Open Access

# The True Identity of the New World Iguanid Lizard Liolaemus chillanensis Müller and Hellmich 1932 (Iguania: Liolaemidae) and Description of a New Species in the Liolaemus elongatus Group 

Jaime Troncoso-Palacios ${ }^{1, *}$, Damien Esquerré ${ }^{2}$, Félix A. Urra ${ }^{3,4}$, Hugo A. Díaz ${ }^{5}$, Carlos CastroPastene ${ }^{6}$, and María Soledad Ruiz ${ }^{7}$<br>${ }^{1}$ Programa de Fisiología y Biofísica, Facultad de Medicina, Universidad de Chile, Independencia 1027, Santiago, Chile<br>${ }^{2}$ Division of Ecology and Evolution, Research School of Biology, The Australian National University 0200, Canberra, Australia<br>${ }^{3}$ Programa de Anatomía y Biología del Desarrollo, Instituto de Ciencias Biomédicas (ICBM), Facultad de Medicina, Universidad de Chile and Fondap Geroscience Center for Brain Health and Metabolism, Independencia 1027, Santiago, Chile<br>${ }^{4}$ Programa de Farmacología Molecular y Clínica, Instituto de Ciencias Biomédicas (ICBM), Facultad de Medicina, Universidad de Chile, Independencia 1027, Santiago, Chile<br>${ }^{5}$ Departamento de Ciencias Ecológicas, Facultad de Ciencias, Universidad de Chile, Las Palmeras 3425, Santiago, Chile<br>${ }^{6}$ Laboratorio de Vida Silvestre y Ecología, Facultad de Ciencias Veterinarias, Universidad de Concepción, Vicente Mendéz 595,Chillán, Chile<br>${ }^{7}$ Instituto de Bio y Geociencias del NOA (IBIGEO)CONICET. 9 de Julio 14, Rosario de Lerma, Salta, Argentina

(Received 27 August 2017; Accepted 19 April 2018; Published 31 May 2018; Communicated by Benny K.K. Chan)
Citation: Troncoso-Palacios J, Esquerré D, Urra FA, Díaz HA, Castro-Pastene C, Ruiz MS. The true identity of the new world iguanid lizard Liolaemus chillanensis Müller and Hellmich 1932 (Iguania: Liolaemidae) and description of a new species in the Liolaemus elongatus group. Zool Stud 57:22. doi:10.6620/ZS.2018.57-22.


#### Abstract

Jaime Troncoso-Palacios, Damien Esquerré, Félix A. Urra, Hugo A. Díaz, Carlos Castro-Pastene, and María Soledad Ruiz (2018) Liolaemus is a particularly species-rich radiation of New World iguanid lizards from southern South America. Thanks to intense systematic and taxonomic research, the knowledge on its specieslevel diversity has increased rapidly over the past several years. The L. elongatus species-complex has received considerable attention and a remarkable case is Liolaemus chillanensis, a name that has been used for two different species that are sympatric in Termas de Chillán, central Chile. Since the holotype is lost, we propose that the first step to identify the true L. chillanensis is through the analysis of the original description. Then we provide a morphological and molecular characterization of $L$. chillanensis based on topotypes and a description of the taxon previously confused with it.


Key words: Chile, Cytochrome b, Liolaemus monticola, Phylogeny, Principal component analysis.

## BACKGROUND

Liolaemus Wiegmann, 1834 is the second most diverse genera of lizards, currently with 257 species (Abdala and Quinteros 2014) and new species described almost every year, especially in the Andean and Patagonian regions of Chile
and Argentina (e.g. Avila et al. 2015; Esquerré et al. 2013; Troncoso-Palacios et al. 2015). Both morphological and molecular data support two main clades that are considered subgenera, roughly separated by the Andes: the Liolaemus (sensu stricto) subgenus or "Chilean" group and the Eulaemus subgenus or "Argentinean" group

[^0](Espinoza et al. 2004; Etheridge 1995; Laurent 1985; Schulte et al. 2000).

There is a group of Andean and Patagonian viviparous and saxicolous lizards within the Liolaemus (sensu stricto) subgenus known as the L. elongatus clade, which is part of a complex of clades known as the L. elongatus-kriegi complex (Cei 1975 1979). Research on this complex has supported four main clades: punmahuida, petrophilus, kriegi and elongatus (Avila et al. 2004 2012; Morando et al. 2003). The L. elongatus clade comprises species almost exclusively confined to rocky environments along the eastern slope of the Andes and south of the Mendoza River basin (Argentina), extending to the volcanic hills of Patagonia in Chubut Province, Argentina, and the Araucanía Region in Chile (Avila et al. 2015; Escobar-Huerta et al. 2015b; Minoli et al. 2013; Morando et al. 2003). Members of the L. elongatus clade are medium to large sized Liolaemus (maximum snout vent length = 107.8), long-tailed, with absent or reduced sexual dichromatism, viviparous, insectivorous, and almost exclusively saxicolous, with a high amount of midbody, ventral and dorsal scales (Abdala et al. 2010; Avila et al. 2015). Currently, this clade is comprised of nine species: L. antumalguen Avila et al. 2010, L. burmeisteri Avila et al. 2012, L. choique Abdala et al. 2010, L. crandalli Avila et al. 2015, L. elongatus Koslowsky 1896, L. janequeoae Troncoso-Palacios et al. 2016, L. Ionquimayensis Escobar-Huerta et al. 2015b, L. shitan Abdala et al. 2010, L. smaug Abdala et al. 2010; and possibly L. carlosgarini Esquerré et al. 2013 (fide Esquerré et al. 2013) and L. cristiani Núñez et al. 1991 (fide Medina et al. 2014); but recently L. Ionquimayensis and L. shitan were suggested as junior synonyms of L. elongatus in Avila et al. (2015) and TroncosoPalacios et al. (2016), respectively.

One of the most confusing issues in the Liolaemus elongatus clade is the identity of the true L. chillanensis Müller \& Hellmich 1932, a name that has been used indistinctly for two different species of Liolaemus that inhabit Termas de Chillán, Biobío Region, Chile. This species has been both included (Avila et al. 20102012 2015; Escobar-Huerta et al. 2015b; Medina et al. 2017; Torres-Pérez et al. 2009) and excluded from the L. elongatus clade (Lobo et al. 2010; TroncosoPalacios et al. 2015). One major problem is that the holotype of L. chillanensis is lost (Franzen and Glaw 2007). Here, we analyze the original description (Müller and Hellmich 1932), review several vouchers used by Torres-Pérez et al. (2009)
and several specimens from different collections, and provide an identification for the species that we conclude is the true L. chillanensis. We use both a molecular (our data and data from GenBank) and a morphological characterization. We also provide a description for the taxon previously confused with $L$. chillanensis.

## MATERIALS AND METHODS

## Morphological data and analyses

We collected specimens in the field by hand or noose. Specimens were dissected to extract a sample of liver/muscle for DNA extraction, and fixed in 99\% ethanol. These and all examined specimens ( $n=80$ ) are indicated in Appendix 1. Morphological characters were examined according to Etheridge (1995), Lobo (2005) and Avila et al. (2010 2012 2015). Body measurements were taken using a digital Vernier calliper ( 0.02 mm precision). Body measurements are provided as mean $\pm$ standard deviation. The stomach and intestinal contents of the individuals were observed under a binocular microscope for a preliminary description of the species' diet. Data for: L. choique and L. smaug were taken from Abdala et al. (2010); L. antumalguen were mainly taken from Avila et al. (2010), plus some specimens reviewed by us; L. burmeisteri were taken from Avila et al. (2012); L. crandalli were taken from Avila et al. (2015); L. lonquimayensis were taken from Escobar-Huerta et al. (2015b); and L. cristiani were taken from Núñez et al. (1991) and two specimens that we reviewed.

As a visualization and exploratory analysis, we performed a Principal Component Analysis (PCA) on the morphometric and meristic variables with the $R$ package FactoMineR (Lê et al. 2008). These analyses were performed in the R statistical environment version 3.2.3 (RDCT 2016). We used the residuals of a regression between the snoutvent length (SVL) and the following six variables: head length (distance between anterior edge of auditory meatus and tip of the snout), head width, head height, axilla-groin distance, arm length and foot length. Missing data from some measurements of some individuals were imputed using the imputePCA function from the MissMDA R package (Josse and Husson 2012).

