An overview of the appendicular skeletal anatomy of South American titanosaurian sauropods, with definition of a newly recognized clade

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Abstract: In the last two decades, the number of phylogenetically informative anatomical characters recognized in the appendicular skeleton of titanosaurian sauropod dinosaurs has increased dramatically with the discovery of new and comparatively complete specimens. Here we provide an overview of the appendicular skeletal morphology of South American titanosaurians and discuss its significance for phylogenetic reconstruction. The appendicular skeletal diversity of South American titanosaurians is substantially greater than was initially appreciated. Moreover, some regions of the appendicular skeleton, such as the pes, exhibit remarkable variability in form. Multiple synapomorphies of Titanosaura and the less inclusive clades Lithostrotia and Saltasauridae consist of characters of the girdles and limbs. Although the phylogenetic definitions of titanosaurian clades such as Saltasaurinae and Lognkosauria are stable, the taxonomic content of these clades has varied in recent analyses depending on the phylogenetic topology recovered. Within Titanosaura, the results of four recent, largely independent analyses support the existence of a derived titanosaurian lineage distinct from the ‘Saltasaurinae line,’ which is herein termed Colossosauria. At present, this clade is mainly comprised by taxa within Lognkosauria and Rinconsauria, and is useful in discussions of titanosaurian lower-level relationships.

Key words: Titanosaura, South America, appendicular skeleton, osteology, phylogeny, Colossosauria.
INTRODUCTION


Anatomical and phylogenetic analyses of titanosaurians are crucial for deciphering the evolutionary history and paleobiology of these singular animals, but such studies have been hampered by missing data, as the osteology of many titanosaurian species is not well understood. Fortunately, this situation is beginning to change with recent discoveries and descriptions of well-preserved, comparatively complete specimens of taxa such as *Tapuiasaurus* from the Early Cretaceous of Brazil (Zaher et al. 2011, Wilson et al. 2016), *Bonitasaura* (Apesteguía 2004, Gallina 2011, Gallina and Apesteguía 2011, 2015), *Dreadnoughtus* (Lacovara et al. 2014, Ullmann and Lacovara 2016, Voegele et al. 2017), *Epachthosaurus* (Martínez et al. 2004), *Futalognkosaurus* (Calvo et al. 2007a, b), *Mendoza­saurus* (González Riga 2003, 2005, González Riga et al. 2018), *Overosaurus* (Coria et al. 2013), and *Patagotitan* (Carballido et al. 2017) from the mid- and Late Cretaceous of Argentina; *Rapetosaurus* (Curry Rogers and Forster 2001, 2004, Curry Rogers 2009) from the latest Cretaceous of Madagascar; *Mansourasaurus* (Sallam et al. 2018), *Rukw­ititan* (Gorscak et al. 2014), and *Shingopana* (Gorscak et al. 2017) from the Late Cretaceous of Africa; *Diamantinasaurus* (Hocknull et al. 2009, Poropat et al. 2015, 2016) and *Savannasaurus* (Poropat et al. 2016) from the Late Cretaceous of Australia, and *Lohuecotitan* (Diez Díaz et al. 2016) from the Late Cretaceous of Europe. Nevertheless, some areas of the titanosaurian skeleton remain poorly understood. For instance, the morphology of the appendicular skeleton has been extensively documented in only a few taxa, such as *Bonitasaura* (Apesteguía 2004, Gallina and Apesteguía 2015), *Diamantinasaurus* (Hocknull et al. 2009, Poropat et al. 2015), *Dreadnoughtus* (Lacovara et al. 2014, Ullmann and Lacovara 2016), *Epachthosaurus* (Martínez et al. 2004), *Neuquensaurus* (Otero 2010), *Opisthocoelicaudia* (Borsuk-Bialynicka 1977), *Rapetosaurus* (Curry Rogers 2009), and *Saltasaurus* (Powell 1992, 2003). More specifically, the distal forelimb is not well known in
most titanosaurs, and in some taxa, the presence or absence of ossified carpals and manual phalanges is controversial (see, e.g., Apesteguía 2005, Mannion and Otero 2012, Poropat et al. 2015). Similarly, the pes of titanosaurs is poorly known; until recently, complete, articulated pedes were known only in Epachthosaurus and Opisthocoelicaudia. Fortunately, knowledge of titanosaurian pedal osteology has recently been enhanced by the discovery of the giant titanosaur Notocolossus from the Late Cretaceous of Mendoza Province, Argentina (González Riga et al. 2016) plus two other, as-yet unnamed species from Mendoza (the ‘Agua del Padrillo taxon,’ UNCUYO-LD 313, González Riga et al. 2015) and Neuquén (the ‘La Invernada taxon,’ MUCPv-1533, González Riga et al. 2008a) provinces, respectively (Figure 1). Furthermore, nearly complete pedes are known for a few other titanosaurs as well, such as Bonitasaura (Gallina and Apesteguía 2015), Mendozasaurus (González Riga et al. 2018), Rapetosaurus (Curry Rogers 2009), and possibly Alamosaurus (the latter based on NMMNH P-49967, a questionably referred specimen from the Upper Cretaceous of the southwestern U.S.A., D’Emic et al. 2011). The pes is very incompletely known in most other titanosaurs.

As noted above, phylogenetic analyses of titanosaurs are important as a basis for further evolutionary and paleobiogeographic interpretations pertaining to this widespread sauropod clade (e.g., Upchurch 1995, 1998, Salgado et al. 1997, Wilson and Sereno 1998, Wilson 2002, Upchurch et al. 2004, Curry Rogers 2005, D’Emic 2012, Mannion et al. 2013, González Riga et al. 2016, 2018, Gorscak and O’Connor 2016, Poropat et al. 2016, Sallam et al. 2018). Appendicular character states often comprise a significant component of the data employed in such analyses (e.g., D’Emic 2012, Mannion et al. 2013, González Riga et al. 2016, Gorscak and O’Connor 2016); moreover, some of these characters constitute synapomorphies of particular titanosaurian clades.

Many aspects of the girdle and limb morphology of titanosaurs (plus additional taxa within the more inclusive neosauropod clade Macronaria, Wilson and Sereno 1998) have been interpreted as being related to the acquisition of wide-gauge posture, where the manus and pedes are located at a considerable distance from the sagittal midline. This interpretation has been supported by many Late Cretaceous sauropod trackways attributed to derived titanosaurs (e.g., Farlow 1992, Lockley et al. 1994, Calvo 1999, Wilson and Carrano 1999, Wilson 2006, González Riga and Calvo 2009). However, several exceptions to this potential correlation between appendicular morphology and posture have also been recognized in the ichnological record (e.g., Lockley et al. 2002, Stevens et al. 2016), and as such, it is now customary for well-preserved trackways to be carefully analyzed, including studies of gait (e.g., Vila et al. 2008, González Riga and Tomaselli 2019). Additionally, computer-aided biomechanical studies are casting new light on the stance and locomotion of sauropods in general (Klinkhamer et al. 2018).

Another complex paleobiological aspect of Titanosauria is the gigantism attained by some lineages and its relationship to appendicular osteology. Wilson and Carrano (1999) and Carrano (2005) argued that many appendicular features seen in titanosaurs and other sauropods—such as graviportal, columnar limb posture, increased limb bone robusticity, shortened distal limb segments, and increased femoral midshaft eccentricity—appear intimately related to the acquisition of large body size. As in other quadrupedal dinosaurs, the manus and pes of sauropods exhibit phalangeal reduction (Osborn 1904, Coombs 1975, Upchurch 1995, 1998, Wilson and Sereno 1998). Typically, such reduction occurred primarily in terms of length, with individual phalanges becoming compact and
Figure 1 – Recent discoveries of appendicular skeletal elements of Late Cretaceous titanosaurian sauropods from the Neuquén Basin of southern Argentina. (a) complete and articulated left hind limb of the unnamed La Invernada taxon (MUCPv-1533), (b) complete and articulated distal left hind limb of the unnamed Agua del Padrillo taxon (UNCUYO-LD 313), (c) right humerus of *Notocolossus gonzalezparejasi* (UNCUYO-LD 301), (d) complete and articulated right pes of *Notocolossus gonzalezparejasi* (UNCUYO-LD 302). Scale bar equals 10 cm in (d).
often disc-like. However, titanosaurs continued this trend, with many taxa reducing and in some cases apparently eliminating ossified manual phalanges. Furthermore, titanosaurs also exhibit the most reduced pedal phalangeal formulae seen within Sauropoda (Borsuk-Bialynicka 1977, Salgado et al. 1997, Wilson and Sereno 1998, Martínez et al. 2004, Apesteguía 2005, González Riga et al. 2008a, 2016).

In the present contribution, we provide an overview of titanosaurian appendicular skeletal anatomy, focusing on the many representatives of this clade that have been recovered from the Cretaceous of South America. We also discuss several of the principal appendicular skeletal characters that have been used in previous phylogenetic analyses of Titanosauria and its subclades, with the goal of identifying areas of agreement and conflict. Finally, we formally define a new clade of derived South American titanosaurs, recognized on the basis of results of recent cladistic analyses.

INSTITUTIONAL ABBREVIATIONS


THE PECTORAL AND FORELIMB SKELETON OF SOUTH AMERICAN TITANOSAURS

Although the fossil record of South American titanosaurs is rich in terms of numbers of specimens and species, taxa with completely known pectoral girdles and forelimbs are scarce (Table I). Exceptions include Epachthosaurus (Martínez et al. 2004) and the saltasaurines Neuquensaurus (Otero 2010) and Saltasaurus (Powell 1992, 2003), for which the pectoral girdle and forelimb are almost completely known, and to a lesser extent Argyrosaurus (Mannion and Otero 2012), Dreadnoughtus (Lacovara et al. 2014, Ullmann and Lacovara 2016), Mendozaaurus (González Riga 2003, 2005, González Riga et al. 2018), Muyelensaurus (Calvo et al. 2007c), Patagotitan (Carballido et al. 2017), Tapuiasaurus (Zaher et al. 2011), and Uberabatitan (Salgado and de Souza Carvalho 2008). The pectoral girdles and forelimbs of South American titanosaurs exhibit the basic anatomical plan of sauropods in general, but also possess unique features that differentiate them from those of non-titanosaurian taxa.
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<td>Calvo et al. 2007a, b, Calvo 2014</td>
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South American titanosaur species for which appendicular elements are preserved. Abbreviations: Sc, Scapula; Co, Coracoid; St, Sternal plate; Hu, Humerus; Ra, Radius; Ul, Ulna; Mc, Metacarpals; Il, Ilium; Pu, Pubis; Is, Ischiium; Fe, Femur; Ti, Tibia; Fi, Fibula; As, Astragalus; Mt, Metatarsals; Pp, Pedal phalanges. Y = bone(s) is/are preserved in the taxon in question; -- = bone(s) not preserved; ? = bone(s) questionably preserved.
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Well-preserved scapulae are reported in *Aeolosaurus rionegrinus* (Powell 2003), *Antarctosaurus wichmannianus* (Huene 1929), *Choconsaurus* (Simón et al. 2018), *Dreadnoughtus* (Ullmann and Lacovara 2016), *Elaltitan* (Mannion and Otero 2012), *Mendozaaurus* (González Riga et al. 2018), *Muyelensaurus* (Calvo et al. 2007c), *Neuquensaurus* (Otero 2010), *Patagotitan* (Carballido et al. 2017), *Pitekunsaurus* (Filippi and Garrido 2008), and *Saltasaurus* (Powell 1992, 2003) among other South American titanosaurs (Table I, Figure 2). Additionally, although the scapula is poorly preserved in the type and only known specimen of *Drusilasaura*, its overall shape is evident (Navarrete et al. 2011).

