New toxodontid (Notoungulata) from the Early Miocene of Mendoza, Argentina

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Abstract We describe a new toxodontid species, Nesodon taweretus sp. nov., from the Aisol Formation in Mendoza Province, central-west Argentina. Nesodon is a frequently found Toxodontidae, member of the Notoungulata, an extinct endemic group of Cenozoic South American mammals that are ecologically similar to current hoofed ungulates. The holotype of N. taweretus sp. nov. is a skull, and we tentatively assign some mandibular fragments and postcranial bones. N. taweretus sp. nov. differs from the other Nesodon species in several cranial and dental features, and close comparisons were made with the Patagonian Nesodon imbricatus, common in the Santa Cruz Formation (Santacrucian Age, Early Miocene). The material is of a similar size to N. imbricatus, with a body mass estimation of about 550 kg. The phylogenetic analysis groups N. taweretus sp. nov. with other species of Nesodon. The absolute age of the Aisol Formation has been established at ca 19.480 ± 0.025 Ma (Burdigalian; Early Miocene). The vertebrate association is encompassed by the Santacrucian Age. Latitudinal separation between Mendoza and Patagonia in the south would have favored taxonomic differences, as reflected in the species of Nesodon.

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Keywords Nesodon · South America · Cenozoic · Early Miocene · U–Pb (CA-ID-TIMS) Tuff age

Kurzfassung Wir beschreiben eine neue Art der Toxodontiden, Nesodon taweretus sp. nov., aus der Aisol-Formation in der Provinz Mendoza, im Zentralwesten von Argentinien. Nesodon ist ein häufig vorkommendes Taxon der Toxodontidae, die zu den Notoungulaten, einer ausgestorbenen Gruppe von endemischen Säugetieren des Känozoikums in Südamerika, gehören und den rezenten gehuften Ungulata ökologisch ähnlich waren. Der Holotyp von N. taweretus sp. nov. ist ein Schädel, dem wir vorläufig einige Unterkieferfragmente und postkraniale Knochen zuweisen. N. taweretus sp. nov. unterscheidet sich von den anderen Nesodon-Arten in mehreren Merkmalen des Schädels und der Bezahlung. Enge Vergleiche wurden mit dem aus Patagonien stammenden und in der Santa-Cruz-Formation (Santacruzium, frühes Miozän) verbreiteten Nesodon imbricatus gemacht. Das Material ist von ähnlicher Grösse wie N. imbricatus, Schätzungen der Körpermasse ist etwa 550 kg. Die phylogenetische Analyse gruppiert N. taweretus sp. nov. mit anderen Arten von Nesodon. Das absolute Alter der Aisol-Formation wurde mittels der U–Pb Zirkondatierung auf etwa 19.480 ± 0.025 Ma (Burdigalium, Unteres Miozän) gesetzt. Die Wirbeltiervergesellschaftung in das Santacruzium eingeschlossen worden. Latitudinale Trennung zwischen Mendoza und Patagonien im Süden hätte taxonomischen Unterschiede begünstigt, wie das anhand der Arten von Nesodon widergegeben ist.

Schlüsselwörter Nesodon · Südarmerika · Känozoikum · frühes Miozän · U–Pb (CA-ID-TIMS) Tuff Datierung

Introduction

For more than a century, Patagonia in Southern Argentina has provided rich fossil localities for the study of Cenozoic mammals. Age calibrations, coupled with the quality and the quantity of the Patagonian fossil record, provide excellent material to diagnose the evolutionary history of the South American biota. The north of South America has provided a less fecund fossil record and has demonstrated a different taxonomic composition to the southern part of the continent (e.g., the Paleogene from Santa Rosa, Peru; the Oligocene from Salla, Bolivia; the Miocene from La Venta, Colombia, and Urumaco, Venezuela; Kay et al. 1997; Campbell 2004; Sánchez Villagrá et al. 2010; Shockey and Anaya 2008). Mendoza Province in central-western Argentina lies to the north of the windswept plains of Patagonia, and is in a suitable geographical location to provide data linking both biogeographic scenarios of the South American vertebrate evolution.

Mendoza Province has been the subject of new paleontological studies in the last few years, with a great deal of information on the evolution and distribution of Cenozoic mammals coming to light (Cerdeño 2011). One of the areas studied and prospected is the Aisol Formation outcropping in the San Rafael Department, in the center of the province (Fig. 1a) (Forasiepi et al. 2009, 2011; Vucetich et al. 2011). The Aisol Formation outcrops in isolated deposits that are exposed on the San Rafael Block and mainly associated with the Atuel and Diamante rivers (Dessanti 1954, 1956; González Díaz 1972; Sepúlveda et al. 2001, 2007). Several fossiliferous localities have been found in the area, with the richest being the Arroyo Seco de la Frazada (Fig. 1a). The stratigraphy of the Aisol Formation has been subject of recent revisions (Forasiepi et al. 2011). Here, the Aisol Formation corresponds to the lower section (LS) described by Forasiepi et al. (2011; Fig. 1b). The fossil vertebrate association includes Anura, Testudininae, Xenarthra, Notoungulata, Litopterna, and Rodentia (Forasiepi et al. 2011). To date, the chronology of the Aisol Formation has been controversial. Early authors suggested that the unit was deposited during the late Early Miocene (Santacrucian Age; Dessanti 1954, 1956; González Díaz 1972). Later fossil mammal finds suggested a middle Miocene age (Friasian s.l. sensu Soria 1983 or Colloncuran sensu Forasiepi et al. 2011). This contribution presents a new radiometric dating of a 19.480 ± 0.025 Ma tuff bed, which constrains the absolute age of the mammalian assemblage at these latitudes to the Early Miocene.

The Notoungulata is a diverse and abundant clade of extinct South American ungulates recorded from the Paleocene to the Pleistocene (Bond et al. 1995; Bond 1999). Following the formation of the Panama Bridge, the distribution of the group increased from South America to Central and North America (Polaco et al. 2004; Lundelius et al. 2013). Their phylogenetic affinities are still unclear. Notoungulates were related to other South American native ungulates (Astrapotheria, Litopterna, Pyrotheria, and Xenungulata) in the group Meridiungulata (McKenna 1975). However, recent analyses dealing with the main phylogenetic relationships of placental mammals recorded affinities of notoungulates and Afrotheria (O’Leary et al. 2013).

Toxodontids are medium to large Notoungulata of the Suborder Toxodontia, characterized by a specialized anterior dentition, with hypselodont incisor tusks and hypsodont molars. Toxodontids diversified during the Cenozoic and presented the richest diversity in the Miocene (Cifelli 1985; Marshall and Cifelli 1990; Nassif et al. 2000). During the Early and Middle Miocene, toxodontids were represented by Nesodon imbricatus, N. conspurcatus, N. cornutus (Santacrucian SALMA in Argentina and Chile); Adinothereium ovinum and A. robustum (Santacrucian SALMA in Argentina); Proadinothereium muensteri (Colhuehuapian...
SALMA in Argentina); *Palyeidodon obtusum* (Friasian s.s. and Colloncuran SALMAs in Argentina, Bolivia, and Chile), *Hyperoxtodon speciosus* (Santacrucian–Colloncuran SALMAs in Argentina and Chile), *Pericotoxodon plaginatathus* (Laventan SALMA in Colombia) (e.g., Pascal 1954; Marshall et al. 1983; Madden 1990, 1997; Nassif et al. 2000; Croft et al. 2003, 2004, 2009; Kramarz et al. 2005; Croft 2007). In addition, Quebrada Honda (Laventan SALMA in Bolivia) produced material that was referred to cf. *Paratrigodon* (Croft 2007); Cojitambo (Santacrucian? SALMA in Ecuador) produced *Pericotoxodon* sp. (Madden 1990); Chucal (Santacrucian SALMA in Chile) produced ?*Palyeidodon* (Bond and García 2002; alternative interpretation in Croft et al. 2004, 2009), and Río Collón Curá and Meseta de la Banda Negra (Colloncuran SALMA in Neuquén Province, Argentina) produced *Nesodon* sp. (Kramarz et al. 2011b). The genus *Nesodon* was abundant in the late Early Miocene and it was considered a taxon guide for the Santacrucian Age (Croft et al. 2003), although its record in the Colloncuran (Kramarz et al. 2011b) demonstrated that the genus continued to later ages.

