



Inferring habitat and feeding behaviour of early Miocene notoungulates from Patagonia

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Notoungulates, native fossil mammals of South America, have been usually studied from a taxonomic point of view, whereas their palaeobiology has been largely neglected. For example, morpho-functional or eco-morphological approaches have not been rigorously applied to the masticatory apparatus to propose hypothesis on dietary habits. In this study, we generate inferences about habitat and feeding preferences in five Santacrucian genera of notoungulates of the orders Typotheria and Toxodontia using novel computer techniques of knowledge discovery. The Santacrucian (Santa Cruz Formation, late-early Miocene) fauna is particularly appropriate for this kind of studies due to its taxonomic richness, diversity, amount of specimens recorded and the quality of preservation. Over 100 extant species of ungulates, distributed among 13 families of artiodactyls and perisodactyls, were used as reference samples to reveal the relationships between craniodental morphology and ecological patterns. The results suggest that all Santacrucian notoungulates present morphologies characteristic of open habitats' extant ungulates. Although the Toxodontia exhibits the same morphological pattern of living mixed-feeders and grazers, the Typotheria shows exaggerated traits of specialized grazer ungulates. □ *Craniodental morphology, ecomorphology, fossil ungulates, knowledge discovery, South America.*

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Notoungulata constitutes the most abundant and diverse clade of endemic South American ungulates, both taxonomically and morphologically (Simpson 1936; Patterson & Pascual 1972; Cifelli 1993). Notoungulates are recorded throughout the Cenozoic, from the late Palaeocene to the late Pleistocene (Bondesio 1986; Croft 1999). The group reaches its greatest taxonomic richness (at the genus level) during the Palaeogene, gradually diminishing throughout the Neogene with forms that become progressively more specialized, until their extinction by the late Pleistocene–early Holocene (see Bond *et al.* 1995).

Following Billet (2010), the clade comprises two monophyletic groups: Toxodontia and Typotheria – plus the basal notoungulate family Henricosborniidae, and a clade comprising *Notostylops* as the sister group of Pyrotheria (before considered as a separate order). Toxodonts include large (i.e. above 44 kg, *sensu* Martin & Steadman 1999) to very large animals (including strictly megamammals, i.e. 1000 kg or more), and are sometimes compared with hippos or rhinos, due to

their inferred general appearance and the grinding pattern of molar crowns (Ameghino 1907; Scott 1912; Bond 1999). Typotheres are small-to-medium size mammals, mostly described as rodent-like in overall form, although different families resemble living capybaras (Mesotheriidae), hares (Hegetotheriidae), hyraxes (Intertheriidae) and small deer (Ameghino 1889; Sinclair 1909; Bond *et al.* 1995; Croft 1999; Reguero *et al.* 2007).

In both toxodonts and typotheres, there is an apparent tendency to evolve from generalized masticatory apparatus with complete dentition, without diastema and brachyodont cheek teeth, to very specialized forms including, for instance, hypertrophied incisors, simplified crown patterns and ever-growing (hypsodont) cheek teeth (Ameghino 1887, 1894; Scott 1937; Simpson 1967; Cifelli 1985). Some phylogenetic studies suggest that hypselodonty probably evolved at least four times within the Notoungulata – in the Toxodontidae, Intertheriidae, Mesotheriidae and Hegetotheriidae (Croft & Weinstein 2008). The

palaeobiological implications of this morphological diversity and the relationship between form and function have not been analysed rigorously before, and the dietary inferences have been mostly limited to broad generalizations.

The Santacrucian mammalian fauna (Santa Cruz Formation, late–early Miocene) along the Atlantic coast of southern Patagonia is particularly appropriate for this kind of study due to its taxonomic richness, diversity, amount of specimens recorded and quality of preservation (see Vizcaíno et al. 2006, 2010; and references therein). Among notoungulates, the Toxodontia are represented by three genera: *Nesodon*, *Adinotherium* (Toxodontidae, Nesodontinae; Fig. 1A, B) and *Homalodotherium* (Homalodotheriidae). Calculating body mass using the mean of output of the allometric equations for ‘all ungulates’ proposed by Janis (1990a), *Homalodotherium* and *Nesodon* are among the largest ungulates from the

Santa Cruz beds (~350 and 550 kg respectively), only being surpassed by the astrapothere *Astrapotherium* (~1000 kg), whereas *Adinotherium* is rather smaller (~120 kg). *Nesodon*, *Adinotherium* and *Homalodotherium* species have complete dentition without diastema and large masseteric and temporal muscles attachment areas. *Nesodon* and *Adinotherium* have hypertrophied lateral incisors and high-crowned of finite growth cheek teeth (protohypsodonty, *sensu* Mones 1982). The Santacrucian Typotheria are small-sized animals represented by *Protypotherium*, *Interatherium* (Interatheriidae; Fig. 1C, D), *Hegetotherium* and *Pachyrukhos* (Hegetotheriidae; Fig. 1E, F). Calculating body masses as described above, *Hegetotherium* and *Pachyrukhos* are respectively the largest (~14 kg) and the smallest (~2.8 kg) examples. Among interatheriids, *Protypotherium* species (~4.5 kg for the smaller *Protypotherium attenuatum* and approximately 8.5 kg for the largest *Protypotherium*

