The morphofunctional design of *Montealtosuchus arrudacamposi* (Crocodyliformes, Upper Cretaceous) of the Bauru Basin, Brazil

Sandra Aparecida Simionato Tavares \(^a,^b\), Fresia Ricardi Branco \(^a\), Ismar de Souza Carvalho \(^c,^*\), Lara Maldanis \(^d,^e\)

\(^a\) Universidade Estadual de Campinas (UNICAMP), Instituto de Geociências, Cidade Universitária, Rua João Pandiá Calógeras, 51, 13083-870, Campinas, SP, Brazil
\(^b\) Museu de Paleontologia “Prof. Antonio Celso de Arruda Campos”, Praça do Centenário-çin — Centro, Monte Alto, SP, Brazil
\(^c\) Universidade Federal do Rio de Janeiro, Centro de Ciências Matemáticas e da Natureza, Instituto de Geociências, Departamento de Geologia, Cidade Universitária, Ilha do Fundão, Av. Athos da Silveira Ramos, 274, Bloco C, 21041-916 Rio de Janeiro, RJ, Brazil
\(^d\) Universidade de São Paulo, Instituto de Física de São Carlos, Av. Trabalhador sáo-carlense, 400, Pq. Arnold Schmidt, 13566-590, São Carlos, SP, Brazil
\(^e\) Laboratório Nacional de Luz Síncrona, Centro Nacional de Pesquisas em Energia e Materiais, Rua Giuseppe Máximo Scolfaro, 10.000 — Polo II de Alta Tecnologia de Campinas, 13083-970, Campinas, SP, Brazil

**A R T I C L E  I N F O**

Article history:
Received 19 April 2017
Received in revised form 22 June 2017
Accepted in revised form 9 July 2017
Available online 11 July 2017

Keywords:
Biomechanics
Crocodyliformes
Montealtosuchus arrudacamposi
Bauru Basin

**A B S T R A C T**

*Montealtosuchus arrudacamposi*, a Peirosauridae from the Upper Cretaceous of the Bauru Basin, was a Crocodyliformes of terrestrial habits. The fossils analyzed in this study belong to the pectoral girdle (scapula and coracoid) and anterior appendicular skeleton (humerus, radius, ulna, carpals, radiale, ulnale, metacarpals and phalanges). In this study we infer the locomotion habits of *M. arrudacamposi*. A morphometric, morphofunctional and 3D reconstruction of the elements of the pectoral girdle and the anterior limbs of *M. arrudacamposi* were performed. For a better understanding of the most plausible pectoral girdle and anterior limb posture, the studied bones were virtually disarticulated and articulated on 3D reconstruction.

The herein results obtained indicate the structures present a relatively thin and elongated aspect, thus allowing an interpretation that *M. arrudacamposi* possessed more slender anterior limbs than living crocodyliforms. This condition allowed for an adducted stance and cursorial habits that would enable movement through terrestrial environments for prey searching.

© 2017 Elsevier Ltd. All rights reserved.

1. Introduction

In order to understand the way of life of an extinct species, it is necessary to consider all the evidences preserved in a bone structure, and thus understanding its functionality. Benton (2010) states that there are several ways to research the behavior of a species through fossils. One of them is to evaluate the bones’ joints and make inferences regarding the shape of the extremities on preserved structures.

Selden (1990) also argues that through the morphology of the bony structures of vertebrates, and the features preserved in them, it is possible to test hypotheses about their functionality. Several authors (Bakker, 1971; Charig, 1972; Bonaparte, 1986; Sennikov, 1987; Kischlat, 2000) have based their work on, among other aspects, the graduation in angles to determine the abduction of the femoral axis and understand the posture and locomotion of tetrapods. However, information contained in the bones of the anterior limbs, such as the humerus, can also provide data regarding the posture of parts of a given species.

Fossil Crocodyliformes are well-represented in sedimentary deposits of Bauru Basin in the municipality of Monte Alto, State of São Paulo, Brazil. Among them, is the Peirosauridae *Montealtosuchus arrudacamposi* (Carvalho et al., 2007), that presents excellent preservation of the bony elements that compose its skeleton. It has the skull, mandible, postcranial elements and dermal shield preserved practically in their original positions (Fig. 1).

In this study, a morphological and morphofunctional description of the bones, constituting the pectoral girdle and anterior...
appendicular elements of *M. arrudacamposi*, was performed to understand its locomotory habits.

