A new capybara from the late Miocene of San Juan Province, Argentina, and its phylogenetic implications

ESPERANZA CERDEÑO, MARÍA E. PÉREZ, CECILIA M. DESCHAMPS, and VÍCTOR H. CONTRERAS


A new hydrochoerine rodent, Cardiatherium calingastaense sp. nov. (Caviidae), is described based on the specimen INGEO-PV 87. It was recovered from the late Miocene Las Flores Formation, cropping out at the Puchuzum locality, San Juan Province, Argentina. The new species is based on a particular combination of characters, among which the following can be highlighted: p4 with internal fissures equally deep as in Cardiatherium paranense, secondary external fissure as in Cardiatherium patagonicum, and lacking the fifth internal fissure and supernumerary internal fissure, as in C. paranense; m3 with a conspicuous labial column in the posterior ramus of the second prism; very deep primary and secondary external fissures in upper cheek teeth, the former producing a labial strong step-shaped profile in M2; sagittal crest on the parietals; bullae small in ventral view; scars of the origin of the masseter medialis muscle with an anterior projection up to the level of the incisive foramen and the maxilla-premaxilla suture. The phylogenetic analysis supports the taxonomic proposal of creating a new species of Cardiatherium and shows C. calingastaense sp. nov. as the sister group of the other species of the genus. The lineage leading to the clade Cardiatherium + largest capybaras would have originated at least during the Chasicoan SALMA (early late Miocene). Cardiatherium calingastaense sp. nov. adds to the previous record of Cardiatherium chasicoense and Cardiatherium paranense in the late Miocene of San Juan and Mendoza provinces, respectively, thus increasing the diversity of capybaras in central-west Argentina.

Key words: Mammalia, Rodentia, Caviidae, Cardiatherium, systematics, phylogeny, Miocene, Argentina.

Esperanza Cerdeño [espe@mendoza-conicet.gob.ar], Instituto Argentino de Nivología, Glaciología y Ciencias Ambientales (IANIGLA), Centro Científico Tecnológico, CONICET Mendoza, Av. Ruiz Leal s/n, 5500, Mendoza, Argentina. María E. Pérez [mperez@mef.org.ar], Museo Paleontológico Egidio Feruglio, Av. Fontana 140, U9100GYO, Trelew, Argentina.

Cecilia M. Deschamps [ceci@fcnym.unlp.edu.ar], Comisión de Investigaciones Científicas de la provincia de Buenos Aires, Argentina; División Paleontología Vertebrados, Unidades de Investigación Anexo Museo de La Plata, Facultad de Ciencias Naturales y Museo, Universidad Nacional de La Plata, 122 and 60, 1900, La Plata, Argentina.

Víctor H. Contreras [vcontre@unsj-cuim.edu.ar], Instituto de Geología Dr. Emiliano P. Aparicio (INGEO) y Departamento Geología, FCEFHN, Universidad Nacional de San Juan, Ignacio de la Roza y Meglioli, 5400 Rivadavia, San Juan, Argentina.

Received 5 September 2018, accepted 19 October 2018, available online 26 November 2018.

Copyright © 2018 E. Cerdeño et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License (for details please see http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Introduction

The Neogene continental outcrops in the San Juan Province, central-west Argentina (Fig. 1), encompass two localities that have yielded important late Miocene vertebrate remains. These localities are Puchuzum (Calingasta Valley), whose faunal assemblage is assigned to the Huayquerean SALMA (late Tortonian–Messinian; see Deschamps et al. 2013, for further details in bio- and chronostratigraphic equivalences), and Loma de Las Tapias, where two different levels are recognized as corresponding to the Chasicoan (Tortonian) and the Huayquerean South American Land Mammal Ages (SALMAs; Contreras et al. 1995; Peralta et al. 1995; Contreras 1996; Cerdeño and Contreras 2000; Ciancio et al. 2006; León and Bracco 2008; Bracco et al. 2011; Contreras and Baraldo 2011; Suvires and Contreras 2011).

Concerning the fossils from Puchuzum, different levels of the section (distinct architectural elements) have provided remains of molluscs, ostracods, insects, fishes, and palynomorphs (found within lacustrine sediments), on the one hand, and mammals and birds (derived from flood plain sediments), together with footprint tracks, on
the other hand (Contreras 1996; Contreras and Peralta 1998; Prămparo et al. 1998; Contreras and Cerdeño 2010; Bracco et al. 2011). Among mammal remains, only the specimens of a Hegetotheriidae (Notoungulata) were studied in detail (Cerdeño and Contreras 2000). However, the faunal assemblage also includes xenarthrans, litopterns and rodents (Contreras and Cerdeño 2010; Bracco et al. 2011). The hegetotheriid *Hemihegetotherium achataleptum* indicated a Huayquerian SALMA for this fauna (Cerdeño and Contreras 2000), but no absolute age has been obtained for the unit. Herein we present the study of a well-preserved skull and mandible of a capybara, associated with some fragments of postcranial bones.

Capybaras are the largest living rodents, represented by the single genus *Hydrochoerus*, reaching a body length of 100–130 cm, and adult males weighting up to 91 kg (Mones and Ojasti 1986). They have semiaquatic habits, and live in groups usually ranging in size from two to 30 individuals (Dunnun 2015). They have characteristic euhypodont, multilaminated cheek teeth (Novak 1991; Deschamps et al. 2007). The phylogenetic relationships of capybaras with other cavioid rodents and the status of hydrochoerids have been revised during the last years (Rowe and Honeycutt 2002; Vucetich and Pérez 2011; Upham and Patterson 2015; Pérez et al. 2018, and references therein). After these authors, capybaras would be within the Family Caviidae as a subfamily, Hydrochoerinae, together with the extinct cardiomyines and the small *Kerodon* (rock-dwelling mocos), the other living genus of the subfamily (Rowe and Honeycutt 2002; Pérez and Pol 2012; Pérez et al. 2018).

The fossil record of capybaras (that is, hydrochoerines excluding ‘cardiomyines’) goes back to the late Miocene (Ameghino 1883a, b; Rovereto 1914; Kraglievich 1930; Rusconi 1933, 1935, 1944; Pascual and Bondesio 1963; Francis and Mones 1965a, b; Bondesio 1985a, b; Mones 1991; among others), but revisions such as those by Mones (1984) and, more recently, Vucetich et al. (2005, 2012, 2014a, b, 2015) and Deschamps et al. (2007, 2009) have evidenced a great taxonomic overestimation due to ontogenetic variation that had not been previously evaluated; consequently, these authors drastically reduced the known diversity of hydrochoerines, especially for the late Miocene–Pliocene interval.