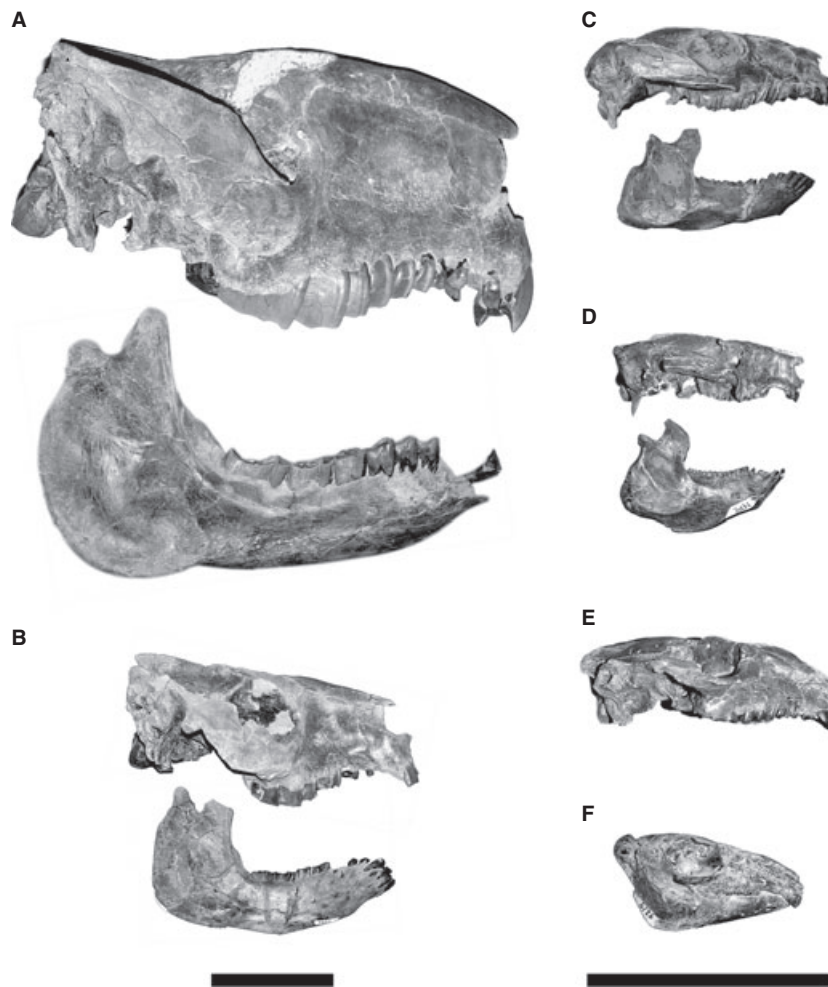


Fig. 1. Skull and mandible of Santacrucian notoungulates genera included in this study. Toxodontids: A, *Nesodon* MPM-PV 3659; B, *Adinotherium* MPM-PV 3532 and 3666. Typotheres: C, *Protypotherium* AMNH 9868; D, *Interatherium* MPM-PV 3471; E, *Hegetotherium* MPM-PV 3526; and F, *Pachyrukhos* AMNH 9219. Scale bar 10 cm.

australe) are larger than *Interatherium* (~3.5 kg). Interatheriids include species with complete dentition and without diastema or hypertrophied incisors. Thus, their incisors are all the same width. In hegetotheriids, the upper and lower central incisors are hypertrophied, and the lateral incisors and canines are atrophied, resulting in a functional diastema in *Hegetotherium*, and an anatomical diastema with missing teeth in *Pachyrukhos*. In all these taxa, cheek teeth are high-crowned and ever-growing (eu-hypsodonty, *sensu* Mones 1982), and the masseteric muscle areas of attachment are larger than the temporal ones.

All genera analysed in this study have been traditionally considered mostly herbivores. Following Scott (1937), and based primarily on the high-crowned cheek teeth, they were considered inhabitants of open plains and other open-habitat, eating mostly grasses (Patterson & Pascual 1968; Cifelli 1985; Billet *et al.* 2009). Bond (1986) and Tauber (1997b) also supported the grazers hypothesis based on the degree of hypsodonty and incisor morphology. With reference to *Nesodon* and *Adinotherium*, Madden (1997) proposed that the most complex molar crowns would have provided great shearing ability among toxodontids allowing them to break down the grasses more easily. However, Townsend & Croft (2008) using a micro-wear approach for three genera of Santacrucian notoungulates disagreed the preceding dietary inferences. They conclude that *Nesodon imbricatus* was a leaf browser that focused more on hard browsing, including, potentially, bark consumption; *Adinotherium ovinum* was a pure leaf browser; and *Protypotherium* was a traditional browser that took both soft browse and soft fruits. Since Townsend & Croft (2008), no other ecomorphological studies on cranio-mandibular morphology have been carried out in this group.

Contrarily, the relationship between habitat preference, diet and craniodental morphology in living ungulates has been intensively studied using different statistical approaches (Gordon & Illius 1988; Solounias & Dawson-Saunders 1988; Janis 1990b; Spencer 1995; Pérez-Barbería & Gordon 1999; Mendoza *et al.* 2002). Mendoza & Palmqvist (2008) show that novel machine-learning techniques of knowledge discovery (i.e. decision trees) were powerful tools for ecomorphological approaches. In this study, we apply the same techniques of knowledge discovery to identify new morphological patterns and infer the habitat and feeding behaviour of the notoungulates from the Santacrucian mammal assemblage. In doing so, species from two orders of living ungulates, Artiodactyla and Perissodactyla, were used as a reference. We assumed then that the morphological solutions present in both

Artiodactyls and Perissodactyls were also achieved by notoungulates.

The Principle of Actualism is applied, assuming that some biomechanical laws have constrained in the same manner the evolution of the craniodental structure of these three taxonomic groups, leading to the convergence of common morphological patterns. However, when the phylogenetic legacy is different, and the starting morphology varies, the same biomechanical laws can be upheld with different strategies, associated to different morphological patterns.

Material and methods

Abbreviations

AMNH: American Museum of Natural History, New York, USA; MACN: Museo Argentino de Ciencias Naturales 'Bernardino Rivadavia', Buenos Aires, Argentina; MLP: Museo de La Plata, La Plata, Argentina; MPM-PV: Museo Regional Provincial Padre 'M. J. Molina', Río Gallegos, Argentina; YPM-PU: Yale Peabody Museum, New Haven, USA.

Data

Forty-seven specimens of notoungulates from the Santa Cruz Formation (late-early Miocene) were analysed. Only specimens with no or little apparent deformation were measured. They include 20 toxodontids (13 specimens of *Nesodon* and seven *Adinotherium*), and 27 tyotheres (10 specimens of *Protypotherium*, 11 *Interatherium*, 3 *Hegetotherium* and 3 *Pachyrukhos*) (see Table 1).

One-hundred and nineteen extant species of artiodactyls and perissodactyls, distributed among 13 families, taken from Mendoza *et al.* (2002), were used to analyse the relationship between the craniodental morphology of ungulates and their ecological adaptations (i.e. habitat and feeding behaviour).

Fourteen craniodental measurements, and the hypsodonty index (HI) were used in the analyses (Fig. 2). The length of diastema (JLB) was measured from the base of the third incisor to the first premolar (see Mendoza *et al.* 2002, table 2) in extant ungulates and the only notoungulate with diastema *Pachyrukhos*. The total length of the jaw (JAW) was obtained summing JLB (when present), JMA, LMRL and LPRL (Fig. 2).

The HI of notoungulates was achieved following Janis (1988), i.e. crown height ratio of m3 calculated as the crown height (taken from RX-radiography) divided by the labio-lingual width of the tooth. For the HI, only the well-preserved and complete

Table 1. List of Notoungulata skulls examined.