We performed a nonparametric KruskalWallis test on the meristic variables with a Dunn test for post hoc comparisons (Dunn 1964) of
the following: midbody scales, dorsal scales (counted between the occiput and the level of the anterior border of the hind limbs), ventral scales, supralabial scales, infralabial scales and fourth toe lamellae; using data from the following species: L. antumalguen, L. carlosgarini, L. chillanensis, L. curis, L. elongatus, L. janequeoae, L. scorialis, the candidate species $L$. sp. Chillán and $L$. sp. 6.

We excluded species without a published dataset from the PCA and Kruskal-Wallis test, but we performed a diagnosis based on scale count and SVL ranges following the diagnosis previously published for the description of Liolaemus included in this work (Abdala et al. 2010; Avila et al. 2010 2012 2015; Escobar-Huerta et al. 2015b; Esquerré et al. 2013). Color pattern features were used as qualitative features of diagnosis for all species.

## Molecular laboratory procedures

Genomic DNA was extracted using a saltextraction protocol (slightly modified from Miller et al. 1988). We amplified the mitochondrial gene cytochrome b (Cyt-b) with Polymerase Chain Reaction (PCR) using the IguaCytob_F2 and IguaCytob_R2 primers (Corl et al. 2010), with the following thermocycle: denaturation at $94^{\circ} \mathrm{C}$ for 5 minutes, then cycle 35 times at $94^{\circ} \mathrm{C}$ for 30 seconds, $50^{\circ} \mathrm{C}$ for 30 seconds, $72^{\circ} \mathrm{C}$ for 1.5 minutes, and then a final hold at $72^{\circ} \mathrm{C}$ for 5 minutes. Sequencing reactions were done using a Big Dye Kit (Applied Biosystems, Foster City, CA). Sequencing was run on an ABI 3130xl Genetic Analyzer. Sequences were edited on Geneious 9.0.4 (Biomatters, Auckland, New Zealand, 2015).

## Phylogenetic reconstruction

Accession numbers of the Cyt-b mitochondrial loci sequences generated in this study and the sequences obtained from GenBank are indicated in Appendix 2. Sequences of the L. lonquimayensis type specimens ( $n=4$ ) were provided by G. Escobar-Huerta. The 164 nucleotide sequences involved in the analysis were aligned using MUSCLE (Edgar 2004). We used JModelTest v2.1.7 (Darriba et al. 2012; Guidon and Gascuel 2003) to select a substitution model (HKY + G), considering both BIC and AIC information criteria. We identified the non-redundant haplotypes of each terminal taxa using DnaSP v5.10 (Librado and Rozas 2009) and performed a Bayesian inference (BI) analyses with MrBayes v3.2.6 (Ronquist et al. 2012). Two independent analyses,
each consisting of two groups of four chains that run independently, were run for $10 \times 10^{6}$ generations, sample frequency $=1,000$. Priors were left by default. Phymaturus vociferator Pincheira-Donoso, 2004 was selected as out group because Phymaturus is the sister genus of Liolaemus (Schulte et al. 2000). The initial 25\% of the samples were discarded as burn-in when calculating the convergence diagnostic, assessed by examining values of average standard deviation of the Potential Scale Reduction Factor (PSRF $=1.000$ for all parameters; Gelmar and Rubin 1992) and the minimum and average Estimated Sample Size (ESS > 5000 for all parameters). Then, to estimate speciation, we used Bayesian implementation of the Poisson Tree Processes (bPTP, Zhang et al. 2013) for species delimitation using the non-ultrametric consensus tree obtained through the BI analyses described above, in the online server (http://species.h-its.org/) run with 50000 MCMC generations (all other parameters by default). Additionally, we generated a maximum clade credibility tree (MCC) using TreeAnnotator v1.8 (Drummond et al. 2012), from the posterior distribution of an MCMC Bayesian phylogenetic inference with BEASTv1.81 (Drummond and Rambaut 2007), which was run for $100 \times 10^{6}$ generations. Relative divergence times (branch lengths) were calculated using the logNormal relaxed clock in BEAST. We then used this ultrametric tree to run the Generalized Mixed Yule Coalescent model (GMYC, Pons et al. 2006). The two sequences of $L$. aff. chillanensis from Avila et al. (2015) were not included in the GMYC analysis.

## RESULTS

## Identification of the true Liolaemus chillanensis

Müller and Hellmich (1932) described $L$. monticola and included three subspecies, all these with allopatric distributions: L. monticola monticola, L. monticola chillanensis and L. monticola villaricensis. In the same publication, they referenced an additional L. monticola ssp. sympatric with L. monticola chillanensis, but did not describe it. Currently, these three described taxa are accepted at the species level (Abdala and Quinteros 2014; Lobo et al. 2010).

The identification of Liolaemus chillanensis is problematic because this species name has been used for two different species of Liolaemus that occur sympatrically in Termas de Chillán and
share two features: a fragmented vertebral line and a black lateral band. However, one of these Liolaemus species has an olive or bluish dorsal and ventral color, lacks precloacal pores and does not belong to the L. elongatus clade. The other does not have an olive or bluish dorsal and ventral color, its males have precloacal pores and it belongs to the L. elongatus clade. Unfortunately, the holotype of $L$. chillanensis, collected in Termas de Chillán at 1700 m , is lost (Franzen and Glaw 2007) and was not illustrated in the original description by Müller and Hellmich (1932). However, the authors did provide a black and white dorsal picture of one male paratype. Müller and Hellmich (1932) indicated that the color pattern of the holotype's dorsum is olive-brown ("Färbung der Oberfläche ein dunkles, leicht ins Oliv gehendes Rehbraun", p. 184) and the ventral color is olive ("Gesamte Unterseite olivgrün", p. 185). They do not describe the coloration of the male paratypes, but stated that the female paratypes have green or blue shades ("dunkel-bis hellblaugrün", Müller and Hellmich 1932:186). Later, Hellmich (1950 1952) stated that this species has a dark "gray-blue" dorsal and ventral color.

At this point it is clear that the true $L$. chillanensis is the lizard from Chillán with olive or bluish dorsal and ventral color. However, Müller and Hellmich (1932) point out that the holotype of L. chillanensis has four precloacal pores, although these are difficult to see ("schwer sichtbar", p. 184), and according to Hellmich (1950) the precloacal pores in L. chillanensis are extraordinarily difficult to see ("auBerordentlich schwer sichtbar", p. 147). Note that in the time of L. chillanensis description Müller and Hellmich (1932), the absence of precloacal pores in male Liolaemus species was unknown, being first reported by Cei and Scolaro (1982), so it is very likely that the mention of precloacal pores in the holotype of L. chillanensis was a mistake or that precloacal pores in $L$. chillanensis males are present at an extremely low frequency (we failed to find a topotype with them). Based on the original description (Müller and Hellmich 1932) and the posterior publications of one of the species authorities (Hellmich 1950 1952), we conclude that the true L. chillanensis is the lizard that occurs in Termas de Chillán (Chile) and that it can be differentiated by its olive or bluish dorsal and ventral coloration (Fig. 1; see more comparative features in Table 1). This has already been noted by other authors (DonosoBarros 1966; Pincheira-Donoso and Núñez 2005). In fact, a photograph of the first record of $L$.
chillanensis from Argentina (Avila et al. 2013:228) clearly displays the olive coloration.

In a recent phylogenetic work based on Cyt-b, Torres-Pérez et al. (2009) included both "monticola" taxa from Chillán: L. monticola chillanensis (currently L. chillanensis) and the taxon that they identified as the undescribed $L$. monticola ssp. However, we recently reviewed part of the vouchers used as L. chillanensis (MZUC 28249 and 28251, Fig. 2) and determined that these do not belong to L. chillanensis because they lack olive or bluish dorsal and ventral color and the male has precloacal pores. Thus, we conclude that these belong to an undescribed species. We also reviewed the vouchers MZUC 28257-59 and 28263, used by Torres-Pérez et al. (2009) as L. monticola ssp., and noted that they have olive or bluish dorsal and ventral color and the males lack precloacal pores (Fig. 3). To confirm the identification of these two taxa in Torres-Pérez et al. (2009), we sequenced Cyt-b for the two species from our own samples, the undescribed Liolaemus sp. and L. chillanensis. In our Cyt-b phylogeny, our sample of L. chillanensis is nested with the L. monticola ssp. from Torres-Pérez et al. (2009) and our undescribed species is nested with L. monticola chillanensis from Torres-Pérez et al. (2009), so the obvious conclusion is that TorresPérez et al. (2009) used the name "L. chillanensis" for the undescribed species from Termas de Chillán (= L. sp. Chillán) and used "L. monticola ssp." for L. chillanensis.

