

New data on the endemic Patagonian long-clawed mouse *Notiomys edwardsii* (Rodentia: Cricetidae)

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Abstract

The monotypic *Notiomys* is the most poorly-known genus of Abrotrichini, a recently erected tribe of South American sigmodontine rodents. Based on a large sample, mostly recovered from owl pellets but including trapped individuals, we refine the known geographic distribution of this mouse (providing 20 new contemporary recording localities and four fossil occurrences), relate this to the principal Patagonian environmental units, add new data on the type locality of the species, and comment on its conservation status. In addition, we present a detailed external and craniodental description and provide data showing geographic variation of some skull features. Finally, we provide preliminary data on genetic variation (cytochrome *b* gene sequences) that demonstrate the existence of a low degree of observed differentiation (ca. 0.7%) between the populations analyzed.

Keywords: Abrotrichini; Argentina; *Chelemys*; *Geoxus*; *Pearsonomys*.

Introduction

Abrotrichines form a small group of genera of South American cricetid rodents. For many decades, abrotrichines were considered to be part of Akodontini; however, the use of molecular evidence and a reappraisal of morphologic variation promoted their separation and consolidation as a new tribe of Sigmodontinae (D'Elía et al. 2007). Within Abrotrichini two main clades are recognized, one constituted solely by the polytypic *Abrothrix* and the other including four long-clawed genera, *Chelemys*, *Geoxus*, *Notiomys*, and *Pearsonomys* (D'Elía et al. 2006).

Knowledge about abrotrichines is markedly uneven and is strongly biased towards *Abrothrix* (Liascovich et al. 1989, Spotorno 1992, Lozada et al. 1996, Smith et al. 2001, Pearson 1995). With the partial exception of *Chelemys*, the other genera of long-clawed mice remain poorly known. Current knowledge of the monotypic *Notiomys*, an endemic species of Argentinean Patagonia, is limited to scattered data on morphology and natural history and the compilation of recording localities (Pearson 1984, Reig 1987, Patterson 1992, Pardiñas and Galliari 1998, Teta et al. 2002, Martin and Archangelsky 2004).

Toward partially filling the gap in our knowledge of the long-clawed mouse *Notiomys edwardsii* Thomas 1890, the aim of the present study was to provide a description of its molar occlusal pattern, skull, stomach, and external morphology, and a preliminary assessment of its genetic variation. Based on our compilation of pertinent information, we restrict the location of the type locality of *N. edwardsii*; in addition, we expand its known geographic distribution through the addition of 20 contemporary and four fossil new recording localities. Finally, we discuss the association between this mouse and the main environmental units recognized for Patagonia, and comment on its conservation status.

Materials and methods

Most of the specimens studied were recovered from owl pellets. A fresh individual was found dead in the field and was prepared as skin and skeleton and tissues preserved in ethanol. We also studied four specimens housed in collections (Table 1). It is important to emphasize that in spite of intensive fieldwork using snap and Sherman traps in different places and different seasons where *Notiomys edwardsii*, as indicated by owl pellet analyses, is abundant, no specimen was trapped. Acronyms for institutions and collections mentioned in the text and tables are: BM, Natural History Museum of London, London, UK; CNP, Colección de Mamíferos del Centro Nacional Patagónico, Puerto Madryn, Chubut, Argentina; CNP-E, Colección de Materiales de Egagrópilas y afines "Elio Massoia" del Centro Nacional Patagónico, Puerto Madryn, Chubut, Argentina; LER.E, Laboratorio de Ecología de Roedores, Departamento de Ecología, Genética y Evolución, Facultad de Ciencias Exactas y Naturales, Universidad de Buenos Aires, Argentina; LIEB M, Laboratorio de Investigaciones en Evolución y Biodiversidad, Facultad de Ciencias Naturales Sede Esquel, Universidad Nacional de la Patagonia, Esquel, Chubut, Argentina; MACN, Museo de Ciencias Naturales "Bernardino Rivadavia", Buenos Aires, Argentina; and MVZ, Museum of Vertebrate Zoology, Berkeley, California, USA.

Table 1 Contemporary and fossil recording localities for *Notiomys edwardsii* (arranged by increasing latitude).

| # | Locality | Latitude S | Longitude W | Altitude (m) | Province | Collection and/or voucher | Main reference | Observations |
|---------------------|----------------------------------|------------|-------------|--------------|------------|---------------------------|--------------------------------|--|
| Contemporary | | | | | | | | |
| 1 | Campo Anexo Pilcaniyeu | 41.020000° | 70.470000° | | Río Negro | MVZ 163065 ^s | Pearson (1984) | Trapped individual |
| 2 | 10 km W-SW Comallo | 41.050000° | 70.410000° | | Río Negro | MVZ 163067 | Pearson (1984) | Trapped individual |
| 3 | Cerro Puntudo | 41.304722° | 66.906111° | | Río Negro | Unknown | Andrade (2008) | Owl pellet material |
| 4 | Laguna La Fariña | 41.323333° | 66.962778° | | Río Negro | Unknown | Andrade (2008) | Owl pellet material |
| 5 | Laguna del Paraguay | 41.351111° | 66.962222° | 1307 | Chubut | CNP-E 73 | Pardiñas and Teta (2007) | Owl pellet material |
| 6 | Cerro Corona Chico | 41.369167° | 66.921111° | | Río Negro | Unknown | Andrade (2008) | Owl pellet material |
| 7 | Laguna Buñuelo | 41.382222° | 66.951111° | | Río Negro | Unknown | Andrade (2008) | Owl pellet material |
| 8 | Laguna Blanca | 41.426389° | 66.957500° | 1300 | Río Negro | CNP 1* | This paper | Found dead individual |
| 9 | Cerro Corona | 41.455833° | 66.914444° | 1567 | Río Negro | CNP-E 31 | Pardiñas and Teta (2007) | Owl pellet material |
| 10 | 3 km N Cuatro Cerros on RP 60° | 41.466667° | 66.983333° | 1340 | Río Negro | CNP-E 80 | Pardiñas and Teta (2007) | Owl pellet material |
| 11 | Cerro Somuncurá Chico | 41.535278° | 66.883889° | | Río Negro | Unknown | Andrade (2008) | Owl pellet material |
| 12 | Laguna Maciega | 41.586944° | 67.097500° | | Río Negro | Unknown | Andrade (2008) | Owl pellet material |
| 13 | Cerro Mimbre | 41.597778° | 66.880278° | | Río Negro | Unknown | Andrade (2008) | Owl pellet material |
| 14 | Subida del Naciente ^s | 41.673333° | 67.154167° | 1042 | Chubut | CNP-E 27 | Canevari et al. (1992) | Owl pellet material |
| 15 | Arroyo Pinturas | 41.701111° | 66.703611° | | Río Negro | Unknown | Andrade (2008) | Owl pellet material |
| 16 | Estancia Calcatreo | 41.733333° | 69.366667° | | Río Negro | Unknown | Andrade et al. (2002) | Owl pellet material |
| 17 | Cerro Castillo | 41.966667° | 69.266667° | | Chubut | CNP-E 191 | Pardiñas and Galliari (1998) | Owl pellet material |
| 18 | Near Salina Grande | 42.053889° | 70.105833° | | Chubut | CNP-E 255 | This paper | Owl pellet material |
| 19 | Estancia Talagapa | 42.137778° | 68.254722° | 1414 | Chubut | CNP-E 204 | This paper | Owl pellet material |
| 20 | Sierras de Talagapa | 42.214178° | 68.222114° | | Chubut | Unknown | Teta and Andrade (2002) | Owl pellet material |
| 21 | 2 km NW Gastre | 42.233333° | 69.200000° | | Chubut | CNP-E 57 | Jayat et al. (2006) | Owl pellet material |
| 22 | Sierras de Talagapa | 42.233822° | 68.240269° | | Chubut | CNP-E 192 | Teta and Andrade (2002) | Owl pellet material |
| 23 | Fofo Cahuel | 42.400000° | 70.516667° | 531 | Chubut | CNP-E 116 | This paper | Owl pellet material |
| 24 | Near Piedra Parada | 42.637222° | 70.223611° | 570 | Chubut | CNP-E 125 | This paper | Owl pellet material |
| 25 | Tres Cuevas | 42.647222° | 70.152778° | 435 | Chubut | CNP-E 33 | This paper | Owl pellet material |
| 26 | Cañadón de la Buitrera | 42.649444° | 70.103333° | 438 | Chubut | CNP-E 8 | This paper | Owl pellet material |
| 27 | 8 km W Paso del Sapo | 42.680556° | 69.674167° | 338 | Chubut | CNP-E 224 | This paper | Owl pellet material |
| 28 | Paso del Sapo | 42.683333° | 69.716667° | | Chubut | CNP-E 62 | Pardiñas and Galliari (1998) | Owl pellet material |
| 29 | 3 km S RP 12 to Colan Conhue | 42.695556° | 70.121667° | 591 | Chubut | CNP-E 234 | This paper | Owl pellet material |
| 30 | Campo de Cretón | 42.733333° | 70.050000° | 809 | Chubut | CNP-E 122 | This paper | Owl pellet material |
| 31 | 4 km S Tres Banderas on RP 11 | 42.808611° | 68.015556° | 956 | Chubut | CNP-E 36 | This paper | Owl pellet material |
| 32 | Colonia Nahuel Pan | 42.950000° | 71.166667° | | Chubut | LIEB M 681 | Martin and Archangelsky (2004) | Trapped individual |
| 33 | Cabaña Arroyo Pescado | 43.025278° | 70.792778° | 568 | Chubut | CNP-E 237 | This paper | Owl pellet material |
| 34 | Campo de Pichiñán | 43.563889° | 69.067222° | 282 | Chubut | CNP-E 210 | This paper | Owl pellet material |
| 35 | 30 km W José de San Martín | 44.033333° | 70.850000° | | Chubut | MVZ 182132 | Pardiñas and Galliari (1998) | Trapped individual |
| 36 | Establecimiento La Ollada | 44.746750° | 69.618806° | 946 | Chubut | CNP-E 291 | This paper | Owl pellet material |
| 37 | Puesto El Chango | 45.464306° | 69.431667° | 240 | Chubut | CNP-E 290 | This paper | Owl pellet material |
| 38 | Estancia Valle Huemules | 45.948000° | 71.507000° | | Chubut | BM | Thomas (1919) | Valle del Lago Blanco restricted by Pardiñas et al. (2003) |
| 39 | Piedra Clavada | 46.583333° | 68.566667° | | Santa Cruz | BM | Thomas (1929) | Trapped individual |
| 40 | Cueva de las Manos | 47.183333° | 70.583333° | | Santa Cruz | CNP-E 47 | This paper | Owl pellet material |
| 41 | Laguna del Diez | 47.883333° | 67.850000° | | Santa Cruz | CNP-E 21 | Jayat et al. (2006) | Owl pellet material |