Toxodontia	
Toxodontidae	
<i>Adinotherium</i>	AMNH: 9141; 9517; MACN-SC: 4355; MPM-PV: 3667; 3668 YPM-PU: 15003; 15136
<i>Nesodon</i>	AMNH: 9128; 9168; 9192; 9510; MACN-A: 775; 5145; MLP: 12-250 YPM-PU: 15000; 15215; 15256; 15260; 15336; 15492
Typotheria	
Interatheriidae	
<i>Interatherium</i>	AMNH: 9483; MPM-PV: 3469; 3471; 3527; 3528 YPM-PU: 15100; 15296; 15300; 15401; 15554
<i>Protypotherium</i>	AMNH: 9187; 9565; MACN-A: 3920; 3991 MPM-PV: 3470; 3530; 3531; 3661; YPM-PU: 15386; 15828
Hegetotheriidae	
<i>Hegetotherium</i>	AMNH: 9159; MPM-PV: 3481; YPM-PU: 15298; 15542
<i>Pachyrhokos</i>	AMNH: 9219; 9283; YPM-PU: 15744

mandibles were used, which correspond to 17 MPM-PV specimens. Janis (1988) procedure implies measuring fully erupted but still unworn m3. However, in the fossil record, it is highly improbable to find specimens with a tooth crown fully formed but unworn, making nearly impossible to examine variation in crown height between unworn and worn m3. Therefore, we assumed that, in euhypsodont notoungulates, the HI does not vary significantly during the lifespan of m3, being comparable with the HI from Janis (1988).

The HI is a size-independent variable, but the other measurements used were size-adjusted by dividing each of them by the lower molar tooth row length, measured along the base of the teeth (see details in Mendoza & Palmqvist 2006).

Analyses

Two computer techniques of knowledge discovery were used: (1) discriminant analysis; and (2) classification trees. Discriminant analysis is a classical technique of multivariate statistics that allows classifying new samples within predefined groups using the discriminant functions adjusted for maximizing the between-groups to within-groups ratio of variance. In this contribution, we mainly use discriminant analysis as a technique of knowledge discovery (Cios *et al.* 1998). Classification trees, developed by Breiman *et al.* (1984), are a knowledge discovery technique (Larose 2004) that stems from the realm of machine learning (see Michie *et al.* 1994). Classification trees are non-parametric models. Some of their advantages are that they look at variables hierarchically rather than simultaneously, they are easy to interpret and they do not assume that the dependent variable follows any given distribution.

In this study, we follow the approach applied by Mendoza (2007) and Mendoza & Palmqvist (2008), with the goal of better understanding the relationship

between the skeleton morphology and their ecological adaptations in living ungulates. The 'discovered' knowledge is then used to infer the habitat and feeding behaviour of extinct ungulates not closely related to living forms. In the extinct taxa, the morphological evidence is restricted to the measurements available on the specimens; therefore, depending on the variables related to each ecological pattern, different specimens could be analysed.

For the first set of analyses, performed to characterize types of habitat, species were classified among the following three categories (Mendoza *et al.* 2005): (1) open habitats (i.e. treeless or scarcely wooded savannas, grasslands, dry deserts and semi-desert steppes); (2) mixed habitats (i.e. wooded savannas, bush land, open forests and species dwelling both in closed and open habitats); and (3) closed habitats (i.e. closed woodlands, riverine, moist deciduous and evergreen forests). Fifty-eight of the 119 living ungulate species included in this study were categorized as open habitats, 30 as mixed habitats and 31 as closed forested habitats. Mendoza & Palmqvist (2008) interpreted JLB to be the best measurement of the jaw, together with HI, for discriminating species from open and mixed habitats. As mentioned above, Santacrucian notoungulates do not have diastema (JLB). Hence, the other variables that also contribute to the length of the jaw, as well as the total length of the jaw, were analysed to evaluate whether they could be used instead of JLB.

As all Santacrucian notoungulates studied were characterized as preferring open or mixed habitats (see Results), we carried a second set of analyses in which extant species from closed habitats were not included. The remaining 88 species from open and mixed habitats were further subdivided according to their feeding behaviour as grazers or non-grazers. Only 31 living species that consume a high percentage of grass, which does not highly fluctuate through the year or between different localities, were classified as

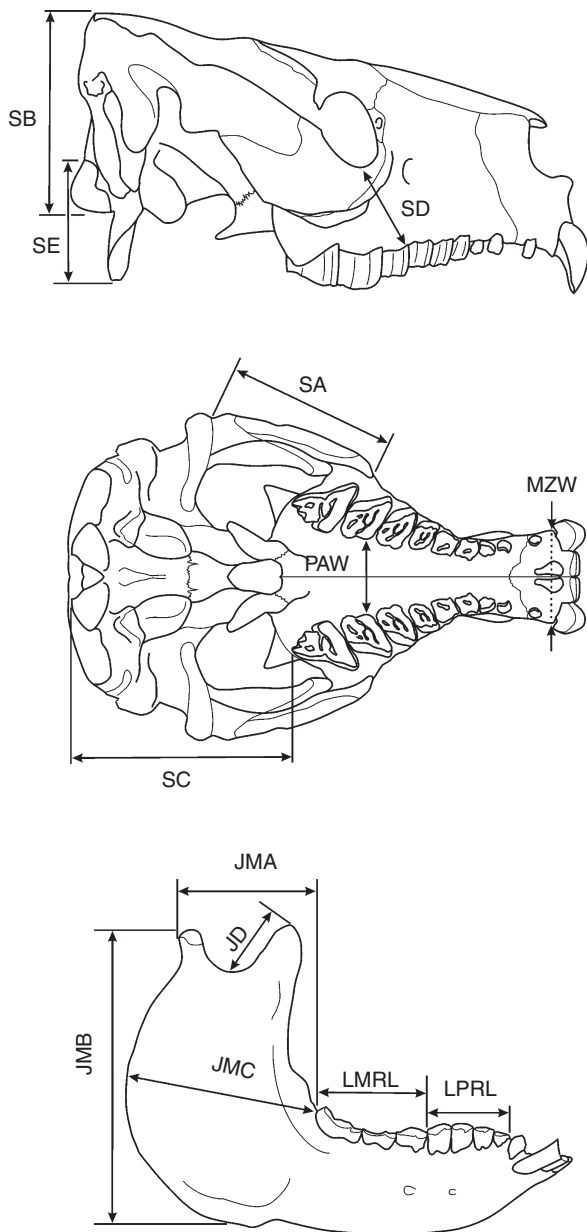


Fig. 2. Skull of *Adinotherium* with selected measurements used in this study. SA, length of the masseteric fossae; SB, occipital height; SC, length of the posterior portion of the skull; SD, depth of the face under the orbit; SE, length of the paraoccipital process; MZW, muzzle width; PAW, palatal width; JMA, posterior jaw length; JMB, depth of mandibular angle; JMC, maximum width of the mandibular angle; JD, length of the coronoid process; LMRL, lower molar tooth row length; LPRL, lower premolar tooth row length (see details in Mendoza *et al.* 2002).

grazers. Among the remaining species, 52 are mixed feeders that consume grass and dicotyledonous plants depending on their availability and five species mainly browse, from which three dwell in open habitats, in spite of being browsers.

