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Anopheles (Nyssorhynchus) atacamensis (Diptera: Culicidae), a new species from Northern Chile

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Anopheles (Nyssorhynchus) atacamensis, a new species in the subgenus Nyssorhynchus, is described and validated using morphological characters of the male and female adult, male genitalia and immature stages. Molecular characterization employing sequences of the ITS2 rDNA and COI mtDNA are provided. The new taxon is compared with Anopheles (Nyssorhynchus) pictipennis (Philippi) from central Chile based on morphological features of the adults, male genitalia and larva. Illustrations of the diagnostic characteristics of the male genitalia, fourth-instar larva and pupa are provided.

Key words: new species - Nyssorhynchus - An. pictipennis - COI - ITS2 - III Región de Atacama - Chile

Anopheles (Nyssorhynchus) atacamensis, a new species, is the third species of the genus Anopheles Meigen reported in Chile and the second of the subgenus Nyssorhynchus Blanchard. This new species was identified as Anopheles (Nyssorhynchus) pictipennis (Philippi) in Northern Chile (Rueda et al. 2008). However, morphological comparisons of characteristics of the male and female adult, male genitalia, fourth-instar larva and pupa between the neotype and specimens of An. pictipennis from III Región de Atacama, Northern Chile, revealed that there are two species under the name An. pictipennis: An. pictipennis s.s. and a new species. The major objectives of this study are to describe and validate the new species from the III Región de Atacama, Chile. Morphological comparisons between An. atacamensis n. sp. and An. pictipennis from the type locality are discussed. Illustrations of diagnostic characteristics of the adults, male genitalia, fourth-instar larva and pupa are provided.

MATERIALS AND METHODS

Mosquitoes were collected in the III Región de Atacama, Chile, as follows: Provincia de Huasco, Carrizarillo (29°06′S 71°24′W), Quebrada Honda, El Sarco (28°13′S 70°07′W) and Provincia de Copiapó, Perales (27°52′S 70°54′W), Sector Totoral (27°53′S 70°56′W), Las Salinas (27°18′S 70°55′W). Larvae were taken from small ground pools. The water was fresh, temperature varying from 14-17°C, deeply shaded, in oasis localities in the Atacama Desert. Immature stages were taken from the habitat with a pipette. Some larvae were preserved in 80% ethanol, whereas others were pooled and transported to the laboratory and reared to obtain adults. The morphological terminology for the adults and immature stages follows Harbach and Knight (1980), for the wing veins we adopted Belkin (1962) and Wilkerson and Peyton (1990) for the wing pale and dark spots. Fourth-instar larvae and pupae were cleared in 10% KOH and mounted on microscope slides. Male genitalia were removed from the adult and macerated in a solution of 20% KOH, stained with acid fuchsin, dehydrated in ETOH from 70-100% and subsequently dissected and mounted on a microscope slide in Canada balsam. Images of the male genitalia, larvae and pupae were recorded with a microscope using the Axiovision software version 3.0.6 SP4 (Carl Zeiss MicroImaging GmbH, Thornwood, NY).

DNA was extracted from two mosquito specimens following the insect DNA extraction protocol provided by the QiAgen DNeasy® Blood and Tissue Kit (Qiagen Ltd, Crawley, UK). Both specimens were collected in Provincia de Huasco, Quebrada Honda, El Sarco, C. Araya coll, in 25th May 2006.

ITS2 ribosomal DNA - Amplification of the ITS2 region was carried out using the 5.8SF (5′-ATC ACT CGG CTC GTG GAT CG - 3′) and 28SR (5′-ATG CTT AAA TTG AGG GGG TAG TC - 3′) primers (Djadid et al. 2007). PCR products were amplified in 25 μL reaction mix containing: 1 μL of DNA of the first elution; 10 mM Tris-HCl, pH 8.3; 50 mM KCl; 1.5 mM MgCl2; 2.5 μL DMSO; 5 pmol of each primer; 200 μM each dNTPs and 2.5 U New England Biolabs® Taq polymerase. PCR protocol consisted of a 2-min denaturation at 94°C, 34 cycles at 94°C, 57°C and 72°C for 30 sec each, followed by a 10-min extension at 72°C. ITS2 PCR amplions obtained from two individuals were purified using PEG precipitation (20% polyethylene glycol 8000/2.5 M NaCl) and cloned into pGem-T Easy Vector (Promega, Madison, WI, USA). Three to five positive clones were sequenced per individual specimen.

