Bioacoustics and systematics of the *Poecilimon heroicus*-group
(Orthoptera: Phaneropteridae: Barbitistinae)

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Abstract. The species of the *Poecilimon heroicus*-group occur around the Caucasus (from north-eastern Turkey to south-eastern Ukraine). We describe the diagnostic morphological characters of all these species and the male calling song of three of the four species. Based on this data the following phylogenetic relationship is derived (*P. tschorochensis* (*P. tricuspis* (*P. heroicus*, *P. bifenes-tratus*))). Within the genus *Poecilimon*, the species can be recognised by a relatively wide pronotum and large tegmina. In one species, *Poecilimon tschorochensis* Adelung, 1907 (type species of the monotypic genus *Artvinia* Karabag, 1962, syn. n.; *P. rammei* Miram, 1938, syn. n.), the tegmina are very large and the song has unusually low spectral components. This species produced di-syllabic schemes at intervals of about 10 s. In two other species of the group, *P. heroicus* and *P. bifenes-tratus*, the calling song of males consists of an uninterrupted dense sequence of long syllables (syllable duration around 0.5 s; ca. 1 syllable/s at 20°C). In these species the auditory spiracles are reduced in size in both sexes, and the females have extremely small tegmina and are unable to respond to the male song acoustically, which would be typical for Phaneropteridae. The change in communication from acoustically responding to mute females has not been previously documented within a group of closely related species.

INTRODUCTION

Bush-crickets produce their calling songs by stridulation using their fore wings. In *Poecilimon* and other short-winged members of the Barbitistinae this seems to be the only function of the tegmina, which therefore should be modified to optimise sound production. Despite being their only function, the size of the tegmina varies greatly. In some species they are of moderate size, in others small to very small and completely concealed under the pronotum (e.g. Heller, 2004). In *P. tschorochensis* (including *P. rammei*), however, they are huge, covering – according to the description – nearly the whole abdomen. Details of the unusual song of this species are presented.

Evidence is presented that the loss of the acoustical females occurred within this group, in contrast to previous studies of other species groups of *Poecilimon*, which are homogeneous in this respect (Heller & Lehmann, 2004; Heller & Sevgili, 2005; Lehmann, 1998). The group is also interesting from another aspect.

MATERIAL AND METHODS

We studied specimens preserved in the following collections: CH (followed by the specimen code) – Collectio Heller; HUZOM – Hacettepe Üniversitesi Zooloji Müzesi, Ankara, Turkey; MSPU – Moscow State Pedagogical University, Moscow, Russia; ZIN – Zoological Institute, Russian Academy of Sciences, St. Petersburg, Russia; ZMHB – Museum für Naturkunde der Humboldt-Universität, Berlin, Germany; ZMUM – Department of Entomology, Moscow University, Moscow, Russia. Other material is deposited in ZMUH – Zoologisches Museum Universität Hamburg, Hamburg, Germany.
Morphological structures were examined and measured using an Olympus stereomicroscope. For sound recording in the field an Uher tape recorder 4200 IC or a Sony WM-D3 cassette tape recorder with a Uher M645 microphone was used (frequency response flat up to 20 kHz resp. 15 kHz), in the laboratory a Racal store 4 D tape recorder with Briel & Kjaer 4133 and 4135 microphones (frequency response flat up to 40 resp. 70 kHz), or a modified tape recorder “Yupiter-202 Stereo” with linear characteristic in the range 0.063-70 kHz, or a PC with A–D converter card L-305 (L-card) (0–300 kHz) and 00 017 RFT microphones (frequency response flat up to 70 kHz) was used. Songs of some specimens of *P. tschorochensis* were recorded in the laboratory using a Sony WM-GX688 Walkman and a stereo microphone (50 Hz to 18 kHz) at 29°C. After digitising the songs on a PC or an Apple computer, oscillograms (after filtering) and sound analysis were made using a PC and the programs Turbolab, Amadeus (Apple) and CoolEdit. Time measurements are given as mean ± standard deviation if not otherwise stated.