The probability of obtaining a good discrimination between the groups compared merely by chance increases with the number of variables involved.

Therefore, only patterns that were found consistently based on limited combinations of variables were considered here.

Starting from 14 measurements, there are 455 possible combinations involving three or fewer of them. Knowledge discovery techniques were used then to identify those combinations of three or less variables that allow a better characterization of the ecological groups. The region occupied by each group in the space defined by a combination of few variables, subspaces of the full morphospace defined by whole 14 variables, constitute the morphological patterns that characterize each group (Mendoza 2007). The position of each extinct species in these subspaces, optimized for the task of group characterization, was carefully analysed to infer their habitat and feeding behaviour.

Results

Inferences about habitat

From the whole variables of the jaw that could be used instead of JLB to discriminate habitat preference, the relative length of the whole jaw (JAW) was the best to characterize species adapted to forage in open and mixed habitats. In the same way that JLB, the HI that allows distinguishing open-habitat species from those of mixed habitats, increases with JAW (see Fig. 3). Consequently, we develop here a Habitat Preference Index ($HPI = HI/1.4 JAW - 2.4$), which combined with HI alone allows the characterization of habitat preference of the extant ungulates.

Mendoza & Palmqvist (2008) showed that living ungulates from closed forested habitats have values of HI lower than 2. We find now that: (1) species dwelling mixed habitats have values of HI between 2 and 4, and values of $HPI < 1$; and (2) species from open habitats also have values of $HI > 2$ and values of $HPI > 1$. There are very few exceptions (see Fig. 3 and Discussion and conclusions section), in which species dwelling open habitats have HPI values lower than 1, six of seven species of *Equus* included in the database were not depicted in Fig. 3 in order to increase its resolution, given that they have very long jaws (5.54–6.12). The three species with the longest jaw have HPI values slightly lower than 1 (0.94–0.97), but they could not be mischaracterized as mixed habitat species because their HI values (5.79–5.83) are much higher than 4, the maximum value observed in the living ungulates dwelling mixed habitats.

Among tpyotheres, *Protypotherium* specimens show a little higher hypsodonty ($HI \sim 4$; Table 2), but most of them are also placed on the left-bottom region of

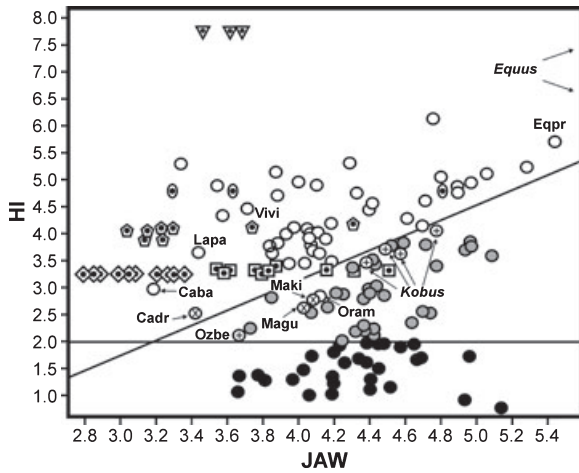


Fig. 3. Distribution of 119 extant ungulates and 26 notoungulates in the morphospace depicted by the hypsodonty index (HI) and the relative length of the jaw (JAW), which allows the characterization of the craniodental morphology of species from open habitats (white symbols), mixed habitats (grey symbols) and closed habitats (black symbols). Notoungulata (dotted symbols): *Interatherium* (squares), *Nesodon* (rhombus), *Protypotherium* (pentagons), *Adinotherium* (ovals) and *Hegetotherium* (inverse triangles); extant ungulates: *Camelus bactrianus* (Caba), *Camelus dromedarius* (Cadr), *Lama pacos* (Lapa), *Lama guanicoe* (Lagu), *Vicugna vicugna* (Vivi) and *Oreamnos americanus* (Oram); water-associated habitat species (symbols with a plus sign): *Ozotoceros bezoarticus* (Ozbe), and four species from the genus *Kobus*; browsers from open habitats (symbols with a cross): *Madoqua guentheri* (Magu) and *Madoqua kirkii* (Maki).

the morphospace depicted by HI and JAW, near camelids (Fig. 3). Two *Protypotherium* specimens, however, have longer jaws, showing a combination of jaw length and hypsodonty more similar to many extant ungulates from open habitats. *Interatherium* specimens have an intermediate hypsodonty (HI ~ 3.3; Table 2) but they show a wide range in the relative length of their jaws, which translates as a wide range of HPI values (0.8–1.3). Most of them show combinations of jaw length and hypsodonty similar to extant ungulates from both open and mixed habitats. Finally, the most hypsodont are two specimens of *Hegetotherium* (HI ~ 7.8; Table 2), comparable with *Equus asinus* (HI ~ 8.7). Their HPI values are very high (2.92 and 3.18), so their HPI are much higher than the threshold that separates open-habitat species from those of mixed habitats. However, there are no living species with such a short jaw combined with so high hypsodonty. Thus, they occupy an exclusive top-left region in the morphospace depicted by HI and JAW.

Among toxodontids, the specimens of *Adinotherium* are very hypsodont (HI ~ 4.9; Table 2), and their jaws length are variable, so their HPI are also very different (1.1, 2.2 and 1.8). All of them, however, show combinations of jaw length and hypsodonty similar to extant ungulates from open habitats. *Nesodon* specimens

Table 2. Summary statistics [mean ± 1 standard deviation (n)] of craniodental measurements of Notoungulata database in millimetres.

