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Panoploscelis scudderi Beier, 1950 and
Gnathoclita vorax (Stoll, 1813):
two katydids with unusual acoustic,
reproductive and defense behaviors
(Orthoptera, Pseudophyllinae)

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COUVERTURE / *COVER*:

View of the Massif of Mitaraka in French Guiana. In médaillon, disturbed male *Gnathoclitia vorax* (Stoll, 1813) escaping from the shelter. Photographs: Sylvain Hugel.

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***Panoploscelis scudderi* Beier, 1950 and *Gnathoclitia vorax* (Stoll, 1813): two katydids with unusual acoustic, reproductive and defense behaviors (Orthoptera, Pseudophyllinae)**

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KEY WORDS

Insecta,
Ensifera,
Pseudophyllinae,
mate guarding,
acoustic communication,
French Guiana,
new synonym.

ABSTRACT

Two species of Eucocconotini Beier, 1960 were collected during the “Our Planet Revisited, Mitaraka 2015” survey in the Mitaraka Mountains belonging to Tumuc-Humac mountain chain in French Guiana: *Gnathoclitia vorax* (Stoll, 1813) and *Panoploscelis scudderi* Beier, 1950. Calling songs of both species are described for the first time, as well as the mandibular and tegminal protest signals from *P. scudderi* males and females. The structures involved in these signals are described and illustrated. The peculiar acoustic and mate guarding behaviors of *Gnathoclitia vorax* are described and illustrated. The synonymy of *Panoploscelis scudderi* Beier, 1950 and *Panoploscelis angusticauda* Beier 1950 n. syn. is discussed and proposed, based on specimens reared from samples collected in Mitaraka.

RÉSUMÉ

Panoploscelis scudderi Beier, 1950 et *Gnathoclitia vorax* (Stoll, 1813): deux sauterelles aux comportements acoustiques, reproducteurs et de défenses remarquables (Orthoptera, Pseudophyllinae).

Deux espèces de Eucocconotini Beier, 1960 ont été collectées au cours de l’expédition “La planète revisitée, Mitaraka 2015” dans le massif du Mitaraka appartenant à la région des Tumuc-Humac en Guyane: *Gnathoclitia vorax* (Stoll, 1813) et *Panoploscelis scudderi* Beier, 1950. Les chants d’appel des deux espèces sont décrits pour la première fois, ainsi que les signaux de protestation que mâles et femelles de *P. scudderi* produisent avec leurs élytres et leurs mandibules. Les structures impliquées dans ces signaux sont décrites et illustrées. Les comportements acoustiques et de garde très particuliers de *Gnathoclitia vorax* sont décrits et illustrés. La synonymie de *Panoploscelis scudderi* Beier, 1950 et *Panoploscelis angusticauda* Beier 1950 n. syn. est discutée et proposée en nous appuyant sur les spécimens élevés à partir des collectes au Mitaraka.

MOTS CLÉS

Insecte,
Ensifère,
Pseudophyllinae,
comportement de garde,
communication acoustique,
Guyane,
synonyme nouveau.

INTRODUCTION

Nine genera of Eucocconotini Beier, 1960 are currently known, all restricted to the northern half of South America (Cigliano *et al.* 2018). Whereas Eucocconotini is among the smallest Pseudophyllinae tribe, it is one of the most morphologically diverse, as pointed by Braun (2011). Most of Eucocconotini display large leg spines and/or sexually dimorphic characters on mouthparts. Specimens belonging to the two Eucocconotini genera *Gnathoclita* Haan, 1843 and *Panoploscelis* Scudder, 1869 have been collected during the “Our Planet Revisited, Mitaraka 2015” survey in the Mitaraka Mountains. These mountains are localized in the southwestern-most corner of French Guiana (Touroult *et al.* 2018). With three weeks on site, such relatively long survey not only aims at performing large scale sampling, it also allows spending time for biological observations of the species in their environment. Among the numerous observations, video and sound records of Orthoptera made during this survey, the ones related to the two species of Eucocconotini observed in Mitaraka are of peculiar interest, and are the object of the present article.

MATERIAL AND METHODS

MATERIAL EXAMINED

Samples collected

The material used in this work was collected during the “Our Planet Revisited, Mitaraka 2015” survey in the Mitaraka Mountains (22.II.2015 to 11.III.2015). This large-scale biodiversity survey was co-organized by the Muséum national d’Histoire naturelle (MNHN) and the NGO Pro-Natura international (Touroult *et al.* 2018). Mitaraka Mountains are a little known area sitting in the southwestern-most corner of French Guiana bordering Surinam and Brazil. Mitaraka belongs to the Tumuc-Humac mountain chain, extending east in Amapa region and west in southern Surinam. Most of the area is covered by a tropical lowland rain forest from which few inselbergs and hills emerge. The biological sampling benefited from the access and benefit sharing agreement “APA973-1”, which is reported in each sample label. All specimens collected during the survey are deposited in MNHN collections.

Museum specimens

High quality pictures of *Panoploscelis scudderi* Beier, 1950 and *Panoploscelis angusticauda* Beier 1950 n. syn. holotypes were kindly provided by Dr Harald Bruckner, Naturhistorisches Museum, Wien, Austria. High quality pictures of *Panoploscelis scudderi* allotype were kindly provided by Anna Liana and Przemysław Szymroszczyk, Museum and Institute of Zoology, Polish Academy of Sciences, Warsaw, Poland.

SAMPLING METHODS

All *Panoploscelis* Scudder, 1869 and *Gnathoclita* De Haan, 1842 specimens were collected during night hours, either by sight with a headlamp or thanks to their conspicuous acoustic behavior. All specimens of these species have been collected

by the author; both species were absent from the numerous traps used by the survey team.

ACOUSTIC ANALYSES AND TERMINOLOGY

Sound recordings were either directly performed on the field, or in studio-like conditions. Sounds were acquired with an Audiotechnica AT822 stereo microphone, on a NIKON D5200 camera (sampling rate: 96 kHz) with a red illumination (to follow the insect behaviour). Song analysis has been performed with Clampfit 10.2 software. Song recordings are deposited in both coll. S.H. and MNHN acoustic databases: <https://sonotheque.mnhn.fr/>. Acoustic terminology is from Stumpner *et al.* (2013).

SUPPLEMENTARY VIDEOS

Supplementary video 1: Panoploscelis scudderi mandibular protest signal of an adult female.

Both males and females respond to air puff, water spraying or simple touch by lifting their forelegs and producing repeated protest signals. Under gentle stimulation, *P. scudderi* respond mostly by elevating their forelegs and by producing mandibular protest signals. Such protest signals involves peculiar structures described in the article. http://museumedia.mnhn.fr/index.php?urlaction=doc&cid_doc=109649&rang=3

Supplementary video 2: Gnathoclita vorax adult male singing in natura (same as in figure 8D).

Adult males of *Gnathoclita vorax* sing by night hours, from the undergrowth from plant-made cavities. The call consists of long (> 1 min) and relatively regular repetitions of phonotomes. http://museumedia.mnhn.fr/index.php?urlaction=doc&cid_doc=109650&rang=2

Supplementary video 3: Courtship behavior of males of Panoploscelis scudderi.

Two males and two virgin females are transferred in the same box (30 min before the sequence). Note that neither males nor females call during the mating sequence. http://museumedia.mnhn.fr/index.php?urlaction=doc&cid_doc=109651&rang=1

ABBREVIATIONS

Institution

MNHN Muséum national d’Histoire naturelle, Paris.