For the spectral measurements a 10 ms portion (sampling rate 160 kHz) of the respective song was analysed and in the figures the mean of 524 point FFT frames was presented. Wing movements were registered by an opto-electronic device (Helversen & Elsner, 1977; modified as in Heller, 1988). Due to the heating effect of the registration lamps, only a lower limit for the body temperature can be given (see Heller, 1988). The actual body temperature may be up to 4 degrees higher than the ambient air temperature (measurements marked by “>”). Some sound data for *P. heroicus* are available at the taxonomic database SysTax.

The stridulatory files were studied using scanning electron microscopes S-405 A (Hitachi) and JSM 840A (JEOL).

The map was produced using the program Versamap (http://www.versamap.com/)

Bioacoustic terminology

**Calling song:** spontaneous song produced by an isolated male.

**Syllable:** the sound produced by one opening-closing movement cycle of the tegmina. Opening/closing hemisyllable: the sound produced by one opening or closing movement of the tegmina.

**Syllable duration:** time from beginning of the first to end of the last impulse. **Syllable interval:** time from end of last impulse to beginning of first impulse of the next syllable. **Syllable period:** syllable duration + syllable interval. **Impulse:** a simple, undivided, transient train of sound waves generated by the impact of one tooth of the stridulatory file. **Pulse:** a long train of sound waves, resulting from the fusion of several impulses. **Afterclick:** an isolated impulse after the end of the main impulse group.

RESULT

**Genus Poecilimon Fischer, 1853**

Type species: *Poecilimon jonicus* (Feiber, 1835) by subsequent designation (Kirby 1906: 375, as *Poecilemon*).

*Artvinia* Karabag, 1962, syn. n. Type species: *Poecilimon tschorochensis* Adelung, 1907.

In 1962, Karabag established the genus *Artvinia* for *P. tschorochensis*. This species has certainly some unique characters, but is without doubt quite closely related to the species of the *Poecilimon heroicus*-group (see above) and should therefore be considered as an aberrant member of this genus. It is the only *Poecilimon* species of...
with huge, balloon-like tegmina (Fig. 1). In the structure of the pronotum, another character used by Karabag (1962) to define *Artvinia*, it comes close to *P. heroicus* and allied species. Large dorsal glands on the first or second tergite have evolved independently in several species of *Poecilimon* (Heller & Lehmann, 2004). In *P. tschorochensis* a large, gland-like structure covered with hairs is situated on the second tergite and smaller ones on the first tergite and even metanotum. Small glands, indicated by an elevation of the central part of the posterior border, can also be found on the first tergite of *P. heroicus*. They occur, however, also in other species like *P. affinis* (own observations) and are probably often overlooked because they are covered by the tegmina. The relationship between *P. tschorochensis* and the genus *Phonochorion*, another group of species with dorsal glands, is unclear. *Phonochorion* is also endemic to Georgia and north-eastern Anatolia, but is very different from *P. tschorochensis* (and other *Poecilimon*) in morphology (compare figures in Karabag 1956 and Karabag 1962) and song (*Phonochorion*: continuous song (Heller 1990), *P. tschorochensis*: isolated pairs of syllables, see below).

**Characteristics of the *Poecilimon heroicus*-group**

From a direct comparison of most European and Anato-
lion *Poecilimon* species (e.g., Heller, 1984; Sevgili, 2001; Heller & Lehmann, 2004; Heller et al., 2005; unpubl. studies) and the literature, it became evident that this group of species of this genus can be separated and recognised by the following characters:

1. pronotum broader posteriorly (pronotum at its widest more than 1.4 times width of the head; Fig. 1)
2. body size relatively large to medium compared to other *Poecilimon* species
3. hind femora with ventral spines
4. song consisting of long syllables (ca. 0.3–0.6 s at 18°C; Figs 2–3)

Members of the *Poecilimon heroicus*-group are recognised by a combination of characters. But by far the most characteristic feature is the posteriorly widened pronotum (1), which can be considered as a unique autapomorphic character of the group. Some or even all of the other characters are possibly plesiomorphic. Body size (2) is a character difficult to assign as either an apomorphic or plesiomorphic state. The possession of spines on the ventral edges of the hind femora (3) is also difficult to evaluate. The spines may be helpful to free entangled hind legs in dense vegetation (e.g. *Cuscuta*: personal observations KGH) and therefore more important for species living near the ground than for bush or tree inhabiting forms. It is certainly a plesiomorphic character for Orthoptera, but the situation for the Barbitistinae is unclear. A slow stridulatory movement (4) was assumed to be a plesiomorphic character of the subfamily Barbitistinae by Heller (1990).