Within the late Miocene, the oldest species is *Cardiatherium chasicoense* Pascual and Bondesio, 1968 from the Chasicoan SALMA, which is known from the type locality Arroyo Chasicó (Buenos Aires Province) as well as from the lower levels of the Loma de Las Tapias Formation in San Juan Province (Contreras 2000; Contreras et al. 2001; Deschamps et al. 2007, 2009; Contreras and Baraldo 2011). In turn, the Huayquerian SALMA localities with hydrochoerines are more abundant and the material was re-evaluated by Vucetich et al. (2005). These authors described *Cardiatherium patagonicum* as the southernmost record, from the Puerto Madryn Formation (Chubut Province). At the same time, they proposed that all material from the Ituzaingó Formation (Entre Ríos Province, northeast Argentina) would correspond to only one species, *C. paranense*, which was later confirmed by Vucetich et al. (2014b). These authors also recognized *C. paranense* in Tupungato, Mendoza Province (central-west Argentina), from levels of the Río de Los Pozos Formation, referred to the late Miocene (Yrigoyen 1993). Another species assigned to this genus, *Cardiatherium orientalis* Francis and Mones, 1963b, was recorded in the late Miocene of Uruguay, and tentatively recognized later in the Cerro Azul Formation, La
The present contribution focuses on the description of a new species of *Cardiatherium* based on well-preserved cranial remains, previously mentioned as *Cardiatherium* sp. (Contreras and Cerdeño 2010), found in the province of San Juan. On the basis of a cladistic analysis, the phylogenetic position of the new species and the monophyly of the genus *Cardiatherium* within Caviidae were tested.

**Institutional abbreviations.**—GHUNLPam, Cátedra de Geología Histórica, Universidad Nacional de La Pampa, Santa Rosa, Argentina; INGEO-PV, Vertebrate Palaeontological collection, Instituto de Geología Dr. Emiliano P. Aparicio, Universidad Nacional de San Juan, San Juan, Argentina; MACN, Museo Argentino de Ciencias Naturales, “B. Rivadavia”, Buenos Aires, Argentina; MLP, Museo de La Plata, Argentina; MMH-CH, Museo Municipal de Ciencias Naturales de Monte Hermoso, Argentina; MMP, Museo Municipal de Ciencias Naturales “Lorenzo Scaglia”, Mar del Plata, Argentina; MNHN-AYO, Ayo-Ayo collection, Muséum national d’Histoire naturelle, Paris, France; MPEF-PV, Palaeontological collection; Museo Paleontológico “Egidio Feruglio”, Trelew, Argentina; PVSJ, Vertebrate Palaeontological collection, Instituto y Museo de Ciencias Naturales, Universidad Nacional de San Juan, San Juan, Argentina; SPV-FHC, Sección Paleontología Vertebrados, Departamento de Paleontología, Facultad de Humanidades y Ciencias, Montevideo, Uruguay.

**Other abbreviations.**—ch., character; he, external fissure; HFI, fundamental internal fissure; hfe, fundamental external fissure; hi, internal fissure; HPE, primary external fissure; hpi, primary internal fissure; HSE/hse, secondary external fissure; hsi, secondary internal fissure; hsi, internal supernumerary fissure; hti, tertiary internal fissure; M/m, upper/lower molar; P/p, upper/lower premolar; pr, prism; prIIa/b, anterior and posterior lobes of pr II; prsa, anterior secondary prism; SALMA, South American Land Mammal Age.

**Nomenclatural acts.**—This published work and the nomenclatural acts it contains, have been registered in ZooBank: urn:lsid:zoobank.org:pub:2A1D2DDE-7100-4BC7-B0B1-8862CC1A064E.

**Material and methods**

The studied material, INGEO-PV 87, includes a single specimen composed of a well-preserved skull with associated mandible, and some fragmentary postcranial remains of the same individual, housed at the Vertebrate Palaeontological Collection of the INGEO. It was compared with the holotypes and referred materials of *Cardiatherium chasicoense*, *C. paranense*, *C. orientalis*, and *C. rosendoi*, housed in the following Argentinean museums: IMUSI, MACN, MLP, MMH, MPEF, and MPEF. The specimens of *C. patagonicum* used for comparisons and the phylogenetic analysis were taken from Vucetich et al. (2005) and Dozo et al. (2010). *Cardiatherium aff. orientalis* from the Cerro Azul Formation in La Pampa Province was also used in this study. The detailed list of revised remains is in Appendix 1.

Dental nomenclature is given in Fig. 2A–E (flexus/flexid were originally named as fissures by Mones 1991, and are here indistinctly used) and follows partially Vucetich et al. (2005) and modifications according to Pérez et al. (2018). However, some modifications have been introduced in this paper, in order to fit with a new character observed in the p4 of *Cardiatherium patagonicum* and the new species here described. This new feature is the external flexid of the posterior lobe of prII of p4 (Fig. 2A), here referred to as hse or h2e according to the nomenclature of m1–m3 (h.s.e. sensu Mones 1991; Fig. 2B–D). Hence, the external fissure of the anterior supernumerary prism of p4 (prsa) is named here h3e (h.s.e. or h.2e sensu Mones 1991; Fig. 2A, D). Lobes of caviids are referred to as prisms in hydrochoerines. Cranial nomenclature follows Vucetich et al. (2014b) and Pérez et al. (2018).

Skull measurements follow Vucetich et al. (2014b): (i) posterior width of the incisive foramen; (ii) rostral width at the anterior margin of the anterior root of the zygomatic arch.

---

![Fig. 2. Dental nomenclature (occlusal view). A–C, E. *Cardiatherium calingastense* sp. nov. A. Right p4. B. Right m1. C. Right m3. E. Right M1. D. *Cardiatherium patagonicum*, right p4. Abbreviations: c, column; he, external fissure (or flexid); HFI, fundamental internal fissure (hypoflexus); hfe, fundamental external fissure; hi, internal fissure; HPE, primary external fissure; hpi, primary internal fissure; HSE/hse, secondary external fissure; hsi, secondary internal fissure; hsi, internal supernumerary fissure; hti, tertiary internal fissure; pr, prism; prIIa/b, anterior and posterior lobes of pr II; prsa, anterior secondary prism.](image-url)
(complete or up to the midline in incomplete specimens); (iii) width between the anteriori”most points of the scars marking the origin of the masseter medialis muscle; (iv) rostral width at the level of P4, measured at the alveolar margin of prism I; (v) rostral width at the posterior margin of the anterior root of the zygomatic arch (complete or up to the midline in incomplete specimens); (vi) rostral width at the level of M2, measured at the alveolar margin of pR1; (vii) distance between the posterior margin of the incisive foramen and P4; (viii) P4–M1 anteroposterior length; and (ix) angle of the beginning of the ventral anterior zygomatic arch and the alveolar series, in lateral view. In order to explore the phylogenetic relationships of the taxon studied, we scored the new anatomical information throughout the last version of the combined Cavioid matrix (Madozzo Jaén et al. 2018), originally published by Pérez and Pol (2012). The combined dataset of morphological and molecular characters (SOM 1 and 2; Supplementary Online Material available at http://app.pan.pl/SOM/app64-Cerdeno_etal_SOM.pdf) was modified. Some multistate characters are ordered, nine characters have been modified and eight characters are new (see SOM 2). The new dataset was analysed under equally weighted parsimony in TNT (Goloboff and Catalano 2015), using a heuristic search of 1000 replicates of Wagner trees, followed by tree bisection and reconnection branch swapping. Node supports were calculated using absolute GC bootstrap frequencies calculated after 1000 pseudoreplicates. The modified Stratigraphic Manhattan Measure (Pol and Norell 2001) was used to calibrate the most parsimonious phylogenetic trees and the chronostratigraphic information for fossil taxa using TNT.

