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1 **SYSTEMATIC REVISION AND EVOLUTIONARY HISTORY OF *ACARECHIMYS***
2 **PATTERSON IN KRAGLIEVICH, 1965 (RODENTIA, CAVIOMORPHA,**
3 **OCTODONTOIDEA)**

4
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21 53 pages, 7 figures, 3 tables, 3 Supplementary Appendices

22 ARNAL *ET AL.*: REVISION AND HISTORY OF *ACARECHIMYS* (RODENTIA,
23 HYSTRICOGNATHI).

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25 **Abstract.** The octodontoid rodent *Acarechimys* was abundant during the early Miocene and
26 had the widest temporal and geographic distribution of any extinct caviomorph. Despite
27 this extensive fossil record *Acarechimys* has not been well characterized. In this work, we
28 systematically revise *Acarechimys*, describe new early–middle Miocene fossils from
29 Argentina and Bolivia, corroborate its monophyly, and study its evolutionary history.
30 *Acarechimys* has brachydont molars, retained deciduous premolars, four crests on upper
31 molars, lowers with variably developed mesolophid and metalophulid II, and absence of
32 mental foramen in the mandible. *Acarechimys* includes: *Acarechimys leucotheae* (late
33 Oligocene, Chubut, Argentina), *A. gracilis* and *A. constans* (early Miocene, Chubut and
34 Santa Cruz, Argentina), and *A. minutus* and *A. minutissimus* (early–middle Miocene of
35 Patagonia Argentina, Bolivia, and Colombia). The temporal and geographic distributions
36 suggest that *Acarechimys* could have evolved in Patagonia, by the early late Oligocene. Its
37 acme was during the late early Miocene in Southern Patagonia. By the middle Miocene,
38 *Acarechimys* decreased in diversity and was last recorded in high latitudes of South
39 America (Patagonia). In lower latitudes, the oldest record is from the late early Miocene of
40 Chucal, northern Chile, and during the late middle Miocene, the genus is recorded in
41 localities of Colombia, Bolivia, and Peru. The available evidence suggests that
42 *Acarechimys* was probably not present in lower latitudes (N of ~ 30° S) before the early
43 Miocene. The reasons *Acarechimys* dispersed northward at this time remain to be
44 elucidated, but the timing coincides with a massive disappearance of other octodontoids
45 from Patagonia.

46 **Key Words.** Octodontoid. Oligocene. Miocene. South America. Evolutionary history.

47

48

49 **Resumen. REVISIÓN SISTEMÁTICA E HISTORIA EVOLUTIVA DE**

50 **ACARECHIMYS PATTERSON EN KRAGLIEVICH, 1965 (RODENTIA,**

51 **CAVIOMORPHA, OCTODONTOIDEA).** El roedor octodontoideo *Acarechimys* fue

52 abundante durante el Mioceno temprano y tuvo la distribución geográfica y temporal más

53 amplia para un caviomorfo viviente. A pesar de su amplio registro fósil *Acarechimys* nunca

54 fue caracterizado correctamente. En este trabajo, realizamos la revisión sistemática de

55 *Acarechimys*, describimos nuevos materiales del Mioceno temprano–medio de Argentina y

56 Bolivia, corroboramos su monofilia y estudiamos su historia evolutiva. *Acarechimys*

57 presenta dientes braquiodontes, retención de premolares deciduos, cuatro crestas en molares

58 superiores, desarrollo variable del mesolófido y el metalofúlido II en molares inferiores y

59 ausencia de foramen mentoniano en la mandíbula. *Acarechimys* incluye: *Acarechimys*

60 *leucotheae* (Oligoceno tardío, Chubut, Argentina), *A. gracilis* y *A. constans* (Mioceno

61 temprano, Chubut y Santa Cruz, Argentina), y *A. minutus* y *A. minutissimus* (Mioceno

62 temprano–medio de Patagonia Argentina, Bolivia y Colombia). Su distribución temporal y

63 geográfica sugiere que *Acarechimys* habría evolucionado en Patagonia en el Oligoceno

64 tardío-temprano. Su acmé fue en el Mioceno temprano-tardío en el sur de Patagonia. Para el

65 Mioceno medio *Acarechimys* disminuyó su diversidad y tiene su último registro en

66 latitudes altas de América del Sur (Patagonia). En latitudes bajas, el registro más antiguo

67 proviene del Mioceno temprano-tardío de Chucal, norte de Chile, y durante el Mioceno

68 medio se lo registra en localidades de Colombia, Bolivia y Perú. La evidencia disponible

69 sugiere que *Acarechimys* probablemente no estuvo presente en bajas latitudes (N de

70 30°S) antes del Mioceno temprano. Las causas de su dispersión hacia el norte deben ser

71 todavía estudiadas, aunque la misma coincide con la desaparición masiva de
72 octodontoideos en Patagonia.

73

74 **Palabras clave.** Octodontoideo. Oligoceno. Mioceno. América del Sur. Historia evolutiva.

75

76

77 ACARECHIMYS is an extinct rodent genus, part of the richest and most diverse clade of
78 caviomorphs: Octodontoidea. It represents a successful evolutionary lineage with unusually
79 wide temporal (late Oligocene–late Miocene) and geographic distributions (southern
80 Argentinean Patagonia to Colombia); it was one of the most abundant octodontoids during
81 the Burdigalian (late early Miocene; Santacrucian South American Land Mammal Age,
82 SALMA; Pascual *et al.*, 1965) of Santa Cruz Province, Argentina.

83 Ameghino (1887), while working at the Museo de La Plata (MLP), erected the
84 genera *Acaremys*, *Stichomys*, and *Sciamys*, and recognized the species: *Acaremys minutus*,
85 *A. minutissimus*, *Sciamys tenuissimus*, and *Stichomys constans*. Subsequently, he named
86 additional species of these genera (Ameghino, 1889, 1891, 1894). Scott (1905) performed
87 an exhaustive revision of the rodents from the Santa Cruz Formation based on fossils
88 collected by the Princeton Expeditions to Patagonia (1896–1899; housed at the Yale
89 Peabody Museum, New Haven, USA) and the collections housed in the museums of
90 Buenos Aires and La Plata that he visited in 1902. As a result, he described additional
91 materials and performed detailed descriptions of *Acaremys minutus*, *A. minutissimus*,
92 *Sciamys tenuissimus*, *Stichomys constans*, and *Stichomys diminutus*. Bryan Patterson
93 studied the paleontology collections deposited at the MLP and MACN in the years 1952–
94 1954 (Olson, 1985) and performed an exhaustive investigation of caviomorphs that was
95 never published. Nevertheless, in his unpublished manuscript (UMS, available at the
96 Vertebrate Paleontology Section, MACN), Patterson provided information that was later
97 used by Pascual (1967) and by the authors of this work (see below). The genus
98 *Acarechimys* was first mentioned by Kraglievich (1965) in a footnote, explaining: ‘*Ce nom,*
99 *inédit, a été appliqué par B. Patterson à l’espèce Acaremys minutus Amegh. (Patterson et*

100 *Kraglievich, ms.*)'. Later, based on Patterson UMS, Pascual (1967) mentioned the genus
101 *Acarechimys* with the species *Acarechimys minutus*, *Acarechimys minutissimus*, and
102 *Acarechimys constans*, and provided collection numbers of the lectotype of each species.
103 Finally, Patterson (in Patterson and Wood, 1982) characterized the genus *Acarechimys* for
104 the Santacrucian SALMA of Patagonia, with *Acarechimys minutus* as the type species
105 (synonym: *Stichomys gracilis* Ameghino, 1891) and *Acarechimys minutissimus* as the only
106 other referred species (synonyms: *Stichomys diminutus* Ameghino, 1891, *Sciamys*
107 *tenuissimus* Ameghino, 1894, and provisionally *Stichomys constans* Ameghino, 1887).
108 Since the 1990s, the genus *Acarechimys* has been recognized at numerous localities beyond
109 Santa Cruz Province: Vucetich *et al.* (1993a) extended its biochron by describing
110 *Acarechimys* sp. from the Langhian (early middle Miocene, Colloncuran SALMA) of
111 Neuquén Province, Argentinean Patagonia; Walton (1997) identified *Acarechimys* cf. *A.*
112 *minutissimus* from the Serravallian (late middle Miocene, Laventan SALMA) of La Venta,
113 Colombia, and Croft *et al.* (2011) described new specimens of *Acarechimys* from the
114 Serravallian (late middle Miocene, Laventan SALMA) of Quebrada Honda, Bolivia. Flynn
115 *et al.* (2002, 2008) and Croft *et al.* (2007) mentioned *Acarechimys* for the early to middle
116 Miocene of Chile, Antoine *et al.* (2016) for the early Miocene of Contamana, Peru, Tejada-
117 Lara *et al.* (2015) for middle Miocene of the Fitzcarrald Arch in the Peruvian Amazonia,
118 and Esteban *et al.* (2014) for late Miocene–early Pliocene of the Andalhuala and Chiquimil
119 formations in Catamarca Province, Argentina.

120 Vucetich *et al.* (2010) transferred the species *Protacaremys pulchellus* Ameghino,
121 1902 to *Acarechimys*, erecting the new combination *Acarechimys pulchellus* (Ameghino,
122 1902) for the Aquitanian–Burdigalian (early Miocene, Colhuehuapian SALMA) of
123 Patagonia. This taxonomic assignation was corroborated later by Arnal (2012) with a

147 Autónoma Tomás Frías, Vertebrate Paleontology Collection, Potosí, Bolivia; **YPM-VPPU**,
148 Princeton University Collection of the Yale Peabody Museum, New Haven, USA.

149 ***Systematic revision.*** Recognition of the type series used by Florentino Ameghino (1887,
150 1889) was necessary to identify the specimens recovered by Carlos Ameghino during his
151 filedtrip of 1887 in the cliffs of the Santa Cruz River, Santa Cruz province, Argentina (Fig.
152 2). In performing this investigation, we took into account different sources of information:
153 Ameghino’s catalog at the MACN, the MLP catalog, Patterson’s UMS, collection labels
154 and file cards, and the account of the history of the conflict between F. Ameghino and
155 Moreno when the Santacrucian species were erected (Fericola, 2011). Patterson studied
156 the paleontology collections deposited at the MLP and MACN, but identified lectotypes
157 and syntypes of fossils housed at the MLP [information that was published by Pascual
158 (1967)]. Patterson mentioned and studied specimens figured by Ameghino (1889) and
159 deposited at the MACN but dismissed the possibility that they might be holotypes because
160 he believed that the fossils collected by C. Ameghino in 1887 were deposited exclusively at
161 the MLP (UMS, pers, comm.). Fericola (2011) determined that those fossils figured in the
162 Atlas by Ameghino (1889) and housed in the MACN were collected by his brother Carlos
163 in 1887, and therefore these fossils could be part of the type series of taxa founded in 1887
164 by F. Ameghino. Additionally, we have inferred that Patterson may have had access to
165 labels with Ameghino’s handwriting and other certain provenance information, since his
166 UMS stresses differences among ‘labeled materials’, ‘improperly labeled materials’, and
167 ‘materials that agree with Ameghino’s original descriptions’. Thus, with some exceptions
168 (see below), we accept the information provided by Patterson in his unpublished
169 manuscript and discuss the implications for each particular species.

170 **Statistical analyses.** In order to test for size differences among *Acarechimys* species, we
171 performed statistical analyses using m1 and m2 length, as these allowed for the largest
172 sample sizes. Sixty-nine m1 and 62 m2 measurements were used in the analyses. Statistical
173 analyses were performed using JMP Pro (SAS Institute, Inc., 2013).

174 **Cladistic analysis.** In order to test the monophyly of the genus, we used an expanded and
175 modified version of the data matrix of Arnal and Vucetich (2015) (Supplementary
176 appendices 2 and 3). In total, it consists of 186 morphological characters and 59 taxa. These
177 include 19 morphological characters from Verzi *et al.* (2016) that were added in order to
178 evaluate comparable data matrices. The living *Abrocoma* was included in order to test
179 whether some *Acarechimys* species group within Abrocomidae. The data matrix was
180 analyzed under equally weighted parsimony using TNT 1.5 (Goloboff and Catalano, 2016).
181 A heuristic search of 1,000 replications of Wagner trees (with random addition sequence)
182 followed by tree bisection and reconnection (TBR) branch-swapping algorithm (holding 10
183 trees per replicate) was conducted. The best trees obtained at the end of the replicates were
184 subjected to a final round of TBR branch swapping. Thirty-one characters were treated as
185 ordered. The robustness of the obtained MPTs was calculated with relative and absolute
186 Bremer supports (Bremer, 1994; Goloboff and Farris, 2001).

187

188 **SYSTEMATIC PALEONTOLOGY**

189 **Order RODENTIA** Bodwich, 1821

190 **Suborder HYSTRICOGNATHI** Tullberg, 1899

191 **Parvorder CAVIOMORPHA** Wood, 1955

192 **Superfamily OCTODONTOIDEA** Waterhouse, 1839

193 **Genus *Acarechimys*** Patterson in Kraglievich, 1965

194 **Type species.** *Acaremys minutus* Ameghino, 1887.

195 *Stichomys* Ameghino, 1887 *partim*

196 *Sciamys* Ameghino, 1887 *partim*

197 *Ameghinomys* Verzi *et al.*, 2016

198

199 **Stratigraphic and geographic occurrences.** Upper levels of the Sarmiento Formation at
200 Cabeza Blanca (Chubut, Argentina), late Oligocene, Deseadan SALMA (Vucetich *et al.*,
201 2015 a); Sarmiento Formation (Chubut, Argentina), early Miocene, Colhuehuapian
202 SALMA (Vucetich *et al.*, 2010); Pinturas Formation (Santa Cruz, Argentina), late early
203 Miocene, ‘Pinturan’ age (Kramarz and Bellosi, 2005); Santa Cruz Formation, (Santa Cruz,
204 Argentina), late early Miocene, Santacrucian SALMA (Ameghino, 1887, 1889; Scott,
205 1905; Vizcaíno *et al.*, 2012); Chucal Formation (Región XV, Chile), late early Miocene,
206 Santacrucian SALMA (Croft *et al.*, 2007); unnamed formation (Pampa Castillo, Región XI,
207 Chile), late early Miocene, Santacrucian SALMA (Flynn *et al.*, 2002); Curá Mallín
208 Formation (Región VIII, Laguna del Laja, Chile), early to middle Miocene (Flynn *et al.*,
209 2008); Collon Curá Formation (Neuquén, Argentina), early middle Miocene, Colloncuran
210 SALMA (Vucetich *et al.*, 1993a); Villavieja Formation (Colombia), late middle Miocene,
211 Laventan SALMA (Walton, 1997); unnamed formation (Quebrada Honda, Bolivia), late
212 middle Miocene, Laventan SALMA (Croft *et al.*, 2011); unspecified formation, Fitzcarrald
213 Arch (Peru), middle Miocene, Laventan SALMA (Negri *et al.*, 2010; Tejada-Lara *et al.*,
214 2015); Pebas Formation, Contamana (Peru), late? Miocene (Antoine *et al.*, 2016).

215

216

[FIGURE 2 ABOUT HERE]

217

218 ***Emended diagnosis*** [from Vucetich *et al.* (2015a); autapomorphies marked with an
219 **asterisk**]. Small to very small sized caviomorph. Brachydont cheek teeth* (see
220 Phylogenetic Analysis, below). Clearly evident cusps and thin crests separated by wide
221 flexi/ids. Retention of deciduous premolars throughout life. Upper molars with four straight
222 crests (anteroloph, protoloph, metaloph, and posteroloph); antero- and posteroloph fused
223 labially in juveniles with proto- and metaloph delimiting the antero- and metafossette,
224 respectively. Lower molars with variably developed metalophulid II: interrupted or absent
225 on m1-m2, absent or interrupted on m3; presence of accessory cusp on lowers molars*.
226 Lower deciduous premolar with variably developed metalophulid II and mesolophid united
227 with or near the metaconid. Lower incisor long, with its posterior end located posterior to
228 m3. Well-developed masseteric crest of the mandible, mental foramen generally absent, and
229 masseteric fossa moderately to very deep anteriorly. Skull with a moderately developed
230 groove for the passage of the nerve infraorbitalis *.

231
232 *Acarechimys minutus* (Ameghino, 1887)

233 Figure 3.1–5

234

235 *Acaremys minutus* Ameghino, 1887

236 *Ameghinomys constans* Verzi *et al.*, 2016 *partim*

237

238 ***Type series.*** MLP 15-410a, left mandible with m1-3; MACN-A 237, right mandible with
239 dp4-m3; MACN-A 238, right mandible with dp4-m3; MACN-A 4075, right mandible with
240 dp4-m1.

241 ***Lectotype.*** MLP 15-410a Patterson (UMS, pers, comm.) and Pascual, 1967.

242 **Paralectotypes.** MACN-A 237, MACN-A 238, MACN-A 4075.

243 **Referred material.**(Supplementary appendix 1).

244 **Geographic and stratigraphic occurrences.** Santa Cruz and Neuquén provinces, Argentina;
245 Quebrada Honda, Bolivia. Santa Cruz Formation, late early Miocene, Santacrucian
246 SALMA; Collon Curá Formation, early middle Miocene, Colloncuran SALMA; unnamed
247 formation, late middle Miocene, Laventan SALMA (Supplementary appendix 1).

248 **Emended diagnosis.** Smaller than *A. constans* and *A. gracilis*. Upper deciduous premolar
249 tending toward reduction and loss of the metaloph, unlike *A. constans*. Lower deciduous
250 premolar with variably developed metalophulid II and mesolophid, a combination not
251 present in the remaining species. Lower molars with metalophulid II reduced to a small
252 spur on m1-2 and absent on m3, as in *A. leucotheae*; presence of accessory cups in m1-2 and
253 variable presence of the posterior extension of the metalophulid I in m1-2; presence of
254 posterior arm of the metaconid on m1-3, as in *A. gracilis*. Lower incisors laterally
255 compressed, unlike *A. constans* and *A. gracilis*. Mandible with the notch for the insertion
256 pars infraorbitalis of the masseter muscle (nmmpio) located below dp4-m1, unlike *A.*
257 *constans* and *A. gracilis*.

258 **Remarks.** MLP 15-410a (left mandible with m1-m3) is labeled as ‘lectotype’ and MLP 15-
259 410b (maxilla with right and left M1-M3) as ‘type?’ of *Acarechimys minutus* in the MLP
260 collection. Based on the catalog information of this museum, we confirm that both
261 specimens belong to the ‘old collections’, which include specimens from the expedition of
262 C. Ameghino in 1887 and expeditions performed by other staff of the MLP in subsequent
263 years (Vizcaíno *et al.*, 2013). Unfortunately, the available information does not allow us to
264 know which of these fossils were recovered by C. Ameghino (no labels or catalog written
265 by Ameghino are available). However, according to the unpublished work of Patterson,

266 MLP 15-410a is the lectotype of *Acarechimys minutus* since ‘in fact the only specimen
267 labeled as *minutus* or attributable to the species is M.L.P. n°15-410a, a portion of left ramus
268 with base of I, roots of dm4, m1-3. This individual thus becomes the lectotype.’ Thus,
269 based on the confirmation made by Patterson, we interpret that MLP 15-410a could have
270 had a label made by F. Ameghino. On the contrary, we think that MLP 15-410b was not
271 available for Patterson in the 1950s, since in his UMS he stated that the palatal fragment
272 described by Ameghino for this species was lost. Only later was MLP 15-410b relocated
273 and available for study in the MLP collections. We believe that this took place after the
274 accession of fossils belonging to old assemblages into the MLP collections, an occurrence
275 that has taken place many times since the 1960s (Reguero and Tonni, pers. comm., 2016).
276 The specimen file card of MLP 15-410b includes a note by R. Pascual that it possibly
277 corresponds to the palatal fragment used by Ameghino in 1889 for the description of
278 *Acaremys minutus* (handwriting by Pascual). We agree with Pascual that this specimen
279 matches Ameghino’s description; nevertheless, there is no evidence that it was part of the
280 collection studied by Ameghino. Therefore, we have not included MLP 15-410b among the
281 syntypes of *Acarechimys minutus*. We recognize three additional specimens housed at the
282 MACN collection as part of the original type series: MACN-A238 and MACN-A 4075,
283 which are illustrated in the atlas of Ameghino (Ameghino, 1889; plate IV, figs. 22–23), and
284 MACN-A 237, which is mentioned in the catalog of Ameghino (MACN) as being a
285 member of the same stock as MACN-A238. In summary, we identify four fossils used by
286 F. Ameghino to erect the species *Acaremys minutus*: MLP 15-410a, MACN-A237, MACN-
287 A 238, and MACN-A 4075.