The toxodontid remains recovered from the Aisol Formation (Forasiepi et al. 2011) consist of two well-preserved skulls, mandibular remains, and isolated postcranial elements. The analysis of the skull and tooth morphology allowed the recognition of two taxa: *Palyeidodon* cf. *P. obtusum* and a new Nesodontinae taxon (Forasiepi et al. 2011). This contribution provides a detailed systematic study of this taxon, recognized as a new species of *Nesodon*, the analysis of its phylogenetic relationships, and an estimation of its body mass.

Abbreviations

**AMNH**, American Museum of Natural History, New York, USA. **MACN**, Museo Argentino de Ciencias Naturales “Bernardino Rivadavia” (A. Ameghino Collection; **PV**, Vertebrate Paleontology Collection; **SC**, Santa Cruz Province Collection), Buenos Aires, Argentina. **MHNSR-PV**, Museo de Historia Natural de San Rafael, Vertebrate Paleontology Collection, San Rafael, Mendoza, Argentina. **YPM PU**, Yale Peabody Museum, Collection of Princeton University, New Haven, USA.

In the description of teeth and the corresponding tables, we use the following abbreviations: **L**, length. **M/m**, upper/lower molar. **P/p**, upper/lower premolar. **W**, width.

**Materials and methods**

**Dating**

Tuff samples were collected from the Aisol Formation to determine the absolute age by U–Pb zircon analysis via a combination of laser ablation inductively coupled plasma mass spectrometry (LA-ICPMS) and chemical abrasion isotope dilution thermal ionization mass spectrometry (CA-ID-TIMS). Tuff level 2 (Fig. 1b) was chosen as the most appropriate sample for dating based on its mineralogical composition and grain size. Approximately 3 kg of rock were processed in the Boise State University Isotope Geology Laboratory for zircon separation, cathodoluminescence (CL) images, and isotopic analysis; see the Electronic supplementary material (ESM) 1. An abundant population of large (approximately 100–300 microns in the major dimension), moderate to highly elongate prismatic zircon crystals were separated by conventional density and magnetic methods. The crystals were placed in a muffle furnace at 900 °C for 60 h in quartz beakers to anneal minor radiation damage. Annealing enhances cathodoluminescence (CL) emission, promotes more reproducible interelement fractionation during LA-ICPMS, and prepares the crystals for subsequent chemical abrasion (Mattinson 2005). Following annealing, individual grains were hand-picked and mounted, polished, and imaged by CL on a scanning electron microscope. From these compiled images, grains with consistent and dominant CL patterns were selected for further isotopic analysis.

U–Pb geochronology and trace element analysis by LA-ICPMS followed the methods described in Rivera et al. (2013) using a ThermoElectron X-Series II quadropole ICPMS and a New Wave Research UP-213 Nd:YAG UV (213 nm) laser ablation system. In-house analytical protocols, standard materials, and data reduction software were used for acquisition and calibration of U–Pb dates and a suite of high field strength elements (HFSE) and rare earth elements (REE). U–Pb and 206Pb/238U ratios and dates for unknowns were calibrated via interspersed measurements of the Plešovice zircon standard (Sláma et al. 2008). Signals at mass 204 were indistinguishable from zero following subtraction of mercury backgrounds measured during the gas blank (<1,000 cps 202Hg), so dates are reported without common Pb correction. Radiogenic isotope ratio and age error propagation for all analyses includes uncertainty contributions from counting statistics and background subtraction. A zircon secondary reference material AUSZ2 [38.9 Ma from unpublished chemical abrasion thermal ionization mass spectrometry (CA-TIMS) data, Boise State University] was interspersed as pairs of analyses for every 20 unknown analyses and yielded a weighted mean 206Pb/238U date of 38.6 ± 1.5 Ma (MSWD = 2.1, n = 8).

U–Pb geochronology by isotope dilution thermal ionization mass spectrometry follows the methods described by Davydov et al. (2010) and Schmitz and Davydov (2012). Zircon crystals were subjected to a modified version of the chemical abrasion method of Mattinson (2005), reflecting a preference for preparing and analyzing selected single
crystal fragments. All analyses were undertaken on crystals that had previously been mounted, polished, and imaged by cathodoluminence (CL), and selected on the basis of zoning patterns. U–Pb dates and uncertainties for each analysis were calculated using the algorithms of Schmitz and Schoene (2007) and the U decay constants of Jaffey et al. (1971). Uncertainties are based upon non-systematic analytical errors, including counting statistics, instrumental fractionation, tracer subtraction, and blank subtraction. These error estimates should be considered when comparing our 206Pb/238U dates with those from other laboratories that used tracer solutions calibrated against the EARTHTIME gravimetric standards. When comparing our dates with those derived from other decay schemes (e.g., 40Ar/39Ar, 187Re–187Os), the uncertainties in tracer calibration (0.05 %; Condon et al. 2007) and U decay constants (0.108 %; Jaffey et al. 1971) should be added to the internal error in quadrature. Quoted errors for calculated weighted means are thus of the form ±X(Y)[Z], where X is solely analytical uncertainty, Y is the combined analytical and tracer uncertainty, and Z is the combined analytical, tracer, and 238U decay constant uncertainty.

Systematics

The description of the skull MHNSR-PV 1004 is organized according to the principal regions: facial, palatal, orbital, vault, and base. The nomenclature for the basicranium and orbitotemporal region corresponds to Gabbert (2004), but according to the principal regions: facial, palatal, orbital, vault, and base. The nomenclature for the basicranium and orbitotemporal region corresponds to Gabbert (2004), but

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Zircon populations and geochronological data

CL imaging of 98 zircon crystals separated from the level-2 tuff sample revealed a bimodal population of grains: a predominant subgroup of moderately to brightly luminescent, weakly oscillatory to planar zoned crystals, which includes all elongate and many more equant crystals and a subsidiary subgroup of less intensely luminescent to non-luminescent crystals that exhibit stronger oscillatory and occasional sector zoning. Some crystals of the predominant luminescent subgroup have irregularly shaped, relatively non-luminescent cores overgrown by the luminescent rims, suggesting that this dominant subgroup is the primary magmatic population of the tuff, while the less luminescent grains and cores may be inherited or recycled from earlier volcanism. Seventy-four spot analyses by LA-ICPMS confirmed the presence of inherited cores and grains of Mesoproterozoic to Eocene age, coexisting with what may be interpreted as a single Miocene population of zircon represented by 44 spot analyses with a weighted mean age of 19.4 ± 0.3 Ma.

Six grains were selected for CA-TIMS analysis on the basis of morphology and CL pattern, avoiding those crystals with obvious resorbed non-luminescent cores but including five elongate grains (z1, z3, z4, z5, and z6) from the primary population and one grain (z2) from the secondary population. Four of the six analyses are concordant and equivalent, with a weighted mean 206Pb/238U date of 19.480 ± 0.009 (0.013) [0.025] Ma (MSWD = 0.85), which is interpreted as dating the eruption and deposition of this tuff (Fig. 2). Two other crystals yielded older 206Pb/238U dates of 19.55 and 20.31 Ma, the latter from the more strongly zoned secondary population of grains, supporting the inference from CL imagery and LA-ICPMS analysis of nuclei of inherited zircons in some magmatic grains (ESM 1, which includes a complete list of the analytical data).

Systematic paleontology

Notoungulata
Toxodontidae
Nesodontinae
Nesodon Owen 1846

Nesodon taweretus sp. nov.

Figures 3, 4, 5, 6, 7, 8, and 9

Derivation of name. Egyptian deity Taweret, meaning “she who is great” or “great one,” usually represented by the body of a hippopotamus, which reflects the robust appearance of the new species.