As in non-titanosaurian sauropods, the proximal (i.e., anterior, if the long axis of the scapula is oriented horizontally) end of the scapula is more expanded than the distal (i.e., posterior) end. In most South American titanosaurs (e.g., *Drusilasaura, Mendozaaurus, Patagotitan, Pitekunsaurus, Saltasaurus*), the proximal expansion is less than twice the dorsoventral breadth of the distal end, with the exceptions of *Dreadnoughtus* and *Muyelensaurus*, in which the proximal end is more than twice as broad as the distal. Nevertheless, the scapulae of most South American titanosaurs also exhibit a slight distal expansion, with the dorsal (i.e., acromial) edge of the blade being dorsally deflected and the ventral edge remaining nearly straight. In most titanosaurs, however, the tallest part of the dorsal margin of the scapular blade is still lower than that of the proximal expansion. An exception is seen in *Patagotitan*, in which the most elevated point of the blade is approximately the same height as that of the proximal end (Figure 2h). Among South American titanosaurs, a reversal to an unexpanded distal scapular blade occurs in *Dreadnoughtus*, in which the dorsal and ventral
edges of the blade are subparallel to one another (Figure 2d).

In some South American titanosaurs (e.g., *Patagotitan*, *Pitekunsaurus*), as in other members of the clade (e.g., *Rapetosaurus*), the orientation of the scapular blade relative to the coracoid articulation is roughly 45°. In other forms (e.g., *Dreadnoughtus*, *Muyelensaurus*), however, the long axes of the blade and coracoid articulation are oriented roughly perpendicular to one another.

In titanosaurs, the acromial ridge is generally not as developed as it is in non-titanosaurian neosauropods such as *Diplodocus* (Hatcher 1901), *Giraffatitan* (Janensch 1961), or *Camarasaurus* (McIntosh et al. 1996), in which this ridge clearly delimits a proximal scapular fossa, widely regarded as the origin site of the M. supracoracoideus (Meers 2003, Otero 2018) (Figure 2m-o). In *Patagotitan*, *Pitekunsaurus*, and the recently described, probably basal titanosaur *Choconsaurus*, the acromial region grades much more smoothly into the scapular blade than is the case in some other taxa (e.g., *Aeolosaurus rionegrinus*, *Dreadnoughtus*, *Elaltitan*, *Muyelensaurus*). When examined in lateral view, the scapular blade is dorsoventrally deep in some taxa (e.g., *A. rionegrinus*, *Patagotitan*, *Saltasaurus*) but substantially shallower in others (e.g., *Dreadnoughtus*, *Muyelensaurus*). A ventromedial process on the ventral margin of the blade (*sensu* Carballido et al. 2011) is present in *Dreadnoughtus*, *Elaltitan*, *Neuquensaurus*, *Patagotitan*, and *Saltasaurus*, among other South American titanosaurs.

**CORACOID**

The coracoid is known in a minority of South American titanosaurs, including *Choconsaurus*, *Dreadnoughtus*, *Epachthosaurus*, *Narambuena* (Filippi et al. 2011a), *Neuquensaurus*, *Patagotitan*, *Quetecsaurus* (González Riga and Ortiz David 2014), *Rinconsaurus* (Calvo and González Riga 2003), *Saltasaurus*, *Tapuiasaurus*, and *Uberabatitan* (Table I, Figure 3a-c, e). In Titanosauria, the proximodistal (i.e., anteroposterior, if the long axis of the scapulocoracoid is oriented horizontally) length of the coracoid may be up to twice the length of the scapular articulation. The coracoid was progressively modified through the evolution of Saltasauridae, increasing in proximodistal elongation, becoming ‘squared’ at its anteroventral margin, and ultimately extending to the height of the acromial process (i.e., attaining a flush suture with the scapula). These derived morphologies are reported in many South American titanosaurs, such as the saltasaurines *Neuquensaurus* and *Saltasaurus* and the lognkosaurian *Quetecsaurus*, and are shared with some Asian titanosaurs (e.g., *Opisthocoelicaudia*, ZPAL MgD-I/48, Borsuk-Bialynicka 1977) and the North American *Alamosaurus* (USNM 15560, Gilmore 1946) as well. An infraglenoid lip is present in most or all South American titanosaurs for which the coracoid is preserved. The position of the coracoid foramen varies among South American titanosaurs, being situated immediately adjacent to the scapular articulation in some taxa (e.g., *Quetecsaurus*, Figure 3c) but located further from this articulation in others (e.g., *Saltasaurus*, Figure 3b). Nevertheless, as the position of the coracoid foramen appears to change through neosauropod ontogeny (see, e.g., Ullmann and Lacovara 2016), these distinctions are probably of limited taxonomic and phylogenetic significance.

**STERNUM**

Sternal plates are reported with some frequency among South American titanosaurs, being known in *Bonitasaura* (Gallina and Apesteguía 2015), *Choconsaurus*, *Dreadnoughtus*, *Epachthosaurus*, *Maxakalisaurus* (Kellner et al. 2006), *Mendozasaurus*, *Muyelensaurus*,
Figure 2 – Morphological comparison of right scapulae of titanosaurians (a-k) versus those of other sauropods (l-o) in lateral view. South American (Argentinean) taxa denoted with an asterisk. (a) *Pitekusaurus macayai*, redrawn from Filippi and Garrido (2008), (b) *Rapetosaurus krausei*, redrawn from Curry Rogers (2009), (c) *Choconsaurus baileywillisi*, redrawn from Simón et al. (2018), (d) *Dreadnoughtus schrani* (left, reversed), redrawn from Lacovara et al. (2014), (e) *Muyelensaurus pecheni*, redrawn from Calvo et al. (2007c), (f) *Saltasaurus loricatus*, redrawn from Powell (1992), (g) *Mendozasaurus neguyelap*, redrawn from González Riga et al. (2018), (h) *Patagotitan mayorum*, redrawn from Carballido et al. (2017), (i) *Drusilasaura deseadensis* (reversed), redrawn from Navarrete et al. (2011), (j) *Opisthocoelicaudia skarzenskii*, redrawn from Borsuk-Bialynicka (1977), (k) *Elalititan liloi*, redrawn from Mannion and Otero (2012), (l) *Mamenchisaurus youngi* (left, reversed), redrawn from Ouyang and Ye (2002), (m) *Diplodocus carnegii* (left, reversed), redrawn from Hatcher (1901), (n) *Giraffatitan brancai* (left, reversed), redrawn from Janensch (1961), (o) *Camarasaurus supremus* (left, reversed), redrawn from Osborn and Mook (1921). Not to scale.
Narambuenatitan, Neuquensaurus, Patagotitan, Petrobrasaurus (Filippi et al. 2011b), Rinconsaurus, Saltasaurus, Tapuiasaurus, and Uberabatitan (Table I, Figure 3f-k). Unlike those of non-titanosaurian sauropods (e.g., the brachiosaurid Giraffatitan) the sternal plates of titanosaurs are generally semilunate in shape and at least 65–70% the length of the humerus (McIntosh 1990, Salgado et al. 1997, Upchurch 1998, D’Emic 2012, Mannion et al. 2013). Nevertheless, the sternal plates of Choconsaurus and Mendozasaurus are less mediolaterally expanded at their posterior ends than are those of some other titanosaurs (e.g., Dreadnoughtus, Maxakalisaurus, Narambuenatitan, Petrobrasaurus, probably Tapuiasaurus), rendering them more subtriangular than semilunate in dorsal or ventral view (Figure 3f, g). This shape, with some variation, was described as a semilunar sternal plate with a relatively straight posterior border by González Riga (2003) and González Riga and Ortiz David (2014). Similarly, there is considerable variation in the lateral projection of the anterior and posterior ends—and therefore the concavity of the lateral margin—of the sternal plate between different South American titanosaurian genera. In Choconsaurus and Mendozasaurus, the lateral margin is only slightly concave in dorsal view, whereas in taxa such as Maxakalisaurus, Saltasaurus, and especially Narambuenatitan this margin is deeply concave; forms such as Dreadnoughtus, Petrobrasaurus, Rinconsaurus, and Tapuiasaurus appear to exhibit an intermediate condition (Figure 3f-k). There is a ridge on the ventral surface of the sternal plate in Bonitasaura, Dreadnoughtus, Neuquensaurus, Patagotitan, and Saltasaurus, among other taxa. 

HUMERUS

The humerus is the most frequently preserved appendicular bone among South American titanosaurian taxa, being known in 27 valid species at present (Table I). It is highly variable in proportions, ranging from slender in taxa such as Gondwanatitan (Kellner and de Azevedo 1999), Mendozasaurus, Panamericansaurus (Calvo and Porfiri 2010), Petrobrasaurus, Rinconsaurus, and especially Muyelensaurus, in which the proximal and distal ends are not markedly expanded, to exceedingly robust as in Dreadnoughtus and saltasaurines (Figure 4). For example, the humeral robustness index (sensu Wilson and Upchurch 2003) of Muyelensaurus is 0.18, whereas in specimens of the saltasaurine Neuquensaurus (e.g., MLP-CS 1049) this index may reach values of up to 0.32. In some South American titanosaurs (e.g., Argyrosaurus, Atacamatitan [Kellner et al. 2011], Bonatitan [Martinelli and Forasiepi 2004, Salgado et al. 2015], Futalognkosaurus [Calvo, 2014], Elaltitan, Mendozasaurus, Muyelensaurus, Narambuenatitan, Neuquensaurus, Uberabatitan), the proximal margin of the humerus is straight, or nearly so, in anterior view (e.g., Figure 4a, c). In other titanosaurs, however (e.g., Notocolossus [González Riga et al. 2016], Opisthocoelicaudia, Paralititan, Quetecsaurus, Saltasaurus), the proximal margin of the humerus is sinuous due to the marked proximal deflection of the humeral head relative to the remainder of the bone (Figure 4j). The lateral margin of the humerus is nearly straight through approximately the proximal half of the element. Futalognkosaurus has a relatively robust humerus with an expanded proximal end that reaches 40% of the total length of the bone, as in Saltasaurus, Neuquensaurus, and Opisthocoelicaudia (Calvo 2014).

The humeri of many South American titanosaurs (e.g., the saltasaurines Neuquensaurus and Saltasaurus, the lognkosaurian Futalognkosaurus [Calvo 2014], Patagotitan, and Dreadnoughtus) possess a well-developed posterolateral bulge around the level of the deltopectoral crest, which is frequently regarded as the insertion site of the M. deltoideus clavicularis (Meers 2003, Otero 2018).
Unlike many other South American titanosaurs, Neuquensaurus and Saltasaurus also exhibit a deltopectoral crest that is markedly expanded distally, a feature that is shared with the Laurasian titanosaurs Alamosaurus and Opisthocoelicaudia and that consequently has been regarded as a synapomorphy of Saltasauridae (Wilson 2002, D’Emic 2012). South American titanosaurs also exhibit variability in features such as the distal extent and medial deflection of the deltopectoral crest, the development of a proximolateral process, and the concavity of the medial margin of the shaft in anterior view. In saltasaurines and other lithostrotians (e.g., the non-South American genera Alamosaurus, Isisaurus, Opisthocoelicaudia, and Rapetosaurus), the articular surfaces of the humeral distal condyles are exposed on the anterior surface of the bone. The humeral distal condyles of titanosaurians are generally flat, except in saltasaurids and Epachthosaurus in which they are divided.