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COI mtDNA - COI fragments of ~500 base pairs (bp) were amplified using the primers C1-J-1718 (5'- GGA GGA TTT GGA AAT TGA TTA GTT CC-3') and C1-N-2191 (5'-CCC GTT AAA ATT AAA ATA TAA ACT TC-3') (Simon et al. 1994). Each PCR reaction contained 1 μL DNA; 10 mM Tris-HCl, pH 8.3; 50 mM KCl; 1.5 mM MgCl2; 2.5 μL DMSO; 5 pmol of each primer; 200 μM each dNTPs; and 2.5 U New England Biolabs® Taq polymerase and the remaining volume of ultrapure H2O up to 25 μL. PCR amplification protocol consisted of a 3-min denaturation at 94°C and 35 cycles at 94°C, 55°C and 72°C for 1 min each, followed by a 7-min extension at 72°C. Sequencing reactions were carried out in both directions using ABI Big Dye Terminator Kit v3.1 (PE Applied Biosystems, Warrington, England) with the same primers used for PCR. Sequencing reactions were purified in Sephadex G50® columns (GE Healthcare). Sequences were analyzed on an ABI Prism 3100 - Avant (Gene Codes Corporation, Ann Arbor, USA).

Template DNA from this study is retained at -70°C in the Faculdade de Saúde Pública (FSP-USP) for future reference. DNA vouchers consist of wings and legs mounted on microscope slides (E-12992) and head, thorax, one wing and legs pinned and one wing mounted on microscope slide (E-12993). Sequences were aligned and optimized manually in MacClade version 4.8 (Maddison & Maddison 2000). Sequence similarities of the ITS2 and COI sequences generated in this study with those previously available in GenBank was assessed using the Basic Local Alignment Tool search (http://www.ncbi.nlm.nih.gov/BLAST) and documented using PAUP* 4.0 software (Swofford 2003).

RESULTS

*An. (Nyssorhynchus) atacamensis* González and Sallum, n. sp.

*An. pictipennis* of Rueda et al. (2008): 448.

Etymology - The name, *atacamensis*, is derived from the name of the geographical locality where the specimens were first collected, the Atacama Region of Northern Chile.

Diagnosis - In the adult stage, *An. atacamensis* can be recognized by a combination of characters as follows: dark grayish pruinose, central portion of the head bare. Scutal integument grayish, scutal scales large, with a pair of broad, subdorsal stripes extending posteriorly from anterior scutal angle to scutellum. Acrostichal area with two lines of white spateulate scales, reaching anterior margin of scutellum, two lines of white spateulate scales extending posteriorly along dorsocentral area scales, white spateulate scales scattered on posterior portion of dorsocentral area, scutal fossa with few scattered, white spateulate scales, becoming a line of white spateulate scales on prealar, supraalar areas; lateral borders of scutum covered with line of white spateulate scales reaching posterior end of paratergite; scutal setae brown with golden reflections. Scutellum pale dusted with long dark setae along posterior margin, with white, spateulate scales anterior to setae, intermixed with dark, long setae with golden reflection, white spateulate scales scattered on posterior portion of dorsocentral area, becoming a line of scales at lateral portions of prescutellar area, scutal fossa with few scattered, white spateulate scales, scales more concentrated on prescutal suture, becoming a line of white spateulate scales on prealar, supraalar areas; lateral borders of scutum covered with line of white spateulate scales reaching posterior end of paratergite; scutal setae brown with golden reflections. Scutellum pale dusted with long dark setae along posterior margin, with white, spateulate scales anterior to setae, intermixed with median of median statistic area, lobe brown. Antenna with with patch of long dark scales at dorsomedial portion, with scattered dark long setae. Thoracic integument grayish. Upper proepisternum with 4-6 setae, with few white scales; prespiracular area without setae, with 2,3 white spateulate scales; prealar area with setae intermixed with white spateulate scales; upper mesokatepisternum with dark brown setae, with a patch of white spateulate scale, posterior border of mesokatepisternum with a patch of