(Table 1 continued)

| # | Locality | Latitude S | Longitude W | Altitude (m) | Province | Collection and/or voucher | Main reference | Observations |
|----|------------------------------|------------|-------------|--------------|------------|---------------------------|---------------------------------|------------------------------|
| 42 | Estancia Laguna Manantiales | 47.533333° | 68.300000° | | Santa Cruz | CIPD 589* | This paper | Trapped individual |
| 43 | Near Cerro El Puma | 48.383333° | 69.966667° | | Santa Cruz | Photograph | This paper | Photograph by Gabriel Rojo |
| 44 | N end Lago Cardiel and RN 40 | 48.900000° | 71.016667° | | Santa Cruz | CNP-E 96 | This paper | Owl pellet material |
| 45 | Parador Luz Divina, RN 40 | 49.883333° | 72.033333° | | Santa Cruz | CNP-E 93 | This paper | Owl pellet material |
| 46 | Estancia La Ascención | 49.886389° | 72.041389° | | Santa Cruz | LER-E 002 | Cueto et al. (2007) | Owl pellet material |
| 47 | Puerto Santa Cruz | 50.016667° | 68.516667° | | Santa Cruz | BM 18.12.21.1 | Thomas (in Milne-Edwards, 1890) | Trapped individual; holotype |
| | Fossil | | | | | | | |
| 48 | Epulán Grande Cave | 40.391111° | 70.196667° | | Neuquén | LL E3-F3* | This paper | Archaeological remains |
| 49 | Talagapa Cave | 42.233333° | 68.233333° | | Chubut | CNP [†] | This paper | Paleontological remains |
| 50 | Watkins cave | 42.750278° | 70.873611° | | Chubut | LIEB | This paper | Paleontological remains |
| 51 | Los Altares profile | 43.893056° | 68.389167° | | Chubut | CNP-E 259–270 | This paper | Paleontological remains |

*Two additional specimens came from this locality, MVZ 163066 and MACN 19334.

[†]Erroneously referred to as Los Cuatro Cerros by Andrade (2008).

[‡]Erroneously referred to as Cañadón del Naciente by Andrade (2008).

[§]Housed at Centro de Investigaciones de Puerto Deseado (CIPD), Universidad Nacional de la Patagonia Austral (Santa Cruz, Argentina).

[¶]A still unnumbered mandible temporarily housed at Centro de Estudios Farmacobiológicos y Botánicos (CEFYBO; Buenos Aires, Argentina).

[‡]Several craniodental remains housed at Archaeological and Anthropological Unit (Centro Nacional Patagónico, Puerto Madryn, Argentina).

*Specimens sequenced in this study.

Morphological descriptions follow the concepts of Carleton and Musser (1989) and Patterson (1992) for cranial anatomy and Reig (1977) for molar occlusal structures. Standard external measurements were taken from specimen labels and the literature. Craniodental measurements were made using digital calipers and follow the definitions provided by Tribe (1996).

The analysis of genetic variation is based on 801-bp cytochrome *b* (cyt *b*) sequences gathered from three specimens. One sequence was obtained from GenBank (U03537) and we obtained the other two (Table 1) following the protocol detailed in D'Elia and Pardiñas (2004) and deposited them with GenBank (EU416275 and EU416276). Sequences were aligned by eye with sequences from specimens of *Chelemys*, *Geoxus*, and *Pearsonomys* (U03533, U03531, AF108672) that were used as an outgroup. Reported sequence divergence values correspond to observed (*p*) values calculated with PAUP* (Swofford 2000) ignoring those sites with missing data. The genealogy discussed was obtained by analysis of maximum parsimony (Farris 1982) instrumented as an exhaustive search in PAUP*. Clade support was assessed by 1000 bootstrap replicates with five replicates of random sequence addition each.

Results and discussion

Notiomys Thomas 1890

Type species *Hesperomys (Notiomys) edwardsii* Thomas, 1890

Remarks There is some confusion regarding the year of publication of this genus and species. Most authorities used 1890 (Osgood 1925, Gyldenstolpe 1932, Cabrera 1961, Musser and Carleton 2005). However, 1891 is the year printed on the cover of the “Mission Scientifique du Cap Horn 1882–1883. Tome VI. Zoologie. Première partie,” the publication in which the names were proposed. Similarly, this publication is listed as 1891 in the “Catalogue of the books, manuscripts, maps and drawings in the British Museum (Natural History)” (1904 edition, L. Di Tommaso, personal communication). To add more uncertainty, Thomas (1929: 42) wrote in reference to *Notiomys edwardsii* “...species described by me in 1892.” We follow here the traditional date used, but remark that this issue needs further research to be resolved.