2. Geological context

Bauru Basin is located in the south-central region of the South American Platform and is distributed in the states of São Paulo, Minas Gerais, Mato Grosso do Sul, Goiás and northeastern Paraguay, adding up to an area of approximately 370,000 km² between latitudes 18°S and 24°S, and longitude 47°W and 56°W (Fernandes and Coimbra, 1996). According to Menegazzo et al. (2016) the Basin’s development occurred from the Cenomanian to early Paleocene in the back-bulge province of a retroarc foreland system that came to existence in response to Andean orogenic events. It is divided into two lithostratigraphic units: the Caiuã Group (Paraná River, Goioi Ére and Santo Anastácio formations) and the Bauru Group (Adamantina, Uberaba and Marília formations) (Fernandes, 1998; Dias-Brito et al., 2001).

The sedimentary fill formed by the Bauru Group corresponds to continental deposits. It is composed of fluvial sediments (Adamantina and Uberaba Formations) and alluvium (Marília Formation) (Fernandes and Coimbra, 2000). In the region of Monte Alto there are two lithostratigraphic units of the Bauru Group surface: the Adamantina’s and the Marília’s Formations.

The fossils of the *Montealtosuchus arrudacamposi* were found in rocks from the Turonian-Santonian (Upper Cretaceous) of the Bauru Basin, Bauru Group, Adamantina Formation in the region of Monte Alto (S 21°09’53.9” W 48°29’54.0”), São Paulo State, Brazil (Fig. 2). These specimens were preserved in a succession of thin to medium layers of loosely-cemented reddish sandstone. The strata overlaying this layer are composed of fine sandstone with remains of tetrapods and bivalves. Finally, there is a highly cemented conglomerate layer containing coprolites, as well as teeth and isolated bone elements from dinosaurs, chelonians, and squamata.

According to Carvalho and Bertini (2000), the paleoclimate during the deposition of the Adamantina Formation is characterized as warm with torrential rains and floods. Basilici et al. (2009) states, based on the study of paleosols, that the Marília Formation was characterized during the Late Cretaceous by a semi-arid climate.
According to Carvalho et al. (2010), an arid or a warm climate, dry seasonal climate with alternated wetter periods, could have influenced the wide diversity of crocodilians in Gondwana during the Cretaceous. According to Iori et al. (2016), in the Adamantina Formation of the Monte Alto region, the Morrhinosuchus and Caipirasuchus crocodyliform specimens are autochthonous and occur in typical paleosol deposits, whereas Barreirosuchus and Montealtosuchus are in deposits of fluvial and/or lacustrine environment.

The remains of *M. arrudacamposi* were found in association with fragments of another skull corresponding to the left side, and there are other several postcranial fragments of at least four other individuals that appear to belong to the same species (Tavares et al., 2015). According to Carvalho et al. (2005), the occurrence of almost complete articulated skeletons suggests these animals could dig large deep burrows in soft substrates that allowed thermoregulation, like in living alligators.

The number of cranial peirosaurid specimens, preserved along the *M. arrudacamposi* holotype, suggests these animals lived in groups and that they may have also had the habit of burying themselves in the substrates just like current crocodyliforms.

3. Materials and methods

3.1. Post-cranial elements of *M. arrudacamposi*

The fossil bones analyzed in this study were: coracoids — right and left — and left scapula (pectoral waist); humeri — left and right, and left radius, ulna, distal carpals, radiale, ulnare, metacarpals and phalanges (left appendicular skeleton) of the holotype of *M. arrudacamposi* (Fig. 3B). This specimen is housed in the Museum of Paleontology "Prof. Antonio Celso de Arruda Campos", based in the city of Monte Alto-SP, collection number MPMA-16-0007/04. The measures were taken with MAUb-CH Stainless steel calipers and a tape measure.

3.2. Nomenclature and taxa compared

The terms used for the anatomical nomenclature of osteology are the same as those used by Mook (1921), Romer (1956), Richardson et al. (2002) and Buckley and Brochu (1999), Hoffstetter and Gasc (1973), Tarsitano (1982). To assign postural and locomotory patterns, von Huene's (1913) proposal was used.
and compared with data from living and fossil crocodyliforms (Table 1).

### 3.3. Computed tomography and 3D images

The holotype of *M. arrudacamposi* was scanned at the Institute of Radiology, Faculty of Medicine, São Paulo USP, Brazil, using the Discovery CT750 HD CT Scanner, GE Health Systems, Milwaukee, USA. The CT images were segmented in InVesalius 3.0 software, developed at the Three-Dimensional Technologies Division, (Renato Archer Information Technology Center, city of Campinas, São Paulo—Brazil), and at the Brazilian Synchrotron Light Laboratory (LNLS), CNPEM (Campinas, São Paulo – Brazil), using the Avizo 9.0 program of the FEI Visualization Sciences Group, software that allows separation of the fossilized structures of the matrix using the contrast generated by their difference in density.

