Flexibility along the Neck of the Neogene Terror Bird
Andalgalornis steulleti (Aves Phorusrhacidae)

Claudia P. Tambussi1,2*, Ricardo de Mendoza1, Federico J. Degrange2,3, Mariana B. Picasso1

1 División Paleontología Vertebrados, Museo de La Plata, Paseo del Bosque s/n, La Plata, Argentina, 2 Consejo Nacional de Investigaciones Científicas y Técnicas CONICET, Ciudad Autónoma de Buenos Aires, Argentina, 3 Centro de Investigaciones en Ciencias de la Tierra CICTERRA, Córdoba, Argentina

Abstract

Background: Andalgalornis steulleti from the upper Miocene–lower Pliocene (=6 million years ago) of Argentina is a medium-sized patagornithine phorusrhacid. It was a member of the predominantly South American radiation of ‘terror birds’ (Phorusrhacidae) that were apex predators throughout much of the Cenozoic. A previous biomechanical study suggests that the skull would be prepared to make sudden movements in the sagittal plane to subdue prey.

Methodology/Principal Findings: We analyze the flexion patterns of the neck of Andalgalornis based on the neck vertebrae morphology and biometrics. The transitional cervical vertebrae 5th and 9th clearly separate regions 1–2 and 2–3 respectively. Bifurcate neural spines are developed in the cervical vertebrae 7th to 12th suggesting the presence of a very intricate ligamentary system and of a very well developed epaxial musculature. The presence of the lig. elasticum interespinalle is inferred. High neural spines of R3 suggest that this region concentrates the major stresses during downstrokes.

Conclusions/Significance: The musculoskeletal system of Andalgalornis seems to be prepared (1) to support a particularly big head during normal stance, and (2) to help the neck (and the head) rising after the maximum ventroflexion during a strike. The study herein is the first interpretation of the potential performance of the neck of Andalgalornis in its entirety and we considered this an important starting point to understand and reconstruct the flexion pattern of other phorusrhacids from which the neck is unknown.

Introduction

Neck length and neck posture are both relevant to give the appropriate position of the head during all kinds of behaviors [1,2]. For activities such as mating or defense the neck has the ability to perform more complex movements [3] linked to the cervical morphology. The head centre of gravity over the feet is deeply correlated with the erect posture on land and also depends on the movements of the neck. In normal stance birds (and all amniotes) maintain vertical its cervical column [4,5].

Size, morphology and number of vertebrae are three important variables involved in the motion of the neck [6,7,8]. Birds have a highly variable number of cervical vertebrae (9–11 in some parrots, 23–25 in swans) but, whatever the number, their necks look shorter or longer depending on the neck-folding (Fig. 1). All birds have a flexible “S” shaped neck but the flexion capacity varies between them and for sure, the arrangement is a consequence of an intricate system made by muscles, ligaments and complex articulations. As flexion increases, the distance between the head and the thorax will be shorter and, under these circumstances the external profile of the animal’s neck region shortens [9].

Each vertebra articulates with the adjacent through a saddle-shaped region (articulatio intercorporalis in the sense of [10]) located at the base of the vertebral body (corpus vertebrae) (Fig. 2) and two additional sliding surfaces located in the top (articulatio zygapophy- sialis cranialis and caudalis) [8]. This morphology allows large dorsoventral movements of the neck but prevents rotation, which is generally small.

Boas [6] divided the cervical column of birds in three main regions according to the dorsoventral bending; the most rostral is the region 1 in which ventral flexion is prevalent, region 2 in which dorsal flexion prevails and region 3 in which both dorsal and ventral flexion are limited. The boundaries between these regions are given by transitional vertebrae that have a particular morphology. Knowing the vertebral morphology of each region one can infer what kind of movements could be possible.

Andalgalornis steulleti (Kraglievich 1931), from the upper Miocene–lower Pliocene (=6 million years ago) of Argentina, was a medium-sized patagornithine phorusrhacid of about 40 kg body mass, 1.4 m height, and 370 mm total skull length [11,12]. Together with the long legs, the atypical large skull with high and narrow beak is a characteristic feature of all phorusrhacids [13]. The Phorusrhacidae are often regarded as apex predators of the
South American Tertiary environments sharing this role with large now-extinct marsupials [14]. Popularly known as “terror birds” (a name applied at least to the giant forms) there are no close modern analogs which facilitate interpretations about their biology.

Here we analyze the flexion patterns of the neck of *Andalgalornis* based on the neck vertebrae morphology and biometrics. Complete columns among phorusrhacids remains are scarce and in most cases in those specimens where the column is quite complete, the vertebrae are severely damaged. So far, the column of *Andalgalornis* is the most complete and well preserved known of a terror bird. The study here is the first interpretation of the potential performance of the neck of *Andalgalornis* in its entirety and we considered this an important starting point to understand and reconstruct the entire neck of other phorusrhacids from which the neck is unknown or severely damaged.

The vertebral morphology of *Andalgalornis* has not been extensively described in the literature. In their original description of the only complete column known for the species, Patterson and

---

**Figure 1. Comparison between external neck guises in selected birds.** 1 toucan, 2 owl, 3 eagle, 4 tinamous, 5 seagull, 6 seriema, 7 ibis, 8 egret, 9 greater rhea, 10 swan. doi:10.1371/journal.pone.0037701.g001

**Figure 2. Features of the 5th cervical vertebrae in the seagull *Larus* sp.** A, cranial view; B, lateral view; C, dorsal view. doi:10.1371/journal.pone.0037701.g002
Kraglievich [15] recognized the presence of 17 presynsacral vertebrae (11 cervicals, 2 cervicodorsals and 4 dorsals) of which 13 form the neck. Their description is limited to the size comparison of the vertebral body and only in a few vertebrae did they also compare the height of the neural spine.