Within the Aegaeid genus *Poecilimon* (Ciplak, 2004) the *P. heroicus* group may be related to other groups with large body size like the *P. ornatus/affinis* and *P. sancti-pauli* groups (Heller & Sevgili, 2005). Most members of these three groups share a calling song consisting of long
syllables with numerous impulses, hind femora with ventral spinules and a wide fastigium (narrow in *P. tschorochensis* and *P. tricuspis* of the *P. heroicus*-group). The *P. heroicus*-group can be considered as the Caucasian branch within this relationship, with the *P. ornatus/affinis*-group occurring in the Balkans and the *P. sanctipauli*-group in western Anatolia.

**Distribution.** Caucasus and surrounding regions (from southern Ukraine (Donets Hills) to north eastern Turkey; see Fig. 11).

**Description of species**

(see Table 1 for a review of characters)

*Poecilimon heroicus* Stshelkanovtzev, 1911

_Type:* Ukraine: SE, Provalskii stud farm on the Donets Hills (ZIN, lectotype male).

*P. heroicus*: Stshelkanovtzev, 1911: 175, Fig.14 (description); Uvarov, 1915: 88 (distribution); Stshelkanovtzev, 1921: 135 (distribution); Uvarov, 1928: 77–78 (key) Mushinsky, 1930: 65, 68–69 (agricultural pest, biology); Bey-Bienko, 1930: 55 (distribution), 1932: 5 (biology), 1933: 318 (distribution); Ramme, 1933: 506, 509 (arrangement), 511 (checklist), 518–519 (description); Tarbinsky, 1948: 84 (key); Ramme, 1951: 416 (distribution); Bey-Bienko, 1954: 255 (key), 281–3 (description); Harz, 1969: 99, 113 (key), 141 (description); Misthenko, 1972: 22 (agricultural pest); Warchalowska-Sliwa et al., 1995: 28 (karyotype); Warchalowska-Sliwa, 1998: 157 (karyotype); Otte, 1997, 1999: 123 (catalogue); Heller et al., 1998: 29 (check-list); Naskrecki & Otte, 1999: (catalogue); Zhantiev & Korsunovskaya, 2005: 107, 109 (song). Databanks: SysTax, OSF2, FaEu.


**Remarks.** The description is based on a series of syntypes (13 males, 4 females), which were deposited in the Zoological Museum Moscow. However, today no specimen of this series is found there. Bey-Bienko (1954) apparently examined parts of the type series and mentions one male as type and two other (one male, one female) as paratypes, all of them now deposited in ZIN and labelled as type and paratypes. We accept the selection of Bey-Bienko (1954: 283) and designate these specimens as lectotype and paralectotypes.

The location of the other syntypes is unknown. In any case, they are not, as mentioned for the holotype (!) by Otte (1997), Naskrecki & Otte (1999) and OSF, in Instytut of Zoologii, Polska Akademia Nauk, Warsaw, Poland and have never been there (A. Liana, pers. comm.).


**Previous records.** Russian Federation: Kabardino-Balkar (Warchalowska-Sliwa et al., 1995), for other records see list in Bey-Bienko, 1954.

**Bioacoustics.** The calling song consists of long, homogenous sequences of syllables (Fig. 2). These sequences last many seconds or even minutes (longest recorded sequence 85 s). The mean duration of a syllable
was between about 400 ms (n = 6 males from Ukraine; T = 18.5–22°C) and 500 ms (male from Caucasus: 486 ± 28 ms, n = 8 syllables; T = 22°C), that of the interval between 600–900 ms (n = 12, T = 22°C) and 1218 ± 85 ms, n = 8 syllables; T = 22°C), that of the interval between 8.7 and 15.2 ms (mean values of 2 males, T > 18.5°C; 25 and 33 ms, T = 18.5°C). The syllables are produced only during the closing movement of the tegmina (Fig. 3). The steps during the closing movement probably indicate the contact of the scraper with a single tooth of the stridulatory file resulting in a sound impulse (see also below). The syllables were composed of 20–38 impulses. The impulse period in the second half of the syllable varied between 8.7 and 15.2 ms (mean values of 2 males, T > 24°C), increasing at lower temperatures (e.g. 23 ms, T = 18.5°C; 25 and 33 ms, T = 22°C; n = 8–10 syllables/male). The carrier frequencies of the song had the maximum at 12 kHz (bandwidth 10 dB below peak: 10–17 kHz; n = 2 males; Fig. 4).