The longest humerus known for any titanosaur is that of the holotypic specimen of Notocolossus (UNCUYO-LD 301, Figures 1c, 4j, González Riga et al. 2016). This bone is even longer and more proximally robust than the humerus of Patagotitan (Figure 4h), a titanosaur that was recently described as the largest dinosaur yet discovered (Carballido et al. 2017). The humerus of Notocolossus is also

Figure 3 – Morphological comparison of titanosaurian coracoids (a-e) and sternal plates (f-k). South American (Argentinean) taxa denoted with an asterisk. (a) Right coracoid in medial view of Neuquensaurus australis*, MLP-Ly 14, redrawn from Otero (2010), (b) left coracoid in lateral view of Saltasaurus loricatus*, PVL 4017-100, redrawn from Powell (1992), (c) right coracoid in medial view of Quetecsaurus rusconi*, UNCUYO-LD 300.15, redrawn from González Riga and Ortiz David (2014), (d) left coracoid in lateral view of Opisthocoelicaudia skarzynskii, ZPAL MgD-I/48, redrawn from Borsuk-Bialynicka (1977), (e) left coracoid in lateral view of Dreadnoughtus schrani*, MPM-PV 1156, redrawn from Ullmann and Lacovara (2016), (f) right sternal plate in dorsal view of Mendozaaurus neguyelap*, IANIGLA-PV 067, redrawn from González Riga et al. (2018), (g) right sternal plate in dorsal view of Choconsaurus baileywillist*, MMCh-PV 44/11, redrawn from Simón et al. (2018), (h) left sternal plate in ventral view of Saltasaurus loricatus*, PVL 4017-102, redrawn from Powell (1992), (i) left sternal plate in ventral view of Narambuenatitan palomoi*, MAU-Pv-N-425, redrawn from Filippi et al. (2011a), (j) left sternal plate in ventral view of Dreadnoughtus schrani*, MPM-PV 1156, redrawn from Ullmann and Lacovara (2016), (k) left sternal plate in ventral view of Petrobrasaurus puestohernandezi*, MAU-Pv-N-449/25, redrawn from Filippi et al. (2011b). Not to scale.
Figure 4 – Morphological and size comparison of titanosaurian right humeri in anterior view. South American (Argentinean) taxa marked with an asterisk. (a) *Neuquensaurus australis* (left, reversed), redrawn from Otero (2010), (b) *Rapetosaurus krausei* (left, reversed), redrawn from Curry Rogers (2009), (c) *Muyelensaurus pecheni* (left, reversed), redrawn from Calvo et al. (2007c), (d) *Narambuenatitan palomoi* (left, reversed), redrawn from Filippi et al. (2011a), (e) *Mendozasaurus neguyelap*, redrawn from González Riga et al. (2018), (f) *Angolatitan adamastor*, redrawn from Mateus et al. (2011), (g) *Dreadnoughtus schrani* (left, reversed), redrawn from Lacovara et al. (2014), (h) *Patagotitan mayorum* (left, reversed), redrawn from Carballido et al. (2017), (i) *Paralititan stromeri*, redrawn from Smith et al. (2001), (j) *Notocolossus gonzalezparejasi*, redrawn from González Riga et al. (2016). Scale bar equals 20 cm.
longer than those of other giant titanosaurs such as *Dreadnoughtus* (Figure 4g, Lacovara et al. 2014), *Futalognkosaurus* (Calvo 2014), and *Paralititan* (Figure 4i, Smith et al. 2001). Therefore, despite the incomplete nature of the known remains, *Notocolossus* is likely among the largest titanosaurs discovered thus far. Nevertheless, we recommend caution when estimating the body size of one gigantic titanosaurian taxon versus another. Accurate estimation of body size (e.g., total length, mass, volume) is highly problematic in the largest titanosaurs because, with the exceptions of *Dreadnoughtus*, *Futalognkosaurus*, and *Patagotitan*, most of these taxa (e.g., 'Antarctosaurus' giganteus, *Argentinosaurus*, *Paralititan*, *Puertasaurus*, *Notocolossus*) are represented by very incomplete skeletons (Lacovara et al. 2014). Moreover, given the morphological disparity seen in relatively complete, smaller-bodied titanosaurs (e.g., *Diamantinasaurus*, *Epachthosaurus*, *Isisaurus*, *Mendozasaurus*, *Opisthocoelicaudia*, *Rapetosaurus*, *Saltasaurus*), it is probable that different gigantic species also had markedly different anatomical proportions, such as the relative proportions and robusticity of the limb elements, the lengths of the cervical and caudal series, and the distance from the glenoid to the acetabulum. Indeed, this contention has already been borne out, at least to some degree, by the substantially different humeral proportions of the giant titanosaurs *Dreadnoughtus*, *Notocolossus*, *Paralititan*, and *Patagotitan* (Figure 4g-j), with *Notocolossus* and especially *Dreadnoughtus* possessing exceedingly robust humeri, that of *Paralititan* being more slender, and that of *Patagotitan* exhibiting an intermediate condition. Although issues surrounding body size estimation in the largest titanosaurs have been partly ameliorated by recent discoveries of relatively complete skeletons of *Dreadnoughtus*, *Futalognkosaurus*, and *Patagotitan*, additional, similarly complete giant titanosaur specimens will be needed to definitively assess the widely varying dimensions postulated in recent works (e.g., Lacovara et al. 2014, Bates et al. 2015, González Riga et al. 2016, Carballido et al. 2017).

**RADIUS AND ULNA**

Both the radius and ulna are known in the South American titanosaurs *Aeolosaurus rionegrinus*, *Argyrosaurus*, *Bonitasaura*, *Dreadnoughtus*, *Elaltitan*, *Epachthosaurus*, *Mendozasaurus*, *Muyelensaurus*, *Neuquensaurus*, *Patagotitan*, *Quetecsaurus*, *Saltasaurus*, and *Tapuiasaurus*, the former bone is additionally known in *Uberabatitan* and the latter in *Narambuenatitan* and *Pitekunsaurus* (Table I). With the exceptions of a few taxa (e.g., *Dreadnoughtus*, *Elaltitan*) the distal mediolateral breadth of the radius is approximately twice its breadth at midshaft. In saltasaurid titanosaurs (and probably some other neosauropods, Upchurch et al. 2015), the distal radius is also beveled approximately 20° proximolaterally relative to the long axis of the shaft (Wilson 2002, D’Emic 2012).

Most lithostrotians exhibit a proximally elevated olecranon process of the ulna, which constitutes a reversal to the basal sauropodomorph condition (Wilson and Sereno 1998, Galton and Upchurch 2004, Mannion et al. 2013). The ulnae of saltasaurid titanosaurs are further characterized by their stout proportions (Wilson 2002); this condition is developed to an extreme in the saltasaurines *Neuquensaurus* and *Saltasaurus*. Most other titanosaurs have more gracile ulnae, a condition that is especially true for taxa such as *Mendozasaurus* and *Narambuenatitan*. In titanosaurs, and sauropods in general, the anteromedial process of the proximal ulna is usually longer than the anterolateral process.

**CARPUS AND MANUS**

With the possible exceptions of the carpal elements reported in *Argyrosaurus* and *Neuquensaurus*,...
but currently lost (Lydekker 1893, Huene 1929, Otero 2010, Mannion and Otero 2012), titanosaur appears to lack an ossified carpus (Upchurch 1998, Wilson 2002, D’Emic 2012). The metacarpus is completely or nearly completely known in several South American titanosaurian taxa (Aeolosaurus rionegrinus, Antarctosaurus wichmannianus, Argyrosaurus, Choconsaurus, Epachthosaurus, Mendozasaurus, Quetecsaurus, and Rinconsaurus) and partially preserved in a few others (Table I, Figure 5). A number of these forms (e.g., A. rionegrinus, saltasaurines) have fairly stout metacarpals, but in Argyrosaurus (Mannion and Otero 2012) and especially Choconsaurus (Simón et al. 2018) these bones are more elongate and slender, as is also the case in Andesaurus, for which only metacarpals I and V are represented (Calvo and Bonaparte 1991, Mannion and Calvo 2011). In many titanosaur, the ratio of the proximodistal length of metacarpal I to that of metacarpal II or III is 1.0 or greater (Upchurch 1998, Mannion et al. 2013); also, the proximal end of metacarpal V is often subequal in size to that of metacarpal I (D’Emic 2012). Nevertheless, in taxa such as Choconsaurus, Epachthosaurus, and Quetecsaurus, the proximal end of metacarpal V is clearly smaller than that of metacarpal I, with the latter being much more anteroposteriorly elongate (Figure 5b).

Multiple authors have argued that, in Titanosauria or clades therein, most or even all manual phalanges were absent or unossified (e.g. Salgado et al. 1997, Wilson 2002, Apesteguía 2005, Curry Rogers 2005). Among South American titanosaur, this contention is supported by the condition in Argyrosaurus, which is known from a complete, articulated forelimb that nonetheless lacks direct evidence of manual phalanges (Mannion and Otero 2012) (but see Discussion below). Similarly, an exceptionally complete, articulated postcranium of Epachthosaurus (UNPSJB-PV 920) possesses only a single rudimentary phalanx on manual digit IV (Martínez et al. 2004); an identical condition occurs in the holotypic skeleton of the Mongolian titanosaur Opisthocoelicaudia (ZPAL MgD-I/48, Borsuk-Bialynicka 1977). Nevertheless, manual phalanges are well-documented in the Australian titanosaur Diamantinasaurus and Savannasaurus (Hocknull et al. 2009, Poropat et al. 2015, 2016), indicating that the diversity of manual morphologies within Titanosauria was almost certainly greater than is presently appreciated (see Discussion).

THE PELVIC AND HIND LIMB SKELETON OF SOUTH AMERICAN TITANOSAURS

ILIUM

The titanosaurian ilium exhibits remarkable features. The bone is at least partially preserved in many South American taxa, including Antarctosaurus wichmannianus (Huene 1929, Powell 2003), Brasilotitan (Machado et al. 2013), Dreadnoughtus (Lacovara et al. 2014, Ullmann and Lacovara 2016), Epachthosaurus (Martínez et al. 2004), Futalognkosaurus (Calvo et al. 2007a, b), Gondwanatitan (Kellner and de Azevedo 1999), Muyelensaurus (Calvo et al. 2007c), Narambuenatitan (Filippi et al. 2011a), Neuquensaurus (Salgado et al. 2005, Otero 2010), Overosaurus (Coria et al. 2013), Petrobrasaurus (Filippi et al. 2011b), Rinconsaurus (Calvo and González Riga 2003), Rocasaurus (Salgado and Azpilicueta 2000), Saltasaurus (Powell 1992, 2003), and Trigonosaurus (Campos et al. 2005) (Table I). Among these taxa, the largest complete ilia and sacrum belong to Futalognkosaurus. In titanosaur, the pre- and postacetabular processes of the ilium are expanded anteroposteriorly and dorsoventrally (Otero and Vizcaíno 2008, Otero 2010). Although lateral projection of the iliac preacetabular process characterizes titanosaur as a whole, the orientation of this process (i.e., lateral or anterolateral) varies considerably among titanosaurian taxa (Figure 6a, b). For example, the lateral projection of the preacetabular process...
is more pronounced in *Futalognkosaurus*, *Neuquensaurus* (Figure 6a), and *Saltasaurus* than it is in *Dreadnoughtus*, *Epachthosaurus*, *Overosaurus* (Figure 6b), or *Trigonosaurus*. To more rigorously define this character, Salgado et al. (2005) proposed an estimate of the ratio between the distance of the lateralmost point of the pubic peduncle versus that of the preacetabular process. A strongly laterally directed preacetabular process appears to be associated with elongation of the sacral ribs, as seen in, for example, *Neuquensaurus* (MCS-5/16, B.J.G.R. pers. obs.).

In some South American titanosaurs (e.g., *Overosaurus*, *Rinconsaurus*), the portion of the iliac blade dorsal to the pubic peduncle is dorsoventrally deep, whereas in others (e.g., *Saltasaurus*) it is much shallower. Furthermore, the ilia of saltasaurines exhibit a ‘kink’ on the
ventral margin of the preacetabular process (D’Emic 2012). These morphological changes are probably related to locomotor specializations (Wilson and Carrano 1999) and/or the attachment of pelvic musculature (Otero and Vizcaíno 2008). As in Sauropoda more generally, the ischial peduncle of the titanosaurian ilium is only slightly developed (Wilson 2002), and in derived taxa such as saltasaurines, it is often confluent with the remainder of the bone. Conversely, the pubic peduncle is elongate, relatively gracile, and anteroventrally projected. Interestingly, the ilia of multiple titanosaurs, including South American forms such as Dreadnoughtus (Ullmann and Lacovara 2016), Epachthosaurus (Martínez et al. 2004), and Neuquensaurus (Cerda et al. 2012), display well-developed internal camerae, indicating that they were probably pneumatized by diverticula from the respiratory system. Conversely, the ilia of some other titanosaurs (e.g., the Tanzanian basal form Rukwatitan, Gorscak et al. 2014) appear to have been apneumatic.