After segmentation of the CT slices, 3D models of the structures were generated (Fig. 3), thus allowing a reorganization of the bones by placing them in the most correct position, according to the morphological information available in the fossils.

### 4. Results

#### 4.1. Scapula (Fig. 4)

Only the left scapula, about 90 mm in length, is preserved in the holotype of *M. arrudacamposi*. It has an anteroposterior expansion giving rise to the scapular blade. This structure is very thin, approximately 1 mm thick. In the lateral view a more evident constriction is observed in the distal portion of the scapula. The posterior concavity is posteriorly positioned at the distal end. In lateral view two crests are evident, one lateral anteriorly in the distal portion and the other lateral posteriorly originating from the proximal end, being more evident at the distal end. There is a well-defined concavity in the medial view’s most distal portion. The glenoid cavity is concave and located ventrally in the distal portion of the scapula.

#### 4.2. Coracoids (Fig. 5)

The right and left coracoids were preserved practically in their original position, connected in both, the proximal and distal portions by a set of osteoderms, which make up the pectoral dermal shield of *M. arrudacamposi* (Tavares et al., 2015). They are
approximately 80 mm long and have a posterior concavity in the region of the diaphysis.

In both coracoids the coracoid foramen is preserved centrally at the proximal end. The contact surface with the scapula is relatively flat, and the glenoidal one is posteromedially oriented. The humeral joint surface is slightly concave. Medially, below the joint surface for the humerus, there is a well-defined groove (posterior sulcus). There is distal expansion of the coracoid.

### 4.3. Humerus (Fig. 6)

Both humeri of *M. arrudacamposi* are preserved. They are approximately 110 mm in length. The left humerus is damaged in the distal epiphysis and preserved coupled to the coracoid and scapula, but displaced from its original position. The right humerus is disarticulated from the coracoid and the scapula (Fig. 1).

---

**Fig. 4.** Left scapula of *Montealtosuchus arrudacamposi* (MPMA-16-0007/04). A, position in which the bones in the study were preserved and lateral view of the scapula. B, medial view of scapula. C, anterior view of forelimb and pectoral girdle — possible correct position of the scapula (articulated scapula with the coracoid and humerus). Abbreviations: cor, coracoid; gs, glenoid surface; esc, scapula; hu, humerus; pc, posterior concavity.

**Fig. 5.** Coracoids of *Montealtosuchus arrudacamposi* (MPMA-16-0007/04). A, position in which the studied bones were preserved. B, right and left coracoids in ventral view, joined by the pectoral osteoderms. C, medial view of the right coracoid. D, anterior view — possible in vivo position of the coracoids (coracoids articulated to the humerus). Abbreviations: ash, humeral joint surface; cor, coracoid; cf, coracoid foramen; dec, distal expansion of the coracoid; esc, scapula; gs, glenoid surface; hu, humerus; ost, osteoderms; pg, posterior sulcus.
In the lateral view, both the left and right humeri have a sigmoidal shape. The proximal end is expanded lateromedially and an attenuated concavity is seen the anterior view. The joint surface of the humeral head for the glenoid cavity is convex.

The deltopectoral crest develops dorsolaterally extending distally to the beginning of the diaphysis. In the posterior view a depression is visible at the proximal end.

The diaphysis of the humerus of *M. arrudacamposi* is cylindrical, approximately 40 mm long and 15 mm in diameter. The distal end is preserved in the right side humerus. There is a pronounced concavity anteriorly. The lateral and medial epicondyles are convex, anteriorly projected and separated by the concavity.

### 4.4. Ulna (Fig. 7)

The ulna of *M. arrudacamposi* is articulated to the radius. Only the left ulna of approximately 100 mm long is preserved. Laterally it has a distinct sigmoidal shape. The proximal end is...
elliptical in cross section. In the posterior view, the proximal end is damaged and the olecranon has not been preserved. The diaphysis measures 80 mm and two thirds of it, from the proximal epiphysis, are considerably flattened. In the remainder of its length the diaphysis becomes more cylindrical. The distal epiphysis is also damaged.

4.5. Radius (Fig. 7)

Only the left radius is preserved in this specimen. It has the same length as the ulna (100 mm). The epiphyses were not completely preserved; however, it is possible to observe an expansion medially at both, the proximal and distal ends.

The proximal half of the diaphysis is circular in cross-section. However, towards the distal end it becomes compressed. In the anterior view, an osteoderm of the appendicular shield (Tavares et al., 2015) is observed on the diaphysis.

4.6. Anterior autopodium (Fig. 8)

The bones that make up the front foot (radiale, ulnare, pisiform, carpals, metacarpals and phalanges) are well preserved and practically in life position.

