Four new species of *Stenocercus* Duméril & Bibron, 1837 (Squamata, Iguania) from the Department of Amazonas in northeastern Peru

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Abstract

The tropical Andes are known to be the richest and most diverse place on earth. This mountainous region covers almost one third of the Peruvian territory, and its herpetofauna remains poorly known. The lizard genus *Stenocercus* Duméril & Bibron, 1837 contains 69 species and most of them occur in the tropical Andes, although some exist as high as 4000 m a.s.l. The examination of newly collected material from the Andes of northern Peru in the Department of Amazonas reveals four new species of *Stenocercus* which we describe below. Of these four new species, three inhabit the ecoregion of Peruvian Yungas at elevations of 1460 to 2370 m a.s.l., and one the Marañón dry forest ecoregion at elevations of 1340 to 1470 m a.s.l. Additionally, we provide new data about coloration, natural history and distribution of the poorly known *S. aculeatus* O’Shaughnessy, 1879; and discuss the identity of some populations of *S. prionotus* Cadle, 2001 and *S. scapularis* Boulenger, 1901.

Key Words

Andes, herpetofauna, Marañón dry forests, montane forest, morphology, Peruvian Yungas

Introduction

With the creation of conservation biology, at the end of the seventies, as a new field of research, new and striking terms such as “biodiversity” or “hotspots” were coined to highlight the importance of biological diversity and areas with a high species richness and endemism, respectively. The tropical Andes, a name currently applied to the Andean mountains between Venezuela and Bolivia, are known to be the most diverse region on earth (Myers et al. 2000). This mountainous region covers almost one third of the Peruvian territory, running from north to south with an average height of 4000 m a.s.l. (Peñaherrera del Aguila 1989). The general landscape includes snow-topped mountains, canyons and valleys. The wide altitudinal range of the Andes also houses diverse habitats as altitude changes including tropical rainforests at 500 to 1500 m a.s.l., cloud forests ranging from 800 to 3500 m a.s.l.; and the highest altitudes of 3000 to 4800 m. a.s.l. contain páramos and grasslands extending up to snow (Duellman and Lehr 2009).

New species of reptiles and amphibians are frequently discovered in the Peruvian Andes especially when remote regions are surveyed (e.g. Cadle 1991; Duellman and Wild 1993; Duellman and Venegas 2005; Chávez et al. 2011; Venegas et al. 2013; Lehr et al. 2017). The Department of
Amazonas in northern Peru is located along the northern portion of the central Andes and possesses a complex toponography that in association with its climate pattern results in a wide variety of vegetation formations (Duellman and Pramuk 1999). The herpetological exploration of the Department of Amazonas has been fruitful since the decade of the seventies with the discovery of 38 species of anurans in the genera: *Atelopus* Duméril & Bibron, 1841 (Lötters et al. 2004), *Centrolene* Jiménez de la Espada, 1872 (Twom-ey et al. 2014), *Dendropsophus* Fitzinger, 1843 (Duellman 1982), *Gastrotheca* Fitzinger, 1843 (Duellman 1987, 2013; Duellman et al. 2014; Duellman and Venegas 2016), *Hylocercus* Peters, 1882 (Rivera-Correa et al. 2016), *Hyloxalus* Jiménez de la Espada, 1870 (Rivero 1991; Morales and Schulte 1993; Morales et al. 2004), *Nobellia* Barbour, 1930 (Duellman 1991), *Phyllomedusa* Wagler, 1830 (Cannatella 1982), *Pristimantis* Jiménez de la Espada, 1870 (Duellman 1990; Duellman and Pramuk 1999), *Rhine-ella* Fitzinger, 1826 (Duellman and Schulte 1992), *Telmato-ebius* Wiegmann, 1834 (Wiens 1993), *Scinax* Wagler, 1830 (Duellman and Wiens 1993); and 15 species of reptiles of the genera: *Ameiva* Meyer, 1795 (Koch et al. 2013), *Epictia* Gray, 1845 (Koch et al. 2015), *Moranasaurus* Dunn, 1933 (Köhler 2003), *Petracola* Doan & Castoe, 2005 (Echevarría and Venegas 2015), *Pseudogonatodes* Ruthven, 1915 (Huey and Dixon 1970); *Phyllocautulus* Gray, 1828 (Venegas et al. 2008; Koch et al. 2016), *Phyllocautus* Peters, 1878 (Koch et al. 2006), and *Stenocercus* Duméril & Bibron, 1837 (Fritts 1972; Cadle 1991; Venegas et al. 2016).

Currently the tropical Andes faces many threats and the most important by far are human activities including mining, logging, construction, agriculture and cattle ranching (Myers et al. 2000). The inter-Andean valleys of the Department of Amazonas are severely degraded and fragmented by human migration, agriculture and livestock (Venegas 2007). This is especially conspicuous in montane forests on the mountain slopes, particularly along the Chiriaco, Marañón and Utcubamba valleys. Therefore, documenting the biodiversity of the Department of Amazonas is a hard race against time.

Lizards of the *Stenocercus* clade are one of the most geographically and ecologically widespread taxa currently ranked as a genus in South America (Torres-Carvajal 2007a). The genus contains 71 species that occur at elevations between 0–4000 m a.s.l. in the Andes and adjacent lowland areas from northern Venezuela and Colombia to central Argentina, with some species in the Atlantic lowlands between southern Brazil and central Argentina, and others in northeastern Brazil (Torres-Carvajal 2007b; Uetz et al. 2020; Venegas et al. 2020). Peru is the country with the highest species richness in the genus *Stenocercus* with 59% of the species’ diversity. However, the taxonomy of *Stenocercus* in this country has remained problematic. In the past the major problems in the taxonomy of *Stenocercus* were the use of multiple generic names to describe new species, different specific epithets for the same described species, and some early descriptions are very brief, making identification difficult (Torres-Carvajal 2007a). Although in the last three decades the alpha taxonomy of *Stenocercus* was noticeably improved with high level descriptions and extensive taxonomic revisions (e.g Cadle 1991, 1998, 2001; Torres-Carvajal 2000, 2007a), and the advances in molecular systematics resolved the monophyly of the genus showing its diversification (Torres-Carvajal et al. 2006; Torres-Carvajal 2007b; Teixeira et al. 2016); identifying species of *Stenocercus* from Peru by physical examination remains a challenge. Additionally, the coloration in life and intraspecific variation in most of the Peruvian species remain poorly known. On the other hand, some species like *S. chrysopygus* Boulenger, 1900, are known to possess intraspecific variation among populations that merits careful examination and phylogenetic analysis (Fritts 1974; Cadle 1998; Schluter 2000; Torres-Carvajal 2007b). In fact, several undescribed species remain in herpetological collections awaiting formal description and new species are discovered in the Peruvian Andes continuously (Venegas et al. 2013, 2014a, 2016, 2020; Köhler and Lehr 2015).

In order to contribute to the knowledge of the scarcely known herpetofauna of the Peruvian Andes, we conducted several herpetological surveys of the montane environments of the Andes of northern Peru, especially in the Department of Amazonas. In this paper, we describe four new species of *Stenocercus* from that department and present new data regarding taxonomy, natural history, and distribution of *S. aculeatus* O’Shaughnessy, 1879. This contribution increases the number of *Stenocercus* species known from Peru to 47.

Materials and methods

Ethics and research

This study was carried out in accordance with the guidelines for use of live amphibians and reptiles in field research (Beaupre et al. 2004), compiled by the American Society of Ichthyologists and Herpetologists (ASH) and the Society for the Study of Amphibians and Reptiles (SSAR). Specimens collected for this study are covered by the following research permits (given by the Ministerio de Agricultura) that include permanent scientific collection of live specimens: 110-2007-INRENA-IFFS-DCB, 118-2007-INRENA-IFFS-DCB, 071-2008-INRENAIFFSDCB, 020-2009-AG-DGFFS-DGEFFS, 424-2010-AG-DGFFS-DGEFFS, 287-2011-AG-DGFFS-DGEFFS, 186-2014-MINAGRI-DGFFS/DGEFFS, 0581-2011-AG-DGFFS-DGEFFS, 295-2017-SERFOR/DGGSPPFFS, 299-2017-SERFOR/DGGSPPFFS, and 067-2019-MINAGRI-SERFOR-DGGSPPFFS.

Field techniques

Lizards were captured by hand or noosing with a fishing rod. Coordinates and elevation were taken with a GPS.
(Garmin, WGS84). All collected specimens were euthanized with T61, fixed in 10% formalin for 48 hours and permanently stored in 70% ethanol. Voucher specimens were deposited at the herpetological collection of the Centro de Ornitología y Biodiversidad (CORBIDI) in Lima, Peru.

**Morphological data**

Measurements of snout-vent length (SVL) and tail length (TL) were taken with a ruler and recorded to the nearest 1 mm. All other measurements were made with digital calipers and recorded to the nearest 0.1 mm. Sex was determined by noting the presence of hemipenes and dimorphic phenotypic characters such as height of the vertebral crest and the ventral pattern. Data on scutellation of all species of *Stenocercus* compared herein was taken from Torres-Carvajal (2007a) and Cadle (1991). Specimens reviewed for comparison are housed at CORBIDI and the Museo de Zoología, Pontificia Universidad Católica de Ecuador, Quito (QCAZ), and are listed in Appendix I. The abdominal skeleton was examined by dissection of specimens (all paratypes): CORBIDI 18661, CORBIDI 18876, CORBIDI 21090, and CORBIDI 21367. We follow the terminology of Cadle (1991) and Torres-Carvajal (2000; 2004; 2007a) for characters included in the description. The volume of oviducal eggs was calculated by the formula for a prolate spheroid $V = \frac{4}{3} \pi (\text{length}/2)^2$ (Torres-Carvajal 2007a).

**Species delimitation**

The taxonomic conclusions of this study are based on the observation of morphological features and color patterns as evidence to infer the existence of species (Frost and Kluge 1994; de Queiroz 1998, 2007). This information is considered as species delimitation criteria following a general lineage or unified species concept (de Queiroz 1998, 2007).

**Results**

*Stenocercus catherineae* sp. nov.

http://zoobank.org/ADE1DC5F-09B3-45A5-B107-FBF536F2FC72

Figs 1–3

**Type material.**

PERU • ♂, adult; Amazonas Department, Bongará Province, Florida District, Huembo; 5°51.47′S, 77°58.75′W; 2090 m a.s.l.; 09 Dec. 2019; P.J. Venegas and L.A. García-Ayachi leg.; CORBIDI 21365.

