - agg. *balcanicus*; diamond (*G. turcomanicus*) - agg. komareki. Adequate to Figure 1 in the paper.

**Supplement S4:** Additional taxonomical clarification. (1) Type population of *Gammarus lacustris* Sars, 1863. (2) Identity of ‘*G. balcanicus*’ from Alma-Ata. (3) Corrected classification of ‘*Anisogammarus*’ *madyensis* Chaudri, Ghauri & Mahoon, 1987.