288 Pascual (1967) stated that MLP 15-410a was the lectotype of *Acarechimys minutus*.
289 Although this material corresponds to an old specimen with the occlusal surface very worn,

290 we agree with Patterson's assignation. By default, the remaining specimens of the original
291 type series are paralectotypes.

292 Based on dental morphology (symmetric absence of mure on the M2 delimiting an
293 antero- and posterofossette and a relatively large and fully tetralophodont M3) we can not
294 assign MLP 15-410b to any recognized caviomorph species. It probably represents a new
295 octodontoid (Arnal and Vucetich, personal observation) and will be the subject of future
296 investigations.

297 **Description.** *Upper cheek teeth.* Both tooth rows are parallel to each other but are obliquely
298 implanted with respect to the palatal plane.

299 The upper cheek teeth are wider than long (Tab. 1), slightly terraced, and
300 tetralophodont. Molars have a long anteroloph that reaches the paracone and delimits an
301 anterofossette in juvenile specimens (Fig. 3.1). The protoloph is anteriorly oblique and
302 curved. The metaloph departs from the junction of the mure and the anterior arm of the
303 hypocone, and its labial end, which includes the metacone, turns back to fuse with the
304 posteroloph, delimiting a metafossette in juveniles (Fig. 3.1). The M3 has a labially placed
305 hypocone, and a hypofossette is formed in adult specimens (Fig. 3.1). The anterofossette
306 and metafossette are equally deep and extend further across the occlusal surface than the
307 mesoflexus. The hypoflexus is the deepest flexus and is anteriorly oriented.

308 The DP4 is molarized and resembles the molars but usually differs in the presence
309 of a reduced metaloph that does not reach the labial end of the posteroloph, by the presence
310 of a short protoloph that does not reaches the paracone, and by a less oblique protoloph and
311 hypoflexus (Fig. 3.1).

312 Upper incisors are oval in section. The anterior face is straight and the lingual face
313 is curved.

314 *Skull*. The description is based on an almost complete skull (YPM-VPPU 15806; Fig. 3.2)
315 and maxillary fragments (MACN-A 4070, MPM-PV 15088). The nasals extend posteriorly
316 to the dorsal root of the zygoma. Posteriorly, the frontal bones have concave lateral
317 margins. The premaxillae occupy the anterior half of the lateral wall of the snout (Fig. 3.2);
318 the ascending processes of the premaxillae are slightly exposed on the skull roof and are a
319 little longer than the nasals, unlike in *Acaremys murinus*, in which they are broadly exposed
320 on the skull roof. The rostral masseteric fossa (*sensu* Patterson and Wood, 1982) is shallow,
321 subtriangular, and limited ventrally by the incisor tuberosity (Fig. 3.2), unlike in *Acaremys*
322 *murinus*, *Pseudoacaremys kramarzi*, and *Sciamys principalis*, in which the incisor
323 tuberosity is included in the rostral masseteric fossa (Arnal and Vucetich, 2015). The dorsal
324 root of the zygoma continues ventrally with a robust vertical ramus of the zygoma, similar
325 to *Sciamys principalis* (Fig. 3.2). This vertical ramus is mainly straight, as in *Prospaniomys*
326 *priscus*, rather than posteriorly oblique as in most octodontoids (Arnal and Kramarz, 2011).
327 The ventral root of the zygoma extends slightly in front of the DP4 (Fig. 3.2), and its
328 antero-posterior diameter is similar to its dorso-ventral diameter, unlike *Pseudoacaremys*
329 *kramarzi*, where the antero-posterior diameter is twice the dorso-ventral one (Arnal and
330 Vucetich, 2015). In ventral view, the masseteric tuberosity (for the origin of the masseter
331 superficialis muscle, pars anterior) is conspicuous and continuous laterally with a shallow
332 fossa for the origin of the masseter lateralis muscle. Posterior to the masseteric tuberosity is
333 a small foramen of uncertain homology. On the dorsal face of the ventral root of the
334 zygoma is a faint furrow for the passage of the infraorbitalis nerve. The horizontal ramus of
335 the zygoma is high in lateral view (Fig. 3.2), unlike *Prospaniomys priscus* in which it is low
336 (Arnal and Kramarz, 2011). It is formed mainly by the jugal bone, which lies at the base of
337 the vertical ramus of the zygoma along with the maxillary bone (Fig. 3.2); this suture is

338 straight and oblique. The paraorbital process is conspicuous and formed by the jugal and a
339 small portion of the squamosal. The jugal fossa (for the origin of the posterior masseter
340 muscle) is well-developed; it is antero-posteriorly long, dorso-ventrally high, and medio-
341 laterally deep, unlike *Prospaniomys priscus*, in which it is short and shallow.

342 The diastema is longer than the tooth row and widens posteriorly but is shorter than
343 in *Acaremys murinus* and *Pseudoacaremys kramarzi*. The large incisive foramina are
344 damaged but seem to be wider than long. Posteriorly, they are continuous with the well-
345 developed diastemal furrows that extend to the DP4s. The posterior palatine foramina are
346 conspicuous and located between the M1s. The openings of the posterior nares are
347 positioned near the posterior half of the M2.

348 *Lower cheek teeth.* The molars are of similar size (Tab. 1). The dp4 is longer than wide
349 (Fig. 3.3; Tab. 1). The metalophulid I is curved and joins the protoconid and metaconid.
350 The ectolophid is relatively short and oblique. A short but variably developed metalophulid
351 II extends from the postero-lingual border of the protoconid (Figs. 3.3–4) as in some
352 specimens of *A. minutissimus* and *A. constans*. The mesolophid is also variably developed;
353 it varies from long and fused with the metaconid (MPM-PV 15089), as in *A. constans*, to
354 reduced, forming a spur of the ectolophid, as in *A. gracilis* (Fig. 3.3). The hypolophid is
355 straight and merges with the entoconid. The posterolophid is long and reaches the labial
356 side of the tooth. The anterofossettid/anteroflexid is rounded, shallow, and merged with the
357 mesoflexid when the mesolophid is reduced (Figs. 3.3–4). The posteroflexid is closed in
358 adult specimens. The hypoflexid is the deepest flexid and is posteriorly oblique. In adult
359 specimens, the dp4 becomes simplified with an oval outline (MACN-A 4071).

360 The molars have three main crests; a fourth crest, second in position (metalophulid
361 II), is variably developed (Fig. 3.3). The metalophulid I is straight, unlike in *A. constans*

362 and *A. gracilis*, where it is curved. The metalophulid II is reduced to a spur of the
363 ectolophid near the area of the protoconid in m1-2 and is absent in m3 (Fig. 3.3). An
364 accessory cusp of uncertain homologies is lingually aligned with this crest and often united
365 to the metalophulid I by a posterior extension of the metalophulid I (Fig. 3.3). The posterior
366 arm of the metaconid is present in m1-3. The hypolophid and posterolophid are as in dp4.
367 The lingual end of the posterolophid lies near the entoconid, unlike in the dp4. The
368 antero+mesoflexid is as broad and deep as the posteroflexid.

369 The lower incisors are laterally compressed and long, extending below the tooth row
370 to the base of the coronoid process.

371 *Mandible.* The mandible is robust. The diastema is shorter than the tooth row, unlike the
372 condition in the cranium. The diastema is concave, and the lowest part is immediately
373 anterior to the dp4 (Fig. 3.5). A mental foramen is generally absent; when present, it is
374 small and located below the anterior half of dp4. The nmmpio is conspicuous, antero-
375 posteriorly long, slightly oblique, and the anterior half is located mainly below the dp4
376 (Fig. 3.5), as in *A. minutissimus*. The masseteric crest is well-developed, laterally
377 projecting, and ventro-posteriorly oriented (Fig. 3.5). The masseteric fossa is deep in its
378 anterior part as in *A. gracilis* but shallower than in species of *Acaremys*. This fossa is
379 antero-dorsally limited by a faint furrow that extends from the posterior border of the
380 nmmpio to the base of the coronoid process. The coronoid process extends postero-laterally
381 from the m2 and delimits a retromolar fossa lateral to m3. On the lingual side of the
382 mandible, the mandibular symphysis extends posteriorly to the posterior border of m1. The
383 mandibular chin is at the level of the dp4.

384

385

[FIGURE 3 ABOUT HERE]

386

387 *Acarechimys constans* (Ameghino, 1887)

388 Figure 4. 1–4

389 *Stichomys constans* Ameghino, 1887

390 *Acarechimys pascuali* Verzi *et al.*, 2016 *partim*

391 *Ameghinomys constans* Verzi *et al.*, 2016 *partim*

392

393 **Type series.** MACN-A 246, palatal fragment with left and right DP4-M2; MACN-A 247,
394 left mandible with m1-2 and right mandible with dp4-m2; MLP 15-39, left mandible with
395 m1-3; MLP 15-57, right mandible with m2-3 and broken m1; MLP 15-200, right mandible
396 with dp4-m2; MLP 15-346, left mandible with dp4-m2; MLP 15-391, right mandible with
397 dp4-m3; MLP 15-391a, right mandible with dp4-m2; MLP 15-391b, right mandible with
398 m1-3; MLP 15-391c, left mandible with broken dp4-m1.

399 **Lectotype.** MLP 15-391 Patterson (UMS, pers, comm.) and Pascual (1967).

400 **Paralectotypes.** MACN-A 246, MACN-A 247, MLP 15-39, MLP 15-57, MLP 15-200,
401 MLP 15-346, MLP 15-391a, MLP 15-391b, MLP 15-391c.

402 **Referred material.** (Supplementary appendix 1).

403 **Geographic and stratigraphic occurrence.** Santa Cruz Province. Santa Cruz Formation,
404 late early Miocene, Santacrucian SALMA (Supplementary appendix 1).

405 **Diagnosis.** Larger than *A. minutus*, *A. minutissimus*, and *A. leucotheae*. Lower deciduous
406 premolars with well-developed or reduced metalophulid II; mesolophid conspicuous, unlike
407 in *A. gracilis*. Lower molars with metalophulid II well-developed on m1 and reduced or
408 absent on m2-3, unlike *A. minutus* and *A. minutissimus*; accessory cusp on m1 variable
409 present and conspicuous posterior arm of the metaconid absent, unlike *A. gracilis*. Lower

410 incisors not compressed medio-laterally, unlike *A. minutus* and *A. minutissimus*. Mandible
411 with the masseteric fossa and nmpio located more posteriorly than in *A. minutus*, *A.*
412 *minutissimus*, and *A. leucothaeae*.

413 **Remarks.** We followed the same procedures as for *Acarechimys minutus* in identifying the
414 original type series used by Ameghino to describe *Stichomys constans*. In the MLP
415 collections, the specimen MLP 15-391 is catalogued as lectotype and MLP 15-391a, b, and
416 c are catalogued as syntypes of *Acarechimys constans*. Based on the MLP catalog, we have
417 confirmed that these specimens belong to the ‘old collections’ of the MLP. Patterson stated
418 that this species was based ‘on a series of incomplete mandibles in the Museo de La Plata,
419 nos. 15-39, MLP 15-57, MLP 15-200, MLP 15-346, MLP 15-391 and MLP 15-391a-d
420 (UMS, pers. comm.). Of these, 15-391 agrees most closely with the type description and is
421 therefore designated as the lectotype’. Thus, it is evident that these fossils were available to
422 Patterson, who did not hesitate in considering them as part of Ameghino’s original type
423 series. Additionally, three specimens figured in Ameghino (1889; Atlas: plate VI, figs. 6–8)
424 housed at the MACN also belong to the original type series (MACN-A 246 and MACN-A
425 247). Patterson (UMS, pers. comm.) stated that the palatal fragment described by
426 Ameghino as *Stichomys constans* (MACN-A 246; Atlas 1889: plate VI, fig. 8) should be
427 referred to *Adelphomys candidus* Ameghino, 1887, an assignation with which we agree.

428 Patterson (UMS, pers. comm.) and Pascual (1967) stated that MLP 15-391 was the
429 lectotype of *Acarechimys constans*. We agree with this assignation. By default, the
430 remaining specimens of the original type series constitute the paralectotypes. Nevertheless,
431 among the paralectotypes, MLP 15-200 has been lost since December 1978 (information
432 provided by the specimen label), and we refer MLP 15-346, MLP 15-391, and MLP 15-
433 391b to *Acarechimys gracilis* based on molar and mandibular morphology (see below).

434 Patterson (in Patterson and Wood, 1982) ‘provisionally’ considered *Stichomys*
435 *constans* a synonym of *Acarechimys minutissimus*. However, this assignation is odd, and
436 we think it could have been an error, since Patterson in his UMS considered *Acarechimys*
437 *constans* as a valid species (pers, comm.). Additionally, based on its size and dental
438 morphology, this species has traditionally been considered a distinct species (Vucetich *et*
439 *al.*, 1993 a? b?, 2010, 2015a; Kramarz, 2004; Croft *et al.*, 2011; Arnal, 2012). Recently,
440 Verzi *et al.* (2016) erected the new genus *Ameghinomys* to include this species
441 (*Ameghinomys constans*) based on materials referred to *Acarechimys minutus*, *Acarechimys*
442 *pulchellus*, *Acarechimys constans*, and other previously undescribed specimens. The
443 conclusions of our systematic revision of *Acarechimys* differ from those of this publication.
444 Verzi *et al.* (2016) associate the palatal fragment labeled as ‘type?’ (MLP 15-410b) of
445 *Acarechimys minutus* mentioned in the previous section with the lectotype of *Acarechimys*
446 *constans* (MLP 15-391). They argue that MLP 15-410b is proportionally larger than
447 *Acarechimys minutus*, that it has a short mure comparable to the short ectolophid present in
448 lower molars of *Acarechimys constans*, and that although Ameghino (1887, 1889) did not
449 mention cranial fragments for *Stichomys constans*, he figured one in 1889 (Verzi *et al.*,
450 2016). Nevertheless, the mentioned palatal fragment (Ameghino, 1889; Atlas: plate VI, fig.
451 8) is MACN-A 246 (Fericola 2011; pag. 52), and not MLP 15-410b as indicated by Verzi
452 *et al.* (2016). Ameghino (1889) did provide the following description for *Stichomys*
453 *constans*: [‘la primera muela superior tiene también tres raíces, como en la especie
454 anterior (*Stichomys regularis*); y las cuatro muelas superiores ocupan un espacio
455 longitudinal de 9 milímetros, como las cuatro inferiores’] [‘the first upper molar also has
456 three roots, as in the preceding species (*Stichomys regularis*); and the four upper cheek
457 teeth occupy a longitudinal space of 9 millimeters, like the four lowers’] (1889; pag. 247).

458 Additionally, MLP 15-410b is considerably smaller than *S. constans* [M1-3 are 6.85mm
459 long, more than 2mm smaller than the value listed for *Stichomys constans* (9mm)]. Lastly,
460 since relatively short ectolophids are common in small rodents, we do not think that the
461 superficial resemblance to a ‘short mure’ on the upper molars (absent on both M2 of MLP
462 15-410b) can be used to refer unassociated upper and lower dentitions to the same species.

463 **Description.** *Upper cheek teeth.* As in *A. minutus*, both tooth rows are parallel to each
464 other (Fig. 4.1) and are obliquely implanted labially with respect to the palatal plane. The
465 upper cheek teeth are tetralophodont and wider than long (Fig. 4.1–2; Tab. 1). All molars
466 and the DP4 have a subquadrangular occlusal outline. The molars have a short anteroloph
467 that does not reach the paracone (Fig. 4.2), unlike in *A. minutus*, where it is long and fused
468 with this crest. The protoloph is oblique anteriorly and curved. The metaloph is straight,
469 departs from the junction of the mure and the anterior arm of the hypocone, and its labial
470 end fuses with the posteroloph, delimiting a metafossette in juveniles (Fig. 4.2). The
471 anterofossette and metafossette are equally deep and penetrate the occlusal surface slightly
472 further than the mesoflexus. The hypoflexus is the deepest flexus and is posteriorly
473 oriented.

474 The DP4 has a conspicuous metaloph, unlike in *A. minutus*, in which it is usually
475 reduced (Fig. 4.2). It is fully molarized.

476 *Skull.* The description is based on MPM-PV 15002, a partial palatal fragment.

477 The ventral root of the zygoma extends slightly in front of the DP4 (Fig. 4.1), and its antero-
478 posterior diameter is similar to its dorso-ventral diameter, as in *A. minutus*. In ventral view,
479 the masseteric tuberosity is conspicuous. Posterior to the masseteric tuberosity is a small
480 foramen of uncertain homology (Fig. 4.1), as in the type species. On the dorsal face of the
481 ventral root of the zygoma is a faint furrow for the passage of the infraorbitalis nerve. In

482 palatal view, well-developed diastemal furrows are evident that extend posteriorly to
483 M1(Fig. 4.1), farther than in *A. minutus*. The posterior palatine foramina are conspicuous
484 and located between the M1s. The posterior nares open opposite the anterior half of the M2.
485 *Lower cheek teeth.* The dp4 is the longest tooth (Tab. 1). It has a curved metalophulid I. A
486 well-developed (MACN-A 4058; MACN-A 4061; MACN-A 4064; MLP 15-391) (Fig. 4.3)
487 or reduced (MACN-A 247a; MACN-A 4075) metalophulid II extends postero-lingually
488 from the protoconid. The ectolophid is oblique and extends posteriorly from the posterior
489 border of the protoconid. Near its posterior end, a well-developed mesolophid extends
490 lingually and usually reaches the metaconid (Fig. 4.3), as in *A. minutissimus*. The
491 hypolophid is straight or curved and reaches the entoconid. The posterolophid is long,
492 anteriorly concave, and does not contact the entoconid (Fig. 4.3). The anterofossettid is
493 rounded and relatively shallow, the metaflexid is narrow, and the posteroflexid is the
494 largest and deepest of the lingual flexids/fossettids. The hypoflexid is posteriorly oriented
495 and is deeper than the lingual flexids.

496 Lower molars have three main crests and a fourth, second in position, variably
497 developed (Fig. 4.3). The metalophulid I is curved, unlike in *A. minutus* and *A.*
498 *minutissimus*, where it is straight. A conspicuous metalophulid II is present in m1 but is
499 reduced or absent on m2-3 (Fig. 4.3). In some cases, a labio-lingually aligned accessory
500 cusp (Fig. 4.3) and/or a posterior extension of the metalophulid I is observed. These two
501 structures are generally reduced or absent in m2-3. The ectolophid is generally longer than
502 in the remaining species of the genus. The hypolophid and the posterolophid of the molars
503 resemble the condition in the dp4. In juveniles, the hypoflexid is confluent with the
504 posteroflexid (m2-3 in Fig. 4.3). The antero+mesoflexid is as broad and deep as the
505 posteroflexid.

506 Lower incisors are not laterally compressed, unlike in *A. minutus* and *A.*
507 *minutissimus*. They are long, with their posterior end located at the base of the coronoid
508 process.
509 *Mandible*. The mental foramen, when present, is small and located opposite the anterior
510 half of dp4 or slightly anteriorly (Fig. 4.4). The nmmpio is more poorly developed than the
511 remaining species, antero-posteriorly short, and located mainly below m1 (Fig. 4.4), as in
512 *A. gracilis*. The masseteric crest is well-developed and is continuous with the posterior end
513 of the nmmpio; it extends laterally as in the remaining species of the genus. The masseteric
514 fossa is as deep as in *A. minutus* and *A. gracilis*; it is posteriorly positioned, with its anterior
515 border opposite m2 or the posterior end of m1 (Fig. 4.4). The furrow that delimits the
516 masseteric fossa antero-dorsally is poorly developed. The base of the coronoid process
517 extends anteriorly to a point between m2 and m3 (Fig. 4.4), unlike in the remaining species
518 of the genus in which it extends to m2. The mandibular symphysis extends posteriorly to
519 the level of m1, and a poorly developed chin is present just anterior to the anterior part of
520 the dp4.