**Paratypes:**

PERU • 1 ♂, 1 ♀, adults, collected with the holotype; CORBIDI 21366-67 • 1 ♀, juvenile; Amazonas Department, Bongará Province, Cuispes District, Cuispes; 5°55.49′S, 77°56.94′W; 1850 m a.s.l.; 8 Mar. 2017; G. Chávez leg.; CORBIDI 18662 • 1 ♀, 1 ♂, adults; Amazonas Department, Bongará Province, Valera District, Co cachimba; 6°2.76′S, 77°53.55′W; 1460 m a.s.l.; 22 May 2007; by P.J. Venegas leg.; CORBIDI 501-02.

**Diagnosis.** *Stenocercus catherineae* sp. nov. differs from other species of *Stenocercus* except for *S. aculeatus*, *S. angulifer* Werner, 1901, *S. huancabambae* Cadle, 1991, *S. philmayi* sp. nov., *S. prionotus* Cadle, 2001, and *S. scapularis* Boulenger, 1901 by having: (1) projecting-angulate temporals; (2) laterally oriented nostrils; (3) dorsal and lateral scales of body similar in size, and (3) scales on posterior surface of thighs keeled and imbricate.

*Stenocercus aculeatus*, *S. angulifer*, *S. prionotus*, and *S. scapularis* can be easily distinguished from *S. catherineae* sp. nov. by having strongly keeled ventrals, whereas in the new species ventrals are feebly to moderately keeled. *Stenocercus aculeatus*, *S. angulifer* and *S. scapularis* also have a dorsolateral crest (absent in *S. catherineae* sp. nov.); and *S. prionotus* lacks a postfemoral mite pocket (present in *S. catherineae* sp. nov.). *Stenocercus catherineae* sp. nov. differs from *S. philmayi* sp. nov. by having smaller dorsal scales with 43 to 53 vertebrales and 46 to 59 scales around midbody versus 32 to 38 vertebrales and 34 to 45 scales around midbody in *S. philmayi* sp. nov. The gular region in adult males of *S. catherineae* sp. nov. is black, whereas in *S. philmayi* sp. nov. it is cream.

The new species is most similar to *S. huancabambae* (Fig. 4A, B) with which it shares ventral scales strongly keeled, like the rest of the aforementioned species, and a strongly compressed tail. Furthermore, both species are geographically close at the northern extreme of the central Andes in the Department of Amazonas (Cadle 1991; Torres-Carvajal 2007b). Nevertheless, the new species can be distinguished from *S. huancabambae* (characters in parentheses) by having the parietal eye not visible through the interparietal cornea (visible); postfemoral mite pocket as a distinct, deep slit-like opening (shallow slit-like opening, Fig. 3); dark patch covering most of the ventral surface of neck absent (present as a circular or elongate blotch, Fig. 4B); and a strongly compressed tail in adult males (very strongly compressed tail as a tape-like along its proximal two thirds).

At the locality of Cuispes, and probably also in Huembo, *S. catherineae* sp. nov. and *S. flagracanthus* sp. nov. exist in sympatry; however, both species are strikingly different. *S. flagracanthus* sp. nov. has a relatively short tail armed with projecting spines, whereas *S. catherineae* sp. nov. has a long tail, compressed laterally, and a projecting vertebral crest.

**Definition.** (1) Maximum SVL in males 83 mm ($n = 3$); (2) SVL in females 75 mm ($n = 2$); (3) vertebrales 43–53; (4) paravertebrales 62–73; (5) scales around midbody 46–59; (6) supraoculares 4; (7) internasales 4–6; (8) postrostrales 2–4; (9) gulars 5–6; (10) gulars 22–26; (11) subdigitals on Finger IV 18–21; (12) subdigitals on Toe IV 25–30; (13) posthumeral mite pocket present as a deep
depression with a wide opening [Type 3 of Torres-Carvajal (2007b)]; (14) postfemoral mite pocket present as a distinct pocket with a posterodorsally oriented slit-like opening [Type 2 of Torres-Carvajal (2007b)]; (15) parietal eye not visible through interparietal cornea in any specimens \( (n = 6) \); (16) scales on occipitoparietal region large, multicalcinate, not imbricate; (17) projecting angulate temporals present; (18) row of enlarged supraoculars present, occupying most of supraocular region; (19) scales on frontonasal region and supraoculars slightly imbricate, multicalcinate; (20) preauricular fringe present, short; (21) neck folds absent; (22) lateral and dorsal nuchals similar in size; (23) posterior gulars rhomboidal, projected posteriorly, keeled and imbricate, not notched; (24) lateral and dorsal body scales similar in size; (25) vertebrae larger than adjacent paravertebrae, forming a distinct vertebral crest; (26) dorsolateral crest absent; (27) ventral keeled, imbricate, mucronate; (28) scales on posterior surfaces of thighs keeled, imbricate, mucronate; (29) inguinal granular pocket absent; (30) inguinal groove absent; (31) preanals not projecting; (32) tail strongly compressed laterally in adult males; (33) tail length 65–70% of total length; (34) caudal whorls per autotomic segment three; (35) caudals not spinose; (36) dark brown stripe extend-
Figure 2. Dorsolateral and ventral views of *Stenocercus catherineae* sp. nov. in life: (A, B) holotype, adult male, SVL 82 mm (CORBIDI 21365); (C, D) adult male, SVL 79 mm (CORBIDI 21366); (E, F) adult female, SVL 75 mm (CORBIDI 21367); (G, H) adult female, SVL 75 mm (CORBIDI 501). Photographs by Pablo J. Venegas.
Description of the holotype. Adult male (Fig. 1); SVL 82 mm; TL 156 mm; maximum head width 15.7 mm; head length 18.7 mm; head height 12.8 mm; occipitals, parietals, interparietals, and postparietals large, multicarinate, slightly imbricate; parietal eye not visible; supracarinals in four rows, multicarinate, slightly imbricate, subequal in size; one canthal; canthal not in contact with the nasal; scales on frontonasal region slightly imbricate and multicarinate; internasals four; postrostrals three, two most lateral wider than long on each side, medial postrostrals as long as wide; supralabials four; infralabials five; loreals four; lorilabials in one row; preocular one, in contact with canthal; lateral temporals keeled, imbricate; gulars in 22 rows between tympanic openings; all gulars keeled, imbricate, apical pit absent; second infralabial not in contact with second and third sublabials; mental in contact with first pair of infralabials; lateral and dorsal scales of body and neck keeled, imbricate, mucronate; lateral and dorsal body scales similar in size; scales around midbody 46; vertebrals larger than dorsals, 43 scales on vertebral row, serrate vertebral crest present; paravertebrals 65; ventrals broad, rhomboidal, keeled, imbricate, preauricular fringe short, composed of three enlarged scales, all similar in size; antegular, gular, postauricular, oblique, supraauricular, longitudinal and antehumeral neck folds absent; limb scales keeled, imbricate; ventral scales of hindlimbs keeled and ventral scales of upper arms keeled, mucronate; lamellae on Finger IV 18; lamellae on Toe IV 27; tail strongly compressed laterally; caudals keeled, imbricate, mucronate; basal subcaudals keeled, imbricate; tail length 1.9 times SVL; posthumeral mite pocket present as a deep depression with a wide opening; postfemoral mite pocket present as a distinct deep pocket with a posterovertrally oriented slit-like opening; postfemoral region composed of imbricate, smooth scales that become keeled towards the tail.

Coloration in life (Fig. 2A, B). Dorsum pale brown with the first two chevrons over the vertebral line black and the rest slightly darker than the background; cream line extending vertically from the arm insertion to the scapular region surrounded by a black blotch; dorsal surface of limbs darker than the dorsum with faint dirty cream transverse bars; flanks reddish brown, including the tail, becoming red toward the venter and dotted with white; subocular and loreal regions creamy white; dorsal surface of head black with the superciliaries and rostral cream; labials, sublabials and mental black, extending as an irregular longitudinal stripe to the neck; gular region black with cream irregular blotches to the sides; a black irregular stripe extends from the gular region ventrolaterally to the arm insertion; ventral surface of neck, chest and forelimbs are dirty cream with a black spot ventrolaterally in the arm; belly and ventral surface of tail pink; pelvic region and ventral surface of hindlimbs dirty cream. The iris is reddish brown.

Coloration in preservative (Fig. 1D, E). It is similar to the coloration in life with the dorsal surface of trunk,

Figure 3. Lateral view of the postfemoral mite pocket in three species of Stenocercus: (A) S. catherineae sp. nov. (CORBIDI 21365), (B) S. philmayi sp. nov. (CORBIDI 21092), and (C) S. huancabambae (CORBIDI 19747). Photographs by Luis A. Garcia-Ayachi. Scale bars: 5 mm.
Figure 4. Dorsolateral and ventral views of adult males of three species of *Stenocercus*: (A, B) *S. huancabambae*, (C, D) *S. aculeatus*, (E, F) *S. prionotus* from northern-central Peru, and (G, H) *S. prionotus* from southern Peru. Photographs by: (A-D) Pablo J. Venegas, (E) Andy Barboza, (F) Diego Vasquez, and (G, H) German Chávez.
limbs and tail darker than in life. Moreover, the reddish coloration of the flanks almost disappears.

**Intraspecific variation.** Measurement and scutellation characters of *Stenocercus catherineae* sp. nov. are presented in Table 1. The second infralabial is in contact with the third sublabial in all specimens, and the first pair of postmentals are not in contact medially in two specimens (CORBIDI 501 and 21367). The three male paratypes are identical to the holotype, including a juvenile (CORBIDI 18662), varying only by having few scattered white dots on the head and, the black patch on the gular region extends to the ventral surface of the neck as a bold band (CORBIDI 21366) (Fig. 2C, D). Ventral scales in the juvenile male are strongly keeled and mucronate.

Sexual dimorphism is evident in adult individuals. In two adult female paratypes (Fig. 2E, H) dorsal coloration is dusky brown with cinnamon vertebral chevrons along the back and tail, and cinnamon blotches along the flanks; hindlimbs with scattered dark brown transverse stripes; head in both specimens are darker than the rest of body, being dark gray (CORBIDI 501) or dark brown (CORBIDI 21367); sides of head grayish white (CORBIDI 21367) or dusty cream with the gular region dark brown (CORBIDI 501) but both specimens have a faint pink hue on belly and base of tail (Fig. 2F, H).