Geological setting

The specimen herein studied was collected in 2006 in levels of the Las Flores Formation, in the area of Puchuzum in the Calingasta Valley (Fig. 1). This unit is widely developed both geographically and in thickness in the Iglesia valley, where its type locality has been recognized (Wetten 1975; Contreras et al. 2013 and references therein). The stratigraphic section at Puchuzum was previously described by Peralta et al. (1995) (see also Bracco et al. 2011), and corresponds to the Neogene sediments cropping out at both margins of the Candelaria Creek, in the piedmont of the Manrique Mountains, Cordillera Frontal. According to geological and palaeontological studies at Puchuzum (Peralta et al. 1995; Contreras et al. 1995; Contreras 1996; Contreras and Peralta 1998; Cerdeño and Contreras 2000; Bracco et al. 2011; Bogan et al. 2018), the bony remains recorded in this locality, which come from the middle of the exposed section, were assigned to the latest Miocene. This mid-section is characterized by brown-reddish to greyish psammo-pelitic deposits, reddish silty to silty-sandy beds, with some carbonaceous and conglomeratic levels, and with cut and fill structures. Three mantiform interbeddings of bentonitic clay, 3 to 10 m thick, are distinguished by their yellowish coloration. These bentonitic packages clearly correspond to sedimentary events that originated in lakes, evidenced by the recorded fossils: bivalves, gastropods, insects, and fish (Peralta et al. 1995; Contreras 1996; Contreras and Peralta 1998; Bracco et al. 2011; Bogan et al. 2018). In turn, the mammals from this mid-section at Puchuzum represent a Huayquerian faunal association within the late Miocene (Peralta et al. 1995; Contreras 1996; Cerdeño and Contreras 2000).

The general palaeoenvironment of the mid-section of Puchuzum is interpreted as a fluvial system with great development of flood plains and rivers, more or less channelled and strongly sinuous. The lacustrine interbeddings correspond to increasing and decreasing events of the lacustrine basin, influenced by tectonic and hydrologic activity. A similar situation has been interpreted for the Las Flores Formation in the Iglesia Basin (Milana 1994; Jordan et al. 1997).

Systematic palaeontology

Rodentia Bowdich, 1821
Hystricognathi Tullberg, 1899
Cavioida (Fischer, 1817) sensu Kraglievich 1930
Caviidae (Fischer, 1817) sensu Waterhouse 1839
Hydrochoerinae (Gray, 1825) Gill, 1872
Genus Cardiatherium Ameghino, 1883b
Type species: Cardiatherium doeringi Ameghino, 1883b (= Cardiatherium paranense [Ameghino, 1883a] in Vucetich et al. 2014b); Paraná River cliffs, Entre Ríos, Argentina, Ituzaingó Formation, Late Miocene.
Species included: Cardiatherium patagonicum Vucetich, Deschamps, Olivares, and Dozo, 2005, Cardiatherium orientalis (Francis and Mone, 1965b), Cardiatherium chasicoense (Pascual and Bondesio, 1968), Cardiatherium rosendoi Bondesio, 1985b, Cardiatherium talicai Francis and Mones, 1965a, and Cardiatherium isseli Rovereto, 1914.
Stratigraphic and geographic range.—Late Miocene (Chasicoan and Huayquerian SALMAs) of Argentina, Brazil, Peru, Uruguay, and Venezuela.

Cardiatherium calingastaense sp. nov.
Figs. 3, 4.
2010 Cardiatherium sp.; Contreras and Cerdeño 2010: 8R.
ZooBank LSID: urn:lsid:zoobank.org:act:343DF38E-F84D-43C7-9687-05C91FF26257
Etymology: After the Calingasta Valley, where the locality of Puchuzum is located.
Holotype: INGEO-PV 87, skull and mandible of an adult individual, together with associated fragments of vertebrae and long bones.
Type locality: Right side of the Candelaria Creek, Puchuzum area, Calingasta Valley, San Juan Province, central-west Argentina.
Type horizon: Mid stratigraphic section of Las Flores Formation, late Miocene.
Diagnosis.—Cavioid diagnosed by the following unique combination of characters (autopomorphies marked with an
Fig. 3. Holotype (INGEO-PV 87) of the hydrochoerine rodent Cardiatherium calingastaense sp. nov., late Miocene, Puchuzum, San Juan Province, Argentina. A. Skull in dorsal (A1) and ventral (A2) views, and detail of the scars for masseter medialis muscle (A3); correspondence indicated by lines. B. Mandible in occlusal (B1) and left lateral (B2) views.
asterisk): euhyodont cheek teeth; p4 with three prisms and P4 with two-lobed prisms as in other Hydrochoerinae; p4 with hfe, h3e, hpi, h2i, and h3i as in Hydrochoeropsis, Phugatherium, Neochoerus, and Hydrochoerus; h2i and h3i equally deep as in Cardiatherium paranense, Hydrochoeropsis dasseni, but differing from Hydrochoerus and Neochoerus in which h2i is shallower than h3i and its apex is behind the hfe; p4 with hse (h2e) as in C. Pattonicus; p4 lacks hsi and hsn, as in C. paranense; hpi, hti, HFI, and hfe of m1/M1–m2/M2 not splitting the teeth, as in other Cardiatherium and differing from Hydrochoeropsis, Phugatherium, Neochoerus, and Hydrochoerus, in which those flexi/ids completely cross the tooth; *m3 with a conspicuous labial column in the posterior ramus of pR1a in contrast to other species; HPE and HSE very deep, the latter shorter than the former but deeper than in other species and oriented backwards; M3 with nine prisms; sagittal crest on the parietals, unlike Hydrochoeropsis, Phugatherium, Neochoerus, and Hydrochoerus, in which the area between both temporal fossae is plane, without forming the sagittal crest; bullae small in ventral view as in Neochoerus and Hydrochoerus; *scars marking the origin of the masseter medialis muscle on the beginning of the anterior zygomatic arch (ventral face), with an anterior projection up to the level of the incisive foramen and the maxilla-premaxilla suture.