From Fig. 2 it can be recognised that uniquely the male opened and closed the tegmina producing only a weak sound. Closer examination revealed that in this case, just before sound production – during the opening phase of the tegmina – a male of another species (Fig. 9). It is necessary to obtain additional material to verify the taxonomic status of this form.

Poecilimon bifenestratus Miram, 1929

Type: Russian Federation: Krasnodar area, Western Caucasus, mountains near Gaiduk, 8 km northeast (NO in German) of Novorossijsk, 28.vi.1928, leg. Novickij (ZIN, holotype male; in addition in ZIN 1♀, 1♂ paratypes).


Remarks. In the description of P. bifenestratus Miram (1929) surprisingly did not compare it with or refer to P. heroicus. She explicitly stated that there is no species similar in size and morphology to P. bifenestratus in the Russian Federation and compared it only with Balkan and Aegean species. P. bifenestratus and P. heroicus, however, are very similar. The males can be recognised by small, but distinct differences in the shape of the cerci (Fig. 5), observed in all specimens examined. As described by Bey-Bienko (1954) the pronotum is slightly narrower in P. bifenestratus than in P. heroicus, but the tegmina are longer. Females of P. bifenestratus have distinctly longer hind legs and ovipositor than P. heroicus.

In addition, Bey-Bienko (1954) mentions differences in coloration. The colour, however, is very variable and we saw green specimens without the typical black “window” (bifenestratus) pattern on the pronotum, pinkish-yellow males and males with typical window and black colour on abdomen in the same habitat (OK; same observations by M. Stolyarov, pers. comm.).

One male, collected by M. Fyodorova in Northern Caucasus near Pjatigorsk, in July 1989, differs slightly in morphology and song from the Ukrainian animals. Zhan-tiev & Korsunovskaya (2005) refer to it as Poecilimon sp. cf. heroicus. Its song had distinctly larger syllable intervals (ca 5 s at 26°C) than that of other specimens, but otherwise similar syllable structure (Fig. 3). The number of teeth on the stridulatory file is lower (62; n = 1) than in P. heroicus (Fig. 9). It is necessary to obtain additional material to verify the taxonomic status of this form.

Poecilimon bifenestratus tschorochensis

Type: Ukraine: Western Carpathians, Tschoroch area, 850 m, 19.xii.1937, leg. Novickij (ZIN, holotype male; in addition in ZIN 3♀, 1♂ paratypes).

P. bifenestratus tschorochensis: Miram (1929)

Previous records. Russian Federation: Type locality Miram, 1926; Gelendzhik, Ramme, 1933 (this locality is cited by Shchelkanovtsev, 1911 were heard singing simultaneously with P. bifenestratus, even during the day.

Poecilimon tricuspis Miram, 1926


P. tricuspis: Miram, 1926: 276–277 (description); Ramme, 1933: 506, 509 (arrangement), 511 (check-list), 518–9 (description); Ramme, 1951: 416 (distribution); Bey-Bienko, 1954: 255 (key), 280–281 (description); Otte, 1997: 127 (catalogue); Heller et al., 1998: 30 (check-list); Chernyakhovskiy, 1994: 106 (check-list); Naskeccki & Otte, 1999: (catalogue); Stolyarov, 2005: 66 (distribution); Özbeg & Aslan, 1996 (misidentification, see below). Databanks: OSF2.

Remarks. P. tricuspis is easy to separate from other members of the group by the shape of its cerci (compare Fig. 5 and 6).

In contrast to P. heroicus and P. bifenestratus, the tegmina overlap dorsally in the female so that she is able to produce sounds.