Holotype. MHNSR-PV 1004, mostly complete skull with right M1–M3 badly preserved and remains of the left molars. Alveoli of the remaining teeth are preserved, but the area of I1 is damaged.

Referred material. MACN-PV 18609, isolated upper molar.

Tentatively assigned material. MHNSR-PV 360, right mandible with part of the symphysis, remains of right i1 and i3, and p4 (broken)–m3; MHNSR-PV 1000, incomplete lower jaw with right m1–m3 and left m1 and m3. MHNSR-PV 122, distal fragment of left femur;

Fig. 2 U–Pb concordia diagram for zircon analyses from dated tuff (level 2 sample). The four red filled ellipses are included in weighted mean calculations.

Fig. 3 Skull MHNSR-PV 1004 of Nesodon taweretus sp. nov. in lateral (a), dorsal (b), and ventral (c) views. crn Crista meatus, frp frontal process of jugal, gf glenoid fossa, inf incisive foramen, ioj infraorbital foramen, lacj lacral foramen, mmp minor palatine foramen, mps major palatine foramen, mpf major palatine foramen, mpl major palatine process of premaxilla, mpl major maxillary foramen, ncr nuchal crest, nf nutrient foramen, npfl narial platform, oc occipital condyle, pdp palatine foramen, pp paracondylar process of exoccipital, ppb postglenoid process, plpx postero lateral process of premaxilla, pop postorbital process of frontal, pp postpalatine apophysis, pt pterygoid, ptp post tympanic process of squamosal, ram temp foramina for rami temporales, scr sagittal crest, sp styliform process, tl temporal line.
MHNSR-PV 1008, incomplete right ulna; MHNSR-PV 1009, left ulna lacking the distal epiphysis; MHNSR-PV 1010, complete left ulna.

Remarks. The mandibular fragments MHNSR-PV 360 and MHNSR-PV 1000 are assigned to the new species in contrast with two other fragments from the Aisol Formation, MHNSR-PV 1001 (right dentary fragment with m3 and partial m2) and MHNSR-PV 1075 (left dentary with the posterior part of the symphysis and p4–m3; ESM 2), whose larger size and morphological features (see description) promotes them as Palaeoiodon, the second toxodontid recovered from the Aisol Formation. However, some characters are inconclusive for a precise determination.

The postcranial bone morphologies of specimens MHNSR-PV 122, MHNSR-PV 1008, MHNSR-PV 1009, and MHNSR-PV 1010 agree with the morphology of the postcranium described for Nesodon (Scott 1912). No postcranial material of Palaeoiodon is currently identified, so our tentative assessment is to equate the new specimens with the new taxon.

Occurrence. The specimens MHNSR-PV 1000, MHNSR-PV 1004, MHNSR-PV 1008, MHNSR-PV 1009, MHNSR-PV 1010, and MACN-PV18609 come from the Arroyo Seco de la Frazada (the holotype, MHNSR-PV 1004, comes from 34°53'57.3"S/68°28'52.6"W); MHNSR-PV 360 and MHNSR-PV 122 come from Cuesta de los Terneros, Arroyo Cochicó; San Rafael Department, Mendoza Province, Argentina. Aisol Formation (Santacrucian Age).

Diagnosis. Species of Nesodon with a robust skull. N. taweretus differs from N. imbricatus in the following features: longer dorsal convexity, especially at the posterolateral border of frontals; W-shaped instead of U-shaped and slightly more retracted anterior border of nasals behind the anterior border of premaxilla; zygoma very deep at the level of the frontal process and with its maximum width at its median part instead of at the back; border of the choana at the level of the posterior border of M3; proportionally longer basicranium; lower basisphenoid–basioccipital tuberosities and lower median keel; more vertical occipital condyle; laterally instead of medi ally oriented paracondylar process; I2–I3–C without diastema; and molars with less persistent folds and fossettes. N. taweretus differs from N. cornutus by the lower and wider occiput.

Description and comparisons

Skull and upper dentition

The skull MHNSR-PV 1004 is slightly laterally compressed, but this does not affect the morphology or dimensions of different cranial sections. Similar to other toxodontids, the skull is narrow and tall. Several sutures are fused, including some in the facial region of the skull (e.g., lacrimal–frontal, jugal–maxilla), which usually remain open in late ontogenetic stages of other placental (Krogman 1930; Rager et al. 2014). Consequently, MHNSR-PV 1004 can be considered a late adult or senile specimen.

In lateral view (Fig. 3a), the upper profile of MHNSR-PV 1004 is shallowly sigmoid-shaped: it is convex at the level of the nasals and frontals, depressed at the level of the temporal region, and raised again at the level of the nuchal crest. In N. imbricatus, the dorsal convexity does not include the posterior part of the frontals and it varies from an almost flat upper profile (Scott 1912) to steeply rising from the forehead to the occiput (Scott 1912; per. obs.). After Scott (1912: figs. 35–36), N. cornutus has a more raised sagittal crest and less convex nasals; unfortunately, the type specimen YPM PV 16012 now lacks its postero dorsal region. In dorsal view (Fig. 3b), MHNSR-PV 1004 is roughly triangular in outline, with shallower prezygomatic constrictions and less expanded zygomatic arches than in N. imbricatus.

Face. The face is high, as is typical for herbivores, and formed by the premaxilla, maxilla, and nasal. In lateral view,
the premaxilla is slightly wider ventrally than dorsally. A very short posterolateral process interposes between the nasal and the maxilla. There are tiny foramina close to the premaxillo-maxillary suture and a large nutrient foramen on the right side of the skull (Fig. 3a). The narial platform is exposed in dorsal view. Similar to *N. imbricatus*, it has a small, blunt and dorsally projected anterior process.

The anterior nasal aperture is diamond shaped, narrower ventrally than dorsally and wider at the mid-dorsal half. The nasals are dorsally convex, with their anterior tip somewhat retracted from the anterior end of the premaxilla. The anterior border of the nasals is W-shaped. The nasals extend posteriorly to the level of the orbit and contact the frontal by means of a broad W-shaped suture;
this suture is an acute W in *N. imbricatus* and U-shaped in *N. cornutus*.

The maxilla is high and the main component of the snout in lateral view. It has a shallow groove at the level of the premolars, similar to *N. imbricatus* (Scott 1912). The infraorbital foramen is small and opens at the level of the anterior border of M2, as in *N. imbricatus*. The maxillary foramen is compressed against the zygomatic arch.

**Palate**

In ventral view, the premaxilla, maxilla, palate, and pterygoids are exposed (Fig. 3c) with feebly distinguished sutures. The maxillo-palatine suture is V-shaped, with the palate narrower and longer than in *N. imbricatus*. The palate is triangular, narrow, and deeply concave at the level of the premolars, becoming shallower and wider backward, especially at the level of M3. According to Scott (1912), the palate increases in depth with age, which supports the advanced ontogenetic stage suggested for MHNSR-PV 1004. At the midline of the palate, there is a thin crest that rises at the level of the posterior half of the P2 and reaches the choana. The small oval incisive foramina open in the premaxilla at the level of I2–I3. Lateral and medial incisive foramina are separated by a crest-like medial palatine process, similar to *N. imbricatus*. In MHNSR-PV 1004, a pair of large foramina open in the maxilla, at the level of the anterior part of M2, recognized as the major palatine foramina for the passage of the major palatine artery and nerve (Evans and de Lahunta 1972). The palate also has three small apertures at the anterior level of M3, one on the right side and two on the left side of the palate, as well as an extra aperture on the left palato-maxillary suture at the level of the posterior border of M3. These accessory openings probably transmitted bundles of the major palatine nerve and accompanying vessels, as occurred in some placental (Wible and Gaudin 2004). The most posterior foramen on the horizontal process of the palatine likely corresponds to the minor palatine foramen for the passage to the soft palate of the minor palatine artery and nerve (Evans and de Lahunta 1972).