	HI	SA	SB	SC	SD	SE	MZW	LPRL	LMRL	JMA	JMB	JMC	JD
<i>Adinotherium</i>	4.92 ±0.42 (2)	105.29 ±6.35 (4)	90.03 ±2.45 (3)	112.64 ±1.29 (4)	52.92 ±4.60 (4)	49.83 ±0.07 (2)	39.20 ±5.56 (3)	37.47 ±10.38 (7)	55.97 ±17.16 (7)	95.72 ±19.70 (3)	115.66 ±26.89 (3)	82.08 ±17.82 (7)	33.70 ±8.42 (4)
<i>Nesodon</i>	3.24 (1)	177.91 ±8.91 (5)	139.75 ±4.54 (3)	195.82 ±10.43 (5)	92.15 ±10.83 (7)	64.92 ±12.13 (5)	70.64 ±5.58 (6)	64.24 ±8.32 (12)	123.53 ±9.45 (13)	170.51 ±14.54 (10)	224.09 ±16.47 (9)	159.87 ±23.82 (13)	55.15 ±6.69 (8)
<i>Interatherium</i>	3.33 ±0.05 (3)	36.96 ±3.66 (5)	22.74 ±2.31 (6)	31.81 ±2.95 (5)	15.10 ±0.79 (6)	13.28 ±2.07 (6)	12.12 ±1.57 (7)	11.72 ±1.20 (10)	15.46 ±1.50 (10)	25.31 ±2.54 (9)	35.32 ±2.54 (9)	29.56 ±2.43 (10)	11.94 ±2.02 (10)
<i>Protypotherium</i>	4.05 ±0.13 (4)	42.41 ±4.51 (6)	26.19 ±4.64 (4)	38.68 ±3.60 (6)	15.30 ±1.53 (7)	16.71 ±4.27 (5)	16.69 ±2.08 (6)	13.54 ±2.14 (10)	21.28 ±2.75 (10)	31.39 ±8.00 (7)	40.45 ±4.74 (8)	36.18 ±7.88 (7)	15.63 ±6.05 (7)
<i>Hegetotherium</i>	7.76 (1)	57.37 ±0.18 (2)	31.23 ±1.16 (2)	41.69 ±1.47 (2)	27.65 ±4.39 (2)	13.18 ±1.46 (3)	19.00 (1)	16.59 ±1.46 (3)	24.02 ±0.83 (3)	40.29 ±1.88 (2)	56.91 ±0.80 (2)	41.05 ±11.70 (3)	4.43 ±0.86 (2)
<i>Pachyrhukhos</i>		37.32 (1)	17.83 ±1.75 (2)	29.63 ±0.14 (2)	19.63 ±1.27 (3)	14.40 ±3.07 (3)	11.64 (1)	10.80 ±0.23 (3)	14.66 ±0.63 (3)	27.32 ±3.26 (3)	39.24 (1)	26.36 ±3.61 (3)	

Diastema length (DLB) was measured only in *Pachyrhukhos* [8.41 ± 0.74 (3)].

show HI values around 3.2 (Table 2) and their HPI ranges from 1.5 to 2.2, so they are clearly over the limits of HI and HPI that separate species dwelling open habitat from those dwelling mixed or closed habitat. They lie on the left-bottom region in the morphospace depicted by HI and JAW, without overlapping with any extant ungulates, but surrounded by open-habitat camelids.

According to their HPI and HI values, most Santacrucian notoungulates foraged in open habitats (Fig. 3). Only three specimens of *Interatherium* show a combination of HI and JAW characteristic of living ungulates that forage in mixed habitats (but see Discussion and conclusions section).

Inferences about diet

All the Santacrucian notoungulates were treated as specimens from open or mixed habitats to perform the analyses about their feeding behaviour. Species from these habitats can be classified as grazers, mixed feeders or browsers.

The classification tree shown in Fig. 4 allowed to discover that the relative width of the muzzle (MZW) and the maximum width of mandibular angle (JMC) play an essential role in the adaptation of ungulates for feeding mainly on grass, allowing a very good characterization of grazers in comparison with other feeding groups (see Fig. 5). This tree correctly reclassifies 94.3% (83 of 88) of species involving only JMC

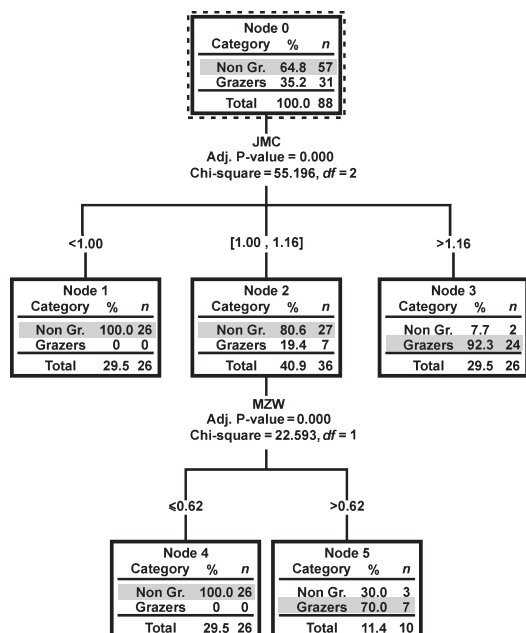


Fig. 4. Classification tree that permitted the discovery of the relationship between both the relative width of the muzzle (MZW) and the relative maximum width of mandibular angle (JMC), and the adaptation of ungulates for feeding on grass.

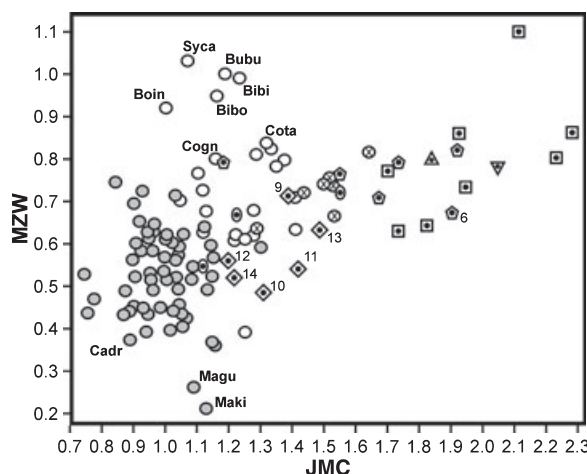


Fig. 5. Distribution of the 88 extant ungulates (circles), inhabitants of open and mixed habitats and 24 notoungulates (dotted symbols) in the morphospace depicted by MZW and JMC, which allows the morphological characterization of grazers (white circles) and non-grazers (grey circles). Notoungulata (dotted symbols): *Interatherium* (squares), *Nesodon* (rhombus), *Protypotherium* (pentagons), *Hegetotherium* (inverse triangles), *Adinotherium* (ovoids), *Pachyrhynchos* (right-angled triangles). Extant ungulates: *Equus* species (crossed circles), *Camelus dromedarius* (Cadr), *Madoqua guentheri* (Magu), *Madoqua kirkii* (Maki), *Connochaetes gnou* (Cogn), *Connochaetes taurinus* (Cota), *Bison bison* (Bibi), *Bos indicus* (Boin), *Syncerus caffer* (Syca), *Bubalus Bubalus* (Bubu) and *Bison bonasus* (Bibo).

and MZW. The importance of these two morphological traits also became evident with discriminant analysis. Curiously enough, the discriminant function that involves the same two variables ($6.253 \text{ MZW} + 5.886 \text{ JMA} - 10.197$) reclassifies correctly the same percentage of species, 94.3%.