## Phylogenetic relationships

Our Bayesian inference found that neither Liolaemus chillanensis nor the undescribed taxon from "Termas de Chillán" (L. sp. Chillán) are closely related to L. monticola (50\% consensus tree) (Fig. 4); this was also found by Torres-Pérez et al. (2009). The inference also concluded that $L$. chillanensis is not part of the L. elongatus clade, but is instead sister to L. cristiani - a species that also features olive dorsal color and lacks precloacal pores - with strong support. We propose that the name "L. chillanensis clade" refer to this group in the future. Liolaemus sp. Chillán is recovered as part of the L. elongatus clade, with strong support. This clade is also comprised of $L$. antumalguen, L. burmeisteri, L. carlosgarini, L. choique, L. curis, L. leopardinus, L. elongatus, L. scorialis, L. shitan, L. smaug, two candidate species (Liolaemus sp. 6 and Liolaemus sp. 7) proposed by Morando et al. (2003) and Liolaemus sp. from Lircay, this last
referred as L. carlosgarini by Escobar-Huerta et al. (2015a). Liolaemus sp. 7 and L. antumalguen were not found to be reciprocally monophyletic (see below). The same occurs with $L$. shitan and $L$. elongatus, which raises doubts about the validity of the first (see Avila et al. 2015). Liolaemus sp. Chillán is closely related to $L$. antumalguen, $L$. sp. 6 and L. sp. 7 with strong support. On the other hand, L. aff. chillanensis by Avila et al. (2015), listed by Medina et al. (2017) as L. sp. 1, is closely related to L. scorialis, a species with type locality in Laja, 5 km NW from where the samples of $L$. aff. chillanensis were collected (Fig. 4).

The species delimitation through GMYC shows the presence of several species in the data
set (likelihood ratio $=12.08 ; P=0.002$ ). Liolaemus chillanensis was recovered as a full species, while eleven lineages were recognized in the $L$. elongatus clade (Fig. 5), L. sp. Chillán among them. In the MCC tree, $L$. sp. Chillán is sister to the lineage comprised of $L$. antumalguen $+L . s p$. 7, with L. sp. 6 basal to this clade. The GMYC model did not discriminate several species and candidate species as independent lineages: 1) $L$. antumalguen $+L$. sp. 7, therefore considered here as conspecifics, in fact Medina et al. (2017) were also unable to differentiate these two taxa at the genetic level, even with a larger data set. 2) L. elongatus $+L$. shitan $+L$. lonquimayensis, previously thought to be junior synonyms of $L$.


Fig. 1. Variation in the true Liolaemus chillanensis. (A-B) Male from Termas de Chillán (SSUC Re 708). (C) Male from near Aguas Calientes (SSUC Re 710). (D) Male from Termas de Chillán (SSUC Re 707). (E-F) Female from Termas de Chillán (SSUC Re 709).


Fig. 2. Some specimens previously misidentified as L. chillanensis in Torres-Pérez et al. (2009): (A-B) MZUC 28251 and (C-D) MZUC 28249. Here identified as $L$. sp. Chillán.

Table 1. Characteristics of Liolaemus antonietae sp. nov. and the species of the $L$. elongatus clade that occur near it, plus L. chillanensis. Juvenile specimens examined are excluded. $M=$ males; $F=$ females

|  | L. antonietae sp. nov. $(M=6, F=2)$ | L. antumalguen | L. burmeisteri | L. carlosgarini ( $M=6, F=11$ ) | L. chillanensis $(M=9, F=4)$ | L. elongatus $(M=3, F=3)$ | L. scorialis $(M=8, F=3)$ | L. smaug |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Max SVL (mm) | 77.6 | 107.8 | 85.2 | 68.8 | 85.8 | 73.7 | 69.9 | 71.3 |
| Midbody scales range | 86-98 | 72-82 | 70-81 | 80-95 | 82-92 | 76-88 | 76-90 | 73-80 |
| Ventral scales | 118-131 | 105-118 | 99-110 | 112-124 | 113-127 | 119-129 | 115-131 | 119-131 |
| Dorsal scales | 71-78 | 70-78 | 76-85 | 68-82 | 75-88 | 67-73 | 72-81 | 69-83 |
| Dorsal pattern | Fragmented vertebral line, dark paravertebral spots and lateral dark bands | Variable, from patternless to two dorsolateral series of black ocelli sometimes fused longitudinally | Light brown speckled with white spots, flanked by band of dark brown between axilla and groin, with few white spots | Marked or inconspicuous dark occipital band and lateral dark bands | Fragmented vertebral line, dark paravertebral spots and lateral dark bands | Vertebral and lateral bands | Marked dark occipital band and lateral dark bands | Dark occipital band with white dots in males, and lateral dark bands |
| Ventral melanism | Absent | Present | Absent | Absent | Absent | Absent | Absent | Absent |
| Head color | Light brown | Variable, from completely black to light-tan or ochre | Ochre | Light brown | Brown | Dark brown | Brown/Light brown | Gray to brown |
| Body color | Light brown | Light gray to ochre | Light brown/kaki | Yellowish brown or light brown | Olive-brown/ bluish brown | Ochre to almost black | Brown/Gray | Yellowish brown (males) or brown (females) |
| Tail rings | Marked/Weak | Absent | Weak | Marked/Weak | Absent/Weak | Marked/Weak | Marked | Absent |
| Precloacal pores in males | 3-5 | 3-4 | 0-5 | 0-3 | 0 | 4-5 | 3-4 | 3-4 |
| Behavior | Arboreal-saxicolous | Saxicolous | Saxicolous | Saxicolous | Saxicolous | Saxicolous | Saxicolous | Arenicolous |

elongatus in Avila et al. (2015) and TroncosoPalacios et al. (2016) and therefore considered here as part of the L. elongatus variation. 3) $L$. carlosgarini was not found to be monophyletic and instead forms part of two not closely related clades: $L$. smaug + L. choique + L. carlosgarini and $L$. carlosgarini $+L$. scorialis. Our sample of $L$. carlosgarini consists of the type specimens and several topotypes, among which it is not possible to differentiate two morphotypes and should all be assigned to L. carlosgarini according to the features provided by Esquerré et al. (2013). These results suggest hybridization or introgression in L. carlosgarini, as have been recorded for other Liolaemus (Morando et al. 2004 2007). 4) L. choique was not found to be an independent lineage (forming part of the $L$. smaug $+L$. choique + L. carlosgarini clade); therefore, its taxonomic
status should be evaluated in the future. Besides, our bPTP species delimitation analysis recognized thirteen different lineages in the L. elongatus clade, among them L. sp. Chillán as the fourth with the highest posterior probability (Fig. 6). Results only differ from GMYC in that part of L. carlosgarini was recovered as a full species, sister to L. scorialis; and part of $L$. aff. chillanensis was also recovered as a full species.

## Morphologic analyses results

The first three Principal Components (PCs) cumulatively account for $76.86 \%$ of the total variation (Table 2). PC1 is mainly explained by variation in head length, head height and head width (Fig. 7). PC2 mostly represents variation in arm length, foot length and AGD (Fig. 7). PC3


Fig. 3. Some misidentified specimens of Liolaemus chillanensis. Part of the vouchers used as "Liolaemus monticola ssp." in TorresPérez et al. (2009) from Termas de Chillán: (A-B) MZUC 28257, (C-D) MZUC 28259. Part of the specimens of "L. cristiani" from Termas de Chillán deposited in the MZUC collection (E-F).
mostly represents variation in AGD, head width and foot length (Fig. 7).