This species, which was known only from a few localities in Northern Ossetia, was recently recorded from north eastern Turkey (Özbeg & Aslan, 1996). However, after examination of their material (preserved in Atatürk University-Erzurum, determined as P. tricuspis by E. Tazegül) and of new material from the same locality, it became obvious it was not P. tricuspis, but another Poecilimon species from the P. zonatus-group (HS).


Previous records. Russian Federation: Northern Ossetia: type locality Miram 1926; Dargavs, Miram 1926 (as unidentifiable; identification by Bey-Bienko, 1954); Balta, Bey-Bienko, 1954; near village of Dargavs and near village of Saniba, Stolyarov, 2005.

Bioacoustics. Song unknown.

Poecilimon tschorochensis Adelung, 1907

Type: Turkey, Arvin district, “Passhöhe Jalanus-Tscham. Gegen 9000’ Höhe” (= Yalnizcam pass), 1.vi.1898, leg. Deryugin (ZIN, holotype male).


P. tschorochensis: Adelung, 1907: 66–68 (description), plate 1; Ramme, 1933: 506, 509 (arrangement), 511 (check-list), 518–9 (description); Ramme, 1951: 416 (distribution); Bey-Bienko, 1954: 256 (key), 286–287 (description, as P. tschorochensis [sic]); Karabag, 1958: 32 (faunistic catalogue); Karabag, 1962: 9–10 (description; transfer to genus Arvinia
as its type species); Otte, 1997: 126 (catalogue, as *P. tchorochensis* [sic]); Naskrecki & Otte, 1999: (catalogue, as *P. tchorochensis* [sic]). Databanks: SysTax, OSF2 (as *P. tchorochensis* [sic]).


**Remarks.** The great similarity of *P. rammei* and *P. tchorochensis* has been known and discussed for a long time but few people have seen both species together. Miram (1938) noted in her description of *P. rammei* the similarity of this species to *P. tchorochensis*, but gave no diagnostic characters. Ramme (1951) examined only *P. rammei* and concluded from the description of *P. tchorochensis* that the shape of the cerci were different in these species. Bey-Bienko (1954) was obviously the first to examine specimens from both species. In his material (3♂, 4♀ of *P. rammei*, 2♂ of *P. tchorochensis*) he found mainly differences in coloration and the shape of the cerci. Karabag (1962) also did not see specimens of *P. rammei*, but in his study of *P. tchorochensis* he assumed it would be the same species. Studying the descriptions and examining our rich material, mainly of *P. tchorochensis*, we found no distinct differences between them. Coloration and the shape of the cerci (Fig. 6) are nearly identical. There are also no differences in the width of fastigium and structure of the last tergite. *P. rammei* Miram, 1938 is therefore considered to be a synonym of *P. tchorochensis* Adelung, 1907.
Material examined. Turkey: Artvin: holotype (ZIN); 15♂, 2♀ Yalnizcam pass, eastern slope, 2200–2400 m a.s.l., 10.–11.viii.1983, leg. K.-G. Heller (CH) (song heard in many places on both sides of the pass, in the northern end down to about 1700 m); 15♂, 13♀ Yalnizcam pass, 9.viii.1987, leg. K. Reinhold (CH); 1♂, Gurdzhan (determined to be situated at 41°21’40˝N, 41°25’37˝E), two labels, one with Cyrillic letters, one with Latin letters: Guzhang, 28.v.1910, leg. Nesterov (hand written, possibly by Miram or Ramme), paratype P. ramrei (ZMHB); 1♂ Badzgeret (41°22’42˝N, 41°39’17˝E), 8.viii.1909, leg. A. Satunin (ZIN); 2♂, 2♀, Kutul, 2100 m, 2.viii.1972, leg. A. Koçak (HUZOM); 15♂, 6♀, Ardanuç, Kutul Yaylasi, Orman kenari, 41°04.373 N, 42°13.245 E, 2230 m, 26.vii.2002, leg. H. Sevgili (HUZOM); 8♂, 1♀ (on Urtica sp.), Bilbilan yaylasi, 41°03.567 N, 42°16.688 E, 2485 m, 26.vii.2002, leg. H. Sevgili (HUZOM); 7♂, 1♀, Yalnizçam dagları, Kutul geçidi, 2150 m, 2.viii.1973, leg. S. Salman (HUZOM); 1♂, 2♀, Savsat, Meselik köyü yaylasisi, 2000 m, 41°31’N 42°53’E, 18.viii.2005, leg. H. Sevgili (one of these females collected by S.S. Caglar; HUZOM); Ardahan: 2♂, Yalnizcam, Bilbilan Yaylasi, 2400 m, 20.viii.1974, 1♀, 15.viii.1974, 1♂, 1♀, the same locality, 2500 m, 15.ix.1973, leg. S. Salman (HUZOM) (area on the border between Artvin and Ardahan).