Mendoza & Palmqvist (2008) show that grazers have a wider muzzle than other ungulates from open habitats, and the few browsers that live in open habitats have some of the narrowest muzzles among ungulates. We show here that grazers also have a jaw with a wider mandibular angle than other ungulates from open and mixed habitats. The morphospace depicted by the relative width of the muzzle (MZW) and the maximum width of mandibular angle (JMC) allows a very good characterization of grazers in comparison with non-grazers (see Fig. 5).

Although grazers of the family Bovidae (order Artiodactyla) are characterized by wide muzzles, grazers of the family Equidae (order Perissodactyla) mainly have wide mandibular angles (see Fig. 5). Non-grazers in both groups show a common and relatively homogeneous pattern, characterized by both narrow muzzles and mandibular angles.

Santacrucian notoungulates share with equids, a wide mandibular angle (only those specimens for which MZW is also available appear in Fig. 5). Although in equids JMC ranges from 1.29 to 1.64, in

Interatherium it ranges from 1.70 to 2.28, in *Protypotherium* JMC from 1.55 to 2.31, in *Hegetotherium* from 1.13 to 2.05 and in *Pachyrukhos* from 1.47 to 2.11. Notoungulates also share with bovids a wide muzzle. In fact, the highest value of MZW corresponds to an *Interatherium* specimen (see Fig. 5). According to its muzzle and mandibular angle width, *Nesodon* was the less-specialized genus in the consumption of grass, although specimens 9 and 13 have both the muzzle and the mandibular angle as wide as some equids (see Fig. 5).

Other traits also seem to be involved in the morphological patterns that adapt ungulates for feeding mainly grass. As shown in Fig. 6, the morphospace depicted by the depth of the mandibular angle (JMB) and the length of the paraoccipital process (SE) also allows a good characterization of grazers in comparison with non-grazers. Again, grazers from the families Bovidae and Equidae show some differences. Both increased the depth of the mandibular angle (JMB), but much more equids than bovids, and only bovids show a longer paraoccipital process (SE; see Fig. 6). Thus, non-grazer ungulates are again the group of species that show a common pattern, characterized by both a shallower mandibular angle and a shorter paraoccipital process than grazers. Notoungulates share both traits of grazers, a deeper mandibular angle (JMB) and a longer paraoccipital process (SE, see Fig. 6).

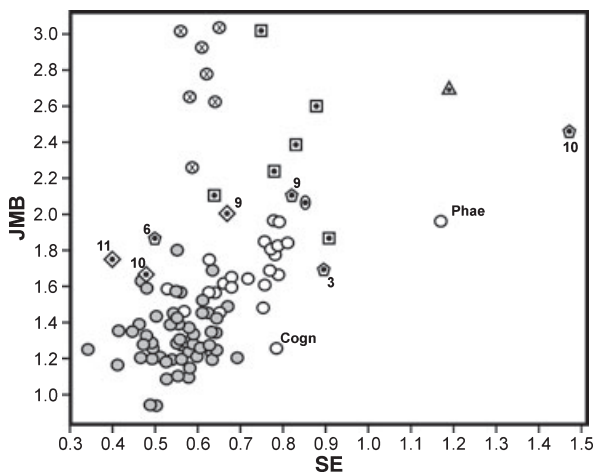


Fig. 6. Distribution of the 88 extant ungulates (circles) inhabitants of open and mixed habitats and 24 notoungulates (dotted symbols) in the morphospace depicted by JMB and SE, which allows the morphological characterization of grazers (white circles) and non-grazers (grey circles). Notoungulata (dotted symbols): *Interatherium* (squares), *Nesodon* (rhombus), *Protypotherium* (pentagons), *Hegetotherium* (inverse triangles), *Adinotherium* (ovoids) and *Pachyrukhos* (right-angled triangles). Extant ungulates: *Equus* species (crossed circles), *Camelus dromedarius* (Cadr), *Madoqua guentheri* (Magu), *Madoqua kirkii* (Maki), *Connochaetes gnou* (Cogn), *Connochaetes taurinus* (Cota), *Bison bison* (Bibi), *Bos indicus* (Boin), *Syncerus caffer* (Syca), *Bubalus Bubalus* (Bubu) and *Bison bonasus* (Bibo).

Among notoungulates, a specimen of *Interatherium* show the deepest mandibular angle, with a JMB value only found in equids. Other specimens show lower values of JMB, and SE values typical of living grazers. There are two specimens of notoungulates, however, with extreme SE values. A specimen of *Protypotherium* shows the longest paraoccipital process, but other specimen of the same genus (Proty 6; Fig. 6), however, has a paraoccipital process shorter than any grazer (SE = 0.5). Its muzzle width is also similar to that of the non-grazers, but its mandibular angle is wider than that of any living non-grazers (Proty 6; Fig. 5). There is also a *Pachyrukhos* specimen with an extreme SE value, although its paraoccipital process is not much longer than that of the only grazer species of the family Suidae (Fig. 6).

JMB is not available for *Nesodon* specimens 12 and 14, which are the closest to the living non-grazers in the morphospace depicted by MZW and JMC (Fig. 5). They could not be so represented in Fig. 6, but both specimens have SE values (0.4 and 0.6) typical of living non-grazers. Both JMB and SE are available, indeed for *Nesodon* 10 and 11, which also show SE values characteristic of non-grazer ungulates, and lower than any living grazer. According to their location in the morphospace depicted by MZW and JMC (Fig. 5), their feeding adaptations are not very clear, but taking also into account their location in the morphospace depicted by JMB and SE (Fig. 6), it seems more probable that both specimens were adapted for grazing and browsing. Neither JMB nor SE is available in specimen 13, but specimen 9 is also located with grazers (Fig. 6).

Discussion and conclusions

The feeding categories of grazing, mixed-feeding and browsing herbivores were initially characterized by Hofmann & Stewart (1972), who correlated feeding behaviour with stomach anatomy. Several authors (Janis & Ehrhardt 1988; Solounias & Moelleken 1993; Janis 1995; Pérez-Barbería & Gordon 1999; Mendoza et al. 2002) have used the correspondences between craniodental morphology and ecological adaptations. However, feeding and habitat morphological adaptations of the craniodental skeleton are not easily distinguishable. In fact, analyses considering the phylogenetic constraints performed by Pérez-Barbería et al. (2001) demonstrated that both habitat and diet are correlated.