Private collection

coll. SH collection of the author, Strasbourg.

Acoustics

Hz Herz;
kHz kilohertz;
min minimal duration;
max maximal duration.

Measurements

H height;
L length;
W width.

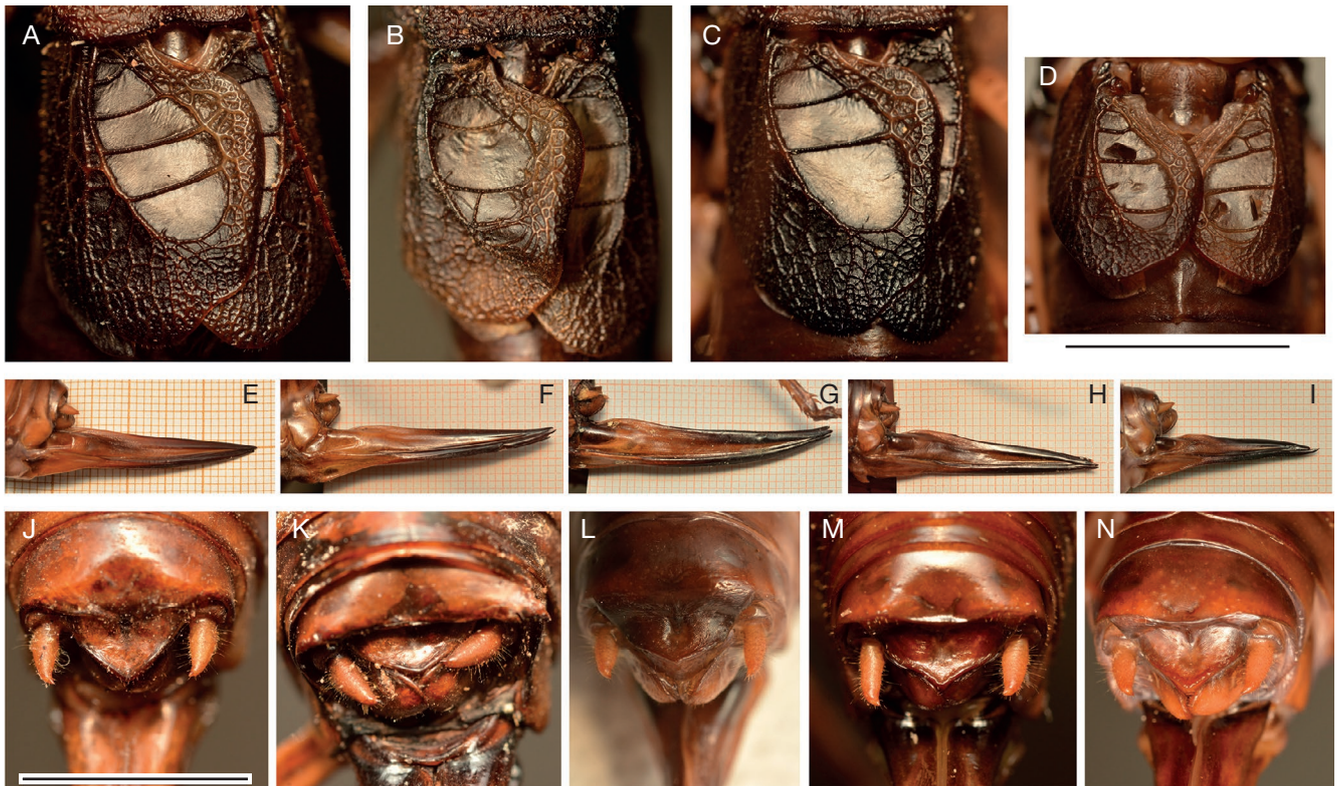


FIG. 1. — Variations in characters used to distinguish *Panoploscelis scudderi* Beier, 1950 and *Panoploscelis angusticauda* Beier 1950 n. syn. females. The specimens pictured are bred from samples collected in Mitaraka (F1 and F2 generations). The frequent-most condition is represented in the left panels (A, E, J). Specimens from Mitaraka display variations in: A–D, the number of tubercle-bearing veins; E–I, the shape of ovipositor in side view (holotype of *P. angusticauda* display the same shape of ovipositor as illustrated in F); J–N, the shape of epiproct hind margin (male juvenile holotype of *P. scudderi* display the same shape of epiproct hind margin as illustrated in N; in *P. angusticauda*, the hind margin is somehow as in K). Scale bars: 10 mm.

RESULTS

Order ORTHOPTERA Latreille, 1793
Family TETTIGONIIDAE Krauss, 1902

Genus *Panoploscelis* Scudder, 1869

PANOPLOSCELIS SCUDDERI BEIER, 1950 AND *PANOPLOSCELIS ANGSTICAUDA* BEIER, 1950 N. SYN.

Both species were described in the same article by Max Beier (1950). *Panoploscelis scudderi* was described from North Brazil after a single subadult male and *P. angusticauda* from Northwestern Brazil after a single adult female. Since holotypes of both species are from different genders and one is immature, only non-sexual characters present in both adults and immatures might help to distinguish *P. scudderi* and *P. angusticauda*, should these really be different. We examined unpublished high quality pictures of both holotype specimens, and the only character that may be used to distinguish them as separate species is the shape of the epiproct: more rounded in *P. angusticauda* holotype and angulose in *P. scudderi* holotype.

Our colony of *Panoploscelis* from Mitaraka displays quite variable shape of epiproct in both males and females (Fig. 1J–N), making impossible to rely on this character to conclude

whether *P. scudderi* and *P. angusticauda* holotypes really belong to different species.

With the currently available characters and since characters previously used to distinguish the species are quite variable in our colony reared from Mitaraka samples, we found no support that *P. angusticauda* and *P. scudderi* are different species and consider at this stage that *Panoploscelis scudderi* Beier, 1950 and *Panoploscelis angusticauda* Beier, 1950 are synonyms.

Interestingly, one additional adult female was included in the original description of *P. scudderi* as female allotype. This female is mentioned from “Stettin” collections by Max Beier (1950) and is now in Warsaw/Łomna. This female is from a distinct locality (British Guiana). Beier probably considered this female as conspecific with the male holotype of *P. scudderi* since both display an angulose epiproct. Although *P. scudderi* holotype cannot be recognized as different from *P. angusticauda* holotype, that female might belong to another yet unnamed species. The epiproct of this female falls within the range of variation of our Mitaraka specimens (similar to Fig. 1J) and does not allow to recognize another species. As pointed by Beier (1950, 1960), this female allotype has a wide ovipositor, 5 times longer than wide, lanceolate, whereas *P. angusticauda* holotype has more slender ovipositor, 6.5 times longer than wide, with regularly

TABLE 1. — Measurements of males of *Panoploscelis scudderi* Beier, 1950 (in mm). Abbreviations: L, length, W, width.