In MHNSR-PV 1004, the posterior border of the palate and the border of the choana are aligned with the posterior border of the M3. This is different from *N. imbricatus*, in which the posterior border of the palate is oblique to the main axis of the skull, with the choana opening behind the level of M3 (Scott 1912). The choana in MHNSR-PV 1004 is single-arched and surrounded by blunt postpalatine apophyses that meet the pterygoid processes. The pterygoid processes are strongly divergent to the back, whereas in *N. imbricatus* they are straighter and much less divergent.

**Orbit**

The orbits are almost central to the skull (Fig. 3a), similar to *N. imbricatus* (Scott 1912). The lacrimal is small and exposed on the orbit and orbital rim, where it has a long spine that directs ventrally, contacting the maxilla. The lacrimo-jugal suture observed in *N. imbricatus* is missing in MHNSR-PV 1004. There is a single and large lacrimal foramen opening inside the orbit. The lacrimal process is rudimentary.

The orbitotemporal fossa is better preserved on the left side of the skull (Fig. 4). Three openings that are aligned and dorsally bordered by the orbital crest are the sphenorbital fissure, the optic foramen, and the ethmoidal foramen. The sphenoral fissure is the largest opening and is located in the posteroventral corner of the fossa. This would convey the maxillary and ophthalmic branches of the trigeminal nerve and the oculomotor, abducens, and trochlear nerves, as in bovids (Schaller 1992). The optic foramen is smaller and rostromedially placed to the sphenobital fissure. The ethmoidal foramen is the smallest. A small foramen opens posteriorly and ventrally to the frontal process, and it is identified as the aperture for the frontal diploic vein (following Wible and Gaudin 2004, describing the anatomy of the yellow armadillo).

The sphenopalatine foramen opens ventral to the ethmoidal foramen and at the level of the posterior border of the M3 (Fig. 4). This aperture is large and directs to the nasal cavity. In lateral view, it faces posteriorly.

**Vault**

The vault comprises the frontal, the parietal, and possibly the interparietal, with a large significance of the frontal (Fig. 3b). At the postorbital level, the frontal bulks, probably due to well-developed frontal sinus, and defines a convex surface, very conspicuous and more pronounced than in *N. imbricatus* and *N. cornutus* (Scott 1912). The postorbital processes are small. MHNSR-PV 1004 has subtle temporal lines, unlike *N. imbricatus* and *N. cornutus*, in which temporal lines are sharper and separate from the sagittal crest ending at the postorbital process (e.g., AMNH 9168) or before reaching the process (Scott 1912). The parietal is narrow and mostly restricted to the posterior part of the skull roof. The sagittal crest is shorter and lower in MHNSR-PV 1004 than in *N. imbricatus*. The squamosal largely contributes to the skull vault by means of a large squamous process that is roughly semicircular in shape, similar to *N. imbricatus*. There are two large foramina in the left and three in the right temporal fossa, on the squamosal and parietal, and close to the nuchal crest. These foramina are recognized as the rami temporales of the
stapedial artery and accompanying veins, following MacPhee (2014).

**Zygoma**

The zygomatic arch is short and robust and includes the jugal, squamosal, and a very short zygomatic process of the maxilla that forms its anterior base (Fig. 3a). The arch is very deep at the level of the frontal process, forming an angle between the front and the rear of the dorsal border of the arch. *N. imbricatus* displays an arch with a more constant depth. In ventral view (Fig. 3c), the maximum width of the arch is at its median part, while *N. imbricatus* displays the widest part of the arch at the back.

**Basicranium**

The good preservation of the specimen permits a detailed description of the basicranium (Fig. 5). Differences from *N. imbricatus* are expressed in absolute size and variations in proportions and the relative development of processes, crests, and tubercles.

The basicranium in MHNSR-PV 1004 is wide and long. Its length (measured from the level of the posterior choana to the posterior border of the condyle) is 39 % of the total skull length, while it is 36.5 % in *N. imbricatus* (Scott 1912: plate 15).

The occipital and sphenoid complexes form the floor of the braincase at midline; sutures between them are not visible. A pair of tuberosities develop close to the midline and medially to the bulla, and extend anteriorly up to the level of the sphenotympanic fissure. The specimen MACN-SC 4355 of *Adinotherium* has stronger tuberosities on the basisphenoid–basioccipital, and the same bone arrangement is expected for MHNSR-PV 1004. These tuberosities converge posteriorly in a feeble and short median crest. *N. imbricatus* has more robust, taller, and parallel tuberosities and a sharper median crest (Scott 1912).

On each lateral side of the sphenoid complex, MHNSR-PV 1004 shows a deep scaphoid fossa that would serve as an area of attachment for the tensor veli palatini muscles (Gabbert 2004). The fossa is surrounded by sharp medial and lateral crests. This fossa is shallower in *N. imbricatus*, with less developed crests.

The small auditory bulla is partially preserved on the right side of the skull. The bulla contacts the sphenoid complex anteriorly, the paracondyilar process posteriorly, and the crista meatus posterolaterally (Fig. 5). The bulla does not contact the lateral edge of the sphenoid along the basicochlear (=basicapsular) fissure, but this space could be broadened by breakage. In *N. imbricatus*, the bulla and the basisphenoid are sutured. The auditory bulla is broken ventrally in MHNSR-PV 1004. In *N. imbricatus*, the bulla projects ventrally beyond the level of the braincase, and it is even further ventrally projected in *Adinotherium*. In toxodontids, the bulla and part of the floor that covers the external acoustic meatus (MAE) ventrally is formed by the ectotympanic (Gabbert 2004). The participation of the entotympanic in the auditory bulla (e.g., Patterson 1936 contra Gabbert 2004), which is apparently the case for some notoungulates (MacPhee 2014), would be resolved with the analysis of juveniles. The crista meatus, a projection of the ectotympanic (Gabbert 2004; MacPhee 2014), is sharp and long in MHNSR-PV 1004, increases in height posterolaterally, and ends in a robust process (Fig. 5). This is similar to *N. imbricatus*, but *Adinotherium* demonstrates a thinner process.

At the right side of the skull, a long vertical styliform process (following van der Klaauw 1931 and Patterson 1932) was preserved fused to the anterior part of the bulla (Fig. 5). Scott (1912) suggested that a similar process in *N. imbricatus* would correspond to the tympanohyal, probably fused to the stylohyal (tympanostyloid in MacPhee 2014), although in notoungulates the cranial end of the hyoid is a narrow bar, lodged within the hyoid recess (MacPhee 2014). Alternatively, the styliform process is located on the anterior end of the auditory bulla and forms the prolongation of the Eustachian tube (van der Klaauw 1931).

The auditory region of *N. imbricatus* has extraordinary specializations (Scott 1912). The MAE forms a long tube, enclosed by the postglenoid process anteriorly, the crista meatus, and the post-tympanic process posteriorly (Gabbert 2004), and opens at an elevated position. In MHNSR-PV 1004, the MAE is at the level of the mid-height of the occiput (Fig. 6); this is slightly lower than in *N. imbricatus* (Scott 1912). Ventral to the MAE, there is a sulcus (“incisura mastoidea” sensu Roth 1903) excavated between the crista meatus and post-tympanic process, and similar to the toxodontians described by Gabbert (2004: fig. 14.2). In MHNSR-PV 1004, anterior to the MAE there is a circular foramen, slightly smaller than the MAE, which
we recognized as the suprameatal foramen following Gabbert (2004).

The glenoid cavity (Fig. 3c) is anterolateral to the bulla and dorsal to the floor of the braincase, as in other nesondontines and other notoungulates (Patterson 1932, 1934). The glenoid cavity is broader than long and saddle-shaped. In lateral view, the cavity is sigmoidal in shape: anteriorly convex and posteriorly concave.

The postglenoid (=retroarticular) process is low and represented by a swelling of the squamosal, similar to N. imbricatus. There is a single aperture for the postglenoid (=retroarticular) foramen on the posterior wall of the postglenoid process. The post-tempypanic (=retrotympanic) process is longer than the postglenoid process. In lateral view (Fig. 3a), the crista meatus interposes between both processes of the squamosal. There is a small foramen on the anterior base of the crista meatus (Fig. 5), identified as the stylomastoid foramen (following the topographic relationships of Cochilius in MachPhee 2014: figs. 13a, b).