**Distribution and natural history observations.** *Stenocercus catherineae* sp. nov. is only known from three proximate localities at Huembo and Cuispes in the northern extreme of the Cordillera Central at the Río Utcubamba basin in the Department of Amazonas, at elevations between 1466 to 2085 m a.s.l. (Fig. 5). According to the terrestrial ecoregions of the world by Olson et al.

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**Table 1.** Variation in scutellation and sexual dimorphism in snout-vent length (mm) of *Stenocercus aculeatus, S. catherineae* sp. nov., *S. dracopennatus* sp. nov., *S. philmayi* sp. nov., and their most similar species *S. angulifer* and *S. huancabambae*. Range followed by mean ± standard deviation is given for quantitative characters if applicable.

| Character                  | S. aculeatus CORBIDI 1712 n = 5 | S. angulifer CORBIDI 11483 n = 6 | S. catherineae CORBIDI 501 n = 3 | S. dracopennatus CORBIDI 21367 n = 32 | S. philmayi CORBIDI 21366 n = 8 |
|----------------------------|---------------------------------|---------------------------------|----------------------------------|---------------------------------------|-------------------------------|
| Scales around midbody      | 32–37                           | 38–50                           | 46–59                            | 39–45                                 | 37–53                         |
| Vertebrals                 | 31–39                           | 37–47                           | 43–53                            | 38–40                                 | 37–51                         |
| Paravertebrals             | 41–48                           | 37–57                           | 62–73                            | 53–57                                 | 48–64                         |
| Gulars                     | 15–18                           | 16–20                           | 22–26                            | 19–20                                 | 18–28                         |
| Supraoculars               | 16.60 ± 1.14                    | 18.47 ± 0.76                    | 24.33 ± 1.63                     | 19.33 ± 0.58                          | 20.93 ± 1.64                  |
| Internasals                | 4–5                             | 4–5                             | 4–5                              | 4–7                                   | 3–5                           |
| Subdigitals Finger IV      | 4–5                             | 4–5                             | 4–5                              | 4–7                                   | 3–5                           |
| Subdigitals Toe IV         | 23–27                           | 19–25                           | 25–30                            | 26–28                                 | 20–28                         |
| Tail length/total length   | 0.70 – 0.72                     | 0.68 – 0.73                     | 0.66 – 0.69 (n=5)                | 0.69 – 0.73                           | 0.67 – 0.70                   |
| Maximum SVL males          | 81                              | 95                              | 103                              | 96                                    | 82                            |
| Maximum SVL females        | unknown                         | unknown                         | 91                               | 82                                    | 75                            |

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![Figure 5. Distribution map of *S. catherineae* sp. nov. (red triangles), *S. dracopennatus* sp. nov. (yellow star), and *S. philmayi* sp. nov. (blue squares), and the similar species: *S. aculeatus* (fuchsia diamonds), *S. angulifer* (blue diamonds), *S. huancabambae* (orange diamonds), *S. prionotus* (turquoise diamonds), and *S. scapularis* (red diamonds). Symbols with a dot in the middle correspond to type localities. White circles with a dot are important cities for reference.](image-url)
(2001), this species occurs in the Peruvian Yungas ecoregion. The habitat at the type locality of \textit{S. catherineae} sp. nov. is a steep area located on the sides of the Río Chido with presence of corn, coffee and fruit plantations, and some patches of secondary forest and shrub vegetation. Several individuals were observed basking on fallen tree trunks or at the base of bushes close to trails between 1000 and 1200 hours. When the lizards were disturbed, they ran to hide inside fern patches. No other lizards or reptiles were observed in sympathy.

At Cuispes only one individual was collected in the croplands close to the village. Other squamate reptiles collected with \textit{S. catherineae} sp. nov. at Cuispes were \textit{Atractus} sp. Wagler, 1828, \textit{Dipsas palmeri} Boulenger, 1912, and \textit{S. flagracanthus} sp. nov. at Cocachimba, the two specimens collected were found basking at 1000 hours on the base of rocky fences with bushes along a trail. An uncollected adult female was observed basking at 900 hours on a fallen wall of an abandoned house. The general landscape at Cocachimba is composed of croplands of corn, fruit and sugar cane bordered by rocky fences and some streams with narrow fringes of riverine forest or shrub vegetation. Other species of squamate reptiles collected with \textit{S. catherineae} sp. nov. at Cocachimba were: \textit{Atractus} sp., \textit{Chironius exoletus} Linnaeus, 1758, \textit{Mastigodryas boddarti} Sentzen, 1796, and \textit{Petrola}c\textit{ola angustisoma} Echevarria & Venegas, 2015.

The female paratype (CORBIDI 21367) collected during the rainy season (December 2019) had 2 well-developed follicles, one in the left and the other in the right ovary. The sizes of these follicles are 10.83 × 9.67 mm and 10.99 × 9.53 mm; their volumes were 530.2 mm³ and 522.6 mm³, respectively.

\textbf{Etymology.} The specific name is a noun in the genitive case and is a patronym for Catherine Dupont, a Peruvian veterinary specialist in One Health, who is actively working searching and monitoring viruses and other zoonotic pathogens. The specific name of this lizard is in recognition of her passion for the natural world and its creatures, and her selfless support of the herpetological division of CORBIDI.

\textit{Stenocercus dracopennatus} \textit{sp. nov.}

\url{http://zoobank.org/0F8C57E0-3F73-48C1-9596-3657BE465B7E}

\textbf{Figs 6–9}

\textbf{Type material.} Holotype:

\textbf{PERU} • ♂; adult; Amazonas Department, Bongará Province, Yambrasbamba District, Yambrasbamba; 5°43.01’S, 77°58.61’W; 2370 m a.s.l.; 07 Sept. 2017; P.J. Venegas leg.; CORBIDI 18875.

\textbf{Paratypes:}

\textbf{PERU} • 2 ♂; adult and juvenile collected with the holotype; CORBIDI 18868, 18876.

\textbf{Diagnosis.} \textit{Stenocercus dracopennatus} \textit{sp. nov.} differs from all species of \textit{Stenocercus}, except for \textit{S. aculeatus}, \textit{S. angulifer}, \textit{S. prionotus}, and \textit{S. scapularis}, by having:

1. projecting angulate temporals,
2. laterally oriented nostrils,
3. dorsolateral crest (distinct on second half of body and base of tail in \textit{S. prionotus}),
4. dorsal and lateral scales of body similar in size,
5. strongly keeled ventrals, and
6. scales on posterior surface of thighs keeled and imbricate.

However, \textit{S. dracopennatus} \textit{sp. nov.} can also be easily distinguished from \textit{S. prionotus} (state of characters in parentheses) by having a low-lying vertebral crest (high and projected, Figs 4E, 9A) and postfemoral mite pocket as a slit-like opening (absent). Furthermore, \textit{S. dracopennatus} \textit{sp. nov.} differs from \textit{S. scapularis} (state of characters in parentheses) by having a thin and inconspicuous subocular stripe (conspicuous and broad subocular stripe, Fig. 16), smooth infralabials and sublabials (keeled), a black patch covering the ventral surface of neck (absent), and 38 to 40 vertebrals (43 to 53).

\textit{Stenocercus dracopennatus} \textit{sp. nov.} can be readily distinguished from \textit{S. angulifer} and \textit{S. aculeatus} by having two canthals and a black patch covering the ventral surface of the neck (one canthal and a black patch extensively covering most of the gular region in the last two species, see Figs 4D, 7B, D). Additionally, \textit{Stenocercus dracopennatus} \textit{sp. nov.} differs from \textit{S. aculeatus} (state of characters in parentheses) by having a longer snout (shorter, Fig. 8A, B), a low-lying vertebral crest (distinctly higher, Fig. 9A, B), a deeper posthumeral mite pocket (less deep, Fig. 8C, D), and cycloidal, smooth or feebly keeled dorsal scales at midbody between the dorsolateral crests with or without minute mucronations (lanceolate, strongly keeled and mucronate, Fig. 9C–F).

\textbf{Definition.} (1) Maximum SVL in males 89 mm \(n = 3\); (2) SVL in females unknown; (3) vertebrals 38–40; (4) paravertebrals 53–57; (5) scales around midbody 39–45; (6) supraoculars 4; (7) internasals 4–5; (8) postrostrals 2–5; (9) loreals 4–5; (10) gulars 19–20; (11) subdigitals on Finger IV 19–21; (12) subdigitals on Toe IV 26–28; (13) posthumeral mite pocket present as a deep depression with a narrow opening [Type 3 of Torres-Carvajal (2007b)]; (14) postfemoral mite pocket present as a distinct pocket with a posteroventrally oriented slit-like opening [Type 2 of Torres-Carvajal (2007b)]; (15) parietal eye not visible through interparietal cornea in any specimens \(n = 3\); (16) scales on occipitoparietal region large, feebly keeled or wrinkled, juxtaposed; (17) projecting angulate temporals present, two; (18) row of enlarged supraoculars occupying most of supraocular region present; (19) scales on frontonasal region feebly keeled, juxtaposed; (20) preauricular fringe present, distinct; (21) neck folds absent; (22) lateral and dorsal nuchals similar in size; (23) posterior gulars lanceolate, projected posteriorly, strongly keeled, mucronate and conspicuously imbricate; (24) lateral and dorsal body scales similar in size; (25) vertebrals larger than adjacent paravertebrals, forming a low vertebral crest; (26) dorsolateral crest present; (27) ventrals strongly keeled, imbricate, mucronate; (28) scales on posterior surfaces of thighs keeled, imbricate, mucronate; (29) inguinal granular pocket absent;
Figure 6. *Stenocercus dracopennatus* sp. nov. preserved holotype, adult male, SVL 79 mm (CORBIDI 18875): dorsal (A), lateral (B), and ventral (C) views of the head; dorsal (D) and ventral (E) views of the entire specimen. Photographs by Luis A. García-Ayachi. Scale bars: 10 mm.

(30) inguinal groove absent; (31) preanal sinus projected; (32) tail compressed laterally in adult males; (33) tail length 68–72% of total length; (34) caudal whorls per autotomic segment three; (35) caudals not spinose; (36) dark brown stripe extending anterodorsally from subocular region to supraciliary present, present only in juveniles; (37) dark patch extensively covering gular region of females unknown; (38) dark patch covering gular region in adult males absent; (39) dark patch on ventral surface of neck in adult males present; (40) dark midventral longitudinal mark such as faint line, conspicuous stripe, or extensive patch in adult males absent; (41) dark patches on ventral surface of thighs in adult males absent; (42) two xiphosternal and three postxiphosternal pairs of inscriptive ribs fused medially, forming three chevrons (Pattern 6A of Torres-Carvajal 2004).