PUBIS

Among South American titanosaurs, the pubis is at least partially preserved in Andesaurus, Antarctosaurus wichmannianus, ‘Antarctosaurus’ giganteus (Huene 1929), Bonitasaura, Dreadnoughtus, Elalititan, Epachthosaurus, Futalognkosaurus, Gondwanatitan, Mendozaaurus, Muyelensaurus, Narambuenatitan, Neuquensaurus, Notocolossus, Patagotitan, Petrobrasaurus, Rinconsaurus, Rocasaurus, Saltasaurus, and Uberabatitan (Table I, Figure 6c-f). In most of these taxa (e.g., Andesaurus, Elalititan, Futalognkosaurus), it is a short, stout bone, but the pubic blade is slenderer in a few other forms (e.g., Muyelensaurus, Petrobrasaurus). The iliac articular surface of the pubis is anteroposteriorly elongate in titanosaurs (Mannion et al. 2013). The obturator foramen is subcircular in many titanosaurs but oval in Andesaurus, Futalognkosaurus, and Muyelensaurus (Mannion and Calvo 2011). The pubes of several titanosaurs (e.g., Dreadnoughtus, Futalognkosaurus) exhibit a longitudinal ventrolateral ridge that is most strongly developed in taxa such as Neuquensaurus, Saltasaurus, and Uberabatitan.

According to many phylogenetic studies (e.g., Salgado et al. 1997), a typical titanosaurian character is the presence of a pubis that is proximodistally longer than the ischium. This feature seems to be present in most or even all titanosaurs, but in taxa such as Futalognkosaurus, it is developed to an extreme, with the pubis being markedly longer and more robust than the ischium (Figure 6f). Furthermore, in this massive Patagonian titanosaur, the bone has a slightly subcircular and expanded distal end and is strongly thickened distally (MUCPv-323, B.J.G.R. pers. obs.). The pubis of the basal titanosaur Andesaurus possesses a proximodistally elongate ischial articulation (Figure 6e); in most other titanosaurian taxa, by contrast, this articular surface is shorter.

ISCHIUM

The ischium is known in many South American titanosaurs, being especially well-preserved in taxa such as Andesaurus, Bonitasaura, Dreadnoughtus, Futalognkosaurus, Muyelensaurus, and Saltasaurus (Table I, Figure 6c-f). The titanosaurian ischium is a short bone with a relatively broad, plate-like blade (Salgado et al. 1997, Otero 2010); this morphology as well as the absence of emargination distal to the pubic articulation are typical of the clade (Wilson 2002, Díez Díaz et al. 2016). A mediolaterally compressed iliac articular surface also appears to be a titanosaurian character (Mannion and Calvo 2011). The pubic peduncle is anteroposteriorly elongate in forms such as Aeolosaurus rionegrinus, Antarctosaurus
wichmannianus, and Muyelensaurus (Figure 6d), but shorter in others (e.g., Andesaurus [Figure 6e], Rinconsaurus, Saltasaurus, Triunfosaurus [de Souza Carvalho et al. 2017]). In many titanosaurian taxa, the acetabular margin of the ischium is strongly concave in lateral view, such that the pubic articular surface forms a proximodorsal projection (D’Emic 2012, Mannion et al. 2013, Figure 6d). The ischial blade is slender in taxa such as A. rionegrinus, Andesaurus, Bonitasaura, Futalognkosaurus, Gondwanatitan, and Muyelensaurus compared to the condition in other forms such as Rinconsaurus and the saltasaurines Rocasaurus and Saltasaurus.

FEMUR

The femur is at least partially preserved in 25 species of South American titanosaurs; after the humerus, it is the most frequently recovered appendicular bone of these sauropods (Table I, Figure 7). The femur is remarkably slender in Atacamatitan (Figure 7a), and to a lesser degree in ‘Antarctosaurus’ giganteus (Figure 7j), Bonatitan (Figure 7k), Mendozaaurus (Figure 7g), Patagotitan (Figure 7h), Petrobrasaurus (Figure 7f), and Rinconsaurus (Calvo and González Riga 2003, fig. 3c), but considerably more robust in taxa such as Dreadnoughtus (Figure 7i), Futalognkosaurus (Calvo 2014), and the saltasaurines Neuquensaurus (Figure 7b, e), Rocasaurus (Salgado and Azpilicueta 2000, fig. 9a), and Saltasaurus (Figure 7e); titanosaurs such as Epachthosaurus (Figure 7d) and Traukutitan (Figure 7l) exhibit an intermediate condition. As with the humerus (see above), the markedly differing robusticities of the femur in giant titanosaur such as ‘A.’ giganteus, Bonitasaura, Dreadnoughtus, Epachthosaurus, Mendozaaurus) but more proximally in others (e.g., Elaltitan, Neuquensaurus, Patagotitan). It is also more prominent in taxa such as Dreadnoughtus (Ullmann and Lacovara 2016) and Patagotitan (Carballido et al. 2017) than it is in others such as saltasaurines (e.g., Figure 7b, e). Saltasaurines exhibit a longitudinal ridge on the anterior surface of the femoral shaft (Otero 2010, D’Emic 2012).

Another important femoral character is the orientation of the long axis of the distal condyles in anterior or posterior view; in many South American titanosaurs, the distal condyles are beveled 10° dorsomedially (Wilson 2002, 2006, D’Emic 2012), though there are exceptions to this condition.
Figure 6 – Morphological comparison of pelvic elements of South American (Argentinean) titanosaurs. Ventral view of sacrum and ilia of (a) *Neuquensaurus australis*, MCS 5/16, redrawn from Salgado et al. (2005) and (b) *Overosaurus paradosorum*, MAUPv-CO-439, redrawn from Coria et al. (2013). Lateral view of left pubis and ischium of (c) *Dreadnoughtus schrani*, MPM-PV 1156, redrawn from Ullmann and Lacovara (2016), (d) *Muyelensaurus pecheni*, MRS-Pv 88 (reversed), redrawn from Calvo et al. (2007c), (e) *Andesaurus delgadoi*, MUCPv-132, redrawn from Mannion and Calvo (2011), (f) *Futalognkosaurus dukei*, MUCPv-323 (reversed), redrawn from Calvo et al. (2007b). Not to scale.

(e.g., *Dreadnoughtus*, Ullmann and Lacovara 2016, *Traukutitan*, Juárez Valieri and Calvo 2011). Moreover, in the femora of many derived titanosaurs, the fibular condyle projects further distally than the tibial condyle (González Riga et al. 2018), and in saltasaurines, these condyles extend onto the anterior surface of the shaft (Wilson 2002, Ullmann and Lacovara 2016).

**Tibia and Fibula**

Both the tibia and fibula are preserved in *Aeolosaurus rionegrinus*, *Antarctosaurus wichmannianus*, *Bonatitan*, *Bonitasaura*, *Dreadnoughtus*, *Elaltitan*, *Epachthosaurus*, *Laplatasaurus* (Gallina and Otero 2015), *Mendozasaurus*, *Muyelensaurus*, *Neuquensaurus*, *Saltasaurus*, and *Uberabatitan*, one or the other of these bones is known in a few other taxa as well (Table I, Figure 8a-l). Like other appendicular elements, the tibia and fibula of South American titanosaurs vary in proportions from relatively gracile to robust. Saltasaurines (Figure 8e) have very stout tibiae with a prominent cnemial crest, whereas in other taxa such as *A. wichmannianus*, *Epachthosaurus* (Figure 8c), and *Mendozasaurus* (Figure 8a) the tibia is considerably slenderer and the crest is weakly developed; other forms such as *Dreadnoughtus* (Figure 8d), *Futalognkosaurus* (Calvo 2014), and *Laplatasaurus* (Figure 8b) exhibit an intermediate condition. The extent of the cnemial fossa on the proximal end of the tibia is also highly variable among South American titanosaurs (Gallina and Otero 2015). In lithostrotian titanosaurs, the mediolateral width...
of the distal end of the tibia is at least twice the diameter of the bone at midshaft (Wilson 2002, Mannion et al. 2013).

The fibulae of South American titanosaur exhibit variability in aspects such as the anteroposterior width of the proximal end, the robusticity and straightness of the shaft, and the development, location, and morphology of the lateral trochanter. The proximal end is anteroposteriorly broad in taxa such as *Dreadnoughtus* (Figure 7 – Morphological comparison of South American titanosaur femora. (a) Right femur in anterior view of *Atacamatitan chilenensis*, SGO-PV-961g, redrawn from Kellner et al. (2011), (b) left femur in posterior view of *Neuquensaurus australis*, MLP-CS 1118, redrawn from Otero (2010), (c) right femur in anterior view of ‘*Neuquensaurus robustus’*, MCS-9, redrawn from Otero (2010), (d) right femur in posterior view (reversed) of *Epachthosaurus sciuttoi*, UNPSJB-PV 920, redrawn from Martinez et al. (2004), (e) right femur in posterior view of *Saltasaurus loricatus*, PVL 4017-79, redrawn from Powell (1992), (f) left femur in posterior view of *Petrobrasaurus puestohernandezii*, MAU-PV-PH 449/8, redrawn from Filippi et al. (2011b), (g) left femur in posterior view of *Mendoza saur neguyelap*, IANIGLA-PV 073/4, redrawn from González Riga et al. (2018), (h) right femur in posterior view (reversed) of *Patagotitan mayorum*, MPEF-PV 4400/26, redrawn from Carballido et al. (2017), (i) left femur in posterior view of *Dreadnoughtus schrani*, MPM-PV 1156, redrawn from Ullmann and Lacovara (2016), (j) right femur in posterior view (reversed) of *Antarctosaurus* giganteus, drawn from unpublished photo by L.D.O.D., (k) left femur in anterior view (reversed) of *Trau kuttin eoca nata*, MUCPv-204, redrawn from Juárez Valieri and Calvo (2011). Not to scale.
and *Uberabatitan* (Figure 8g) but much narrower in many others (e.g., *Epachthosaurus, Laplatasaurus, Mendozasaurus*, Figure 8f, h, j). Similarly, the shaft is relatively robust in *Dreadnoughtus, Uberabatitan*, and saltasaurines (e.g., Figure 8k) but more gracile in *Laplatasaurus* and *Mendozasaurus*. Interestingly, despite the undoubtedly gigantic size of the animal, the fibula of *Argentinosaurus* (initially described as a tibia, Bonaparte and Coria 1993) is considerably more gracile and slender-shafted than the stouter fibulae of taxa such as *A. rionegrinus, Dreadnoughtus, Elaltitan, Laplatasaurus, Uberabatitan*, and saltasaurines. Nevertheless, at a colossal 1.55 m in length, the *Argentinosaurus* fibula remains the longest yet known for a titanosaur by a considerable margin (see Lacovara et al. 2014, table 1). When observed in lateral view, the fibular shaft is slightly sigmoid in *Dreadnoughtus, Epachthosaurus, and Saltasaurus* (Powell 1992) but generally straighter in forms such as *Argentinosaurus, Laplatasaurus, Mendozasaurus*, and *Neuquensaurus*. The lateral trochanter is very well-developed in taxa such as *Dreadnoughtus, Laplatasaurus, Neuquensaurus*, and *Uberabatitan* but less so in others (e.g., *Epachthosaurus, Mendozasaurus*). It is placed slightly proximal to midshaft in most South American titanosauras but more distally in selected taxa from other landmasses (e.g., the Madagascan titanosaur *Rapetosaurus*, Figure 8i). The lateral trochanter takes the form of a single tuberosity in most taxa (e.g., *Dreadnoughtus, Laplatasaurus*) but is comprised of two subparallel ridges in *Epachthosaurus* (Martínez et al. 2004). In *Laplatasaurus, Uberabatitan*, and perhaps a few other titanosauras there is a marked concavity immediately posterior to the lateral trochanter (Salgado and de Souza Carvalho 2008, Gallina and Otero 2015).