We will dwell on the comparison between several extant birds of different size and habits in order to establish a correlation between cervical morphology and mode of life. Such anatomical analysis is not available for any phorusrhacid, while it is a first requirement for any quantitative analysis in comparative, functional or ecological morphology.

**Results**

**Anatomical Description**

We use the term cervical vertebrae to mean the true cervical (without ribs or with fused ribs) and the cervicodorsal vertebrae to refer to vertebrae with free ribs that do not contact with the sternum [16]. Skeletal characters used in the descriptions are shown in Figure 2 while Figure 3 exhibits the measurements and angles analyzed in this paper. Our treatment here follows the concept of regions defined by types of mobility, which are separated by transitional vertebrae: fifth cervical vertebra and ninth cervical vertebra in *Andalgalornis*. Comparisons [between brackets] with other phorusrhacids were made when possible.

**Region 1 (R1)**

This region includes cervical vertebrae one to five. In general, they are robust and short (I2 with lower values than those from the R2), with weakly developed *facies articularis cranialis* (I1 with smaller values than R2), and the *facies articularis caudalis* exposed when viewed dorsally (Fig. 3, 4). This is a more massive (cohesive) region of the neck that is capable of downward (ventral) flexion but only slightly upward (dorsal) flexion is possible (see above and Table 1).

The atlas (first vertebra) of *Andalgalornis* is unknown.

The second cervical vertebra, the axis, is badly deformed mainly in the corpus. It appears to be long and ventrally rough with three ridges, one central and two laterally placed [in *Procariama simplex* Rovereto 1914 the corpus is smooth, and there is only one central ridge]. The *zygapophyses cranialis* are small. Viewed laterally, behind the *zygapophysis*, the *ansa costotransversaria* is located [absent in *P. simplex*] with small *processus costalis*. The *zygapophyses caudalis* are large, and ventrally concave. The *facies articularis caudalis* is dorso-caudally oblique [strongly dorso-cranially oblique in *Psilopterus* Moreno and Mercercat 1891 *sensu* [17]. The *processus spinosus* is high, thick and has a wide posterior fossa for ligament insertion.

The *corpus* of cervical vertebra 3 is narrow and ventrally convex [flat in *P. simplex*]. The *facies articularis cranialis* is thin viewed cranially [broad in *P. simplex*] and extensive in the ventral region. The *zygapophyses cranialis* face craniodorsally and are slightly convex. The *ansa costotransversaria* is robust and craniocaudally disposed, the tuberculum ansae is thick but slightly marked [absent in *P. simplex*]. The *processus costalis* is small. The *facies articularis caudalis* is proportionately smaller and widely dorsally directed. In dorsal view, the general shape of the vertebra is trapezoidal, wider than longer, with the *concavitas lateralis* hidden. The *processus spinosus* is high and wide, caudally located, with an elliptic deep fossa placed caudally which possibly corresponds to the insertion of the *ligamentum elasticum interlaminare*. The high *tori dorsalis* are latero-caudally located above the most lateral sector of the *zygapophysis caudalis*. The latter is large and slightly concave.

Cervical vertebra 4 presents the *corpus* proportionately larger, with a ventrally-directed crest (*processus ventralis corporis*) descending from the midline of the *centrum* [crest absent in *Procariama*]. The *processi caroticus* are low, slender and extended ventrally around the

---

**Figure 3. Schematic vertebrae showing the measurements and angles taken.**

A. cranial view; B. dorsal view; C–D, lateral view. ML maximum length, MH maximum height, LF in dorsal view, distance between the *facies articularis cranialis* and caudalis, L4 in dorsal view, distance between the cranial and posterior borders of the *foramen vertebralis*, L4 in ventral view, distance between the cranial and posterior borders of the *foramen vertebralis*, CW cranial width, L1, L4 and FW1 are following Guinard et al. (2009: Fig. 2), I1 development and cranial projection of *facies articularis cranialis* (*I*1 = L4-L1/L4), I2 elongation index (*I*2 = L4/FW1), the later in the same way as Guinard et al. (2009), da dorsal angle, va ventral angle, c. circa because deformation.

doi:10.1371/journal.pone.0037701.g003
Figure 4. Complete neck of *Andalgalornis steulleti* FMNH - P14357 and selected vertebrae. A, neck with R2 in maximum dorsal flexion. R1 region 1, R2 region 2, R3 region 3, VT cervico dorsal vertebrae; B, vertebra 3 (R1); C, vertebra 6 (R2); D, vertebra 11 (R3). Top line, dorsal view; Midline, right lateral view; Bottom line, cranial view. Scale 1 cm.

doi:10.1371/journal.pone.0037701.g004
facies articularis cranialis. The facies articularis caudalis is vertical [oblique in Polioptera]. Viewed dorsally, the vertebra is trapezoidal. The processus spinosus is high and narrow and is located on the caudal portion of the vertebra. A posterior and very deep fossa is presented on the spine but narrower than that of the vertebra 3. Caudally, there is a small notch in the bar that connects both zygapophysis caudalis. The tori dorsalis are slightly proximal and lateralized and are higher than previous vertebrae.