521

522 [FIGURE 4 ABOUT HERE]

523

524

525 *Acarechimys gracilis* (Ameghino, 1891) **comb.nov.**

526

Figure 4.5–6

527 *Stichomys gracilis* Ameghino, 1891

528 *Protacaremys pulchellus* Ameghino, 1902

529 *Acarechimys pascuali* Verzi *et al.*, 2016 *partim*

530 **Holotype.** MACN-A 4263, left mandible with dp4-m3.

531 **Referred material.** (Supplementary appendix 1).

532 **Geographic and stratigraphic occurrence.** Chubut and Santa Cruz provinces. Sarmiento
533 Formation, Colhué Huapi Member, early Miocene; Santa Cruz Formation, late early
534 Miocene (Supplementary Appendix 1).

535 **Emended diagnosis.** Within the size range of *A. constans* and slightly larger than *A.*
536 *minutus*. Lower deciduous premolars with well-developed metalophulid II and reduced
537 mesolophid, unlike remaining species. Lower molars with metalophulid II poorly
538 developed in m1 and reduced in m2-3; accessory cusp and posterior arm of the metaconid
539 present, as in *A. minutus*. Lower incisors not laterally compressed, as in *A. constans*.

540 Mandible with the nmpio and masseteric fossa located posteriorly, unlike in *A. minutus*,
541 *A. minutissimus*, and *A. leucothecae*.

542 **Remarks.** *Stichomys gracilis* was originally described by Ameghino (1891). Patterson (in
543 Wood and Patterson, 1982) considered *S. gracilis* to be a junior synonym of *Acaremys*
544 *minutus*, upon which he based the genus *Acarechimys* (Patterson and Wood, 1982).

545 Nevertheless, we do not think that MACN-A 4263 (holotype of *S. gracilis*) is assignable to
546 *Acarechimys minutus*; rather, it is indistinguishable from MACN-A 52-128, the holotype of
547 *Acarechimys pulchellus* (Ameghino, 1902). Thus, *Acarechimys pulchellus* is a junior
548 synonym of *Stichomys gracilis*, and we erect the new combination *Acarechimys gracilis*
549 (Ameghino, 1891).

550 **Description.** *Lower cheek teeth.* The m2 is the largest tooth in the dental series (Tab. 1).
551 The dp4 has a curved metalophulid I. The ectolophid and the metalophulid II extend
552 posteriorly from the protoconid. The metalophulid II is conspicuous and postero-lingually
553 oblique, as in *A. minutissimus* and *A. constans* (Fig. 4.5). The mesolophid is reduced to a

554 short spur that extends from the posterior end of the metalophulid II and delimits a fossettid
555 (Fig. 4.5). The hypolophid is long and straight. The posterolophid is long and curved but
556 does not contact the lingual end of the hypolophid (Fig. 4.5). The hypoflexid is the deepest
557 flexid and is posteriorly oblique.

558 The lower molars have three main crests and a fourth, second in position, variably
559 developed (Fig. 4.5). The metalophulid I is curved, as in *A. constans*. The metaconid
560 extends posteriorly to form the posterior arm of the metaconid, which is well-developed in
561 m1-3 (Fig. 4.5), unlike in *A. constans*, in which is absent. The metalophulid II is reduced
562 but more developed in m1 than in m2-3 (Fig. 4.5). An accessory cusp is present, including
563 in m3, unlike in other species of the genus (Fig. 4.5). The hypolophid and posterolophid are
564 as in dp4.

565 The lower incisors are robust and not laterally compressed. They have a thick
566 enamel layer, a straight lingual border, and a curved labial one. This tooth is long, as in the
567 remaining species of the genus, with its posterior end posterior and labial to m3 (Fig. 4.6).
568 *Mandible.* A short, concave diastema is present, as in the remaining species (Fig. 4.6). The
569 mental foramen is generally absent, but a very small foramen is present anterior to the dp4
570 in MPM-PV 17430. The nmmpio is long, conspicuous, and mostly located below m1 (Fig.
571 4.6). The masseteric crest is well-developed, continuous with the nmmpio, and projects
572 laterally. The masseteric fossa is slightly deeper than in *A. constans* but shallower than in *A.*
573 *minutissimus* (Fig. 4.6). The furrow that delimits the masseteric fossa antero-dorsally is
574 poorly developed, as in the remaining species. The coronoid process extends anteriorly to
575 the level of m2; it is postero-laterally extended and delimits a retromolar fossa lateral to
576 m2-3. Its anterior border is straight, and its dorsal tip is dorsal to the cheek teeth.

577 Posteriorly the mandibular notch is shallow. Lingually, the mandibular symphysis extends
578 posteriorly to the level of dp4; posteriorly, the mental process is moderately developed.

579

580 *Acarechimys leucotheae* Vucetich *et al.*, 2015

581 Figure 5. 1

582

583 **Type and only material.** MPEF-PV 10677, left mandibular fragment with dp4-m3 and
584 incisor.

585 **Geographic and stratigraphic occurrence.** Chubut Province, Argentina.

586 Upper levels of Sarmiento Formation at Cabeza Blanca, late Oligocene; Deseadan

587 SALMA.

588 **Diagnosis [modified from Vucetich *et al.* (2015a)].** Very small, within the size range of *A.*
589 *minutissimus*. Cheek teeth brachydont and terraced to a greater degree than in other species
590 of the genus. Lower molars with three crests plus a very short metalophulid II and an
591 accessory cusp in m1-2, as in *A. minutus*; posterior arm of the metaconid and posterior
592 extension of the metalophulid I absent; posterolophid more transverse than in the other
593 species, resulting in a more open posteroflexid. Lower deciduous premolar with the
594 ectolophid conspicuously separated from the protoconid, very oblique, and more
595 perpendicular to the antero-posterior axis of the tooth than in the other species of the genus;
596 metalophulid II reduced, as in some specimens of *A. minutus* and *A. constans*. Mandible
597 with the notch for the insertion of tendon of the masseter medialis pars infraorbitalis more
598 oblique than in the remaining species; mental foramen absent, as in *A. minutissimus* and *A.*
599 *minutus*.

600 **Remarks.** This species was recently described by Vucetich *et al.* (2015a) based on a single
601 specimen (Fig 5.1). No new specimens or additional information are available.

602

603 [FIGURE 5 ABOUT HERE]

604

605 *Acarechimys minutissimus* (Ameghino, 1887)

606 Figure 5. 2–4

607 *Stichomys diminutus* Ameghino, 1891

608 *Sciamys tenuissimus* Ameghino, 1894

609

610 **Type series.** MACN-A 256, left mandible with dp4-m3; MACN-A 257, left mandible with
611 dp4-m3; MACN-A 258, right mandible with incisor and dp4-m3; MLP 15-188, left
612 mandible with dp4-m3.

613 **Lectotype.** MLP 15-188 Patterson in Pascual, 1967.

614 **Paralectotypes.** MACN-A 256, MACN-A 257, MACN-A 258.

615 **Referred material.** (Supplementary appendix 1).

616 **Geographic and stratigraphic occurrence.** Santa Cruz and Neuquén provinces, Argentina;
617 La Venta, Colombia. Pinturas, Santa Cruz, and Collon Curá formations, early Miocene, late
618 early Miocene, and middle Miocene, ‘Pinturan’ age, Santacrucian and Colloncuran
619 SALMAs; Villavieja Formation, middle Miocene, Laventan SALMA (Supplementary
620 appendix 1).

621 **Emended diagnosis.** Very small, within the size range of *A. leucotheae*. Cusps conspicuous
622 in young specimens. Lower deciduous premolar with reduced metalophulid II and well-
623 developed mesolophid, unlike the remaining species of the genus. Metalophulid II

624 generally absent in the lower molars or present in m1 as a bulge into the ectolophid, unlike
625 in the remaining species of the genus; accessory cusp in antero+mesoflexid of m1-2
626 variably present, and posterior arm of the metaconid and posterior extension of the
627 metalophulid I absent, unlike in *A. minutus*. Lower incisors large relative to the mandible
628 size. Mandible with the nmmpio straight and below dp4-m1, as in *A. minutus* and *A.*
629 *leucotheae*.

630 **Remarks.** MLP 15-188 is labeled as the lectotype of *Acarechimys minutissimus*. The MLP
631 collections include several specimens from the ‘old collections’ referable to *A.*
632 *minutissimus* (MLP 15-1, MLP 15-188a, MLP 15-398, MLP 15-408), but unfortunately the
633 available information does not indicate whether they were collected by C. Ameghino (as is
634 the case for *A. minutus* and *A. constans*). Patterson (UMS, pers, comm.) mentioned MLP
635 15-188 as the type of the species. Thus, we can infer that only MLP 15-188 was available
636 to him. Two specimens were figured by Ameghino (1889; Atlas: plate IV, figs. 24, 25).
637 Figure 24 corresponds to MACN-A 257, but the specimen corresponding to figure 25 could
638 not be found [Fericola (2011) erroneously stated the reverse, that the figure 24 specimen
639 could not be found and that figure 25 specimen is MACN-A 257]. Additionally, MACN-A
640 256 and MACN-A 258 are mentioned in Ameghino’s catalog as belonging to the same
641 stock as MACN-A 257. In fact, Ameghino (1889) mentioned that this species was
642 represented by several mandibles. Thus, we conclude that four specimens (MLP 15-188,
643 MACN-A 256, MACN-A 257, MACN-A 258) are part of the original type series used by
644 Ameghino to erect *A. minutissimus*.

645 Patterson (UMS, pers, comm.) and Pascual (1967) determined MLP 15-188 to be
646 the lectotype of the species. As a consequence, the remaining specimens of the original
647 species are considered to be paralectotypes.

648 **Description.** *Upper cheek teeth.* Tooth rows labially obliquely implanted with respect to the
649 palatal plane, as in *A. minutus*. Cheek teeth with four crests (Fig. 5.2). The M2 is slightly
650 larger than DP4 and M1 (Tab. 1). Cusps discernable and molars slightly terraced.

651 The molars have a subquadrangular occlusal outline (Tab. 1) and well-defined
652 paracone and metacone (Fig. 5.2). The protocone area is more rounded than in *A. minutus*.
653 The anteroloph is short, unlike in the type species, but an anterofossette forms with little
654 wear since the paracone does not extend very far anteriorly (M2; Fig. 5.2). The protoloph is
655 slightly curved. The metaloph is straight and ends labially in the metacone, which is
656 posteriorly extended and contacts the relatively short posteroloph. Consequently, the
657 antero- and posterofossettes are delimited in juvenile specimens (Fig. 5.2). The
658 paraflexus/fossette is the smallest and shallowest fossette, and extends across the occlusal
659 surface as far as the mesoflexus. The hypoflexus is the deepest flexus and is anteriorly
660 oriented.

661 The DP4 resembles the molars but differs in its rounded occlusal outline and in the
662 absence of an anteriorly oriented paracone. Unlike in *A. minutus*, the anteroloph is short
663 and oblique (Fig. 5.2), and in some specimens, the metaloph is reduced (MACN-A 4145),
664 as in *A. minutus*.

665 The upper incisors are oval in section, as in *A. minutus*. The enamel is thick, and the
666 anterior face is straight and the labial face is curved.

667

668 *Skull.* The description is based on two small, poorly preserved skull fragments (MACN-A
669 12683; YPM-PVPU 15178). In the ventral aspect of the skull, the incisive foramina are
670 well-developed, as in *A. minutus*. Posteriorly, they are continuous with well-developed
671 diastemal furrows that extend posteriorly to the anterior border of the DP4s. The ventral

672 root of the zygoma extends just in front of the DP4, and its antero-posterior diameter is
673 similar to its dorso-ventral diameter, as in the type species. The masseteric tuberosity is
674 well-developed, and the lateral furrow for the insertion of the lateral masseteric muscle is
675 shallow. Unlike in *A. minutus*, there is no foramen of uncertain affinities posterior to the
676 masseteric tuberosity. On the dorsal face of the ventral root of the zygoma is a faint furrow
677 for the passage of the infraorbital nerve.

678 *Lower cheek teeth.* Juveniles have terraced cheek teeth with the metaconid and entoconid
679 higher than the protoconid and hypoconid, and adults have flat occlusal surfaces. The dp4
680 of this species is most variable: it has a curved metalophulid I whose lingual and labial
681 portions can be separated from each other (MACN-A 4076; MACN-A 4083). The
682 metalophulid II is usually reduced, as in *A. minutus*, *A. leucothaeae*, and some specimens of
683 *A. constans* (Fig. 5.3), but in some cases, it reaches the mesolophid. The ectolophid is
684 lingually concave, and in juveniles (MACN-A 1896; MPM-PV 15098), as well as in those
685 specimens from the Pinturas Formation, it is not connected to the protoconid. The
686 mesolophid is always well-developed, as in *A. constans*, and reaches the metaconid to
687 delimit an anterior fossettid (Fig. 5.3). In some cases, this crest is disconnected from the
688 ectolophid (MACN-A 4094). The hypolophid is straight or curved, and reaches the lingual
689 side of the tooth. The posterolophid is long and curved, unlike in *A. leucothaeae*. The
690 anterofossettid is rounded and shallow and disappears with the posterofossette in adults.
691 The straight mesoflexid and the posteriorly oblique hypoflexid are the deepest flexi and
692 remain open in adults.

693 The molars have three crests. The metalophulid II is absent or reduced to a minute
694 bulge on m1 (Fig. 5.3). The metalophulid I is straight, unlike in *A. constans* and *A. gracilis*,
695 with a labio-lingually aligned protoconid and metaconid or a metaconid that is slightly

696 anterior to the protoconid (Fig. 5.3). The hypolophid and posterolophid are as in the dp4.
697 Antero+mesoflexid are merged owing to the absence of metalophulid II (Fig. 5.3). An
698 accessory cusp is usually present on m1 and m2 (Fig. 5.3). This cusp can be connected to
699 the metalophulid I by a posterior extension of the latter (MLP 15-398). The posterior arm of
700 the metaconid is not present. In some juvenile specimens (MACN-A 4083; MACN-A 4092;
701 MACN-A 4093; MLP 15-398), the hypoflexid is united with the posteroflexid.
702 Antero+mesoflexid and posteroflexid are similar in depth; the hypoflexid is the deepest
703 flexid and is posteriorly oblique.

704 Lower incisors are laterally compressed and very large relative to mandible size
705 (Fig. 5.4). The enamel layer is thick, and the anterior face is straight and the labial face is
706 curved. The incisors are long, extending below the m3 and ending in a bulge on the base of
707 the coronoid process or in a furrow in broken specimens (Fig. 5.4).

708 *Mandible.* The mandible of this species has a conservative morphology that contrasts with
709 the great variability of the lower cheek teeth. It is robust, and the diastema is dorsally
710 concave and shorter than the tooth row as in the remaining *Acarechimys* species (Fig. 5.4).
711 The mental foramen is nearly always absent (only on MACN-A 4081 is a very small
712 foramen located anterior to the dp4). The nmmpio is straight and conspicuous, and extends
713 below the dp4-m1 (Fig. 5.4). The masseteric crest is continuous with the posterior border of
714 the nmmpio and protrudes laterally. The masseteric fossa is deep anteriorly, as in *A.*
715 *minutus* (Fig. 5.4). The furrow that delimits this fossa antero-dorsally is poorly developed.
716 The base of the coronoid process extends forward to the level of m2, as in *A. minutus* and
717 *A. gracilis*, thereby delimiting a retromolar fossa lateral to m3. The mandibular symphysis
718 extends posteriorly to m1, and a moderately developed chin is present anteriorly, which
719 delimits the notch for the insertion of the digastric muscle.

720

721 **Statistical Analysis.** We tested for size differences among *Acarechimys* species using m1
722 and m2 length, as these allowed for the largest sample sizes. ANOVAs of both m1 length
723 (N = 69) and m2 length (N = 62) were highly significant ($p < 0.0001$). Based on Tukey's
724 HSD (Tab. 2), two subgroups of *Acarechimys* are statistically distinguishable: a group of
725 larger species consisting of *A. constans*, *A. gracilis*, and the sample from Quebrada Honda
726 (referred here tentatively to *A. minutus* and *A. minutissimus*), and a smaller group consisting
727 of *A. leucotheae*, *A. minutissimus*, and *A. minutus*. These size groupings are distinct in a
728 bivariate plot of m1 vs. m2 length (Fig. 6; N = 58).

729 Two conclusions can be drawn from this analysis. First, size does not appear to be a
730 useful criterion for distinguishing *A. minutissimus* from *A. minutus*, even though this was
731 the main criterion used by Ameghino (1887) for distinguishing these species. Although the
732 smallest *Acarechimys* specimens do pertain to *A. minutissimus*, and m1 and m2 lengths are
733 statistically different between these species (though only at the $p < 0.05$ to 0.005 level),
734 there is significant size overlap between larger specimens of *A. minutissimus* and smaller
735 specimens of *A. minutus* (Fig. 6). The single known specimen of *A. leucotheae* plots very
736 close to this area of overlap. *Acarechimys constans* and *A. gracilis* show a pattern similar to
737 *A. minutissimus* and *A. minutus* but with even greater size overlap (Fig. 6); the two species
738 show virtually the same range of values for both m1 and m2 length and cannot be
739 distinguished from one another based on size alone.

740 The other noteworthy result of this analysis is that the specimens in the *Acarechimys*
741 sample from Quebrada Honda, tentatively identified as both *A. minutus* and *A.*
742 *minutissimus*, are significantly larger than Patagonian specimens referred to these species,
743 and similar in size to *A. constans* and *A. gracilis* (Fig. 6). This suggests that the Quebrada

744 Honda populations referred to *A. minutus* and *A. minutissimus* evolved in parallel toward
745 larger size in this region during the late middle Miocene, or that they represent different
746 species from those from Patagonia. If the first hypothesis is correct, the differences in size
747 could be the result of similar responses to a common environmental factor such as climate
748 or habitat. Body size change in response to climate has been documented in other extinct
749 species (*e.g.*, Gingerich, 2003; Chew, 2015) and climate change has been proposed to have
750 had a significant effect on the evolution of body size in North American Cenozoic
751 mammals (Lovegrove and Mowoe, 2013). In this case, it is curious that the single specimen
752 of *Acarechimys* from La Venta, Colombia, here referred to *A. minutissimus*, is similar in
753 size to Patagonian specimens of this species (m1 length = 1.56 mm; Walton, 1990, tab. 1).
754 If climate (temperature) were primarily responsible for the larger size of Quebrada Honda
755 specimens, one might expect a similar pattern at La Venta, which is the same age as
756 Quebrada Honda and also located in tropical latitudes. Testing this hypothesis requires
757 studying additional specimens of *Acarechimys* collected at Quebrada Honda since the
758 publication of Croft *et al.* (2011) and incorporating data from ongoing paleoenvironmental
759 studies at the site (Cadena *et al.*, 2015; Catena *et al.*, 2016). It should also be noted that
760 three *A. minutus* specimens from Patagonia are particularly large and fall within the range
761 of specimens from Quebrada Honda (the first two are represented only by m1): MLP 82-
762 XII-1-31 (Santa Cruz Province, exact provenance unknown), MLP 91-IX-1-200 (Collon
763 Curá Formation at Cañadón del Tordillo), and MPM-PV 4193 (Puesto La Costa, costal
764 Santa Cruz Province; Fig. 6). Each of these represents a relatively large individual within
765 an otherwise small-bodied population and provides no systematic explanation for the
766 relatively large size of the Quebrada Honda sample.