4.6.1. Radiale (Fig. 8)

The radiale measures 30 mm and articulates with the distal epiphysis of the radius. The proximal radiale epiphysis is well-expanded medially. The proximal and distal joint surfaces are concave, and the proximal surfaces have a somewhat more defined concavity than the distal one. Dorsomedially, at both the proximal and distal ends, there is a surface for the articulation of this bone at the distal end of the ulna and the ulnar condyles.

4.6.2. Ulnare (Fig. 8)

The ulnare is one-third smaller than the radiale. The articulation surface of the proximal epiphysis is flat and poorly developed. It articulates medially to the proximal epiphysis of the radiale and laterally to the distal epiphysis of the ulna. The distal epiphysis is well-expanded laterally and medially. In the dorsal view, it is in contact with the medial distal portion of the radial epiphysis. Ventrally articulates with the dorsal region of the distal carpus and proximal portion of the pisiform.

4.6.3. Pisiform (Fig. 8)

The pisiform is a bone that articulates proximally to the ulna and medially to the ulnare. It is expanded lateromedially in the distal portion, forming a surface with a slightly concave aspect. Anteriorly, it is in contact with the diaphysis and the proximal end of the ulnar and radial, as well as with the distal epiphysis of the ulna.

4.6.4. Carpi (Fig. 8)

In *Montealtosuchus arrudacamposi* the proximal carpus is preserved. The proximal end is articulated dorsally in the ventrolateral region of the ulnare, and the distal is articulated to digit IV. In the lateral view it is flattened.

The distal carpus is a small bone, approximately 10 mm in diameter. It is rounded dorsally articulating with the ventromedial portion of the distal epiphysis of the ulnare. Ventrally it is flattened and articulated to the proximal epiphyses digits of II and III.

4.6.5. Metacarpi (Fig. 8)

Only four of the five metacarpi present in the crocodyliform are preserved. They are approximately 25 mm long. The digits I, II and III are expanded and convex at the proximal ends and digit IV is more rectilinear along its length. There is a groove at the distal end,
on the dorsal and lateral face of each metacarpal. On the distal surface the condyles for the articulation of the proximal phalanges are located medially and laterally.

4.6.6. Phalanges (Figs. 8, 9)

Only nine phalanges are preserved composing the formula (2, 3, 4, ?, ?). The phalanges of digits I, II and III were preserved. They have an hourglass shape (log glass) and are short. The proximal phalanges are the largest in this series of bones, at approximately 10 mm long, with those of digits I and II at practically the same size, becoming smaller in digit III (8 mm). The medial phalanges of digit I and II are smaller (5 mm) than the proximal phalanges. The most distal phalanx of digit III is 3 mm long, but the smallest phalanx in diameter is displaced and preserved between digits III and IV. Probably it belongs to digit IV or V, which does not have any phalanx preserved in the natural position.

The proximal ends of the phalanges are concave and the distal is convex, forming the articular surfaces. Each phalanx has a rounded cavity on the left side of the distal end. Digits I, II and III preserve the ungual phalanges. These are articulated dorsally to the phalange proximal to them. The ungual phalanx of digit I is larger (20 mm) than those of digits II and III. All have a dorsal curvature that gives them the shape of a scythe. They are laterally flattened and convex. The phalanges of digits II and III have a greater ventral concavity than that of digit I. The tips of the phalanges are sharp.

5. Discussion

5.1. Morphological inferences

The structures of the pectoral girdle and anterior appendicular skeleton of *M. arrudacamposi* have well developed ridges that probably served to fix the scapulocoracoid muscles and brachial regions (*M. triceps, M. scapulohumeralis, M. humeroantebrachialis inferior, M. triceps brachii, M. humeroradialis, M. extensor carpi radialis, M. flexor ulnaris, M. abductor radialis, M. supinator, M. pronator teres, M. extensor carpi radialis brevis, M. pronator quadratus, M. abductor radialis, M. extensor carpi radialis longus*) (Meers, 2003). These muscles implies the possibility of an erect posture.

The bony elements of the appendicular skeleton of *M. arrudacamposi* are arranged vertically below the articulations of the coracoid and scapula. The humerus, ulna and radius are elongated bones, which suggests that this crocodyliform would be able to raise its body from the ground. Therefore, a hollow cylinder structure is stronger in axial compression than in bending, so that in order to have the safety factor requisite, the limb bone of a sprawling posture would be larger (in diameter) than that of an erect posture.

According to Briton (2005), this is also a described characteristic for the living crocodyliforms, which in addition to the aquatic locomotion habits, move in terrestrial environments by means of a
high walk, gallop (that propels the members alternately to the air) and creeping (which allows the trunk contact with the ground and members positioned laterally). Grigg and Kirshner (2015) believe that the galloping gait can be considered an almost perfect symmetric boundary, where the anterior limbs reach the ground sequentially as the hind limbs do.