### TARSUS AND PES

The tarsus of titanosauras is represented exclusively by the astragalus, with the possible exception of *Elaltitan* for which an ossified calcaneum has been reported (Mannion and Otero 2012); a calcaneum was also described for *Neuquensaurus* by Huene (1929) but this bone is now lost (Otero 2010). Among South American titanosauras, the astragalus is known in only 11 taxa: *Aeolosaurus rionegrinus, Bonatitan, Bonitasaura, Dreadnoughtus, Elaltitan, Epachthosaurus, Mendozasaurus, Muyelensaurus, Neuquensaurus, Notocolossus*, and *Uberabatitan* (Table 1, Figure 8m-p). In several of these forms (e.g., *Notocolossus, Uberabatitan*, Figure 8n, p) it is proximodistally tall and pyramidal in shape, but in others (e.g., *Dreadnoughtus* [Figure 8m], *Elaltitan* [Figure 8o], *Epachthosaurus, Mendozasaurus*) it is lower. D’Emic (2012) regarded a mediolaterally narrow astragalus as a synapomorphy of *Saltasauridae*. The element identified as the calcaneum of *Elaltitan* is amorphous and globular, consistent in appearance with the calcanea of non-titanosaurian sauropods (Figure 8q, Mannion and Otero 2012).

Complete pedes are extremely scarce in the fossil record of titanosauras (Table I, Figure 9, see also Curry Rogers 2005, González Riga 2011), and indeed, sauropods as a whole (McIntosh 1990). Intrinsic factors related to the large body dimensions of these dinosaurs coupled with the relatively small size and fragility of their skull bones, posterior caudal vertebrae, and manual and pedal elements evidently led to the early disarticulation, and therefore loss, of these comparatively diminutive bones during the biostratigraphic stage of necrokinesis (González Riga et al. 2008b). In fact, of the well over 70 valid titanosaurian taxa recognized at present (Wilson et al. 2016, M.C.L. pers. obs.), only three are known from complete and articulated pedes: *Opisthocoelicaudia* from Mongolia (Figure 9a, ZPAL MgD-I/48, Borsuk-
Figure 8 – Morphological comparison of titanosaurian tibiae (a-e), fibulae (f-l), and tarsals (m-q). South American taxa denoted with an asterisk. (a)-(e), right tibiae in lateral view: (a) *Mendozasaurus neguyelap*, IANIGLA-PV 073/2, redrawn from González Riga et al. (2018), (b) *Laplatasaurus araukanicus*, MLP-CS 1128, redrawn from Gallina and Otero (2015), (c) *Epachthosaurus sciuttoi*, UNPSJB-PV 920, redrawn from Martínez et al. (2004), (d) *Dreadnoughtus schrani*, MPM-PV 1156, redrawn from Ullmann and Lacovara (2016), (e) *Saltasaurus loricatus*, PVL 4017-84, redrawn from Powell (1992). (f)-(l), left fibulae in lateral view: (f) *Laplatasaurus araukanicus* (reversed), MLP-CS 1127, redrawn from Gallina and Otero (2015), (g) *Uberabatitan ribeiroi*, CPP-1107-UrHo, redrawn from Salgado and de Souza Carvalho (2008), (h) *Epachthosaurus sciuttoi*, UNPSJB-PV 920, redrawn from Martínez et al. (2004), (i) *Rapetosaurus krausei* (reversed), FMNH PR 2209, redrawn from Curry Rogers (2009), (j) *Mendozasaurus neguyelap*, IANIGLA-PV 074/3, redrawn from González Riga (2003), (k) *Neuquensaurus robustus* *, MLP-CS 1265, redrawn from Otero (2010), (l) *Dreadnoughtus schrani*, MPM-PV 1156, redrawn from Ullmann and Lacovara (2016). (m)-(q), tarsal elements in various views: (m) left astragalus in proximal view of *Dreadnoughtus schrani*, MPM-PV 1156, redrawn from Ullmann and Lacovara (2016), (n) right astragalus in anterior view of *Notocolossus gonzalezparejiasi*, UNCUYO-LD 302, redrawn from González Riga et al. (2016), (o) right astragalus in anterior view of *Elaltitan lilloi*, PVL 4628, redrawn from Mannion and Otero (2012), (p) left astragalus in anterior view of *Uberabatitan ribeiroi*, CPP-1082-UrHo, redrawn from Salgado and de Souza Carvalho (2008), (q) calcaneum adhered to medial surface of distal fibula of *Elaltitan lilloi*, PVL 4628, redrawn from Mannion and Otero (2012). Not to scale.
Bialynicka 1977) and Epachthosaurus (Figure 9b, UNPSJB-PV 920, Martínez et al. 2004) and Notocolossus (Figure 9g, UNCUYO-LD 302, González Riga et al. 2016) from Argentina. Two additional titanosaurs with complete, articulated hind feet may be added to this select group: the Agua de Padrillo taxon (UNCUYO-LD 313, González Riga et al. 2015), which preserves both complete pedes and is currently undergoing laboratory preparation, and the La Invernada taxon (MUCPV-1533, González Riga et al. 2008a), which preserves the complete left fore- and hind limbs and is presently under study (Figures 1a, b, 9c, d). Preliminary analyses suggest that both the Invernada and Padrillo forms correspond to new genera (B.J.G.R. pers. obs.). Pedal elements are known for many other titanosaurs, including other South American taxa (see Table I), but none of these preserve the pes completely and in articulation (though a few other specimens approach this condition, e.g., ?Alamosaurus NMMNH P-49967, Figure 9e, D’Emic et al. 2011).

In sauropods, a progressive reduction in both the number and length of the pedal phalanges has been previously documented and is the most apparent evolutionary trend in the structure of the hind foot in these herbivorous dinosaurs (Bonnan 2005, González Riga et al. 2008a, 2016). For example, in the basal eusauroposes Shunosaurus (Zhang 1988) and Omeisaurus (He et al. 1988), a total of 12 pedal phalanges are described. Titanosaurs, by contrast, have fewer pedal phalanges. The possible basal titanosaur Epachthosaurus has a pedal phalangeal formula of 2-2-3-2-0 (nine phalanges total, Martínez et al. 2004), and an even more reduced formula of 2-2-2-2-0 (eight phalanges total) occurs in the Padrillo (UNCUYO-LD 313) and Invernada (MUCPv-1533) taxa, Notocolossus (González Riga et al. 2016), and Mendozaaurus (González Riga et al. 2018). Opisthocoelicaudia was originally described as having a pedal phalangeal formula of 2-2-2-1-0 (Borsuk-Bialynicka 1977), but the shape of the distal condyle of the first phalanx of digit IV suggests the presence of a second ossified phalanx in this digit. Because of this, we herein interpret that this taxon had a phalangeal formula of 2-2-2-2-0, as is the case in all other derived titanosaurs for which the pes is completely represented.

Another relevant aspect of sauropod pedal structure is the development of the distal articular facets of the metatarsals. Indiplocodicids, for example, the articular facets are strongly convex and extend onto the dorsal (= anterior) face of the metatarsals (Bonnan 2005). This condition is clearly visible in Barosaurus (AMNH 6341, B.J.G.R. pers. obs.), in which the distal end of metatarsal I is strongly convex, indicating a wide range of mobility of phalanx I-1. A similar but less pronounced case is observed in Apatosaurus (CM 3018, phalangeal formula 2-3-3-2-1, B.J.G.R. pers. obs.), suggesting that the elevated mass of this taxon may have led to a reduction in the mobility of its phalanges in comparison with the more lightly-built diplodocine diplodocids Barosaurus and Diplodocus. In the macronarian Camarasaurus, two distal articular facets are present in metatarsals I and II (YPM 1901, B.J.G.R. pers. obs.), in accordance with the well-developed pedal phalanges of this taxon (phalangeal formula 2-3-2-2-1, McIntosh et al. 1996).

In titanosaurs, by contrast, the distal articular facets of the metatarsals are less developed than in other sauropods. In Mendozaaurus, for instance, these facets are only slightly convex and only some of them extend onto the dorsal surface of the metatarsal in question (González Riga et al. 2018). An extreme case is observed in Notocolossus, the metatarsals of which have nearly flat distal articular facets, indicating reduced mobility of the digits (González Riga et al. 2016, B.J.G.R. pers. obs.). Interestingly, unlike other titanosaurs, the pedal unguals of this gigantic taxon are small, blunt, and amorphous; although there is some possibility that this condition is pathologic (González Riga et al. 2018).
et al. 2016), it is also consistent with the limited development of the distal articular facets of the metatarsals.

A distinctive character of some titanosauriform pedal unguals is the presence of an elongate tuberosity on the ventral (= plantar) surface. Among South American titanosaurians this structure is present in Bonatitan (Salgado et al. 2015), Brasilotitan (Machado et al. 2013), Dreadnoughtus (Ullmann and Lacovara 2016), Mendozaasaurus (González Riga et al. 2018), the La Invernada taxon (González Riga et al. 2008a), and probably others.

González Riga et al. (2016) preliminarily recognized two primary titanosaurian pedal skeletal morphotypes, which they termed ‘long-footed’ (where, as in non-titanosaurian sauropods, the first four metatarsals exhibit a significant increase in length and a decrease in robusticity from medial to lateral) and ‘short-footed’ (with metatarsals that are all roughly the same length). Further study has led us to recognize considerable variation within González Riga et al.’s (2016) long-footed morphotype, revealing a broad diversity in form (Figure 9). Moreover, within Titanosauria, there is no clear correlation between body size and pedal osteology; instead, the differing hind foot architecture of various titanosaurians is probably more intimately related to evolutionary trends seen within different lineages.

The short-footed pedal morphotype is a massive structure that has thus far been observed only in Notocolossus (Figure 9g). This giant sauropod exhibits the lowest differences between the lengths of the metatarsals of any titanosaurian taxon yet discovered (for instance, the ratio of the length of metatarsal III to that of metatarsal I is only 1.14, González Riga et al. 2016). Moreover, in Notocolossus, differences in robusticity between metatarsal I and metatarsals II–V are less pronounced than in other titanosaurians. This may be quantified using the Metatarsal Robustness Index (MtRI), which is herein defined as the minimum mediolateral breadth of metatarsals II–V divided by that of metatarsal I. In Notocolossus, the MtRI is greater than 0.70 in metatarsals II–V. Moreover, the non-ungual phalanges are relatively long and wide in relation to the metatarsals, and both the size and shape of the unguals are unique, as noted above and as was described by González Riga et al. (2016).

As mentioned above, several variances may be observed within the long-footed titanosaurian morphotype (Figure 9a-f). The medium-sized derived titanosaur Opisthocoelicaudia (body length ~12 m, femoral length 1.39 m, Borsuk-Bialynicka 1977) has a relatively short, compact metatarsus, very short non-ungual phalanges, and large unguals (Figure 9a). Metatarsal V is shorter than metatarsal I (metatarsal V length/metatarsal I length = 0.93, Borsuk-Bialynicka 1977). Metatarsal III is the longest, in contrast to other titanosaurians (metatarsal III length/metatarsal I length = 1.33, González Riga et al. 2016) (Table II).

