Mechanical Analysis of Feeding Behavior in the Extinct “Terror Bird” *Andalgalornis steulleti* (Gruiformes: Phorusrhacidae)

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**Abstract**

The South American phorusrhacid bird radiation comprised at least 18 species of small to gigantic terrestrial predators for which there are no close modern analogs. Here we perform functional analyses of the skull of the medium-sized (~40 kg) patagornithine phorusrhacid *Andalgalornis steulleti* (upper Miocene–lower Pliocene, Andalgalá Formation, Catamarca, Argentina) to assess its mechanical performance in a comparative context. Based on computed tomographic (CT) scanning and morphological analysis, the skull of *Andalgalornis steulleti* is interpreted as showing features reflecting loss of intracranial immobility. Discrete anatomical attributes permitting such cranial kinesis are widespread phorusrhacids outgroups, but this is the first clear evidence of loss of cranial kinesis in a gruiform bird and may be among the best documented cases among all birds. This apomorphic loss is interpreted as an adaptation for enhanced craniofacial rigidity, particularly with regard to sagittal loading. We apply a Finite Element approach to a three-dimensional (3D) model of the skull. Based on regression analysis we estimate the bite force of *Andalgalornis* at the bill tip to be 133 N. Relative to results obtained from Finite Element Analysis of one of its closest living relatives (seriema) and a large predatory bird (eagle), the phorusrhacid’s skull shows relatively high stress under lateral loadings, but low stress where force is applied dorsoventrally (sagittally) and in “pullback” simulations. Given the relative weakness of the skull mediolaterally, it seems unlikely that *Andalgalornis* engaged in potentially risky behaviors that involved subduing large, struggling prey with its beak. We suggest that it either consumed smaller prey that could be killed and consumed more safely (e.g., swallowed whole) or that it used multiple well-targeted sagittal strikes with the beak in a repetitive attack-and-retrait strategy.

**Introduction**

Phorusrhacids were a predominantly South American radiation of gruiform birds known from middle Paleocene to lower Pleistocene deposits [1,2] and most closely related to seriemas (Cariamidae) among extant birds [3-6]. Their often gigantic body sizes, large skulls, and carnivorous lifestyles have resulted in phorusrhacids being popularly termed “terror birds.” *Andalgalornis steulleti* [7], 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. All clades of phorusrhacids (Patagornithinae, Phorusrhacinae, Brontornithinae, Psilopterinae, Mesembriornithinae; [5]) are considered to be flightless ground predators or scavengers and ranged from 0.9 to 3.0 m in height [1,8]. They are often regarded as apex predators that dominated South American Tertiary environments in the absence of carnivorous placental mammals [1]. However, phorusrhacids were largely contemporaneous with the carnivorous borhyaenid marsupial radiation, which also included some large to gigantic species [9]. To date, hypotheses of feeding ecology in terror birds have been based mostly on the presence of large skulls with hooked beaks and have yet to be supported by appropriate biomechanical studies. Here we perform a biomechanical analysis of the skull of *Andalgalornis steulleti* (FMNH P14357) in a comparative context, using comparative anatomy and a powerful engineering tool, Finite Element Analysis (FEA), to predict the mechanical behavior of its skull.

**Results**

Divergent skull morphology and loss of cranial kinesis

The nature of the connections between the skull bones is important to the analysis of mechanical behavior, because these connections influence the relative movement of areas of the skull (i.e., cranial kinesis) [10] and hence the distribution of stress and...
strain [11,12]. Holliday and Witmer [10] recently proposed four criteria for inferences of intracranial mobility in extinct taxa. Extant neornithine birds (including seriema) generally meet all four criteria, and in the vast majority of extant birds part or all of upper bill can be elevated and depressed relative to the braincase, although these movements have been analyzed mechanically in only a few taxa [13,14,15]. The fact that phorusrhacids are phylogenetically embedded within Neornithes indicates that they evolved from fully kinetic ancestors. The skull of *Andalgalornis*, however, shows some features suggesting loss of intracranial immobility. We assessed the detailed morphology of these features in *Andalgalornis* and some other phorusrhacids using CT scanning, supplementing these findings with observations of other taxa.