Transitional Vertebra between R1–R2 (Fig. 5)

The fifth cervical vertebra of Andalgalornis is the last of the first region [identical condition in P. simplex and Mesembriornis milneedwardsi; Moreno 1889] presenting a transitional morphology [18]. The vertebral body is longer and broader [narrower in Procariama] than in the preceding vertebrae widening caudally. The elongation index is greater than that of the cranial vertebrae but lower than that of the vertebrae of R2 (Table 2). Ventrally, the corpus is slightly concave. The facies articularis cranialis disposse more vertically in relation to the cranial vertebrae (II greater than that of vertebrae 3 and 4). The facies articularis caudalis is vertically disposed. The zygopophyses cranialis are vertically disposed, with a slope directed cranio-medially. Caudolaterally to the facies articularis cranialis are the well process transversus. Baumel and Wiemer [10] pointed out that cervical vertebrae have processus transversus indistinguishable from the ansa costotransversaria in mature birds. Skull morphology suggests that Andalgalornis steulleti FM-P14357 is an adult specimen and could even be of advanced maturity. The condition of having the processus transversus distinguishable of the ansa costotransversaria is indeed a feature rare among mature birds [10]. The ansa costotransversaria is robust and long cranio-caudally, with a tuberculum ansae with low development [absent in Procariama]. Dorsally it is possible to observe the concavitas lateralis due to the reduction of the lamina lateralis arcus (that connects zygopophyses cranialis and zygopophyses caudalis). The processus spinosus is centrally placed and is smaller than in the preceding vertebrae. The caudal lacuna interzygapophysialis is more pronounced than in preceding vertebrae. It presents a “V” shape with its apex located at the same level than the small tori dorsalis [absent in Procariama], and ends in the most caudal section of the zygopophyses caudalis. This shape allows the vertebral accommodation during the dorsal flexion. The zygopophyses caudalis are small and their facets face latero-ventrally [similar condition in M. milneedwardsi].

| Region  | Vertebras | Ventral angle | Dorsal angle |
|---------|-----------|---------------|--------------|
| R1      | V3–V4     | –             | –            |
|         | V4–V5     | 30°           | 20°          |
| R2      | V5–V6     | 18°           | 27°          |
|         | V6–V7     | 11°           | 40°          |
|         | V7–V8     | 9°            | 43°          |
|         | V8–V9     | 4°            | 45°          |
| R3      | V9–V10    | 0°            | 36°          |
|         | V10–V11   | 10°           | 30°          |
|         | V11–V12   | 5°            | 27°          |
|         | V12–V13   | 0°            | 10°          |

- Not measured.

doi:10.1371/journal.pone.0037701.t001

Region 2 (R2)

In Andalgalornis steulleti vertebrae of R2 do not exceed 35% of all neck vertebrae (Table 3), including vertebrae 6 to 9.

In the sixth cervical vertebra (the first of this region) dominates the dorsal flexion (between vertebrae 5 and 6, Table 1). The facies articularis cranialis (Fig. 4) is wider than the anterior vertebrae and is perpendicular to the longitudinal axis of the neck (L2 higher than previous vertebrae). The zygapophyses cranialis have a medial slope and are smaller than the ones of the more cranial vertebrae. The processus transversus are vertical and lateral to the zygopophyses cranialis [oblique in Procariama]. These processi are posteriorly connected with the base of the tori dorsalis by thin projections. These projections are present in cervical vertebrae 6, 7, 8 and 9 of A. steulleti [barely pronounced in vertebra 6 and highly developed in vertebrae 8, 9, 10 and 11 in P. simplex; vertebra 7 is broken at the level of the processus and it is not possible to establish the presence of this feature; well developed in vertebrae 6 to 9 in M. milneedwardsi; occur in vertebrae 7, 8, 9 and 10 in the Cariamidae Chunga]. The ansa costotransversaria is robust and has a strong development of the tuberculum ansae and the crista laterales (for the muscle intertransversarii attachment). The processus caroticus is well marked and ventromedially directed. The facies articularis caudalis is wide. Dorsally, the processus spinosus is disposed caudally, allowing the accommodation of the previous vertebra during dorsal flexion. The lamina lateralis is coincident with the vertebral body [not in Procariama] and the concavitas lateralis is widely developed. The processus spinosus is reduced, like the tori dorsalis that constitutes low prominences.

The cervical vertebrae 6, 7 and 8 are very similar in morphology (Fig. 4). However, the 7th is different in that the ansa costotransversaria presents a further development of the tuberculum ansae and the crista lateralis, the processus caroticus are more robust, the zygopophyses cranialis are more inclined medially, the crista transverso-obliqua link the much reduced processus spinosus with the tori dorsalis and the area of insertion of the lig. elasticae interlaminarum is deeper, forming a furrow from the neural arch to the lacuna interzygapophysialis.

The elongation index of the 8th cervical vertebrae is less than those of the 6th and 7th. This vertebra presents the facies articularis caudalis not well extended laterally, adopting a square like form, the processus spinosus is small and is bifurcated [same condition in P. simplex] (see above). The zygopophyses caudalis are greatly separated from each other, increasing the angle of the lacuna interzygapophysialis.