In living crocodyliforms the humerus articulates to the robust bones of the pectoral girdle (Grigg and Kirshner, 2015). The humerus of *M. arrudacamposi* also articulates to the coracoid and the scapula, however, more vertically than in living crocodyliforms which are more adapted to aquatic environments. They aid in lateral stability when a crocodile is swimming or floating. On land, limbs are able to raise and hold the heavy body uplifted from the ground.

According to Klein (2016), the distal end of the coracoid of living crocodyliforms articulates with the scapula and humerus. In *Alligator* the scapula is near and parallel to the spine contributing with the coracoid to the formation of the glenoid cavity. In *M. arrudacamposi*, the distal end of the coracoid also articulates with the scapula and humerus.

In general, the scapula of *M. arrudacamposi* is similar to that of *U. terri*culus which has the same width and height (Vasconcellos, 2006; Turner, 2006). The scapular blade is expanded anteposteriorly, differing from the patterns described for living crocodiles (Meers, 2003). It is much thinner antero-posteriorly than those of *Baurusuchus albertoi*, *Campinasuchus dinisci* and *Mariliasuchus amarali* (Nascimento and Zaher, 2010; Nobre and Carvalho, 2013; Cotts et al., 2017).

The posteriorly positioned concavity at the distal end in *M. arrudacamposi*, is also observed in *B. albertoi* and *C. dinisci*. In *Sinosuchus clarki* and *M. amarali* this concavity is not evident (Nascimento and Zaher, 2010; Sertich and Groenke, 2010; Nobre and Carvalho, 2013; Cotts et al., 2017).

According to Leardi et al. (2015), the distal end of the scapula of *Yacarera ni boliviensis* has an asymmetrical aspect in lateral view, with the anterior extremity more projected and proximally located than the posterior extremity. In *M. amarali* and *S. clarki* this characteristic also occurs; however, in *M. arrudacamposi* this extremity is not very noticeable (Sertich and Groenke, 2010; Nobre and Carvalho, 2013).

In *U. terri*culus, *B. albertoi*, *C. dinisci*, *S. clarki*, *M. amarali*, *Y. boliviensis*, *Caipirasuchus montealtensis*, as in *M. arrudacamposi*, the coracoids are more expanded in the distal than in the proximal portion. The coracoid foramen in these crocodyliforms is circular, but in *C. dinisci* is located in the posterodorsal region and in *C. montealtensis* it is more compressed (Vasconcellos, 2006; Nascimento and Zaher, 2010; Iori et al., 2016; Cotts et al., 2017).

The articular surface for the scapula in *Notosuchus terrestris* has a triangular shape and in *S. clarki* and *M. amarali* is curvilinear (Pol, 2005; Sertich and Groenke, 2010; Nobre and Carvalho, 2013). In *Y. boliviensis* this surface is ‘U’-shaped (Leardi et al., 2015). This surface in *M. arrudacamposi* is straight as in *B. albertoi* (Nascimento and Zaher, 2010).

In the coracoids of *U. terri*culus and *C. dinisci* the glenoid surface was not preserved (Vasconcellos, 2006; Cotts et al., 2017). In *B. albertoi*, *S. clarki*, *Y. boliviensis* is posterolaterally oriented (Nascimento and Zaher, 2010; Sertich and Groenke, 2010; Leardi et al., 2015). *Striatosuchus maxhechti* shows a glenoidal process posteromedially (Riff and Kelner, 2011). In *C. montealtensis* it is aligned interventrally (Iori et al., 2016). In *M. arrudacamposi* the glenoidal surface is aligned posteromedially.

The humeral articular surface is broad and convex in *S. maxhechti*, flat in *M. amarali* and slightly concave in *M. arrudacamposi* (Riff and Kelner, 2011; Nobre and Carvalho, 2013). The humerus of *M. arrudacamposi* articulates vertically to the coracoid, whereas in the present crocodyliform, this structure is directed to a more horizontal position.

In the lateral view the humerus of *M. arrudacamposi* exhibits a sigmoidal shape as in *B. albertoi*, *N. terrestris*, *M. amarali* (Pol, 2005; Nascimento and Zaher, 2010; Nobre and Carvalho, 2013). The proximal end is expanded lateromedially and an attenuated concavity occurs in anterior view. The articulation surface with the glenoid cavity (humeral head) is convex lateromedially as in *B. albertoi* (Nascimento and Zaher, 2010).