Holotype BM 18.12.21.1, an adult individual, unknown sex, collected by Le Brun in December 1882.

Type locality “la Patagonie, au sud de Santa Cruz, vers le 50° degré de latitude Sud” (Thomas in Milne-Edwards 1890: A.25). It is interesting to reconstruct how this rodent captured in Santa Cruz province (continental Argentina) was finally reported as a new species in an account covering the region of Cape Horn. Several scientific expeditions were sent from France to observe the transit of Venus in 1882. Two of them focused on southern localities of South America, Santa Cruz and Cape Horn. The expedition to Santa Cruz was led by the naval officer Georges-Ernest Fleuriais (1840–1895) in the ship

Volage (Mouchez 1882). Le Brun (spelling according to Fleuriais 1882) or Lebrun (according to Milne-Edwards 1890) was the naturalist assigned to this expedition. Le Brun collected some specimens, including the holotype of *Notiomys edwardsii*, in an area around where the Volage crew carried out astronomical observations. The exact location of these places is not known, but likely they were not far from "Santa Cruz", at that time the only village in this large area, no more than a few shanties by Cañadón Misionero, a small creek with its mouth at 49°59' S and 68°33' W. In 1884 Carlos Moyano was designated as the first governor of Santa Cruz territory; he selected "Santa Cruz" as the capital, but moved its placement approximately 4 km south, to another temporary creek (Cañadón del Quemado), approximately 4.5 miles downwater of the conjunction between the rivers

Santa Cruz and Chico (Moyano 1948). During the last century, the use of "Santa Cruz" declined and was replaced by Puerto Santa Cruz, with geographic coordinates at 50°01' S and 68°31' W. Therefore, the "Santa Cruz" visited by Le Brun in 1882 and Puerto Santa Cruz are not the same place, although they are close to each other. Moreover, ambiguity regarding the exact place where Le Brun collected the holotype of *N. edwardsii* must be borne in mind. In this context and for taxonomic and nomenclatural purposes, the type locality of *N. edwardsii* must be restricted to Puerto Santa Cruz (province of Santa Cruz, Argentina), a proposition mentioned briefly by Pardiñas and Galliari (1998), and incorrectly transcribed by Musser and Carleton (2005: 1136). We note that, other than the holotype, no additional specimen is known from the type locality; the closest record-

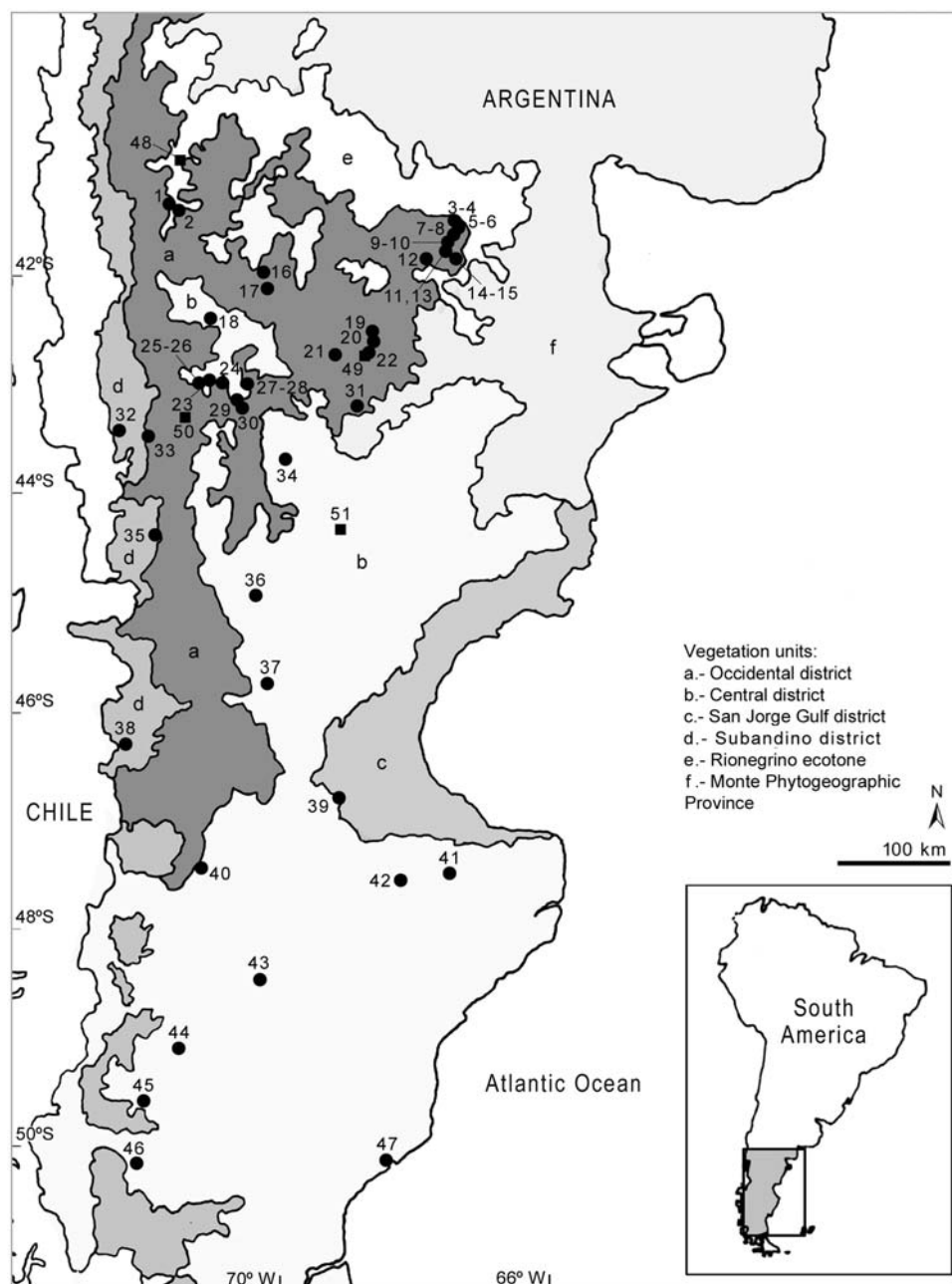


Figure 1 Recording localities for *Notiomys edwardsii* superimposed on the principal phytogeographic units that occur in Argentinean Patagonia. See Table 1 for reference numbers.

ing locality to Puerto Santa Cruz is near Cerro El Puma, approximately 180 km to the northwest.

When the Fleuriais mission ended, Le Brun traveled by land to Punta Arenas (Chile) to meet the second French expedition to South America, commanded by Captain Louis-Ferdinand Martial, destined for Cape Horn (Martial 1888). All the zoological material collected by Le Brun in both French missions was finally studied and published by Milne-Edwards (1890), who invited Oldfield Thomas to describe the cricetids.

Distribution An exhaustive search of the literature and collections retrieved 47 contemporary recording localities for *Notiomys edwardsii* (Table 1; Figure 1), ranging from ca. 41°S to 50°S and encompassing the Argentinean provinces of Santa Cruz, Chubut, and Río Negro. One record based on material recovered from *Galictis cuja* scats from Quilquihue (Neuquén province, Argentina; Delibes et al. 2003) is dubious and can be discarded (A. Travaini personal communication).