The PCA plots (Fig. 7) have 95\% confidence ellipses around the centroid of the species and illustrate the morphometric differences between the species included in the analysis. The PC1 vs PC2 graphic shows a partial overlap between Liolaemus sp. Chillán, L. chillanensis, L. carlosgarini, L. scorialis and $L$. elongatus, but there is no overlap between Liolaemus sp. Chillán and L. chillanensis in the PC1 vs PC3 graphic. Although Liolaemus sp . Chillán partially overlaps with $L$. carlosgarini in the PC1 vs PC3 graphic, the ellipses have different orientation. Liolaemus sp . Chillán, L. scorialis and
L. elongatus partially overlap in all PC analyses, but they are not sister species in our phylogenetic reconstruction. According to the mDNA phylogeny, Liolaemus sp. Chillán, L. antumalguen and L. sp. 6 are closely related, but show no overlap in the PC1 vs PC2 graphic; in the PC1 vs PC3 graphic only $L$. antumalguen and $L$. sp. 6 show partial overlap. Besides, Liolaemus sp. Chillán and L. janequeoae show no overlap in any of the PCs graphics.

A Kruskal-Wallis test on the meristic variables revealed significant differences in three of the six variables analysed: scales around midbody, dorsal scales and ventral scales (Table 3). Liolaemus sp. Chillán differs from L. antumalguen (closely


Fig. 4. Phylogenetic relationships using Bayesian inference ( $50 \%$ consensus tree) based on Cyt-b (HKY + G). In color: Liolaemus sp. Chillán (red, our data from Shangrila, Termas de Chillán and GenBank data from Termas de Chillán), L. chillanensis (blue, our data from Termas de Chillán and data from GenBank) and L. elongatus from Chubut, type locality (green). Red circle on the node denotes posterior probabilities $>0.95$. Scale shows the number of changes per site. Numbers between parentheses indicate the number of sequences of the collapsed nodes.


Fig. 5. Maximum Clade Credibility ultrametric tree (MCC) with species delimitation found through the Generalized Mixed Yule Coalescent model (GMYC). Lineages of the L. elongatus clade and L. chillanensis are in color. Red circle on the node denotes posterior probabilities $>0.95$. Scale shows the number of changes per site. Numbers between parentheses indicate the number of sequences of the collapsed nodes.

Table 2. Eigenvalues, the percentage of the total variance and the cumulative percentage of variance for PC1-3, and the correlation of each variable (residuals of a regression with the SVL)

|  | PCA |  |  |
| :--- | :---: | :---: | :---: |
| Eigenvalues | PC1 | PC2 | PC3 |
| Eigenvalue per Component | 2.41 | 1.25 | 0.94 |
| \% of var. | 40.29 | 20.86 | 15.69 |
| Cum. \% of var. | 40.29 | 61.16 | 76.86 |
| Eigenvectors | PC1 | PC2 | PC3 |
| Head Length (HL) | 0.885 | -0.220 | -0.081 |
| Head Height (HH) | 0.753 | 0.209 | 0.373 |
| Head Width (HW) | 0.790 | -0.133 | 0.153 |
| Axilla-Groin Distance (AGD) | -0.579 | 0.373 | 0.393 |
| Foot Length (FL) | 0.225 | 0.559 | -0.750 |
| Arm Length (AL) | 0.277 | 0.831 | 0.237 |



Fig. 6. Bayesian Poisson Tree Processes (bPTP) with highest posterior probability supported delimitation. Zero in the deep nodes denotes that no samples through MCMC recover all sequences as a single species. Values in the collapsed nodes denote the posterior probability to support each species. Scale shows the number of changes per site. Numbers between parentheses indicate the number of sequences of the collapsed nodes.

Table 3. Results of the Kruskal-Wallis test (H) with degrees of freedom (d.f.) among species of the $L$. elongatus clade and $L$. chillanensis and the post hoc comparison (Dunn's test). Only significant results are provided

|  | Kruskal-Wallis test |  |  | Dunn's test |
| :---: | :---: | :---: | :---: | :---: |
|  | H | $P$ | d.f. |  |
| Midbody scales | 62.15 | $<0.01$ | 8 | $L$. sp. Chillán $>L$. antumalguen, $L$. curis, $L$. sp. 6; $L$. janequeoae $>L$. antumalguen, L. curis, L. sp. 6; L. carlosgarini > L. antumalguen, L. curis, L. sp. 6 |
| Dorsal scales | 58.46 | $<0.01$ | 8 | L. chillanensis $>$ L. curis, L. elongatus, L. sp. 6; L. janequeoae $>$ L. curis, $L$. elongatus, L. sp. 6; L. scorialis > L. curis, L. sp. 6 |
| Ventral scales | 56.51 | $<0.01$ | 8 | $L . s p$. Chillán > L. sp. 6; L. chillanensis >L. sp. 6; L. elongatus > L. sp. 6; L. scorialis > L. sp. 6; L. janequeoae > L. antumalguen, L. curis, L. sp. 6 |

related in our phylogeny) in the number of midbody scales, but does not differ from L. carlosgarini or $L$. scorialis.

Based in this, we proposed that Liolaemus sp. Chillán need to be recognized as full species because the Cyt-b samples of it form a monophyletic lineage; L. sp. Chillán also can be statistically differentiated and shows no overlap in morphological space with regards to its most closely related taxa (L. antumalguen and L. sp. 6).

## Liolaemus antonietae sp. n.

(Figs. 2, 8)
urn:Isid:zoobank.org:pub:A4F4AEED-1AB0-48E7-88DDC2711FBACED2

2009 Liolaemus monticola chillanensis, Torres-Pérez, Méndez, Benavides, Moreno, Lamborot, Palma \& Ortiz. Biol. J. Linn. Soc., 96: 635.
2015 Liolaemus chillanensis, Escobar-Huerta, Santibáñez \& Ortiz. Gayana, 79: 95.
2016 Liolaemus sp. Chillán Troncoso-Palacios, Díaz, Puas, Riveros-Riffo \& Elorza. Zookeys, 632:127
2017 Liolaemus chillanensis Medina, Avila, Sites \& Morando. J. Zool. Syst. Evol. Res., 55: 238.

Material examined: Holotype: SSUC Re 697, male (Fig. 8A-B). Termas de Chillán, Biobío Region, Chile ( $36^{\circ} 54^{\prime} \mathrm{S}, 71^{\circ} 24^{\prime} \mathrm{W}, 1766 \mathrm{~m}$ ). Collected by J. Troncoso-Palacios, H. Díaz and F.A. Urra. January 9, 2014.

Paratypes: SSUC Re 695-96, 698-99 (Fig. 8C-G), male, female, male, female, respectively. Same data as the holotype. SSUC Re 700, male. Laguna del Huemul, Shangrila, Biobío Region, Chile ( $36^{\circ} 52^{\prime} \mathrm{S}$, $71^{\circ} 28^{\prime} \mathrm{W}$, 1955 m ). Collected by F.A. Urra. December 2014.

Specimens used in morphological and color variation: MZUC 38086 and 38090, males. Termas de Chillán.

Specimens used only in color variation: MZUC 28251, 28254 and 28569, three males; and MZUC 28249 one female. Termas de Chillán.

Diagnosis: Here we provide a diagnosis in regards of all species in the L. elongatus clade plus Liolaemus chillanensis - previously confused with L. antonietae - and L. monticola - which has a similar color pattern. Liolaemus antonietae is a medium size Liolaemus (max. SVL $=77.6 \mathrm{~mm}$ ), with many midbody scales ( $86-98 ; 91.5 \pm 3.8$ ), lateral dark


Fig. 7. Principal Component Analysis (PCA) plots. On the left panels, the individuals are colored according to their species as shown on the legend on the top left corner. Ellipses represent the $95 \%$ confidence interval around the centroid for each species. On each axis, the PC is labelled according to its number and the percentage of the total variance that PC explains. On the right panels are the variables graphs, which illustrate the contribution of each variable to the construction of the axes.
band, absence of dorsal melanism or black ventral color. The males have precloacal pores and it is the only member of the L. elongatus clade that can exhibit arboreal behavior.