The approach followed in this study was a hierarchical analysis identifying the correlation between morphology and habitat types, and then feeding

behaviour between certain habitats, *versus* Spencer (1995), who evaluated habitats preference within diet categories.

High crown is the morphological trait that have been most frequently considered as indicative of feeding behaviour in ungulates in relation to grass consumption, and correlated with the spread of grass land habitats (for extensive overviews on the subject, see Janis 1995; MacFadden 2000; Strömberg 2006; Billet *et al.* 2009; Reguero *et al.* 2010). Van Valen (1960) provided the first quantitative approach to these relationships, defining an HI. It was not until the late 1980s that a comparative study by Janis (1988) tested this hypothesis in living ungulates, hyraxes and the African elephant. Further research (Janis 1995) led this author to conclude that HI better indicates open-habitat environmental preference than determining grass consumption in extinct ungulates.

More recently, after removing phylogenetic effects, Williams & Kay (2001) found that tooth crown height in rodents and ungulates was positively correlated with the proportion of monocots (primarily grass) in the diet and was negatively correlated with foraging height preferences. In a study on fossil equids and phytoliths, Strömberg (2006) showed that care should be taken when interpreting diet/habitat from hypsodonty of extinct species. Moreover, Mendoza & Palmqvist (2008) found that, in ungulates, there is no direct relationship between the degree of hypsodonty and the amount of grass consumed, except for the correspondence between grazing and dwelling in open habitats. These authors also proposed that hypsodonty represents a 'key adaptation' of ungulates for feeding in open environments, where the vegetation that grows at ground level accumulates airborne grit (Mendoza & Palmqvist 2008). In a recent contribution, Billet *et al.* (2009) summarized the factors that favoured increasing hypsodonty as two alternative hypotheses, named increasing chewing effort and increase of abrasives consumed.

Generating a hypothesis about habitat preferences was the first step in our analysis. In Mendoza & Palmqvist (2008), all ungulates that dwell in open or mixed habitat has an HI >2, and the HI that allows distinguishing open habitats from mixed habitats increases with the relative length of diastema (JLB). In this contribution, we faced the problems that most notoungulates studied do not have diastema, and therefore JLB was not applicable to our sample, and for *Pachyruchos* – the only one with a diastema – there are not specimens available in the collection studied to calculate HI following the procedure described above. Applying the same knowledge-discovery techniques as that of Mendoza & Palmqvist (2008), we found that the other measurement that shows a

similar pattern is the length of the jaw (JAW). Thus, there are species adapted to open habitats with low hypsodonty, like camels, but all of them have a short jaw, and there are hypsodont species dwelling mixed habitats, which have jaws that are always long.

Using the relative length of the jaw and the HI, we defined a HPI, which allows the inference of whether an ungulate species is more likely to dwell open or mixed habitats. Using that index, combined with the HI, it is possible to characterize the species adapted to forage in open, mixed and closed habitats. Only open-habitat highly selective browsers feeding on heavy shrubbery such as *Madoqua guentheri* and *Madoqua kirkii* the dik-diks and *Oreamnos americanus* the Mountain Goat, which is found only on the steep slopes of Rocky Mountain, and those species dwelling open but water-associated habitats such as *Ozotoceros bezoarticus* the pampas deer and Waterbucks and another three species of the genus *Kobus*, which have a HPI lower than 1, cannot be distinguished from those species dwelling mixed habitats (Fig. 3).

According to their HPI and HI values and the region that they occupy in the morphospace depicted by HI and JAW, all Santacrucian notoungulates foraged in open habitats (Fig. 3). With the exception of two specimens of *Hegetotherium*, they show a combination of HI and JAW that is not very different from the extant ungulate species dwelling open habitats. *Hegetotherium* specimens (HI = 7.76) are almost as hypsodont as the wild ass (*Equus asinus*, HI = 8.73), but their jaws are much shorter, so they occupy a region of the morphospace depicted by HI and JAW, which is not occupied by any extant ungulate species (see Fig. 3). In the case of *Interatherium*, it could have also been from mixed or water-associated open habitats.

Following Tauber (1997a), one peculiarity of the Santa Cruz Formation is the presence of indicators of different climatic and environmental conditions. On one side, the occurrence of Primates, and echimyid and erethizontid rodents has been interpreted as indicators of warm and humid conditions, and forests (Vucetich 1986; Pascual & Ortiz-Jaureguizar 1990). On the other hand, the diversity of armadillos (Vizcaíno *et al.* 2006) and the presence of probable palaeodunes (Bown & Lariestra 1990), gypsum crystals, mud cracks and other sedimentological features (Tauber 1997a) suggest the existence of open environments in relatively dry conditions. Additionally, the Santa Cruz Formation rocks are, to a great extent, a sequence of superimposed mudstones of volcanoclastic origin, although some relatively unaltered tuffs also occur (Bown & Fleagle 1993). The consequent abundance of abrasive volcanic particles may account for much of the grit that the herbivores must have

ingested with the food (Kay *et al.* 1999). The knowledge of the evolution of the early Miocene palaeoflora of Patagonia (Barreda & Palazzesi 2007) is coherent with this scenario, confirming the existence of both close and open environments. Although forests remained, probably developing riparian or gallery forests, across extra-Andean Patagonia until about the middle Miocene, in the late early Miocene drier conditions would have prevailed in lowland areas, and the contraction of humid elements coincides with the expansion of xerophytic taxa.

The next step in our research was to characterize the feeding craniodental traits of ungulates correlated to dwelling open or mixed habitats. The features distinguishing grazers from browsers probably relate to the different physical demands of feeding on grass *versus* leaves. Muzzle width has been considered important for the inference of alimentary styles in extinct herbivores (Solounias & Moelleken 1993; Janis 1995; Solounias *et al.* 1995). Grass is in general more fibrous, and requires a greater intake and degree of mastication than leaves. Grazers show wider muzzles than browsers, which allow the intake of large bites (bulk feeding) while grazing (Owen-Smith & Novellie 1982). Grazers also have bigger masseter muscles than browsers, reflected in a larger and deeper angle of the jaw, and a longer masseteric fossa on the skull (Axmacher & Hofmann 1988; Clauss *et al.* 2008). Browsers are selective feeders (Gordon & Illius 1988), have a narrower muzzle than grazers, which is useful for feeding selectively on certain plants or parts of plants (Solounias *et al.* 1988; Solounias & Moelleken 1993).