Emended morphological diagnosis A member of the tribe Abrotrichini characterized by the following combination of traits: small size (head and body length ca. 80 mm), short tail (ca. 40 mm); head with bright orange spots on the rump sides; very small ears showing a tuft of contrasting white hairs; manus and pes dorsally covered by whitish-orange to white hairs; manus with well-developed sharp claws; pes with conspicuous inner and outer fringes of hairs; skull with shorted rostrum, broad amphoral interorbital region, rounded braincase and expanded zygomatic arches; nasals covering most of the rostrum and surpassing the anterior face of incisors, somewhat trumpeted; inconspicuous zygomatic notches; narrow and tall zygomatic plates without an upper free border, strongly flared with respect to sagittal plane and producing a wide infraorbital foramen associated with a well-depressed and diagonally oriented nasolacrimal region; interparietal bone filiform; well-inflated exoccipitals and mastoids; palate wide and long; incisive foramina typically not reaching the anterior plane of M1; paired posterolateral palatal pits adjacent to the anterior border of the mesopterygoid fossa; mesopterygoid roof excised by large sphenopalatine vacuities; inflated otic capsules with short Eustachian tubes; upper incisors orthodont; molars brachyodont with opposing main cusps; M1 trilophodont with fan-shaped procingulum anteriorly indented by a shallow anteromedian flexus; M2 bilophodont with area of the proto- and paracone clearly broader than that of the hypo- and metacone; M3 reduced and cylindrical in outline; M3/m3 with a single root; mandible gracile with lower border moderately straight, shallow diastema, and mental foramen large and clearly labially positioned; stomach unilocular and hemiglandular.

Description Small sigmodontine with a fusiform body (Figure 2). Ears very reduced and extremely thin, internally and externally covered by long hairs, with a noticeable pure white tuft, which originates inside the pinna notch and surpasses its outer margin. Eyes of medium size (maximum diameter 2.7 mm in CNP 1). Numerous

mystacial vibrissae (15 per side), some totally dark, some totally white, and some basally black and distally whitish. Largest vibrissae (ca. 30 mm) posteriorly surpass the auricular pinna margin. Nose covered by dark hairs conforming a “leather button” (Pearson 1984: 234). Dorsal pelage dense, soft, but not long (mid-dorsum hairs ca. 7–8 mm in length); ventral pelage soft. Lanceolate scale pattern on guard hairs; medullar cells arranged in two parallel rows (see also Chehébar and Martín 1989). Dorsal head and body coloration with agouti effect, produced by almost entirely gray hairs with ochraceous tip; belly coloration gray cream to white, produced by hairs that are basally gray and distally white; lateral line of contact between back and belly marked by hairs with distal half orange. Bright orange spots on rostrum sides. Tail bicolor, moderately haired, proximal hairs ca. 10 mm, short distal tuft; three hairs per scale, each covering three scales in length (dorsal middle region). Manus well covered dorsally with whitish orange hairs; carpal vibrissae shorter than claws, the latter ca. 4 mm in length, powerful, transversally compressed and ventrally ridged except for the distal third tip. Six palmar pads with naked surface; ungual pads rounded. Hind foot short and broad, with an inner and outer fringe of stiff hairs well extended beyond the plantar surface (Figure 3); bicolored, from ankle to middle portion of metatarsal region washed orange and whitish from this point to the end of the fingers; posterior claws short (ca. 1.6 mm) ventrally opened. Six naked plantar pads with smooth surfaces, hypothenar large, first and fourth larger than the second and third.

Skull moderately robust with shortened rostrum, broad interorbital region, well-rounded braincase, expanded zygomatic arches, and flat dorsal profile (Figure 4). The nasals are long, covering most of the rostrum, anteriorly divergent with maximum width just posterior to the anterior end (Figure 5A); in old adult individuals the anterior third of the nasals are noticeably upturned. Nasals and premaxillae surpass the anterior incisive face, appearing “trumpet-like” in lateral view (Figure 5B). In addition, a moderately developed gnathic process is present. The nasofrontal suture is V-shaped and located posteriorly to the plane defined by lacrimal bones (Figure 5A); the premaxilla-frontal suture is well denticulated. Zygomatic notches are inconspicuous, in line with the near absence of an upper free border in the zygomatic plates (Figure 5B). The lacrimal bones are small. Interorbital region with amphoral or “hour-glass shaped” form and frontal sinuses well inflated, frontal external borders squared but not ridged, and coronal suture nearly straight with sharp contacts with parietal bones. Interparietal bone filiform (Figure 5C). In the dorsal view of the braincase, large portions of squamosal and occipital bones are evident. Parietal or lambdoidal crests inconspicuous. Occipital region with well-inflated exoccipitals. Rostrum in lateral view with premaxillo-maxillary suture partially arched and well-developed nasolacrimal foramen. Infraorbital foramen broad and connected with a depressed diagonally oriented nasolacrimal region; a medium-size circular foramen is present on the rostral process of the premaxilla closely or directly opening on the premaxillo-maxillary suture (Figure 5B); this foramen, topographically associated with nasolacrimal structures, is also present in other



Figure 2 External aspect of *Notiomys edwardsii*. Above, an individual photographed alive in the area of Cerro El Puma (Santa Cruz, Argentina); note the white tuft of hair in the ear, a trait unique among Patagonian sigmodontines. Below, the original drawing included in Milne-Edwards (1890: Plate 3, Figure 1).



Figure 3 Ventral surface of the right foot of *Notiomys edwardsii* (CNP 1) showing conspicuous inner and outer fringes of hairs (arrows); also note the ventral opening of the claws.

Abrotrichini (e.g., *Abrothrix*, *Chelemys*, *Geoxus*) and some Akodontini (e.g., *Akodon spegazzinii*; see Blaustein et al. 1992, where it is referred to as the “preorbital foramen II”), but in *Notiomys* it is well developed and consistent (visible with the naked eye). Zygomatic plate tall and narrow, with parallel sides and no upper border. Zygomatic plate inclined nearly 45° with respect to sagittal plane (Figure 5D). Inconspicuous area of insertion of masseter superficialis in the anterior portion of zygomatic plate base. Zygomatic arches delicate, with jugal well developed. Temporal region with strong participation of occipital bone; lateral projection of parietal squared in outline and well developed. Mastoid well inflated, producing the emergence of a squared lower edge; mastoid process well developed and next to large paraoccipital process. Hamular process weak and curved distally on the mastoid promontory of the otic capsule (Figure 5E). Tegmen tympani well developed, attached to squamosal ridge. Postglenoid foramen well developed but subsquamosal fenestra reduced. Auditive meatus large; malleus with an enlarged orbicular process. Incisive foramen



Figure 4 *Notiomys edwardsii* (MVZ 163067): top to bottom, skull in dorsal, ventral, and lateral views, and left mandible in labial view. Scale bars represent 5 mm.

short and broad, with posterior end typically not reaching the plane defined by the anterior face of M1 (a moderate variation of this trait appears to be ontogenetically related; Figure 5F). Palate broad and long (*sensu* Hershkovitz 1962), flat, with minute to very large paired posterolateral palatal pits located near the anterior border of the mesopterygoid fossa. The latter fossa is narrow and parallel-sided with a medium palatine process moderately developed on its anterior border. Roof of mesopterygoid region with large sphenopalatine vacuities. Parapterygoid fossae broader than mesopterygoid fossa, flat, with evident fontanelles and posteriorly well ossified to contact otic capsules (Figure 5G). Presphenoid short and broad. Ectotympanic bullae large and well inflated with short Eustachian tubes; pteriotic totally covered in dorsal view (Figure 5H). Carotid canal only visible in internal lateral view due to ectotympanic development; stapedial foramen well developed, as is the petrotympanic fissure. Large foramen magnum posteriorly located. Alisphenoid strut absent, foramen ovale confluent with alisphenoid canal; a tunnel for secondary arterial connection between internal carotid and orbital-maxillary circulation is present in some individuals (e.g., MVZ 163067); trough for masticatory-buccinator nerve broad but very shallow; squamosal-alisphenoid groove clearly visible with a medial perforation, leads anteriorly to a medium sphenofrontal foramen (Figure 5I). Mandible gracile, with the lower border moderately straight (Figure 4). Lower diastema shallow with the anterior-most point surpassing upwards the alveolar plane. Mental foramen large and located on the base of the diastema, scarcely visible from dorsal view. Masseteric crest moderately developed; its inferior ridge