Liolaemus antonietae differs from L. chillanensis because this species has olivebrown or bluish-brown dorsal color, olive or bluish ventral color and lacks precloacal pores, whereas L. antonietae never has olive or bluish coloration and the males have precloacal pores. Liolaemus antonietae resembles L. monticola in that both species have a brown dorsal color and dark lateral band. However, phylogenetic evidence shows that L. monticola does not belong in the L. elongatuskriegi complex (Fig. 4). Moreover, L. monticola is smaller (max. SVL $=65.6 \mathrm{~mm}$ ) and has fewer midbody scales (54-66) than L. antonietae (max. SVL = 77.6 mm ; midbody scales $=86-98$ ).

Liolaemus antonietae is closely related to $L$. antumalguen, but is smaller (max. SVL $=77.6 \mathrm{~mm}$ vs. max. $\mathrm{SVL}=107.8 \mathrm{~mm}$ ) and has more midbody scales (86-98 vs. 72-82, Table 3). Moreover, L. antumalguen has black ventral coloration, and several specimens have black transversal dorsal spots or a very melanic dorsal coloration, whereas L. antonietae has whitish or gray ventral coloration and totally lacks black transversal dorsal spots.

Liolaemus antonietae has more midbody scales (86-98 vs 76-90), and is larger (max. SVL = 77.6 mm vs max. SVL $=69.9 \mathrm{~mm}$ ) than $L$. scorialis. Dorsal coloration in L. scorialis is noticeably darker than in L. antonietae, with a marked dark occipital band, which is absent in $L$. antonietae.

Liolaemus antonietae differs from L. carlosgarini because this latter species features a yellowish brown dorsal coloration and a conspicuous dark occipital band, traits absent in L. antonietae. All males of L. antonietae have precloacal pores, whereas only $50 \%$ of $L$. carlosgarini males have precloacal pores.

Liolaemus curis has a yellowish dorsal coloration accompanied by black transversal dorsal spots or an overall melanic dorsal coloration, and less midbody scales $(68-76)$ than L. antonietae (Table 3).

Liolaemus antonietae is smaller (max. SVL $=77.6 \mathrm{~mm}$ vs. max. $\mathrm{SVL}=90.7 \mathrm{~mm}$ ) than $L$. choique, and although we have no data of $L$. choique for statistical analysis, the midbody scale count ranges show almost no overlap (86-98 vs. 74-88, Table 1). Also, L. choique has yellowish or black dorsal coloration, whereas L. antonietae has light brown dorsal coloration.

Liolaemus antonietae differs from L. elongatus
because this last features a marked occipital and lateral bands or dorsal melanism, whereas only some specimens of $L$. antonietae have an inconspicuous occipital band and no specimen has dorsal melanism.

The dorsal pattern of Liolaemus antonietae resembles the pattern of $L$. smaug. However, males of $L$. antonietae have light brown dorsum, whereas males of $L$. smaug have a bright golden yellow dorsal color. Females and males of $L$. antonietae have white dots on the dorsum, a trait only found on males of $L$. smaug.

Liolaemus antonietae differs from $L$. janequeoae, because the latter species does not feature any dorsal pattern apart of few black or white dots and is smaller than $L$. antonietae (max. SVL 66.9 vs 77.6 mm ).

Liolaemus antonietae differs from L. leopardinus, because it lacks the leopard-like dorsal spots on dorsum and tail present in $L$. leopardinus.

Liolaemus antonietae is smaller (max. SVL = 77.6 mm ) than L. crandalli (max. SVL $=93.4 \mathrm{~mm}$ ), but both exhibit a similar color pattern. However, L. crandalli has blackish dorsal head coloration, which is light brown in L. antonietae. According to the phylogenetic analysis of Avila et al. (2015) this species is the sister taxon of $L$. smaug $+L$. choique, whereas in our phylogeny $L$. antonietae is not closely related to L. smaug or L. choique.

Although we have no scale count data of $L$. burmeisteri to perform a statistical analysis, the midbody (70-81) and ventral scale count ranges (99-110) show no overlap with L. antonietae (8698, 118-131, respectively). Additionally, almost all specimens of $L$. burmeisteri lack a dorsal pattern, whereas $L$. antonietae always features a dorsal pattern (vertebral line, paravertebral dark spots and white dots).

Description of the holotype: Male. SVL = 61.2 mm . Horizontal diameter of the eye: 2.88 mm. Subocular length: 3.8 mm . Length of the fourth supralabial: 2.5 mm . Head length (from the anterior border of the auditory meatus to the tip of the snout): 14.3 mm . Head height (at the level of ear openings ): 6.1 mm . Head width (distance between the two ear openings): 11.8 mm . Neck width: 11.7 mm . Interorbital distance: 6.3 mm . Eareye distance: 4.8 mm . Internasal scales distance: 1.8 mm . Ear width: 1.4 mm . Ear height: 1.8 mm . Axilla-groin distance: 24.4 mm . Body width: 17.2 mm . Forelimb length: 23.2 mm . Hindlimb length: 38.6 mm . Tail length (not autotomized): 99.2 mm , with relation tail length/SVL $=1.6$. Rostral scale,


Fig. 8. Liolaemus antonietae sp. nov., all from type locality. (A-B) Holotype, male (SSUC Re 697), (C-E) Paratypes males (SSUC Re 698 and 695) and (F-G) Paratype female (SSUC Re 699).
wider ( 2.8 mm ) than high ( 1.1 mm ).
Two postrostrals. Four internasals. Interparietal is pentagonal, with a small central spot marking the position of the parietal eye. The interparietal is smaller than the parietals, and is surrounded by six scales. Nine scales between the interparietal and rostral. Seventeen scales between the occiput and the rostral (Hellmich index; Lobo, 2005). Orbital semicircles are complete and formed by thirteen scales on both sides. Five supraoculars on the right side and four on the left. Six superciliary scales. Frontal area divided into five scales, from back to front: 2, 1 and 2. Preocular separated from the lorilabials by one loreal scale. Two scales between nasal and canthal. Nasal in contact with the rostral, surrounded by six scales (excluding the rostral). One row of lorilabials between the supralabials and the subocular. Six supralabials, the fourth is curved upward without contacting the subocular. Five infralabials scales. Pentagonal mental scale, in contact with four scales. Five pairs of postmental shields, the second is separated by two scales. Temporal slightly keeled, subimbricated. Nine temporal scales between the level of superciliary scales and the rictal level. Four scales on the anterior edge of the ear, slightly projected, which do not cover the auditory meatus. Differentiated auricular scale, which is narrow and elongated. Forty-six gular scales between the auditory meatuses. The lateral neck fold is " $Y$ " shaped. There is a ventrolateral fold running from the axilla to the groin. Midbody scales: 90. Dorsal scales are lanceolated, subimbricated, keeled (without mucrons) and with interstitial granules. Dorsal smaller than the ventrals. Dorsal scales: 74. Ventral scales are rounded, smooth, imbricated and without interstitial granules. Ventral scales: 118. Four precloacal pores. Supra-femoral scales are lanceolated, imbricated and smooth or slightly keeled. Infra-femoral scales are lanceolated or rhomboidal, smooth and imbricated. Supraantebrachials scales are lanceolated, imbricated and keeled. Infra-antebrachials are rounded or rhomboidal, imbricated and smooth. Dorsal scales of the first third of the tail are rhomboidal, imbricated, keeled and mucronated. Ventral scales the first third of the tail are lanceolated, smooth and imbricated. Lamellae of the fingers: I: 10, II: 14, III: 20, IV: 23 and V : 12. Lamellae of the toes: $\mathrm{I}:$ 10, II: 18, III: 22, VI: 29 and V: 19.

Color of the holotype in life: The head has a light brown dorsal color, with few dispersed dark brown spots. The head has a similar shade as the
dorsum. The subocular is whitish with a vertical black line at the posterior end. The cheeks are whitish. The temporal zone is light brown with a dark brown stripe on the middle. The dorsum is light brown. There is a fragmented vertebral line. There are 12 series of black spots on the paravertebral fields, running from the occiput to the base of the tail. The flanks are light brown with a black lateral band that goes from the upper portion of the shoulder to the groin. There are several white dots dispersed on this black lateral band. There are white dots dispersed between the black lateral band and the paravertebral fields. The area between the belly and the dark band on the flank is yellowish with dispersed white dots. The limbs are brown, with dispersed black dots on the fore limbs and dispersed black and whitish dots on the hind limbs. The tail has a light brown dorsal coloration, with dark rings. Ventrally, the throat, the belly, the limbs and the tail are light grey, with some whitish scales dispersed. Inconspicuous yellowish color on the cloaca and thighs. Precloacal pores are orange.