**Female** - Integument dark with grayish pruinose.
<table>
<thead>
<tr>
<th></th>
<th>An. pictipennis</th>
<th>An. atacamensis (F1)</th>
<th>An. atacamensis (F2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flagellomere 1</td>
<td>without scales on dorsal surface</td>
<td>without scales on dorsal surface</td>
<td>with narrow scales on dorsal surface</td>
</tr>
<tr>
<td>Clypeus</td>
<td>oval</td>
<td>square</td>
<td>rounded</td>
</tr>
<tr>
<td>Proboscis, apex at margin of labella</td>
<td>with pale scales</td>
<td>with pale scales</td>
<td>with dark scales</td>
</tr>
<tr>
<td>Scutum</td>
<td>grayish; scales on posterior region of the acrostichal and dorsocentral areas spatulate; prescutellar area with lateral line of white scales</td>
<td>grayish; scales on posterior region of the acrostichal and dorsocentral areas spatulate; prescutellar area with lateral line of white scales</td>
<td>blackish; scales on posterior region of the acrostichal and dorsocentral areas narrow spatulate; prescutellar area with sparse, white scales</td>
</tr>
<tr>
<td>Scutellum</td>
<td>pale dusted with white, spatulate scales on median lobe, long, narrow, truncate white scales on lateral lobes</td>
<td>with scales on lateral lobes and laterally on medial lobe</td>
<td>with scales posteriorly on lateral and medial lobes</td>
</tr>
<tr>
<td>Hindtarsomere 2</td>
<td>dark-scaled on basal 0.7, white-scaled on apical 0.3</td>
<td>dark-scaled on basal 0.8, white-scaled on apical 0.2</td>
<td>dark</td>
</tr>
<tr>
<td>Hindtarsomere 3</td>
<td>entirely white</td>
<td>entirely white</td>
<td>dark-scaled at basal 0.2, white at apical 0.8</td>
</tr>
<tr>
<td>Hindtarsomere 5</td>
<td>entirely white</td>
<td>white, with apical dark ring</td>
<td>white, with apical dark ring</td>
</tr>
<tr>
<td>Scales on dorsal surface of R_{4+5}</td>
<td>mostly pale-scaled, a small postbasal dark patch, and a small preapical dark patch</td>
<td>proximal end with a small white patch, a postbasal dark patch, remaining of vein white-scaled</td>
<td>a dark patch on proximal 0.2, white patch at middle, a dark patch on preapical portion, and a small white patch at apex</td>
</tr>
<tr>
<td>Scales on dorsal surface of Cu</td>
<td>mostly pale-scaled, with dark scales on proximal 0.5 of distal 0.5, pale-scaled on distal 0.5, bifurcation of Cu₁ and Cu₂, pale-scaled</td>
<td>with pale scales on proximal 0.5, distal 0.4 dark scaled, dark-scales not reaching bifurcation of Cu₁ and Cu which is pale-scaled</td>
<td>with pale scales on proximal 0.5, distal 0.5 dark-scales, reaching bifurcation of Cu₁ and Cu, bifurcation of Cu₁ and Cu₂, pale-scaled</td>
</tr>
<tr>
<td>Scales on dorsal surface of Cu₁</td>
<td>proximal 0.5 pale-scaled, interrupted by two submedian dark spots, distal 0.5 dark-scaled, apex pale</td>
<td>proximal 0.5 white-scaled, with a dark and white patches at middle, dark-scaled on distal 0.3, apex with a small white patch</td>
<td>proximal 0.5 dark with pale scales at proximal end, a small submedian pale patch, dark-scaled distally with a few pale scales at distal apex</td>
</tr>
<tr>
<td>Scales on dorsal surface of Cu₂</td>
<td>proximal 0.5 pale-scaled, distal 0.5 dark-scaled, apex pale</td>
<td>mostly white-scaled, a dark preapical patch, apex pale</td>
<td>pale-scaled, distal 0.5 dark-scaled, apex pale</td>
</tr>
</tbody>
</table>

F1, F2: morphological forms of *An. atacamensis* from distinct localities in the III Región de Atacama, Northern Chile.

White scales at level of upper margin of mesomeron; upper mesepimeron with 5-8 setae, a patch of white scales anterior to setae. Legs. Coxas and trochanters grayish; forecoxa with white scale-patches on dorsal and ventral margins of anterior surface; midcoxa with white scales on dorsal anterior and posterior surfaces and on ventral posterior surface; hindcoxa with white scales on dorsal and ventral margins of lateral surface and a patch on ventral margin of posterior surface. Forefemur with white and dark scales, scales at base forming pale and dark rings, anterior surface mostly dark with scattered pale scales along margins extending most of the length of the segment, apex usually with white scales; midfemur with pale and dark scales at base forming pale and dark bands, anterior surface dark-scaled, with white scales at apex, posterior surface mostly pale-scaled, dark-scaled at basal 0.3, lateral surface mostly dark with ovoid, white patch extending to anterior surface; hindfemur with pale and
dark scales at base forming one pale and one dark band, anterior surface dark with narrow line of white scales extending from a large, elongate, subapical white patch, pale scales at apex; posterior surface white-scaled, dark at subapical portion, with white scales at apex. Tibiae with pale and dark golden-brown scales forming longitudinal pale stripes, mid and hindtibiae with longitudinal line of white scales interrupted at subapical portion of both ends, with patches of white scales at apex; tarsi with scales varying from white to golden to black; fore, mid and hindtarsomeres 1 mostly pale-scaled, with dark ring at base, dark at subapical portion, with broad apical band of white scales; foretarsomeres 2,3 dark-scaled, with apical ring of variable length of white scales, foretarsomeres 4,5 dark-scaled; midtarsomere 2 dark-scaled, with white ring at apex, midtarsomeres 3,4 and 5 dark-scaled; hindtarsomere 2 dark-scaled at basal 0.8, white-scaled at apical 0.2, hindtarsomeres 3,4 entirely white-scaled, hindtarsomere 5 mostly white-scaled with dark ring at apex. Wing. Length 4.02-4.85 mm (mean = 4.29; n = 6 wings of 3 specimens). Dark scales brown, pale wing scales white on most proximal pale spots and posterior veins, pale yellowish on subcostal pale, preapical pale and apical pale spots. Costa with prehumeral dark, presector dark, sector dark, preapical dark and apical dark spots, basal and prehumeral pale spots fused, humeral pale, accessory sector pale, subcostal pale, preapical pale and apical pale spots; vein Rs dark-scaled; R_{2,3} dark-scaled at base and apex; R_{1,5} with one or two pale scales at proximal end, dark-scaled on proximal 0.2, pale-scaled on distal 0.8. Vein M mostly dark-scaled, with pale spot at base and medial portion, with a pale spot at basal 0.5, dark-scaled distally, split of veins M_{1,2} and M_{1,4} pale-scaled. Vein Cu pale-scaled on proximal 0.6, dark-scaled distally; vein 1A dark-scaled with small pale spots at proximal and distal ends and along middle. Pale fringe spots at distal end of each vein. Halter. Scabellum and pedicel with pale brown integument, capitellum dark brown-scaled.