Transitional Vertebra between R2–R3 (Fig. 5)

The cervical vertebra 9 also shares the same morphology as the cervical vertebrae of the R2, but is wider (see AW in Table 2). The processus spinosus is bifurcated, the space between both spines is broad and the entire structure is higher than in the cranial vertebrae [18]. The processi costalis are robust being much extended caudally than in vertebrae of R2, almost reaching the end of the vertebral corpus. These processi are present in all the vertebrae previously described, but they are broken or eroded and comparative development can not be established. The caudal lacuna interzygapophysialis is bounded by three sides, taking a trapezoidal shape [triangular P. simplex], with the longest side between the caudal ends of the zygopophyses caudalis.

Region 3 (R3)

This region includes vertebrae 10 to 13. Patterson and Kraglievich [15] described the 10th vertebra as a cervical one because it has an ansa costotransversaria instead of a floating rib. However, the presence of processus vertralis corporis and the
Figure 5. Transitional vertebrae of *Andalgalornis steulleti* FMNH - P14357. A, E dorsal view; B, F ventral view; C, G right lateral view; D, H cranial view. Scale 1 cm.
doi:10.1371/journal.pone.0037701.g005
differences in the *lacuna interzygapophysialis* between this vertebra and the 9th, point out that the 10th vertebra is the first one of the third region.

The 10th cervical vertebra (Fig. 4) is wider than previous vertebrae in dorsal view. The *processus spinosus* is stout, caudally placed and has a bigger area for the insertion of the *lig. elasticum interlaminare*. There is no *processus caroticus*, but *processus ventralis corporis* is present. The angle measurements (Table 1) indicate a limited dorsal and ventral flexion for the R3 (see next section).

The 11th cervical vertebra is very similar to the 10th. It has a compressed *processus ventralis corporis*, well extended and cranially projected. The *processus spinosus* is big and caudally placed. The insertion area of the *lig. elasticum interlaminare* is wide and creased.

The 12th cervicodorsal vertebra was described by Patterson and Kraglievich [15] as the first cervicodorsal vertebra. This one is stouter than the preceding ones and the main features are the absence of fused ribs (which are free) and the wide development of the *processus spinosus*, with bigger and creased areas for ligament insertion.

Vertebrae with Bifurcated Neural Spines (BNS)

The fifth caudal cervical and first cervicodorsal vertebrae [19] of *Andalgalornis steulleti* have low bifurcated neural spines (*metapophyses sensu* [20]; this means cervical vertebrae 7–9 of R2, 10 and 11 of R3, and the cervicodorsal vertebra 12 of R3. Altogether, 38% of the total cervical vertebrae number of the neck has BNS (Fig. 4). For comparative purposes Table 3 shows the distribution and proportions of vertebrae with BNS in other birds.

Vertebrae with Bifurcated Neural Spines (BNS)

The fifth caudal cervical and first cervicodorsal vertebrae [19] of *Andalgalornis steulleti* have low bifurcated neural spines (*metapophyses sensu* [20]; this means cervical vertebrae 7–9 of R2, 10 and 11 of R3, and the cervicodorsal vertebra 12 of R3. Altogether, 38% of the total cervical vertebrae number of the neck has BNS (Fig. 4). For comparative purposes Table 3 shows the distribution and proportions of vertebrae with BNS in other birds.

Vertebrae with Bifurcated Neural Spines (BNS)

The fifth caudal cervical and first cervicodorsal vertebrae [19] of *Andalgalornis steulleti* have low bifurcated neural spines (*metapophyses sensu* [20]; this means cervical vertebrae 7–9 of R2, 10 and 11 of R3, and the cervicodorsal vertebra 12 of R3. Altogether, 38% of the total cervical vertebrae number of the neck has BNS (Fig. 4). For comparative purposes Table 3 shows the distribution and proportions of vertebrae with BNS in other birds.

Table 2. Measurements and indices of 3rd to 13th *Andalgalornis steulleti* cervical vertebrae. Abbreviations in Figure 3.

|        | R1   | R2   | R3   | BNS | T  |
|--------|------|------|------|-----|----|
| V      | 3    | 4    | 5    | 6   | 7  | 8   | 9   | 10  | 11  | 12  | 13  |
| ML     | 68.92| 69.13| 78.92| 79.64| 72.19| 69.63| 69.28| 69.70| 72.55| 67.23| 78.83|
| MH     | 54.25| 57.77| 57.73| 52.46| 53.78| 61.15| 65.65| 73.74| 87.6 | 98.24| 107.27|
| L1     | 44.82| 45.35| 46.49| 40.06| 40.51| 31.63| 34.45| 32.56| 36.87| 39.25| 45.94|
| LF     | 47.94| 54.39| 56.29| 56.77| 56.60| 53.50| 49.78| 54.12| 55.38| 65.71| 58.30|
| L4     | 54.42| 52.47| 56.40| 59.78| 60.63| 59.45| 58.96| 59.20| 57.61| 51.30| 45.91|
| AIW    | 70.23| 60.99| 72.73| 73.31| 69.70| 73.43| 75.06| 78.36| 81.7 | 93.3 | 85.66|
| Fv1    | 70.26| 59.82| 59.55| 57.11| 50.15| 51.83| 54.86| 59.60| 56.97| 62.85| 50.14|
| Cw     | 75.84| 69.4 | 47.27| 43.94| 42.33| 52.52| 56.39| 56.62| 52.85| 52.25| 43.7 |
| I1     | 0.065| 0.166| 0.174| 0.294| 0.284| 0.408| 0.307| 0.398| 0.334| 0.402| 0.212|
| I2     | 0.774| 0.887| 0.947| 1.046| 1.208| 1.014| 0.973| 1.011| 0.816| 0.915|    |