Variation: In six males (including holotype): SVL: 61.2-77.6 mm. Axilla-groin distance: 24.429.3 mm . Head length: 14.3-18.8 mm. Head width: 11.7-15.3 mm. Head height: 6.1-9.6 mm. Leg length: 38.6-45.7 mm. Arm length: 23.226.3 mm . Foot length: 18.9-22.5 mm. Tail length: 99.2-120.0 mm (two specimens, autotomized in the others). Tail length/SVL = 1.6. In two females: SVL: 56.6-70.5 mm. Axilla-groin distance: 24.931.8 mm . Head length: 13.4-15.4 mm. Head width: 10.5-12.2 mm. Head height: 6.3-7.3 mm. Leg length: 36.9-38.7 mm. Arm length: 22.024.7 mm . Foot length: 18.8-20.2 mm. Tail length: 99.0-110.0 mm (no autotomized), with relation tail length/SVL = 1.6-1.8.

The scalation variation in Liolaemus antonietae is as follows. Midbody scales: 86-98 ( $91.5 \pm 3.8$ ). Dorsal scales: 71-78 (74.4 $\pm 2.3$ ). Ventral scales 118-131 (123.6 $\pm 5.4$ ). Fourth finger lamellae: 21-24 (23.0 $\pm 1.1$ ). Fourth toe lamellae: 28-30 (28.8 $\pm 0.8$ ). Supralabial scales: 6-7 (6.2 $\pm$ 0.4 ), the fourth curved upward. Infralabial scales: 4-5 (4.7 $\pm 0.5)$. Interparietal scale pentagonal or hexagonal, bordered by $5-8$ scales ( $5.8 \pm 1.2$ ). Nasal and rostral always in contact. Precloacal pores in males: 3-5. Precloacal pores are absent in females.

Males and females have a similar color patterns to the holotype, with small variations. Some specimens have no distinguishable vertebral line, while others have an inconspicuous occipital
band. Some specimens have diffuse dark rings on the dorsal surface of the tail. In specimens with regenerated tail, there is a vertebral line on the regenerated zone. Some males have no yellowish coloration on the cloaca and thighs and have whitish color on the ventral surface of the throat, belly, limbs and the tail, with inconspicuous gray spots dispersed. Females have marked dark spots on the throat and belly and the white dorsal dots are less abundant than males, even totally absent in one female.

Distribution and natural history: Known from two localities in the Biobío Region, Chile: Termas de Chillán (type locality, $36^{\circ} 54^{\prime} \mathrm{S}, 71^{\circ} 24^{\prime} \mathrm{W}, 1,766 \mathrm{~m}$, Fig. 9) and Laguna del Huemul, Shangrila ( $36^{\circ} 52^{\prime} \mathrm{S}$, $71^{\circ} 28^{\prime} \mathrm{W}, 1,955 \mathrm{~m}$ ), 8 km NW from the type locality. Unlike other species of the L. elongatus clade, which are mainly saxicolous (Morando et al. 2003; Avila et al. 2015), L. antonietae is mostly arboreal, which was seen basking on fallen trunks and on trees up approximately 4 m high. Few specimens were observed basking on rocks. Furthermore, we observed a very peculiar gregarious behavior, in which 18 specimens were observed coming out of the surrounding vegetation and entering to the same hollow of a stand tree (Nothofagus sp.) at sunset, in a time frame no longer than an hour. The vegetation in the type locality is dominated by Chusquea sp. and Nothofagus sp. It is an abundant lizard. At the date of capture (January), one female had three embryos and the other had several small oocytes. No remains were found in the stomach, but remnants of insects and plants along with parasitic nematodes were observed inside the intestine. In both localities, L. antonietae was found in syntopy with snake Tachymenis chilensis and lizards L. chillanensis, L. septentrionalis and L. tenuis, being recently recorded also Phymaturus vociferator (Urra et al. 2017).

Etymology: This species is named after Antonieta Labra Lillo, a prominent Chilean herpetologist who has made significant contributions to the eco-physiology and behavior of lizards, especially to the study of the influence of chemical signals on their behavior. She has also edited the book "Herpetología de Chile", which had a great impact on the Chilean herpetology. We propose the common name "Antonieta's lizard" in English and "Lagarto de Antonieta" in Spanish. JTP gratefully thanks AL for years of teaching and collaboration.

## DISCUSSION

We describe a new species of Liolaemus belonging to the L. elongatus group, Liolaemus antonietae, which was previously identified as L. chillanensis by Torres-Pérez et al. (2009) and followed by Escobar-Huerta et al. (2015b). The species was probably misidentified because the definition of $L$. chillanensis itself was ambiguous. The description was written in German (Müller and Hellmich 1932), while most researchers who worked on this species have been Spanish speakers. It is also difficult to obtain this publication (to our knowledge there is no official digital copy available). Moreover, the holotype of $L$. chillanensis is lost (Franzen and Glaw 2007). Here we provide several color pictures of $L$. chillanensis in life, showing the variation in color, and highlight the features that diagnose it, based on the distinctive characteristics that Müller and Hellmich (1932) describe for the holotype. Additionally, we add a Cyt-b sequence from L. chillanensis to GenBank and we note that the sequences from GenBank currently assigned to L. monticola ssp. (AY850623, AY850624, AY730670, AY529904, AY850625, AY529903) are indeed assignable to $L$. chillanensis.

The identity of several specimens assigned to $L$. chillanensis has been confusing. Avila et al. (2010 2012) used a Cyt-b sequence of "L. chillanensis" (voucher LJAMM-CNP 3434) without locality data, but according to Medina et al. (2017), this specimen was collected in Laguna del Maule, Maule Region, Chile, and is assignable to $L$. carlosgarini. Recently, Troncoso-Palacios et al. (2015) made a mistake regarding this, because they point out that the "L. chillanensis" Cyt-b sequence used by Avila et al. (2010 2012) was generated by Torres-Pérez et al. (2009), which is incorrect because both works used different vouchers. Additionally, Avila et al. (2015) used a Cyt-b sequence of L. aff. chillanensis (vouchers LJAMM-CNP 14027 and 14029) from 14.2 km W Refugio del Laja and assigned it to the $L$. elongatus-kriegi complex, a result contrary to that obtained by us for $L$. chillanensis. One species in the L. elongatus-kriegi complex, L. scorialis, was described recently from Laja by Troncoso-Palacios et al. (2015) and it is possible that this publication was not included as background source of data in Avila et al. (2015). In fact, both sequences of $L$. aff. chillanensis were placed in the same node as L. scorialis in our BI analysis (Fig. 4) and bPTP analysis delimited $L$. scorialis and one sequence


Fig. 9. Distributional map for Liolaemus antonietae sp. nov. along with geographically proximate species of the L. elongatus clade. Stars: Liolaemus antonietae sp. nov (red = Termas de Chillán, type locality; white = Shangrila). Green pentagon: $L . \operatorname{smaug}(1=$ near Las Leñas, $2=$ between Las Loicas and Peteroa Volcano, $3=$ near Las Loicas). Lilac cross: L. carlosgarini $(1=$ Maule Lagoon, $2=$ Lircay). Black diamond: L. choique (Paso el Choique). Gray square: L. antumalguen (1= Domuyo volcano, 2= Tromen Volcano). Pink haxagon: L. burmeisteri (Caepe Malal). White asterisk: L. crandalli (Auca Mahuida Volcano). Green triangle: L. scorialis (1 = Laja Lagoon, 2 = La Mula Lagoon). Orange circle: L. elongatus (formerly L. lonquimayensis, Lonquimay Volcano). Brown circle: L. elongatus used for morphology and DNA analyses (Llaima volcano). Yellow circles: L. elongatus used for DNA analyses ( $1=$ Pampa de Lonco Luan, 2 = Primeros Pinos, 3 = Portal La Atravesada, 4 = Laguna Blanca, $5=$ near Ingeniero Jacobacci, $6=$ San Carlos de Bariloche, 7 $=$ Ojo de Agua, $8=$ El Maiten, $9=$ Esquel, $10=$ Tecka, $11=$ Gobernador Costa and $12=$ Los Manantiales). Blue circles: $L$. shitan ( $1=$ Estancia Piedras Blancas, type locality, 2 = near San Antonio del Cuy). Pink inverted triangle: L. janequeoae (Tolhuaca). Blue ellipse: $L$. sp. 6 (Copahue Volcano).
of $L$. aff. chillanensis as the same species (Fig. 6). This is congruent with Medina et al. (2017), since they list LJAMM-CNP 14027 and 14029 as L. sp. 1 and include " 12 taxa... one being described (L. sp. 1, Esquerré, personal communication)" in their L. elongatus clade analysis (p. 239). In fact, Troncoso-Palacios, Díaz, Esquerré and Urra, described L. scorialis (Troncoso-Palacios et al. 2015) from the Laja surroundings.