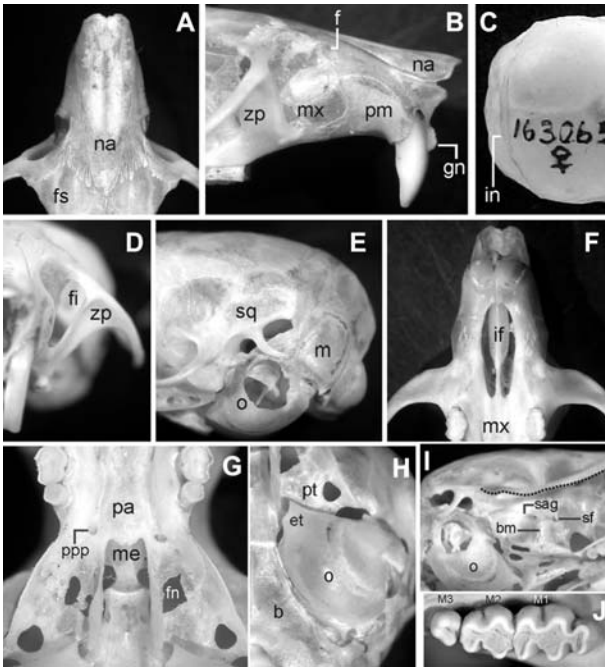


Figure 5 Craniodental morphological details of *Notiomys edwardsii* (MVZ 163067, except as noted). (A) Rostrum in dorsal view showing inconspicuous zygomatic notches and well-inflated frontal sinuses [fs]. (B) Rostrum in lateral view showing the absence of an upper free border in the zygomatic plate, the moderately arched premaxillo [pm]-maxillary [mx] suture, nasal-premaxillary forward projection, circular foramen on the rostral process of the premaxilla [f], and gnathic process [gn]. (C) Partial braincase in dorsal view showing reduced interparietal [in] bone (MVZ 163065). (D) Skull in frontal view showing the wide infraorbital foramen [fi] and the inclination of the zygomatic plate [zp]. (E) Temporal region in lateral view showing the well-inflated mastoid [m]. (F) Rostrum in ventral view showing anterior extension of the nasal bones and the short incisive foramina [if]. (G) Posterior palatal region showing mesopterygoid fossa [me] morphology and large sphenopalatine vacuities. (H) Otic capsule [o] with very short Eustachian tube [et] and largely ossified parapterygoid plate [pt]. (I) Carotid circulatory pattern illustrated by osteological features. (J) Occlusal view of the left upper tooth-row in a young individual (MVZ 163065); note the asymmetry between the anterior and posterior lobes of M2. b, basioccipital; bm, trough for masticatory-buccinator nerve; fn, fontanel; na, nasal; pa, palatine; ppp, posterolateral palatal pits; sag, squamosal-alisphenoid groove; sf, sphenofrontal foramen; sq, squamosal.

meets with the upper ridge, forming a unique crest to the level of the mental foramen. Coronoid process robust and well curved posteriorly, higher than condyloid process, producing a well-excavated sigmoid notch. Angular process broad and shorter with respect to the condyle, producing a lunar notch moderately excised but oblique in profile. Incisor capsule projection medium in size, lying below the coronoid process. Conspicuous retromolar fossa with a medium foramen. Dental foramen large and located over the milohyoid line between coronoid and condyloid processes.

Upper incisors robust and orthodont (Figure 5B); dentine fissure straight (sensu Steppan 1995). Molars brachyodont with some tendency to mesodonty (Figure 5J). Coronal topography bilevel, terraced on the lingual side

and crested (in juvenile specimens) to planate (in adults) on the labial side; main cusps arranged in opposite pattern. M1 trilophodont; procingulum fan-shaped and slightly compressed transversally with shallow but consistently present anteromedian flexus; anteroflexus-parastile absent; proto- and paraflexus subequal and transversal in orientation; their internal ends meet, producing an anterior mure medially located; para- and protocone subequal in size; paracone coalesces with reduced mesoloph; short but broad hypoflexus; metacone anterior-posteriorly compressed, labially projected; short but present posteroloph directed posteriorly. M2 bilophodont with short anteroloph directed forward, paraflexus almost obsolete, as are the hypo- and mesoflexus; area of the proto- and paracone clearly broader than that of the hypo- and metacone; short posteroloph directed posteriorly. M3 reduced and cylindrical in outline, occlusal details typically missing due to wear; mesofosetus present in young individuals (e.g., CNP 1). Lower molars more crested than upper ones. In m1 the procingulum is sub-squared in outline with a marked anteromedian flexus, producing two subequal conules; anterior murid medially located; proto- and hypoconid anterior-posteriorly compressed; broad hypoflexid, bulbous metaconid, lingually projecting entoconid coalescent with a reduced mesolophid; posterolophid well developed and directed anteriorly. m2 similar in shape to m1, but wider, without evidence of anterolophid. m3 reduced and bilophodont, with clear transverse compression of the posterior lobe.

Molar roots (partially inferred from alveolar pattern): M1 with two large and circular roots, one anterior and one posterior, the former occasionally divided in two, and conspicuous inter-root space densely foraminated, where a small accessory labial root may be present (under paracone); M2 with three roots, one lingual elliptical in outline and two labials that are subequal and circular; M3 with a single circular root; m1 with two large roots (anterior and posterior) located along the midline of the tooth, and one reduced root in the labial side; m2 with two roots (anterior and posterior); m3 with a single circular root.

Vertebral count (on CNP 1) indicates 13 thoracic, six lumbar, and 18 caudal vertebrae, and 13 ribs (three free); the same numbers were reported by Steppan (1995) based on specimen MVZ 163065.

Stomach morphology unilocular and hemiglandular (Figure 6), typical of many Sigmodontinae (Carleton 1973). The antrum, which is covered by glandular epithelium, is larger than the corpus, which in turns is mostly covered by cornified epithelium. The glandular epithelium from the antrum extends to the corpus and surpasses to the left of the orifice of the esophagus. Carleton (1973) described two gross morphologies among sigmodontines with a unilocular-hemiglandular stomach. The type reported here for *Notiomys*, in which the glandular epithelium covers a small area of the corpus, is also present in *Geoxus* and *Pearsonomys* (Carleton 1973; D'Elía et al. 2006), which form a clade sister to *Notiomys* in phylogenetic analyses (Smith and Patton 1999; D'Elía et al. 2006). *Abrothrix* has a unilocular-hemiglandular type, in which the glandular epithelium covers a slightly smaller

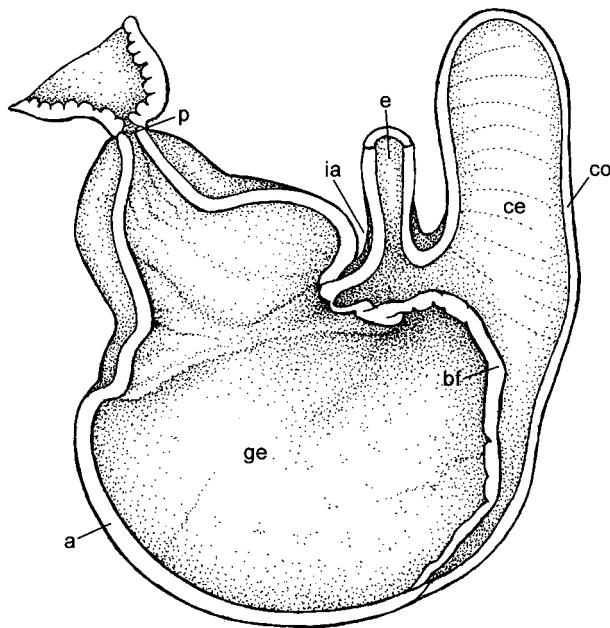


Figure 6 Stomach morphology in *Notiomys edwardsii*: cross-section based on CNP 1. a, antrum; bf, bordering fold; c, corpus; ce, cornified squamous epithelium; e, esophagus; ge, glandular epithelium; ia, incisura angularis; p, pylorus.

area of the stomach (Carleton 1973). The stomach morphology of *Chelemys* remains unknown.