767

[FIGURE 6 ABOUT HERE]

768

769

770 ***Phylogenetic Analysis***

771 The cladistic analysis resulted in 12 Most Parsimonious Trees (MPTs) of 798 steps
772 each (Consistency index=0.318; Retention index=0.544) and the best score hit 57 times out
773 of 1,000. The strict consensus tree (Fig. 7) has a well-resolved topology. Results are in
774 general agreement with previous analyses, but minor differences are present with respect to
775 the relationships of some groups. One interesting aspect is the basal position of
776 *Draconomys verai* (early Oligocene of Chubut province, Argentina) within caviomorphs.
777 The phylogenetic relationships of some species originally described as basal octodontoids
778 (*e.g.*, *D. verai*; *Eosallamys simpsoni*, *Eoespina woodi*, and *Eosachacui lavocati* from the
779 late Eocene?–early Oligocene of Peru; *Changquin woodi* from the late Oligocene of Chubut
780 province; and *Dudumus ruigomezi* from the early Miocene of Chubut province, Argentina)
781 and of those taxa from the late middle Eocene of Contamana, Peru (*Cachiyacuy*
782 *contamanensis* and *Canaanimys maquiensis*) are not clear and vary in different analyses
783 (Antoine *et al.*, 2012; Arnal *et al.*, 2014; Arnal and Vucetich, 2015). Therefore, the
784 relationships of these species will be the subject of a future study focused on basal
785 caviomorphs.

786 Pan-Octodontoidea (node 1) is characterized by six synapomorphies [metaloph on
787 DP4 indistinct, probably fused to the posteroloph (character 13:1); mesostyle on DP4
788 indistinct or absent (character 15:0); mesolophule slightly oblique on M1-M2 (character
789 42:1); presence of an anterior flexid in metalophulid I of p4 (character 70:0); short
790 mesolophid on p4 (character 73:0); nmmpio at the middle of the mandible high

791 (character111:1)]. Nevertheless, node 1 has a low support (Fig. 7), since most of those
792 characters are scored in only a few taxa.

793 Within stem-Octodontoidea, several clades previously recovered in other analyses
794 are also recovered here. Node 2 (Fig. 7) is the sister-clade to remaining octodontoids and
795 includes *Eosallamys simpsoni*, *Migraveramus beatus*, and several other species including
796 the enigmatic *Plesiacaechimys koenigswaldi* from the middle Miocene of Neuquén
797 Province (Argentina). Later-diverging clades include (*Eoespina woodi* + *Sallamys?*
798 *minutus*), Acaremyidae, (*Caviocricetus lucasi* + *Dudumus ruigomezi*), and the five species
799 of *Acaechimys*. The phylogenetic relationships of crown Octodontoidea (Fig. 7) should be
800 considered tentative because the taxonomic sample of this study is not focused on this part
801 of the tree.

802 *Acaechimys* is recovered as a monophyletic genus of stem-octodontoids (Fig. 7)
803 characterized by the presence of brachydont cheek teeth (character 3:0), the presence of an
804 accessory cusp on m1-2 (character 97:0), and a groove for the passage of the nerve
805 infraorbitalis (character 139:1). *Acaechimys minutus*, the earliest-diverging species, is
806 distinguished by a metacone that is slightly lingual to the paracone on M2 (character 39:1).
807 The clade of *Acaechimys gracilis* + *A. constans* is characterized by a lack of compressed
808 lower incisors (character 101:1) and by having the nmmpio positioned beneath m1
809 (character 110:0). Its sister clade, *A. minutissimus* + *A. leucothaeae*, is characterized by the
810 presence of terraced occlusal surfaces in all cheek teeth (character 9:0) and a deep anterior
811 portion of the masseteric fossa (character 113:2). A striking aspect of these results is the
812 reacquisition of very low cheek teeth by the genus (interpreted as a reversal within
813 octodontoids), although such a reversal has never been postulated for octodontoids.
814 Analyses underway will further test this hypothesis. Lastly, unlike the proposal of Verzi et

815 al. (2016) *Abrocoma cinerea* is not directly related to any species of *Acarechimys*; rather, it
816 is the earliest-diverging crown-Octodontoidea.

817 [FIGURE 7ABOUT HERE]

818

819 **Discussion and Conclusions**

820 The systematic and phylogenetic analyses performed here allow us to define the content of
821 the *Acarechimys* group and to better discriminate the included species. *Acarechimys* is
822 characterized by a unique mix of states that are plesiomorphic (*e.g.*, low-crowned cheek
823 teeth) and apomorphic (retention of the deciduous premolar, absence of mental foramen,
824 presence of an accessory cusp on m1-2, and presence of a groove for the passage of the
825 nerve infraorbitalis) among octodontoids. Additionally, *Acarechimys* species differ from
826 each other by the presence/absence of dental and mandibular structures (Tab. 3). Our
827 statistical analyses demonstrate that size is not a relevant feature for distinguishing *A.*
828 *minutus* from *A. minutissimus*. Traditionally, this was the main feature used for
829 distinguishing species, albeit tentatively (Vucetich *et al.*, 1993a), but our metric analyses
830 demonstrate that there are significant overlaps in size among specimens referred to these
831 species (see above; Fig. 5).

832 The temporal and geographic distributions of *Acarechimys* species suggests that the
833 genus could have evolved in Patagonia by at least the early late Oligocene (in the first
834 Patagonian radiation event; Arnal and Vucetich, 2015). The geologically oldest species is
835 the minute *Acarechimys leucotheae*, which has only been identified at the late Oligocene
836 (Deseadan SALMA) site of Cabeza Blanca, in Chubut Province, Argentina (Fig. 2). Cabeza
837 Blanca has produced the greatest diversity of late Oligocene rodents in South America,
838 nearly three times as many species as any other site of this age (Vucetich *et al.*, 2015a). This

839 is likely due, at least in part, to the large number of rodent specimens that have been
840 collected there and the thorough taxonomic investigations that have focused on this site
841 (see Wood and Patterson, 1959; Vucetich *et al.*, 2015a), though only a single specimen of
842 *A. leucothaeae* has been identified from the site thus far. This suggests that *Acarechimys* was
843 relatively rare at that time, a conjecture that is supported by the absence of specimens
844 referable to the genus from La Flecha, the other rich Patagonian locality. Nevertheless, it is
845 noteworthy that no specimens of *Acarechimys* have yet been identified from Salla, Bolivia,
846 which has also produced a rather diverse and rich fauna of late Oligocene rodents (Lavocat,
847 1976; Patterson and Wood, 1982; Vucetich, 1991). Thus, the present evidence suggests that
848 *Acarechimys* could have originated in Patagonia.

849 *Acarechimys* apparently continued to be rare prior to the late early Miocene, as
850 *Acarechimys gracilis*, from the Colhuehuapian SALMA of Chubut Province (Fig. 2), is
851 known only through one specimen, the holotype of *Protacaremys pulchellus* (Ameghino,
852 1902; Vucetich *et al.*, 2010). Kramarz *et al.* (2004) mentioned the presence of *Acarechimys*
853 for the Colhuehuapian beds of Chichinales Formation in Río Negro (Fig. 2), but we have
854 studied this specimen and determined that it does not belong to *Acarechimys*. Despite the
855 rich octodontoid fossil record and the great morphological disparity of this group prior to
856 the late early Miocene, *Acarechimys* remained poorly diversified during this interval (about
857 half of Colhuehuapian caviomorphs are octodontoids; Vucetich *et al.*, 2010, 2015b).

858 By the late early Miocene, *A. minutissimus* is recorded for the ‘Pinturan’ age of Santa
859 Cruz Province (Fig. 2) (Kramarz, 2004; see Kramarz and Bellosi, 2005 and Perkins *et al.*,
860 2012 for a discussion about the overlap of some parts of the ‘Pinturan’ and Santacrucian
861 levels). The acme of the genus was during the Santacrucian SALMA (Santa Cruz
862 Formation), where four of the five recognized species lived in what is today

863 Argentinean Santa Cruz Province (*A. minutus*, *A. constans*, *A. gracilis*, and *A.*
864 *minutissimus*). This great diversity could partly be attributable to the fact that the Santa
865 Cruz Formation exposures have a wide distribution (Fig. 2), and have been broadly
866 prospected since the nineteenth century, resulting in an unparalleled collection of fossils
867 (Ameghino, 1887, 1889; Scott, 1905; Vizcaíno *et al.*, 2012; Fernicola *et al.*, 2014).
868 *Acarechimys* has also been recovered at other Santacrucian localities in Chile (Flynn *et al.*,
869 2002, 2008; Croft *et al.*, 2007), but these materials have not yet been figured nor described
870 in detail. In general, Santacrucian caviomorphs are quite distinct from those of the
871 Colhuehuapian and also from those of the ‘Pinturan’ age (Vucetich *et al.*, 2015b),
872 exhibiting a marked tendency toward increased hypsodonty (Kramarz, 2001; Pérez and
873 Vucetich, 2012; Arnal and Pérez, 2013). This ecological shift in rodents is generally
874 thought to be a consequence of climatic deterioration in Patagonia between Colhuehuapian
875 and Santacrucian intervals (Vucetich, 1986; Pérez and Vucetich, 2012; Arnal and Pérez,
876 2013). However, the evidence for such deterioration is equivocal. Global temperatures
877 remained relatively stable across this interval (Zachos *et al.*, 2008), and open habitats were
878 present at least episodically during both the Colhuehuapian and Santacrucian SALMAs
879 (Dunn *et al.*, 2015), though arid-adapted shrubs only became dominant in Patagonia after
880 the late Miocene (Palazzesi and Barreda, 2012). Kay *et al.* (2012) interpreted the
881 paleoenvironment of coastal Santa Cruz as highly seasonal with a mosaic of vegetation
882 including both forested and more open areas. It is possible that an increase in exogenous
883 grit, such as volcanic ash, may have driven the trend toward increased hypsodonty and a
884 replacement of caviomorph species across this interval, but it is difficult to test such a
885 hypothesis at present due to a paucity of studies that include data from both the
886 Colhuehuapian and Santacrucian SALMAs. Regardless of the precise causes of the

887 ecological shifts in other rodents, *Acarechimys* is noteworthy in being the only octodontoid
888 lineage that retained generalized, brachydont cheek teeth into the late early Miocene in high
889 latitudes.

890 The Patagonian fossil record is scarce for the middle Miocene, and known fossil sites
891 have a more northerly location compared to the early Miocene (Pascual and Ordreman
892 Rivas, 1971; Pascual and Ortiz Jaureguizar, 1990; Pérez, 2010; Arnal and Pérez, 2013).
893 Rodents are represented in few Colloncuran localities (Colloncuran SALMA; earliest
894 middle Miocene) in Neuquén and Río Negro provinces (*e.g.*, Cañadón del Tordillo and
895 Pilcaniyeu Viejo respectively; Bondesio *et al.*, 1980; Vucetich *et al.*, 1993a; Fig. 2). During
896 the Mayoan (latest middle Miocene), rodents have been reported from several small
897 faunules in western Chubut and Santa Cruz provinces (Kraglievich, 1930; Bondesio *et al.*,
898 1980; Vucetich and Pérez, 2011; Pérez *et al.*, 2016), as well as the locality of El Petiso in
899 Chubut Province, whose age is estimated to be post-Colloncuran (Villafañe *et al.*, 2008;
900 Arnal and Pérez, 2013). Among these middle Miocene localities, *Acarechimys* has only
901 been identified at Cañadón del Tordillo and Estancia Collon Curá. In a recent preliminary
902 revision of unpublished caviomorphs of Cañadón del Tordillo and Estancia Collon Curá, a
903 very high octodontoid diversity was identified (Vucetich and Arnal, pers. obs.). However,
904 *Acarechimys* diversity appears to be lower than during the Santacrucian. During this time,
905 northern Patagonia experienced a short period of regreening; forests and more humid
906 conditions are inferred for Cañadón del Tordillo based on the presence of monkeys and a
907 high diversity of porcupines (Candela, 2003; Dunn *et al.*, 2015; Vucetich *et al.*, 2015b), as
908 well as many low-crowned octodontoids (Vucetich and Arnal, pers. obs.). Few late
909 Miocene sites are known from Patagonia. These have yielded only fragmentary remains of

910 rodents and no octodontoids (Pascual and Bondesio, 1985; Vucetich *et al.*, 2005; Dozo *et*
911 *al.*, 2010).

912 A variety of fossil sites are known from central and northern Argentina and lower
913 latitudes of the continent during the Eocene to Miocene interval. Sites of Eocene and
914 Oligocene age are known from Peru, Bolivia, Brazil, and Uruguay. In general, the fossil
915 record of these localities (with the exception of Salla, Bolivia) is very poor, and no
916 *Acarechimys* or closely similar taxa have been described (Lavocat, 1976; Mones and
917 Castiglione, 1979; Patterson and Wood, 1982; Vucetich, 1991; Vucetich *et al.*, 1993b;
918 Bond *et al.*, 1998; Vucetich and Ribeiro, 2003; Frailey and Campbell, 2004; Antoine *et al.*,
919 2012). An unidentified species of *Acarechimys* was mentioned for the late early Miocene of
920 Chucal, northern Chile (~18° S; Croft *et al.*, 2007), and the genus has been recorded at the
921 late middle Miocene of La Venta, Colombia (~3° N; Walton, 1997), Quebrada Honda,
922 Bolivia (~22° S; Croft *et al.*, 2011), and the Fitzcarrald Arch in Peruvian Amazonia (~11°
923 S; Tejada-Lara *et al.*, 2015: fig. 9N) (Fig. 2). Other early and middle Miocene localities
924 have yielded remains of caviomorphs, but there is no record of *Acarechimys* (*e.g.*, Madre de
925 Dios Subandean Zone, Peru; Antoine *et al.*, 2013). For the late Miocene, Campbell *et al.*
926 (2006) mentioned the possible presence of *Acarechimys* in the Madre de Dios Formation in
927 the Amazonia region, and Antoine *et al.* (2016) did the same for the Pebas Formation of
928 Peru. Based on our firsthand study of the material of the Madre de Dios Formation and
929 examination of photos of specimens from the Pebas Formation, we do not believe
930 *Acarechimys* occurs at these sites. Instead, the specimens from the Amazonia region
931 represent a new caviomorph species that is broadly represented in southwest Amazonia
932 (Brazil and Peru) during the late Miocene (Vucetich *et al.*, in prep). The phylogenetic

933 relationships of this new taxon relative to *Acarechimys* will be the subject of future
934 research.

935 Based on available evidence, *Acarechimys* was apparently not present in lower latitudes
936 of the continent (north of 35° S) before the early Miocene; the factors favoring its dispersal
937 after this time remain to be elucidated. The change in its distribution toward low latitudes
938 after the early middle Miocene is broadly reminiscent of a pattern of range contraction seen
939 in several other groups of mammals including vermilinguan xenarthrans, platyrrhine
940 primates, and astrapothere ungulates (Pascual *et al.*, 1996; Ortiz Jaureguizar and Cladera,
941 2006; Croft *et al.*, 2016), and it raises the possibility of a common environmental or
942 ecological cause. Paleoecological studies of *Acarechimys* are necessary to provide
943 additional insights into how and why this tiny caviomorph was able to achieve the widest
944 temporal and geographic distribution of any caviomorph genus, while retaining a
945 persistently brachydont dentition.

946

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960

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1238

1239 **Figure 1.** Dental and mandibular nomenclature used in this work. **1**, upper cheek teeth: **Al**,
1240 anteroloph; **anterofl**, anteroflexus; **H**, hypocone; **M**, metacone; **Mel**, metaloph; **Mr**, mure;
1241 **P**, protocone; **Pa**, paracone; **Prl**, protoloph; **Psl**, posteroloph; **2**, lower cheek teeth: **ac**,
1242 accessory cusp; **ecd**, ectolophid; **et**, entoconid; **hd**, hypoconid; **hld**, hypolophid; **md**,
1243 metaconid; **med I**, metalophulid I; **med II**, metalophulid II; **msd**, mesolophid; **pamd**,
1244 posterior arm of the metaconid; **pemed**, posterior extension of the metalophulid I; **prd**,
1245 protoconid; **psd**, posterolophid; **3**, mandible: **cor**, coronoid process; **d**, diastema; **mac**,
1246 masseteric crest; **maf**, masseteric fossa; **mf**, mental foramen; **nmmpio**, notch for the
1247 insertion of the masseter muscle, *pars infraorbitalis*.

1248

1249 **Figure 2.** Map showing fossil localities where *Acarechimys* has been found or mentioned;
1250 **1**, Cabeza Blanca, Chubut, Argentina; **2**, Gran Barranca, Chubut, Argentina; **3**, Río Pinturas
1251 Valley, Santa Cruz, Argentina; **4**, coastal localities of the Santa Cruz Formation, Santa
1252 Cruz, Argentina; **5**, localities at the cliffs of the Río Santa Cruz, Santa Cruz, Argentina; **6**,
1253 Pampa Castillo, Región XI, Chile; **7**, Chucal Formation, Región XV, Chile; **8**, Laguna del
1254 Laja (Curá Mallín Formation), Región VIII, Chile; **9**, Cañadón del Tordillo, Neuquén,
1255 Argentina; **10**, El Petiso, Chubut, Argentina; **11**, Huemules River (Río Mayo Formation),
1256 Chubut, Argentina; **12**, Quebrada Honda, Bolivia; **13**, La Venta, Colombia; **14**, Fitzcarrald
1257 Arch, Peru; **15**, Puerta de Corral Quemado y Villavil (Andalhuala and Chiquimil
1258 formations), Argentina. A, B, and C refer to dismissed mentions of *Acarechimys* (**A**,

1259 Chichinales Formation, Río Negro, Argentina; **B**; Madre de Dios Formation, Peru; **C**;
1260 Pebas Formation, Peru).

1261

1262 **Figure 3.** *Acarechimys minutus*; **1**, MPM-PV 15088, left DP4-M3 (reversed); **2**, YPM-
1263 VPPU 15806, right lateral view of the skull; **3**, MACN-A 237, right dp4-m3 (paralectotype);
1264 **4**, MLP 91-IX-1-97, right dp4; **5**, MPM-PV 15089, external view of right mandible.

1265 **Abbreviations:** **dzt**, dorsal root of the zygoma; **hramus**, horizontal ramus of the zygoma;
1266 **pmx-mx**, premaxillary-maxillary; **mx-j**, maxillary-jugal; **rmf**, rostral masseteric fossa;
1267 **vramus**, vertical ramus of the zygoma; **vrz**, ventral root of the zygoma. Scale bars = 2mm.

1268

1269 **Figure 4.** **1–4**, *Acarechimys constans*; **1–2**, MPM-PV 15002, palatal fragment with left
1270 and right DP4-M2; **2**, left DP4-M2 (reversed); **3**, MLP 15-391, right dp4-m3 (lectotype); **4**,
1271 MACN-A 4064, external view of left mandible (reversed); **5–6**, *Acarechimys gracilis*
1272 MACN-A 52-128; **5**, right dp4-m3; **6**, external view of the right mandible. Anterior to the
1273 right, except Figure 4.1. **Abbreviations:** **ac**, accessory cusp; **Al**, anteroloph; **maf**,
1274 masseteric fossa; **med II**, metalophulid II; **Mel**, metaloph; **mf**, mental foramen; **msd**,
1275 mesolophid; **nmmpio**, notch for the insertion of the masseter muscle, *pars infraorbitalis*;
1276 **pamd**, posterior arm of the metaconid; **Prl**, protoloph. Scale bars = 2mm.

1277

1278 **Figure 5.** **1**, *Acarechimys leucothaeae*, MPEF-PV 10677, left mandible with dp4-m3
1279 (holotype; reversed); **2–4**, *Acarechimys minutissimus*; **2**, MACN-A 4076, right dp4-m3; **3**,
1280 MACN-A 258, right mandible with dp4-m3 (paralectotype); **4**, MACN-A 4093, left
1281 mandible. Anterior to the right. **Abbreviations:** **ac**, accessory cusp; **Al**, anteroloph; **M**,
1282 metacone; **maf**, masseteric fossa; **med II**, metalophulid II; **Mel**, metaloph; **msd**,

1283 mesolophid; **nmmpio**, notch for the insertion of the masseter muscle, *pars infraorbitalis*;
1284 **Pa**, paracone; **Prl**, protoleph; **PsI**, posteroleph. Scale bars = 1 mm.