Medina et al. (2017) found low Cyt-b distance between the taxon that they refer as " $L$. chillanensis" (here described as L. antonietae) and the lineage composed of $L$. antumalguen + L. sp. 7, but stated that both lineages were recover as full species in the species delimitation analysis. Moreover, Medina et al. (2017) shows in the "Supporting information Table 2" that L. antumalguen + L. sp. 7 do not share Cyt-b haplotypes with "L. chillanensis". Congruent with this, our species delimitation analyses (GMYC and bPTP) found $L$. antonietae (referred as $L$. chillanensis in Medina et al. 2017) as a lineage at the species level, monophyletic in regards to the most closely related lineage ( $L$. antumalguen $+L . s p .7)$. Additionally, the PCA shows that $L$. antonietae and $L$. antumalguen do not share the same morphological space; some meristic variables show statistical differentiation and striking differences in coloration were found. All these support the recognition of this taxon as an independently evolving lineage but the low Cyt-b distance suggests that $L$. antonietae is a young species.

Acknowledgements: This work and the new species name have been registered with ZooBank under urn:Isid:zoobank.org:pub:A4F4AEED-1AB0-48E7-88DD-C2711FBACED2. We thank P. Zavala (Pontificia Universidad de Católica de Chile) for allowing us to review and deposit material into the collection under his care. We thank the following colleagues and institutions for allowing us to review specimens: H. Núñez (Museo Nacional de Historia Natural), J.N. Artigas (Museo de Zoología de la Universidad de Concepción) and M. Lamborot (Colección del Laboratorio de Citología de la Universidad de Chile); M. Morando, F. Breitman and C. Aguilar for providing use useful information on PCR procedures and primers; I. G. Brennan for extensive help in the lab; F. Ferri, D. Vieites, F. Torres-Pérez, L.J. Avila and J.S. Keogh for their comments on the early version of the manuscript; F. Ferri, C.S. Abdala and A. Laspiur for sending literature; G. Feuerhake for providing photographs.
J. Troncoso-Palacios thanks M. Penna for his support and L. Rodríguez for her support in the field. D. Esquerré was supported by a Becas Chile-Conicyt Scholarship. F.A. Urra is supported by FONDECYT postdoctoral fellowship \#3170813. The map (Fig. 9) was made with ArcGis 10.5.1 free trial. We thank the Servicio Agrícola y Ganadero (SAG) for collecting permits ( $\mathrm{N}^{\circ} 4468$ ).

Authors' contributions: JTP conceived and designed the study, drafted the manuscript (except the background section), collected specimens and morphological data, perform morphological and phylogenetic analyses and made some figures. DE drafted the background section, perform DNA laboratory procedures and perform morphological and phylogenetic analyses. FAU and HAD collected specimens and made some figures. CCP collected specimens. MSR collected specimens and morphological data. All authors contributed with the corrections.

Competing interests: JTP, DE, FAU, HAD, CCP and MSR declare that they have no conflict of interest.

Availability of data and materials: Sequences generated in the study have been deposited in GenBank (see Appendix 2).

Consent for publication: The authors give their consent to publish this paper.

Ethics approval consent to participate: Not applicable.

## REFERENCES

Abdala CS, Quinteros AS. 2014. Los últimos 30 años de estudios de la familia de lagartijas más diversa de Argentina. Actualización taxonómica y sistemática de Liolaemidae. Cuadernos de Herpetología 28:55-82.
Abdala CS, Quinteros AS, Scrocchi GJ, Stazzonelli JC. 2010. Three new species of the Liolaemus elongatus group (Iguania: Liolaemidae) from Argentina. Cuadernos de Herpetología 24:93-109.
Avila LJ, Fulvio-Pérez CH, Medina CD, Sites JW Jr, Morando M. 2012. A new species of lizard of the Liolaemus elongatus clade (Reptilia: Iguania: Liolaemini) from Curi Leuvu River Valley, northern Patagonia, Neuquén, Argentina. Zootaxa 3325:37-52.
Avila LJ, Martínez LE, Morando M. 2013. Checklist of lizards and amphisbaenians of Argentina: an update. Zootaxa 3616:201-238. doi:http://dx.doi.org/10.11646/ zootaxa.3616.3.1.
Avila LJ, Medina CD, Fulvio-Pérez CH, Sites JW Jr, Morando
M. 2015. Molecular phylogenetic relationships of the lizard clade Liolaemus elongatus (Iguania: Liolaemini) with the description of a new species from an isolated volcanic peak in northern Patagonia. Zootaxa 3947:67-84. doi:10.11646/zootaxa.3947.1.4.
Avila LJ, Morando M, Pérez CHF, Sites JW Jr. 2004. Phylogenetic relationships of lizards of the Liolaemus petrophilus group (Squamata, Liolaemidae), with description of two new species from Western Argentina. Herpetologica 60:187-203.
Avila LJ, Morando M, Pérez DR, Sites JW Jr. 2010. A new species of the Liolaemus elongatus clade (Reptilia: Iguania: Liolaemini) from Cordillera del Viento, northwestern Patagonia, Neuquén, Argentina. Zootaxa 2667:28-42.
Cei JM. 1975. Herpetología patagónica. X. El conjunto evolutivo de Liolaemus elongatus: análisis serológico. Physis 34:203-208.
Cei JM. 1979. The Patagonian Herpetofauna. In: Duellman WE (ed) The South American Herpetofauna: Its origin, evolution, and dispersal, Museum of Natural History, University of Kansas, Lawrence.
Cei JM, Scolaro JA. 1982. A new species of the Patagonian genus Vilcunia, with remarks on its morphology, ecology and distribution. J Herpetol 16:354-363.
Corl A, Davis AR, Kuchta SR, Comendant T, Sinervo B. 2010. Alternative mating strategies and the evolution of sexual size dimoprhism in the side-blotched lizard, Uta stansburiana: a population-level comparative analysis. Evolution 64:79-96. doi:10.1111/j.1558-5646.2009.00791. x .
Darriba D, Taboada GL, Doallo R, Posada D. 2012. jModelTest 2: More models, new heuristics and parallel computing. Nature Methods 9:772. doi:10.1038/nmeth. 2109.
Donoso-Barros R. 1966. Reptiles de Chile. Ediciones de la Universidad de Chile, Santiago.
Dunn OJ. 1964. Multiple comparisons using rank sums. Technometrics 6:241-252.
Drummond AJ, Rambaut A. 2007. BEAST: Bayesian evolutionary analysis by sampling trees. BMC Evol Biol 7:214.
Drummond A, Suchard MA, Xie D, Rambaut A. 2012. Bayesian Phylogenetics with Beauti and the Beast 1.7. Mol Biol Evol 29:1969-1973.
Edgar RC. 2004. MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Res 32:1792-1797. doi:10.1093/nar/gkh340.
Escobar-Huerta G, Ormazabál GA, Carvajal RI, Ortiz JC. 2015a. Ampliación del área de distribución de Liolaemus carlosgarini (Esquerré, Núñez \& Scolaro, 2013) en la Reserva Nacional Altos de Lircay (Chile). Cuadernos de Herpetología 29:155-156.
Escobar-Huerta G, Santibáñez J, Ortiz JC. 2015b. Liolaemus lonquimayensis (Squamata: Liolaemidae), a new lizard species for Chile without precloacal pores. Gayana 79:94101.