Other traits of the soft anatomy (such as penis morphology and gall bladder presence/absence) of *N. edwardsii* remain unknown. The same is true for karyological data, with this being the only genus of Abrotrichini for which this information is lacking.

Measurements External measurements (in mm) for the few available complete adult specimens (BM Piedra Clavada, LIEB M-681, MVZ 163065–163067, see Thomas 1929, Pearson 1984, Martin and Archangelsky 2004) provide the following mean and standard deviation values: total length, 129.9 ± 6.95 ($n=5$); tail length, 41.9 ± 2.83 ($n=5$); hind foot (with claw), 19.1 ± 1.24 ($n=5$); ear length, 7.4 ± 0.82 ($n=5$); and weight (in g), 21.3 ± 3.33 ($n=3$). Values for a young specimen (CNP 1) are: 115, 35, 19, 7 and 17, respectively. Craniodental measurements are provided in Table 2.

Morphological variation A consistent geographically structured pattern of variation was observed for the size of paired posterolateral palatal pits (ppp; Figure 7). Samples (localities #41, 44–46; Table 1) south of Río Deseado (ca. $46^{\circ}45'$ S) have large to very large ppp, in some cases with internal subdivisions, whereas samples north of Río Deseado (localities #1, 2, 5, 8–10, 14, 17–19, 23–27; Table 1) show small to minute or even absent ppp. In addition, although comparisons of external traits are clearly limited by the small sample sizes, individuals from southern populations appear to be more distinctly bicolored, including more pure white on the belly and dorsal surface of the manus and pes than those from northern populations, which are generally characterized by gray-cream ventral coloration. One-way ANOVA and

Tukey tests performed on craniodental measurements of comparable samples (Table 2) revealed significant differences between southern (Santa Cruz) and northern (Somuncurá) populations in zygomatic plate length and both mandibular and maxillary molar alveolar lengths.

Genetic variation and genealogy of *Notiomys edwardsii* haplotypes Partial cyt b gene sequences are available for three specimens of *N. edwardsii* captured at different localities. Each specimen showed a different haplotype. Haplotypes recovered in Campo Anexo Pilcaniyeu (locality #1) and Estancia Laguna Manantiales (#42), some 760 km apart, are 0.64% divergent. The haplotype recovered in Laguna Blanca (#8) diverges 0.75% from that of Pilcaniyeu (300 km) and 0.74% from the specimen from Manantiales (700 km). The genealogical analysis recovered three shortest trees of 201 steps ($CI=0.920$; $RI=0.853$), consensus of which (data not shown) shows a highly supported ($BS=100$) *N. edwardsii* clade in which haplotypes from Laguna Blanca (#8) and Pilcaniyeu (#1) are sisters to each other ($BS=75$).

Fossil record Craniodental remains of *Notiomys edwardsii* were recovered in four Holocene deposits in Patagonia (Table 1; Figure 1). In the Watkins cave sequence, which covers approximately the last 1000 years, *N. edwardsii* is recorded at high frequencies in all stratigraphical levels. The same is true for the Talagapa sequence and Los Altares profile. Both the Los Altares profile and Epullán Grande cave are outstanding for *N. edwardsii* because they fall in areas where this rodent is not currently found (Figure 1). At the former site, *N. edwardsii* is present from lower levels (dated approx. 2200 years) to upper levels (<100 years), suggesting very recent local extinction for the species. The case of Epullán Grande cave is somewhat different; a single *N. edwardsii* was recorded in levels dated as 7060 ± 90 radiocarbon years before the present (Crivelli Montero et al. 1996). Taphonomically, microvertebrate assemblages at all these sites were mainly produced by the raptor activities of owls (Pardiñas 1999).

Reproductive data Almost nothing is known about the reproduction of *Notiomys edwardsii*. A young female (CNP 1) with an imperforate vagina was recorded at the end of the summer season (March).

Natural history Hershkovitz (1966) discussed the degree of fossoriality reached by sigmodontine rodents. He noted two morphologic extremes in this tendency, one convergent with moles (exemplified by *Blarinomys* within sigmodontines) and the other with gophers (exemplified by *Kunsia*). *Notiomys* has a morphologic bauplan that was assumed to be indicative of a fossorial or semi-fossorial habit (Pearson 1984) and appears to be in line with mole-like adaptations. This bauplan includes, in general terms, reduced eyes and ears, shortened tail, short and velvety pelage, well-developed manus claws, narrow zygomatic plate, rounded and expanded braincase with reduced interparietal bone, wide interorbital region, and reduced molars. In *Notiomys*, the degree of reduction of the ears is outstanding for sigmodontine

Table 2 Measurements (in mm) of *Notiomys edwardsii* individuals and populational groups.