According to our results, non-grazers show a common and relatively homogeneous morphological pattern, which characterize them in comparison with grazers. They show a narrow muzzle, a shallow and narrow mandibular angle, and a short paraoccipital process. Thus, in the four-dimensional morphospace defined by these variables, all the non-grazers would occupy a region around 1 of the 16 vertices of a tesseract (the four dimensional analogue of a cube) containing all the species. Grazers are not so homogeneous, as there are some differences in the way that bovids and equids adapted to feed mainly on grass. Both groups widened their muzzle, but bovids much more than equids, which mainly widened their mandibular angle (see Fig. 5). Both groups also increased the depth of the mandibular angle, but much more equids than bovids, and only bovids enlarged the paraoccipital process (see Fig. 6).

Among the assemblage of notoungulates of the Santa Cruz Formation, the tyotheres *Interatherium*, *Protypotherium*, *Hegetotherium* and *Pachyrukhos* show some characteristic of living grazers, i.e. a wide muzzle, an extremely wide and deep mandibular angle,

and a large paraoccipital process (see Figs 5, 6). Thus, they show a morphological pattern similar to the one of those extant ungulates adapted to feed mainly on grass, and far from the one of those that feed mainly on browse, or grass and browse depending of availability. Tyotheres, however, do not fit with the general pattern observed in extant ungulates, as small forms are browsers (Pérez-Barbería & Gordon 2001). Based on the incisors morphology and implantation, Tauber (1996) proposed that the larger species of *Protypotherium* (*P. australe*) was adapted to forage on harder grasses than the other two (*P. praerutilum* and *P. attenuatum*). Among hegetotherids, Reguero *et al.* (2007) concluded that, according to their muzzle and masseteric morphology, Oligocene pachyrukinines were selective grazers.

Rodent-like tyotheres (especially *Interatherium* and *Pachyrukhos*) exhibit morphological features that exaggerate the patterns shown by modern grazing ungulates, and some (especially *Hegetotherium*) exaggerated even more the features (HI) that indicate dwelling in open habitats. The last two statements suggest that both the peculiar general rodent-like morphology and other than feeding habits, like digging, may be influencing their position in the respective morphospaces. A locomotory morphofunctional analysis by Elissamburu (2004), comparing with living caviomorph rodents, reveals that certain indices allow to characterize *Pachyrukhos* and *Protypotherium* as occasional diggers and *Interatherium* and *Hegetotherium* as diggers. Billet *et al.* (2009) proposed that their 'grit hypothesis' applied to other notoungulates like the mesotheriid *Trachytherus* from the Oligocene of Bolivia, which could be a fossorial animal that ingested a great quantity of exogenous grit along with its food (Shockey *et al.* 2007). It is known that, in caviomorph rodents, adaptation to digging habits also correlates with craniodental traits (Verzi & Olivares 2006) and that digging behaviour usually implies the ingestion of large quantities of abrasive soil elements playing a role in the development of hypsodonty in xenarthrans (Bargo *et al.* 2006). As it was mentioned above, abundance of abrasive volcanic particles in the Santa Cruz Formation may account for much of the grit that the herbivores must have ingested with the food. Considering their small body size, general likeness with non-ungulate herbivores (glires, hyraxes), that they have no close relatives among living ungulates and that they do not match exactly the same morphospace, other living herbivores analogues should be used in further analyses to contrast these inferences.

The toxodontid *Adinotherium* also shows the craniodental morphology characteristic of living grazers, i.e. a wide muzzle, an extremely wide and deep mandibular angle, and a large paraoccipital process

(see Figs 5, 6). *Nesodon*, however, seems to be less specialized in the consumption of grass than the others notoungulates. Most specimens of *Nesodon* are similar to living ungulates classified as a mixed feeder, although there are at least two specimens of *Nesodon* showing a morphological pattern close to the typical one of living equids. Toxodontids similarity to other ungulates has been highlighted since the earliest studies of Patterson (1932). They attained body sizes comparable with living ungulates and share the morphospace occupied by living grazing and mixed-feeders ungulates in open habitats.

There are some other traits, which have not been represented here, that also seem to be involved in the morphological patterns that adapt ungulates for feeding mainly on grass. The posterior jaw length (JMA), for example, also allows a good characterization of grazers in comparison with non-grazers when it is represented in combination with the length of the paraoccipital process (SE). Non-grazer ungulates are characterized by both a shorter posterior jaw and paraoccipital process. Among notoungulates, only *Nesodon*'s three specimens occupy a region between grazers and non-grazers, whereas the fourth is located in the middle of typical grazers.

Grazers also show a longer posterior portion of the skull (SC), but this is the only trait that Santacrucian notoungulates do not share with them. *Phacochoerus aethiopicus* the warthog, the only grazer of family Suidae, is also the only extant grazer that shares a short posterior portion of the skull (SC) with notoungulates.

In summary, our results indicate that the Santacrucian notoungulates show similar, or even exaggerated, morphological patterns to extant ungulates adapted for feeding mainly on grasses, and far from those mixed-feeders or browsers. This implies interesting consequences in the consideration of the evolution of craniodental traits to grazing habits in ungulates. Following Janis *et al.* (2004), it was not until the start of the late Miocene, around 10 Ma, that ungulates with morphologies consistent with more specialized grazing adaptations appeared. The fact that typotheres could be classified as grazers implies that native South American ungulates evolved grazing specializations at least by the late-early Miocene at ca. 16 MA, i.e. much before than extant main lineages in northern hemisphere. This hypothesis had been proposed before on the basis that hypsodonty among ungulates has its first appearance in South American lineages rather than in northern hemisphere ungulates (see Patterson & Pascual 1972; MacFadden 2000; Billet *et al.* 2009).

Using species from two living orders of ungulates, artiodactyls and perissodactyls, as a reference sample, to infer the ecological behaviours of a third extinct

order of ungulates, we are assuming that notoungulates resolved the same biomechanical problems with convergent morphological solutions. However, the majority of living ungulates are ruminant artiodactyls, and quantitative correlations of morphology and behaviour derived from ruminants may not be directly applicable to other types of ungulates, although general qualitative observations may still hold true (Janis 2008). This is particularly remarkable when dealing with forms that developed so different morphologies as typotheres. Although a validation of our results through other approaches (e.g. morpho-functional, biomechanical, biogeochemical and tooth micro- and mesowear analyses) remains pending, the fact that different morphological traits consistently point to the same adaptations strengthens them.

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