Anterior to the bulla, there is a broad opening, the sphenotympanic fissure (Fig. 4) following Gabbert (2004) or piriform fenestra following MacPhee (2014). In the vicinity of the sphenotympanic fissure, independent carotid foramen, vidian foramen, and foramen ovale are not identified. All the soft tissues that these individual openings convey in other mammals would pass together through the sphenotympanic fissure in MHNSR-PV 1004, in common with other toxodontids (Gabbert 2004). On the sphenoid, there is a narrow groove that directs from the nasopharyngeal passage to the auditory cavity, ending at the anteromedial border of the sphenotympanic fissure that would contain the Eustachian tube. This groove is deeper in N. imbricatus.

Behind the bulla, the large jugular fossa is posteriorly bordered by the occipital complex. It is confluent with the basicochlear fissure. Lateral to the jugular fossa, bounded by the bulla anteriorly, the crista meatus anterolaterally, and the paracondylar process posteriorly, there is a deep depression identified as the hyoid recess for the tympanic-hyal and stylohyoid articulation (Gabbert 2004). The outline of the hyoid recess is slightly larger than the outline of the jugular fossa and oval in section, similar to N. imbricatus. There is a single hypoglossal foramen opening immediately posterior to the jugular fossa. This foramen is smaller than in N. imbricatus and Adinotherium.

In posterior view (Fig. 6), the occipital face is wide and low and defines a roughly semicircular outline, similar to N. imbricatus and different from the higher occiput of N. cornutus, which also has a higher sagittal crest (Scott 1912). The nuchal crests draw a semicircular curve, and they do not project further than the level of the occipital condyles. No sutures are observed in this area with the exception of part of the exoccipital–squamosal suture at the posterolateral corner of the occiput. There is no evidence of the mastoid process in the occipital view of MHNSR-PV 1004. According to Gabbert (2004), and differing from previous interpretations (Roth 1903; Scott 1912; Patterson 1932), the petrosal is not exposed in occipital view. The occiput in MHNSR-PV 1004 and other toxodontians is formed by the occipital complex and squamosal (Gabbert 2004; see also MacPhee 2014 for other notoungulates).

The foramen magnum is oval, with the main axis horizontal. The occipital condyles are oval and close to each other in ventral view (Figs. 3c, 5). The notch between the condyles is very deep. In lateral view, the condyles (Fig. 3a) project backwards from the occiput, but slightly less than in N. imbricatus. In posterior view (Fig. 6), the main axis of the condyle is vertical and the paracondylar process is laterally oriented instead of medially as in N. imbricatus, with a greater separation than in this taxon.

The occiput in MHNSR-PV 1004 (Fig. 6) has a deep depression, dorsal to the occipital condyle and at the level of the exoccipital-squamosal suture, with an aperture. In other notoungulates and pyrotheres, a similar opening was interpreted to be the mastoid foramen (Gabbert 2004; Billet 2010), but the possibility of being the posttemporal canal could not be discarded for some native ungulates (Kramarz et al. 2011a). In the specimen MACN 5305 of N. imbricatus, the aperture opens intracranially in the transverse sinus, and this would link the opening with the venous system. A similar relationship was recorded for other notoungulates (e.g., Paedotherium and Cochilius) discovered with detailed micro-CT images (MacPhee 2014: figs. 9d, 12e, 15d, respectively). However, the petrosal anatomy of a probable Itaboraian notoungulate evidenced the presence of an arteria diploe¨tica magna (Billet and de Muizon 2013). This suggests homologies of the posterior occipital opening with the posttemporal canal for the passage of the arteria diploe¨tica magna and accompanying vein (MacPhee 2014). Lateral and dorsal to the depression, the occiput bulges, probably due to the well-developed epitympanic sinus. Those internal cavities were recognized in other toxodontians (Gabbert 2004; MacPhee 2014).
Upper dentition

The dental formula of the skull is I1–3 C1 P1–4 M1–3, as in other toxodontids (Scott 1912; Madden 1997). The anterior alveoli are tightly packed and lack diastemata between I2–I3 and C, which are present in *N. imbricatus*. Molars are moderately imbricated. The size increases rapidly from M1 to M3 (Fig. 7a, b). The layer of enamel covers the labial and anterolingual surfaces of the teeth. The occlusal outline is slightly anterolabial–posterolingually elongated. The original Y-shaped lingual fold is lost with wear, isolating a central fossette. It is ovoid and rather reduced in M1, but much longer and narrower in M2–M3. In the M3, the Y-shape is still visible in the fossette, with a long anterior

**Fig. 7** Upper dentition of *Nesodon taweretus* sp. nov. and dentaries tentatively assigned to it. MHNSR-PV 1004, right maxilla with M1–M3 in occlusal view (a, b). MHNSR-PV 1000, lower jaw in lateral (c) and occlusal view (d), and detail of left m2, and right m1 and m3 (e) (not to scale). MHNSR-PV 360, lower jaw in lateral view (f) and detail of m1–m3 in occlusal view (g, h) (not to scale). *af* Accessory fossettid, *con* mandibular condyle, *cor* coronoid process, *ehf* ento-hypoconid fold or fossettid, *h* hypoflexid, *k* median symphyseal labial keel, *mef* meta-entoconid fold or fossettid, *mf* mental foramen
ramus and a short posterior one. The M3 also has a postfossette and a shallow lingual groove limiting the protocone. The ectoloph is more concave at its anterior third.

Comparing with other Miocene toxodontids, the molars of MHNSR-PV 1004 are much larger than those of Adinotherium and about the same size as those of N. imbricatus. Adinotherium has molars with a straighter ectoloph and more persistent folds and fossettes than those of N. imbricatus. The M3 also has a postfossette and a shallow lingual groove limiting the protocone and a short posterior one. The M3 also has a postfossette and a shallow lingual groove limiting the protocone. The ectoloph is more concave at its anterior third.

The preserved molars in MHNSR-PV 360 (Fig. 7g, h) show some differences in m3 from that of MHNSR-PV 1000. This molar has two main lingual folds that are larger than those of N. taweretus, and there are diastemata between i3–C–P1 (Scott 1912). The molars of N. taweretus have less persistent folds and fossettes than those of N. imbricatus, which in extremely worn molars still show the posterior ramus of the lingual fold. All three taxa show closed fossettes by the confluence of protoloph and metaloph forming lingual columns, and Hyperoxodontodon, with more elongated molars and without postfossette (Roth 1899; Kraglievich 1930; Cabrera and Kraglievich 1931; Madden 1990, 1997), show that the lingual fold remains open.

Mandible and lower dentition

Mandible

Four toxodontid mandibular fragments that were recovered from the Aisol Formation proved difficult to assign taxonomically. MHNSR-PV 1001 is larger than the other three mandibular fragments and MHNSR-PV 1075 is more suitable to Palyeidodon than Nesodon (ESM 2). We provisionally assign MHNSR-PV 1000 and MHNSR-PV 360 to Nesodon, and MHNSR-PV 1001 and MHNSR-PV 1075 to Palyeidodon, with the following comparative description outlining these assignations.

MHNSR-PV 1000 is an incomplete mandible lacking the anterior part of the symphysis and the vertical ramus (Fig. 7c, d). The teeth have advanced wear, comparable to that of the upper teeth in the skull and corresponding to a more aged individual than MHNSR-PV 360. The horizontal ramus is robust.

The ventral border is convex (less evident in MHNSR-PV 360 at the level of m2–m3). The posteroventral inflexion of the dentary shown in N. imbricatus (Scott 1912: pl. 13) is absent in MHNSR-PV 1000 and very shallow in MHNSR-PV 360. The symphysis is fused and lacks any vestiges of the suture. It extends back to the level of p4–m1 embrasure in MHNSR-PV 1000 or to the level of p4 in MHNSR-PV 360. Both morphologies were observed in specimens of N. imbricatus, which was related to the large individual variations in the extension of the mandibular symphysis observed for this species (Scott 1912: 187). Both specimens have two mental foramina in lateral view and one additional foramen in anterior view and below i3. An interesting feature preserved in MHNSR-PV 360 is a thin, median symphyseal labial keel below the alveolar border. This area is not preserved in the remaining specimens from the Aisol Formation and therefore it is not comparable. A low median symphyseal keel is present in N. imbricatus, Xotodon, and Calchaquitherium (Nassif et al. 2000).