Abdomen: integument dark brown, terga II-VII with dense covering of whitish scales medially, without caudolateral scale-tufts, tergum VIII covered with scales, terga with long pale brown and yellowish setae. Sternum I bare, sterna II-V with longitudinal patches of pale scales lateral of longitudinal midline, sterna VI-VII pale-scaled medially, sternum VIII mostly pale-scaled.

Male (Figs 1, 2) - Similar to female except for the following sexual differences: ratio of maxillary palpus and proboscis length 0.90-1.12 (mean = 1.01; n = 3), mostly dark-scaled, with a white band at apex of palpomere 2, palpomere 3 dark-scaled with a broad white band at subapical portion extending laterally and ventrally and wide white band at apex; palpomere 4 dark with white band at apex; palpomere 5 dark basally, white-scaled on apical 0.5; palpomere 3 with tufts of long, strong, dark setae at apex of ventral surface, palpomeres 4 and 5 some-

Fig. 1: Anopheles (Nyssorhynchus) atacamensis, male genitalia. A: segment IX; B: proctiger; C: dorsal accessory seta; D: apex of ventral accessory seta.
what fused, enlarged, palpomere 4 with long, dark setae extending from base to apex of dorsal and ventral margins of inner surface. Genitalia. Tergum VIII with few short setae distally, with long, spatulate scales scattered on segment, more numerous on basal 0.5; sternum VIII with scattered short to long setae and few sparse, slender, spatulate scales, distal margin convex. Segment IX spiculate (Fig. 1A); tergum poorly developed, membranous, developed laterally, sternum well developed, subrectangular, distal margin superficially emarginated, with shallow depression at middle, apodeme absent. Tergum X, paraproct and lateral portions of cerci spicate, spicules extending to posterior end of proctiger (Fig. 1B). Gonocoxite cylindrical, mesally straight, dorsolateral surface with long, spatulate scales from base to apex, tergal surface with short to long setae, with 4,6 (n = 4) tergomesal setae; tergomesal surface with two accessory setae (Fig. 1C, D) arising from two well-developed prominent tubercles, dorsal seta arising 0.45-0.63 (mean = 0.56; n = 4) from base of gonocoxite, ventral seta arising 0.5-0.69 (mean = 0.61; n = 4) from base of gonocoxite, accessory setae cylindrical, sharply pointed at apex, straight or slightly curved apically, dorsal and ventral setae similar in length and development; internal seta arises from a small tubercle, 0.58-0.69 (mean = 0.64; n = 4) from base of gonocoxite; parabasal setae strong, short, hooked at apex, inserted in a short tubercle. Gonostylus 0.82-0.86 (mean = 0.92; n = 4) length of gonocoxite, curved mesally along entire length, slightly broad at base ending in a somewhat truncate apex, gonostylar claw short, strong arising from apex of ventral surface; ventromesal surface with short, slender setae. Dorsal claspette (Fig. 2A) with two large setae curved at apical 0.5 and 1 smaller, more slender, straight seta, pedicel somewhat subrectangular, with rounded base and one protuberance at base of ventral surface. Ventral claspette (in ventral view) (Fig. 2B) entirely bare, somewhat rounded, tapering to apex, apex narrow, ventrally with numerous fine laterally directed striations, median sulcus with distal opening straight, long, strongly sclerotized with more sclerotized area approximately at mid-length, claspette with two well-developed lobes well expanded ventrally (Fig. 2C, D).
2C), continuing to basal lobules which are expanded ventrally, semicircular along margin; refringent structure absent. Aedeagus (Fig. 2D, E) with apex longer than wide, apex curved dorsally, strongly sclerotized laterally, weakly sclerotized at apex and on lateral margins; ventromesal subtriangular projections strongly sclerotized, meeting on midline forming a subapical collar, with a pair of well-developed subapical leaflets; leaflets straight, strongly sclerotized, dorsolaterally projecting, inner margin straight, smooth, dorsal, ventral and lateral surfaces strongly serrated, ending in a narrow point, leaflets arising subapically from a membranous lateral area at apex of aedeagus.