| | MVZ 163067 | | | MVZ 163065 | | | GNP 1 | | | Santa Cruz (localities #40, 41, 44, 45) | | | Somuncurá plateau (localities #5, 9, 10, 14, 31) | | | Northwestern Chubut (localities #21, 22, 25, 28) | | |
|---------------------------------------|------------|----|-------|------------|-------|---|-------|-------|-------|--|-----|---|---|------|----|---|-------|----|
| | x | SD | n | x | SD | n | x | SD | n | Min | Max | n | x | SD | n | Min | Max | n |
| Occipito-nasal length | 24.23 | | 23.12 | 20.68 | 22.62 | | | | | | | | | | | | | |
| Condyle-incisive length | 21.93 | | 20.95 | 20.68 | 20.68 | | | | | | | | | | | | | |
| Interorbital breadth | 4.83 | | 4.77 | 5.03 | 4.93 | | 0.12 | 4.78 | 5.14 | 9 | | | 4.96 | 0.21 | 6 | 4.70 | 5.25 | 6 |
| Zygomatic breadth | 13.51 | | 12.26 | 11.84 | 11.84 | | | | | | | | | | | | | |
| Braincase breadth | 11.52 | | 11.14 | 11.16 | 11.16 | | | | | | | | | | | | | |
| Rostral breadth | 5.02 | | 4.24 | 4.34 | 4.34 | | 0.18 | 4.93 | 5.49 | 9 | | | 4.89 | 0.31 | 4 | 4.45 | 5.13 | 4 |
| Rostral length | 8.40 | | 7.85 | 7.85 | 7.85 | | 0.52 | 7.78 | 9.01 | 4 | | | 8.51 | - | 2 | 8.37 | 8.65 | 2 |
| Rostral height | 4.00 | | 3.54 | 3.95 | 3.95 | | 0.21 | 3.75 | 4.20 | 4 | | | 3.70 | - | 2 | 3.53 | 3.87 | 2 |
| Nasal length | 9.39 | | 9.29 | 9.34 | 9.34 | | 0.32 | 9.42 | 10.15 | 4 | | | 9.43 | - | 2 | 9.33 | 9.53 | 2 |
| Incisive foramen length | 4.73 | | 4.41 | 4.32 | 4.41 | | 0.24 | 4.52 | 5.17 | 9 | | | 4.61 | 0.34 | 12 | 4.02 | 5.32 | 12 |
| Incisive foramen breadth | 1.76 | | 1.80 | 1.49 | 1.49 | | 0.10 | 1.58 | 1.86 | 9 | | | 1.95 | 0.21 | 4 | 1.77 | 2.21 | 4 |
| First molar breadth | 1.14 | | 1.09 | 1.05 | 1.05 | | 0.07 | 0.88 | 1.08 | 8 | | | 1.03 | 0.05 | 13 | 0.92 | 1.12 | 13 |
| Temporal fossa length | 7.30 | | 6.56 | 5.87 | 5.87 | | | | | | | | | | | | | |
| Diastema length | 6.65 | | 6.30 | 5.89 | 5.89 | | 0.43 | 6.47 | 7.76 | 8 | | | 7.02 | 0.45 | 13 | 6.08 | 7.63 | 13 |
| Palatal length | 11.91 | | 11.27 | 11.00 | 11.00 | | 0.31 | 11.52 | 12.39 | 8 | | | 11.18 | 0.73 | 3 | 10.35 | 11.72 | 3 |
| Postpalatal length | 8.02 | | 7.97 | 7.83 | 7.83 | | | | | | | | | | | | | |
| Palatal bridge length | 4.96 | | 4.44 | 4.09 | 4.09 | | | | | 1 | | | 4.76 | 0.24 | 3 | 4.53 | 5.01 | 3 |
| Palatal breadth at first molar | 3.28 | | 2.90 | 3.33 | 3.33 | | | | | 1 | | | 3.47 | 0.49 | 3 | 2.98 | 3.95 | 3 |
| Palatal breadth at third molar | 3.21 | | 2.96 | 2.79 | 2.79 | | | | | | | | | | | | | |
| Zygomatic plate length | 1.91 | | 1.70 | 1.65 | 1.65 | | 0.19 | 1.31 | 1.86 | 18 | | | 1.80 | 0.18 | 19 | 1.55 | 2.24 | 19 |
| Skull height | 7.38 | | 7.08 | 7.08 | 7.08 | | 0.20 | 7.07 | 7.69 | 9 | | | 7.36 | 0.26 | 3 | 7.07 | 7.56 | 3 |
| Bullar width | 4.45 | | 4.53 | 4.59 | 4.59 | | | | | | | | | | | | | |
| Bullar length | 3.81 | | 3.79 | 3.92 | 3.92 | | | | | | | | | | | | | |
| Breadth across incisor tips | 1.81 | | 1.63 | 1.59 | 1.59 | | 0.13 | 1.65 | 2.01 | 8 | | | 1.84 | 0.10 | 3 | 1.73 | 1.91 | 3 |
| Mesopterygoid fossa breadth | 1.28 | | 1.29 | 1.17 | 1.17 | | 0.20 | 1.40 | 1.80 | 4 | | | 1.24 | - | 1 | - | - | 1 |
| Molar row, crown length | 3.35 | | 3.29 | 3.35 | 3.35 | | | | | | | | | | | | | |
| Molar row, alveolar length | 3.44 | | 3.45 | 3.37 | 3.37 | | 0.28 | 3.60 | 4.58 | 10 | | | 3.73 | 0.22 | 13 | 3.30 | 3.99 | 13 |
| Greatest length of mandible | 12.97 | | 11.92 | 11.87 | 11.87 | | 0.84 | 11.38 | 13.98 | 16 | | | 12.84 | 0.64 | 36 | 11.50 | 13.93 | 36 |
| Depth of ramus | 2.80 | | 2.66 | 2.51 | 2.51 | | 0.13 | 2.96 | 2.85 | 17 | | | 2.58 | 0.14 | 43 | 2.34 | 2.89 | 43 |
| Mandibular molar row, alveolar length | 3.51 | | 3.29 | 3.48 | 3.48 | | 0.22 | 3.53 | 4.31 | 17 | | | 3.81 | 0.18 | 47 | 3.32 | 4.25 | 47 |

x, mean; SD, standard deviation; min, minimum; max, maximum; n, sample size. Locality numbers refers to those of Table 1.

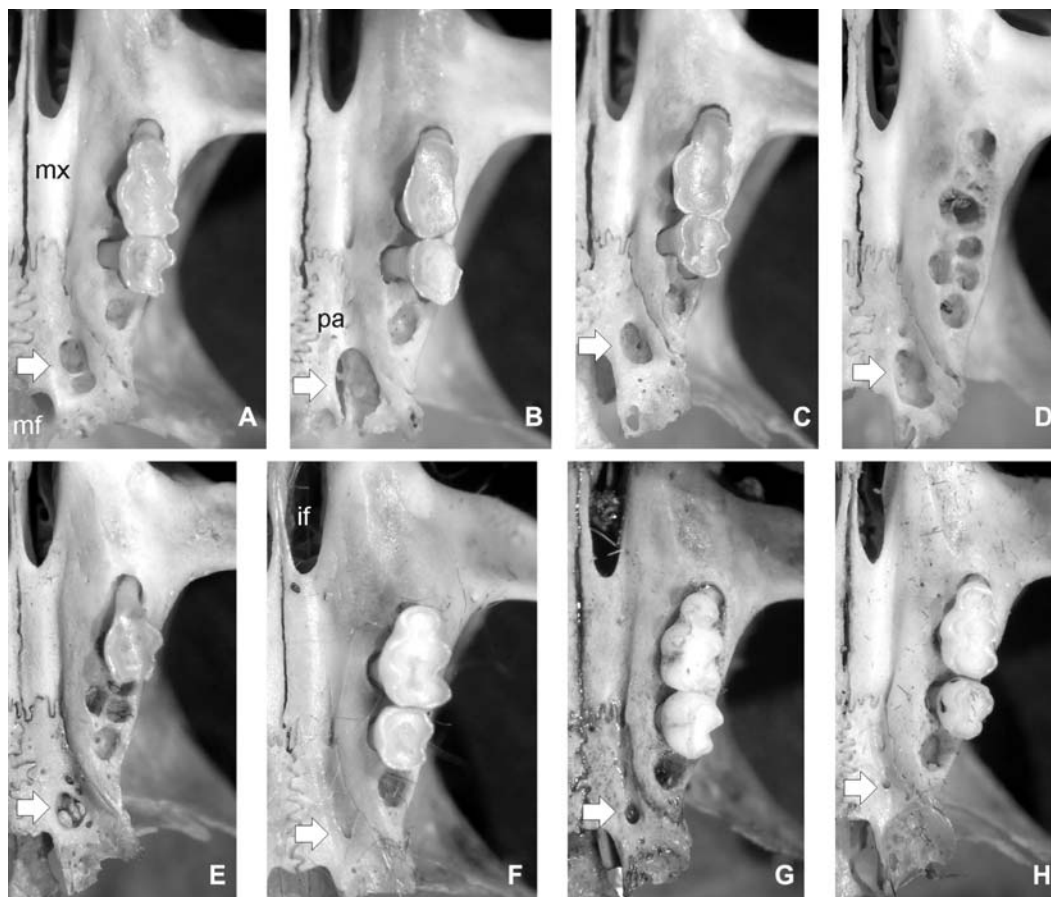


Figure 7 Morphological variation in the size of posterolateral palatal pits (indicated by arrows) in *Notiomys edwardsii* from southern populations. (A–D) N end Lago Cardiel and RN 40 (locality #44 in Table 1); (E) Laguna del Diez (#41), and northern populations; (F) Laguna del Paraguay (#5); (G) Paso del Sapo (#28); (H) Cerro Castillo (#17). Abbreviations: mx, maxillary; mf, mesopterygoid fossa; pa, palatine; if, incisive foramina.

rodents, and are even more reduced than those of *Blarinomys* (Matson and Arawaya 1977). In addition, the tail length (32% of total length, $n=5$) places *Notiomys* among the sigmodontines with the shortest tails. However, the eye of *Notiomys* is medium in size, suggesting cursoriality to a greater extent than for other mole-like sigmodontines. The same could be said with respect to the intriguing brightness of its pelage and contrasted bicolored pattern. Regrettably, almost nothing is known about the natural history of *Notiomys* (Pearson 1984) and its potential capacity to make burrows itself, beyond that few individuals were obtained in traps settled in entrances of *Ctenomys* burrows (Pearson 1984, R. Sage personal communication). Regarding diet, analyses of three stomachs revealed insects, arthropods, and vegetal seed remains (Pearson 1984, Martin and Archangelsky 2004). Some morphological traits, such as the obsolescence of the upper free border in the zygomatic plate, the rapid decay in molar occlusal structures, and the reduction in both size (especially M3/m3) and root number of molar teeth, may be linked to a diet dominated by soft items. In summary, the evidence at hand, although tenuous, appears to indicate that *Notiomys* is possibly a subfossorial small rodent with a capability to dig in search of invertebrate prey. The adaptations to this niche involve a particular mosaic of morphological traits largely shared with strict fossorial forms; this mosaic has been achieved

independently several times (e.g., *Blarinomys*, *Oxymycterus*, and *Kunsia* within Akodontini; D'Elía 2003) within the sigmodontine radiation.