Description.—As it is reflected in the diagnosis of the new proposed taxon, the specimen INGEO-PV 87 (Figs. 3, 4) may be referred to the genus Cardiatherium, discarding its assignment to other genera of capybaras such as Phugatherium, Hydrochoeropsis or the modern lineages represented by Hydrochoerus and Neochoerus (see also Phylogenetic analysis and Discussion). Therefore, the following description is focused on the comparison of the studied material with the known species of Cardiatherium.

Skull: It is somewhat mediolaterally deformed and lacks the left zygomatic arch, the left parietal-occipital area, and the left M3 (Fig. 3A1, A2).

In dorsal view (Fig. 3A1), the nasal is anterolaterally broken but the preserved portion indicates that this bone is long. The nasal is a little longer than the frontal in the new species as in C. paranense and the biggest capybaras (Phugatherium, Hydrochoeropsis, Hydrochoerus). Laterally, the nasal is in contact with the dorsal projection of the premaxilla and posteriorly with the frontal. The nasal is dorsally slightly convex, differing from the strongly bulging condition of C. paranense and from Hydrochoerus, in which the nasal is slightly bulgy in the middle portion. The posterior border of the nasal is anteriorly slightly concave, differing from the strongly concave border of C. paranense. The nasal and frontal (Fig. 3A1) are proportionately narrower than in C. paranense, Phugatherium, and Hydrochoerus. The frontal and the anterior portion of the parietal are dorsally flat as in most Hydrochoerinae. The posterior portion of the parietal has a temporal fossa posterolaterally concave and the temporal lines join, although not completely fusing, in a short sagittal crest that diverges again posteriorly at the nuchal crest. This condition contrasts with that of the largest capybaras, which present an interposed area between the temporal fossae (which are shallower than in C. calingastaense sp. nov.) instead of the sagittal crest. The postorbital apophyses are incomplete and would have been short.

In right lateral view, the ventral portion of the zygomatic arch is preserved and is similar to the condition of Hydrochoerus. The maxillary anterior root of the arch is dorso-ventrally flattened and narrows posteriorly, being dorso-ventrally elongated like the jugal, which is antero-posteriorly short. The squamosal forms the posterior portion of the arch and is posteriorly extended above the tympanic bulla. The external auditory meatus is above the level of the molar series. The angle formed by the anterior root of the zygomatic arch and the alveolar plane in Lateral view is <20°, similar to C. paranense, but different from C. aff. orientalis from the Cerro Azul Formation, in which it is 24–26° (Vucetich et al. 2014b).

In ventral view (Fig. 3A2), the diastema is longer than the molariform series (Table 1), like in C. paranense, C. Pattonicus, H. dasseni, and P. novum. The scars for the insertion of the masseter superficialis muscle are anteriorly convergent as in the other species of Cardiatherium; however, in INGEO-PV 87 there is an additional scar in this area. One is oval, transversely concave, placed between the root of the zygomatic arch and the P4, but extended anteriorly beyond this tooth, as observed in the other species of the genus; and the other one is another smooth surface at an obtuse angle with the first scar, which is directed to the sagittal plane, reaching the border of the incisive foramen, where the premaxilla-maxilla suture is located (Fig. 3A2, A3). The incisive foramen is ovoid and broad as in C. paranense whereas it is narrow in C. aff. orientalis (Table 1). The base of P4 does not project laterally and therefore does not form the bulge observable in ventral view in C. paranense. The ratio between the distance from the base of the incisive foramen up to the middle point between both P4s and the upper tooth series length is close to the minimum values of Cardiatherium, but different from the large capybaras (Phugatherium) and the modern lineages (Hydrochoerus–Neochoerus), in which the tooth series is proportionally longer (Table 1). The palate is rather shallow; from the level of M1 to that of the third prism of M3, there is a medium, narrow crest with a long and narrow palatal foramen at each side (M2–M3 level). The mesopterygoid fossa is narrow in outline, with a rounded anterior margin at the level of the posterior portion of the M3, as in C. paranense and C. Pattonicus. The tooth rows are divergent, but with a smooth curvature with M3 aligned with P4–M2, as in C. paranense. In C. Pattonicus and C. aff. orientalis, the M3 are nearly parallel. The basicranium is badly preserved, although part of the basisphenoid and the right tympanic bulla are preserved. The bulla is relatively small. The occipital face is incomplete and displaced; the right paraoccipital apophysis is distally incomplete, but it would have been relatively long (Fig. 3A1).
Mandible: The mandible preserves the symphysis, with the incisors, most of the left horizontal ramus, with p4–m3, and the anterior part of the right ramus (Fig. 3B). The right cheek teeth were recovered separately, but complete. The symphysis is narrow and procumbent, with a smooth median keel on the labial side; the posterior end of the symphysis does not reach the premolar level. The horizontal rami are well divergent; they are wide and vertically convex; ventrally, the area from the level below m1 to the base of the symphysis is flattened. The mental foramen is located anteriorly to p4 and at the dorso-ventral midpoint of the lateral surface of the dentary. The bottom of the i1 alveolus reaches internally the anterior half of the m2 and widens the lingual side of the ramus (clearly visible on the left side). The posterior border of the symphysis in C. calingastaense sp. nov. is below p4; according to Vucetich et al. (2005) and Deschamps et al. (2007), its position depends on the age of individuals, located anterior of p4 in juveniles and below the midpoint of p4 in larger (older) individuals. Main mandibular measurements are provided in Table 2.

Dentition: Both upper and lower incisors are narrow (I1, 8.1 mm; i1, 7.4 mm and 7.6 mm) and strongly curved, with bevelled occlusal surface and enamel only on the labial side (Fig. 3B).

Table 1. Comparison of skull dimensions (in mm, except for 9, 12) of Cardiatherium calingastaense sp. nov. following Vucetich et al. (2014b) with other late Miocene Cardiatherium and Recent Hydrochoerus hydrochaeris. Abbreviations: 1, posterior width of the incisive foramen; 2, rostral width at the anterior margin of the anterior root of the zygomatic arch (complete or up to the midline in incomplete specimens); 3, width between the anteriormost points of the scars marking the origin of the masseter medialis muscle; 4, rostral width at the level of P4, measured at the alveolar margin of prism I; 5, rostral width at the posterior margin of the anterior root of the zygomatic arch (complete or up to the midline in incomplete specimens); 6, rostral width at the level of M2, measured at the alveolar margin of pRfI; 7, distance between the posterior margin of the incisive foramen and P4; 8, P4–M1 anteroposterior length; 9, angle of the beginning of the ventral anterior zygomatic arch and the alveolar series, in lateral view; 10, length of upper diastema; 11, length of P4–M3; 12, relationship (%) between dimensions 10 and 11; 1 holotype of C. paranense and MLP 71-VI-16-1; 2 GHUNLPam 27389 (the smallest) and GHUNLPam 5274 (the largest).