Body	Pronotum	Left forewing	Fore tibia	Mid tibia	Hind tibia	Fore femur	Mid femur	Hind femur	Hind femur
L	L	L	L	L	L	L	L	L	W
50-66	13.5-15.5	11-14	20.5-23.5	20-21.5	29-35.3	18.5-23	18-20.5	28-32.5	5-5.5

converging margins. Females in our colony of *Panoploscelis* from Mitaraka have very variable ovipositor shape and length (Fig. 1B), and ovipositor of the female considered by Beier as *P. scudderi* allotype is only slightly less slender than extreme specimens in our colony, raising doubts on reliability of this character. Interestingly, the left forewing of this specimen has 5-6 transverse tubercle-bearing veins on the mirror whereas *P. angusticauda* female holotype has 3-4 tubercle-bearing veins on the mirror. In addition, the shape of the mirror of that female considered by Beier as *P. scudderi* allotype is very unusual, much narrow distally, and tubercle bearing veins as well as large cells occur outside of the mirror area and might suggest an abnormality. Our colony of *Panoploscelis* from Mitaraka typically displays 3-4 tubercle-bearing veins on the mirror (Fig. 1A, Fig. 3A), but a significant proportion displays variation (2-5 veins, Fig. 1B-D) indicating that this character might not be very stable and/or subject to abnormalities. At this stage, it is difficult to define whether this specimen corresponds to *Panoploscelis scudderi* with abnormal wings or to another yet unnamed species.

Panoploscelis scudderi Beier, 1950.

Panoploscelis scudderi Beier, 1950: 112.

Panoploscelis angusticauda Beier, 1950: 113, n. syn.

TYPE MATERIAL. — “Coll. Br. V. W. [Printed], Rio Branco, Peru [manuscript; in reality Brazil], Staudinger [manuscript]; 24.875 [manuscript]; Type [printed, red label], *Panoploscelis armata* Scudd. [manuscript] determ. Karny [printed]; *Panoploscelis scudderi* n. sp. Type ♂ [manuscript]; NHM, Wien, Austria, high quality pictures examined. — Type [printed, red label]; Natt: 12; Rio-Negro [manuscript]; *Panoploscelis armata* Scudd. [manuscript]; *Panoploscelis angusticauda* n. sp. Type ! ♀ [manuscript] det. Beier [printed], NHM, Wien, Austria, high quality pictures examined.”

TYPE LOCALITY. — Rio Branco, Brasil.

OTHER MATERIAL EXAMANED. — **French Guiana.** Planète revisitée Guyane 2015, Monts Tumuc-Humac, Massif du Mitaraka, Layon D, 54.4509°O, 2.2357°N, 280 m a.s.l. – 54.4517°O, 2.2338°N 293 m a.s.l., 23.II-10.III.2015, nuit, 1 ♂, 1 ♀, MNHN. — Planète revisitée Guyane 2015, Monts Tumuc-Humac, Massif du Mitaraka, vers sommet en Cloche, 54.4541°O, 2.2349° N, 370 m – 54.4646°O, 2.2329°N 470 m a.s.l., 23.II-10.III.2015, nuit, 1 ♀, MNHN. These specimens were reared, and F1-F2 generations were also examined for the present work to assess the variability of diagnostic characters.

DISTRIBUTION. — South America: North Brazil, Guiana, French Guiana.

DESCRIPTION

Male

Head. Fig. 2A, B. All as in female. Fastigium verticis spine-shaped, reaching antennal sockets. Fastigium frontis spine-shaped. Carina lateralis interna distinct at the basis, carina lateralis externa complete. Scapus with one distal inner spine and one distal dorsal tubercle.

Thorax. Fig. 2B. Anterior margin with a distinct median tubercle; in addition to the two transverse sulci, sagittal sulci at least distinct at the level of posterior transverse sulcus (as in female); posterior margin broadly rounded with an inconspicuous median emargination of the rim. Metazona elevated.

Legs. All as in female, anterior tibiae and femora longer than mid tibiae and femora. Fore femur (Fig 2C): distal spine of inner ventral margin perpendicular to femur axis; dorsally with 4-5 spines. Posterior femora dorsally with 5-7 spines (Fig. 2D); mid femora unarmed dorsally. Inner genicular lobe of fore femora unarmed. Wings. Left forewing shorter than right forewing (this very unusual asymmetry is possibly a generic character). Left mirror area triangular, right mirror somewhat oval. Anal field posterior margin of left forewing with a concavity; posterior margin of both forewing with an emargination distal to the anal field. File with c. 66 teeth. Abdomen. Suranal distal margin variable, usually with an angle (as in female), sometimes very obtuse. Cercus short, stout, with a single inner ventral spine-shaped projection.

Subgenital plate broad at basis, moderately tapering distally; with a more or less rounded V-shaped emargination between the styli.

Measurements. See Table 1.

Female

Female has been described under *Panoploscelis scudderi* and *Panoploscelis angusticauda* in Beier (1950, 1960). Variation in characters are illustrated Fig. 1. Forewings of a female specimen from Mitaraka are illustrated in Fig. 3A, B. Female forewings of *P. scudderi* lack the scraper lobe pointed by Montealegre-Z *et al.* (2003) in females of *Panoploscelis specularis* Beier, 1950.

BIOACOUSTICS

Male call (Fig. 4).

In rearing conditions, with many males and females together, the call of *Panoploscelis scudderi* males is very rarely heard. Single males sing only by night hours. At 24 °C, the song is made of 3-12 verses repeated every 31.7 ± 3.1 s (min: 18.44 s; max: 43.60 s). The first verse is usually softer than the subsequent ones. The duration of one verse is 7.5±0.6 s (min: 4.4 s; max:



FIG. 2. — Male of *Panoploscelis scudderi* Beier, 1950, after a specimen collected in Mitaraka: **A**, face in front view; **B**, Head and pronotum in dorsal view; **C**, right fore femur in front view. Note that the distal most ventral anterior spine is on the axis of the picture; **D**, hind femur in side view; **E**, apex of the abdomen in dorsal view; **F**, cercus in dorsal view; **G**, subgenital plate in ventral view. Scale bars: A-E, G, 10 mm; F, 1 mm.