The other long-footed titanosaurians analyzed herein are Epachthosaurus (Figure 9b, Martínez et al. 2004), Mendozaasaurus (Figure 9f, González Riga et al. 2018), the unnamed Invernada and Padrillo taxa (Figure 9c, d, González Riga et al. 2008a, 2015), and NMMNH P-49967, an isolated pes provisionally attributed to Alamosaurus (Figure 9e, D’Emic et al. 2011). In contrast to Opisthocoelicaudia, in all of these taxa, metatarsal V is longer than metatarsal I and metatarsals I–IV show a progressive increase in length; because of this, metatarsal IV is the longest (the length ratio of metatarsal IV/metatarsal I is 1.39–1.57 in these taxa) (Table II). The unguals are relatively large in relation to metatarsal length. Of all titanosaurians for which the pes is completely known, Epachthosaurus is unique in retaining nine phalanges. This accords with the basal position of this genus that is frequently recovered by phylogenetic analyses (e.g., Carballido et al. 2017) and the hypothesis of progressive reduction of the...
Figure 9 – The best-preserved pedes of titanosaurian sauropods. South American (Argentinean) taxa marked with an asterisk. (a) *Opisthocoelicaudia skarzynskii*, ZPAL MgD-I/48, right pes, redrawn from Borsuk-Bialynicka (1977), (b) *Epachthosaurus sciuttoi*, UNPSJB-PV 920, right pes, redrawn from Martínez et al. (2004), (c) La Invernada taxon*, MUCPv-1533, left pes, redrawn from González Riga et al. (2008a), (d) Agua del Padrillo taxon*, UNCUYO-LD 313, left pes, redrawn from González Riga et al. (2015), (e) ?*Alamosaurus sanjuanensis*, NMMNH P-49967, right pes, redrawn from D’Emic et al. (2011), (f) *Mendozasaurus neguyelap* *, IANIGLA-PV 077/1–10, 078/1–2, 079, right pes, redrawn from González Riga et al. (2018), (G) *Notocolossus gonzalezparejasi*, UNCUYO-LD 302, right pes, redrawn from González Riga et al. (2016). Scale bars equal 10 cm.
pedal phalanges within Titanosauria (González Riga et al. 2008a, 2016).

Among this second group of long-footed titanosaurs, there are, as yet, no definitive correlations between pedal structure and body size, though some possible trends are evident. In the smaller-bodied taxa within this group (Epachthosaurus and the Invernada and Padrillo taxa, with body lengths of up to approximately 10 m, Martínez et al. 2004, González Riga et al. 2008a), metatarsals IV and V are relatively slender (Figure 9b-d). In contrast, in ?Alamosaurus (NMMNH P-49967) and Mendozasaurus, metatarsal V is relatively robust and longer than metatarsals I and II (Figure 9e, f). Both of these latter animals were very large: in ?Alamosaurus, metatarsal IV is 29.1 cm in length, and the femoral length of this individual has been estimated at 1.6–2.0 m (D’Emic et al. 2011, González Riga et al. 2016). Similarly, metatarsal III of an undescribed specimen of Mendozasaurus (UNCUYO-LD 356) is 29.2 cm in length, and as such, the individual in question was probably comparable in size to that represented by NMMNH P-49967. Further discoveries of relatively complete titanosaurian pedes are needed to further evaluate the potential relationships between body size and pedal morphology discussed herein.

### DISCUSSION

The skeletal structure of sauropods has traditionally been interpreted as being relatively conservative in comparison to that of other dinosaurs. This is documented in, for example, Wilson and Curry Rogers’ (2005) summary of the history of sauropod discoveries. In an early stage of the study of these iconic herbivorous dinosaurs, Romer (1968) lamented the difficulty in achieving a classification of sauropods due to their relatively incomplete fossil record. Thankfully, however, the sauropod record has improved dramatically in recent decades, leading to significant advances in knowledge of the anatomy, evolution, and paleobiology of these animals. As Wilson and Curry Rogers (2005) pointed out, “The improvement in our understanding of sauropod phylogeny is the result of an improved sauropod fossil record.”

Many sauropod species are primarily defined on anatomical characters derived from the presacral, sacral, and/or anterior caudal vertebrae, and therefore, many authors have justifiably

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### TABLE II

Proximodistal lengths and relative proportions of metatarsals of titanosaurian sauropods with the most completely preserved pedes. Roman numerals indicate metatarsal number. Columns I–V indicate lengths in mm; remaining columns are length ratios of various metatarsals.

<table>
<thead>
<tr>
<th>Taxon</th>
<th>Specimen</th>
<th>I</th>
<th>II</th>
<th>III</th>
<th>IV</th>
<th>V</th>
<th>II / I</th>
<th>III / I</th>
<th>IV / I</th>
<th>V / I</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Opisthocoelicaudia skarzynskii</em></td>
<td>ZPAL MgD-I/48</td>
<td>150</td>
<td>180</td>
<td>200</td>
<td>180</td>
<td>140</td>
<td>1.20</td>
<td>1.33</td>
<td>1.20</td>
<td>0.93</td>
<td>Borsuk-Bialynicka (1977)</td>
</tr>
<tr>
<td><em>Epachthosaurus sciuttoi</em></td>
<td>UNPSJB-PV 920</td>
<td>125</td>
<td>153</td>
<td>177</td>
<td>185</td>
<td>153</td>
<td>1.22</td>
<td>1.41</td>
<td>1.48</td>
<td>1.22</td>
<td>Martínez et al. (2004)</td>
</tr>
<tr>
<td>Padrillo taxon</td>
<td>UNCUYO-LD 313</td>
<td>109</td>
<td>138</td>
<td>146</td>
<td>152</td>
<td>130</td>
<td>1.26</td>
<td>1.33</td>
<td>1.39</td>
<td>1.19</td>
<td>González Riga et al. (2016)</td>
</tr>
<tr>
<td>Invernada taxon</td>
<td>MUCPv-1533</td>
<td>120</td>
<td>137</td>
<td>168</td>
<td>172</td>
<td>127</td>
<td>1.14</td>
<td>1.40</td>
<td>1.43</td>
<td>1.05</td>
<td>González Riga et al. (2008)</td>
</tr>
<tr>
<td>?<em>Alamosaurus sanjuanensis</em></td>
<td>NMMNH P-49967</td>
<td>195</td>
<td>245</td>
<td>270</td>
<td>291</td>
<td>281</td>
<td>1.25</td>
<td>1.38</td>
<td>1.49</td>
<td>1.44</td>
<td>D’Emic et al. (2011)</td>
</tr>
<tr>
<td><em>Mendozasaurus neguyelap</em></td>
<td>IANGLA-PV 077</td>
<td>135</td>
<td>158</td>
<td>181</td>
<td>212</td>
<td>169</td>
<td>1.17</td>
<td>1.34</td>
<td>1.57</td>
<td>1.25</td>
<td>González Riga et al. (2018)</td>
</tr>
</tbody>
</table>
focused much of their attention on the axial skeleton (e.g., Bonaparte 1999, Wilson 1999, 2012). Knowledge of other regions of the sauropod skeleton, especially the skull, manus, and pes, has lagged behind understanding of vertebral anatomy, and as such, the extent of morphological variability in these parts of the skeleton has not been as thoroughly characterized. Nevertheless, there is no definitive evidence that certain structures changed significantly more than others through sauropod evolutionary history. As more discoveries have been made, it has become apparent that some parts of the sauropod skeleton deserve more attention from researchers than they had previously been afforded. This is certainly the case as regards the appendicular skeleton of titanosaurs.

At first glance, the appendicular anatomy of titanosaurs may appear fairly homogeneous. However, as more well-preserved specimens have come to light, it has become clear that there is considerable variation in the size and morphology of the girdle and limb elements within the clade. Accordingly, a significant number of appendicular skeletal characters have been incorporated into recent phylogenetic analyses of Titanosauria and more inclusive clades such as Titanosauriformes, Macronaria, and Neosauropoda (e.g., Curry Rogers 2005, D’Emic 2012, Mannion et al. 2013, González Riga et al. 2016, 2018, Gorscak and O’Connor 2016, Carballido et al. 2017, Sallam et al. 2018). Although the percentages of appendicular skeletal characters in these analyses have varied considerably, the absolute number of characters has generally increased over the years. For example, in a study of the evolutionary history of Titanosauriformes, D’Emic (2012) included 51 appendicular characters out of a total of 119 (42%), whereas in their analysis of the relationships of Notocolossus, González Riga et al. (2016) modified the dataset of Carballido and Sander (2014) to include 119 appendicular features, comprising 34% of the total.

Here, we follow the node-based phylogenetic definition of Titanosauria proposed by Salgado et al. (1997) and subsequently modified by Wilson and Upchurch (2003). Many titanosaurian clades exhibit considerable diversity. One of these appears to be Lognkosauria, which is defined as the most recent common ancestor of Mendozaasaurus neguyelap and Futalognkosaurus dukei and all descendants (Calvo et al. 2007a). Whereas prior studies (e.g., González Riga and Ortiz David 2014, González Riga et al. 2016) restricted Lognkosauria to two species, more recent analyses by Carballido et al. (2017) and González Riga et al. (2018) have suggested a more diverse clade that also includes Argentinosaurus, Patagotitan, and possibly Drusilasaura, Notocolossus, Puertasaurus, and/or Quetecsaurus.

In previous phylogenetic analyses, Titanosauria has been diagnosed by various appendicular synapomorphies, some of which were initially proposed for the titanosaurian subclades Titanosauroidea or Titanosauridae (which have since been abandoned due to the invalid status of the genus Titanosaurus, Wilson and Upchurch 2003). A review of existing phylogenetic analyses of Titanosauria demonstrates that the clade has not always been supported by the same suite of appendicular skeletal characters; in other words, there is no universal agreement among researchers as to which morphologies of the girdles and limbs are diagnostic of Titanosauria. In recent years, phylogenetic studies have included additional taxa and characters, and as a result, some previously-proposed titanosaurian synapomorphies are now thought to characterize either more inclusive or less inclusive clades. The appendicular skeletal character states that were proposed as synapomorphies of Titanosauria and several of its subclades in the
analyses of D’Emic (2012), Mannion et al. (2013), and González Riga et al. (2016) are summarized in Table III. Other recent, comprehensive phylogenetic analyses of titanosaurs include, among others, those of Gorscak and O’Connor (2016), Poropat et al. (2016), Carballido et al. (2017), Gorscak et al. (2017), González Riga et al. (2018), and Sallam et al. (2018).

Titanosauria

The clade Titanosauria has historically been supported primarily by vertebral characters, with appendicular synapomorphies playing a secondary role. For example, Salgado et al. (1997) postulated only one appendicular synapomorphy for Titanosauria: the presence of a pubis that is considerably longer than the ischium (Salgado et al. 1997:character 24). This feature is unknown in some titanosaurs and therefore has an ambiguous distribution, as seen, for example, in the dataset of Salgado et al. (2015).

Wilson (2002), in an extensive study of sauropod phylogeny, proposed four appendicular synapomorphies of Titanosauria: crescentic sternal plates (his character 158), prominent ulnar olecranon process (reversal) (character 168), ischial blade plate-like, no emargination distal to pubic peduncle (character 193), and distal tibia expanded mediolaterally to twice midshaft breadth (character 205). Subsequently, D’Emic (2012), in a large-scale analysis of Titanosauriformes, recovered only a single appendicular synapomorphy of Titanosauria, character 193 of Wilson’s (2002) matrix, referring to the morphology of the ischial blade. In D’Emic’s (2012) analysis, a pubis considerably longer than the ischium (the lone appendicular synapomorphy of Titanosauria according to Salgado et al. 1997, their character 24) is regarded as diagnostic of a more inclusive clade that includes Sauroposeidon + (Tastavinsaurus + (Euhelopodidae + (Chubutisaurus + Titanosauria)))) (D’Emic 2012:character 102).

In the phylogenetic analysis of Carballido et al. (2011), Titanosauria was supported by one unambiguous appendicular synapomorphy: the absence of a strongly developed ventromedial process on the scapula (character 202, state 0). According to Carballido et al. (2011), this feature is present only in titanosaurian outgroups (e.g., Chubutisaurus, Ligabuesaurus, Wintonotitan).

In the phylogenetic study of González Riga and Ortiz David (2014), Titanosauria was supported by two unambiguous appendicular synapomorphies: absence of well-developed distal phalangeal articular facets on metacarpals (character 71, state 1), and humerus/femur length ratio less than 0.9 (character 77, state 1). Both of these traits are absent in the non-titanosaurian titanosauriform Ligabuesaurus (Bonaparte et al. 2006). In their revision of another non-titanosaurian titanosauriform, Chubutisaurus, Carballido et al. (2011:104) estimated a humerus/femur ratio of 0.86 for this taxon, proposing that a value of less than 0.8 was characteristic of titanosaurs.