<table>
<thead>
<tr>
<th>Skull dimension</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardiatherium calingastaense sp. nov. (INGEO-PV 87)</td>
<td>5.8</td>
<td>29.0</td>
<td>13.5/21.8</td>
<td>18.1/25.1</td>
<td>29.9</td>
<td>32.6</td>
<td>19.0</td>
<td>19.3</td>
<td>18</td>
<td>71.1</td>
<td>56.6</td>
<td>75.3</td>
</tr>
<tr>
<td>Cardiatherium paranense (N = 12)</td>
<td>3.2</td>
<td>22.6</td>
<td>–/16.0</td>
<td>10.7/-</td>
<td>21.4</td>
<td>24.0</td>
<td>9.0</td>
<td>12.5</td>
<td>18</td>
<td>132.0</td>
<td>31.0</td>
<td>96.8</td>
</tr>
<tr>
<td>maximum</td>
<td>8.0</td>
<td>32.4</td>
<td>–/28.0</td>
<td>19.4/-</td>
<td>37.7</td>
<td>43.9</td>
<td>22.5</td>
<td>22.5</td>
<td>19</td>
<td>75.0</td>
<td>59.0</td>
<td>78.6</td>
</tr>
<tr>
<td>Cardiatherium aff. orientalis (N = 5)</td>
<td>2.5</td>
<td>21.8</td>
<td>–/16.0</td>
<td>12.9/-</td>
<td>22.1</td>
<td>30.5</td>
<td>15.1</td>
<td>10.2</td>
<td>24</td>
<td>74.2</td>
<td>41.5</td>
<td>93.8</td>
</tr>
<tr>
<td>minimum</td>
<td>3.0</td>
<td>29.0</td>
<td>–/27.0</td>
<td>17.4/-</td>
<td>28.2</td>
<td>36.8</td>
<td>18.8</td>
<td>18.8</td>
<td>26</td>
<td>58.0</td>
<td>53.0</td>
<td>91.3</td>
</tr>
<tr>
<td>maximum</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>75.0</td>
<td>57.0</td>
<td>76</td>
</tr>
<tr>
<td>Cardiatherium patagonicum (MPEF-PV 2521)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>25.1</td>
<td>37.2</td>
<td>148.2</td>
</tr>
<tr>
<td>Hydrochoerus hydrochaeris (N = 8)</td>
<td>5.0</td>
<td>16.2</td>
<td>26.2</td>
<td>11.0</td>
<td>20.2</td>
<td>23.6</td>
<td>9.2</td>
<td>11.6</td>
<td>13</td>
<td>25.1</td>
<td>37.2</td>
<td>148.2</td>
</tr>
<tr>
<td>minimum</td>
<td>10.2</td>
<td>40.0</td>
<td>34.1</td>
<td>24.6</td>
<td>43.2</td>
<td>52.0</td>
<td>24.2</td>
<td>27.2</td>
<td>17</td>
<td>67.9</td>
<td>79.3</td>
<td>116.8</td>
</tr>
</tbody>
</table>
Among upper cheek teeth (Fig. 4A1–A4; Table 3), P4, M1, and M2 are formed by two heart-shaped prisms, joined by a short isthmus. As in all species of Cardiatherium, HPE is deeper than HSE, unlike Phugatherium in which these flexi are almost similar in depth (Fig. 5A–E). In INGEO-PV 87, the HPE is very deep and curved backward, in such a way that in M2 it reaches the labial side and produces a deformation of this wall resulting in a strong step-shaped profile (Figs. 4A2, A3, 5A). The HSE is shorter than HPE, but is deeper than in the other species and oriented slightly backward (Fig. 5A–C); only the largest specimen of C. patagonicum has an HSE similar in depth to C. calingastaense sp. nov., but anteriorly concave. P4 has undulated walls of HPE. The HFI is so deep that it almost splits prI from prII. M3 has nine prisms; prI is bilobed with a deep HPE similar to those of the other upper cheek teeth; the other eight prisms are laminar with a shallow, open V-shaped labial fissure in prI to prVII whereas in prVIII there is a barely marked labial concavity; as usual in capybaras, prIX is transversally shorter with a convex posterior margin and no labial fissure (Figs. 4A9, 5A).

Among lower cheek teeth (Figs. 4A5–A9, 6A; Table 4), p4 has a deep h3e, bifurcated at the end, and anteriorly directed. C. calingastaense sp. nov. lacks h4i and h5i as C. paranense (Fig. 6C), whereas h5i is extended up to 25% of the occlusal surface in C. patagonicum (Fig. 6B), C. orientalis (Fig. 6D) and C. chasicoense (Fig. 6E). C. calingastaense sp. nov. and C. aff. orientalis have no supernumerary internal flexid (hsni) typical of C. orientalis and C. patagonicum (Fig. 6B, D). The hfe is also deep, wide and transversely directed whereas in other species it is oblique. Lingually, the hpi is transversely directed and very deep, reaching almost the labial side, producing a deformation on this margin; at the same time, the third external flexid (h2e) is markedly concave (Figs. 4A8, 6A) whereas in C. patagonicum (also with very deep hpi) the h2e is only incipient (Fig. 6B). The h2i and h3i are long

Table 2. Mandibular dimensions (in mm) of Cardiatherium calingastaense sp. nov. (INGEO-PV 87).

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Symphysis length</td>
<td>51.5</td>
</tr>
<tr>
<td>Anterior transversal diameter of symphysis</td>
<td>23.6</td>
</tr>
<tr>
<td>Transversal diameter symphysis at foramen level</td>
<td>31.9</td>
</tr>
<tr>
<td>Anterior height of symphysis</td>
<td>c. 23.4</td>
</tr>
<tr>
<td>Height symphysis at foramen level</td>
<td>24.0</td>
</tr>
<tr>
<td>Transversal diameter between p4s, anterior to alveolus</td>
<td>30.6</td>
</tr>
<tr>
<td>Height horizontal ramus ahead left p4</td>
<td>20.3</td>
</tr>
<tr>
<td>Maximum width anterior to lateral crest</td>
<td>67.1</td>
</tr>
</tbody>
</table>