Previous records. Turkey: Artvin: type locality (Adelung, 1907); Badzgeret (Bey-Bienko, 1954); Gurdzhan (= Gurjian (?) in Karabag, 1958), type locality of P. ramrei (Miram, 1938); Kutul yaylasi, ca. 2500 m a.s.l. (Karabag, 1958) (1♂ in ZMHU, SysTax); Kutul geçidi (Salman, 1978); Ardahan: Bilbilan Yaylasi (Salman, 1978).

Bioacoustics. The calling song consists of pairs of syllables separated by intervals of several seconds (Fig. 2; mean/median 10/8.5 s, n = 25 intervals/12 males, T = 29°C). Each pair has a relatively complicated structure in time and frequency domain. After the first opening movement which is mostly but not always silent, the male closes the tegmina slowly at the beginning and quickly at the end. During the slow phase a series of faint impulses is produced (Fig. 7; due to the very low amplitude compared to the loud parts of the song they are difficult to recognise in Fig. 3). During the fast closing phase the sound becomes much louder and the previously separated impulses fuse into one or a few irregular pulses. At the beginning of the opening movement of the second syllable two to three, rarely one short, but quite loud pulses are produced (Fig. 3). The remaining opening movement is silent, but immediately at the beginning of the closing movement a long series of impulses can be heard (mean 84 impulses, range 50–120, n = 49 syllables/17 males). This series is again produced during a slow closing movement, which is followed by a fast phase from which one or a few pulses result. After an interval of 40–80 ms (at T = 29°C) often, but not always one or two afterclicks are produced. In recordings taken at night mostly a longer series of afterclicks (2–5 afterclicks; n = 10 syllables, ≥2 males; T = 17°C) could be heard. The duration of the first closing hemisyllable was 136 ± 27 ms, that of the second 184 ± 15 ms and that of the whole second movement cycle 225 ± 14 ms (n = 31 syllables, 1–3 from 12 males; T = 29°C). At lower temperatures the duration of all parts can, of course, be much longer (Fig. 7).

In the analysis of the frequency composition (Fig. 8; for the position of the spectra within the song see Fig. 7) it can be seen that during the impulse series of the first and second syllable audible components dominate, mainly over a quite narrow frequency band of between 5–10 kHz (mean of frequency maximum: 6.7 ± 0.8 kHz, 10 dB below maximum: 5.6–8.1 kHz; n = 18 males). During the fast closing movements and in the opening hemisyllable, however, much louder, broadbanded ultrasound components were registered together with low frequency components. Due to these ultrasound components, the amplitude of these parts is strongly underestimated in recordings made using audio tape recorders (compare Fig.

Fig. 7. Oscillograms of the calling song of P. tschorochensis from different localities and at different temperatures; A, B, E – Yalnizcam pass; C – Bilbilan yaylasi; D – Kutul yaylasi. Frequency range of recording A: 1–100 kHz, B–E: 1–20 kHz. Horizontal marks on recording A indicate the position of the spectra presented in Fig. 8.
3 and the different recordings in Fig. 7). In the afterclick mainly high frequency components were observed.

The occurrence of the different frequencies may be explained by the assumption of at least two resonators within the tegmina, which are activated differently. This hypothesis is supported by an examination of the stridulatory file, which is made up of two parts.