Table 3. Comparison of the number of vertebrae per region of *Andalgalornis* with selected taxons with and without bifurcated neural spines.

|        | R1   | R2   | R3   | BNS | T  |
|--------|------|------|------|-----|----|
| Andalgalornis | 5v  | 4v  | 4v  | 5v  | 13  |
| Rhea   | 5v  | 9v  | 3v  | 6v  | 17  |
| Chunga | 5v  | 6v  | 2v  | absent | 13 |
| Theristicus | 5v  | 7v  | 5v  | 8   | 17  |
| Buteo  | 5v  | 4v  | 5v  | absent | 14 |
| Tyto   | 5v  | 4v  | 4v  | absent | 13 |
| Ramphastos | 5v  | 4v  | 4v  | absent | 14 |
| Cygnus | 12v | 9v  | 2v  | absent | 23 |
| Nycticorax | 5v  | 8v  | 4v  | absent | 17 |
| Rhynchotus | 5v  | 8v  | 3v  | absent | 16 |
| Larus  | 5v  | 5v  | 3v  | absent | 13 |
| BNS vertebrae with bifurcated neural spines, T total vertebrae of the neck, v vertebrae. |

do:10.1371/journal.pone.0037701.t003

do:10.1371/journal.pone.0037701.t002
arrangement is very similar to those of carnivorous birds like the Hawk *Buteo* and the Barn Owl *Tyto*, but very distinct to the seriema *Chunga* a very striking considering that Cariamidae are the closest extant relatives to the Phorusrhacidae [25,26]. On the contrary, in the neck of *Rhea americana*, R2 represents the 53% of the total number of the cervical vertebra and flexion between R2 and R3 is located more caudally, between dorsal vertebra 14 and 15.

Along the neck of *Andalgalornis steulleti*, the corpus length (CL) remains approximately constant recording the lowest values in the 15th vertebra and the highest in the 6th, 7th and 8th vertebrae. The maximum length (ML) noticeably increases in the 5th and 6th vertebrae. However, their value caudally decreases from there. Maximum height (MH) increases from the 6th vertebra, but in the vertebrae 11, 12 and 13 it becomes really high. The anterior width (AW) remains relatively constant with slight increase in the last three vertebrae. Only at the vertebrae 3rd and 4th the anterior width is minor than the caudal width (CW).

Metric properties of the neck of *Andalgalornis* are also similar to that observed in *Buteo* and *Tyto*. In *Tyto*, in the vertebrae 5 and 6 the maximum length dominates over the anterior width. In *Buteo* instead, the anterior width is larger than the posterior one. The last *Andalgalornis’* vertebrae behave like in *Buteo*: the maximum height is major than the anterior width. None of the measured variables behaves in a similar way to the long necks of *Struthio* (see length dimension, Figure 4 [3]) or *Rhea*. Unexpectedly, vertebrae of the neck of the Black Legged seriema *Chunga* metrically behaves very different from the terror bird *Andalgalornis*. The measured variables in the seriemas however, show a pattern similar to *Rhea* and *Rhyynchotus*: the maximum length predominates in all central vertebrae from the fourth to the 12th (from the third to the 13th in *Rhyynchotus* and from the fourth to the 15th in *Rhea*). Both in *Rhea* and *Chunga* the cranial width exceeds the value of vertebra body length only in the last vertebrae (the 13th in *Chunga* and the 14th in *Rhea*). The Gull *Larus* (Charadriiformes, Laridae) shares with *Andalgalornis* the same number of neck vertebrae (13) but the metric behavior of *Larus* is remarkably different from all the species compared; the maximum length is larger in the first nine vertebrae and the cranial width predominates in the remaining ones.

In relation to the angles between vertebrae (Table 1), the only reliable angles values obtained on R1 is between vertebrae 4 and 5 being the ventral angle larger than the dorsal one. The R2 shows significantly low ventral angles values but very high dorsal angles. The dorsal values gradually increase until the articulation between vertebrae 8 and 9. Meanwhile, ventral angles decrease until the articulation of vertebrae 8 and 9. In the R3 the dorsal angles values caudally decrease whereas the ventral angles are variable but low.

**Discussion**

Morphologically, it is relatively easy to distinguish which vertebra corresponds to each region of the neck of *Andalgalornis steulleti*. Cervical vertebrae from R1 ventrally have very developed *facies articularis cranialis*, the *arcus vertebrae* is cranially extended as the *corpus*, the *zygapophysis cranialis* are mediocranially sloping and the *lucana interzygapophysialis* is absent. R2 cervical vertebrae have a low *processus spinosus*, the *concavis lateralis* are deep, the *lucana interzygapophysialis* is very well developed adopting a wedge or “V” shape and ventrally, the *corpus* had *processus caroticus*. On the contrary, vertebrae from R3 have a *processus spinosus* high and laterally expanded, the *lucana interzygapophysialis* is poorly developed and adopts a trapedozoidal shape and the *corpus* had a *crista ventralis corporis*.