Fig. 6. Schematic occlusal view of lower teeth of Cardiatherium. A. Cardiatherium calingastaense sp. nov. (late Miocene, Puchuzum, San Juan Province, Argentina), holotype, INGEO-PV 87, right p4–m2 and reversed left m3. B. Cardiatherium patagonicum (late Miocene, Estancia Rincón Chico, Chubut Province, Argentina), MPEF-PV 740/9, reversed left p4; MPEF-PV 740/24, right m1–m2; MPEF-PV 740/7, reversed left m3. C. Cardiatherium paranense (late Miocene, Ituzaingó Formation, Entre Ríos Province, Argentina), MLP 50-40-XI-15-1, right p4–m3. D. Cardiatherium orientalis (late Miocene, Camacho Formation, San Gregorio, Uruguay), SPV-FHC-27-XI-64-20, right p4–m3. E. Cardiatherium chasicoense (late Miocene, Arroyo Chasicó, Buenos Aires Province, Argentina), MMP 300-M, right p4–m2; MMH-CH 88-6-71, reversed left m3.
and also bifurcated at the end, and both delimit a conspicuous internal column (c3). In C. patagonicum, h2i is much shallower than h3i; in C. paranense (Fig. 6C) and C. chasicoense (Fig. 6E), these flexids are shallower and similar in depth, those of the latter being the shallowest. The m1–m2 are similar in morphology (Figs. 4A 6, 6A). The hsi penetrates less than 50% of the width of prI as in the other species of Cardiatherium, whereas hti is deep (crossing the prism but not splitting as in C. patagonicum; Fig. 6B) and bifurcated at the end, producing a deformation of the labial margin at this point. The hse is deep and wide. The m3 is the longest molar of the series, and flexids are not bifurcated. This molar has a conspicuous labial column in the posterior ramus of prIIa, delimited by two accessory fissures between hse and hfe (Figs. 4A7, 6A).

The previous comparative description has been focused on the best-known species, but some comparison can also be provided for two other poorly known taxa. In C. isseli (holotype and only specimen MACN 6354; middle Miocene, Río Negro Province; Rovereto 1914), the p4 is quite similar, but the m3 differs because the anterior margin of prI is straight and the hsi is very deep. In C. talicei (holotype SPV-FHC-10-VIII-63-1; Pliocene, San Gregorio, Uruguay; Francis and Mones 1965a; Mones 1991, who adds two other specimens), the flexids are anteroposteriorly wider; the p4 has deeper h4i and no h2e; h2i and h3i are shallower, especially the former; and the m3 shows a deep hsi, more anteriorly directed than in C. calingastaense sp. nov., which is reflected in a longer and more triangular first prism.

Stratigraphic and geographic range.—Late Miocene, middle section of the Las Flores Formation. Right side of the Candelaria Creek, Puchuzum area, Calingasta Valley, San Juan Province, central-west Argentina.

Phylogenetic analysis

We assessed the phylogenetic affinities of C. calingastaense sp. nov. using the dataset and protocols presented in previous studies (Pérez et al. 2018; Madozzo Jaén et al. 2018; see SOM 1–3). The parsimony analysis of the combined dataset (including morphological and molecular partitions) resulted in 333 most parsimonious trees (MPTs) of 3364 steps. The strict consensus of all MPTs is shown in SOM 3. Figure 7 displays the node Hydrochoerinae in the strict consensus.

Table 3. Comparison of upper tooth dimensions (in mm) of Cardiatherium. a minimum and maximum values after Vucetich et al. (2014b); * M1 and M2 not differentiated; AW, anterior width, L, length; PW, posterior width; W, width; approximate values in parentheses.

<table>
<thead>
<tr>
<th></th>
<th>P4</th>
<th>M1</th>
<th>M2</th>
<th>M3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>L</td>
<td>AW</td>
<td>PW</td>
<td>L</td>
</tr>
<tr>
<td>Cardiatherium calingastaense sp. nov. (INGEO-PV-87) right</td>
<td>10.0</td>
<td>6.3</td>
<td>7.6</td>
<td>8.8</td>
</tr>
<tr>
<td>Cardiatherium patagonicum* minimum</td>
<td>8.0</td>
<td>7.2</td>
<td>7.5</td>
<td>8.1</td>
</tr>
<tr>
<td>Cardiatherium paranense* minimum</td>
<td>6.0</td>
<td>4.6</td>
<td>4.6</td>
<td>5.1</td>
</tr>
<tr>
<td>Cardiatherium aff. orientalis* minimum</td>
<td>4.8</td>
<td>3.0</td>
<td>3.6</td>
<td>4.0</td>
</tr>
<tr>
<td>Cardiatherium paranense* maximum</td>
<td>12.0</td>
<td>8.2</td>
<td>9.7</td>
<td>11.0</td>
</tr>
<tr>
<td>Cardiatherium paranense* maximum</td>
<td>12.0</td>
<td>8.2</td>
<td>9.7</td>
<td>11.0</td>
</tr>
<tr>
<td>Cardiatherium paranense* maximum</td>
<td>12.0</td>
<td>8.2</td>
<td>9.7</td>
<td>11.0</td>
</tr>
<tr>
<td>Cardiatherium aff. orientalis* maximum</td>
<td>12.0</td>
<td>8.2</td>
<td>9.7</td>
<td>11.0</td>
</tr>
<tr>
<td>Cardiatherium orientalis (holotype)* minimum</td>
<td>11.2</td>
<td>4.6</td>
<td>5.9</td>
<td>9.6</td>
</tr>
<tr>
<td>Cardiatherium aff. orientalis* minimum</td>
<td>9.9</td>
<td>3.8</td>
<td>4.8</td>
<td>8.0</td>
</tr>
<tr>
<td>Cardiatherium patagonicum* maximum</td>
<td>23.0</td>
<td>8.5</td>
<td>10.6</td>
<td>11.2</td>
</tr>
<tr>
<td>Cardiatherium paranense* maximum</td>
<td>12.0</td>
<td>4.6</td>
<td>5.9</td>
<td>12.3</td>
</tr>
<tr>
<td>Cardiatherium chasicoense* minimum</td>
<td>6.3</td>
<td>2.1</td>
<td>3.6</td>
<td>6.1</td>
</tr>
<tr>
<td>Cardiatherium chasicoense* maximum</td>
<td>13.7</td>
<td>4.8</td>
<td>6.8</td>
<td>11.5</td>
</tr>
</tbody>
</table>

Table 4. Comparison of lower tooth dimensions (in mm) of Cardiatherium. a minimum and maximum values after Vucetich et al. (2005); b values correspond to the holotype MLP 40-XI-15-1; c values after Deschamps et al. (2009), a very small m1–m2 of Cardiatherium patagonicum was not here considered; * m1–m2 not differentiated; AW, anterior width; L, length; MW, medium width; PW, posterior width; approximate values in parentheses.