**Description of the holotype.** Male (Fig. 6); SVL 79 mm; TL 210 mm; maximum head width 16.4 mm; head length 20.5 mm; head height 13.5 mm; parietals, interparietals and postparietals large; interparietals keeled, parietals and postparietals barely spinose, juxtaposed; occipital small, barely wrinkled; parietal eye not visible; supracroculos in four rows, keeled, slightly imbricate, subequal in size; canthals two; canthal not in contact with the nasal; scales on frontonasal region slightly imbricate, keeled; internasals five; postrostrals five, the three on the middle longer than wide and one third longer than postrostrals on the sides; supralabials
Figure 7. Dorsolateral and ventral views of *Stenocercus dracopennatus* sp. nov. in life: (A, B) holotype, adult male, SVL 79 mm (CORBIDI 18875); (C, D) adult male, SVL 88 mm (CORBIDI 18868); (E, F) juvenile male, SVL 56 mm (CORBIDI 18876). Photographs by Pablo J. Venegas.

five; infralabials seven; loreals five; lorilabials in one row; preocular one, in contact with second canthal; lateral temporalis keeled, some of these with a minute mucron, imbricate; gulars in 19 rows between tympanic openings; all gulars keeled, mucronate, imbricate, posteriorly projected, apical pit absent; second infralabial in contact with first to third sublabials; mental in contact with first pair of infralabials; lateral and dorsal scales of body and neck keeled, imbricate, mucronate; lateral and dorsal body scales similar in size; scales around midbody 39; vertebrals larger than dorsals, 38 scales on vertebral row, low serrate vertebral crest present; paravertebrals 53; ventrals broad, rhomboidal, strongly keeled, mucronate, imbricate; preauricular fringe short, indistinct, composed of six small scales, all similar in size; antegular, gular, postauricular, oblique, supraauricular, longitudinal and antehumeral neck folds absent; limb scales strongly keeled, imbricate, mucronate; ventral scales of hindlimbs and upper arms strongly keeled and mucronate; lamellae on Finger IV 21; lamellae on Toe IV 28; tail compressed laterally; caudals keeled, imbricate, mucronate but without projected spines; basal subcaudals strongly keeled, imbricate; tail length 2.6 times SVL; posthumeral mite pocket present as a deep depression with a narrow opening; postfemoral mite pocket present as a distinct deep pocket with a curved slit-like opening bordering the thigh insertion; postfemoral region composed of imbricate, keeled scales.
**Coloration in life** (Fig. 7A, B). Dorsal surface of the body is dusty brown with a greenish yellow hue on the pelvic region and tail; the vertebral and dorsolateral crests are yellowish with faint gray vertebral chevrons on the back, the posterior margin of each chevron is yellowish as well; antehumeral region with a dark brown vertical stripe with the anterior margin yellowish; flanks with diagonal rows of yellowish dots; limbs and proximal half of tail with transverse yellowish stripes; ventrolateral region turquoise; sides of head sepia and ocular region black. Ventrally, gular region sepia, ventral surface of neck covered by a black patch; chest, pelvic region and ventral surface of hindlimbs, and base of tail dirty cream with the sides of belly turquoise; proximal half of the tail is also dirty cream with transverse paler bands. The iris is dark brown.

**Coloration in preservative** (Fig. 6D, E). Similar to the coloration in life however the dorsal background and marks are paler than in life with a long patch of depigmentation on the back. The ventral surface turns bluish gray with scattered pale blotches on the belly.

**Intraspecific variation.** Measurements and scutellation characters of *Stenocercus dracopennatus* sp. nov. are presented in Table 1. The first pair of postmentals are not in contact medially in one specimen (CORBIDI 18868). The adult male paratype is larger than the holotype with 89 mm of SVL and its dorsal coloration is pale compared to the holotype, lacking the dark dorsal chevrons present in the holotype (Fig. 7C). The ventral pattern is identical to the holotype (Fig. 7D). The second paratype is a juvenile male (CORBIDI 18876) with the dorsum cinnamon and with the vertebral chevrons more contrasting than in the holotype; a distinct dark brown stripe extending anterodorally from subocular region to supraoculars; dots on the flanks of this juvenile specimen are yellow with the dots on the axillary region whitish and lacking the turquoise hue of the adult specimens (Fig. 7E). Ventrally, the gular region is dark sepia and the rest of body whitish cream with the sides of belly dark gray and not turquoise like in the adult (Fig. 7F). Females are unknown.

**Distribution and natural history observations.** *Stenocercus dracopennatus* sp. nov. is only known from the type locality, a summit near Yambasamba village at 2370 m elevation, located on the eastern slope of the Cordillera de Colán, at the Rio Chiriaco basin, Department of Amazonas, Peru (Fig. 5). According to Peñaherrera del Aguila (1989) and Olson et al. (2001), the distribution of this new species occurs within the Yungas and Peruvian Yungas ecoregions, respectively. The new species inhabits...
Figure 9. Lateral and dorsal views of adult males of Stenocercus for comparison: (A, C, E) S. dracopennatus (CORBIDI 18875) and (B, D, F) S. aculeatus (CORBIDI 1712). Photographs by Luis A. Garcia-Ayachi. Scale bars: 10 mm.
a mountain top covered by a dwarf montane forest full of terrestrial and arboreal bromeliads on a white sand soil. Six individuals were observed basking on a sunny morning, between 1000 and 1100 hours, on the sand and on fallen branches and running to find refuge in patches of terrestrial bromeliads and long grasses. No other reptile species were found in sympatry with *S. dracopennatus* sp. nov.

**Etymology.** The specific epithet “*dracopennatus*” is a noun derived from two words in Latin, “*draco*” that means dragon, the mythological being, and “*pennatus*” that means feathered. The specific name is a noun in apposition and refers to the similarity between lizards and dragons, which in both Western and Chinese cultures are beings similar to reptiles like crocodiles or serpents. Moreover, due to the big scales of this new species that give it the appearance of being covered by feathers, we decided to name *S. dracopennatus* sp. nov. for its resemblance to an imaginary feathered dragon.

**Stenocercus flagracanthus** sp. nov.

http://zoobank.org/C2AE661B-A0E9-4E88-B3E3-E412E6F1E414

**Type material.** Holotype: PERU • ♂, adult; Amazonas Department, Bongará Province, Cuispes District, Cuispes village; 5°55.49’S, 77°56.94’W; 1850 m a.s.l.; 7 Mar. 2017; G. Chávez leg.; from farms adjacent to Cuispes village; CORBIDI 18658.

**Paratypes:**

![Images of *Stenocercus flagracanthus* sp. nov.]

**Figure 10.** *Stenocercus flagracanthus* sp. nov. preserved holotype, adult male, SVL 75 mm (CORBIDI 18658): dorsal (A), lateral (B), and ventral (C) views of the head; dorsal (D) and ventral (E) views of the entire specimen. Photographs by Luis A. García-Ayachi. Scale bars: 10 mm.
Figure 11. Dorsolateral and ventral views of *Stenocercus flagracanthus* sp. nov. in life: (A, B) holotype, adult male, SVL 75 mm (CORBIDI 18658); (C, D) adult male, SVL 72.9 mm (CORBIDI 18659); (E, F) adult female, SVL 68 mm (CORBIDI 18661); (G, H) sub adult male, SVL 61 mm (CORBIDI 18660). Photographs by German Chávez.

PERU • 2 ♂, 1 ♀, adults; collected with the holotype; CORBIDI 18659–61 • 2 ♂, adults, 1 juvenile; Amazonas Department, Bongará Province, Shipasbamba District, Canta Gallo; 5°55.43’S, 77°59.03’W; 1720 m a.s.l.; 27 Aug. 2017; A. García-Bravo leg.; CORBIDI 18748, 18749, 18750 • 1 ♀, adult; Amazonas Department,
Bongará Province, Cuispes District, Cuispes village; 5°55.79’S, 77°56.66’W; 1880 m a.s.l.; 25 Sept. 2019; L.A. García-Ayachi leg.; CORBIDI 22035.

**Diagnosis.** *Stenocercus flagracanthus* sp. nov. differs from all other species of *Stenocercus*, except from *S. arndti* Venegas, Echevarria & Alvarez, 2014, *S. bolivarensis* Castro & Ayala, 1982, *S. carrioni* Parker, 1934, *S. chlorostictus* Cadle, 1991, *S. crassicaudatus* Tschudi, 1845, *S. empetrus* Fritts, 1972, *S. eunetopsis* Cadle, 1991, *S. torquatus* Boulenger, 1885, and *S. simonsii* Boulenger, 1899 by having granular lateral scales with strongly projected mucrons but differ by having the scales on the dorsal surface of neck and posterior surface of thighs imbricate and keeled.

**Definition.** (1) Maximum SVL in males 76.8 mm \((n = 5)\); (2) maximum SVL in females 68.3 mm \((n = 1)\); (3) vertebrae 83–97; (4) paravertebrals 95–111; (5) scales around midbody 96–104 (6) supraoculars 4–6; (7) internasals 4–6; (8) postrostrals 4–5; (9) loreals 4–7; (10) gulars 55–62; (11) lamellae on Finger IV 26–29; (12) lamellae on Toe IV 30–33; (13) posthumeral mite pocket present as one or more vertical folds or ridges \([\text{Type 1 of Torres-Carvajal (2007b)}]\); (14) postfemoral mite pocket distinct with slit-like opening \([\text{Type 2 of Torres-Carvajal (2007b)}]\); (15) parietal eye absent; (16) occipital scales small, smooth, juxtaposed; (17) projecting angular temporal absent; (18) row of enlarged supraoculars occupying most of supraocular region absent; (19) scales on frontonasal region juxtaposed, smooth; (20) preauricular fringe short; (21) antangular, antehumeral, gular, longitudinal, oblique, postauricular, and supra-auricular neck folds present; (22) lateral nuchals and dorsals similar in size; (23) lateral body scales, granular, smaller than dorsals, becoming slightly imbricate toward the groin; (24) vertebrae slightly enlarged, forming a distinct row of scales from forelimbs to hindlimbs; (25) dorsolateral crest absent; (26) paravertebrals from the second third of dorsum, like the adjacent dorsals, becoming gradually larger, imbricate, keeled and mucronate toward the hindlimb insertion; (27) ventral scales smooth, imbricate; (28) scales on posterior surface of thighs granular; (29) prefemoral fold present; (30) inguinal groove present; (31) preanals not projected; (32) tail not compressed laterally; (33) tail relatively short (tail length 50–54% of total length); (34) caudal whorls per autotomic segment two; (35) tail strongly spinose; (36) postocular stripe present; (37) gular region in males grayish with cream dots; (38) gular region in females grayish with pale dots; (39) black patch on ventral surface of neck in adult males absent; (40) dark midventral stripe in adult males absent; (41) dark patch on ventral surface of thighs, vent and tail in adult males absent; (42) background color of dorsum pale brown in males and gray in females, but with distinct black transversal stripes in both sexes; (43) two post-xiphisternal pairs of inscriptive ribs, one long (not in contact midventrally) and the other short \((\text{Pattern 1B of Torres-Carvajal 2004)}\).