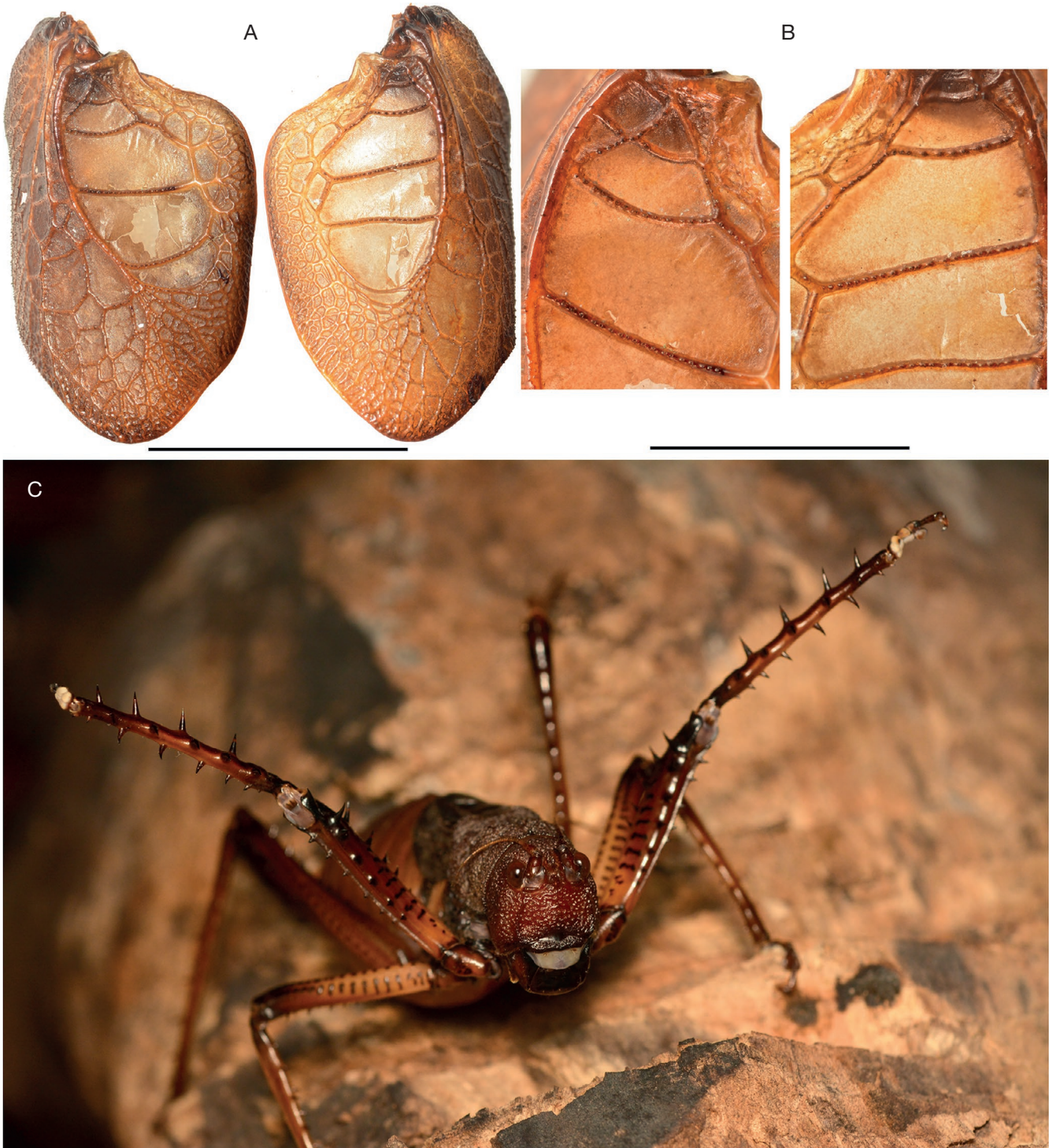


FIG. 3. — Female of *Panoploscelis scudderi* Beier, 1950, after a specimen collected in Mitaraka: **A**, forewings in dorsal view; **B**, detail of forewings showing tubercles in dorsal view; **C**, defensive position. Scale bars: A, 10 mm; B, 5 m.

10.0 s). Verses are made of 29.4 ± 1.9 phonatomes (min: 18; max: 34) repeated every 251 ± 2 ms (min: 233 ms; max 271 ms). The duration of one phonatome is 60 ± 2 ms (min: 30 ms; max: 87 ms). At 24°C, a first peak was recorded at $c.$ 10.3–11.0 kHz, and a secondary broad peak was recorded at 30.5 kHz.

Isolated males seldom produce isolated verses (1–3 per nights).

Female call

In our initial rearing conditions (30 adult pairs in a 0.3 m³ cage at 24°C by night), we failed to record any call from females (at F3 generation, a total of 50 adult females have been reared). Since males only rarely sing in the same conditions, this lack of spontaneous calls might simply be linked to specimen density.

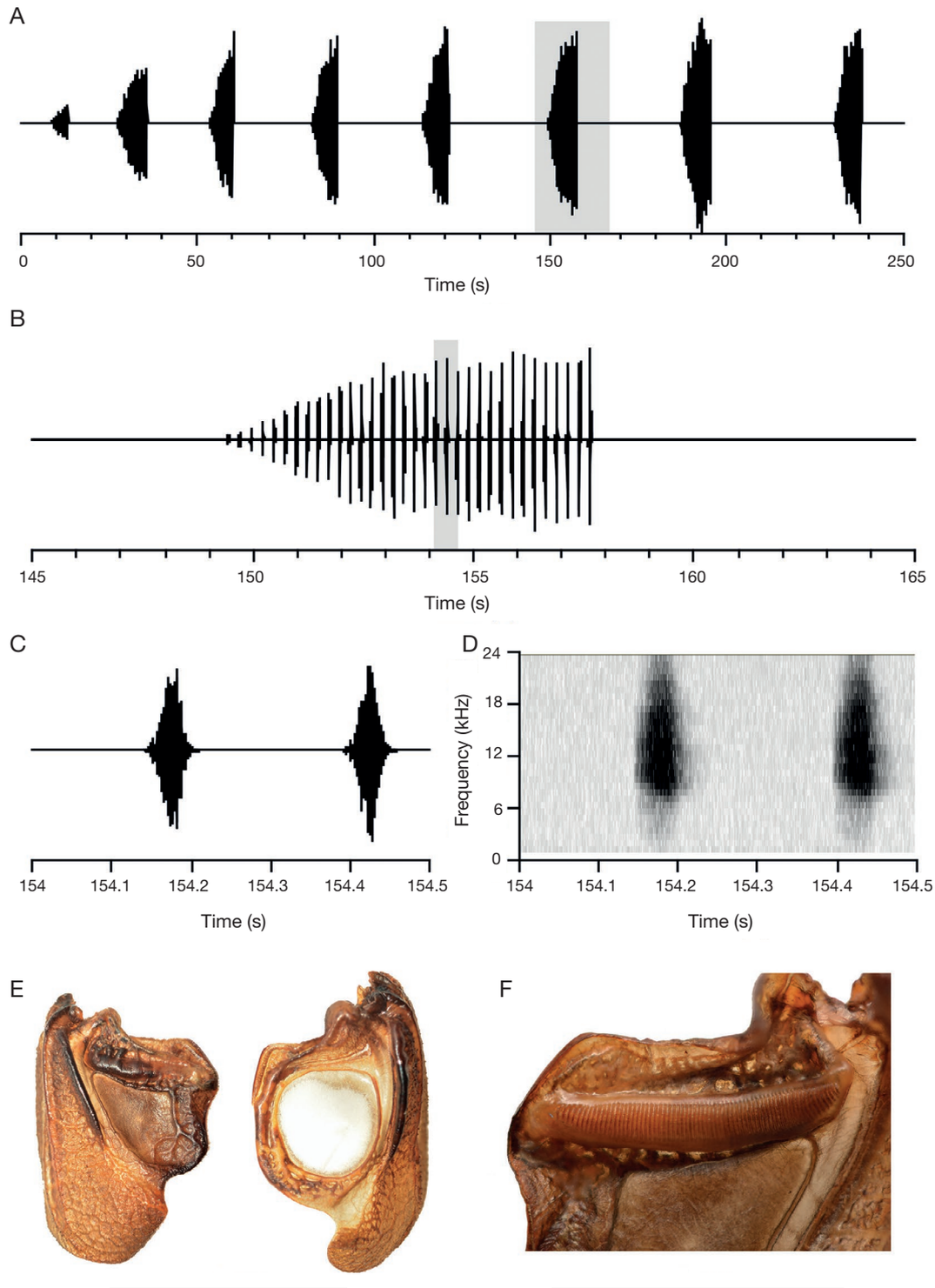


FIG. 4. — Male of *Panoploscelis scudderii* Beier, 1950, call and stridulatory apparatus. **A-D**, call of a wild male in studio conditions at 24°C after a specimen collected in Mitaraka; **E**, male forewings in dorsal view; **F**, file in ventral view. Scale bars: 10 mm.