<table>
<thead>
<tr>
<th></th>
<th>p4</th>
<th>m1</th>
<th>m2</th>
<th>m3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>L</td>
<td>AW</td>
<td>MW</td>
<td>PW</td>
</tr>
<tr>
<td>Cardiatherium calingastaense sp. nov. (INGEO-PV-87) left</td>
<td>14.4</td>
<td>5.3</td>
<td>7.0</td>
<td>6.8</td>
</tr>
<tr>
<td>Cardiatherium paranense* minimum</td>
<td>9.7</td>
<td>4.0</td>
<td>5.1</td>
<td>5.1</td>
</tr>
<tr>
<td>Cardiatherium paranense* maximum</td>
<td>19.8</td>
<td>6.4</td>
<td>7.8</td>
<td>8.8</td>
</tr>
<tr>
<td>Cardiatherium orientalis (holotype)*</td>
<td>11.2</td>
<td>4.6</td>
<td>5.9</td>
<td>9.6</td>
</tr>
<tr>
<td>Cardiatherium aff. orientalis* minimum</td>
<td>9.9</td>
<td>3.8</td>
<td>4.8</td>
<td>8.0</td>
</tr>
<tr>
<td>Cardiatherium aff. orientalis* maximum</td>
<td>23.0</td>
<td>8.5</td>
<td>10.6</td>
<td>11.2</td>
</tr>
<tr>
<td>Cardiatherium patagonicum* maximum</td>
<td>12.0</td>
<td>4.6</td>
<td>5.9</td>
<td>13.3</td>
</tr>
<tr>
<td>Cardiatherium chasicoense* minimum</td>
<td>6.3</td>
<td>2.1</td>
<td>3.6</td>
<td>6.1</td>
</tr>
<tr>
<td>Cardiatherium chasicoense* maximum</td>
<td>13.7</td>
<td>4.8</td>
<td>6.8</td>
<td>11.5</td>
</tr>
</tbody>
</table>
Fig. 7. Stratigraphic adjustment of the strict consensus tree for the phylogenetic analysis of Hydrochoerinae (333 MPTs of 3364 steps). Upper case letters indicate the nodes within Hydrochoerinae, as described in the main text. Silhouettes reflect external aspect and relative size of representatives of the main clades within Hydrochoerinae.
Our phylogenetic analysis recovers the monophyly of *Cardiatherium* (Fig. 7: node B and SOM 3). At the same time, this clade is the sister group of the largest capybaras (a clade including *Hydrochoeropsis*, *Phugatherium*, *Neochoerus* and *Hydrochoerus*; Fig. 7). *Cardiatherium calingastaense* sp. nov. is the sister group of the other species of the genus (Fig. 7: node B). *Cardiatherium* and the largest capybaras (Fig. 7: node A) share the following unambiguous synapomorphies: root of the lower incisors extending up to the level of the anterior lobe of m2 (ch. 20); concave palatal surface (ch. 55); presence of h2i on p4 (ch. 86); anterior location of h3i (ch. 90); hpi deeper than h5i (ch. 96); c3 present on prl of p4 (ch. 98); oblique h3e (ch. 101); posterior prism of m1–m2 complex heart-shaped (ch. 106); presence of hti on m1–m2 (ch. 109); hsi shallower than hpi on m1 (ch. 114); hpi in m1–m2 reaching the labial end (ch. 115); presence of hse on m1–m2 (ch. 116); p4–m1 shorter than m2–m3 (ch. 121); m1<m2<m3 (ch. 122); m3 complex (ch. 123); HPE deeper than HSE (ch. 138); laminar posterior prisms in M3 with three or more prisms, in which the first prism is heart-shaped (ch. 141); only the first prism is heart-shaped or lanceolate-shaped and the others are laminar in M3 with three or more prisms (ch. 142).

On the other hand, the *Cardiatherium* clade (Fig. 7: node B) is supported by two unambiguous synapomorphies: antero-posterior length of the posterior portion of the upper diastema more than 20% of the antero-posterior length of the maxilla (measured from the premaxilla-maxilla suture to the posterior border of maxilla at the level of the posterior projection of M3) (ch. 52), and hsi in m1–m2 is extended less than 50% of the occlusal surface (ch. 108). The other species of the genus (*C. patagonicum*, *C. paraanaense*, *C. aff. orientalis*, and *C. chasicoense*; Fig. 7: node C) share a shallow h2i on the occlusal surface of p4 (ch. 88) and a normal development of c3 on p4 (ch. 99), while *C. calingastaense* sp. nov. and the largest capybaras have a deep h2i on p4 and a long c3 on p4. The clade that includes *C. paraanaense*, *C. aff. orientalis* and *C. chasicoense* (Fig. 7: node D) shares hti extended up to 50% of the prism on m1 (ch. 110) and m2 (ch. 111), while in *C. calingastaense* sp. nov. and *C. patagonicum* the hti on m1–m2 is crossing the prism but not splitting as in *Phugatherium* and *Hydrochoeropsis*; in *Neochoerus* and *Hydrochoerus*, the hti on m2 is crossing and dividing the prism. Finally, the clade formed by *C. aff. orientalis* and *C. chasicoense* (Fig. 7: node E) is supported by a shallow hsi in complex m3 (ch. 125) and a laminar prl in complex m3 (ch. 126). The other species of *Cardiatherium* present a deep hsi in complex m3.

The most parsimonious hypothesis places *C. calingastaense* sp. nov. as the sister group of a clade including all the other species of the genus because it shares some characters with the largest hydrochoerines (ch. 88[1], 99[2], 110[1], 111[1]). When *C. chasicoense* is forced as the earliest branching species of the genus, the most parsimonious hypothesis costs three more steps.

The analysis with stratigraphic adjustment (Fig. 7) suggests that the lineage leading to the clade *Cardiatherium* + largest capybaras (*Hydrochoeropsis*, *Phugatherium*, *Neochoerus*, and *Hydrochoerus*) would have originated at least during the Chasicoan SALMA, in view of the phylogenetic position of *C. chasicoense*. The evolutionary novelties such as the increasing complexity of molariforms (e.g., presence of h2.i, h5.i, h2.e, c3 on p4, presence of hse on m1–m2, addition of laminar prisms in M3) would have been achieved during the Chasicoan SALMA. In turn, the novel traits that characterize the clade including the largest capybaras (e.g., deepening of flexi/flexids splitting transversally the prisms) would have been acquired at some point in time between the Chasicoan and Montehermosan SALMAs, suggested by the presence of an extensive ghost lineage (Fig. 7; see Pérez et al. 2018 for discussion on basal forms of “cardiomyines”).

Concluding remarks

The specimen INGEO-PV 87 is included in the genus *Cardiatherium* on the basis of cranial and dental characteristics, as detailed above, some of the latter being close to *Cardiatherium patagonicum*; however, this specimen shows an exclusive combination of both cranial and dental features that leads to propose the new taxon *Cardiatherium calingastaense* sp. nov.