**Description of holotype.** Male \((\text{Fig. 10)}\); SVL 75.0 mm; TL 87.0 mm; maximum head width 16.0 mm; head length 19.0 mm; head height 12.4 mm; scales on parietal and occipital regions small, smooth, juxtaposed, subequal in size; parietal eye not visible: supraoculars smooth, juxtaposed; circumorbitals absent; canthals two; loreals six; postrostrals four; internasals five; supralabials five; infralabials five; lorilabials in one row; preocular divided into two scales, most dorsal in contact with poste...
rior canthal; lateral temporals granular; gulars in 56 rows between tympanic openings; all gulars cycloid, smooth, imbricated; second infralabial in contact with first and second sublabials; first pair of postmentals in contact; mental in contact with first pair of infralabials and first pair of postmentals; dorsal and lateral scales of neck granular until the level of arm insertion, becoming gradually enlarged, imbricate, keeled to strongly keeled, and mucronate toward the hindlimbs insertion; lateral scales of body granular becoming slightly imbricate and feebly keeled toward the groin; scales around midbody 101; vertebrals 97 enlarged, keeled on the second half of body, imbricate, forming indistinct vertebral row; paravertebras adjacent to vertebras row larger than dorsals, keeled, imbricate becoming larger and mucronate toward hindlimbs insertion; paravertebras 111; vertrals smooth, imbricate, nearly twice the size of the dorsals, only paravertebras near hindlimbs insertion are twice the size of ventrals; preauricular fringe short composed of five enlarged, granular scales; suprauricular, antehumeral, gular, longitudinal, oblique, antegular, postauricular and rictal neck folds present; dorsolateral, ventrolateral and prefemoral folds present; dorsal scales of forelimbs imbricate, feebly keeled; dorsal scales of hindlimbs imbricate, strongly keeled and mucronate; ventral humeral scales granular becoming imbricate toward to the forearm; ventral scales of forelimbs and hindlimbs smooth, imbricate; palmar and plantar scales imbricate, keeled; lamellae on Finger IV 29; lamellae on Toe IV 33; tail rounded (tail length 53% of total length); caudal scales keeled, strongly mucronate, imbricate; basal subcaudal scales smooth, imbricate; posthumeral mite pocket present as one or more vertical folds or ridges; postfemoral mite pocket distinct with slit-like opening.

Coloration in life (Fig. 11A, B). Dorsal surface pale brown sptattered with dirty cream dots bearing a distinct black collar (incomplete dorsally) with dirty cream borders, broad white stripes without pale interspaces along dorsum, finely blotched with black on neck and limbs, and a middorsal triangular black blotch posterior to ociput; dorsal surface of head with black flecks; side of head with the loreal region, subocular scale, and jaws gray, supralabials and temporal region pale brown with a postocular black stripe; tail black with the distal quarter brown. Ventral surface creamy gray with the gular region gray with faint cream blotches better defined on the sides; tail surface at the distal half gray. Iris pale brown.

Coloration in preservative (Fig. 10D, E). Dorsal coloration gray, except on the head that remains brown, dots whitish cream, the borders of the collar white, and the black marks as in life. Ventral surface whitish cream with the gular region darker than in life.

Intraspecific variation. Measurements, scutellation, and other morphological characters of Stenocercus flagracanthus sp. nov. are presented in Table 2. Loreals 4–7; supralabials 4–6; infralabials 5; second infralabials in contact with third sublabials in 75% of specimens; first pair of postmentals in contact medially in all specimens. In one dissected paratype the pattern was two xiphiternal and two postxiphiternal pairs of inscriptive ribs, one long but not in contact midventrally and the other short [Pattern 1B; Torres-Carvajal (2004)].

The adult male paratypes (n = 4) are identical to the holotype (Fig. 11). Sexual dimorphism is noticeable in size, the single collected female (CORBIDI 18661) is smaller than males (Table 2). The black collar can be complete or incomplete and is present in both sexes, gray in the single female paratype and black in males. The dorsum in the female paratype is grayish brown; the dorsal black marks along the back of males are faint gray and black on the pelvic region and tail (Fig. 11E, F). The single juvenile specimen has the dorsum gray including the tail with a well-defined complete dark gray collar but without the transverse black stripes of the adult individuals and dorsal surface of head brown.

Distribution and natural history observations. Stenocercus flagracanthus sp. nov. is only known from two close localities, Cuispes village and Canta Gallo, both on the Amazon versant of the extreme northern portion of the central Andes in the Rio Ucubamba basin (Department of Amazonas), at elevations of 1720 and 1880 m (Fig. 12). According to Peñaherrera del Aguila (1989) and Olson et al. (2001), the distribution of this new species occurs within the Yungas and Peruvian Yungas ecoregions, respectively. The habitat of Stenocercus flagracanthus sp. nov. lies within agricultural lands with a mixture of corn, fruit trees and coffee plantations, and also pastures for livestock. The area east of Cuispes village has some montane forest remnants in steep areas but no individual of Stenocercus flagracanthus sp. nov. was observed there. Also, the road between Cuispes and Shipasbamba pass-

Table 2. Variation in scutellation and sexual dimorphism in snout-vent length (mm) of Stenocercus flagracanthus sp. nov., and the similar species S. crassicaudatus and S. torquatus. Range followed by mean ± standard deviation is given for quantitative characters if applicable. Scale counts and measurements of S. crassicaudatus and S. torquatus were taken from Torres-Carvajal (2007a).

| Character                  | S. flagracanthus | S. crassicaudatus | S. torquatus |
|---------------------------|------------------|-------------------|-------------|
| n                         | 8                | 31                | 46          |
| Scales around midbody     | 96–104           | 97–121            | 102–137     |
| Vertebrals                | 83–97            | 83–97             | 83–115      |
| Paravertebras             | 95–111           | 107–166           | 103–151     |
| Gulars                    | 55–62            | 44–55             | 47–67       |
| Supraoculars              | 4–6              | 6–8               | 6–8         |
| Internasals               | 4–6              | 4–7               | 4–6         |
| Subdigitals Finger IV     | 26–29            | 23–32             | 22–29       |
| Subdigitals Toe IV        | 30–33            | 26–38             | 26–32       |
| Tail length/total length  | 0.51–0.54 (n = 7) | 0.57–0.62         | 0.47–0.54   |
| Maximum SVL males         | 77               | 95                | 84          |
| Maximum SVL females       | 68               | 85                | 74          |
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Stenocercus philmayi sp. nov.

http://zoobank.org/82F392EA-A164-46F6-9461-1B197FE52804

Figs 13, 14

Type material. Holotype:

PERU • ♂, adult; Amazonas Department, Luya Province, Pisuquía District, Las Corontas; 6°28.75’S, 78°8.52’W; 1470 m a.s.l.; 13 Dec. 2019; I. Wong and A. García-Bravo leg.; CORBIDI 21077.

Paratypes:

PERU • 1 ♀, adult, 1 juvenile; collected with the holotype; CORBIDI 21090, 21093 • 2 ♂, adults, 2 juveniles; Amazonas Department, Luya Province, Pisuquía District, Las Corontas; 6°28.75’S, 78°8.52’W; 1470 m a.s.l.; 13 Dec. 2019; I. Wong and A. García-Bravo leg.; CORBIDI 21074, 21078, 21075, 21087 • 1 ♂, adult; Amazonas Department, Luya Province, Pisuquía District, Las Corontas; 6°28.79’S, 78°8.61’W; 1390 m a.s.l.; 13 Dec. 2019; I. Wong and A. García-Bravo leg.; CORBIDI 21077.

Diagnosis. Stenocercus philmayi sp. nov. differs from other species of Stenocercus except for S. aculeatus, S. angulifer, S. catherineae sp. nov., S. dracopennatus sp. nov., S. huancabambae, S. prionotus, and S. scapularis by having: (1) projecting-angled temporalis, (2) laterally oriented nostrils; (3) dorsal and lateral scales of body similar in size, and (3) scales on posterior surface of thighs keeled and imbricate. Stenocercus aculeatus, S. angulifer and S. scapularis differs from S. philmayi sp. nov. by having a dorsolateral crest (absent in S. philmayi sp. nov.). Stenocercus prionotus and S. philmayi sp. nov. share a prominent vertebral crest, however the former lacks a postemoral mite pocket (present in S. philmayi sp. nov.). Adult males of S. aculeatus and S. angulifer can be easily distinguished from S. philmayi sp. nov. by having the gular region covered by a black patch (absent in the new species) and fewer gulars (15 to 18 in S. aculeatus and 16 to 20 in S. angulifer versus 20 to 24 in S. philmayi sp. nov.). The new species differs from S. scapularis by having fewer scales around midbody (34 to 45 in S. philmayi sp. nov. versus 52 to 70 in S. scapularis according to Torres-Carvajal (2007b)).