A key element of extant avian kinesis is the transformation of bony connections that are immobile in nonavian outgroups into thinned “flexion zones” that permit bending movements [10,13–17]. These flexion zones, satisfying the criterion of “permissive kinematic linkages” [10], typically are found in three locations in extant birds where the bones become strongly dorsoventrally flattened, permitting elevation/depression movements of the upper jaw segment: (1) the craniofacial hinge (within or between the premaxillae, nasals, and frontal), (2) the palatine bones, and (3), the jugal bar. Although the available skull of *Andalgalornis* (FMNH P14357) is somewhat damaged in the region corresponding to the craniofacial flexion zone of other birds, CT scans reveal that enough structure is preserved to show remarkably thick bone in this region relative to that present in extant seriema or other kinetic birds. Moreover, the lacrimal bone is fused rostrally with the upper bill (nasal and maxillary bones) and caudally with the frontal bone, effectively spanning and further restricting movement at the craniofacial flexion zone. Likewise, the flexion zone in the palatine bone clearly has been transformed into a rigid structure in that the bone has become thickened and folded into a U-shaped conformation. Finally, the flexion zone in the jugal bar has also been apomorphically transformed into a thickened beam that is mediolaterally elliptical rather than dorsoventrally flattened. The absence of any of these permissive kinematic linkages individually would be enough to limit or prevent kinesis, but the absence of all three clearly reflects enhanced rigidity and no movement of the upper bill relative to the braincase. Additionally, the lacrimal bone bears an expanded descending ramus that is firmly sutured to the jugal bar, further locking these units together (Figure 1, Text S1, Figure S1), rather than being kinematically separated as in birds retaining cranial kinesis. Other phorusrhacids appear to show a mosaic of secondarily a kinetic attributes, suggesting that, although they all seem to have had reduced or no kinetic mobility, they vary in the extent to which they have rigidified the former flexion zones. For example, specimens of most of the taxa studied reveal the thickened and folded palatines, but some of the smaller-bodied taxa (e.g., psilopterines) may have retained craniofacial and jugal flexion zones closer to the plesiomorphic condition. But all available evidence suggests that large-bodied taxa, such as other patagornithines (e.g., *Andresornis*, *Patagonornis*) and the truly gigantic phorusrhacids (e.g., *Kelenken*, *Devornix*), resembled *Andalgalornis* in transforming all three flexion zones into thickened, reinforced, and immobile junctions.

Zusi [17] regarded cranial kinesis as a universal feature of all birds, but various birds have been considered as akinetic (e.g., ratites [18], penguins [19], colbies [20], toucans [21], hawfinch [22]). This is the first time that akinesis of a gruiform has been reported, and may be among the best documented case for any bird based on anatomical grounds. It is likely that more morphological analyses like the present one will reveal more cases of secondarily reduced or lost kinetic ability in other birds.

**Bite force**

*Published in vivo* bite force data for birds is largely restricted to Passeriformes [23,24], a derived clade of generally small-bodied animals. These studies show that bite force is related to skull morphology and geometry, as well as the capacity of contraction of the jaw muscles [23,25,26]. In Galápagos finches, beak size and especially head width are strongly correlated with bite force, and head size closely correlates with jaw muscle dimensions [27,28]. Passeriforms are obviously a poor model for the estimation of bite forces in phorusrhacids, due to differences not only in phylogeny but also in skull size and morphology.

To predict bite force in *Andalgalornis*, we constructed a regression line ($R = 0.9123; R^2 = 0.8598; p = 0.0000$) using three different data sources: published avian bite force data [23], our own bite force data obtained with a transducer designed to measure forces *in vivo*, and body mass data [23,29,30] (see Materials and Method and Table S1). Based on regression analysis (Figure S2) including the seriema (50 Newtons) and eagle (50 N), we estimate the bite force of *Andalgalornis* as 133 N at the bill tip.

**Finite element analysis**

Finite Element Analysis (FEA) is an engineering approach initially developed to predict stress/strain distributions in manmade objects with relatively simple geometry. With advances in computer technology FEA has emerged as a powerful tool in the investigation of mechanical behavior in complex biological structures, its nondestructive nature rendering it of particular value in paleontological studies [31,32].

The *Andalgalornis* skull model recorded lower mean brick stress (load per unit area) than either *Cariama* or *Haliaeetus* under three extrinsic (i.e., movements of the prey in relation to the head [33]) loads (see Materials and Method and Table 1). We consider it likely that the skull of *Andalgalornis* could have supported larger bite loads than the one tested in this study (133 N), although validation would be required to firmly establish this. In a comparative context, the mechanical behavior of *Andalgalornis* differs further in that mean “brick” element stress values (Von Mises, VM) for the lateral shake simulation were four times greater than those recorded for pullback values. The pullback loading generated the least stress in each of the three skulls. In both *Cariama* and *Haliaeetus*, mean stresses under lateral-shake and normal bite loadings were similar. For *Andalgalornis*, however, the lateral-shake loading induced mean brick stresses that were double those generated by the normal bite.