The specimen MHNSR-PV 1075 assigned to Palyeidodon (ESM 2) shows a higher vertical ramus than MHNSR-PV 360, with its posterior border more projected backward from the condyle. The ventral border of the horizontal ramus is straighter than in MHNSR-PV 1000 and MHNSR-PV 360. Both features are observed on the holotype of Palyeidodon obtusum (Madden 1990: 155, fig. 29b).

Lower dentition

Lower molars of MHNSR-PV 1000 (Fig. 7e) present a wide, labially convex trigonid; they have meta-entoconid and ento-hypoconid fossettids. The latter is small and has almost disappeared in m2. The m1 is wider at the occlusal surface than at the alveolar level. The hypoflexid is deeper in m3.

The preserved molars in MHNSR-PV 360 (Fig. 7g, h) show some differences in m3 from that of MHNSR-PV 1000. This molar has two main lingual folds that are largely open and, as a consequence, meta-entoconid and ento-hypoconid fossettids are not yet formed (the meta-entoconid fold is still barely open in m2). Instead, two other fossettids are well defined: one close to the labial wall and centrally placed in the talonid, equivalent to the accessory fossettid defined by Madden (1990), and another is posteriorly placed (Fig. 7h). The accessory fossettid is commonly present in Nesodon (Scott 1912) but is absent in Palyeidodon (Madden 1990), which supports the view that MHNSR-PV 360 belongs to the new species of Nesodon. However, the specimen MHNSR-PV 1075 assigned to Palyeidodon (ESM 2) has a small accessory fossettid in m3. Given the difference between the vertical and horizontal rami of these two specimens, having accessory fossettids could reflect a variable character in Palyeidodon.

Postcranium

Among the abundant isolated postcranial elements from the Aisol Formation (Forasiepi et al. 2011), a distal fragment of the left femur (MHNSR-PV 122) and three ulnae (MHNSR-PV 1008, MHNSR-PV 1009, and MHNSR-PV 1010) are tentatively assigned to the new species.

The femur is massive and anteroposteriorly thick (Fig. 8a, b). The trochlea is shallow, irregular in shape, with the medial border taller than the lateral. A blunt
vertical crest develops on the diaphysis from the medial border of the trochlea. The lateral epicondyle is more robust and protruded than the medial epicondyle. In posterior view, the medial condyle is narrower, taller, and more posteriorly projected than the lateral condyle. The intertrochanteric fossa is narrow and very deep.

The ulna is robust and slightly posteriorly curved (Fig. 8c–e). MHNSR-PV 1009 is smaller and more slender than the other two, probably due to ontogenetic differences. It lacks the distal epiphysis, indicating that it was not completely ossified, therefore corresponding to a juvenile. The olecranon is robust, laterally concave, and long. Its proximal surface has deep scars, suggesting a strong triceps musculature. The articular area for the humerus is deep and surrounded by protruding anconeal and medial coronoid processes. The radial facet is concave and medially faced. The diaphysis is lateromedially compressed in MHNSR-PV 1008 and MHNSR-PV 1009, whereas it is more circular in section in MHNSR-PV 1010 and strengthened by vertical crests on its medial and lateral sides. The distal epiphysis is preserved only in MHNSR-PV 1010 (Fig. 8c). The styloid process is distolaterally projected and has a convex articular facet.

Bone histology

The three ulnae were sampled for histological analysis (Fig. 9a–d). The bone samples are characterized by a compact cortex and a dense network of fine trabeculae in the medullary region. The tissue of the cortex is well vascularized by mostly longitudinally oriented vascular canals. Also, very few irregularly oriented canals can be found. The vascular canals are arranged randomly. Osteocyte lacunae are large and very abundant (Fig. 9a). The
cortices of the three ulnae are all heavily remodeled. Areas of primary bone matrix are visible between secondary osteons, which suggest a mostly parallel-fibered to lamellar organization of the tissue with elongated and flattened osteocyte lacunae (Fig. 9b). Bundles of fibers are arranged randomly. Localized areas of woven bone characterized by round osteocyte lacunae are also found. The most external layer of the cortex with parallel fibers is partially preserved in MHNSR-PV 1008 and MHNSR-PV 1010. It contains only very few secondary osteons and clearly contrasts with the heavily remodeled inner cortex (Fig. 9b). The anterior portion of the cortex of MHNSR-PV 1008 additionally shows some laminar organization, and four circumferential lines of parallel-fibered bone tissue can be made out in the remodeled primary matrix (Fig. 9c, d). These lines do not represent lines of arrested growth (LAGs). The layer of bone lining the medullary cavity and the trabeculae show lamellar tissue organization.

Discussion
Polanski (Dessanti 1954, 1956; González Díaz 1972) identified Nesodon imbricatus in the deposits of the Aisol Formation but, unfortunately, the material was not figured and is now missing from the collections (Soria 1983). The preceding comparative description makes it evident that MHNSR-PV 1004 presents clear similarities with the genus Nesodon. MHNSR-PV 1004 is conspicuously larger than Adinotherium (see below; Fig. 10) with a very shallow or an absent angled posteroventral border of the mandible. It differs from Palyeidodon in an unreduced sagittal crest, the presence of a postfossette in M3, the isolation of primary lingual fold as fossette in later wear stages in M2 and M3, and the presence of accessory fossettids in lower molars (revised diagnosis of Nesodon in Croft et al. 2004). There are some noticeable cranial differences with respect to N. imbricatus and other Nesodon species (e.g., N.
Table 1 Measurements of *Nesodon taweretus* sp. nov. from holotype and tentatively referred material from the Aisol Formation. The measurements used for body mass estimation (ESM 2) are shown with abbreviations in parentheses.

| Skull | MHNSR-PV 1004 |
|-------|---------------|
| Total skull length (TSL) | 440 |
| Posterior skull length (PSL = SC) | 187 |
| Premaxillary height, middle line | ~ 53.5 |
| Muzzle (premaxilla–nasal) height | ~ 129 |
| Muzzle width (MZW) | 60.5 |
| Palatal width, level of P1 | 36 |
| Palatal width, level of M1 | 63 |
| Palatal width, level of M2 (PAW) | 75 |
| Palatal width, level of M3 | 76.2 |
| Frontal width at postorbital processes | 141.7 |
| Depth of the face under the orbit (SD) | 110 |
| Anterior nasal width | 74.2 |
| Premaxilla-orbit length | 210 |
| Nasal aperture-nuchal crest length | 354.5 |
| Nasal aperture-orbit length | 167 |
| Length of the ridge for the origin of masseter (SA) | 129 |
| Occipital width | ~ 211.5 |
| Occipital height (OCH = SB) | 138 |
| Length of the paraoccipital process (SE) | 80 |
| Nuchal crest–foramen magnum height | 88.5 |
| Foramen magnum | 26 × 42 |
| Basicranial length (BL) | 175.5 |
| Tip of nasal-occipital crest length | 343.2 |
| Sagittal crest length | 55.5 |
| Zygomatic arch maximum height | 82 |
| Zygomatic arch posterior height | 47.9 |
| Length M1–M3 | 125.7 |
| Length I1–M3 (broken area of I1 alveoli) | ~ 266.3 |
| Length P1–M3 (alveoli P) | ~ 199.4 |
| Length P1–P4 (alveoli) | ~ 79.2 |

| Mandible | MHNSR-PV 360 | MHNSR-PV 1075 | MHNSR-PV 1000 | MHNSR-PV 1001 |
|-----------|--------------|--------------|--------------|--------------|
| Total jaw length (TJL) | ~ 340 | – | ~ 360 | – |
| Symphyseal length | 66.0 | – | – | – |
| Posterior jaw length (JMA) | 107.5 | – | – | – |
| Height of the horizontal ramus at the level of p4 (labial view) | 65.0 | 65.0 | 75.0 | – |
| Height of the horizontal ramus at the level of m3 (labial view) | 99.0 | 106 | 104 | 132 |
| Mandible width at the level of p4 | 23.0 | 29.0 | 30.00 | – |
| Mandible width at the level of m3 | 29.5 | 28.0 | 36.0 | 32.0 |
| Height from ventral border of the horizontal ramus to condyle | ~ 36 | 50.0 | – | – |
| Width of the condyle | – | – | – | – |
| Lower molar tooth row length | 119.8 | 132.2 | 136.4 | – |

| Upper dentition | MHNSR PV 1004 |
|-----------------|---------------|
| M1 L | 35.2 |
| M1 W | 28.2 |
| M2 L | 42.6 |
conspurcatus), which are interpreted as being of specific significance.