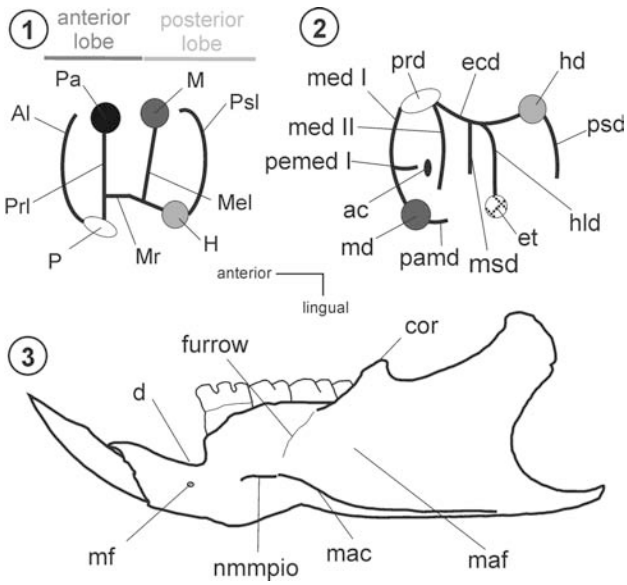
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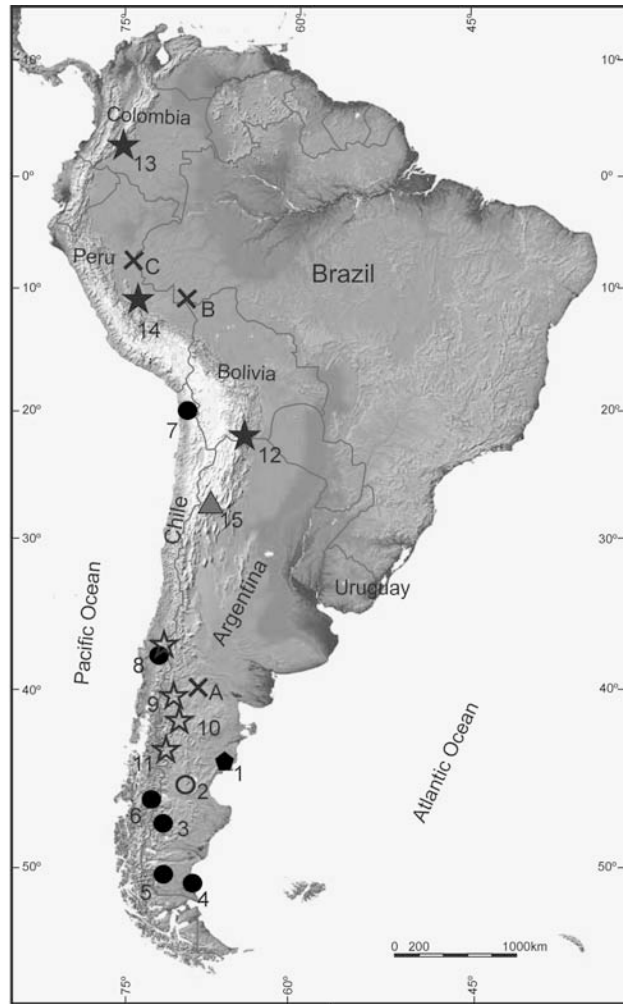
1286 **Figure 6.** Bivariate plot showing the relationship between m1 and m2 length (measured in
1287 mm) of the five species of *Acarechimys*.

1288

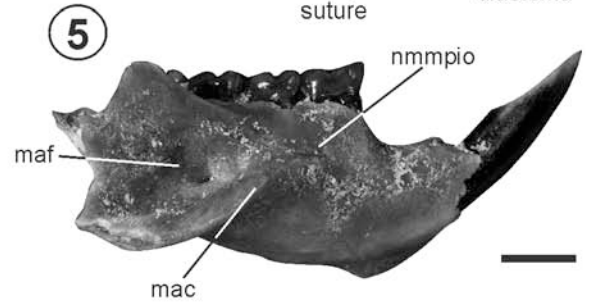
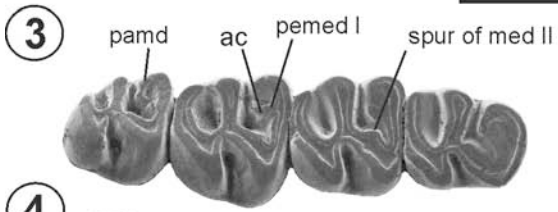
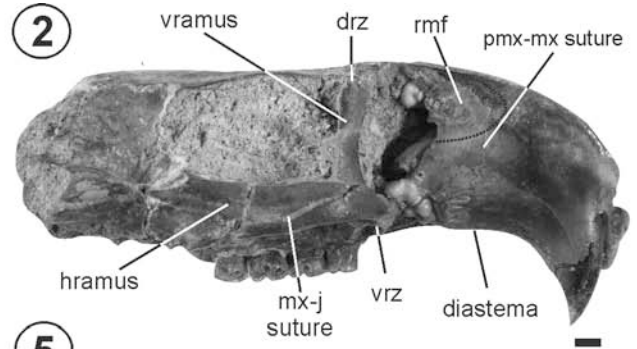
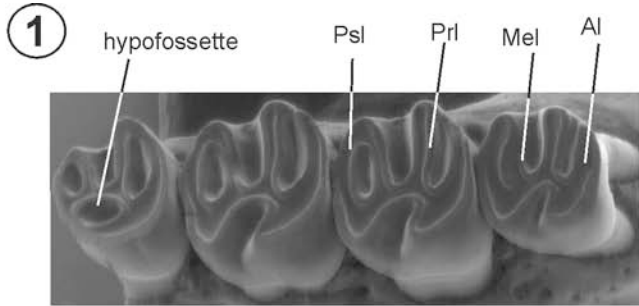
1289 **Figure 7.** Consensus tree of 12 MPTs showing the phylogenetic relationships of
1290 *Acarechimys* species (highlighted in green). Numbers at nodes separated by slash refer to
1291 absolute (anterior) and relative (posterior) Bremer supports values. Numbers in circles are
1292 nodes: 1- Pan Octodontoidea, 2- Basal octodontoid clade.

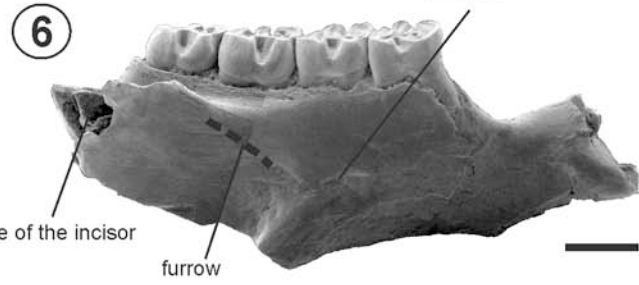
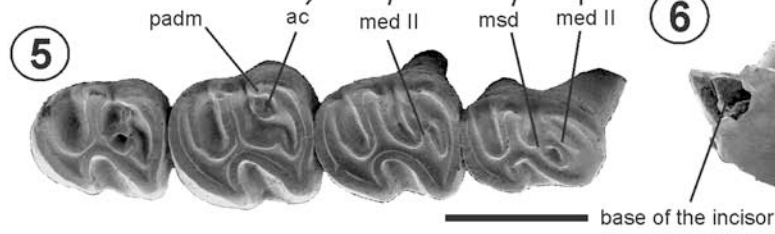
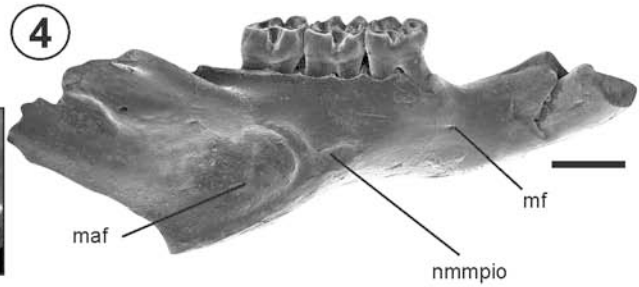
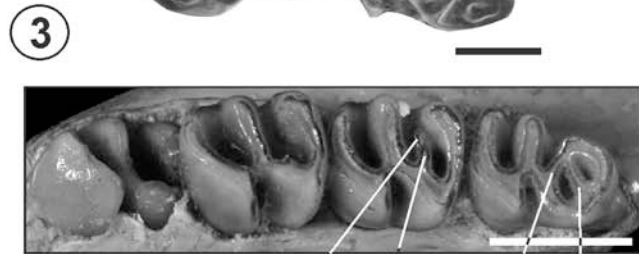
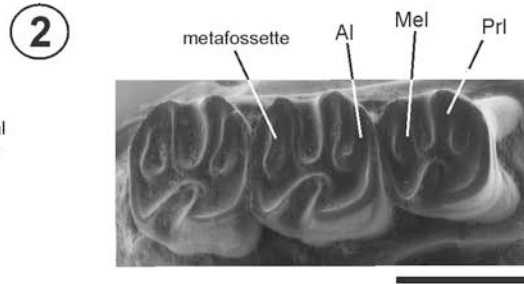
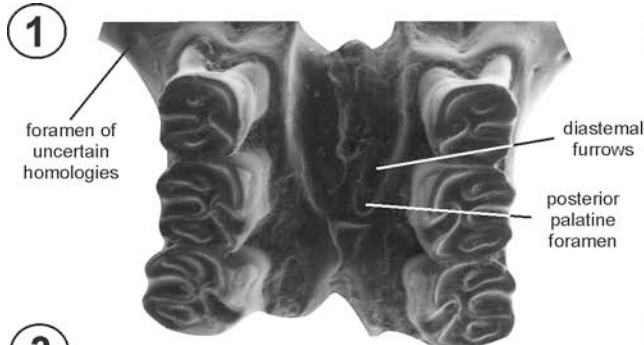
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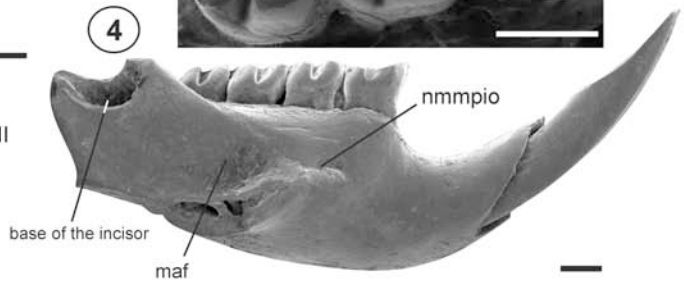
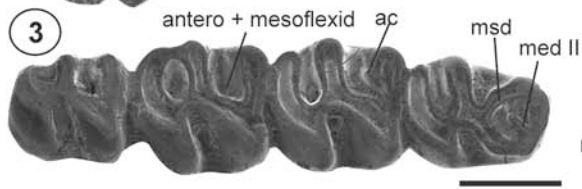
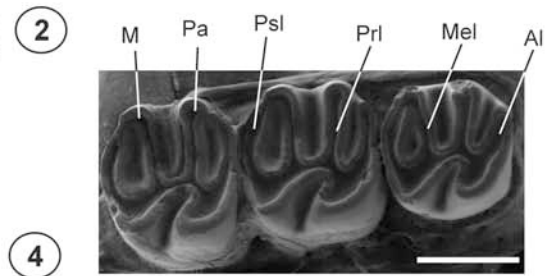
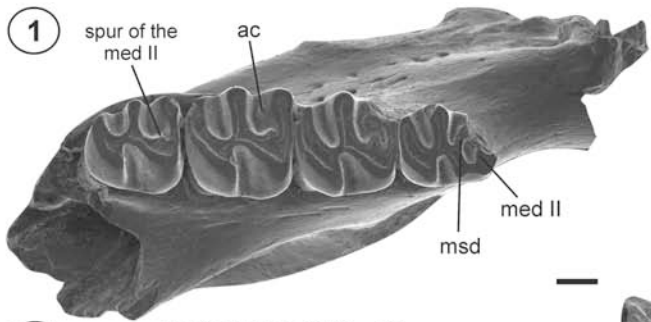


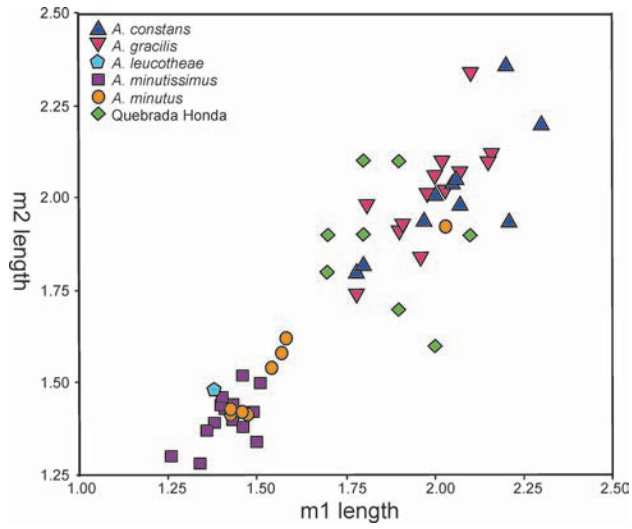


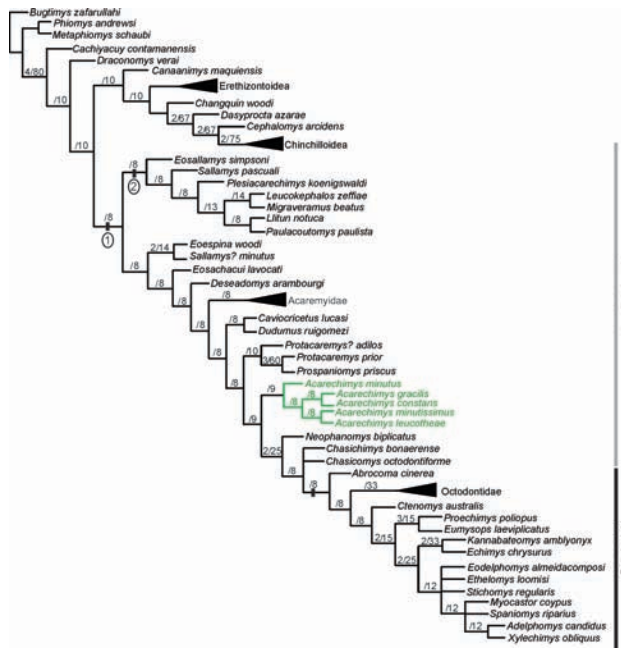
- ◆ late Oligocene (Deseadan SALMA)
- early Miocene (Colhuehuapian SALMA)
- late early Miocene (Santacrucian SALMA and Pinturan Age)
- ☆ middle Miocene (Colloncuran SALMA)
- ★ middle Miocene (Laventan SALMA)
- ▲ late Miocene/ early Pliocene
- ✕ dismissed mentions of *Acarechimys*











Order: Caviomorpha

Order: Castorimorpha

Table 1. *Acarechimys* dental measurements. Abbreviations: apl, antero-posterior length; mw, maximum width

<i>A. minutus</i>	MACN-A 237		MACN-A 4070		MACN-A 4073		MLP 15-410a		MPM-PV 15039	
	apl	mw	apl	mw	apl	mw	apl	mw	apl	mw
dp4	1.62	1.30			1.56	1.18	-	-	1.44	1.22
m1	1.57	1.52			1.46	1.39	-	1.58	1.48	1.40
m2	1.58	1.48			1.50	1.49	1.62	1.62	1.42	1.38
m3	1.24	1.40			-	-	1.56	1.58	-	-
DP4-M3			6.92	-						
DP4			1.62	1.90						
M1			1.84	2.06						
M2			1.90	2.10						
M3			1.56	1.84						

<i>A. constans</i>	MLP 15-391		MACN A 247a		MACN A 247b		MPM-PV 15002		MPM-PV 15092	
	apl	mw	apl	mw	apl	mw	apl	mw	apl	mw
dp4	2.22	1.64	-	-	2.36	1.64	-	1.56	2.33	1.53
m1	2.00	1.88	2.20	2.01	2.30	2.08	2.12	1.82	1.97	1.67
m2	2.01	1.94	2.36	2.16	2.20	2.16	-	-	1.94	1.96
m3	1.92	1.76	-	-	-	-	-	-	-	-
DP4-M2							5.04	-		
DP4							1.46	1.78		
M1							1.78	2.01		
M2							1.80	2.20		

<i>A. leucothaeae</i>	MPEF-PV 10677	
	apl	mw
dp4	-	1.02
m1	1.38	1.34
m2	1.48	1.34
m3	1.20	1.18

<i>A. minutissimus</i>	MACN-A 258		MACN-A 1894		MACN-A 12683		MLP 15-1			
	apl	mw	apl	mw	apl	mw	apl	mw		
dp4	1.22	1.11	1.51	1.12			1.41	1.19	1.32	1.12
m1	1.25	1.26	1.41	1.30			1.43	1.34	1.26	1.28
m2	1.46	1.46	1.43	1.31			1.40	1.32	1.30	1.30
m3	1.28	1.21	1.20	1.20			-	-	1.00	1.12
DP4-M2					3.96	-				
DP4					1.30	1.30				
M1					1.28	1.46				
M2					1.38	1.53				

<i>A. gracilis</i>	MACN-A 52-128		MACN-A 4060		MACN-A 4263		MLP 15-391a		MPM-PV 17430	
	apl	mw	apl	mw	apl	mw	apl	mw	apl	mw
dp4	1.88	1.54	2.32	1.70	1.76	1.48	1.90	1.59	2.08	1.54
m1	1.96	1.74	2.10	1.98	1.91	1.89	1.81	1.84	2.15	1.87
m2	2.01	1.89	2.34	2.10	1.93	1.79	1.98	1.96	2.10	2.03
m3	1.82	1.68	2.16	1.78	-	-	-	-	1.86	1.57

Table 2. Results of Tukey's HSD tests of mean m1 length (to left of diagonal) and mean m2 length (right of diagonal) among *Acarechimys* samples. Only p-values for significant differences ($p < 0.01$) are listed. Abbreviations: N = number of specimens.

	A. constans (N = 11)	A. gracilis (N = 13)	A. leucotheae (N = 1)	A. minutissimus (N = 14)	A. minutus (N = 9)	<i>Quebrada Honda</i> (N = 14)
A. constans (N = 15)	-	(none)	0.0170	< 0.0001	< 0.0001	(none)
A. gracilis (N = 14)	(none)	-	0.0143	< 0.0001	< 0.0001	(none)
A. leucotheae (N = 1)	0.0004	0.0014	-	(none)	(none)	(none)
A. minutissimus (N = 15)	< 0.0001	< 0.0001	(none)	-	0.0346	< 0.0001
A. minutus (N = 11)	< 0.0001	< 0.0001	(none)	0.0042	-	0.0017
<i>Quebrada Honda</i> (N = 13)	0.0047	(none)	0.0345	< 0.0001	0.0080	n/a

Table 3. Main dental and mandibular characters of *Acarechimys*. Abbreviations: see Figure 1.

	<i>A. minutus</i>	<i>A. constans</i>	<i>A. gracilis</i>	<i>A. leucotheae</i>	<i>A. minutissimus</i>
<i>dp4</i>					
<i>Metalophulid II</i>	Variably-developed (usually reduced)	Variably-developed (conspicuous or reduced)	Well-developed	Reduced	Reduced to a spur
<i>Mesolophid</i>	Usually reduced	Well-developed	Reduced	Well-developed	Well-developed
<i>m1</i>					
<i>Metalophulid II</i>	Reduced	Variably developed (usually conspicuous)	Variably developed	Reduced	Reduced to a spur or absent
<i>Pamd</i>	Present	Absent	Present	Absent	Absent
<i>Pemed I</i>	Variably Present	Variably present	Variably-developed	Absent	Absent
<i>Accessory cusp</i>	Present	Variably present	Present	Present	Present/absent
<i>m2</i>					
<i>Metalophulid II</i>	Reduced	Reduced	Reduced	Reduced	Absent
<i>Pamd</i>	Present	Absent	Present	Absent	Absent
<i>Pemed I</i>	Variably present	Variably present	Variably present	Absent	Absent
<i>Accessory cusp</i>	Present	Variably present	Present	Present	Present/absent
<i>m3</i>					
<i>Metalophulid II</i>	Absent	Reduced	Reduced/absent	Reduced	Absent
<i>Pamd</i>	present	Absent	present	Absent	Absent
<i>Pemed I</i>	Absent	absent	Variably present	Absent	Absent
<i>Accessory cusp</i>	Absent	Variably present	Present	present	Present
<i>Lower incisor</i>					
<i>Laterally compress</i>	yes	no	no	-	yes
<i>Mandible</i>					
<i>Anterior border</i>	Below <i>dp4</i>	Below <i>dp4-m1</i>	Below <i>dp4-m1</i>	Below <i>dp4</i>	Below <i>dp4</i>
<i>nmmpio</i>					
<i>Mental foramen</i>	Absent	Usually absent	Usually absent	Absent	Absent

Supplementary Appendix 1. List of specimens assigned to *Acarechimys*. Abbreviations: CL, coastal localities; CSCr, cliffs of the Santa Cruz river; Fm, Formation; Mb, Member; SCP, Santa Cruz Province. Within parentheses are previous systematic assignments. For the 'old collections' localities we translate what is written in the collection labels. For a discussion about the localities names see Vizcaíno *et al.* (2012) and Fernicola *et al.* (2014).