Therefore at F4 generation, subadult males and females were separated and kept in isolation for two months after their last molting. Under these conditions, males were calling very frequently (at least one call every 10 min for 25 males together in a cage). We were nevertheless unable to record any song

from females, even with overnight recordings (8 full night recordings). Since females may only respond to male call, we play-backed male calls (one call every 10 min, during the whole night): no responses were recorded (two overnight recordings). We then checked whether females responded to males during

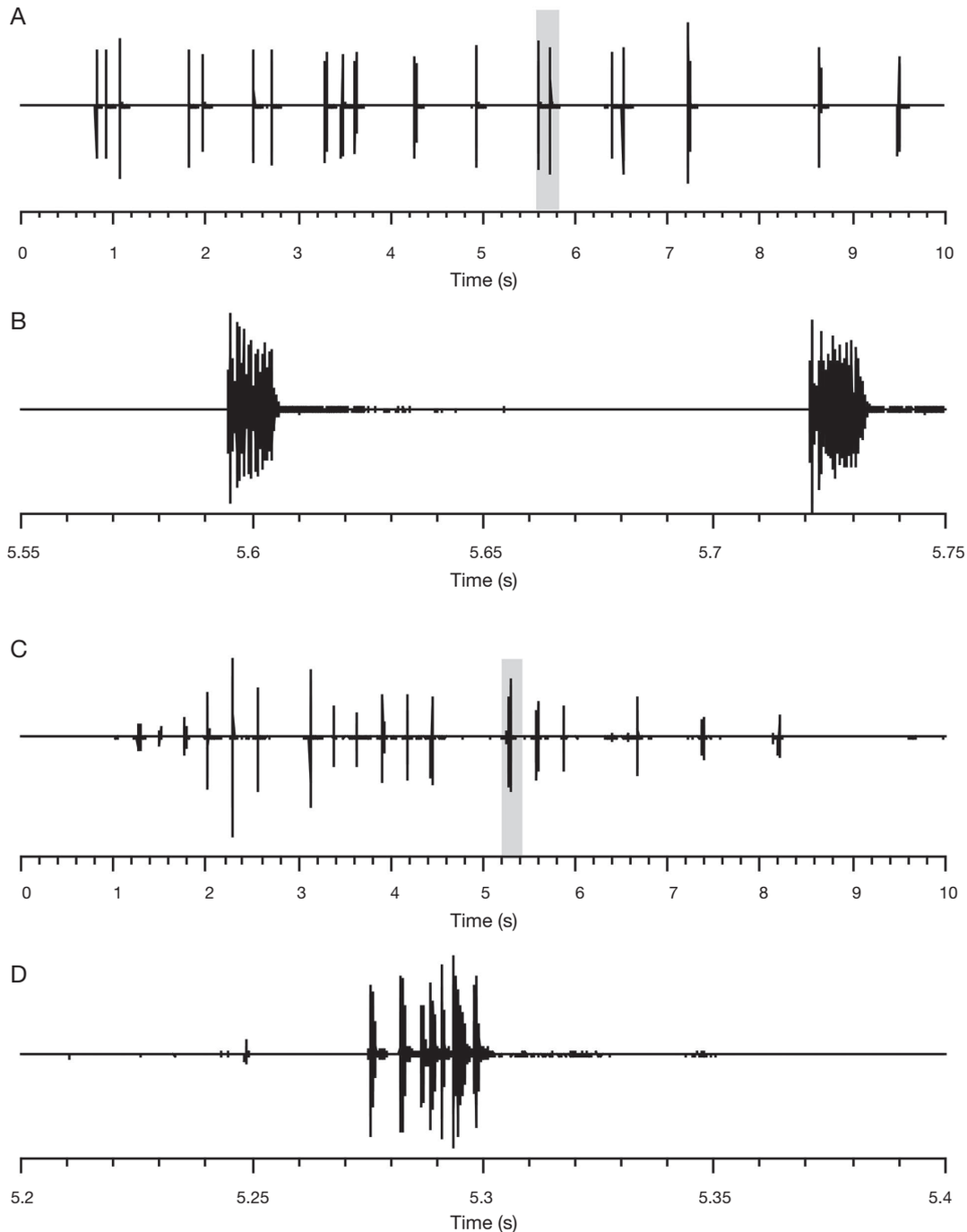


FIG. 5. — Tegminal protest signals of *Panoploscelis scudderi* Beier, 1950: **A, B**, male; **C, D**, female. All from wild specimens in studio conditions at 24° C.

the mating process: we put virgin females together with males in one single empty cage and imaged/recorded them during the whole night. Under these conditions, all females mated within the first 30 minutes, and neither males nor females produced any song (Supplementary video 3).

These data suggest that call of *P. scudderi* females is not required in the reproductive behavior of that species. It cannot be ruled out that *P. scudderi* females do not call at all since the only tegminal sounds we recorded were produced as protest signals.

Interestingly, *P. scudderi* females lack the scraper lobe pointed by Montealegre-Z *et al.* (2003) in *P. specularis* (Fig. 3A). This might speculatively suggest that intraspecific female signaling might be less developed in *P. scudderi*.

Male and female protest signals

Both males and females respond to air puff, water spraying or simple touch by lifting their forelegs and repeated protest signals (Fig. 3C). These protest signals are produced by tegminal stridulatory organs as well as mandibles/labrum.