The deltoplecostal crest develops dorsolaterally extending distally to the beginning of the diaphysis, differing from *B. albertoi* and *S. maxhechti*, in which it expands anteriorly (Nascimento and Zaher, 2010; Riff and Kelner, 2011). In *M. amarali* and *N. terrestris* this crest is slightly developed and does not exhibit a marked medial deviation (Pol, 2005; Nobre and Carvalho, 2013). It is similar to that of *U. terri*culus and *A. tsangatsangana*, which in the latter extends distally by about one third of the humerus length (Vasconcellos, 2006; Turner, 2006).

In the posterior view a depression occurs in the proximal end of the humerus, however, it is not markedly circular as in *S. maxhechti*, *N. terrestris*, *M. amarali*, *Y. boliviensis*, *C. montealtensis* (Riff and Kelner, 2011; Nobre and Carvalho, 2013; Leardi et al., 2015; Iori et al., 2016).

The diaphysis of the humerus of *M. arrudacamposi* is cylindrical, approximately 40 mm long and 15 mm in diameter. The distal end is preserved in the right humerus, however, is damaged in the most peripheral part of the distal epiphysis in the left humerus. There is a pronounced concavity anteriorly. The lateral and medial epicondyles are convex, projected anteriorly and separated by a concavity. The characteristics described for the distal humerus of *M. arrudacamposi* are also present in *U. terri*culus, *B. albertoi*, *N. terrestris*, *A. tsangatsangana*, *S. clarki* and *M. amarali* (Pol, 2005; Vasconcellos, 2006; Turner, 2006; Nascimento and Zaher, 2010; Sertich and Groenke, 2010; Nobre and Carvalho, 2013). In *C. dinisci* the humeral diaphysis is more elongated and slightly thinner than in *S. maxhechti* (Riff and Kelner, 2011; Cotts et al., 2017).

In general, the ulna of *M. arrudacamposi* is similar to the ulna of *B. albertoi*, *P. sera* and *C. dinisci* (Nascimento and Zaher, 2010; Godoy et al., 2016; Cotts et al., 2017). The proximal end differs from that of *S. clarki*, and is posteriorly more inclined towards the radius (Sertich and Groenke, 2010). The diaphysis is more flattened than in *B. albertoi* and *M. amarali* (Nascimento and Zaher, 2010; Nobre and Carvalho, 2013).

The *M. arrudacamposi* radius is a straight and elongated bone as in other crocodyliforms (eg *U. terri*culus, *B. albertoi* and *M. amarali*) (Vasconcellos, 2006; Nascimento and Zaher, 2010; Nobre and Carvalho, 2013). In *C. dinisci* an anterior torsion occurs for the diaphysis portion of the radius (Cotts et al., 2017). In general, the ulna of *M. arrudacamposi* is similar to the ulna of *B. albertoi*, *P. sera* and *C. dinisci* (Nascimento and Zaher, 2010; Godoy et al., 2016; Cotts et al., 2017). The proximal end differs from that of *S. clarki*, and is posteriorly more inclined towards the radius (Sertich and Groenke, 2010). The diaphysis is more flattened than in *B. albertoi* and *M. amarali* (Nascimento and Zaher, 2010; Nobre and Carvalho, 2013).
As in *M. arrudacamposi*, the *B. albertoi* ulnar is smaller than the radial, but the distal end is more expanded than the proximal one and has a triangular shape due to a narrow anterior extension. This form is also present in *U. terrificus* (Vasconcellos, 2006; Nascimento and Zaher, 2010). In *C. dinizii* the radiate and ulnar are longer than they are wide, but the radiate is more robust and higher than the ulnar, as in *M. arrudacamposi* (Cotts et al., 2017).

Vasconcellos (2006) described for *U. terrificus* and Nascimento and Zaher (2010) for *B. albertoi* only the distal carpus, which is also preserved in *M. arrudacamposi*. In *C. dinizii* the distal carpus is articulated to the ulnar in a ventral region to the articular facet of the distal end of the bone (Cotts et al., 2017). In *M. arrudacamposi* the articular surface with the ulnare is rounded and the contact surface with the metacarpals is concave. In *U. terrificus*, it is in direct contact with III and IV (Vasconcellos, 2006). In *B. albertoi*, the contact of the distal carpus is with III, IV and V metacarpi (Nascimento and Zaher, 2010). In *M. arrudacamposi* it is only in contact with digits II and III.

Only four (I, II, IV, IV) out of the five metacarpals present in crocodylians are preserved (Romer, 1956). They are moderately short compared to those of *B. albertoi* (Nascimento and Zaher, 2010). They are more similar to those of *U. terrificus*, which are moderately elongated and robust (Vasconcellos, 2006). In all the peirosaurid, *M. arrudacamposi*, *U. terrificus* and *B. albertoi*, the greatest expansion in the proximal portion occurs in digit I. All have a distal expansion that allows the proximal phalanges to articulate.