Environments and conservation The 47 contemporary recording localities known for *Notiomys edwardsii* (Table 1; Figure 1) are strongly associated with two districts of the Phytogeographical Province (PP) Patagónica (*sensu* León et al. 1998); 12 recording localities fall in the Central district and 33 are in the Occidental district. The remaining two localities are located in the Subandino district (also belonging to the PP Patagónica). Central and Occidental districts include adverse habitats, basaltic plateaus, and hilly landscapes, dominated by poorly-developed shrubby steppes (León et al. 1998). Differences in the number of recording localities for both districts may be attributable to asymmetric sample efforts. In contrast, the absence of *N. edwardsii* in contiguous districts, such as Golfo de San Jorge, Ecotone Monte Patagonia and PP del Monte are based on thousands of studied samples (Pardiñas et al. 2003, Pardiñas and Udrizar Sauthier unpublished data). Therefore, it is safe to consider *N. edwardsii* as an endemic species of PP Patagónica.

Numerous collections of *Notiomys edwardsii* from Somuncurá (localities #3–13; Table 3), a large basaltic plateau covering ca. 15,000 km² in north central Patagonia (Croce 1963), invite exploration of microscale hab-

Table 3 Minimum number of *Notiomys edwardsii* individuals (and percentage of the total number of individuals of the sample) represented in owl pellets at specific Patagonian localities.

| # | Locality | <i>N. edwardsii</i> | Total sample | % |
|----|----------------------------------|---------------------|--------------|------|
| 31 | 4 km S Tres Banderas on RP 11 | 4 | 1312 | 0.3 |
| 26 | Cañadón de la Buitrera | 1 | 242 | 0.4 |
| 30 | Campo de Cretón | 1 | 113 | 0.9 |
| 23 | Fofo Cahuel | 1 | 81 | 1.2 |
| 41 | Laguna del Diez | 1 | 79 | 1.3 |
| 40 | Cueva de las Manos | 2 | 122 | 1.6 |
| 25 | Tres Cuevas | 1 | 57 | 1.8 |
| 21 | 2 km NW Gastre | 1 | 51 | 2.0 |
| 45 | Parador Luz Divina, RN 40 | 2 | 47 | 4.3 |
| 9 | Cerro Corona | 8 | 178 | 4.5 |
| 44 | N end Lake Cardiel and RN 40 | 15 | 213 | 7.0 |
| 14 | Subida del Naciente | 18 | 201 | 9.0 |
| 5 | Laguna del Paraguay | 8 | 78 | 10.3 |
| 10 | 3 km N Cuatro Cerros on route 60 | 11 | 101 | 10.9 |

itat conditions there (Figure 8). The landscape is dominated by a herbaceous steppe (“coironal”) composed of *Poa ligularis* (“coirón poa”) and *Festuca pallens* (“coirón blanco”), approximately 30% bare ground (Beeskov et al. 1982), and several basaltic outcrops ringing temporary small lagoons.

In the Argentine Red Book (SAREM 2000), *Notiomys edwardsii* is categorized as Vulnerable B1 (severely fragmented or known to exist at no more than ten localities). This categorization was correct at that time, because there were only eight recording localities for *N. edwardsii* (Pardiñas and Galliari 1998). The data provided in the present paper clearly indicate that *N. edwardsii* is widely distributed in Patagonia, reaching moderate frequencies at some localities and environments. This suggests that *N. edwardsii* should be categorized as Lower risk/least concern, which is in fact the category proposed earlier, albeit without explanation, by the International Union for Conservation of Nature (IUCN) for the species (Contreras 1996).

Remarks Pearson (1984: 233) suggested a potential connection between *Notiomys edwardsii* and *Hespero-*



Figure 8 Typical environment occupied by *Notiomys edwardsii* in the Somuncurá plateau (Río Negro, Argentina). The foreground shows *Festuca* and *Poa* grasslands (>1300 m) that surround Laguna Blanca (second plane, Cerro Corona).

mys (Acodon) michaelsoni Matschie 1898, a relatively obscure form described on the basis of a specimen from Punta Arenas (southern Chile) and generally allocated under *Geoxus*. The sole information on *michaelsoni* is, regrettably, that originally published by Matschie (1898); the type appears to be lost. Judging by the pictures and data provided by Matschie (1898: 5–6 and Figure 1), *Hesperomys (Acodon) michaelsoni* does not resemble *N. edwardsii*; *michaelsoni* has a more slanted zygomatic plate, enlarged rostrum, developed interparietal bone, and large manus claws that clearly contrast with the state characters of *N. edwardsii*. All these traits suggest a clear connection with *Geoxus*, as previously discussed and accepted by Osgood (1925).

Final considerations

When Osgood (1925) subsumed *Chelemys*, *Geoxus*, and *Notiomys* in a single genus (to which the name *Notiomys* applied for priority reasons), the generic distinctiveness of *N. edwardsii* was emphatically defended by Thomas (1927: 656): “broad muzzle and interorbital region, squared supraorbital edges, widely splayed anteorbital foramina, narrow, vertical, non-projecting zygomatic plate, short palatal foramina, and minute brachyodont molars make up an ensemble to which one cannot do otherwise than give generic rank.” Despite the arguments put forward by Thomas, and in line with the paradigm that dominated sigmodontine systematics during the decades 1940–1970, *Notiomys* remained broadly defined until the revisionary work of Pearson (1984). Our ongoing studies on Abrotrichini (D’Elía et al. 2006, 2007) highlight that, even when *Notiomys sensu lato* is monophyletic (i.e., phylogenetically a broad *Notiomys sensu* Osgood 1925 is supported), in accordance with current classification it is best to keep the genus *Notiomys* restricted to *N. edwardsii* alone and to keep using *Chelemys*, *Geoxus*, and *Pearsonomys* as valid genera. In fact, morphological traits displayed by *Notiomys* are clearly more trenchant than those that distinguish *Chelemys*, *Geoxus*, or *Pearsonomys*.

Departing from most broadly distributed sigmodontines, *Notiomys edwardsii* does not have any taxon under its synonymy. Moreover, there are no studies assessing

patterns of geographic variation of this species. The results presented here indicate that across its large distribution (approx. 9° of latitude), *N. edwardsii* exhibits geographic variation, both at the morphological and genetic level. Morphologically, specimens from south of the Río Deseado are qualitatively and quantitatively different from those of the north. Although available sampling precludes an in-depth comparison, genetic variation appears to follow a similar geographic pattern. Even when divergence values are similar for all haplotype pairs (range 0.64–0.75%), the recovered genealogy shows that haplotypes from localities north of the Río Deseado are more closely related to each other than to the haplotype recovered at the single locality south of the river we analyzed. Clearly, considering the few samples available and our poor understanding of intrapopulation variation within *N. edwardsii*, it is premature to hypothesize about the biological meaning, if any, of these patterns of geographic variation. However, these results invite further exploration of species limits among populations of *N. edwardsii*.

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