Fourth-instar larva (Fig. 3) - Head: with dark pigmented areas, posteriorly on dorsal apodeme, posteriorly on ecdisial line and lateralia, posteriorly on labiogula. Collar well developed, moderately large, strongly sclerotized. Dorsomentum well developed, strongly sclerotized. Ventromentum well developed, strongly sclerotized with three lateral teeth. Hypostomal suture short, not extending to tentorial pit. Median labral plate moderately developed, with small lateral lobe bearing seta 1-C. Seta 2-C widely spaced, single with few moderately developed spicules distally, 3-C shorter than 2-C, spiculate, distance between 2-C equal distance 2-C to 3-C, clypeal index 1.0, 4-C strongly developed, similar to 3-C, single, simple, extending beyond base of 2-C. Setae 5-7-C plumose, long, 5-C with 12 branches, extending well beyond base of 2-C, 6-C with 13 branches, extending to base of 3-C, 7-C with 15 short branches, inserted in line with insertions of 5,6-C, 8-C with two or three branches, 9-C with three branches, 10-C triple, 11-C plumose, number of branches not counted, 12-C with four or five branches, 13-C with three branches, 14-C min, branched, 15-C with two or three branches, mesad of hypostomal suture. Antenna. Flagellum strongly aciculate, with strong spicules on dorsal, ventral and mesal surfaces, seta 1-A longer than width of flagellum at the point of insertion, moderately large, arising from dorsomesal surface, about 0.3 from base.

Thorax: setae 9-12-P,M,T inserted on large common tubercle, with a large, moderately pigmented, lateral spine. Prothorax. Seta 1-P fanlike (Fig. 3A) with thin branches, not inserted on common tubercle with 2-P (Fig. 3B), 2-P plumose with nine branches inserted on large tubercle closer to 3-P than 1-P, 3-P single, simple, short, inserted on tubercle with 2-P, 4-P plumose with 25 branches, 5-P plumose, 6-P long, single, simple, 7-P plumose, 8-P plumose with nine or 10 branches inserted on a moderately large tubercle lateral of setae 9-12-P, 14-P with three branches, moderately long. Mesothorax. Seta 1-M plumose, moderately long, inserted on large tubercle, 2-M single, simple or forked, 3,5-M single, simple, 4-M with three branches, shorter than 3-M, 6-M with three branches, moderately long, 7-M with four branches similar to 4-M, 8-M plumose, 14-M pectinate, small with six branches arising near ventral midline. Metathorax. Seta 1,2-T single and simple, short, 1-T similar to 2-T, 3-T fanlike with six thin branches (Fig. 3B), 4-T small with three or four branches, 13-T double.

Abdomen: median tergal plates present, moderately large, strongly sclerotized, arising on anterior margin of segments I-VIII, accessory median and submedian tergal plates arising posterior of median tergal plates, subspherical, small. Seta 1-I fanlike with seven branches (Fig. 3C), 1-II-VII palmate with serrated moderately broad, lanceolate branches (Fig. 3D, E). Seta 2-I small, bifid or triple, 2-II moderately large with six branches, 2-III...
large, triple, 2-IV single, simple, 2-V double or triple, 2-VI triple, 2-VII triple. Seta 3-I single, 3-II single, 3-III single, 3-IV single, 3-V single, 3-VI single, 3-VII triple. Seta 6-I-III plumose, inserted on distinct tubercle, 6-IV-VI long, branched (Fig. 3F), 6-VII branched. Seta 7-I-II plumose, long, inserted on tubercle with 6-I,II, 7-III with four branches, 7-IV triple, 7-V triple, 7-VI triple, 7-VII with five branches. Seta 8-III double, 8-IV double, 8-V double, 8-VI double, 8-VII with four or five branches. Seta 9-I with five branches, 9-II with six branches, 9-III with six branches, 9-IV with four branches, 9-V with three branches, 9-VI triple, 9-VII with five branches. Seta 10-I single, 10-II single, 10-III single, 10-IV single, 10-V single, 10-VI single, 10-VII with four branches. Seta 11-I triple, 11-II single, 11-III single, 11-IV single, 11-VI single, 11-VII single. Seta 12-I with three branches, 12-II single, 12-III single, 12-IV double, 12-V single, 12-VI single, 12-VII single. Seta 13-I with three branches, 13-II with four branches, 13-III with three branches, 13-IV with five branches, 13-V with three branches, 13-VI with five branches, 13-VII with four branches. Segment VIII. Seta 1-VIII triple, 2-VIII with five branches, 3-VIII with five branches, 4-VIII single, 5-VIII with five branches. Spiracular lobe. Pecten with seven long teeth intermixed with 12 short teeth, teeth length variable, teeth usually curved dorsally with serrations on base. Median plate strongly pigmented without lateral arms, anterior median process of median plate strongly sclerotized and short. Seta 1-S strongly developed with four branches, inserted on caudal surface of posterolateral spiracular lobe near posterodorsal margin of pecten, 6-S with four branches, 7-S single, 8-S with five branches, 9-S double, 10-13-S single, small. Segment X. Saddle incomplete, 1-X single, well developed, long, inserted outside the saddle, 2-X strongly plumose, seta 4-X with 8.5 pairs of long, strongly developed, plumose setae.