In all three load cases applied to *Andalgalornis*, three areas concentrated the highest stress: the bill tip (i.e., the hook), the occipital condyle, and the contact between the palatines and pterygoids with the interorbital septum (Figure 2A, B, C). Skull stress was clearly highest under the lateral-shake simulation (Figure 2A). The same is true for *Cariama*, but, in the case of the eagle, the stress was highest in the normal bite than in the lateral-shake extrinsic load. Moreover, in *Andalgalornis*, the stress was highest in the ventrolateral rims of the beak (the toma), the dorsal surface of the nasal bones, the rostral edge of the antorbital fenestra (below the naris), the body of the quadrate bone, in the temporal fossa, and, the lower margin of the zygomatic arch (jugal bar).

**Discussion**

Our comparative FE analyses show that the skull of *Andalgalornis* was best optimized to resist rostrocaudally and dorsoventrally...
directed loads, but less well-adapted to resist laterally directed loadings. VM stress values obtained for *Andalgalornis* are consistent with a very high vertical/transverse index of the rostrum. Moreover, as noted above, the plesiomorphically dorsoventrally flattened flexion zones have been transformed into thickened bony connections that are reoriented into more vertically disposed, transversely compressed bars that would resist sagittal loads well but be relatively weak with regard to lateral loads. In birds with a vertical/transverse index of approximately 1.0 (i.e., beaks as wide as high), such as *Cariama* or *Haliaeetus*, the skull would be expected to be equally resistant to forces directed laterally and vertically, which is consistent with our results. Likewise, both seriema and eagle retain the dorsoventrally flattened flexion zones required for kinesis.

The loss of cranial kinesis in *Andalgalornis* and other large-bodied phorusrhacids is a remarkable derived character, but the interpretation of this loss is complicated by persistent uncertainties about the functional role of cranial kinesis [10,34]. Recent FE studies have revealed that the soft tissues within sutures may function as shock absorbers, acting to lower potentially damaging tensile stress and that loss of kinesis enhances skull rigidity and robusticity, but potentially at the cost of local increases in strains.

**Figure 1. Skull of *Andalgalornis steulleti* (FMNH P1435).** A, dorsal view, B, ventral view, and C, left lateral view, based on volume rendering of CT scan data. Fossil bone is shown in light brown, and rock matrix and plaster restoration are shown in grey. doi:10.1371/journal.pone.0011856.g001
mustelid in part explained by the comparatively longer distance between the palatines, pterygoids, and the septum. This region may act as a stress releaser, compensating at least in part for loss of the kinetic system. A carnivorous lifestyle for phorusrhacids has been widely assumed [1,3,5,36], but more detailed hypotheses of predatory behavior have never been adequately tested. Previous reconstruction and biomechanical analysis of the jaw adductor musculature of Andalgalornis [37] shows that the jaw apparatus is optimized for strength at the expense of speed, and again, our discovery of loss of cranial kinesis is consistent with increased skull rigidity. Any adaptation that results in increased stiffness of the jaw apparatus, such as transformation of mobile flexion zones into rigid struts, will result in some increase in maximum theoretical bite force [31,38]. On the other hand, there may be complex trade-offs between safety factors, the presence or absence of kinesis, and the bone and muscle masses required to achieve any given performance limits. For example, a stiffer and more brittle structure (as in akinesis) may result in higher yield points and some reduction in the muscle mass required to achieve a given bite force, but it may also limit the response time available to modify behavior and avoid catastrophic failure where an organism bites into unexpectedly resistant materials [31]. Regarding the loss of cranial kinesis, a net consequence might be that more bone or bone of greater density (i.e., greater bone mass), might be needed to maintain effective safety margins. In clearly akinesis phorusrhacids such as patagornithines and phorusrhacines, the requisite increased bone mass was presumably tolerated because their large body masses precluded flight and hence the premium on ultra-light structures. Future FE based analyses could aid in the quantification of such interrelated factors.