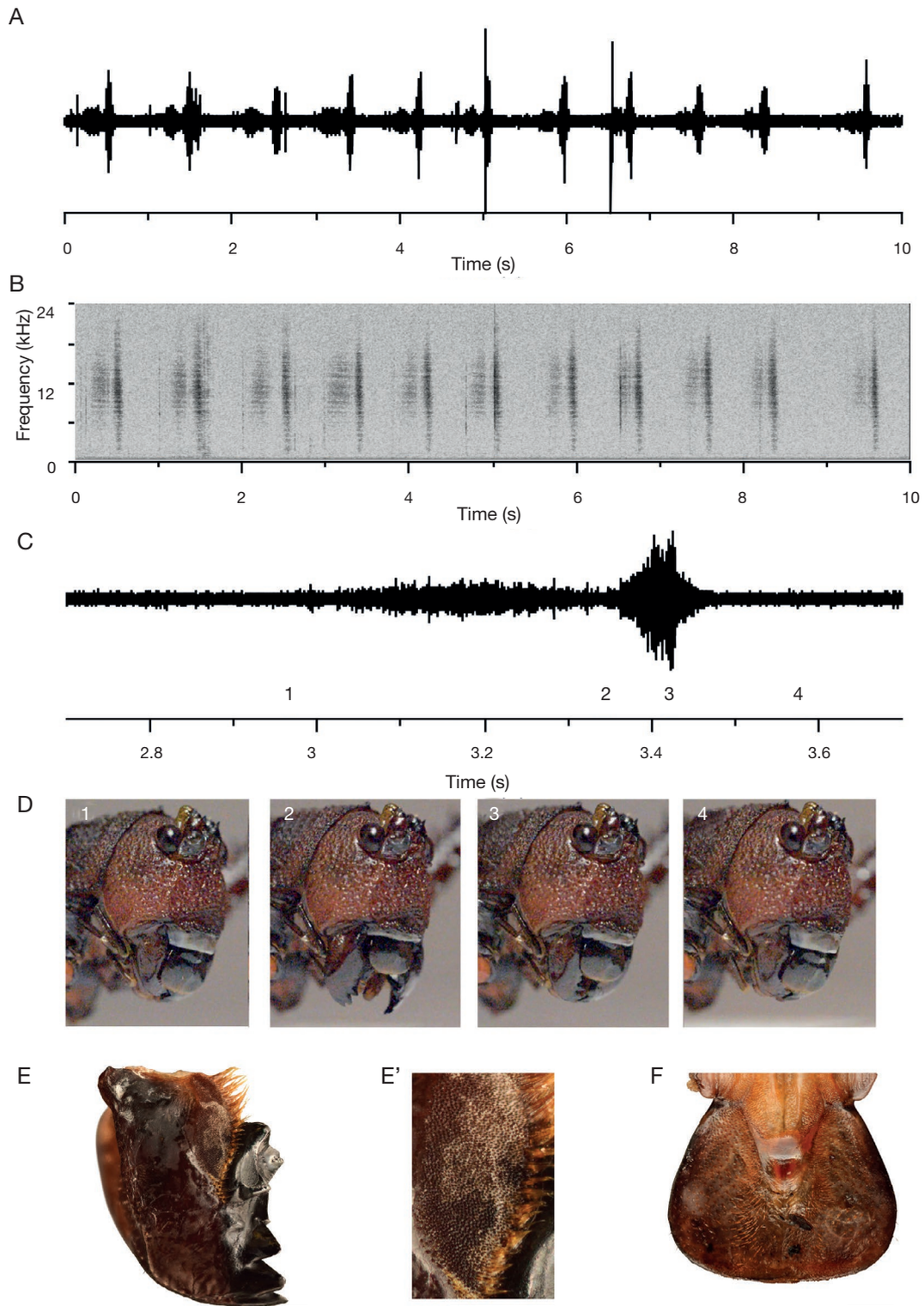


FIG. 6. — Mandibular protest signals of *Panoploscelis scudderi* Beier, 1950: **A-C**, mandibular protest signals produced by a F2 adult female; **D**, pictures corresponding to the time points given in **C** illustrating the motion of mandibles, clypeus and labrum; **E**, right mandible (note the area with tubercles in **E'**; **F**, clypeus and labrum, inner view. Scale bars: 1 mm.

At 24 °C, male tegminal protest signal (Fig. 5AB) is made of phonotomes isolated or grouped (2-3) repeated every 420 ± 78 ms (min: 108 ms; max: 1335 ms). The frequency peaks at 9.4 kHz, a frequency significantly lower than male

call. At 24°C, female tegminal protest signal (Fig. 5CD) is made of verses repeated every 348 ± 56 ms (min: 246 ms; max: 836 ms). Verses are made of 5 ± 1 phonotomes (min: 2; max: 12) of irregular duration and spacing. The power spectrum

of female tegminal protest is broad and displays a double peak: one at 10 kHz and another at 14 kHz.

Male and female mandibular protest signal (Fig. 6, Supplementary video 1)

Mandibular protest signals are similar in males and females. These are the first protest signals produced when a specimen is stimulated, the tegminal signal requiring more stimulation. This signal has two phases: the first phase is produced while opening the mandibles and up lifting the labrum by clypeus retraction, the second is produced by the return to the resting state. The total duration of a cycle is very variable, lasting *c.* 0.5 s, with a first phase usually longer than the second one. These signals are repeated every 2.3 ± 0.2 s (min: 0.7; max: 3.6). The power spectrum of this signal displays a broad peak between 10.3 kHz and 14.6 kHz

Protest signals appear broader and emitted at lower frequencies than male call. This might speculatively be linked to the correlation between signal valence and signal frequency in intraspecific communication of some rodent species where lower frequencies are associated to unpleasant situations (Knutson *et al.* 2002).

Comparison of P. specularis and P. scudderii male call

Whereas the call of *P. specularis* males is made of two phonotomes separated by *c.* 130 ms and regularly repeated every *c.* 3 s, the song of *P. scudderii* is more complex in structure, with phonotomes organized in verses (see above). Alternatively, since *P. scudderii* sometimes produces isolated verses, one cannot rule out that *P. specularis* could do the same, and the full call of this species with organized verses may speculatively have been missed. The frequency of the maximally intense peak is relatively lower in *P. scudderii* than in *P. specularis* (10–11 kHz *vs.* 11–13 kHz).

Genus *Gnathoclita* De Haan, 1842

Gnathoclita vorax (Stoll, 1813)

Gryllus (Acheta) vorax Stoll, 1813: plate 4c, figs 19, 20.

Gryllus (Gnathoclita) vorax De Haan, 1843: 208.

TYPE MATERIAL. — Lost?

TYPE LOCALITY. — America

OTHER MATERIAL EXAMINED. — **French Guiana.** Planète revisitée Guyane 2015, Monts Tumuc-Humac, Massif du Mitaraka, 54.450137°O, 2.233883°N (GF DZ), 315 m a.s.l.; 23.II-10.III.2015, nuit, 1 ♂, MNHN. — Planète revisitée Guyane 2015, Monts Tumuc-Humac, Massif du Mitaraka, Layon D, 54.4509°O, 2.2357°N, 280 m a.s.l. – 54.4517°O, 2.2338°N, 293 m a.s.l., 23.II-10.III.2015, nuit, 1 ♂, 1 ♀, MNHN. — Planète revisitée Guyane 2015, Monts Tumuc-Humac, Massif du Mitaraka, vers sommet en Cloche, 54.4541°O, 2.2349°N, 370 m a.s.l., - 54.4646°O, 2.2329°N, 470 m a.s.l., 23.II-10.III.2015, nuit, 1 ♂, 1 ♀, MNHN.

DISTRIBUTION. — Surinam, Guiana, Brazil North [Cigliano *et al.*, OSF, 2018], French Guiana.

REDESCRIPTION

Males and females of this species have been comprehensively re-described (Willemsse 1954, Beier 1960, De Jong 1971).

Stridulatory apparatus (Fig. 7D, E)

Left FW mirror infuscated, with no concavity, oval shaped except the anterior proximal angle; about 2 times as high (maximal height) as wide (maximal width). File with *c.* 101 lamellar teeth (Fig. 7E).

BIOACOUSTICS (FIG. 7A-C, SUPPLEMENTARY VIDEO 2)

Gnathoclita vorax sings by night hours, from the undergrowth from plant-made cavities. The call consists of long (> 1 min) relatively regular repetitions of syllables. At 26°C, syllables are repeated every 3.818 ± 0.625 s (syllable duration: 50.8 ± 4.3 ms). The frequency peaks between 8750–8900 Hz.

Comparison of G. vorax and G. sodalis Brunner von Wattenwyl, 1895 male call

The call of *G. sodalis* is made of phonotomes repeated every 180–480 ms, depending on the temperature (Montealegre-Z & Morris 1999) whereas *G. vorax* phonotomes are repeated every *c.* 4 s. The call of *G. sodalis* peaks at about 15–16 kHz, about twice the frequency of *G. vorax* peak (Montealegre-Z & Morris 1999). Interestingly, in addition to acoustic communication, *G. sodalis* males use tremulatory vibration to communicate with other males and females (De Souza *et al.* 2011). This tremulatory behavior may not occur in *G. vorax* since males of this species sing in a very narrow environment (see below).