**Pupa** (Fig. 4): cephalothorax. Seta 2-C double. Seta 3-C triple. Seta 8-C single, well developed, heavily pigmented, inserted at level of base of trumpet. Seta 9-C with 2-5 branches, small, inserted posterior to base of trumpet. Trumpet. Gradually broadening from base to apex, angusticorn. Meatal cleft large. Metanotum. Seta 10, 11-C subequal, 12-C longer. Seta 10-C single to triple. Seta 11-C with 3,4 branches. Seta 12-C single or double. Abdomen. Weakly pigmented, light yellowish brown, concolorous with cephalothorax. Usually with more strongly pigmented area on anteromesal portion of terga II-VII. Seta 1-V strong, long, single, extending slightly beyond apex of following segment, similar to 1-V. Seta 1-VI strong, long, single, extending beyond apex of following segment. Seta 5-VI similar to 1-VI, strong, long, single, extending beyond apex of following segment. Seta 9-IV-VI spiniform, inserted on caudola-

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**Fig. 4: Anopheles (Nyssorhynchus) pictipennis and Anopheles (Nyssorhynchus) atacamensis**, pupa. A: paddle outer basolateral serrations in *An. pictipennis*; B: seta 1-Pa of *An. pictipennis*; C, D: paddle outer basolateral serrations in *An. atacamensis*; E: seta 1-Pa of *An. atacamensis*; F: seta 9-VIII of *An. pictipennis*; G: seta 9-VIII of *An. atacamensis*. 
teral margin of segment, 9-VII, inserted on caudalateral margin of segment, single, 9-VIII spiniform, single, inserted on caudalateral margin of segment (Fig. 4G), seta 9-IV-VIII heavily pigmented, seta 9-V-VIII long, strong, curved. Paddle. Lightly tanned, outer basolateral serrations prominent (Fig. 4C), filamentous spicules on outer margin prominent; seta 1-Pa single, darkly pigmented, single, curved (Fig. 4E), 2-Pa single or double; toothed margin index 3.8-4.5, paddle teeth well developed (Fig. 4D), teeth broad at base tapering to apex.

**DNA sequences** - The ITS2 and COI sequences generated during this study are available in GenBank under the following accession numbers: ITS2 (GQ902959-GQ902965), COI (F1, GQ902966; F2, GQ902967). Seven clones of the entire ITS2 were sequenced for one individual of *An. atacamensis* (F1, GQGQ902959-GQ902961) and one individual of *An. atacamensis* (F2, GQ902962-GQ902965). The latter is designated the melanic form because the adult has hindtarsomere 3 dark-scaled on the basal 0.2, whereas it is entirely white-scaled in F1. The fragment amplified included 119-123 bp of the flanking 5.8S gene and 31-32 bp of the 28S gene. The cloned sequences were aligned with a single sequence of *An. pictipennis* from the Atacama Region, Northern Chile (EU433947; Rueda et al. 2008). The length of the ITS2 sequences, excluding the 5.8S and 28S, varied from 426-432 bp with length variation due to a 6-bp indel (Table II). Two cloned sequences of one individual showed a 6-bp insertion, whereas one cloned sequence of the same individual had a 6-bp deletion. Four cloned sequences of the second individual and EU433947 showed the same 6-bp deletion (Table II). A few base substitutions were observed; however, they were not consistent among the clones. We estimated mean sequence divergence of ITS2 variants within individuals using uncorrected P distance in PAUP (Swofford 2003). Sequence divergence among the clones of *An. atacamensis* F1 and F2 and EU433947 ranged from 0.00000-0.00769 whereas those of *An. atacamensis* F1 ranged from 0.00170-0.00520 and *An. atacamensis* F2 ranged from 0.00172-0.00516. One sequence derived from F1 shares 100% similarity with a sequence derived from F2 and F1 and EU433947. The nucleotide bases of the ITS2 fragment consist of 22% A, 19% T, 27% C and 31% G. A single mtDNA haplotype was detected in the 501-bp COI fragment for both F1 and F2. The nucleotide bases of the COI fragment consist of 30% A, 38% T, 17% C and 15% G. Among sequences available in GenBank, the COI gene of *An. atacamensis* shares the highest similarity (92% in query coverage of 99%) to that of *Anopheles muneztorvari* (AF368115), whereas in query coverage of 100%, the highest similarity is shared with *Anopheles albitarsis* E (DQ076231).