It is clear that the external appearance of the neck does not unequivocally reflect the number of vertebrae that compose it, nor does it reflect folding capability. An externally short neck like in *Buteo* has a high capability of folding in the sagittal plane. Because the possibility of high folding is untenable in the terror bird *Andalgalornis*, it should be considered not a short but a medium-long necked bird.

In all birds there is a discontinuous *ligamentum elasticum interlaminare* that connects the *laminae* of adjacent vertebrae [22 and literature cited therein]. This ligament is more developed in the cranial (R1) and caudal (R3) cervical region and helps to keep the head erect when the animal is in neutral position [21]. A second ligament, the *ligamentum elasticum inter-spinale*, is present only in all the birds that have BNS and occupies the space between both neural spines. Based on a topological criterion, we infer the presence of this ligament in *Andalgalornis* (Fig. 6). This ligament helps to support the weight of the neck when the posterior part descends and falls below the level of origin of the ligament.

Concerning the neck myology (Fig. 6), the *m. biventer cervicis* filled in part the space between the neural spines. Both *m. biventer cervicis* (inserted on the dorsal tubercles or dorsal processes below the *zygapophyses caudalis*) and *m. longus colli dorsalis pars caudalis* are the main epaxial muscles of the neck in all extant birds.

Therefore, the presence of bifurcate neural spines in *Andalgalornis* suggests the presence of a very intricate ligamentary system (with both the *lig. elasticum inter-spinale* and *ligamentum elasticum inter-laminare*) and a very well developed epaxial musculature.

So far, bifurcated neural spines have been reported in birds from different lineages and different trophic habits with large skulls (e.g. *Andalgalornis*, *Diatryma*, *Geryornis*) or relatively small size skulls (*Rhea*). Therefore, the presence of BNS in birds is puzzling and warrants further investigation. Outside the clade Aves, the bifurcation of the cervical and dorsal neural spines characterizes several clades of saurischian dinosaurs (e.g., *Flagellicaudata*), as well as several different genera (*Camarasaurus*, *Opisthocoelicaudia*), all of which evolved independently [22]. The biological significance of the BNS is still unknown but this diversity is an interesting topic to explore phylogenetic constraints and possible adaptive significance of the *lig. elasticum inter-spinale* [19].

In *Andalgalornis*, the *occiput* and *condylus occipitalis* position indicates that at least the first two cervical vertebrae of R1 could only be located in line with the major axis of the skull. Based on the osteology, other positions would be anatomically and mechanically untenable. R2 and R3 show a marked prevalence of the ventral flexion (high dorsal angles). Also, the maximum potential dorsiflexion of the cervical vertebrae on the sagittal plane (Table 1) suggests that the head could not be located over the back of the body (Fig. 7). In other words, a position of the head over the level of the scapulae exceeds the possible range of motion during life. From the previously mentioned characters as well as the manual manipulation of the cervical vertebrae, it is possible to infer that the neck does not take a complete vertical upright position in a stationary situation (Fig. 7) as in *Rhea* or *Struthio* (see Figure 11 for comparison in [3]).

Throughout the cervical column, the neural spines vary showing the different possibilities of bending of the three sections. If the spine is longer, the lever arm length (the perpendicular distance from the line of action to the pivot) increases magnifying the effective force on the corpus [27]. When the neck is flexed downward, the neural spines transmit tension stresses whereas the vertebral centra resist compression stresses. Because in the cervical column of *Andalgalornis* the neural spines are evident in R1, lower in R2, and conspicuously increases in the R3, it seems possible that during down strokes, R3 concentrates the major stresses.
Figure 6. Schematic hypothetical lateral reconstruction of neck muscles and ligaments in *Andalgalornis* based on sites of muscle origin and insertions identified in the holotype and according [10,22,30]. A–C represents successively superficial layers. Arrow indicates cranial direction.
doi:10.1371/journal.pone.0037701.g006
It is known that the dorsal neck muscles are related with the dissipation of tension stresses during a neutral position of the animal [28]. Judging by the highly developed neck system, it can be supposed that the neck of *Andalgalornis* could be related to two main functions: supporting a particularly big head during normal stance and helping the neck (and the head) to rise after the maximum ventroflexion during a down stroke in the attack of prey. A finite element analysis applied to *Andalgalornis steulleti* [12] suggests that the skull would be prepared to make sudden movements in the sagittal plane to subdue prey. Based on these results, it is interpreted that during an attack phase -downstroke- the head of the animal would quickly move to achieve the maximum ventro-flexion. To start a new phase it is essential that the head and the neck quickly and effectively recover their initial position in height. Piecing altogether, the *lig. elasticum interespinale* and *lig interlaminare* as well as the epaxial muscle (mainly the *m. biventer cervicis*) helped the neck (and the head) to rise after the maximum ventroflexion during a strike.

From the morphology of the vertebrae as a whole and particularly of the transitional vertebrae the cervical flexion patterns can be established. Morphological correlates with activity and capability are difficult to pin down. But combining our results with other evidence is possible to infer some paleobiological aspects of *Andalgalornis*. Our study also indicates which movements are not possible or restricted mainly by mechanical constraints. But those movements that have no restrictions were not necessarily employed by the animal. In other words, our results indicate what kind of movements *Andalgalornis* was able to do (in a vertical plane), but this does not necessarily imply that they were done.