GUARDING BEHAVIOR OF *GNATHOCLITA VORAX*

A total of five males of *Gnathoclita vorax* were observed during the Mitaraka survey in 2015. All these observations occurred late by night hours (11 pm–2 am). These males were sheltering in narrow tube-shaped hollow plant sections, their head facing the aperture of the cavity (Fig. 7A-D). The diameter of these shelters was hardly larger than a male's head, preventing turn-overs (Figs 7D, 8). The shelter of all but one specimen was within hollow dead stems of *Astrocaryum* sp. These observations are in line with a previous report of *Gnathoclita vorax* singing from within *Gadua* sp. bamboo stems (Naskrecki 2008).

Three of the five males observed were located owing to their loud syllables repeated every 5 s (Fig. 7A-C). These males were stridulating alone within their shelter, their antennae protruding from the entrance.

The two other males were found owing to their conspicuous antennae protruding from the aperture. None were stridulating. In both cases, an adult female was localized at the bottom of the tube-shaped shelter, behind the male who blocked the entrance (Fig. 8A'). None of the females had a spermatophore attached.

These specimens were kept separated for one week, and subsequently placed altogether in a large cage containing wood-made tube-shaped shelters of similar measurements to those observed in the field. Every morning, the localization of the females was checked. During the five first days together

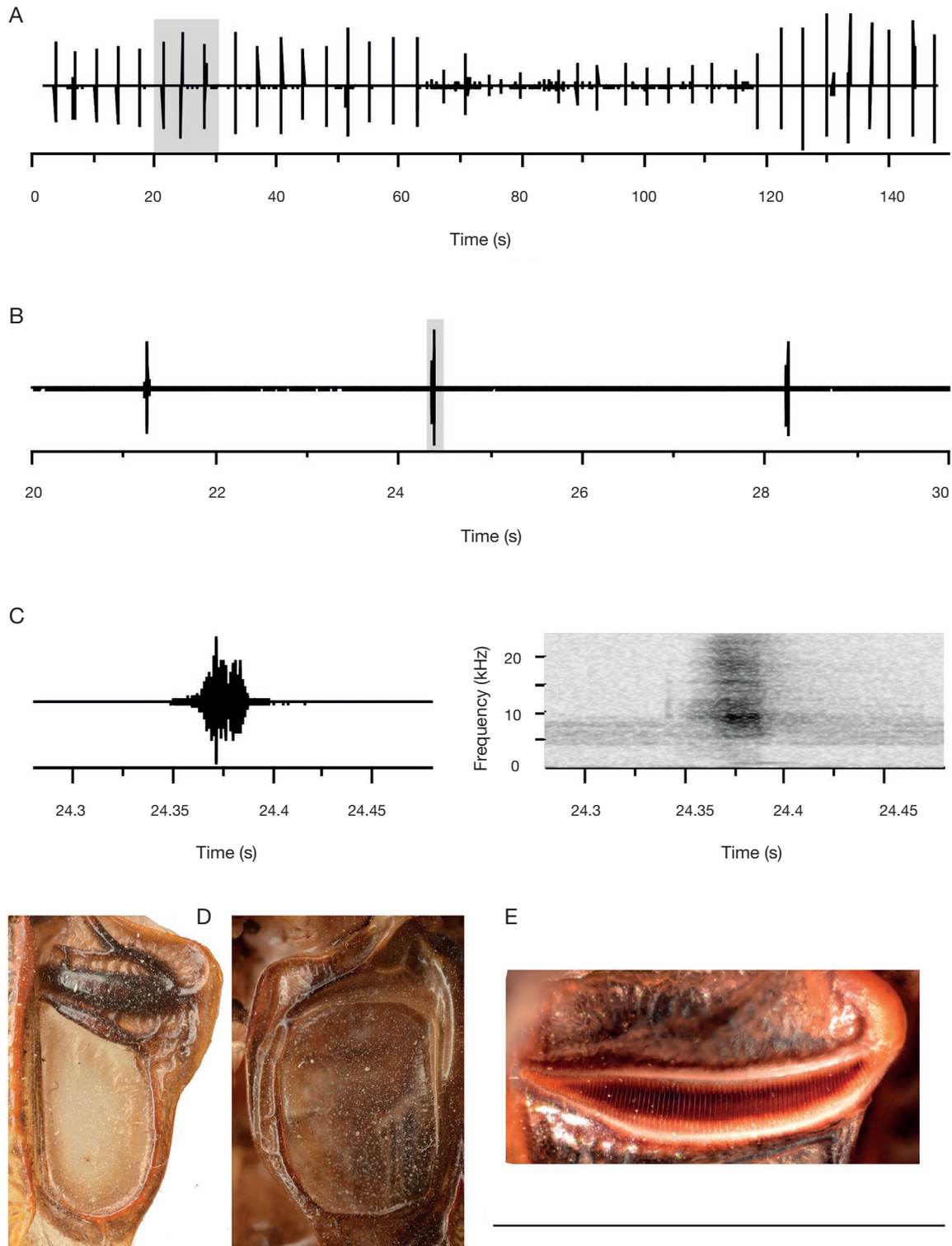


FIG. 7. – Call of *Gnathoclitia vorax* Beier, 1950: **A-C**, Field recording of a male calling from a cavity (26°C); **D**, male stridulatory apparatus in dorsal view; **E**, male file in ventral view. Scale bar: D, 5 mm; E, 2.5 mm.

with the males, both females were observed as in the field: at the bottom of the shelter, a male blocking the entrance. Unfortunately, we have not been able to observe whether the female entered the shelter after mating, and whether the male was actively driving her. After five days, both specimens were

forced out of the shelter; only males returned to the shelter during the subsequent days and restarted to stridulate during nights.

These observations suggest that males of *Gnathoclitia vorax* display a form of mate guarding. Such a behavior was never



FIG. 8. — Shelters of *Gnathoclitia vorax* (Stoll, 1813) in the field: **A**, the face of a male is visible within a hollow shaft of *Astrocaryum* Meyer, 1818 leaf, antennae protruding from the entrance; **A'**, when an aperture is made on the shaft, a female is visible behind the male (at the time of the picture, the male was already collected); **B**, entrance of a shelter (male removed); **C**, disturbed male escaping from the shelter; **D**, picture of the male singing in **A-C** during the call.

formally observed in Tettigoniidae (Gwynne 2001), although a possible type of male guarding behavior was recently reported in *Oncodopus* and *Colossopus* (Unal & Beccaloni 2017).

The function of mate guarding has extensively been discussed in Grylloidea (Simmons 1990; Sakaluk 1991; Hockham & Vahed 1997; Bateman & MacFadyen 1999; Wynn & Vahed 2004; Bussiere *et al.* 2006; Parker 2009; Parker & Vahed 2009) and this behavior was shown to prevent the female from removing the spermatophore of a Gryllidae species (Hockham & Vahed 1997). Since females had no spermatophore attached within the shelter, the mate guarding of *Gnathoclitia vorax* was rather recalling the burrow sharing occurring in some Grylloidea, such as *Gryllus campestris* (Rodriguez-Munoz *et al.* 2011). In this latter species, the burrow sharing was shown to protect from predation and increase mating of paired males).