<i>Acarechimys minutus</i>	Description	Locality	Horizon
MACN-A 237 (syntype of <i>Acaremys minutus</i>)	Right mandible with dp4-m3	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 238 (syntype of <i>Acaremys minutus</i>)	Right mandible with dp4(broken)-m3	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4070	Left maxilla with DP4-M3	Seuhen, SCP	Santa Cruz Fm, late early Miocene
MACN-A 4071	Right mandible with dp4-m3 and incisor	Seuhen, SCP	Santa Cruz Fm, late early Miocene
MACN-A 4072a	Right mandible with dp4-m3	Monte Observación, SCP	Santa Cruz Fm, late early Miocene
MACN-A 4073	Left mandible with dp4-m2	Monte Observación, SCP	Santa Cruz Fm, late early Miocene
MACN-A 4082	Right mandible with dp4-m3	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4264	Left mandibular fragment with dp4	Seuhen, SCP	Santa Cruz Fm, late early Miocene
MACN-PV SC4098	Right mandible with dp4-m1	Karaiken, SCP	Santa Cruz Fm?, late early Miocene
MLP 15-410a (syntype of <i>Acaremys minutus</i>)	Left mandibular fragment with m1-m3	CSCr, SCP	Santa Cruz Fm, late early Miocene
MLP 82-XII-1-31	Right mandible with m1	Santa Cruz Province	Santa Cruz Fm, late early Miocene
MLP 91-IX-1-199a	Right m1 or m2	Cañadón del Tordillo, Neuquén Province	Collon Curá Fm, middle Miocene
MLP 91-IX-1-200	Left m1 or m2	Cañadón del Tordillo, Neuquén Province	Collon Curá Fm, middle Miocene
MMP-PV 4193	Left mandible with dp4-m3	Puesto La Costa, SCP	Santa Cruz Fm, late early Miocene
MMP-PV 15039	Right mandible with dp4-m2	Cañadón Silva, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15087	Right m2	Estancia el Tordillo, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15088	Left maxilla with DP4-M3	Estancia el Tordillo, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15089	Right mandible with dp4-m3 and incisor	Rincón del Buque, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15090	Right mandible with dp4-m3	Rincón del Buque, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17435	Right mandible with m1-m2	Rincón del Buque, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17436	Right mandibular fragment with dp4	Estancia El Tordillo, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17437	Left dp4	Estancia El Tordillo, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17440	Right maxilla with DP4	Rincón del Buque2, SCP	Santa Cruz Fm, late early Miocene
UATF-V-001262	Skull with left and right DP4-M3	Quebrada Honda, Bolivia	Unnamed Fm, middle Miocene

YPM-PU 15806	Skull with left and right DP4-M3 and incisors	Killik Aike, SCP	Santa Cruz Fm, late early Miocene
<i>Acarechimys</i> cf. <i>A. minutus</i> MLP 91-IV-1-15	Left mandible with m1-m2	Cañadón del Tordillo, Neuquén Province	Collón Curá Fm, middle Miocene
UATF-V-000934	Right mandible with dp4-m2 and incisor	Quebrada Honda, Bolivia	Unnamed Fm, middle Miocene
UATF-V-000935	Right mandible with dp4-m3	Quebrada Honda, Bolivia	Unnamed Fm, middle Miocene
UATF-V-000950	Right mandible with dp4(broken)-m3	Quebrada Honda, Bolivia	Unnamed Fm, middle Miocene
<hr/> <i>Acarechimys constans</i> <hr/>			
MACN-A 247 (syntype of <i>Stichomys constans</i>)	Left mandible with m1-m2 and right mandible with dp4-m2	SCP	Santa Cruz Fm, late early Miocene
MACN-A 4058	Right mandible with dp4-m1	Monte Observación, SCP	Santa Cruz Fm, late early Miocene
MACN-A 4061	Left mandible with dp4-m3	Monte Observación, SCP	Santa Cruz Fm, late early Miocene
MACN-A 4064	Left mandible with dp4-m2 and broken incisor	Monte Observación, SCP	Santa Cruz Fm, late early Miocene
MACN-A 4074	Left mandible with dp4-m1	Monte Observación, SCP	Santa Cruz Fm, late early Miocene
MACN-A 4075 (syntype of <i>Acaremys minutus</i>)	Right mandible with dp4-m1	Monte Observación, SCP	Santa Cruz Fm, late early Miocene
MLP 15-57 (syntype of <i>Stichomys constans</i>)	Right mandible with m1 (broken)-m3	Santa Cruz	Santa Cruz Fm?, late early Miocene?
MLP 15-200 (syntype of <i>Stichomys constans</i>)	Right mandible with dp4-m2 (lost specimen)	CSCr	Santa Cruz Fm, late early Miocene
MLP 15-319	Right mandible with m1-m3	CSCr	Santa Cruz Fm, late early Miocene
MLP 15-391 (syntype of <i>Stichomys constans</i>)	Right mandible with dp4-m3	CSCr, SCP	Santa Cruz Fm, late early Miocene
MLP 15-391c (syntype of <i>Stichomys constans</i>)	Left mandible with broken dp4-m1	Suppose 'curso inferior del río Santa Cruz'	Santa Cruz Fm, late early Miocene
MPM-PV 4223	Right mandible with dp4-m2	Puesto Estancia La Costa, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15002	Broken right mandible with dp4-m1, palatal fragment with both DP4-M2, and isolated vertebra	Puesto Estancia La Costa, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15091	Right mandible with dp4-m2	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15092	Right mandible with dp4-m2 and incisor	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15093	Right mandible with dp4-m2	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15094	Left mandible with dp4-m1	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15095	Right mandible with m1-m3	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15096	Left mandible with dp4-m2	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15097	Right mandible with m2-m3	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
<hr/> <i>Acarechimys gracilis</i> <hr/>			
MACN-A 52-128 (holotype of <i>Protacaremys pulchellus</i>)	Left mandible with dp4-m3	Barranca sur del Colhue Huapí Lake, Chubut	Sarmiento Fm, Colhue Huapí Mb, early

MACN-A 4060	Left mandible with dp4-m3 and incisor	Province Monte Observación, Santa Cruz	Miocene Santa Cruz Fm, late Early Miocene
MACN-A 4068	Right mandible with dp4-m2	Monte Observación, SCP	Santa Cruz Fm, late Early Miocene
MACN-A 4263 (holotype of <i>Stichomys gracilis</i>)	Left mandible with dp4-m2	Seuhen, SCP	Santa Cruz Fm, late Early Miocene
MLP 15-39 (syntype of <i>Stichomys constans</i>)	Left mandible with m1-m3	CSCr	Santa Cruz Fm, late Early Miocene
MLP 15-346 (syntype of <i>Stichomys constans</i>)	Left mandible with dp4-m2	Santa Cruz river?	Santa Cruz Fm?, late early Miocene
MLP 15-391a (syntype of <i>Stichomys constans</i> and holotype of <i>Acarechimys pascuali</i>)	Right mandible with dp4-m2	Suppose 'curso inferior del río Santa Cruz'	Santa Cruz Fm, late early Miocene
MLP 15-391b (syntype of <i>Stichomys constans</i>)	Right mandible with m1-m3	Suppose 'curso inferior del río Santa Cruz'	Santa Cruz Fm, late early Miocene
MPM-PV 17430	Left mandible with dp4-m3	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17431	Right mandible with m1-m3	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17432	Right mandible with dp4(broken)-m3	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17433	Left mandible with m1-m2	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17434	Left mandible with dp4-m2	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17439	Left m1 in mandibular fragment	Estancia El Tordillo, Santa Cruz	Santa Cruz Fm, late early Miocene
<i>Acarechimys leucotheae</i>			
MPEF-PV 10677	Left mandible with dp4-m3 and incisor	Cabeza Blanca, Chubut Province	Sarmiento Fm, late Oligocene
<i>Acarechimys minutissimus</i>			
IGM 184229	Left mandible with dp4-m3	La Venta, Colombia	Villa vieja Fm, middle Miocene
MACN-A 256 (syntype of <i>Acaremys minutissimus</i>)	Left mandible with dp4-m3	SCP	Santa Cruz Fm, late early Miocene
MACN-A 257 (syntype of <i>Acaremys minutissimus</i>)	Left mandible with dp4-m3	SCP	Santa Cruz Fm, late early Miocene
MACN-A 258 (syntype of <i>Acaremys minutissimus</i>)	Right mandible with dp4-m3 and incisor	SCP	Santa Cruz Fm, late early Miocene
MACN-A 1894	Right mandible with dp4-m3	Exact provenance unknown	
MACN-A 1896	Left mandible with dp4-m3	Exact provenance unknown	
MACN-A 1897	Right mandible with m1-m2	Exact provenance unknown	
MACN-A 1898	Left mandible with m2	Exact provenance unknown	
MACN-A 4072b	Right mandible with dp4-m2	Monte Observación, Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4076	Right mandible with dp4-m3 and incisor	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4077	Mandíbula izquierda casi	Santa Cruz	Santa Cruz Fm, late

MACN-A 4079	completa con m1-m3 e incisivo Right mandible with m1-m3 and incisor	Santa Cruz	early Miocene Santa Cruz Fm, late early Miocene
MACN-A 4080	Right mandible with dp4-m3	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A4081	Right mandible with dp4-m1	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4083	Left mandible with dp4-m2	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4084	Edentulous left mandible with incisor	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4085	Right edentulous mandible and a m1 or m2 stick on the mandible	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4086	Left mandible with m1-m2 and a m3 stick on the mandible	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4087	Right mandible with dp4-m2 and incisor	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4088	Left mandible with m1-m3	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4089	Left mandible with m1-m3	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4090	Right mandible with m1-m3 and incisor	Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4091	Left mandible with dp4-m1 and incisor	Territorio de Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4092	Right mandible with dp4-m2 and incisor	Territorio de Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4093	Left mandible with m1-m3	Territorio de Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4094	Right mandible with dp4-m2 and incisor	Territorio de Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4145 (syntype <i>Sciamys tenuissimus</i>)	Right maxillary fragment with DP4	Monte Observación, SCP	Santa Cruz Fm, late early Miocene
MACN-A 4146 (syntype <i>Sciamys tenuissimus</i>)	Left mandibular fragment with dp4 and right mandibular fragment with m2	Monte Observación, SCP	Santa Cruz Fm, late early Miocene
MACN-A 4265 (holotype <i>Stichomys diminutus</i>)	Right dp4-m1	Monte Observación, SCP	Santa Cruz Fm, late early Miocene
MACN-A 4266	Fragmento mandibular derecho con dp4-m1	Monte Observación, SCP	Santa Cruz Fm, late early Miocene
MACN-A10093	Fragmento mandibular derecho con dp4-m2	Yacimiento Dipilus 1897- 98, SCP	Santa Cruz Fm, late early Miocene
MACN-A 12683	Right mandible with dp4-m2; right maxilla with DP4-M2	Santa Cruz Province?	Santa Cruz Fm, late early Miocene
MACN-PV SC2158	Right mandible with dp4 and incisor	Portezuelo Sumich Norte, SCP	Pinturas Fm, late early Miocene
MACN-PV SC2589	Right dp4-m2	Toldos Sur, SCP	Pinturas Fm, late early Miocene
MACN-PV SC2590	Right maxillary with DP4-M1	Toldos Sur, SCP	Pinturas Fm, late early Miocene
MACN-PV SC4045	Right mandible with m1-m3	Gobernador Gregores, SCP	Pinturas Fm, late early Miocene
MACN-PV SC4091	Left mandible with dp4-m1	Lago Cardiel, SCP	Pinturas Fm, late early Miocene
MLP 15-1	Left mandible with dp4-m3 and	Santa Cruz	Santa Cruz Fm?, late

MLP 15-188 (syntype of <i>Acaremys minutissimus</i>)	incisor		early Miocene?
MLP 15-188a	Left mandible with dp4-m3	CSCr, SCP	Santa Cruz Fm, late early Miocene
MLP 15-398	Right mandible with dp4-m2 and incisor	CSCr	Santa Cruz Fm, late early Miocene
MLP 15-408	Left mandible with dp4-m2	CSCr	Santa Cruz Fm, late early Miocene
MLP 91-IX-1-95	Left mandible with m1-m3	CSCr	Santa Cruz Fm, late early Miocene
MLP 91-IX-1-199b	Left dp4	Cañadón del Tordillo, Neuquén Province	Collon Curá Fm, middle Miocene
MLP 91-IX-1-201a	Right m3	Cañadón del Tordillo, Neuquén Province	Collon Curá Fm, middle Miocene
MLP 91-IX-1-201b	Right dp4	Cañadón del Tordillo, Neuquén Province	Collon Curá Fm, middle Miocene
MLP 91-IX-1-201c	Right m1-m2	Cañadón del Tordillo, Neuquén Province	Collon Curá Fm, middle Miocene
MLP 91-IX-2-168	Left M1 or M2	Cañadón del Tordillo, Neuquén Province	Collon Curá Fm, middle Miocene
MPM-PV 15098	Left dp4	Estancia Collon Curá, Neuquén Province	Collon Curá Fm, middle Miocene
MPM-PV 15099	Left mandible fragment with dp4 and incisor	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15100	Left mandible with m1-m2 and incisor	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15101	Right mandible with dp4(broken)-m2	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 15102	Right mandible with dp4-m2	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17426	Left mandible with m1	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17427	Right mandible with dp4-m1	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17428	Right mandible with dp4-m2 and incisor	Estancia Cordón Alto, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17438	Left mandible with dp4-m2	Estancia El Tordillo, CSCr, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17429	Left mandible with dp4-m2	Rincón del Buque 2, SCP	Santa Cruz Fm, late early Miocene
MPM-PV 17441	Left dp4	Rincón del Buque 2, SCP	Santa Cruz Fm, late early Miocene
YPM-PU 15178	Partially preserved skull with right DP4-M2 and left DP4-M1	10 miles South of the Coy Inlet, SCP	Santa Cruz Fm, late early Miocene
<i>Acarechimys</i> cf. <i>A. minutissimus</i>			
UATF-V-000960	Right mandibular fragment with dp4-m1	Quebrada Honda, Bolivia	Unnammed fm, middle Miocene
<hr/>			
<i>Acarechimys</i> sp.			
MACN-A 1895	Left mandible with m1	Santa Cruz Province?	Santa Cruz Fm, late early Miocene
MACN-A 4062	Left mandible with dp4-m3	Monte Observación, Santa Cruz	Santa Cruz Fm, late early Miocene
MACN-A 4067	Left mandible with dp4-m3	Monte Observación, Santa Cruz	Santa Cruz Fm, late early Miocene

Supplementary appendix 2. List of characters used in the phylogenetic analysis: an expanded version of the data matrix used by Arnal and Vucetich (2015). New characters are at the end of the list (168–186) and those from Verzi *et al.* (2016) are indicated as such; characters marked with an asterisk are modified with respect to the original version. The following characters are treated as ordered: 3, 14, 15, 22, 31, 38, 39, 44, 52, 54, 55, 59, 61, 67, 69, 75, 80, 83, 84, 99, 103, 104, 107, 109, 113, 118, 119, 128, 148, 161, and 164.

- (1) P3/DP3: present (0), absent (1).
- (2) Cingula: present (0), absent (1).
- (3) Hypsodonty: brachydont (0), mesodont (1), protohypsodont (2), euhypsodont (3).
- (4) Roots number on upper molars: three (0), four (1).
- (5) Tooth row: straight (0), convex (1).
- (6) Tooth rows: parallel (0), anteriorly slightly convergent (1), anteriorly very convergent (2), anteriorly slightly divergent (3).
- (7) Crest thickness in occlusal view: slenderer than flexi/ids (0), equal or broader than flexi/ids (1).
- (8) Deciduous premolars (Marivaux *et al.*, 2004): normal replacement (0), retention (1).
- (9) Terraced occlusal surface: present (0), absent (1).
- (10) Cusp differentiation: yes, labial cusps of upper molars and lingual cusps of lower molars wider than their associated crests (0), no, cusps indistinct, entirely submerged in their associated crests (1).
- (11) Enamel on upper molars: complete and uniformly distributed (0), complete and not uniformly distributed (1), interrupted on the labial side (2), interrupted on the posterior wall of the posterior lobe (3).

- (12) Mesolophule on DP4: present (0), absent (1).
- (13) Metaloph on DP4: present and joined lingually to the anterior arm of the hypocone (0), present and joined lingually to the posteroloph (1), indistinct, probably fused to the posteroloph (2), connected to the metaconule (3).
- (14) Anterior arm of the metacone on DP4 (Antoine *et al.*, 2012): absent (0), weakly pronounced (1), high (2).
- (15) Mesostyle on DP4 (Antoine *et al.*, 2012): indistinct or absent (0), moderate (1), strong (2).
- (16) Hypocone on DP4: small (0), moderate (as large as the protocone) (1).
- (17) Size of P4 respect M1: $P4 \geq M1$ (0), $P4 < M1$ (1).
- (18) Crown outline of P4: transverse (0), oval (1), heart shape (2), quadrangular (3), reversed heart shape (4), subtriangular (5).
- (19) Anterocingulum on P4: small, short (0), long (1).
- (20) Anterocingulum on P4 (Antoine *et al.*, 2012): low (0), high (anteroloph) (1).
- (21) Anterocingulum (or anteroloph) – paracone connection (Antoine *et al.*, 2012): absent (0), present (via a parastyle or not) (1).
- (22) Mesolophule on P4 (Antoine *et al.*, 2012): absent (0), short (1), reaches the buccal side (2).
- (23) Metaloph on P4: present (0), absent (1).
- (24) Metaloph on P4: connected to the metaconule (0), connected to the anterior arm of the hypocone (1), connected to the posteroloph (2).
- (25) Metacone on P4: small (0), strong (1).
- (26) Hypocone on P4: absent (0), present (1).
- (27) Hypocone on P4: labial to protocone (0), lingually aligned to protocone (1).

- (28) Anterior arm of the hypocone on P4 (Antoine *et al.*, 2012): absent (0), present (1).
- (29) Posteroloph – metacone connection on P4 (Antoine *et al.*, 2012): absent (0), present (1).
- (30) Hypoflexus on P4: absent (0), present (1).
- (31) Hypoflexus on P4: as a superficial lingual groove (0), as a moderately deep lingual groove separating protocone and hypocone, less penetrating than in molars (1), very deep, as in molars (2).
- (32) Endoloph on P4: absent (0), present (1).
- (33) Figure eight dental pattern in upper molars: absent (0), present (1).
- (34) Crest obliquity on M1-M3: transversal to the anteroposterior axis of the teeth (0), anterolabially-posterolingually oblique (1). All crests should be oblique to consider character state 1.
- (35) Anteroloph on M1-M3: moderately high (0), high (1).
- (36) Anteroloph on M1-M3: short, not reaching the paracone (0), reaches the paracone (1), long, reaches the labial border of the crown but not connected to the paracone (2).
- (37) Paracone on M1-M3: larger than metacone (0), equal to metacone (1).
- (38) Posterior arm of the paracone (Antoine *et al.*, 2012): absent (0), weakly pronounced (1), strong and high (2).
- (39) Paracone – metacone position on M2 (Antoine *et al.*, 2012): mesiodistally opposed (0), metacone slightly lingual (1); metacone strongly lingual (2).
- (40) Anterolingual angle of the tooth on M1-M3: rounded or forming an obtuse angle (0), forming a right angle (1).
- (41) Mesolophule on M1-M3: present (0), absent (1).