The study performed here was uniplane and movements of the neck are multidimensional by nature. Future studies should consider all possibilities of the behavior of *Andalgalornis*. Also, this
study is the first one to analyze the potential functioning of a phorusrhacid's neck in its entirety. In a broad sense, we may assume that the necks of other large phorusrhacids (big headed birds with high and compressed beaks) would respond similarly to that of Andalgalornis. We are now one step closer to understanding the phorusrhacids' lifestyle.

Materials and Methods

Cervical vertebrae of Andalgalornis steulleti were studied at División Paleontología Vertebrados of Museo de La Plata (cast MLP 604-I) and Field Museum of Natural History (FM-P14525, Mesembriornis milneedwardsi MAGN 5944 and Psilopterus lemoinei AMNH 9157, AMNH 9257 and YPM-PU15402 were studied for comparison. Nine modern species representing a wide range of external neck length were evaluated: long necks Cygnus melancoryphus MLP 688, Nycticorax nycticorax MLP 69, Theristicus caudatus MLP 474, and Rhea americana MLP 650; medium neck length Rynchotus rufescens MLP 30 and Lanius sp. MLP 577, short necks Buteo magnirostris MLP 362, Tyto alba MLP 33 and, Ramphastos toco MLP 314 (Fig. 2). Also, the medium neck length Cariamidae Chunga burmeisteri MAGN 2351a, MLP 52 was included in the analysis because of its close phylogenetic relationships with phorusrhacids. The osteological samples for comparison studied come from the Museo de La Plata (MLP) and Museo de Ciencias Naturales “Bernardino Rivadavia” (MACN).

Descriptions and observations are organized around the three regions proposed by Boas [6]. Two transitional vertebrae (TV) separated region 1 (R1) from region 2 (R2) and region 2 from region 3 (R3). Studying penguins Guinard et al. [9] applied an alternative approach identifying biomechanical zones within the cervical system they call “modules” [following 29]. In the same paper they propose that their method can be applied to any other avian taxa [9]. However, we prefer to apply here the classical proposal of Boas [6] because regions are easy to delimit and to compare between the set of birds chosen here.

The anatomical terminology is given in the text with their Latin denomination (as used by Baumel and Witmer [10]) and English equivalents.

Ten biomeetric measurements were taken directly from the material using a digital caliper (Fig. 3). Each measurement was taken three times, averaged, and the standard error was calculated. When the error was greater than 0.5 the three measurements were repeated. Two indices were calculated: the development and cranial projection of facies articularis cranialis and the elongation index, the latter in the same way as Guinard et al. [9]. The former gives an idea of the dorsal folding of the neck, e.g. vertebrae with low values have been weakly folded and the latter is an expression of the cohesion of the neck, e.g. regions formed by vertebrae with low values will be more cohesive.

We applied the zygapophyseal alignment model [24] to establish the possible range of movement of the neck of Andalgalornis. The cervical vertebrae were articulated and manually manipulated on a flat horizontal surface parallel to the sagittal plane, to recreate the range of allowable dorsoventral flexion. Therefore, for the purpose of modeling flexion and motion we reduced the head-neck complex to a planar multi-element system. The intervertebral space was estimated as 1 mm and the minimum functional overlap of the zygapophysis as 3 mm. However, there is no direct way of knowing how much cartilage would originally have been present in a neck of a phorusrhacid. In this sense, our reconstructions of possible neck poses are conservative. Photographs were taken showing the maximum neck flexion capacity between regions, concentrating on the bending at the level of transitional vertebrae. We measured with a goniometer the dorsal and ventral angles between one vertebra and the next one when vertebrae of the neck are assembled directly on the material. During the dorsal flexion (caudal loop or kyphotic), dorsal angles exhibit low values. On the contrary, during ventral flexion (rostral loop or lordotic), the ventral angles show low values.

The material was photographed using a reflex photographic camera. The images were processed with Adobe Photoshop CS3 software to produce the final figures.

Acknowledgments

We are indebted to J. Wilson who provided helpful comments on earlier versions of the manuscript. We also thank A. Evans and two anonymous reviewers for their comments. We thank W. Simpson and J. Holstein (FMNH), L. Chiappe and P. Johnson (NHM), A. Kramarz (MACN) and M. Reguero (MLP) are thanked for allowing access to collections under their care.

Author Contributions

Conceived and designed the experiments: CPT RdM FJD. Performed the experiments: CPT RdM FJD. Contributed reagents/materials/analysis tools: CPT RdM FJD MP. Wrote the paper: CPT RdM FJD MPB.

References

1. Zweers G, Bout R, Heidweiller J (1994) Motor organization of avian head-neck system. In: Davies M, Green P, eds. Perception and Motor control in birds. Berlin: Springer-Verlag pp 201-221.

2. Bout R (1997) Postures of the avian cranio-cervical column. J Morph 231: 207-295.

3. Drezinski G, Christian A (2007) Flexibility along the neck of the ostrich (Struthio camelus) and consequences for the reconstruction of dinosaurs with extreme neck length J Morph 268: 701-714.

4. Vidal PP, Graf W, Berthoz A (1986) The orientation of the cervical vertebral column in unrestrained awake animals. Exp Brain Res 61: 549-539.

5. Taylor MP, Wedel MJ, Nash D (2008) Head and neck posture in sauropod dinosaurs inferred from extinct animals. Acta Palaeontol Pol 53: 215-220.