**Material examined** - Holotype - Pinned adult male with associated genitalia on a microscope slide, bearing the following collection data: Chile, III Región de Atacama, Provincia de Copiapó, Los Perales 30/06/2007 C Reyes Coll, deposited at FSP-USP (E-12984). Paratypes: Chile, III Región de Atacama, Provincia de Copiapó, one male Los Perales, 30/06/2007 C Reyes Coll (FSP-USP, E-12985), one female Totoral 30/06/2007 C Reyes Coll (FSP-USP, E-12994), one male Las Salinas, 02/03/2009 C Reyes Coll (IE-UMCE), one female Las Salinas, 02/03/2009; Provincia de Huasco, one female Quebrada Honda, El Sarco 25/05/2006 C Araya Coll (FSP-USP, E-12995). Three fourth-instar larvae mounted on a separate microscope slides, Provincia de Huasco, Carrizalillo 28/02/2009, C Reyes Coll (FSP-USP E-12986, E-12987, E-12988); three fourth-instar larvae mounted on separate microscope slides, Provincia de Copiapó, Las Salinas 02/03/2009, C Aguilera Coll (FSP-USP, E-12989, E-12990, E-12991).

**Other material** - Nine specimens of *An. pictipennis* with the following data: Santiago, Rio Mapocho, A. Neghme Coll., as follow: one pinned adult male with associated male genitalia on microscope slide and two separate slides, one with the right wing and the second with four legs; one male genitalia mounted on microscope slide, not associated with the adult; one pinned adult female; one fourth-instar and one first-instar larva mounted on a separate microscope slides; three pupal exuviae mounted on a single microscope slide; eggs mounted on one microscope slide. Collection 6000-6002; 6126-6128; 6251; 6742-6744. Six fourth-instar larvae from Santiago, location not specified, A. Neghme Coll. 1946, Lane and Neghme Det. 1947. Collection FSP-1 6857-6862. Two females of *An. atacamensis* for DNA studies (FSP-USP, E-12992, E-12993).

**Distribution** - *An. atacamensis* is known from Chile, III Región de Atacama, Provincia de Huasco: Carrizalillo (29°06’S 71°24’W), Quebrada Honda El Sarco (28°13’S 71°07’W); Provincia de Copiapó: Perales (27°52’S 70°54’W), Sector Totoral (27°53’S 70°56’W), Las Salinas (27°18’S 70°55’W) and Puerto Viejo (27°19’S 70°5’W) (Rueda et al. 2008).

**Bionomics** (Fig. 5A, B) - Larvae and pupae were found in small, permanent ground pools in oasis areas in the Atacama Desert. The source is ground water that emerges as small fountains in the oasis in Carrizalillo (Provincia de Huasco) (Fig. 5A). The water from the fountains is used to irrigate olive tree plantations, the major crop in the area. The water is clear, fresh, moderately cold (varying from 14-17°C), partially shaded with mud and organic matter on the bottom. Immatures stages were also collected in the Copiapó River (Las Salinas Sector) (Fig. 5B) among Juncaceae vegetation, in a slow current, about 5 km from the mouth of the river to the Pacific Ocean.

**DISCUSSION**

In considering that *An. atacamensis* is morphologically more similar to *An. pictipennis* than to any other species of the subgenus *Nyssorhynchus*, this new species is included in the Pictipennis Subgroup of the Argyritarsis Section of the subgenus. Because of the morphological similarity, specimens of *An. atacamensis* from the Atacama Region of Northern Chile were misidentified as *An. pictipennis* by Rueda et al. (2008). However, these species can be distinguished based on characters of the
TABLE II
Alignment of the 5.8S, ITS2 and 28S sequences of *Anopheles atacamensis* from distinct localities in the III Región de Atacama, Northern Chile

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- either an indel or missing data at the 5' and 3' ends.