To compare Nesodon taweretus sp. nov. with other nesodontines, measurements of the lower dentition of specimens MHNSR-PV 360 and PV 1000 were added to the bivariate plot presented by Croft et al. (2004: fig. 3) and reproduced in Fig. 10. It shows that the m1 of MHNSR-PV 1000 exceeds the size of N. imbricatus, while that of MHNSR-PV 360 falls within the range of this species; a similar situation occurs with m2. The m3 of MHNSR-PV 360 is wider than the specimens of N. imbricatus, and the other m3 has a similar size to the rest of the sample. Therefore, dental measurements of N. taweretus sp. nov are similar to those of N. imbricatus, but they are clearly larger than those of N. conspurcatus and Adinotherium ovinum.

The body mass of Nesodon taweretus sp. nov. was estimated as about 550 kg, using the average of the results obtained using different algorithms (4.1, 4.2, 4.3, and 5.1 from Mendoza et al. 2006) and equations (TSL, OCH, SLML, PSL, and TJL from Janis 1990) following Cassini et al. (2012, tables 14.1 and 14.2; ESM 2). The weighted

| Table 1 continued |
|-------------------|-------------------|
| **Upper dentition** | MHNSR PV 1004 |
| M2 L | 33.9 |
| M3 L | 60.0 |
| M3 W | 37.7 |

| Lower dentition | MHNSR-PV 1000 (right side) | MHNSR-PV 1000 (left side) | MHNSR-PV 360 | MHNSR-PV 1075 | MHNSR-PV 1001 |
|-----------------|-----------------------------|-----------------------------|---------------|---------------|---------------|
| p4 L | ~35 | ~35 | ~23 | 25 | – |
| p4 W | ~18 alveolar size | ~18 alveolar size | – | – | – |
| m1 L | 33.0 | ~31 | 28.0 | 33.0 | – |
| m1 Wa | 18.0 | – | 15.0 | – | – |
| m1 Wp | 18.6 | – | 16.0 | 17.0 | – |
| m2 L | ~34 | 40.0 | 36.0 | 40.0 | – |
| m2 Wa | – | – | 15.6 | 17.3 | – |
| m2 Wp | – | 18.0 | 15.5 | 17.0 | – |
| m3 L | 68.0 | ~66 | 51.0 | 50.0 | 61.0 |
| m3 Wa | 20.0 | 21.0 | 16.0 | 16.0 | 19.6 |
| m3 Wp | 15.7 | 16.5 | 12.7 | 12.8 | 16.0 |

**Fig. 10** Bivariate plot of lower molar dimensions of *Nesodon* and *Adinotherium* (modified from Croft et al. 2004)
mean of Janis’s equations (which includes the percent prediction error in the estimation) provides a slightly larger value of about 570 kg. Both estimations yield similar values. The estimated body mass for *N. taweretus* sp. nov. is similar to that for *N. imbricatus* (around 640 kg; range between 500 and 800 kg; Cassini et al. 2012).

Studies of bone histology are infrequent for notoungulates, and this study contains detailed information and analysis for further work and comparison. The histology of the ulnae tentatively assigned to *N. taweretus* sp. nov. is in accordance with the structure described for *Toxodon* and *Mesotherium* (as *Typotherium*; de Ricqlès et al. 2009) and more generally for eutherian mammals (Hurum and Chinsamy-Turan 2012), and suggests a decrease in growth rate during ontogeny. Remodeling is a great factor in *N. taweretus* sp. nov., which is a common feature in mammalian bones (Enlow and Brown 1958)—particularly in large taxa, probably due to increased loading (Straehl et al. 2013).

**Phylogenetic analysis**

A parsimony analysis was performed based on 25 taxa and 59 morphological cranial and dental characters (ESM 2) modified from the matrix presented by Nassif et al. (2000). Taxa included four outgroups (two Notohippidae and two Leontiniidae; alternatively see Cerdeño and Vera 2010 for the phylogenetic support for these families) and 21 Toxodontidae.

Some codifications were modified from the contribution of Nassif et al. (2000). A low median symphyseal keel is present in *N. imbricatus* and *N. taweretus* sp. nov. [character 14 (1)]. A new revision of the material referred to *Trigodon* indicates that the lingual enamel in m1 is placed between the anterior fold and the hypoconulid, without covering the lingual face of the hypoconulid completely [character 54 (1)]. The specimen MHNSR-PV 1075 here assigned to *Palyeidodon* reveals variation in the presence of accessory fossettids in lower molars [character 46 (0 and 1)]. New studies in *Dinotoxodon* (Schmidt 2013) allowed us to complete the codifications for the characters 27, 30, 43–48, and 54–56 and to modify others as follows: the horizontal ramus may or may not have a ventral extension [character 8 (0 and 1)]; the upper molars have a median cristae and an incipient Y-shaped median valley [character 29 (0)]; and the lingual enamel on M3 covers only the posterior groove [character 32 (2)].

The data matrix (ESM 2) was analyzed using maximum parsimony with equally weighted characters and under implied weighting with the computer program TNT 1.1

![Phylogenetic relationships of Toxodontidae derived from the analysis performed under implied weights (K = 3)](image-url)
(Goloboff et al. 2008b). All characters were treated as non-additive. Equally weighted parsimony analysis was conducted performing a heuristic search of Wagner trees with 500 random addition sequences, followed by TBR (tree bisection reconnection), and saving 20 trees per round, which improves the searches and ensures that all optimal trees are found. The program provided the 16 most parsimonious topologies of 183 steps, with a consistency index (CI) of 0.46 and a retention index (RI) of 0.65. The strict consensus and Bremer index of each node are shown in ESM 2.

Analysis performed under implied weighting ($K = 3$) provided one tree (Fig. 11). This method uses evidence of homoplasy to estimate character reliability and finds trees with the maximum total fit (optimal topologies) (Goloboff 1993; Goloboff et al. 2008a). The trees constructed under implied weighting have the most reliable characters (Goloboff 1993). The topologies of the consensus from the analysis under equally weighted characters and the tree obtained under implied weights were in general agreement. Monophyly of the major groups (Toxodontidae, Nesodontinae, Toxodontinae) is supported in both analyses. Differences are in the positions of some toxodontines and consequently the re-arrangement of some parts of the tree. The clade comprising Calchaquitherium, Nonotherium, and Xotoodon that is found with equally weighted characters (ESM 2), as in the analysis by Nassif et al. (2000), is reorganized under implied weighting, mainly due to the different positions of Calchaquitherium (Fig. 11). We base the discussion on the tree topology and the supporting node characters with the tree obtained under implied weighting because it is better resolved and the characters have been demonstrated to be more reliable than equally weighted characters are (Goloboff 1993; Goloboff et al. 2008a). The tree obtained (Fig. 11) is compared with previous phylogenetic results (Nassif et al. 2000).