The stridulatory file of *P. tschorochensis* has a quite unusual shape, not observed in any other tettigoniid (Figs 9, 10). The distal, about 2.5 mm long and straight part of the file bears about 110 small stridulatory teeth (n = 1; several other tegmina similar). While closing the tegmina the scraper, which is situated on the edge of the right tegmen, has to run at first over this part of the file. Then the file bends sharply at a 50 degree angle towards the articulation of the wing. This proximal, again straight part bears 36 large teeth and is used for sound production just before closing the wings completely (Fig. 10). From the file structure and stridulatory movement it seems possible that during the first part of the closing hemisyllable the outer, heavily enlarged parts of the tegmen, which are bordered by the distal part of the file, vibrate. This accords well with the low frequency recorded. During the final fast closing movement more proximal and smaller areas of the tegmina may vibrate producing higher frequencies. The higher sound amplitude of this part of the syllable might be explained by the increased muscle power near the articulation. The pulses of the opening hemisyllable and the faint afterclicks may also be produced by stridulation using this part of the file.

The female tegmina overlap dorsally and the right one bears many stridulatory pegs on its dorsal surface as is typical for acoustically responding phaneropterids. The prothoracic spiracles of males and females are as large as those, for example, in *P. laevissimus* (Stumpner & Heller, 1992).

**Remarks on behaviour and ecology**

During our short visits we did not notice acoustical duetting between males and females. Accidentally, however, we did observe parts of the courtship behaviour. After physical contact between male and female, the male elevated its huge tegmina and offered the female access to his dorsal glands (Fig. 1C). Initially the female was very interested in the glands (Fig. 1D), however, after some time she lost interest and left without mating. Dorsal glands of very different sizes are probably present in most or all species of *Poecilimon* and related genera, like *Isopha* (Engelhardt, 1915). In most species they are exposed and easily accessible for a female after she has mounted the male. Elevation of the tegmina was not observed in any short winged phaneropterid species, but is reported for long-winged species of that family (e.g. *Scudderia furcata*; own observations KGH) and tree crickets (e.g., Brown, 1999). Remarkably, in the females
the first tergites are pale like the huge tegmina of the males (Fig. 1C). Some females, however, were completely green (except for reddish brown markings on the pronotum).

*P. tschorochensis* was found together with the phaneropterids *Poecilimon similis richteri* Ramme, 1933, *Iso- phyia zernovi* Miram, 1938 and *Isophya schneideri* Brunner von Wattenwyl, 1878 and the tettigoniids *Psorodonotus specularis* (Fischer-Waldheim, 1846) and *Parapholidoptera distincta* (Uvarov, 1921). In Bilbilan, Yalnizcam Mts, they were mainly found on *Urtica* sp.

**DISCUSSION**

**Phylogeny**

The assumption of a close relationship between the four species mentioned above is based mainly on the unusual shape of the pronotum. It is also supported by their circum-Caucasian distribution. Within the group, *Poecilimon tschorochensis* does not show any synapomorphies with the other species and has to be considered as its most basal member. The other three species possess cerci, which have at least two teeth at their tip (Fig. 5), whereas *P. tschorochensis* shows the ancestral cercal shape, ending in one simple tooth (Fig. 6). Among these three species, the parapatric *P. heroicus* and *P. bifenestratus* share a complex of characters, all associated with non acoustical females. The tegmina of the females are small and do not overlap dorsally, and the auditory spiracles are strongly reduced in size. The systematic relationships can thus described as (*P. tschorochensis* (*P. tricuspis* (*P. heroicus*, *P. bifenestratus*))).

The origin of the group may therefore be north eastern Turkey (Caucasian part), from where it spread northwards to the Caucasus. Probably *P. heroicus* occupied the steppes of the foothills in the north of the Caucasus after the last ice age.

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**Bioacoustics**

The most unusual character of the song of the species studied is its spectral composition, especially that of *P. tschorochensis*, which has the lowest peak frequency of all *Poecilimon* species, when the maxima of sound spectra are compared (Fig. 12). In the genus *Poecilimon*, the frequency maximum is negatively correlated with body size, but *P. tschorochensis* has neither a wider head or longer hind femora than other *Poecilimon* species. Regarding the huge size of its tegmina, however, the low frequency of song is not unexpected. *P. heroicus* and *P. bifenestratus* also have very low peak frequencies (Fig. 12). Both species are distinctly larger than *P. tschorochensis* but the deviation in the frequency expected due to their size is much smaller. However, the results indicate that a low frequency is a basal feature of the group, probably linked with the posterior widening of the pronotum and the enlargement of tegmina.