- (42) Direction of the mesolophule: straight (transverse) (0), slightly oblique (distobuccally oriented) (1).
- (43) Metaloph on M1-M3: lingually joined to the anterior arm of the hypocone (0), lingually joined to the posteroloph (1), indistinct, probably fused to the posteroloph (2), lingually joined to the mesolophule (3).
- (44) Anterior arm of the metacone: absent (0), weakly pronounced (1), high (2).
- (45) Hypocone on M1: labial to protocone (0), lingually aligned to protocone (1).
- (46) Hypocone on M2: labial to protocone (0), lingually aligned to protocone (1).
- (47) Posteroloph – metacone connection (Antoine *et al.*, 2012): absent (0), present (1).
- (48) Mure in juveniles/sub-adults: absent (0), present (1).
- (49) Mure connection (Antoine *et al.*, 2012): on the protoloph (central to the tooth) (0), on the protoloph more lingually (1).
- (50) Metacone size on M3 (Antoine *et al.*, 2012): distinct cusp (as large as the paracone) (0), reduced but distinct (1), crestiform (2).
- (51) Hypocone position in relation to the protocone on M3: more labial (0), strongly more labial (1), at the same level (2).
- (52) Hypoflexus orientation on M1-M3: anteriorly oblique (0), slightly anteriorly oblique or transverse to the anteroposterior axis of the tooth (1), posteriorly oblique (2).
- (53) Mesoflexus groove respect metaflexus groove in M1-M2: mesoflexus groove deeper than metaflexus groove (0), mesoflexus groove equal than metaflexus groove (1).
- (54) Size of M1 / M2: M1 < M2 (0), M1 = M2 (1), M1 > M2 (2).

- (55) Size of M3 / M2 (Antoine *et al.*, 2012): M3 < M2 (0), M3 = M2 (1), M3 > M2 (2).
- (56) M1 length / width proportions: length = width (0), length < width (1), length > width (2).
- (57) M2 length / width proportions: length = width (0), length < width (1), length > width (2).
- (58) Upper incisors: laterally compressed (0), laterally no compressed (1).
Laterally compressed when anteroposterior diameter / transverse diameter ≥ 1.5 .
- (59) Posterior arm of the metaconid (metastylar fold) on dp4 (Marivaux *et al.*, 2004): absent (0), weak and low (1), well-developed and high (2).
- (60) Anteroconid on dp4 (Marivaux *et al.*, 2004): present (0), absent (1).
- (61) Metaconid position respect protoconid on dp4: anterior (0), aligned (1), posterior (2).
- (62) Metalophulid I on dp4: present (0), absent (1).
- (63) Metalophulid II on dp4: present (0), absent (1).
- (64) Metalophulid II on dp4: conspicuous (0), reduced (1).
- (65) Mesolophid on dp4: present (0), absent (1).
- (66) Mesolophid on dp4: conspicuous (0), reduced (1).
- (67) Ectolophid on dp4: absent (0), mesially interrupted (unconnected to the protoconid) (1), complete (connected to the protoconid) (2).
- (68) Metaconid position respect protoconid on p4: anterior (0), aligned (1).
- (69) Posterior arm of the metaconid (metastylar fold) on p4: absent (0), weak and low (1), well-developed, high, and long (2).
- (70) Flexid on anterior aspect of metalophulid I on p4 (Vucetich and Kramarz, 2003): present (0), absent (1).

- (71) Metalophulid II on p4: absent (0), present (1).
- (72) Mesolophid on p4: absent (0), present (1).
- (73) Mesolophid on p4 (Antoine *et al.*, 2012): short (0), long, reaches the lingual side (1).
- (74) Hypolophid on p4 (Marivaux *et al.*, 2004): absent (0), present (1).
- (75) Anterior arm of the hypoconid on p4 (Antoine *et al.*, 2012): absent (0), thin (1), strong (2).
- (76) Hypoconulid on p4 (Antoine *et al.*, 2012): minute to absent (0), moderate (1).
- (77) Posterolophid – entoconid connection (Antoine *et al.*, 2012): absent (0), present (1).
- (78) Talonid on p4: sub-equal (0), wider than the trigonid (1).
- (79) Size of p4 / m1: $p4 \geq m1$ (0), $p4 < m1$ (1).
- (80) Proportion of p4: length clearly $>$ width (0), length $>$ width (1), length \geq width (2).
- (81) Enamel on lower molars: complete and uniformly distributed (0), complete and not uniformly distributed (1), interrupted on the anterior face and anterior face of hypoflexid (2).
- (82) Eight occlusal pattern on upper molars: absent (0), present (1).
- (83) Anterofossettid and metafossettid on lower molariforms: persistent (0), ephemeral (1), absent (2). Fossetids are considered ephemeral when they are lost in juvenile/sub-adult specimens.
- (84) Metaconid position respect protoconid on m1-m3: anterior (0), aligned (1), posterior (2).
- (85) Metalophulid II on m1-m2: present (0), absent (1).
- (86) Metalophulid II on m1-m2: complete (0), reduced (1).

- (87) Metalophulid II connection: mesiobuccally to the metaconid (0), distolingually to the metaconid, on the posterior arm of the metaconid (1), distolingually to the metaconid, on the mesostylid (2), do not contact neither (3).
- (88) Metalophular spur : absent (0), short (1).
- (89) Entoconid position respect hypoconid on m1-m3: aligned (0), anterior (1).
- (90) Posterolophid on m1-m2: short (0), long (1).
- (91) Posterolophid – entoconid connection (Antoine *et al.*, 2012): absent (0), present (1).
- (92) Crest obliquity on m1-m3: transversal to the anteroposterior axis of the teeth (0), anterolabially-posterolingually oblique (1). All crests should be oblique to consider character state 1.
- (93) Anterior arm of the hypoconid: absent in unworn or little worn teeth (0), present in all stages (1).
- (94) Mesiodistal pinch of the hypoconid (Antoine *et al.*, 2012): absent (0), present (1).
- (95) Hypoconulid on m1-m3: recognizable (0), indistinct (1).
- (96) Spur of the posterior margin of metalophulid I on m1-m2: present (0), absent (1).
- (97) Accessory cusp posterior to metalophulid I on m1-m2: present (0), absent (1).
- (98) Posterior arm of metaconid on m1-m2: present (0), absent (1).
- (99) Hypoflexid orientation on m1-m3: posteriorly oblique, or opposed to the metaflexid or hypolophid (0), transverse or opposed to the hypolophid or mesoflexid (1) anteriorly oblique or opposed to the anteroflexid (2).
- (100) Size of m3 / m2 (Marivaux *et al.*, 2004): m3 = m2 (0), m3 < m2 (1).
- (101) Lower incisors: laterally compressed (0), laterally no compressed (1).
Laterally compressed when anteroposterior diameter is at least 1.5 the width.

- (102) Anterior face of lower incisors: curve (0), forming a right lingual border and a curved labial one (1), plane (2).
- (103) Lower incisors: long, passing beneath m3 (0), the base reaching m3 (1), short, the base does not reaches m3 (2).
- (104) Lower incisors enamel microstructure: multiseriate HSB with acute IPM (0), multiseriate HSB with transitional IPM (1), multiseriate HSB with rectangular IPM (2).
- (105) Diastema length: shorter than the p4(dp4)-m1 distance (0), equal or larger than the p4(dp4)-m1 distance (1).
- (106) Mental foramen: present (0), absent (1).
- (107) Development of the mental foramen: small (0), conspicuous (1), large (2).
- (108) Orientation of the mental foramen: externally oriented (0), anteriorly oriented (1), dorso-anteriorly oriented (2).
- (109) Position of the mental foramen: anterior to the lowest part of the diastema (0), at the lowest part of the diastema (1), beneath p4/dp4 (2).
- (110) Position of the anterior border of notch for the masseter muscle pars infraorbitalis (nmmpio): beneath m1 (0), anterior to m1 (1).
- (111) Position of the nmmpio respect mandible high: above the mid high (0), at the middle of the mandible high (1).
- (112) Origin of the masseteric crest: includes the notch for the masseter muscle pars infraorbitalis (0), does not include the notch for the masseter muscle pars infraorbitalis (1).
- (113) Depth of the anterior portion of the masseteric fossa: shallow or flat (0), moderately deep (1), deep (2).
- (114) Anterior margin of the coronoid process: convex (0), straight (1), concave (2).

- (115) Mandibular notch: conspicuous (0), poorly developed or absent (1).
- (116) High of the coronoid process respect the mandibular condyle: same high (0), ventral to the condyle (1).
- (117) High of the mandibular condyle: higher than the occlusal surface (0), as the occlusal surface (1).
- (118) Posterior border of the mandibular symphysis: anterior to the premolars (0), at the level of the premolars (1), posterior to the premolars (2). It is measure with the tooth row horizontal to the floor.
- (119) Posterior extension of the premaxillaries related to nasals: shorter (0), equal (1), longer (2).
- (120) Frontal extension between nasals and premaxillaries: absent (0), present (1).
- (121) Nasals shape: parallel lateral margins (0), lateral margins wider anteriorly (1).
- (122) Incisor included into the rostral masseteric fossa: no (0), yes (1).
- (123) Incisor foramina: length (0), short (1). They are considered long when its length is equal or larger than the half of the length of the diastema.
- (124) Incisor foramina shape: laterally narrow (0), anteriorly narrow (1), posteriorly narrow (2), both extremes acute (3). They are considered narrow when they are equally wide along the length of the foramina.
- (125) Premaxillary-maxillary suture: at the posterior border of the incisor foramina (0), at the middle length of the incisor foramina (1); posterior to the incisor foramina (2).
- (126) Diastemal ridges: absent (0), present (1).
- (127) Diastemal ridges: poorly developed (0), conspicuous (1).
- (128) Frontals: wider anteriorly than posteriorly (0), straight lateral margins (1), narrower anteriorly than posteriorly (2); concave lateral margins (3).
- (129) Post-orbital process: absent (0), present (1).

- (130) Post-orbital process: small (0), conspicuous (1).
- (131) Conformation of the post-orbital process: build only by the frontal (0), build by the frontal and parietal (1).
- (132) Post-orbital constriction: absent (0), present (1).
- (133) Post-orbital constriction: small (0), conspicuous (1).
- (134) Position of the zygomatic dorsal root (ZDR): anterior to M1 (0), at the level of M1 or M1-M2 (1).
- (135) Exposition of the lacrimal onto the vertical or ZDR: little exposed (0), conspicuous (1), no exposed (2).
- (136) Ventral root of the zygomatic arch: similar anteroposterior and dorsoventral diameters (0), anteroposterior diameter twice dorsoventral or more (1).
- (137) ZVR respect the palatal level: at the same level (0), ZVR dorsal (1).
- (138) ZDR respect ZVR: aligned (0), posterior (1).
- (139) Groove for the infraorbitalis nerve within the infraorbital foramen: absent (0), present (1).
- (140) Groove for the passage of the infraorbitalis nerve within the infraorbital foramen: present with a small lateral rim (0), present with a large lateral rim (1).
- (141) Masseteric tuberosity (for the insertion of the masseteric superficial muscle): poorly developed (0), well developed (1).
- (142) Paraorbital process: present (0), absent (1).
- (143) Paraorbital process: build by the jugal and squamosal (0), build by the squamosal (1), build by the jugal (2).
- (144) Jugal fossa: present (0), absent (1).
- (145) Depth of the jugal fossa: superficial (0), deep (1).
- (146) High of the jugal fossa: low (0), high (1).

- (147) Length of the jugal fossa: antero posteriorly short (0), antero posteriorly long (1). It is considered long when its length is equal or longer than the length of the horizontal ramus of the zygoma.
- (148) Etmoidal foramen: at the level of M3 (0), at the level of M2-M3 (1), at the level of the M2-M1 (2); at the level of the M2 (3).
- (149) Sphenopalatine foramen: anterior to the M1 (0), at the level of M1 (1), posterior to M1 (2), groove located at the DP4-M1 level (3), groove located at the M1-M2 level (4).
- (150) Posterior palatine foramina: between palatines and Mx at the M1 (0), into the Mx at the level of the premolar (1), into the Mx at the level of M1.
- (151) Posterior palatine foramina: small (0), conspicuous (1).
- (152) Posterior nares: at the level of M3 (0), at the level of M2 (1).
- (153) Spheno-palatine vacuities: absent (0), present (1).
- (154) Spheno-palatine vacuities: small (0), conspicuous (1).
- (155) Buccinator and masticatory foramina: separated (0), fused (1).
- (156) Oval foramen: bounded by the aliesfenoid (0), bounded posteriorly by the tympanic bulla (1).
- (157) Ventral extension of the lateral process of the supraoccipital: exceeds the dorsal border of the bulla (0), extends until the dorsal border of the bulla (1); do not reach the dorsal border of the bulla (2).
- (158) Parietals on the skull roof: reach or are close to the occiput (0), retracted and not near the occiput (1).
- (159) Dorsal extension of the mastoid exposure: do not exceeds the dorsal border of the bulla (0), exceeds the dorsal border of the bulla (1), mastoid exposed on the skull roof (2).

- (160) Mastoid exposure on the occiput: absent (0), present (1).
- (161) Shape of the mastoid exposure: concave (0), plane (1), convex (2).
- (162) Dorsal exposition of the petrosal: absent (0), present (1).
- (163) Epitympanic sinus: small (0), conspicuous (1).
- (164) Hypotympanic recess: small (0), inflated (1), hypertrophied (2).
- (165) Tympanic fenestra below MAE: absent (0), present (1).
- (166) Tympanic fenestra below MAE: small (0), conspicuous (1).
- (167) Paraoccipital process: ventrally oriented with its tip separated from the bulla and well developed (0), ventrally oriented, with its tip fused to the bulla (1), short, laterally oriented and completely fused to the bulla (2), short, ventrally oriented and completely fused to the bulla (3).
- (168) Premaxillary septum separating incisive foramina (Verzi *et al.*, 2016): with posterior ends of premaxillae joined medially, forming a pointed or rounded projection which may join an anterior apophysis of the maxilla (0), with posterior ends of premaxillae divergent, each one forming a small lateral apophysis (1).
- (169) Lacrimal foramen (Verzi *et al.*, 2016): opens into the orbital portion of the lacrimal (0); opens into the maxilla (1).
- (170) Portion of maxilla surrounding foramen into nasolacrimal canal (Verzi *et al.*, 2016)*: with a suture posterior to the foramen (0); continuous around the foramen (1); no portion of the mx surrounding the lacrimal foramen (2)
- Character stated 2 was included in order to score *Dasyprocta azarae*.
- (171) Foramen into nasolacrimal canal (Verzi *et al.*, 2016): opens into maxilla only (0); surrounded posteriorly by lacrimal (1).

(172) Relationship between zygoma and orbital region (Verzi *et al.*, 2016): dorsal margin of the zygoma concave, not restricting orbital region (0), dorsal margin of the zygoma very slightly concave or straight, restricting orbital region (1).

We do not clearly understand the last part of each character state (not restricting orbital region/ restricting orbital region) since we interpreted that a concave dorsal margin of the zygoma should restrict, or delimit (if this is the sense), the orbital region. Nevertheless, we described this character as was done in Verzi *et al.* (2016) but scored them taking into account if the zygoma is concave or very slightly concave.

(173) Contact among maxilla, lateral palatine plate and alisphenoid in basitemporal region (Verzi *et al.*, 2016): located posterior to the M3 alveolus (0); lateral to the M3 alveolus (1).

(174) Pterigoid fossa in ventral view (between alisphenoid bridge and anterior margin of lateral palatine plate) (Verzi *et al.*, 2016)*: subcircular, with anteroposterior and transverse diameters subequal (0), suboval, with anteroposterior diameter greater than transverse one (1), suboval, with labiolingual diameter greater than anteroposterior one (2).

Character stated 2 was included in order to scored *Myocastor coypus*.

(175) Lateral margin of pterygoid fossa (Verzi *et al.*, 2016): oriented posterodorsally and not forming a flange extending posteriorly (0); forming a flange level with the medial margin and extending posteriorly toward the bulla (1).

(176) Ventral margin of posterior process of squamosal (Verzi *et al.*, 2016): not laterally deflected (0); laterally deflected, thus forming a shelf (1).

(177) Posterior process of squamosal (Verzi *et al.*, 2016): straight and deep at its origin, with its posterior portion wide (0); lower (dorso-ventrally narrow) at its

origin and with posterior portion narrow due to development of the epitympanic recess (petrosal bulla) (1).

- (178) Lateral process of supraoccipital (Verzi *et al.*, 2016)*: short, located dorsal to mastoid process (0); long, ventrally extended overlapping the mastoid process or below the level of the latter (1); reaches the middle dorso-ventral length of the mastoid exposure (2).

Character stated 2 was included in order to modified some scores of Verzi *et al.*, 2016.

- (179) Orientation of distal portion of paroccipital process (Verzi *et al.*, 2016)*: on a plane parallel or subparallel to occipital plane (0); rotated so that its external margin becomes posterolateral or posterior (1); rotated so that its external margin becomes anterolateral (2).

Character stated 2 was included in order to score *Proechimys poliopus*.

- (180) Lateral crest (Verzi *et al.*, 2016)*: descending toward masseteric notch following same direction as the anterior margin of coronoid apophysis, or nearly so (0); with trajectory uncoupled from that of the anterior margin of the coronoid apophysis, more ventral along the mandibular body, forming a markedly descending curve that rises at its anterior extreme, which corresponds to the anterior end of the notch (1); more ventral along the mandibular body (2).

Original character 19 of Verzi *et al.* (2016), which includes features of the lateral crest and the nmmpio, was decoupled in two characters (here characters 181 and 182).

- (181) Mandibular notch for tendon of medial masseter muscle: subhorizontal (0); horizontal (1).

- (182) Anterior margin of base of coronoid apophysis: close to the alveolar edge of molars (0); more lateral and ventral with respect to alveolar edge of molars, even extending anteriorly as a more or less marked rim distinct from the lateral crest (1).
- (183) Lingual extreme of the protocone area (or posterior outgrowth of the protocone) in M1-2 of non-senile adults: oriented posteriorly (0); oriented more lingually (1).
- (184) Anteroloph on DP4: with no inflection on anterior or anterolingual surface (0); with weak inflection on anterior or anterolingual surface, from which point the orientation of this loph changes slightly to become more anterolabial (1).
- (185) Mesoflexus (or meso- + metaflexus) of M1-2: with bottom (lingual extreme) and labial extreme approximately equal in depth (0); markedly deeper at lingual extreme (1).
- (186) Metalophulid II of m1-2: originating from the protoconid area (0); originating from the metalophulid I (1); originating from the anterior part of the ectolophid (2).

Supplementary appendix 3. Character taxon matrix used for the phylogenetic analysis. The letters represent polymorphic scorings (*i.e.*, A = states 0 and 1; B = states 1 and 2; C = states 1 and 3; D = 0 and 2; E = 1 and 3).