Espinoza RE, Wiens JJ, Tracy CR. 2004. Recurrent evolution of herbivory in small, cold-climate lizards: Breaking the ecophysiological rules or reptilian herbivory. P Natl Acad Sci USA 101:16819-16824. doi:10.1073/ pnas. 0401226101.
Esquerré D, Núñez H, Scolaro JA. 2013. Liolaemus carlosgarini and Liolaemus riodamas (Squamata: Liolaemidae), two new species of lizards lacking precloacal pores, from

Andean areas of central Chile. Zootaxa 3619:428-452.
Etheridge R. 1995. Redescription of Ctenoblepharys adspersa Tschudi, 1845, and the taxonomy of Liolaeminae (Reptilia: Squamata: Tropiduridae). Am Mus Novit 3142:1-34.
Franzen M, Glaw F. 2007. Type catalogue of reptiles in the Zoologische Staatssammlung München. Spixiana 30:201274.

Gelman A, Rubin DB. 1992. Inference from iterative simulation using multiple sequences. Statistical Science 7:457-511.
Guindon S, Gascuel O. 2003. A simple, fast and accurate method to estimate large phylogenies by maximumlikelihood. Syst Biol 52:696-704.
Hellmich W. 1950. Beiträge zur kenntnis der herpetofauna Chiles XIII. Die eidechsen der ausbeute Schröder (gattung Liolaemus, Iguan.). Veröffentlichungen der Zoologischen Staatssammlung München 1:129-194.
Hellmich W. 1952. Ensayo de una clave para las especies chilenas del género Liolaemus. Investigaciones Zoológicas Chilenas 1:10-14.
Josse J, Husson F. 2012. Handling missing values in exploratory multivariate data analysis methods. Journal de la Société Française de Statistique 153:79-99.
Koslowsky J. 1896. Sobre algunos reptiles de Patagonia y otras regiones argentinas. Revista del Museo de La Plata 7:447-457.
Laurent RF. 1985. Segunda contribución al conocimiento de la estructura taxonómica del género Liolaemus Wiegmann (Iguanidae). Cuadernos de Herpetología 1:1-37.
Lê S, Josse J, Husson F. 2008. FactoMineR: An R Package for multivariate analysis. J Stat Softw 25:1-18.
Librado P, Rozas J. 2009. DnaSP v5: A software for comprehensive analysis of DNA polymorphism data. Bioinformatics 25:1451-1452.
Lobo F. 2005. Las relaciones filogenéticas dentro del grupo chiliensis (Iguania: Liolaemidae: Liolaemus): sumando nuevos caracteres y taxones. Acta Zoológica Lilloana 49:67-89.
Lobo F, Espinoza RE, Quinteros S. 2010. A critical review and systematic discussion of recent classification proposals for liolaemid lizards. Zootaxa 2549:1-30.
Medina CD, Avila LJ, Sites JW Jr, Morando M. 2014. Multilocus phylogeography of the Patagonian lizard complex Liolaemus kriegi (Iguania: Liolaemini). Biol J Linn Soc 113:256-269. doi:10.1111/bij. 12285.
Medina CD, Avila LJ, Sites JW Jr, Morando M. 2017. Phylogeographic history of Patagonian lizards of the Liolaemus elongatus complex (Iguania: Liolaemini) based on mitochondrial and nuclear DNA sequences. J Zool Syst Evol Res 55:238-249. doi:10.1111/jzs. 12163.
Miller SA, Dykes DD, Polesky HF. 1988. A simple salting out procedure for extracting DNA from human nucleated cells. Nucleic Acids Res 16:1215.
Minoli I, Medina CD, Frutos N, Morando M, Avila LJ. 2013. A revised geographical range for Liolaemus elongatus Koslowsky, 1896 (Squamata: Liolaemini) in Argentina: review of reported and new-data based distribution with new localities. Acta Herpetol 8:159-162.
Morando M, Avila LJ, Sites JW Jr. 2003. Sampling strategies for delimiting species: genes, individuals, and populations in the Liolaemus elongatus-kriegi complex (Squamata: Liolaemidae) in Andean-Patagonian South America. Syst Biol 52:159-185.
Morando M, Avila LJ, Baker J, Sites JW Jr. 2004. Phylogeny and phylogeography of the Liolaemus darwinii complex
(Squamata: Liolaemidae): Evidence for introgression and incomplete lineage sorting. Evolution 58:842-861.
Morando M, Avila LJ, Turner CR, Sites JW Jr. 2007. Molecular evidence for a species complex in the patagonian lizard Liolaemus bibronii and phylogeography of the closely related Liolaemus gracilis (Squamata: Liolaemini). Mol Phylogenet Evol 43:952-973.
Müller L, Hellmich W. 1932. Beiträge zur Kenntnis der Herpetofauna Chiles. IV. Liolaemus monticola, ein weiterer neuer Rassenkries aus den Hochanden Chiles. Zool Anz 99:177-192.
Núñez H, Navarro J, Loyola J. 1991. Liolaemus maldonadae y Liolaemus cristiani, dos especies nuevas de lagartijas para Chile (Reptilia, Squamata). Boletín del Museo Nacional de Historia Natural de Chile 42:79-88.
Pincheira-Donoso D, Núñez H. 2005. Las especies chilenas del género Liolaemus Wiegmann, 1834 (Iguania: Tropiduridae: Liolaeminae): Taxonomía, sistemática y evolución. Publicación Ocasional del Museo Nacional de Historia Natural de Chile 59:1-486.
Pons J, Barraclough TG, Gomez-Zurita J, Cardoso A, Duran DP, Hazell S, Kamoun S, Sumlin WD, Vogler AP. 2006. Sequence-based species delimitation for the DNA taxonomy of undescribed insects. Syst Biol 55:595-609.
R Development Core Team. 2016. R: A language and environment for statistical computing [Online]. Vienna, Austria: R Foundation for Statistical Computing. Available from: http://www.R-project.org. Retrieved 11 November 2014.

Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Höhna S, Larget B, Liu L, Suchard MA, Huelsenbeck JP. 2012. MrBayes 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space. Syst Biol 61:539-542.
Schulte JA II, Macey JR, Espinoza RE, Larson A. 2000. Phylogenetic relationships in the iguanid lizard genus Liolaemus: multiple origins of viviparous reproduction and evidence for recurring Andean vicariance and dispersal. Biol J Linn Soc 69:75-120.
Torres-Pérez F, Méndez MA, Benavides E, Moreno RA, Lamborot M, Palma RE, Ortiz JC. 2009. Systematics and evolutionary relationships of the mountain lizard Liolaemus monticola (Liolaemini): how morphological and molecular evidence contributes to reveal hidden species diversity. Biol J Linn Soc 96:635-650. doi:10.1111/j.10958312.2008.01140.x.

Troncoso-Palacios J, Díaz HA, Esquerré D, Urra FA. 2015. Two new species of the Liolaemus elongatus-kriegi complex (Iguania, Liolaemidae) from Andean highlands of southern Chile. ZooKeys 500:83-109. doi:10.3897/ zookeys.500.8725.
Troncoso-Palacios J, Díaz HA, Puas GI, Riveros-Riffo E, Elorza AA. 2016. Two new Liolaemus lizards from the Andean highlands of Southern Chile (Squamata, Iguania, Liolaemidae). ZooKeys 632:121-146. doi:10.3897/ zookeys.632.9528.
Urra FA, Díaz H, Werning H, Eisenberg T, Troncoso-Palacios J. 2017. Phymaturus vociferator Pincheira-Donoso, 2004 (Squamata: Liolaemidae): new records and updated geographic distribution. Check List 13:2137. doi:10.15560/13.3.2137.
Zhang J, Kapli P, Pavlidis P, Stamatakis A. 2013. A general species delimitation method with applications to phylogenetic placements. Bioinformatics 29:2869-2876.

Appendix 1. Specimens examined. Museum codes are as follow: MNHNCL (Museo Nacional de Historia Natural, Chile), MRC (Museo Regional de Concepción), MZUC (Museo de Zoología of Universidad de Concepción) and SSUC (Colección de Flora y Fauna Patricio Sánchez Reyes, Pontificia Universidad Católica de Chile). (download)

Appendix 2. Specimens used for phylogenetic analysis. (download)


[^0]:    *Correspondence: E-mail: jtroncosopalacios@gmail.com