The new species shares the presence of two canthals with the geographically close S. catherineae sp. nov. and S. dracopennatus sp. nov. (all from the northern extreme of the central Andes in the Department of Amazonas). However, S. philmayi sp. nov. possesses conspicuously larger dorsal scales than S. catherineae sp. nov., resulting in 32 to 38 vertebrales and 34 to 45 scales around midbody (43 to 53 vertebrales and 46 to 59 scales around midbody in S. catherineae sp. nov.). Adult males of S. catherineae sp. nov. have a black patch covering most of the gular region (Fig. 2B, D) and S. huancabambae has a black elongate or circular patch covering the ventral surface of the neck (Fig. 4B), and both species share a pink coloration on the belly and the base of the tail (Fig. 2B, 4B); whereas the new species lacks a black or pink coloration on ventral surfaces (Fig. 14B, D). Additionally, S. huan-
Figure 13. *Stenocercus philmai* sp. nov. preserved holotype, adult male, SVL 95 mm (CORBIDI 21092): dorsal (A), lateral (B), and ventral (C) views of the head; dorsal (D) and ventral (E) views of the entire specimen. Photographs by Luis A. García-Ayachi. Scale bars: 10 mm.

cabambae possesses a very strongly compressed tail laterally compared to the compressed tail of *S. philmai* sp. nov. In the case of *S. dracopennatus* sp. nov. (character state in parentheses) by having three occipitals (one), a black patch on the ventral surface of neck in adult males (dark coloration absent on ventral surface), and strongly keeled scales on the belly (keeled).

**Definition.** (1) Maximum SVL in males 95 mm \( (n = 4) \); (2) SVL in females 74 mm \( (n = 1) \); (3) vertebrae \( 32–38 \); (4) paravertebrals \( 49–59 \); (5) scales around midbody \( 34–45 \); (6) supraoculars \( 4–5 \); (7) internasals \( 3–5 \); (8) postrostrals \( 2–4 \); (9) loreals \( 4–6 \); (10) gulars \( 20–24 \); (11) subdigitals on Finger IV \( 18–21 \); (12) subdigitals on Toe IV \( 28–31 \); (13) posthumeral mite pocket present as a deep depression with a narrow opening [Type 3 of Torres-Carvajal (2007b)]; (14) preauricular mite pocket present as a distinct pocket with a posterovertrally orientated slit-like opening [Type 2 of Torres-Carvajal (2007b)]; (15) parietal eye not visible through interparietal cornea in any specimens \( (n = 8) \); (16) scales on occipitoparietal region large, keeled, not imbricate; (17) projecting, anagulate temporals present; (18) row of enlarged supraoculars occupying most of supraocular region present; (19) scales on frontonasal region not imbricate; (20) preauricular fringe present, distinct; (21) neck folds absent; (22) lateral and dorsal nuchals similar in size; (23) posterior gulars rhomboidal, projected posteriorly, keeled and con-
Figure 14. Dorsolateral and ventral views of *Stenocercus philmayi* sp. nov. in life: (A, B) holotype, adult male, SVL 95 mm (CORBIDI 21092); (C, D) adult male, SVL 86.0 mm (CORBIDI 21077); (E, F) adult female, SVL 74 mm (CORBIDI 21090); (G) juvenile male, SVL 55 mm (CORBIDI 21075); and (H) hatchling, SVL 33 mm (CORBIDI 21093). Photographs by Iván Wong.
spicuously imbricate, not notched; (24) lateral and dorsal body scales similar in size; (25) vertebrals larger than adjacent paravertebrals, forming a prominent vertebral crest; (26) dorsolateral crest absent; (27) ventrals keeled, imbricate, not mucronate; (28) scales on posterior surfaces of thighs keeled, imbricate, mucronate; (29) inguinal granular pocket absent; (30) inguinal groove absent; (31) preanalcs projected; (32) tail compressed laterally in adult males; (33) tail length 71–72% of total length; (34) caudal whorls per autonomic segment three; (35) caudals not spinose; (36) dark brown stripe extending anterodorsally from subocular region to supraciliaries absent; (37) dark patch extensively covering gular region of females absent; (38) dark patch covering gular region in adult males absent; (39) black patch on ventral surface of neck in adult males absent; (40) dark midventral longitudinal mark such as faint line, conspicuous stripe, or extensive patch in adult males absent; (41) dark patches on ventral surface of thighs in adult males absent; (42) two xiphisternal and three postxiphisternal pairs of inscriptions ribs fused medially, forming three chevrons (Pattern 6A of Torres-Carvajal 2004).

Description of the holotype. Male (Fig. 13); SVL 95 mm; TL 246 mm; maximum head width 17.5 mm; head length 22.5 mm; head height 15.4 mm; parietals, interparietals and postparietals large, parietals rugose and the rest of scales keeled, not imbricate, occipitalis three, small, keeled; parietal eye not visible; supraoculars in five rows, keeled, slightly imbricate, subequal in size; canthals two; canthal in contact with the nasal; scales on frontonasal region slightly imbricate, keeled; internasals four; postrostrals four, both wider than long; supralabials five; infralabials six; loreals five; lorilabials in one row; preocular one, in contact with second canthal; lateral temporals keeled, imbricate; gulars in 20 rows between tympanic openings; all gulars keeled, imbricate, apical pit absent; second infralabial not in contact with second and third sublabials; mental in contact with first pair of infralabials; lateral and dorsal scales of body and neck keeled, imbricate, mucronate; lateral and dorsal body scales similar in size; scales around midbody 40; vertebrals larger than dorsals, 35 scales on vertebral row, prominent serrate vertebral crest present; paravertebrals 59; ventrals broad, rhomboidal, keeled, imbricate; preauricular fringe short, composed of four enlarged scales, all similar in size; antegular, gular, postauricular, oblique, supraauricular, longitudinal and antehumeral neck folds absent; limb scales keeled, imbricate, mucronate; ventral scales of hindlimbs and upper arms keeled and mucronate; lamellae on Finger IV 18; lamellae on Toe IV 31; tail compressed laterally; caudals keeled, imbricate, mucronate; basal subcaudals strongly keeled, imbricate; tail length 3.42 times SVL; posthumeral mite pocket present as a deep depression with a narrow opening; postfemoral mite pocket present as a distinct shallow pocket with a posterovertrally orientated slit-like opening; postfemoral region composed of imbricate, keeled scales.

Coloration in life (Fig. 14A, B). Dorsal surface of body greenish gray with dark brown chevrons and narrow greenish white interspaces over the vertebral line; body flanks dusty brown splattered with whitish dots; dorsal surface of limbs olive green with scattered faint brown specks; dorsal surface of tail dusty brown with the crest greenish cream and faint cream transversal stripes; dorsal surface of head pale green with some scattered cream dots; sides of head greenish cream with the ocular region dark green; sides of neck greenish cream as the sides of head. Ventrally, gular region pale greenish cream; neck and chest paler than gular region; belly and tail cream with a tan hue; pelvic region pale cream and hindlimbs tan. The iris is dark brown.

Coloration in preservative (Fig. 13D, E). Similar to the life coloration however the greenish hue on the dorsum is pale brown and dorsal surface of the head is dark gray. Ventrally, gular region and chest dusty gray with a bluish hue, the rest of body is dark tan with the pelvic region dark cream.

Intraspecific variation. Measurements and scutella of Stenocercus philmaiayi sp. nov. are presented in Table 1. Second infralabial not in contact with third sublabial in any specimens, and first pair of postmentals not in contact medially in one specimen. The other adult male paratypes (n = 3) are identical to the holotype (Fig. 14C, D). Two juvenile males (CORBIDI 21087 and 21075) have the same dorsal pattern as adults (Fig. 14G) However, the dorsal surface of the head is brown, the greenish hue of the sides of head, neck and forearms is absent, and they possess a cream dorsolateral stripe that extends from the loreal region to the scapular region in CORBIDI 21087 and to the base of tail in CORBIDI 21075. Ventrally juvenile males are cream with scattered elongate pale gray blotches on neck, chest and sides of belly. The single hatchling paratype (CORBIDI 21093) has the dorsal surface dark brown with narrow black chevrons over the vertebral line and a longitudinal cream stripe from the loreal region to the scapular region of CORBIDI 21087 and to the half of tail in CORBIDI 21075. Ventrally juvenile males are cream with scattered elongate pale gray blotches on neck, chest and sides of belly. The single hatchling paratype (CORBIDI 21093) has the dorsal surface dark brown with narrow black chevrons over the vertebral line and a longitudinal cream stripe from the loreal region to the scapular region (Fig. 14H). The dorsal surface of the hindlimbs also presents thin black bars. Ventral surface is cream without marks.

Sexual dimorphism is evident in adults. Dorsal coloration in a single female paratype (CORBIDI 21090) is dark brown with thin darker brown chevrons and darker thin brown bars on hindlimbs (Fig. 14E, F). Also present a longitudinal cream stripe from the loreal region to scapular region becoming faint from the temporal region. Ventral surface is completely cream without marks.

Distribution and natural history observations. Stenocercus philmaiayi sp. nov. is only known from Las Coronatas in the northern portion of the central Andes at elevations of 1340–1470 m within the Rio Marañón basin (Fig. 5). According to the terrestrial ecoregions of the world by Olson et al. (2001), this species inhabits the Marañón dry forests ecoregion and following the ecoregions of Brack-Egg (1986), the equatorial dry forest ecoregion. The general landscape in the habitat of S. philmaiayi sp. nov. is the ecotone between dry forest and
humid montane forest. The dry forest in this zone has high trees with a canopy between 4 and 6 m, dense understory vegetation and scattered patches of cacti. Individuals of *S. philmayi* sp. nov. were observed during sunny days between 800 and 1400 hours basking on fallen logs close to trails that border or cross patches of forest. One adult male specimen was collected basking in the understory vegetation at 1 m in height. Other individuals were observed basking on rocks in patches of cacti and also on rocky fences with bushes near houses. Additionally squamate reptile species collected with *S. philmayi* sp. nov. were *Ameiva aggerecusans* Koch, Venegas, Rödder, Flecks & Böhme, 2013, *Microlophus stolzmanni* Stein-dachner, 1891, *Phyllodactylus pachamama* Koch, Flecks, Venegas, Bialke, Valverde & Rödder, 2016, *Epictia septemlineata* Koch, Venegas & Böhme, 2015, and *E. antoniogarciai* Koch, Venegas & Böhme, 2015.

Cadle (2001), reported an undescribed species of *Stenocercus* (represented by a single specimen), from 17 km ENE of Balsas village (6°49.00’S, 78°0.00’W) (Fig. 4) with similar features to *S. philmayi* sp. nov. The location of this specimen is 40.5 km to the south of the type locality of *S. philmayi* sp. nov. at an elevation of 1477 m, and lies also in the Marañón dry forests ecoregion.

The single female paratype collected during the rainy season (December 2019) had 2 eggs, one in the left and one in the right ovary. The sizes of these follicles are 19.71 × 9.44 mm and 19.81 × 8.40 mm; their volumes were 919.6 mm³ and 731.8 mm³, respectively.