In *M. arrudacamposi* a displaced phalanx is preserved, which is between the digits III and IV, and nine articulated phalanges distributed as follows: digits I-2, II-3, III-4, IV-7, V-7. This formula differs from *U. terrificus* (3-4-5-3-?) and that of *C. dinizii* (2-3-4-5-3) but is identical for digits I, II and III of *B. albertoi* (2-3-4-3-4) (Vasconcellos, 2006; Nascimento and Zaher, 2010; Cotts et al., 2017).

In *M. arrudacamposi* the proximal phalanges of metacarpals I and II are the robust. This condition also occurs in *U. terrificus* (Vasconcellos, 2006). As in *B. albertoi*, there is a decrease in the length of the proximal phalanx compared to the distal phalanges. The ungual phalanges are present in the digits I, II and III of *M. arrudacamposi* as well as in *U. terrificus* and *B. albertoi* (Vasconcellos, 2006; Nascimento and Zaher, 2010).

According to Vasconcellos (2006), the claws of *Baurusuchidae* are robust and almost twice as curved as the *U. terrificus* claws which are thin and have laminated ventral edges. The ungual phalanges of *M. arrudacamposi* are similar to those of *U. terrificus*, but they do not appear to articulate laterally with each other, as suggested by Vasconcellos (2006).

### 5.2. Morphofunctional inferences

In order to understand the locomotion of an animal in the morphofunctional analysis of the post-cranial structures of paleovertebrates, the supporting articular surfaces need to be evaluated. This evaluation is essential because their conformation is the way they are constructed thus, becoming an important aspect for understanding the articulation of limb bones. New methodologies, such as 3D reconstructions in life position of adjacent articulated fossilized bones, contribute in the morphofunctional studies of paleovertebrates.

Authors such as Duarte et al. (2011) used a virtual three-dimensional model of *B. salgadoensis*, obtained through a CT scan to virtually manipulate the appendicular bone elements of this crocodyliform, which by the way were fully articulated. After separation, it was possible to virtually allocate them to a position as close as possible to the original in life.

Like *B. salgadoensis*, the appendicular bony elements of *M. arrudacamposi* were also separated and virtually aligned in a more plausible life-like position. The 3D realignment of the bones, the pectoral girdle and anterior appendicular skeleton of *M. arrudacamposi*, showed that this crocodyliform had a more vertical position than of its anterior limbs.

The pectoral girdle of living crocodylimorphs does not connect the axial skeleton and is held in place by the muscles and fascias (Grigg and Kirshner, 2015). The scapula is larger than the coracid; the latter articulates in the ventral region of the scapula and along with the glenoid process, forms the glenoidal cavity, that is an articular surface for the head of the humerus (Vieira et al., 2016). The scapula in living crocodylimorphs is arched over the coracid (Grigg and Kirshner, 2015). Also, in *M. arrudacamposi*, the bones of the pectoral girdle do not articulate to the axial skeleton.

The scapula and the coracoids have been preserved almost in their original position. The coracoids are connected by a set of osteoderm and the scapula of *M. arrudacamposi* is not arched as in living crocodylimorphs, suggesting a more vertical articulation of the humerus to the elements of the pectoral girdle (Fig. 5A–D).

Clark et al. (2004) believed that the alignment of the glenoid fossa, the convex head on the humerus, the lack of rotation around the distal end of the ulna, the compact metacarpals, and the reduced outer digits, all indicate that limb moved in a vertical-parasagittal plane. This position is classified by von Huene (1913) as adducted, in which the animal's body assumes a more upright position (Blob, 2001; Berman and Henri, 2003).

In this study we observed that the characteristics described by Clark et al. (2004) are also present in the structure of *M. arrudacamposi*. In this crocodyliform the glenoidal surface is postero-medially aligned, and the head of the humerus, which articulates with the glenoid cavity, is convex.

The left humerus of *M. arrudacamposi* was preserved invertedly from its position in life, suggesting there was a rotation of this bone around the articular surface of the coracid during the process of preservation. This structure was virtually disarticulated and repositioned in the most suitable living position. The virtual articular adjustment between the coracid, scapula and the positioning of the humerus next to the coracid, also allowed the repositioning of the radius, ulna and elements of the front autopodium. For Clark et al. (2004), the conformation of the shoulder and wrist joints and the reduction in the size of the external digits may be evidence that the anterior limbs are positioned directly under the body. These authors have analyzed the pectoral girdle fossils and the anterior limb bones of *Jungarsuchus sloanii*. They concluded its humerus is different from any other Crocodyliformes because it has a well-developed hemispherical head projecting perpendicular to the long axis of the diaphysis, with a convex proximal articulation surface.