6. Boas JEV (1929) Biologisch-anatomische Studien Huber den Halb der Vogel. D Kg Danse Vidensk Selk Skrifter Naturvidensk Mathem 9: 105-222.

7. Van der Leeuw AHJ, Bout RG, Zweers GA (2001) Evolutionary morphology of the neck system in ratites, fowl, and waterfowl. Neerh J Zool 51: 243-296.

8. Van der Leeuw AHJ, Bout RG, Zweers GA (2001) Evolutionary morphology of the neck system in ratites, fowl, and waterfowl. Neerh J Zool 51: 243-296.

9. Guinard G, Marchand D, Courant F, Gauthier-Clerc M, Le Béhére C (2010) Morphology, ontogenesis and mechanics of cervical vertebrae in four species of penguins (Aves: Spheniscidae). Polar Biol 33: 907-922.

10. Bauml JJ, Witmer LM (1993) Osteología. In: Bauml JJ, King AS, Brezaile JE, Evans HE, Van den Barge JC, eds. Handbook of Avian Anatomy: Nomina Anatomica Avium. Cambridge: Nuttall Ornithological Club. pp 65-132.

11. Tambussi CP, Chiappe ML, Porro M (2010) Alguns aspectos biomecânicos de a locomoción dos forróceros (Aves, Griformes). Ameghiniana 49: 141-165.

12. Degrange FJ, Tambussi CP, Moreno K, Witmer LW, Wroe S (2010) Mechanical Analysis of Feeding Behavior in the Extinct “Terror Bird” Andalgalornis steulleti (Griformes: Phorusrhacidae). PLoS ONE 5: e11356.

13. Degrange FJ, Tambussi CP (2011) Re-examination of Psilopterus lemoinei (Aves, Phorusrhacidae), a late early Miocene terror bird from Patagonia (Argentina). J Vert Paleon 31: 1080-1092.

14. Tambussi CP, Noriega JI (1996) Summary of the Avian Fossil Record from Southern South America. In: Arragón G, ed. Contributions of the southern south America to vertebrate paleontology. Münster: Verlag. pp 287-295.
15. Patterson B, Kraglievich L (1960) Sistemática y nomenclatura de las aves fororracoideas del Plioceno Argentino. Publ Mus Mun Cien Nat Trad Mar del Plata 1: 1–51.
16. Zusi RL (1962) Structural Adaptations of the Head and Neck in the Black Skimmer. Cambridge: Nuttall Ornithological Club. 101 p.
17. Sinclair W, Farr M (1932) Aves of the Santa Cruz beds. In Scott W, editor. Reports of the Princeton University expeditions to Patagonia (1896–1899). Volume 7. Princeton University, New Jersey. pp 157–191.
18. De Mendoza RS, Tambussi CP, Degrange FJ (2009a) Morfología de las vértebras cervicales de transición entre regiones en Andalgalornis steulleti (Kraglievich, 1931) (Aves: Phorusrhacidae). Ameghiniana 46: 73R.
19. De Mendoza RS, Tambussi CP, Degrange FJ, Picasso MBJ (2009b) El arco neural bifido del cuello de Aves fosiles: inferencias acerca del sistema ligamentario asociado. Ameghiniana 46: 19R.
20. Osborn HF, Mook CC (1921) Camarasaurus, Amphicoelias, and other sauropods of Cope. Memoirs Am Mus Nat Hist 3: 247–387.
21. Wedel MJ, Sanders RK (1999) Comparative anatomy and functional morphology of the cervical series in rats and sauropods. J Vert Paleont 19: 83.
22. Tsuihiji T (2004) The ligament system in the neck of Rhea americana and its implication for the bifurcated neural spines of sauropod dinosaurs. J Vert Paleont 24: 165–172.
23. Dozo T, Bouza P, Moni A, Palazzesi L, Barreda V, et al. (2010) Late Miocene continental biota in northeastern Patagonia (Península Valdés, Chubut, Argentina). Palaeog Palaeoclim Palaeoecol 297: 100–109.
24. Christian A, Dzemski G (2007) Reconstruction of the cervical skeleton posture of Brachiosaurus brancai Jannench, 1914 by an analysis of the intervertebral stress along the neck and a comparison with the results of different approaches. Fossil Rec 10: 37–48.
25. Andrews C (1896) Remarks on the Stereornithes, a group of extinct birds from Patagonia. Bia 7: 1–12.
26. Alvarenga HMF, Höfling E (2003) Systematic revision of the Phorusrhacidae (Aves: Ralliformes). Papéis Avul Zool 43: 55–91.
27. Vogel S (2000) Ancas y palancas, mecánica natural y mecánica humana. Barcelona: Tusquets. 392 p.
28. Klingenberg CP (2002) Morphometrics and the role of the phenotype in studies of the evolution of developmental mechanisms. Gene 287: 3–10.
29. Vanden Berge JC, Zweers GA (1993) Myologia. In: Baumel JJ, King AS, Breazile JE, Evans HE, Vanden Berge JC, eds. Handbook of Avian Anatomy: Nomina Anatomica Avium. Cambridge: Nuttal Ornithological Club. pp 180–247.
30. Degrange FJ (2012) Morfología del cráneo y complejo apendicular posterior en aves fororracoideas: implicancias en la dieta y modo de vida. Ph.D. Dissertation. La Plata: Universidad Nacional de La Plata. 390 p.