**Etymology.** The specific epithet *philmayi* is a noun in the genitive case and is a patronym for Philip May (1946–2017), an American lichenologist and philanthropist, who was passionate about protecting biological diversity. During his life-time, his generous support of Nature and Culture International was instrumental to the protection of endangered ecosystems and endemic species in the Amazonas, Cajamarca, and La Libertad departments of Peru. Even after his death in 2018, his generosity has continued to protect Latin America’s biodiversity through charitable bequests. This new species was discovered in one of the departments that May supported during his life, and naming it after him, honors May’s enduring legacy as a champion of biodiversity.

**Stenocercus aculeatus O’Shaughnessy, 1879**

Figs 4C, D, 8B, 9B, D, F, 15

**Note.** An elusive species represented by five specimens in museum collections (Cadle 2001; Torres-Carvajal 2007a). Currently it is known from five localities at elevations of 723 and 1311 m between Zamora, in the Andes of southern Ecuador, and its southernmost record in Pampa Seca at Department of La Libertad in northern Peru (Cadle 2001; Torres-Carvajal and Carvajal-Campos 2009). No data about its natural history and coloration in life exist and probably due to its disjunct distribution more than one species is represented under its specific epithet.

Collecting specimens in the Andes of northern Peru for almost two decades, we acquired only two specimens of *S. aculeatus* in two locations of Abra Patricia at elevations of 1700 and 1990 m, just at the limits of the departments of Amazonas and San Martin (Fig. 5). Extending its altitudinal range by 679 m above its previously known record, Pampa Seca in the Mishollo Valley, at an elevation of 1311 m (Cadle 2001), one of the specimens (CORBIDI 1712) is also the first report for the Department Amazonas. Both specimens have scale counts similar to the description of Torres-Carvajal (2007a).

Abra Patricia is the pass between the Río Chiriaco and Río Mayo basins located in the Peruvian Yungas, according to Olson et al. (2001). It is a steep area mostly covered by humid montane forest with an abundance of orchids, bromeliads, lichens and *Chusquea* spp. Kunth, 1882. The forest at the base and on the slopes is 3 to 5 m of high, decreasing in height toward the tops. Some slopes are rocky and very steep with only shrubby vegetation or long grasses. The weather is rainy most of the year and hours with sun are usually limited.

One specimen (CORBIDI 11483) was collected by ornithologists of the Louisiana State University during an ornithological expedition to Abra Patricia in August 2002. The specimen is an adult male of 95 mm SVL preserved in ethanol with an overall dark gray coloration. Only a black patch extensively covering the gular region is dark enough to observe. Although no collecting data is available for this specimen the area where it was collected is a steep slope covered by montane forest near a cleared area for cattle ranching.

The second individual (Fig. 15) is another adult male (CORBIDI 1712) of 80 mm SVL and was collected by P.J. Venegas in October 2008. This specimen was encountered during the morning at 1000 hours on the top of long grass at a height of 50 cm in a flat area with scattered scrubs and grasses close to the road. The specimen was lethargic due to lack of sun, being easy to capture. The same day at 1200 hours three more individuals were observed: one male basking at the base of long grasses on a steep rocky wall close to the road and a couple were observed on a trail along a summit covered by scrub and scattered boulders to 200 m from the road. These individuals ran to hide at the base of dense scrub when they detected PJV getting closer, and were impossible to capture.

Coloration in life of adult males (CORBIDI 1712): dorsal surface (Fig. 4C) is dusty brown with a sepia hue on dorsum, bearing dark brown transverse bands with narrow cream interspaces; vertebral crest is greenish yellow with the tip of some spines brown; limbs and tail with faint cream transverse stripes; coloration of flanks similar to dorsum but with the ventrolateral region bluish; side of neck and arm insertion cream; dorsal surface of head dark brown with the occipital region darker; sides of head dark brown with the loreal and subocular region white, interrupted by a diagonal dark brown subocular stripe, snout dark brown, tympanic region and ventrolateral region of neck black. Ventrally (Fig. 4D), the anterior half
of the gular region dirty cream and the rest of the gular region and ventral surface of neck black; chest, forearms and belly brownish cream with a thin dark brown stripe from the chest to the end of belly, sides of belly bluish gray; pelvic region and ventral surface of hindlimbs tan, a cream blotch on the cloacal region; proximal half of tail creamy tan with faint brown transverse bands the rest of tail brown.

Discussion

Observations on the taxonomy and distribution of *Stenocercus aculeatus*, *S. prionotus* and *S. scapularis*.

*Stenocercus aculeatus* was a species briefly described in the past that generated confusion with other taxa, such as *S. angulifer*, due to the lack of specimens available in museum collections and the lack of data about its coloration in life (Cadle 2001; Torres-Carvajal 2000, 2007b); resulting in a species difficult to identify and to compare with similar species. Our two specimens of *S. aculeatus* reported here have similar scale counts to specimens reported by Torres-Carvajal (2007a), presenting a little variation in some counts, attributable to the low sample (*n* = 5) of *S. aculeatus* in the literature (see Table 1).

The type locality of *S. aculeatus*, Moyobamba, at an elevation of 723 m (Cadle 2001; Torres-Carvajal 2007a) is the lower altitudinal range for the species. However, the other known localities lie at elevations between 915 and 1311 m (Cadle 2001; Torres-Carvajal 2007a; Torres-Carvajal and Carvajal-Campos 2009), and the specimens reported here from Abra Patricia are from 1700 to 1987 m,
making the exact location of the type locality in O’Shaughnessy (1879), doubtful. Moyobamba is a city located in the valley of the Río Mayo at low elevation between two major mountain ridges. The closer mountains to Moyobamba are located 5 and 8 km from the city, reaching heights of 900 and 1500 m, and are similar to the other places where *S. aculeatus* specimens were collected (e.g. Cerro Calzada is the most striking mountain near to Moyobamba). Although our new records of *S. aculeatus* are higher than the rest of previously known localities, the closer known localities from Moyobamba for *S. aculeatus* (i.e. the range between Moyobamba and Cahuapanas, and Icuta on the Balsapuerto-Moyobamba trail; see Cadle (2001)) lie in the Cordillera Escalera, a mountain ridge to the east of Moyobamba that divides San Martin and Loreto departments (Pitman et al. 2014). The habitats of the Cordillera Escalera at elevations above 1000 m have a similar herpetofauna to Abra Patricia and the upper basin of the Río Mayo (see Venegas et al. 2014b). In fact, summits of the Cordillera Escalera reach 2700 m (Pitman et al. 2014) a similar elevation to the mountains of Abra Patricia.

The geographical distribution of *S. aculeatus* still seems peculiar with a gap of approximately 276 km to the northermost locality in Zamora in southern Ecuador (Torres-Carvajal and Carvajal-Campos 2009), and a gap of approximately 256 km to the southermost locality in Pampa Seca (Río Mishollo valley) in the Department of La Libertad (Cadle 2001; Torres-Carvajal 2007a). Nevertheless, except for *S. aculeatus*, no other species of *Stenocercus* occurs to north and south of the Río Marañón (the main tributary of the upper Amazon) (see Torres-Carvajal 2007a), and according to Cadle (2001), no *Stenocercus* species was reported in herpetofaunal surveys in northern Amazonas Department on the Río Cenepa and the Río Santiago or in biological rapid assessment of the Cordillera del Condor of southern Ecuador and northern Peru; or in the Cordillera de Kampankis, a narrow mountain ridge between the Río Santiago and Río Morona (see Catenazzi and Venegas 2012). For the gap toward the southermost locality in Pampa Seca, no evident barriers occur and montane habitats between 700 and 2000 m in this space are poorly known in herpetological terms. Although Torres-Carvajal and Carvajal-Campos (2009) did not show details or photographs of the specimen reported from Ecuador, the low sample size of specimens currently available for *S. aculeatus* in museums and the lack of DNA samples remains a barrier for understanding the variation and identity of this elusive species through its range.

In the process of diagnosing the new species described herein, we examined the morphology and distribution of *S. prionotus* and *S. scapularis*, both from the Peruvian Amazon foothills, and adjacent Amazon lowlands, including Bolivia for *S. prionotus* (Cadle 2001; Torres-Carvajal 2007b). As was shown by Cadle (2001), we also observed a conspicuous variation in the vertebral crest height and ventral coloration, between adult males of *S. prionotus* from populations of north-central Peru (Fig. 4E, F) and southern Peru and Bolivia (Fig. 4G, H). Both populations are separated by a gap between the Río Pachitea (Department of Huánuco) and Cocha Cashu (Department of Madre de Dios) (Cadle 2001). Although Cadle (2001) did not find significant statistical support for separating both populations based in a lower or higher number of vertebral scales as a reflection of their size, we suggest that populations from north-central Peru, southern Peru and Bolivia need to be treated as two allopatric and different species.

The sympatric relationships between species such as *S. aculeatus*, *S. prionotus* and *S. scapularis* were discussed by Cadle (2001) and remain poorly understood. Although the distributions of *S. aculeatus* and *S. prionotus* overlap in northern Peru, both species possess different altitudinal ranges, and *S. prionotus* occurs in lower elevations than *S. aculeatus*. Cadle (2001), mentioned possible places where *S. prionotus* and *S. scapularis* could be sympatric in central Peru but did not include southern Peru. However, the presence of *S. scapularis* in central Peru includes the distribution gap of *S. prionotus* (see Fig. 5) and we found clear variation in ventral coloration and head shape between populations of *S. scapularis* from central and southern Peru. Although we were not able to review adult males from central Peru, adult females from central and southern Peru (Fig. 16) can be distinguished (respectively): brown irises versus reddish irises, dark brown gular region versus pale lavender, brown belly with cream blotches vs. creamy tan belly with irregular transverse bands, and frontal region in profile gently raised versus abruptly raised (see Fig. 16E, F). The geographic gap between the central and southern populations lies in the Río Urubamba valley, which could well be a barrier, separating two different species.

Distributional patterns and morphological affinities between *S. aculeatus*, *S. catherineae* sp. nov., *S. dracopennatus* sp. nov., *S. huancabambae*, and *S. philmayi* sp. nov.