The absence of ossified manual phalanges was proposed as a synapomorphy of Opisthocoelicaudinae by Wilson (2002:character 181, state 2), although Opisthocoelicaudia possesses at least one vestigial phalanx on manual digit IV (Borsuk-Bialynicka 1977:31). Previously, Salgado et al. (1997) had proposed this morphology as diagnostic of their ‘Titanosauridae’ (a clade that is largely similar to what is now known as Lithostrotia). Salgado et al. (1997) indicated that the absence of manual phalanges should be evaluated based on the morphology of the distal articular facets of the metacarpals, due to the likelihood that such phalanges could easily be lost due to the taphonomic process of necrokinesis. Similarly, Giménez (1992) proposed to examine the distal ends of metacarpals to assess the presence of manual phalanges, since in several derived titanosaurs the metacarpals exhibit roughened, flattened distal surfaces rather than convex articular
TABLE III  
Proposed synapomorphies of the appendicular skeleton within Titanosauria and some of its subclades according to selected recent works (D’Emic 2012, Mannion et al. 2013, González Riga et al. 2016). Synapomorphies are ordered by anatomical region. (*) Description of synapomorphy modified from original source; (#) synapomorphy obtained from a published phylogenetic analysis but not explicitly mentioned previously.

<table>
<thead>
<tr>
<th>Proposed synapomorphy</th>
<th>Diagnostic of</th>
<th>Source</th>
<th>Character originally proposed by</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scapula, ventral margin with well-developed ventromedial process (#)</td>
<td>Titanosauria</td>
<td>González Riga et al. 2016</td>
<td>Carballido et al. 2011</td>
</tr>
<tr>
<td>Coracoid proximodistally long*</td>
<td>Lithostrotia</td>
<td>D’Emic 2012</td>
<td>Wilson 2002</td>
</tr>
<tr>
<td>Sternal plate length ≥65–70% humerus length (*)</td>
<td>Lithostrotia</td>
<td>D’Emic 2012; Mannion et al. 2013</td>
<td>McIntosh 1990; Upchurch 1998</td>
</tr>
<tr>
<td>Humerus length less than 80% femur length (*)</td>
<td>Saltasauridae</td>
<td>D’Emic 2012</td>
<td>Wilson 2002</td>
</tr>
<tr>
<td>Humeral deltopectoral crest strongly expanded distally (*)</td>
<td>Saltasauridae</td>
<td>D’Emic 2012</td>
<td>Wilson 2002</td>
</tr>
<tr>
<td>Humerus with strong posterolateral bulge around level of deltopectoral crest</td>
<td>Saltasauridae</td>
<td>D‘Emic 2012</td>
<td>D’Emic 2012</td>
</tr>
<tr>
<td>Humeral radial and ulnar condyles divided distally (*)</td>
<td><em>Alamosaurus</em> + <em>Saltasaurini</em></td>
<td>D’Emic 2012</td>
<td>Wilson 2002</td>
</tr>
<tr>
<td>Anterior surface of distal lateral condyle of humerus undivided</td>
<td>Lithostrotia</td>
<td>Mannion et al. 2013</td>
<td>D’Emic 2012</td>
</tr>
<tr>
<td>Radius distal end beveled ~20° proximolaterally relative to shaft</td>
<td>Saltasauridae</td>
<td>D’Emic 2012</td>
<td>Wilson 2002</td>
</tr>
<tr>
<td>Prominent ulnar olecranon process, projecting well above proximal articulation (*)</td>
<td>Lithostrotia</td>
<td>Mannion et al. 2013</td>
<td>McIntosh 1990, Wilson and Sereno 1998</td>
</tr>
<tr>
<td>Metacarpal I:metacarpal II/III proximodistal length ratio ≥1.0</td>
<td>Lithostrotia</td>
<td>Mannion et al. 2013</td>
<td>Upchurch 1998</td>
</tr>
</tbody>
</table>
### TABLE III (continued)

<table>
<thead>
<tr>
<th>Proposed synapomorphy</th>
<th>Diagnostic of</th>
<th>Source</th>
<th>Character originally proposed by</th>
</tr>
</thead>
<tbody>
<tr>
<td>Metacarpal V proximal end subequal in size to that of metacarpal I</td>
<td>Saltasauridae</td>
<td>D’Emic 2012</td>
<td>D’Emic 2012</td>
</tr>
<tr>
<td>Ilium with kink on ventral margin of precacetabular process</td>
<td>Alamosaurus + ‘Saltasaurini’</td>
<td>D’Emic 2012</td>
<td>D’Emic 2012</td>
</tr>
<tr>
<td>Anteroposterior to mediolateral width ratio of iliac articular surface of pubis ≥2.0</td>
<td>Titanosauria</td>
<td>Mannion et al. 2013</td>
<td>Mannion et al. 2013</td>
</tr>
<tr>
<td>Acetabular margin of ischium strongly concave in lateral view such that pubic articular surface forms proximodorsal projection (*)</td>
<td>Titanosauria (Mannion et al. 2013) or Lithostrotia (D’Emic 2012)</td>
<td>D’Emic 2012, Mannion et al. 2013</td>
<td>D’Emic 2012</td>
</tr>
<tr>
<td>Femur with longitudinal ridge on anterior face of shaft</td>
<td>Alamosaurus + ‘Saltasaurini’</td>
<td>D’Emic 2012</td>
<td>Otero 2010</td>
</tr>
<tr>
<td>Femoral distal condyles beveled 10° dorsomedially relative to shaft</td>
<td>Saltasauridae</td>
<td>D’Emic 2012</td>
<td>Wilson 2002</td>
</tr>
<tr>
<td>Ratio of mediolateral width of distal end of tibia:long axis of a cross-section horizontally through the midshaft ≥2.0 (*)</td>
<td>Lithostrotia</td>
<td>Mannion et al. 2013</td>
<td>Wilson 2002</td>
</tr>
<tr>
<td>Astragalus mediolaterally narrow</td>
<td>Saltasauridae</td>
<td>D’Emic 2012</td>
<td>D’Emic 2012</td>
</tr>
</tbody>
</table>

As was described by Apesteguía (2005), there seems to have been a progressive reduction of the manual phalanges from basal titanosauriforms to derived titanosaurids.

We observe at least two manual morphologies in titanosaurids that are pertinent to discussions of the presence or absence of ossified phalanges in these sauropods: (1) metacarpals with well-defined distal articular facets and curved metacarpal I, as evidence of manual phalanges, and (2) metacarpals with poorly-defined distal articular facets, with or without vestigial ossified manual phalanges. In the first case, the presence of manual phalanges is supported both by well-defined distal articular facets on the metacarpals and the discovery of manual phalanges associated with the specimens in question. This is the case for two early Late Cretaceous Australian taxa, *Diamantinasaurus* and *Savannasaurus*, described by Hocknull et al. (2009) and Poropat et al. (2015, 2016), respectively. In *Diamantinasaurus*, the manual elements were not preserved in articulation but have been tentatively
interacted as indicative of a manual phalangeal formula of 2-1-1-1-1. In Savannasaurus, Poropat et al. (2016) recognized at least two manual phalanges, though these authors did not specify to which digits these bones pertained. These important discoveries hint at the existence of diverse manual structures within Titanosauria, though articulated specimens are needed to confirm certain aspects of their anatomy (e.g., the proposed phalangeal formula of Diamantinasaurus). Additional titanosaurian taxa such as Andesaurus and Argyrosaurus exhibit indirect evidence of manual phalanges, though these bones have yet to be discovered in these taxa (Mannion and Calvo 2011, Mannion and Otero 2012). In these two large Patagonian titanosaurs, metacarpal I is curved and somewhat ‘banana-shaped,’ similar to that of the basal titanosauriform Janenschia (Apesteguía 2005:334), thereby suggesting the possible presence of one or more diminutive manual phalanges (e.g., Figure 5c, Argyrosaurus).

In the second case, the metacarpals have poorly developed distal articular facets, and in two genera, Epachthosaurus and Opisthocoelicauda, a rudimentary phalanx is present on digit IV. Though both of these taxa are represented by fully articulated postcranial skeletons, there is no evidence of other ossified manual phalanges (Borsuk-Bialynicka 1977, Martinez et al. 2004). Similarly, in the unnamed Invernada taxon (González Riga et al. 2008a), no manual phalanges were discovered, although much of the skeleton was exquisitely preserved, including the articulated left fore- and hind limbs with all metacarpals and the complete pes (B.J.G.R. pers. obs.). An important feature of these taxa is that the metacarpals are in contact distally, forming a structure that is more tubular than the metacarpus of other neosauropods (Figure 5c, Epachthosaurus). In this context, the absence of ossified manual phalanges cannot be used as positive evidence (i.e., that these bones were present and subsequently removed by taphonomic processes, as suggested by Poropat et al. 2016:1012), nor as negative evidence of genuine absence (e.g., Salgado et al. 1997, Wilson 2002). To more rigorously evaluate this character in a given titanosaurian taxon, one must consider the presence or absence of distal articular facets on the metacarpals and the taphonomic context of known specimens.

Lognkosauria

Lognkosauria is a node-based clade defined as the most recent common ancestor of Mendozasaurus neguyelap and Futalognkosaurus dukei and all descendants (Calvo et al. 2007a). Many previous studies (e.g., Calvo et al. 2007a, b, González Riga and Ortiz David 2014, González Riga et al. 2016) have restricted Lognkosauria to these two taxa, but more recent analyses (Carballido et al. 2017, González Riga et al. 2018) have postulated Argentinosaurus and Patagotitan as lognkosaurians as well; the analysis of Carballido et al. (2017) additionally positioned Drusilasaura, Puertasaurus, and Quetecsaurus within this clade (Figure 10b), whereas that of González Riga et al. (2018) also included Notocolossus (Figure 10d). Intriguingly, most lognkosaurians or putative lognkosaurians are exceedingly large-bodied animals, with Argentinosaurus, Futalognkosaurus, Notocolossus, Patagotitan, and Puertasaurus all being among the largest titanosaurians yet discovered.

In the analysis of González Riga et al. (2018), Lognkosauria was diagnosed by eight synapomorphies, although none of these were regarded as unique to the clade. Two of these morphologies (a deep spinodiapophyseal fossa on the lateral surface of the base of the neural spine in posterior cervical vertebrae and laterally expanded posterior cervical neural spines resulting from expansion of the lateral lamina) are also present in the North American titanosaur Alamosaurus (Tykoski and Fiorillo 2017). From their analysis of
the data matrix of González Riga and Ortiz David (2014), Tykoski and Fiorillo (2017) obtained a single resolved tree in which Alamosaurus was postulated as the sister taxon of Lognkosauria (Figure 10c).

RECOGNITION OF A NEW TITANOSAURIAN CLADE

Although many aspects of titanosaurian phylogeny remain unresolved, recent studies have shed considerable light on the interrelationships of taxa within the clade. The phylogenetic definitions of groups such as Saltasaurinae and Lognkosauria are stable, but their taxonomic content has varied depending on the phylogenetic hypothesis recovered. The recent study of González Riga et al. (2018) employed a data matrix of 84 taxa scored for 423 characters. Analysis of this matrix recovered a clade of South American lithostrotian titanosaurs pertaining to Rinconsauria and Lognkosauria. The other principal lithostrotian clade postulated by this analysis primarily includes taxa from Asia and North America plus the Brazilian titanosaur Tapuiasaurus and the Indian Isisaurus (Figure 10d). In the phylogenetic hypothesis of González Riga et al. (2018), Lognkosauria is comprised by Argentinosaurus, Futalognkosaurus, Mendozasaurus, Notocolossus, and Patagotitan.