The long elements of the anterior appendages of *Uberabasuchus terrificus* are generally rectilinear and robust with well developed articular surfaces and condyles, similar to those of *M. arrudacamposi*. According to Vasconcellos (2006) the Peirosauridae could have had a more parasagittal posture on land than that of recent Crocodyliformes. Vasconcellos et al. (2007) analyzed the appendicular bones of *B. salgadoensis* and concluded they are long and robust structures. The anterior appendicular bones are almost straight and have broad articular facets. These aspects suggest a more upright posture for this crocodyliform. *M. arrudacamposi* has elongated appendicular bones as well; however, the articular facet contact between the coracid and humerus is respectively concave and convex.

Another baurusuchid that has elements of the anterior limbs preserved is the *Striatitosuchus maxhechti*. Riff and Kellner (2011) affirm the humerus of *S. maxhechti* has a broad and convex head which articulates to the glenoid process posteroventrally, thus enabling movement of the anterior limbs in the parasagittal plane. The aspects of the glenoid cavity and humeral head of *S. maxhechti* are similar to those of *M. arrudacamposi*.1357460817362213
The characteristics of the structures preserved in the bony elements of the pectoral girdle and anterior appendages of *Mar"iliasuchus amarali* differ from those observed in *M. arrudacamposi*. The articular facet of the scapula with the coracoid of *M. amarali* is located along the anterior half of the distal end. In the coracoid, the articular surface for the humeral head is flat, whereas in *M. arrudacamposi* is convex. Nobre and Carvalho (2013) concluded that *M. amarali* did not have an upright posture, or semi-erect as proposed for *M. arrudacamposi*, but rather a sprawling posture, similar to the present crocodyliforms.

In addition to the orientation of the glenoid surface, the convex head of the humerus led Clark et al. (2004) to suggest that the lack of rotation around the distal end of the ulna, compact metacarpals, and reduced exterior digits was a sign that the anterior limb could have moved vertically in the parasagittal plane.

In *M. arrudacamposi* the distal extremity of the ulna and the radius were not preserved, so it is not possible to infer the form of rotation around this extremity. However, the metacarpals are short and close to each other, a condition similar to those of *S. maxhechti*, which are compressed together and not spread laterally as in living crocodyliform. These aspects suggest a digitigrade posture of the manus, where only the distal portions of the digits rest on the ground (Clark et al., 2004; Riff and Kellner, 2011) (Fig. 10). Both *M. arrudacamposi* and *U. terrificus* have short robust metacarpals. According to Vasconcellos (2006) these characteristics indicate the metacarpals could have been adapted to support the inherent loads from terrestrial locomotion.

6. Conclusion

The exceptional preservation of the bony elements of the pectoral girdle and the anterior appendicular skeleton of the *M. arrudacamposi* holotype, the virtual disarticulation with a subsequent 3D reorganization of these structures, articulating them in the position closest to the original, suggest this Peirosauridae species inhabited terrestrial environments and moved around by means of a high walk.

The right and left coracoids joined by the pectoral osteoderms indicate these structures were preserved essentially in their original position. In conjunction with these features, the articulation surfaces, mainly of the scapula, coracoid and humerus, showed that the anterior limbs of *M. arrudacamposi* had a more upright position compared to the living crocodyliforms. The anatomical arrangement of long bones and autopodal elements, such as the elongated metacarpals close to each other, indicate that *M. arrudacamposi* had cursorial habits, allowing them to wander in terrestrial environments in search of prey.

Acknowledgements

We thank to the crew of Museu de Paleontologia “Prof. Antonio Celso Arruda Campos”, Antonio Celso de Arruda Campos (in memoriam), Cledinei A. Francisco, Fabiano Vidoi Iori, Deerson da Silva (Pepi), Paulo Gilberto da Rocha Tavares, José Augusto Bugarelli, Angélica Fernandes, Fernando Henrique Morais da Rocha and Hellen Tavares, for the help in the field and laboratory work. The paleoartist Pepi for the permission to use the reconstruction image of *Montealtosuchus*. The geologist Fabio Branco, Prof. Paulo Hilário Nascimento Saldiva and Prof. Edson Amaro Jr. supported the tomography analysis in Institute of Radiology, Faculty of Medicine, São Paulo USP, Brazil. Jorge Vicente Lopes da Silva and crew to the analysis in the Three-Dimensional Technologies Division, Renato Archer Information Technology Center, Campinas, São Paulo, Brazil. Special thanks are due to the Editor-in-Chief, Eduardo A.
Koutoukos and two anonymous reviewers contribution, by the depth of their comments, in their respective attempts to help to improve the paper. This study was supported by Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES), Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq) and Fundação Carlos Chagas Filho de Amparo a Pesquisa do Estado do Rio de Janeiro (FAPERJ).

References