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REFERENCES

- BATEMAN P. W. & MACFADYEN D. N. 1999. — Mate guarding in the cricket *Gryllodes sigillatus*: influence of multiple potential partners. *Ethology* 105: 949-957. <https://doi.org/10.1046/j.1439-0310.1999.00484.x>
- BEIER M. 1950. — Das Genus *Panoploscelis* Scudder (Orthot.-Pseudophyllinae). *Proceedings of the 8th International Congress of Entomology, Stockholm 1948*: 111-115.
- BEIER M. 1960. — Orthoptera Tettigoniidae (Pseudophyllinae II), in MERTENS R., HENNIG W. & WERMUTH H. (eds), *Das Tierreich*, 74. Walter de Gruyter & Co., Berlin, p. 396.
- BRAUN H. 2011. — *Ottotettix*, a new Katydid genus and species from the rainforest of southern Ecuador (Orthoptera, Tettigoniidae, Pseudophyllinae, Eucocconotini). *Journal of Orthoptera Research* 20 (1): 39-42. <https://doi.org/10.1665/034.020.0103>
- BUSSIERE L. F., HUNT J., JENNIONS M. D. & BROOKS R. 2006. — Sexual conflict and cryptic female choice in the black field cricket, *Teleogryllus commodus*. *Evolution* 60: 792-800. <https://doi.org/10.1554/05-378.1>
- CIGLIANO M. M., BRAUN H., EADES D. C. & OTTE D. — Orthoptera Species File. Version 5.0/5.0. 01/02/2018. <http://Orthoptera.SpeciesFile.org>
- DE HAAN W. 1843. — Bijdragen tot de kennis der Orthoptera. Temminck [Ed.] Verhandelingen over de Natuurlijke Geschiedenis der Nederlandsche Overzeesche Bezittingen 19/20: 208.
- DE JONG C. 1971. — Notes on the types of *Gnathoclitia vorax* (Stoll) (Tettigoniidae, Pterophyllinae). *Zoologische Mededelingen uitgegeven door het Rijksmuseum van Natuurlijke Historie te Leiden* 45 (26): 303-311.
- DE SOUZA L. R., KASUMOVIC M. M., JUDGE K. A. & MORRIS G. K. 2011. — Communicating male size by tremulatory vibration in a Columbian rainforest katydid, *Gnathoclitia sodalis* (Orthoptera, Tettigoniidae). *Behaviour* 148: 341-357. <https://doi.org/10.1163/000579511X559418>
- HOCKHAM L. R. & VAHED K. 1997. — The function of mate guarding in a field cricket (Orthoptera: Gryllidae; *Teleogryllus natalensis* Otte and Cade). *Journal of Insect Behavior* 10: 247-256. <https://doi.org/10.1007/BF02765557>
- GWYNNE D. T. 2001. — *Katydid and Bush-Crickets: Reproductive Behavior and Evolution of the Tettigoniidae*. Cornell Univ. Press, Ithaca, New York, 2001.
- KNUTSON B., BURGDORF J. & PANKSEPP J. 2002. — Ultrasonic vocalizations as indices of affective states in rats. *Psychological Bulletin* 128 (6): 961-977. <https://doi.org/10.1037/0033-2909.128.6.961>
- MONTEALEGRE Z. F. & MORRIS G. K. 1999. — Songs and systematics of some Tettigoniidae from Columbia and Ecuador I. Pseudophyllinae (Orthoptera). *Journal of Orthoptera Research* 8: 163-236.
- MONTEALEGRE Z. F., GUERRA P. A. & MORRIS G. K. 2003. — *Panoploscelis specularis* (Orthoptera: Tettigoniidae: Pseudophyllinae): extraordinary female sound generator, male description, male protest and calling signals. *Journal of Orthoptera Research* 12 (2): 173-181. [https://doi.org/10.1665/1082-6467\(2003\)012\[0173:PSO TPE\]2.0.CO;2](https://doi.org/10.1665/1082-6467(2003)012[0173:PSO TPE]2.0.CO;2)
- NASKRECKI P. 2008. — Katydid of Selected Sites in the Konashen Community Owned Conservation Area (COCA), Southern Guyana. *RAP Bulletin of Biological Assessment*: 25-30.
- SAKALUK S. K. 1991. — Postcopulatory mate guarding in decorated crickets. *Animal Behaviour* 41: 207-216. [https://doi.org/10.1016/S0003-3472\(05\)80472-5](https://doi.org/10.1016/S0003-3472(05)80472-5)
- PARKER D. J. 2009. — Pre and postcopulatory mate choice in *Platygyryllus primiformis*: cryptic female choice and sexual conflict. *Biosci Horizons* 2: 161-171. <https://doi.org/10.1093/biohorizons/hzp019>
- PARKER D. J. & VAHED K. 2010. — The intensity of pre- and postcopulatory mate guarding in relation to spermatophore transfer in the cricket *Gryllus bimaculatus*. *Journal of Ethology* 28: 245-249. <https://doi.org/10.1007/s10164-009-0176-6>
- PASCAL O., TOUROULT J. & BOUCHET P. 2015. — *Expédition “La Planète Revisitée” Guyane 2014-2015. Synthèse des premiers résultats*. Muséum nationale d’Histoire naturelle; Pro-Natura International, 280 p.
- RODRIGUEZ-MUNOZ R., BRETMAN A. & TREGENZA T. 2011. — Guarding Males Protect Females from Predation in a Wild Insect. *Current Biology* 21 (20): 1716-1719. <https://doi.org/10.1016/j.cub.2011.08.053>
- SIMMONS L. W. 1990. — Postcopulatory guarding, female choice and the levels of gregarine infections in the field cricket, *Gryllus bimaculatus*. *Behav Ecol Sociobiol* 26: 403-407. <https://doi.org/10.1007/BF00170897>
- STUMPNER A., DANN A., SCHINK M., GUBERT S. & HUGEL S. 2013. — True katydids (Pseudophyllinae) from Guadeloupe: Acoustic signals and functional considerations of song production. *Journal of Insect Science* 13 (157): 1-16. <https://doi.org/10.1673/031.013.15701>
- TOUROULT J., POLLET M. & PASCAL O. 2018. — Overview of Mitaraka survey: research frame, study site and field protocols, in TOUROULT J. (ed.), “Our Planet Reviewed” 2015 large-scale biotic survey in Mitaraka, French Guiana. *Zoosystema* 40 (13):

- 327-365. <https://doi.org/10.5252/zoosystema2018v40a13>. <http://zoosystema.com/40/13>
- ÜNAL M. & BECCALONI G. W. 2017. — Revision of the Madagascan genera *Oncodopus* Brongniart and *Colossopus* Saussure (Orthoptera: Tettigoniidae: Conocephalinae; Euconchophorini), with description of *Malagasopus* gen. nov. *Zootaxa* 4341 (2): 193-228 <https://doi.org/10.11646/zootaxa.4341.2.2>
- WILLEMSE C. 1954. — *Gnathoclita vorax* (Stoll), a little known Pseudophyllid from Dutch Guiana (Orthoptera). *Ano del Libertador Gral. San Martin* 1: 239-245.
- WYNN H. & VAHED K. 2004. — Male *Gryllus bimaculatus* guard females to delay them from mating with rival males and to obtain repeated copulations. *Journal of Insect Behavior* 17: 53-66. <https://doi.org/10.1023/B:JOIR.0000025132.02196.da>

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