male and female adults, male genitalia and fourth-instar larvae. Differential characters of the adult stages of *An. atacamensis* and *An. pictipennis* are listed in Table I. Based on the male genitalia, *An. atacamensis* can be distinguished from *An. pictipennis* by having the apex of the aedeagus tapered to a blunt point (Fig. 2E), ventromesal subtriangular projection extending ventromedially from the middle length of the aedeagus to form a broad fused arch (Fig. 2D), ventral claspette well projected ventrally, forming two hyaline sacciform, somewhat oval lobes (Fig. 2C) bordering the mesal cleft, basal lobe is elongate, small and narrow and the median sulcus is narrowed at the distal and basal ends and is slightly broader at the middle (Fig. 2B). In *An. pictipennis* the apex of the
The morphological variant found in the Atacama Region may represent a melanic form of *An. atacamensis*. A similar polymorphism is known in *An. albitarsis* Lynch Arribálzaga, i.e., *An. albitarsis imperfectus* Corrêa and Ramos was defined based on the presence of a dark band on hindtarsomere 3 (Corrêa & Ramos 1943). However, this characteristic was found to be a morphological variation and thus *An. imperfectus* was synonymized with *An. albitarsis* by Lane (1953). Wilkerson et al. (1995) proposed that in the subgenus *Nyssorhynchus*, those individuals with a long dark basal band on hindtarsomere 2 are more likely to have a dark basal band on hindtarsomere 3. This seems to be the case of the melanic form of *An. atacamensis*, which has a long dark basal band on hindtarsomere 2 and also a basal dark band on hindtarsomere 3. The COI sequences tell us that these two forms are conspecific because they share a unique haplotype. The ITS2 sequences show singleton polymorphisms and a 6-bp indel among the clones. However, none were consistent and one clone of form F1 and one of F2 share 100% similarity, suggesting that the variation observed in the ITS2 represents intragenomic variation. It is noteworthy that F1 and F2 were from distinct localities in the Atacama Desert. We expected to find COI variation between these forms, because the populations are isolated in distinct oases in the desert. However, the sample size (n = 2) was too small, consequently we cannot draw conclusions about the affinities between these individuals.

Although included in the Argyritarsis Section, *An. atacamensis* and *An. pictipennis* share morphological characters with members of the Myzorhynchella Section (Peyton et al. 1992) of the subgenus *Nyssorhynchus*. Diagnostic characters include the following: in adults, posterolateral scale-tufts are absent from the abdominal segments, in the male genitalia, the ventral claspette lacks spicules on both the ventral and dorsal surfaces, the protiger is spicate, the dorsal claspette possesses one subapical and two apical setae, both dorsal and ventral accessory setae end in an acute tip, the aedeagus is strongly developed with serrated lateral leaflets that arise subapically and the ventromesal subtriangular projection extends ventromesally from mid-length of the abdominal setae, forming a distinct arch; in larvae abdominal seta 6-IV-VII is branched.

The adults of *An. atacamensis* and *An. pictipennis* differ from those of the Myzorhynchella Section in having scales on the abdominal terga and the scutum covered with large scales. It is noteworthy that *An. pictipennis* and *An. atacamensis* differ from other species of the subgenus *Nyssorhynchus* by having larval seta 1-II-VII with lanceolate leaflets with notched margins. These species share this type of leaflets with members of the subgenera *Anopheles*, *Lophopodomyia* and *Stethomyia*. Consequently, when using Linthicum’s (1988) larval key to identify subgenera of *Anopheles*, larvae of *An. pictipennis* and *An. atacamensis* can be keyed out erroneously to those subgenera.

Edwards (1932) considered *Myzorhynchella* as a subgenus of *Anopheles* and *An. pictipennis* a species of that group. Forattini (1962) adopted *Myzorhynchella* as
a subgenus of *Anopheles*, but *An. pictipennis* was not assigned to any subgenus. Later, Peyton et al. (1992) defined Myzorhynchella as a section of *Nyssorhynchus* and retained *An. pictipennis* in the Argyritarsis Section of *Linthicum* (1988). Moreover, it is noteworthy that *An. pictipennis* was not included in the phylogenetic analyses of the Anophelinae conducted by Sallum et al. (2000, 2002), Krzywinski et al. (2001a, b) and Harbach and Kitching (2005). Consequently, the position of the Pictipennis Subgroup within *Anopheles* is uncertain.

*An. pictipennis* was collected in the geographically isolated desert zone of Northern Chile, characterized by hyper-aridity that developed progressively with the uplift of the Andes as they reached elevations between 1,000-2,000 m, coupled with the intensification of a cold, upwelling Humboldt Current (Hartley & Chong 2002, Houston & Hartley 2003). The climate of this zone is controlled by its location (15° and 30°S) and the Humboldt current (Houston 2006). This isolation suggests an independent evolution of *An. pictipennis*, in the South, with respect to *An. atacamensis*, as it has apparently occurred with others species in the Atacama Desert (Frias et al. 1998). Probably, *An. atacamensis* was isolated from other populations by the climatic events that maintain the hyper-aridity of the Atacama Desert. The biota of the Northern Chile is more closely related to Neotropical species than the rest of Chilean insects (Morrone 2006). The southern South American entomofauna is more closely related to taxa of the Australasian Region than to Neotropical taxa (Gondwana elements) (Morrone 1994, Bourdon et al. 2009, Ribeiro 2009), suggesting that the southern biota may have a composite origin. This could explain the morphological similarity observed among the adults and larvae of *An. atacamensis* that is shared with species of the Myzorhynchella Section. This section of the subgenus *Nyssorhynchus* has a geographic distribution that includes regions of Brazil, Bolivia, Peru and Northern Argentina (Galvão 1941, Forattini 1962), but is mainly restricted to Brazil.

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**REFERENCES**