Concerning the skull, *C. calingastaense* sp. nov. is closer to *C. paranense* than to more derived capybaras, but differs from the former in: nasal dorsally slightly convex (strongly bulging in *C. paranense*); posterior margin of the nasal slightly concave anteriorly (strongly concave in *C. paranense*); the scar for the insertion of the masseteric muscle with two differentiated areas (single scar in *C. paranense* and other species); and absence of bulge at the base of P4 (present in *C. paranense*).

With respect to the dentition, *C. calingastaense* sp. nov. is characterized by the great depth of flexi/flexids, which makes it different from most of the species of *Cardiatherium*. Vucetich et al. (2005) concluded that deeper flexi/flexids correlate with larger size, which, in turn, is directly related to age. The great depth observed in flexi/flexids of INGEO-PV 87, resulting in a particular morphology with a step-shaped profile, suggests that it would be an adult individual. In addition, the bifurcation of flexi/flexids has been also observed in adult (large) specimens of *Cardiatherium* (e.g., p4, MPEF 740/34, *C. patagonicum*; m2, MLP 40-XI-15-1, *C. paranense*; m3, MMH-CH 88-6-41 and MLP 55-IV-28-15, *C. chasicoense*), as well as in other hydrochoerines (m3, MNHN-AYO 226, *Phugatherium saavedrai*; p4, MLP 15-232, *P. cataclisticum*) and some cardiomyines (*Caviodon cuyano*) (Vucetich et al. 2011, 2014a, b).

Some characters observed in *C. calingastaense* sp. nov. resemble the molariforms of Ploiocene capybaras (such as *Phugatherium* or *Hydrochoeropsis*), although the latter are different in many others (see above). One of those characters is the deep HSE slightly oriented backwards and another one is the c3, although this column is present in all the species of *Cardiatherium* (see Deschamps et al. 2013: fig. 4),
in *C. calingastaense* sp. nov. the c3 is very conspicuous, delimited by very deep h2i and h3i.

A particular character of the studied specimen is the conspicuous labial column in the posterior ramus of prIIa in m3, which is not known in the other species of *Cardiatherium*. Even though molariforms of hydrochoerines have strong individual variability (Mones 1991), this feature of m3 has not been previously described. Only more material from Puchuzum could reveal the existence of variation on this character within *C. calingastaense* sp. nov.

The phylogenetic analysis supports the taxonomic proposal of a new species of *Cardiatherium* and places *C. calingastaense* sp. nov. as the sister group of a clade including all the other known species of the genus.

*Cardiatherium calingastaense* sp. nov. increases the known diversity of capybaras in the late Miocene of San Juan Province. Up to now, the only record of capybaras from this area was that of *C. chasicoense* (the oldest capybara) from the lower levels of Loma de Las Tapias Formation, corresponding to the Chasicoan SALMA. At the same time, the species *C. paranense* had been recognized in the near Mendoza Province, in levels of Huayquerian age. Therefore, the new taxon corresponds to the first record of capybaras in the Huayquerian SALMA of San Juan Province and increases the general diversity of this group in the late Miocene of central-west Argentina.

Acknowledgements

The specimen INGEO-PV 87 was collected by Mariano Hidalgo (San Juan, Argentina) during a field season at Puchuzum within the project CICT-UNSJ 21E/649, Universidad Nacional de San Juan. Special thanks are due to Paula Villagra (Mendoza, Argentina) for taking photographs of the fossil; Natalia Monaco (Centro Científico Tecnológico CONICET, Mendoza, Argentina) prepared Fig. 1, and Santiago Hernández Del Pino (Instituto Argentino de Nivología, Glaciología y Ciencias Ambientales, Mendoza, Argentina) helped with Figs. 3 and 4. The authors thank the editor, Olivier Lambert (Royal Belgian Institute of Natural Sciences, Brussels, Belgium) and reviewers, Alvaro Mones (Augsburg, Germany) and an anonymous reviewer, for their useful comments on the manuscript. This paper contributes to the Research Project CICTCA-UNSJ, Programación 2018–2019: Estudio bio-cronoestratigráfico y correlación regional de las unidades neógenas de Loma de La Dehesa, valle de Matagusano, San Juan, Argentina (V.H. Contreras).

References


Appendix 1

List of revised material of *Cardiatherium*.

*Cardiatherium chasicoense*: MMP 300-M (holotype of *Procardiatherium chasicoense*), fragment of right mandible with p4–m3; MMH-CH 85-4-40, almost complete right mandible with i1, p4, m1, and m3; Las Barrancas Member (= FSC facies) of the Arroyo Chasicó Formation, Buenos Aires Province, Argentina; PV-SJ-537, palate fragment with right and left P4–M3 and associated mandible with p4–m3; Arenisca Albardón Member of the Loma de Las Tapias Formation, San Juan Province, Argentina.


*Cardiatherium orientalis*: SPV-FHC 27-XI-64-20 (holotype of *Kiyutherium orientalis*), right mandible with p4–m3; San Pedro Member (or Kiyú Lithofacies) of the Camacho Formation (see Perea et al., 2013); Barrancas de San Gregorio, San José, Uruguay.

*Cardiatherium aff. orientalis*: GHUNLPam 139, damaged skull fragment with both M1–3; GHUNLPam 2009, skull fragment with both P4–M3; GHUNLPam 5236, anterior fragment of palate with anterior root of the zygomatic arch, both P4 and left M1; GHUNLPam 5274; skull with both P4–M3, rostrum, right orbital area and damaged skull roof; GHUNLPam 8978, small fragment of palate with left M1–2; GHUNLPam 14452, palate with both P4–M3 and posterior portion of rostrum; GHUNLPam 14661, distorted skull with both P4–M3; GHUNLPam 14985, damaged rostrum with right incisor, right P4–M2 and left P4; GHUNLPam 27389, skull with both P4–M3, incisors, a left mandible fragment with p4–m3, and six cervical vertebrae articulated to skull; MLP 62-XII-4-17, left mandible fragment with p4–m3 and palatal fragment with left P4–M3 and right P4–M1 (the palatal fragment is currently lost).

*Cardiatherium rosendoi*: MLP 29-X-8-26 (holotype and single specimen of *Kiyutherium rosendoi*), left mandibular fragment with p4, m1 and part of m2; Puerta de Corral Quemado, Andalhuala Formation (“Araucanian”, late Miocene), Catamarca Province, Argentina.

*Cardiatherium patagonicum*: MPEF-PV 740/1, holotype, right mandibular fragment with m1–m2; MPEF-PV 740/2, isolated right m1 or m2; MPEF-PV 740/9, left p4; MPEF-PV 740/11, incomplete left p4; and MPEF-PV 740/24, isolated right m1 or m2; Estancia Rincón Chico, upper levels of the Puerto Madryn Formation, Peninsula Valdés, Chubut Province. MPEF-PV 2521, complete skull; La Pastosa, Puerto Madryn Formation, Peninsula Valdés, Chubut Province (data after Dozo et al. 2010).