	10	20	30	40	50	60	70
<i>Bugtimys zafarullahi</i>	000???00?0	0031111200	110010--10	-100111200	0030001100	1001122?20	1100002120
<i>Metaphiomys schaubi</i>	000???0100	000101????	???????????	??00121100	00111?0101	10-0100?11	01001-2???
<i>Phiomys andrewsi</i>	100???0000	0?????1100	0001110101	0000001110	00111?010?	?000?11?01	1100002101
<i>Garridomys curunuquem</i>	11200?1011	2?????0011	1002111111	2000111000	1-101110--	2001011???	????????121
<i>Eoviscaccia boliviana</i>	?120??1011	2?????0011	1???111111	2100111000	1-101?10-?	0101011???	??????????1
<i>Eoviscaccia frassinettii</i>	?120??1011	???????????	???????????	???????????	???????????	???????????	??????????1
<i>Dasyprocta azarae</i>	1120031011	00122?0311	120B111111	2000111200	00C2111112	011102212?	?000002??1
<i>Cephalomys arcidens</i>	112???101?	??????0511	00??10--10	-?00121000	1-001?1102	?101?20???	????????101
<i>Eosteiromys homogenidens</i>	1110000010	0?????0311	?001010111	2000100010	0010001111	00000001?0	20000A2111
<i>Steiromys detentus</i>	1110000010	0010?10311	0001111111	2000110010	0010001111	00000001?1	10000021?1
<i>Sallamys? minutus</i>	?1????0010	???????????	???????????	???????????	???????????	???????????	????????120
<i>Paulacoutomys paulista</i>	?11???0011	???????????	???????????	???????????	???????????	???????????	????????100
<i>Migraveramus beatus</i>	?11???0011	???????????	???????????	???????????	???????????	???????????	????????001
<i>Llitun notuca</i>	?11???0011	???????????	???????????	???????????	???????????	???????????	????????120
<i>Leucokephalos zeffiae</i>	?11???0011	???????????	???????????	???????????	???????????	???????????	????????100
<i>Changquin woodi</i>	?12???0?11	???????????	???????????	???????????	???????????	???????????	???????????
<i>Dudumus ruigomezi</i>	1110??0100	002001----	-----	--001000A0	0A20A11111	1000D11?00	2001002---
<i>Prospaniomys priscus</i>	1110000110	010001----	-----	--00101010	1-00001110	10A0000100	10A1002---
<i>Protacaremys? adilos</i>	?1????0???	???????????	???????????	???????????	???????????	???????????	???????????
<i>Protacaremys prior</i>	1110110110	010001----	-----	--01111010	1-00111111	1001021?01	1000002---
<i>Acarechimys minutus</i>	1100000110	010D01----	-----	--00111010	1-00111111	1000011?21	100A0A2---
<i>Acarechimys leucotheae</i>	?10???0100	???????????	???????????	???????????	???????????	???????????	?001001---
<i>Acarechimys minutissimus</i>	11000?0100	010001????	???????????	??00111000	1-0011111-	-00?-11?01	10AA00B---
<i>Acarechimys constans</i>	1100000110	010001????	???????????	??00101000	1-0011111-	-000-11?21	100A002---
<i>Acarechimys gracilis</i>	?10???0110	???????????	???????????	???????????	???????????	?????????01	1000012---
<i>Platypittamys brachyodon</i>	1100000000	0?????1201	001-00-010	-000100010	1-00001111	00000010??	????????001
<i>Galilelomys baios</i>	?1????0000	???????????	???????????	???????????	???????????	???????????	????????100
<i>Galileomys antelucanus</i>	1110??0000	0?????1401	00A1010111	1010110010	1-00111111	000?000???	????????100
<i>Acaremys murinus</i>	1110000010	00200114A1	00A1111111	1010111010	1-00111111	0000011001	?01-002100

<i>Caviocricetus lucasi</i>	1100??0100	002001----	-----	--00001010	0120111111	10000D2?01	A100012---
<i>Plesiacarechimys koenigswaldi</i>	11100?0110	002001----	-----	--00100010	0110101111	0000011?01	10000A2---
<i>Stichomys regularis</i>	1110011111	010001----	-----	--0110-001	1-00111112	010A0DD101	10001-2---
<i>Xylechimys obliquus</i>	?11???1?11	???????????	???????????	???????????	???????????	???????????	???????????
<i>Spaniomys riparius</i>	1110001111	010001---	-----	--0012-00A	1-00100102	1100000101	00001-2---
<i>Sallamys pascuali</i>	?11???0010	0?????1101	00A2010?10	-100101010	0AE00?1102	0000011??0	2100001100
<i>Adelphomys candidus</i>	1110011111	010001----	-----	-?0112-001	1-00001102	1100022?01	20???1-2---
<i>Ethelomys loomisi</i>	?11???0?11	???????????	???????????	???????????	???????????	???????????	???????????
<i>Deseadomys arambourgi</i>	?11???0011	0?????110?	001-110011	0?0010-000	1-00011112	1000002???	????????000
<i>Draconomys verai</i>	?100?0011	0?????????	???????????	??00121000	0011001112	100?111???	????????101
<i>Eosallamys simpsoni</i>	?10???00?0	0?????1101	000?110?10	-100121010	01E00011A2	0000011???	?00000????
<i>Eospina woodi</i>	?10???0000	0?????1111	000?10-???	-100101000	1-00000112	000?0???	1100001110
<i>Eosachacui lavocati</i>	?10???00?0	0?????1111	1???100111	1000101000	1-00000102	200?011?01	10??002100
<i>Eodelphomys almeidacomposi</i>	?1???1?11	???????????	???????????	???????????	???????????	???????????	???????????
<i>Cachiyacuy contamanensis</i>	010??00?0	0012210110	0202111100	-100101000	00E0A?A101	1000111?01	200000110?
<i>Canaanimys maquiensis</i>	?10??0?0?0	0?????????	???????????	??00101100	00320?A100	0000011?2?	???002???
<i>Eumysops laeviplicatus</i>	1120011111	001001---	-----	--0012-000	0AB0101112	200A0111?1	?0001-2---
<i>Myocastor coipus</i>	1120021111	010001----	-----	--0112-000	1-00001102	20001021?1	?000002---
<i>Echimys chrysurus</i>	1111111111	010001----	-----	--0012-000	1-0010A0-2	0100022001	0000?0---
<i>Proechimys poliopus</i>	1120001111	010?01----	-----	--0011-000	1-0?AA1102	00010000?1	?0001-2---
<i>Kannabateomys amblyox</i>	1111011111	010001----	-----	--0112-000	1-00100102	0111011101	00001-2---
<i>Neophanomys biplicatus</i>	?12???1111	??????----	-----	--?????????	???????????	???????????	10001-2---
<i>Chasicomys octodontiforme</i>	1120??1111	0?????1----	-----	--1011-0?0	??00??10?	100???????	????????--
<i>Chasichimys bonaerense</i>	?12???1111	??????----	-----	--1???-???	???????????	???????????	?0????2---
<i>Pithanotomys columnaris</i>	113-02?111	0?????----	-----	--101?-???	????11?10?	01010221??	?0????2---
<i>Plataeomys brevis</i>	113-02?111	0?????----	-----	--101?-???	????11?10?	01010221??	?0????2---
<i>Ctenomys australis</i>	113-01?111	0?????----	-----	--0?1?-?20	????11?1??	--?20201??	?0????2---
<i>Octomys mimax</i>	113-00?111	0?????----	-----	--101?-?00	????11?10?	11010220??	?0????2---
<i>Octodontomys gliroides</i>	113-03?111	0?????----	-----	--101?-?20	????11?1??	01010220??	?0????2---
<i>Abrocoma cinerea</i>	113-011?11	1?????????	???????????	??101?-?1?	---10-10-	02002111??	???????????

	80	90	100	110	120	130	140
<i>Bugtimys zafarullahi</i>	1110211111	0000001110	0011011010	?0???02?10	?0????????	??????????	??????????
<i>Metaphiomys schaubi</i>	??????????	?000001110	0011011000	10?2001?10	001???????	??????????	?????0????
<i>Phiomys andrewsi</i>	10-0211111	0000013110	0011011100	0000?01?11	001???????	??????????	??????????
<i>Garridomys curunuquem</i>	0A01000100	1011011011	000?111001	100??02011	000?????2?	??????????	?????01?0-
<i>Eoviscaccia boliviana</i>	?????0?100	202?????011	10011???1?	?2???0??1?	??????????	??????????	??????????
<i>Eoviscaccia frasinettii</i>	?????0?110	102??????1	1?001???0?	??????????	??????????	??????????	??????????
<i>Dasyprocta azarae</i>	1111101001	000100A001	10111A?000	1000100201	0000010201	011320-111	111010110-
<i>Cephalomys arcidens</i>	10-1001100	?11100?001	1011111110	1000?0?11	100???????	??????????	?????0?0-
<i>Eosteiromys homogenidens</i>	0111201101	0001001001	101111100?	10?0?0?10	001?????1?	10?110011	1100?0010-
<i>Steiromys detentus</i>	0111201101	0001001001	1011111000	1200001110	001?????210	0003110010	1101?0110-
<i>Sallamys? minutus</i>	10-10011??	0?????????	??????????	??????????	??????????	??????????	??????????
<i>Paulacoutomys paulista</i>	10-1201101	0001001011	00111?000	?????0??01	?0????????	??????????	??????????
<i>Migraveramus beatus</i>	10-1200111	0001003011	001110000?	100?10211?	?00???????	??????????	??????????
<i>Llitun notuca</i>	1101201111	0001001001	001110000?	100?102111	?0????????	??????????	??????????
<i>Leucokephalos zeffiae</i>	10-1200111	0101003011	00111?001	110?102111	?00?????1??	??????????	??????????
<i>Changquin woodi</i>	??????????	0001001001	001111100?	??0??????1	?01???????	??????????	??????????
<i>Dudumus ruigomezi</i>	-----	000A000010	0011111101	1102?00011	101?????2?	??????????	?????00?0-
<i>Prospaniomys priscus</i>	-----	0001001000	0A11111011	110?000011	A011000110	000110-110	011010000-
<i>Protacaremys? adilos</i>	??????????	0001000001	101111110?	??????????	??????????	??????????	??????????
<i>Protacaremys prior</i>	-----	0001001001	0010111001	01020A0?11	100?????1??	????11????	?????00?0-
<i>Acarechimys minutus</i>	-----	0001013011	00111A0001	01021A0021	10B?????2?	?00?11130-	-0-0?00?10
<i>Acarechimys leucotheae</i>	-----	0001013011	0011110101	??0??1---1	102???????	??????????	??????????
<i>Acarechimys minutissimus</i>	-----	000AA13011	A01111A101	0?0211---1	102?????2??	?????11???	?????00?10
<i>Acarechimys constans</i>	-----	00010A3011	00111AA101	110?11---0	101?????2??	?????11???	?????00?10
<i>Acarechimys gracilis</i>	-----	0001A13011	00111A0001	110??A0010	?01?????2??	??????????	??????????
<i>Platypittamys brachyodon</i>	10-0?01011	000A001010	00111A1A01	0102?0?01	00?????0?00	0?????20-	-0-??0????
<i>Galilelomys baios</i>	?????01111	0101000000	0011????0?	010?101111	?02???????	??????????	??????????
<i>Galileomys antelucanus</i>	10-0?01111	01010A0010	001110010?	010??01011	003???????	??????????	?????0?10
<i>Acaremys murinus</i>	A0-0?01112	01A1000011	00111A1101	010?101011	00311??20?	1111111211	0111?01111
<i>Caviocricetus lucasi</i>	-----	000001-010	0010010101	010111---1	101?????1??	??????????	?????00?0-
<i>Plesiacarechimys koenigswaldi</i>	-----	00010A1010	0011100001	0101?1---1	101???????	?0????????	?????????10
<i>Stichomys regularis</i>	-----	0001A1-000	0111111111	0022100010	10020??100	1??111111?	01??00?0-

<i>Xylechimys obliquus</i>	??????????	0001014000	0111101111	?11??????0	?00???????	??????????	??????????
<i>Spaniomys riparius</i>	-----	000000-000	0111111111	0122100A11	1002???11A	011311111?	01?1?11110
<i>Sallamys pascuali</i>	10-1200101	00011--011	0011100101	1001?0?11	?0????????	??????????	??????????
<i>Adelphomys candidus</i>	-----	0001A1-000	0111111111	120201--0	100???01??	??????????	?????????0-
<i>Ethelomys loomisi</i>	??????????	00011--00A	011111111?	112??01011	?01???????	??????????	??????????
<i>Deseadomys arambourgi</i>	10-1??1111	00010A0011	0011111101	111?101?10	10????????	??????????	??????????
<i>Draconomys verai</i>	10-1101111	0001003001	0011111001	10???02010	??????????	??????????	??????????
<i>Eosallamys simpsoni</i>	10-12001?1	0001001001	0011111001	??????????	??????????	??????????	??????????
<i>Eospina woodi</i>	1???001112	0001001001	001111100?	??0?02?11	?0????????	??????????	??????????
<i>Eosachacui lavocati</i>	1101101111	000100E011	00111AA001	??????????	??????????	??????????	??????????
<i>Eodelphomys almeidacomposi</i>	??????????	00011--000	111111111?	??????????	??????????	??????????	??????????
<i>Cachiyacuy contamanensis</i>	10-1100112	0001002101	00111A0000	??????????	??????????	??????????	??????????
<i>Canaanimys maquiensis</i>	??????????	0001012001	0011110001	??????????	??????????	??????????	??????????
<i>Eumysops laeviplicatus</i>	-----	00011--011	00111??101	110211--1	001??11120	010111110-	-0-100010-
<i>Myocastor coipus</i>	-----	000200-001	0111111110	1022?1--0	1001010111	0111110110	011101110-
<i>Echimys chrysurus</i>	-----	00011--010	0110111121	11220A0011	1010010110	11?111120-	-0-001010-
<i>Proechimys poliopus</i>	-----	0001A13011	1011111121	110211--1	1020010200	010111120-	-0-121110-
<i>Kannabateomys amblyox</i>	-----	00011--011	0111111121	000201--1	1010010201	111101010-	-10001110-
<i>Neophanomys biplicatus</i>	-----	0111A13011	001111?101	0102?1--1	?01???????	??????????	??????????
<i>Chasicomys octodontiforme</i>	-----	01111--01?	00111???01	120???????	??1???????	??????????	??????????
<i>Chasichimys bonaerense</i>	-----	01111--011	001111?10?	120??1--1	101???????	??????????	??????????
<i>Pithanotomys columnaris</i>	-----	0121?--?0?	?01?????11	120?01--0	001?110?10	110110-111	010110110-
<i>Plataeomys brevis</i>	-----	0121?--?0?	?01?????11	120211--0	011?1??111	?10110-???	???1?0110-
<i>Ctenomys australis</i>	-----	002???????	??1?????1	120211--1	1100000210	110110-110	010001110-
<i>Octomys mimax</i>	-----	0121???0?	?01?????11	110211--1	0110110121	101101120-	-101001111
<i>Octodontomys gliroides</i>	-----	0122???0?	?01?????11	100211--1	0111110210	101111120-	-101011111
<i>Abrocoma cinerea</i>	??????????	102?1--?11	011?111120	?22211--1	1110010010	010011?310	00-110110-

	150	160	170	180	186
<i>Bugtimys zafarullahi</i>	??????????	??????????	??????????	??????????	???1?0
<i>Metaphiomys schaubi</i>	??????????	??????????	??????????	??????????2	001112
<i>Phiomys andrewsi</i>	??????????	??????????	??????????	??????????	0?1112

<i>Garridomys curunuquem</i>	1?????????	???????????	???????????	???????????	0?10?-
<i>Eoviscaccia boliviana</i>	???????????	???????????	???????????	???????????	??10??
<i>Eoviscaccia frasinettii</i>	???????????	???????????	???????????	???????????	??10??
<i>Dasyprocta azarae</i>	11-0101110	0010122110	1011110002	-000000010	011?02
<i>Cephalomys arcidens</i>	1???????????	???????????	???????????	???????????	???????
<i>Eosteiromys homogenidens</i>	?????????02?	?1?????????	???????????	???????????	0?1?02
<i>Steiromys detentus</i>	1????????042	11?????1???	?00?????????	???????????	011102
<i>Sallamys? minutus</i>	???????????	???????????	???????????	???????????	???????
<i>Paulacoutomys paulista</i>	???????????	???????????	???????????	???????????	0?1??0
<i>Migraveramus beatus</i>	???????????	???????????	???????????	???????????	0????0
<i>Llitun notuca</i>	???????????	???????????	???????????	???????????	0????0
<i>Leucokephalos zeffiae</i>	???????????	???????????	???????????	???????????	01????0
<i>Changquin woodi</i>	???????????	???????????	???????????	???????????	0?????2
<i>Dudumus ruigomezi</i>	1???????????	???????????	???????????	???????????	0-1102
<i>Prospaniomys priscus</i>	0000001?00	1111?1111	21111120??	?????01202	101102
<i>Protacaremys? adilos</i>	???????????	???????????	???????????	???????????	?????0
<i>Protacaremys prior</i>	1???????????	???????????	???????????	???????????	101102
<i>Acarechimys minutus</i>	1000111?1	11?????????	???????????	???????????	111002
<i>Acarechimys leucotheae</i>	???????????	???????????	???????????	???????????	0?????D
<i>Acarechimys minutissimus</i>	1???????????	1???????????	???????????	???????????	111002
<i>Acarechimys constans</i>	1???????????	11?????????	???????????	???????????	011002
<i>Acarechimys gracilis</i>	???????????	???????????	???????????	???????????	0111?2
<i>Platypittamys brachyodon</i>	???????????	?1?????????	???????????	???????????	0?1??2
<i>Galilelomys baios</i>	???????????	???????????	???????????	???????????	01????2
<i>Galileomys antelucanus</i>	0???????????	???????????	???????????	???????????	011?12
<i>Acaremys murinus</i>	1???????????	00?????????	?????????0??	???????????	1111?2
<i>Caviocricetus lucasi</i>	1???????????	???????????	???????????	???????????	011102
<i>Plesiocarechimys koenigswaldi</i>	???????????	???????????	???????????	???????????	0?1000
<i>Stichomys regularis</i>	0???????????	11?????????	???????????	???????????	011002
<i>Xylechimys obliquus</i>	???????????	???????????	???????????	???????????	01????0
<i>Spaniomys riparius</i>	?????????1?2	11?????????	???????????	???????????	011000
<i>Sallamys pascuali</i>	???????????	???????????	???????????	???????????	0?1?02
<i>Adelphomys candidus</i>	???????????	11?????????	???????????	???????????	01?????
<i>Ethelomys loomisi</i>	???????????	???????????	???????????	???????????	?????-

<i>Deseadomys arambourgi</i>	??????????	??????????	??????????	??????????	0?1?02
<i>Draconomys verai</i>	??????????	??????????	??????????	??????????	0?1?02
<i>Eosallamys simpsoni</i>	??????????	??????????	??????????	??????????	??1?02
<i>Eospina woodi</i>	??????????	??????????	??????????	??????????	0?1?02
<i>Eosachacui lavocati</i>	??????????	??????????	??????????	??????????	??1?02
<i>Eodelphomys almeidacomposi</i>	??????????	??????????	??????????	??????????	??????
<i>Cachiyacuy contamanensis</i>	??????????	??????????	??????????	??????????	??1102
<i>Canaanimys maquiensis</i>	??????????	??????????	??????????	??????????	??A?02
<i>Eumysops laeviplicatus</i>	0000111020	10???10110	00000-0010	000??10102	01100-
<i>Myocastor coipus</i>	0000100022	10111?0110	00000-0000	1102100112	001000
<i>Echimys chrysurus</i>	0000111211	100-?10010	0000110000	10?0?00100	01100-
<i>Proechimys poliopus</i>	0010111300	1110120110	0000100011	0101110120	00101-
<i>Kannabateomys amblyox</i>	0020111211	100-??0010	10000-100?	?0???00??0	11100-
<i>Neophanomys biplicatus</i>	??????????	??????????	??????????	??????????	11???-
<i>Chasicomys octodontiforme</i>	??????????	??????????	??????????	??????????	01???-
<i>Chasichimys bonaerense</i>	??????????	??????????	??????????	??????????2	A0???-
<i>Pithanotomys columnaris</i>	1?????????	?0????????	???1??????	??????????0	11-0--
<i>Plataeomys brevis</i>	1?????????	?1????????	??????????	??????????0	11-0--
<i>Ctenomys australis</i>	1020111130	1111110111	21000-2000	100??00200	11-0--
<i>Octomys mimax</i>	1020111231	1111?20111	21110-210?	?1?0001?02	11-0--
<i>Octodontomys gliroides</i>	1000111242	1111?20111	2111112100	1001000200	11-0--
<i>Abrocoma cinerea</i>	1010011341	0111?01101	21110-2010	1000001200	011???