The monophyly of the Toxodontidae is supported by two synapomorphies (node 1): i1 triangular in section [35 (1)], and the enamel on the lingual side of i3 being narrower than on the labial side [39 (3)]. The nesodontines consist of Nesodon and Adinotherium, with Proadinothere- rium as the sister group of the remaining toxodontids; node 2 is supported by any synapomorphy. In node 3, Adinotherium and Nesodon share a sigmoid zygomatic arch [6 (1)] and symphysis without a well-differentiated chin angle [15 (0)]. Nesodon taveretus sp. nov. forms a monophyletic group (node 4) with N. imbricatus sharing a rounded postero-ventral border of the vertical ramus [11 (0)] and the symphysis with a median labial keel [14 (1)]. This result supports the generic determination of the new species. Node 5 is supported by eight synapomorphies: [3 (1)], [28 (1)], [30 (1)], [31 (1)], [39 (1)], [54 (1)], [55 (1)], and [56 (1)], and includes all toxodontines (sensu Nassif et al. 2000). The first divergence corresponds to Paley eidodon, followed by Hyperoxotodon (node 6). Node 7 groups the remaining taxa with three synapomorphies: a triangular and buccolingually flattened cross-section of I1 [19 (4)], P2 without groove or fossette [24 (1)], and an absence of lingual enamel in i1–i2 [38 (2)]. This group splits into two clades. The first is formed by Nonotherium and Xotoodon (node 8), which share a sigmoid zygomatic arch [6 (1)], P3–P4 without a groove or fossette [26 (1)], and very concave ectoloph of the molars [33 (1)]. The second, major clade (node 9) is supported by two synapomorphies: p1 absent [42 (2)] and reduced lingual enamel of p2–p4 [43 (1)], and gathers two large groups. One group is formed by [Andinotoxodon ((Dinotoxodon Toxodon) (Gyrinodon (Ocnerotherium Hoffstetterius)))] and is supported by two synapomorphies (node 10): alveolar border of the symphysis straighter [13 (1)] and anterior fold (paracoonid-metaconid) of m1–m2 at the same level as the buccal fold [48 (1)]. The other clade (node 15) is composed of seven taxa which share intraorbital foramen close to the zygomatic apophyses of the maxilla [5 (0)] and coronoid apophyses and condyles convergent [10 (1)]. Within this large clade, Posnanskytherium, Pisanodon, and Pericot otoxodon appear as successive basal taxa (nodes 15, 16, and 17, respectively), and as sister taxa of two small monophyletic groups supported by three synapomorphies (node 18: [40 (1)], [42 (1)], [58 (1)]). One group is composed of Calchaquitherium and Mixotoxodon (node 19: [41 (0)], [43 (2)], [47 (0)], [52 (1)] and the other by Parartragodon and Trigodon (node 20: [32 (1)] and [49 (1)]).

The topology obtained (Fig. 11) differs from that in Nassif et al. (2000) mainly in the arrangement of the Toxodontinae (node 7). Some clades such as (Nonotherium and Xotoodon), (Ocnerotherium and Hoffstetterius), and (Parartragodon and Trigodon) are recovered in both analyses but their positions are different within the cladogram. Moreover, Dinotoxodon appears as sister taxon of Toxodon (Fig. 11, node 12), sharing a well-developed anterior fold in m1–m2 ([47 (0)]), in contrast to the results in Nassif et al. (2000), where Dinotoxodon is related to Pericotoxodon, and Toxodon to Andinotoxodon.

Final remarks

The vertebrate association from the Aisol Formation includes anurans, turtles, carnivorous birds, and a large set of mammals (Soria 1983; Forasiepi et al. 2011). These preliminary studies did not provide precise taxonomic determinations, and initially suggested affinities with associations from the Middle Miocene of Patagonia (Soria 1983; Forasiepi et al. 2011). Among toxodontids, Paley eidodon cf. P. obtusum would relate the Aisol Formation to the Collón Curá and Río Frías formations (Roth 1899;
Pascual 1954; Madden 1990). The presence of cf. *Pal-yeidodon* was suggested for the extrapatagonian Santacru-
cian locality of Chucal, Chile (Bond and García 2002),
which is older than 17.5 Ma (Croft et al. 2007), based on
poorly preserved material. If this find is correct, the pre-

cence of *Palyeidodon* in late Early Miocene outcrops may
not be surprising, but this assignation was questioned
(Croft et al. 2004, 2009) and should be considered quite
tentative. The recognition of the genus *Nesodon* suggests
that the vertebrate association from Aisol is no older than
Santacrucian, although the new species *N. taweretus* diff-

erentiates the Aisol assemblage from the other Santacruc-
cian faunas. Other mammals from Aisol are not conclusive
in determining the age.

U–Pb zircon dating of the tuff sample from the base of the
Aisol Formation suggests a late Early Miocene age of
19.480 ± 0.025 Ma (Burdigalian). A similar accuracy has
not yet been established for the top of the sequence. Tec-
tosedimentary evidence from the Andean orogenic basins
from south-central Argentina suggests that the sequence may
extend to the Langhian, but not younger than that. During
the late Burdigalian and early Langhian, a major tectonic
event occurred, with a consequent strong angular uncon-
formity in the sedimentary sequences, associated with
intense volcanism (Garrido et al. 2012 and literature cited
therein). We do not find evidence of this sedimentary hiatus
within the Aisol Formation, but a continuous succession
including the U–Pb dated tuffs. Consequently, we associate
the vertebrate assemblage with the late Early Miocene.

The Santacrucian Age covers the time span between
~18 and 16 Ma and typically includes outcrops of the
Santa Cruz Formation in the eastern region of the Santa
Cruz Province on the Atlantic coastal plain (Vizcaïno et al.
2012; Perkins et al. 2012). Slightly older vertebrate asso-
ciations from the Adean foothills (e.g., Karaike) dated
~19–14 Ma (Perkins et al. 2012) are possibly included in
the Santacrucian (see discussion in Vizcaïno et al. 2012
and referenced literature). At intermediate South American
latitudes, the Santacrucian Age may extend to earlier times
considering the mammalian association from Chucal (north
Chile) dated at ~19–17 Ma (Croft et al. 2007). Assuming
that the absolute dating of ca. 19.5 Ma is correct for the
base of the Aisol Formation and that the vertebrate asso-
ciation corresponds to the Santacrucian Age, paleoenvi-
rmental conditions could explain the taxonomic


distinction between the Aisol and Santa Cruz formations in
southern Patagonia. The large latitudinal separation
between both areas would have impacted the different

evolutions of the faunas. This hypothesis has already been
proposed for the early Oligocene assemblage from Tingu-
iririca at a similar latitude in Chile (Croft et al. 2008)
and the late Oligocene assemblage from Quebrada Fiera in
southern Mendoza (Cerdeño et al. 2010; Cerdeño 2011).

Similarly, the Santacrucian assemblage from Chinches
(San Juan Province; López et al. 2011) and the Santacruc-
cian assemblage from Chucal (north Chile; Flynn et al.
2002; Croft et al. 2004) are distinguishable from their
Patagonian equivalents. Furthermore, analysis of Middle
Miocene fossil associations from middle latitudes of
Bolivia (Croft et al. 2009) has already suggested an evident
latitudinal provincialism that could have begun to develop
by at least the Early Miocene, deriving from the Andean
orogeny and other paleoenvironmental factors (Croft et al.
2004).

Conclusions

The present study proposes the existence of a new tox-
odontid notoungulate in the Aisol Formation (Mendoza,
Argentina), recognized as *Nesodon* sp. nov.

*Nesodon taweretus* sp. nov. differs from the other
*Neosodon* species in several cranial and dental features.
Close comparisons were made with the Patagonian *N. im-

bricatus*, a common toxodontid in the Santa Cruz Forma-
tion (Santacrucian Age). *N. taweretus* was similar in size to
*N. imbricatus*, with a body mass of about 550 kg. Phylo-
genetic analysis supports the assignation of the new species
to the genus *Nesodon*.

A tuff sample from the base of the Aisol Formation pro-
vided a high-precision U–Pb zircon age of 19.480 ±
0.025 Ma, which places the base of the unit in the late Early
Miocene—Burdigalian. The vertebrate association would be
included in the Santacrucian Age. Latitudinal separation could
explain taxonomic differences, as exhibited by the species of
*Nesodon* found in the Mendoza and Santa Cruz areas.

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