There are two advantages of low frequencies. First, sound signals with low frequencies have generally a larger range than those with high frequencies, a well known phenomenon in bioacoustics (for tettigoniids see Zhantiev, 1981; Römer & Lewald, 1992). High frequencies, for example, are strongly attenuated in dense vegetation. Properties of the habitat of these *Poecilimon* species possibly favour the use of long sound wave lengths (low frequencies). This idea is supported by finding in the background of some field recordings of *P. tschorochensis* the song of *Psorodonotus specularis*, which has a similar spectral composition. *Psorodonotus* are large bush-crickets typically found near to the ground in moist habitats. *P. bifenestratus* sings on the ground or not very high up in vegetation, *P. heroicus* and *P. tricuspis* occur in grassland with great plant diversity, but typically not very
tall plants. However, *P. tschorochensis* is found higher up in vegetation than *Psorodonotus* and appears at least not to be strictly bound to moist conditions (own observations).

A second reason for the use of low frequencies may be sexual selection. In many studies females have been shown to prefer larger than average males. If females judge size on the basis of properties of acoustical signals, they should prefer signals of lower than average frequency, because these might indicate larger males. Although the results from the few experiments addressing this question are far from clear (see Gerhardt & Huber, 2002), the hypothesis may hold in some cases. The huge size of the tegmina in *P. tschorochensis* may be a consequence of a runaway process—a kind of acoustical peacock tail (Andersson, 1994).

Another interesting character change is the loss of an acoustical response by females within the *P. heroicus*-group. The combination of characters related to this loss is known also in *Poecilimon* species of other groups (e.g., *P. propinquus*-group (Lehmann, 1998), *P. ampliatus*-group (Heller & Lehmann, 2004), which also have mute females. The *P. heroicus*-group, however, is the first group, where within a group this change in acoustic communication system can be traced. In both the above mentioned groups the females of all species have strongly reduced tegmina, too small to produce sounds. However, the systematic relations of the groups to each other as well as to acoustical responding species are unknown.

In all species similar changes are observed in the acoustical behaviour and hearing organs. *P. tschorochensis* with acoustical females produces schemes (combination of two syllables) at intervals of about 10 s, whereas *P. heroicus* and *P. bifenestratus* produce long sequences of syllables at intervals of only about one second. Intervals of about the same difference in duration were recorded for *Poecilimon* species with acoustically responding and mute females, respectively (Heller & Helversen, 1993), and this corresponds to the different functions of the song. In *P. tschorochensis* and similar species, the male song has to elicit an acoustical response from females, which is used by the male for his approach, whereas in *P. heroicus* and other species with mute females the female has to walk to the male guided by his song. *P. heroicus* and *P. bifenestratus* also show a reduction in the acoustic spiracles as observed in other *Poecilimon* species with mute females (Stumpner & Heller, 1992). In *P. heroicus* and *P. bifenestratus* this change is more easily understood than in the smaller species, because the reduction in size of the prothoracic spiracle does not significantly decrease the auditory sensitivity at low frequencies (below about 10 kHz) (Stumpner & Heller, 1992). In terms of the frequency of the male song, a reduction in the size of the spiracles might also be expected in *P. tschorochensis*. However, the female response includes ultrasonic components judging by the size of her tegmina (similar to the situation in *Ancistrura nigrovittata* (Dobler et al., 1994; Stumpner, 2002). Decoupling the size of male’s and female’s spiracles is less likely to occur during evolution than equally sized spiracles in both sexes.

For a deeper understanding of the evolution of this communication system more data on the response of females and the acoustic behaviour of *P. tricuspidis* are urgently needed. A detailed study of all species of the group may uncover the reasons for the existence of two different acoustic communication systems in bush-crickets.

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Databanks

SysTax (a Database System for Systematics and Taxonomy): http://www.biolgie.uni-ulm.de/systax/daten/index.html


FaEu (Fauna Europaea) http://www.faunaeur.org

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