The interspecific phylogenetic relationships in *Stenocercus* are complex and morphologically heterogeneous. However, we discuss some conjectures originating on the basis of the morphological resemblance and the geographical proximity between *S. aculeatus*, *S. catherineae* sp. nov., *S. dracopennatus* sp. nov., *S. huancabambae*, and *S. philmayi* sp. nov. These five species are structurally similar (see Table 1); all of these species possess two xiphisternum and three postxiphisternal pairs of inscriptive ribs fused medially, forming three chevrons (Pattern 6A of Torres-Carvajal 2004) and are altitudinally segregated through the upper basin of the Río Marañón and its tributaries in the Huancabamba Deflection and the northern limit of the central Andes (see Fig. 4). While *S. huancabambae* inhabits dry deciduous forest at elevations of 200 to 1318 m along the Río Marañón and its tributaries, like the Río Utcubamba (Cadle 1991; Koch et al. 2018); *S. catherineae* sp. nov. occurs, like the former species, in the basin of the Río Utcubamba, but at higher elevations of 1460 and 2090 m, inhabiting montane forest; and *S. philmayi* sp. nov. occupies the ecotone be-
Figure 16. Dorsolateral and ventral views of *Stenocercus scapularis* in life: (A, B, E) adult female (CORBIDI 21983) from San Ramón, Department of Junín, Central Peru; (C, D, F) adult female (CORBIDI 17739) from La Convención, Department of Cusco, southern Peru; (G, H) adult male, (CORBIDI 17738) from La Convención, Department of Cusco, southern Peru. Photographs by Juan C. Chávez-Arribasplata.
between dry Marañon forest and montane forest in terrain with denser understory than the habitat of *S. huancabambae*, at a close altitudinal range between 1340 to 1470 m. On the other hand, *S. dracopennatus* sp. nov. inhabits the highest elevation at 2370 m in the humid montane forest of the Río Chiriaco basin (a tributary of the Río Marañon), whereas *S. aculeatus* is nearby inhabiting the same type of forest up to 1970 m elevation on the divide between the Río Chiriaco and Río Mayo basins.

Although these species apparently possess allopatric distributions, they are distinguished externally by sexually dimorphic characters (e.g. height of crest, marks on flanks and ventral coloration on gular region and neck; see Cooper and Burns 1987; Watkins 1998) with a potential role in either intra or interspecific communication that suggests possible contact zones or sympathy in some areas. *Stenocercus huancabambae* and *S. catherineae* sp. nov. are very similar in squamation (e.g. both species possess one canthal in contrast to two canthals in the other two new species), structure and color pattern, and distinguished basically by the form and size of the black patch in the gular-neck region and the height of the tail. The flanks in the four species are characterized by bright flecks with different levels of contrast between species being subtle in *S. philmayi* sp. nov. and conspicuous in *S. dracopennatus* sp. nov. Additionally, height of the crest is identical in *S. catherineae* sp. nov., *S. huancabambae*, and *S. philmayi* sp. nov. but strikingly low in *S. dracopennatus* sp. nov. On the other hand, the height of the tail and its compressed form in *S. huancabambae* are strongly pronounced in contrast to the rest of the species. Although not well understood, the relation of tail height and visual signals in lizards could be involved in intrasexual aggression as is the height of the crest in iguanas (e.g. Watkins 1998). The aforementioned species are similar in size; however, the strikingly high tail of *S. huancabambae* makes it distinctively bigger in lateral view.

We suggest that these specific distinctions shown by males, are due to the abundance of closely related species in the same region—the northern portion of the central Andes—likely with contact zones. Morphological resemblance and particular distinctions concerning features related to inter-sexual selection in this assemblage of lizards are a result of close phylogenetic relationships owing to a recent process of vicariance driven by constant geological changes in the formation of a complex landscape like the Huancabamba Deflection and the central Andes (Vuilleumier 1969; Cadle 1991; Duellman and Pramuk 1999).

Conservation status

We describe four new species of *Stenocercus* only known either from a single locality or up to a maximum of three localities. *Stenocercus catherineae* sp. nov. is known from three geographically proximate localities in the Río Utcubamba basin. This species was abundant in the type locality, where several individuals were found basking close to a trail on a slope of second growth vegetation along the Río Chido, a tributary of the Río Utcubamba. The general landscape at the type locality is strongly impacted by agriculture and cattle ranching with some scattered patches of forest. The same degree of fragmentation is observed in the locality of Cocachimba where several individuals of *S. catherineae* sp. nov. were observed in bushes and on rocky fences delimiting croplands. However, at the locality of Cuispes only one individual of *S. catherineae* sp. nov. was collected whereas several individuals of *S. flagracanthus* sp. nov. were observed. The latter species is known from two close localities, both in the Río Utcubamba basin. However, all individuals observed were found in croplands and were using human construction as refuges (e.g. houses, rocky fences and piles of firewood).

*Stenocercus philmayi* sp. nov. is known only from its type locality and an unconfirmed locality (see Fig. 5) reported by Cadle (2001). We found that *S. philmayi* sp. nov. matches the identification key of Cadle (2001) as “Stenocercus new species”, that is based on a single specimen that occurs 40.5 km to the south of the type locality of *S. philmayi* sp. nov., at a similar elevation and also in the Marañon dry forests ecoregion. Although we were unable to examine the specimen (ROM 16458) reported by Cadle (2001), we consider it possible that this specimen belongs to *S. philmayi* sp. nov., although this needs to be confirmed by examining the specimen or collecting more specimens at the same locality. The general habitat of this new species is better conserved than the habitat of *S. catherineae* sp. nov. and *S. flagracanthus* sp. nov.; and the main cause of habitat fragmentation in the type locality of *S. philmayi* sp. nov. are the clearings for coca (*Erythroxylum* sp. Browne, 1756), mango (*Mangifera indica*, Linnaeus, 1753a), and cacao (*Theobroma* sp. Linnaeus, 1753b). Nevertheless, *S. philmayi* sp. nov. is locally abundant and some individuals were observed near houses and croplands. Nevertheless, the distribution of this species remains known with certainty for a single locality. Finally, *S. dracopennatus* sp. nov. is also known from a single locality where only few individuals were observed. The general landscape is only impacted by a road and its habitat is on white sand soils so neither agriculture nor cattle ranching occur in the area; however, some slopes seem to have been burned in the past.

*Stenocercus catherineae* sp. nov., *S. flagracanthus* sp. nov., and *S. philmayi* sp. nov. inhabit croplands and use human-made buildings as refuges, as is well known for other species of *Stenocercus* (Dávila-Jativa and Cisneros-Heredia 2017). Although these habits showed a high degree of resilience in these species, we prefer to be cautious in giving them a conservation status, especially due to their poorly known geographical distribution. Thus, following the IUCN categories and criteria (IUCN 2012), we categorize these three species as Data Deficient (DD). On the other hand, in *S. dracopennatus* sp. nov., we have not observed adaptations for living in anthropic conditions and inasmuch as this species is known from a single locality, it must be considered as Data Deficient, too.
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Appendix I

Material examined.

Stenocercus aculeatus – PERÚ: Amazonas: Bongará: Abra Patricia, CORBIDI 01712; San Martín: Rioja: Cerro Patricia, CORBIDI 11483.

Stenocercus arndti – PERÚ: Cajamarca: Chota: Quebrada Checos, CORBIDI 01680,01681,01685, Quebrada La Irica, CORBIDI 01682–01684, 01686–01692.

Stenocercus carrioni – ECUADOR: Loja: Celica, Huajala QCAZ 10319, 10324, 10329.

Stenocercus chlorostictus – PERÚ: Piura: Ayabaca: Caserio de toronche – Comunidad de Olleros, CORBIDI 16891; Huancabamba: Canchaque, CORBIDI 19361–19366, 19368, 19369, 19382, 19383; Cajamarca: Santa Cruz: Agua Azul, MUSM 25821.

Stenocercus crassicaudatus – PERÚ: Cusco: Urubamba: Machu Picchu, CORBIDI 9058, MUSM 4905, 4906, 8691.

Stenocercus empetrus – PERÚ: La Libertad: Huamachuco MUSM 8658, 8675, 8677–8680; Santiago de Chuco: Shulgomopampa, CORBIDI 20534, 20535, 20536, 20537; Cajamarca: Cajamarca: MUSM 4909–10, La Colmena MUSM 8659, San Vicente MUSM 8676, Cerca al Cañon de Zangal, CORBIDI 01454, Ayllambo, CORBIDI 06532, Cataratas de Llacanora, CORBIDI 08638.

Stenocercus eunetopsis – PERÚ: Cajamarca: Santa Cruz, Udima, MUSM 4018, 4022–4029.

Stenocercus huancabambae – PERÚ: Amazonas: Bagua: Chonza Alta (Bagua Chica), CORBIDI 00737, Comunidad Nativa Copallin, CORBIDI 05605, Utcubamba: Cumba, CORBIDI 05791, 05792, Luya: Playa Jumet, CORBIDI 11941, Huaranguillo, CORBIDI 11976; Cajamarca: Jaén: El Tupire (Santa Rosa de la Yunga), CORBIDI 01014–01027, Santa Rosa de la Yunga, CORBIDI 01035, 01889, 01920, Jaén, CORBIDI 01861, 01864, Bellavista, CORBIDI 05790, Bosque de Yanahuana, CORBIDI 14844,14845, Chota: La Granja, CORBIDI 01693, Cutervo: Gotas de Agua, CORBIDI 01903, San Ignacio: Perico, CORBIDI 01930, 01931, 14061, Los Llanos, CORBIDI 19744, 19746, 19747, 19775.

Stenocercus prionotus – PERÚ: Huánuco: Huánuco: Dos Aguas, CORBIDI 6895, Malgotingo, CORBIDI 15001; Madre de Dios: Tambopata, CORBIDI 18599.

Stenocercus scapularis – PERÚ: Cusco: La Convención: Echarati, CORBIDI 06043, 17738, 17739, 17761, Santa Rosa de Cutivireni, CORBIDI 5431, Alto Shima, CORBIDI 9698, 9699, Chimparina, CORBIDI 18828; Junín: Satipo: Bosque rivereño, CORBIDI 21413, Chanchamayo: San Ramón, CORBIDI 21983.

Stenocercus simonsii – ECUADOR: Azuay: Giron, sendero El Salado, QCAZ 09636.

Stenocercus torquatus – PERÚ: Junín: Alto Yurinaqui, CORBIDI 21432, 21433, 21436, 21437, 21675–21677; Pasco: Oxapampa: Huampal, CORBIDI 7211, Bosque de Shollet, CORBIDI 9912.