Carballido et al. (2017), in their study of the giant titanosaur Patagotitan, analyzed a dataset of 405 characters and 87 sauropodomorph taxa (including 28 titanosaurs) that was modified from the matrix of Carballido and Sander (2014). In contrast to most previous studies, Malawisaurus was not recovered as a comparatively basal titanosaur, but instead was placed in a position more derived than Lognkosauria and Rinconsauria. Because Malawisaurus is included in the definition of Lithostrotia (Upchurch et al. 2004), under this phylogenetic hypothesis, members of both Lognkosauria and Rinconsauria would be considered to be non-lithostrotian titanosaurs (Figure 10b). In Carballido et al.’s (2017) topology, the colossal Argentinosaurus and Patagotitan are again nested within Lognkosauria, this time accompanied by the similarly gigantic Puertasaurus. Another enormous titanosaur, Notocolossus, is placed as the sister taxon of Lognkosauria, and two main lineages are recovered within Carballido et al.’s (2017) Eutitanosauria: a ‘lithostrotian line’ that includes Saltasauridae and a second lineage that includes Lognkosauria and Rinconsauria.

The existence of a clade that includes undisputed members of Lognkosauria and Rinconsauria was previously recovered by Tykoski and Fiorillo (2017) based on the data matrix of González Riga and Ortiz David (2014) (Figure 10c). Similarly, Gallina and Apesteguía (2011) also recovered this clade, termed ‘node A’ in their analysis (Figure 10a). These authors based their study on the dataset of Calvo et al. (2007a) and González Riga et al. (2009), but they added new cranial and postcranial characters, as well as some taxa.

Relevant taxa such as Futalognkosaurus, Mendozasaurus, Muyelensaurus, and Rinconsaurus have been excluded from other recent and pertinent phylogenetic analyses, and as such, it is difficult to further evaluate the existence of the new clade proposed herein. However, although Mendozasaurus and Rinconsaurus were not included in the phylogenetic study of the bizarre Australian titanosaur Savannasaurus (Poropat et al. 2016, dataset of 297 characters and 72 taxa), the new group is supported by the recovery of a Muyelensaurus + (Epachthosaurus + Futalognkosaurus) clade. In this case, the new clade is independent from Nemegtosauridae and Saltasauridae (Poropat et al. 2016, fig. 7). Similarly, although Muyelensaurus and Rinconsaurus were not included in González Riga et al.’s (2016) phylogenetic analysis of Notocolossus (dataset of 350 characters and 33 taxa), a distinct, well-defined clade that includes
that genus plus *Dreadnoughtus*, *Futalognkosaurus*, *Mendoza saurus*, and *Tapuiasaurus* was recovered. Finally, though Lacovara et al. (2014, dataset of of 341 characters and 70 taxa) excluded *Muyelensaurus* and *Rinconsaurus* from their analysis of *Dreadnoughtus*, *Malawisaurus* was recovered as more derived than lognkosaurians, as was also the case in the analysis of *Patagotitan* (Carballido et al. 2017).

**REVISED PHYLOGENETIC ANALYSIS WITH ADDITIONAL CHARACTERS**

We executed a slightly revised phylogenetic analysis that employed the dataset of González Riga et al. (2018, 426 characters and 84 taxa), which in turn was based on that of Mannion et al. (2017) with the addition of some characters (see González Riga et al. 2018 for details). At present study, we added three characters (C424–C426) to

Figure 10 - Four cladistic hypotheses of relationships of titanosaurian taxa. Cladograms have been redrawn and are limited to Titanosauria for ease of comparison. Hypotheses depicted are those of (a) Gallina and Apesteguía (2011), (b) Carballido et al. (2017), (c) Tykoski and Fiorillo (2017), and (d) González Riga et al. (2018). Abbreviations: LO, Lognkosauria, RI, Rinconsauria.
the matrix; one of these is new and the other two were originally proposed by González Riga et al. (2016) (see Appendix). These added characters are as follows:

C424. Number of phalanges in pedal digit II: three (0), two (1) (González Riga et al. 2016).
C425. Number of phalanges in pedal digit IV: three or more (0), two (1) (modified from González Riga et al. 2016).
C426. Number of phalanges in pedal digit V: two (0), one (1), zero (2) (this paper).

The matrix was analyzed under equal character weighting using TNT (Tree analysis using New Technology) v. 1.1 (Goloboff et al. 2008). The multistate characters 11, 14, 15, 27, 40, 51, 104, 122, 147, 148, 177, 195, 205, and 259 were treated as ordered. In addition, eleven highly incomplete taxa (and therefore unstable) were excluded prior to the analysis (specimen AODF 836, Astrophocaudia, Australodocus, Brontomerus, Fukuititan, Fusiisaurus, Huanghetitan, ’Huanghetitan’ ruyangensis, Liubangosaurus, Mongolosaurus and Tendaguria). First, the data matrix was analyzed using New Technology Search with the functions ‘sectorial searches’, ‘drift’ and ‘tree fusing’. It was also used ‘get tree’ from ‘driven search’ and ‘find minimum length’ three times. Second, the resultant trees were searching by Traditional Search using the option ‘tree bisection-reconstruction’. This process resulted in 660 MPTs of 1741 steps and produced a fairly well-resolved strict consensus tree (Consistency Index, 0.248; Retention index, 0.560). The strict consensus of these generates a polytomy of basal titanosaurians but recovers the phylogenetic relationships of lithostrotian taxa within two primary clades: a lineage containing Lognkosauria and Rinconsauria, herein termed Colossosauria, and a clade containing Saltasaurus and other derived taxa.

PHYLOGENETIC DEFINITION OF COLOSSOSAURIA

Four recent phylogenetic analyses based on largely independent datasets (Gallina and Apesteguía 2011, Carballido et al. 2017, Tykoski and Fiorillo 2017, and González Riga et al. 2018) recovered a clade that includes Lognkosauria and Rinconsauria. Moreover, the results of an amended phylogenetic analysis based on the dataset of González Riga et al. (2018) that includes three additional appendicular characters also supports the existence of this group. Accordingly, we herein propose a new taxon following the tenets of phylogenetic taxonomy (Sereno 2005, Cantino and de Queiroz 2010), as follows:

Colossosauria new taxon

Etymology. From the ancient Greek colossos, colossus, giant, in reference to the gigantic size of some genera within the clade; from the Greek saurus, lizard, reptile.

Definition. Colossosauria is phylogenetically defined as the most inclusive clade containing Mendozasaurus neguyelap but not Saltasaurus loricatus or Epachthosaurus sciuttoi (stem-based).


Taxa. Following González Riga et al. (2018) and this paper, Colossosauria includes Argentinosaurus, Futalognkosaurus, Mendozasaurus, Muyelensaurus, Notocolossus, Patagotitan, and Rinconsaurus. After Carballido et al. (2017), the clade comprises the aforementioned taxa plus Aeolosaurus, Bonitasaura, Drusilasaura, Overosaurus, Puertasaurus, and Quetecsaurus.

Diagnosis. Following this paper, Colossosauria is diagnosed by seven ambiguous synapomorphies: humerus, minimum mediolateral width divided by proximodistal length less than 0.15 (Character 42, state 1), middle–posterior dorsal neural arches with posterior centroparapophyseal lamina (character
middle–posterior dorsal neural spines vertical or slightly posterodorsal (character 164, state 0), middle–posterior dorsal neural arches, neural canal in anterior view enclosed in a deep fossa in the dorsal surface of the centrum (character 338, state 1), glenoid does not expand strongly laterally relative to the lateral surface of the coracoid (character 361, state 0), humerus, deltopectoral crest, mediolateral thickness of anterior attachment surface with distal half mediolaterally expanded relative to proximal half (character 369, state 0), metacarpal V with dorsomedial margin of distal third forming a prominent ridge or flange (character 420, state 1).

**Observations.** The name proposed herein does not affect the previous nomenclature of other titanosaurian clades. Figure 11 shows the position of the clade Colossosauria.

**CONCLUSIONS**

Systematic and phylogenetic studies of titanosaurian sauropod dinosaurs have often focused greater attention on the postcranial axial skeleton than on the appendicular skeleton. Nevertheless, this practice has changed in recent years alongside

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**Figure 11** – Strict consensus cladogram (limited to Titanosauria) generated from a revised analysis of the data matrix of González Riga et al. (2018) with the addition of three characters (this paper), showing the position and taxonomic content of the newly-recognized stem-based clade Colossosauria. Abbreviations: Lo, Lognkosauria, Ri, Rinconsauria, Sa, Saltasauridae.
discoveries of new specimens that preserve the appendicular skeleton in its entirety, or nearly so. These fossils have provided a wealth of new anatomical and paleobiological information on titanosaurians, a group that is characterized by marked variation in both body size and morphology.

An overview of the appendicular skeletal morphology of titanosaurian taxa from South America indicates that this part of the skeleton exhibits greater anatomical diversity than was initially appreciated. Detailed comparative studies of each of these appendicular elements and skeletal regions would likely yield additional character information useful for phylogenetic analyses. From a systematic point of view, the present review shows that there is no definitive consensus on the appendicular character states that serve as synapomorphies for Titanosauria and Lithostrotia. However, within Titanosauria, recent phylogenetic analyses confirm the presence of a titanosaurian lineage that differs from the ‘Saltasaurinae line.’ Four recent cladistic studies based on largely independent datasets have recovered this new clade, which is herein termed Colossosauria. At present, this clade is mainly comprised by taxa belonging to Rinconsauria and Lognkosauria, the latter including several exceptionally gigantic species.

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AUTHOR CONTRIBUTIONS

The goals of this paper were proposed by B.J.G.R. Anatomical description was performed by B.J.G.R., M.C.L., A.O., and L.M.I. Phylogenetic analysis was performed by L.D.O.D., A.O., and B.J.G.R. The discussion was written by B.J.G.R., M.C.L., A.O., L.M.I., L.D.O.D., and A.W.A.K. The English revision was completed by M.C.L. The figures were done by L.D.O.D and B.J.G.R. The tables were made by M.C.L., B.J.G.R., and L.M.I.

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ZAHER H, POL D, CARVALHO A, NASCIMENTO P, RICCOMINI C, LARSON P, JUÁREZ VALIERI R,
APPENDIX

SCORES FOR ADDITIONAL PHYLOGENETIC CHARACTERS

The phylogenetic analysis in this paper is based on the dataset of González Riga et al. (2018), with the addition of three characters (C424–C426), the scores for which are as follows:

  *Omeisaurus*: 424 (0), 425 (0), 426 (0),
  *Shunosaurus*: 424 (0), 425 (0), 426 (0),
  *Cedarosaurus*: 424 (0), 425 (0), 426 (0),
  *Apatosaurus*: 424 (0), 425 (0), 426 (0),
  *Diplodocus*: 424 (0), 425 (0), 426 (0),
  *Camarasaurus*: 424 (0), 425 (0), 426 (0),
  *Janenschia*: 424 (0), 425 (0), 426 (0),
  *Gobititan*: 424 (1), 425 (1), 426 (?),
  *Epachthosaurus*: 424 (0), 425 (1), 426 (1),
  *Mendozasaurus*: 424 (1), 425 (1), 426 (1),
  *Notocolossus*: 424 (1), 425 (1), 426 (1),
  *Opisthocoelicaudia*: 424 (1), 425 (1), 426 (1),
  *Alamosaurus*: 424 (1), 425 (1), 426 (?).

NOTE ADDED IN PROOF

Since this paper went to press, another South American titanosaur that preserves appendicular skeletal remains has been formally described: *Kaijutitan maui* from the Upper Cretaceous (Coniacian) Sierra Barrosa Formation of Neuquén Province, northern Patagonia, Argentina (Filippi et al. 2019). The type material of this large-bodied taxon includes multiple, mostly incomplete appendicular bones, in addition to cranial and postcranial axial remains. The preserved appendicular elements are as follows: incomplete scapula and coracoid, sternal plate, humerus, radius?, ulna, metacarpals II and III, incomplete ?ilium and femur, tibia, astragalus, and incomplete metatarsal II. Though the incompleteness of many of these bones renders comparisons difficult, most (e.g., the ulna and tibia) suggest a stout-limbed animal, comparable in this regard to *Dreadnoughtus* from Upper Cretaceous sediments in southern Patagonia.