Table 1. Mean brick element stresses (von Misses, VM) in solved FE models under three load cases; L = lateral shake, N = normal bite, P = pullback.

| Element number | Andalgalornis | Haliaeetus | Cariama |
|----------------|--------------|------------|---------|
| Lateral shake  | 1,080,137    | 860,757    | 775,698 |
| Normal Bite    | 0.570        | 2.412      | 3.235   |
| Pullback       | 0.234        | 0.678      | 0.833   |
| L/N ratio      | 1.803        | 0.941      | 1.015   |
| L/P ratio      | 4.393        | 3.351      | 3.943   |

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[11,12,35]. Thus, we consider it probable that the high stresses recorded in the interorbital septum in our FE model of Andalgalornis are artificial, because the model did not incorporate sutures between the palatines, pterygoids, and the septum. This region may act as a stress releaser, compensating at least in part for loss of the kinetic system.

The skulls of the patagornithines Andalgalornis steullet (= A. ferox [44]; FMNH P14357) and Andrewsonornis abbotti (FMNH P13417) and of the psilopterine Psilorhaptor lemoinei (FMNH P13257) were scanned at O’Bleness Memorial Hospital in Athens, Ohio, using a General Electric (GE) LightSpeed Ultra Multislice CT scanner equipped with the Extended Hounsfield option. Specimens were scanned helically at a slice thickness of 625 μm, 120 kV, and 200–300 mA. Moreover, another skull of Psilorhaptor lemoinei (AMNH 9927) and a skull of the extant red-legged seriema (Cariama cristata, FMNH 105635) were scanned at the Ohio University MicroCT Scanning Facility (OUiCT) on a GE eXplore Locus MicroCT Scanner at a slice thicknesses of 92 μm and 45 μm, 80 kV, and 500 μA. Data were output from the scanner in DICOM format and then imported into 4.1.2 (Mercury-TGS, Chelmsford, MA) for viewing, analysis, and visualization. The CT scan data were analyzed to assess the internal architecture of bony regions relevant to assessments of cranial kinesis, which were supplemented with gross observation of the skulls of these specimens and others.

Materials and Methods

CT scanning

The skulls of the patagornithines Andalgalornis steullet (= A. ferox [44]; FMNH P14357) and Andrewsonornis abbotti (FMNH P13417) and of the psilopterine Psilorhaptor lemoinei (FMNH P13257) were scanned at O’Bleness Memorial Hospital in Athens, Ohio, using a General Electric (GE) LightSpeed Ultra Multislice CT scanner equipped with the Extended Hounsfield option. Specimens were scanned helically at a slice thickness of 625 μm, 120 kV, and 200–300 mA. Moreover, another skull of Psilorhaptor lemoinei (AMNH 9927) and a skull of the extant red-legged seriema (Cariama cristata, FMNH 105635) were scanned at the Ohio University MicroCT Scanning Facility (OUiCT) on a GE eXplore Locus MicroCT Scanner at a slice thicknesses of 92 μm and 45 μm, 80 kV, and 500 μA. Data were output from the scanner in DICOM format and then imported into 4.1.2 (Mercury-TGS, Chelmsford, MA) for viewing, analysis, and visualization. The CT scan data were analyzed to assess the internal architecture of bony regions relevant to assessments of cranial kinesis, which were supplemented with gross observation of the skulls of these specimens and others.

Body mass estimation

The body mass estimate of 40 kg for Andalgalornis was made following previously published methodology [45] and submission of scale plastic models [43, unpublished data].

Bite force

Bite force data were obtained in vivo from captive birds: an adult Cariama cristata at the Zoological Garden of La Plata, Argentina, and an adult black-chested eagle (Geranoaetus melanoleucus) from the Horco Molle Reserve, Argentina. G. melanoleucus is similar in size to H. albicilla, and its bite force was used as a proxy in the FEA of H. albicilla. For each animal, multiple bites at the bill tip were recorded with the transducer, and we use the maximum bite force in the analysis. The safety standards related to health, hygienic, and dietary plans were followed for the proper management of wildlife, issued by the
application and enforcement authority (Dirección de Fauna of Ministerio de Asuntos Agrarios of Buenos Aires Province, Argentina) within the framework of the Provincial Law Decree 12.238/98 and 2308/01. Measurements followed a protocol made in conjunction with the authorities of the establishment which involved measurements taken in the morning before the daily intake, no admission to the environments in which birds are housed and, in all cases, with the assistance of the zookeeper for each animal.

Finite element analysis
Finite element analysis (FEA) has emerged as a powerful tool in the prediction of biomechanical behavior among both extant and
extinct animals [38,46-52]. The protocols implemented in the present study largely follow recently developed methodologies [31,38,51,52].

Homogeneous models (assuming constant material properties throughout) were constructed using the CT data for Andalgalornis (FMNH P14357) with segmentation performed using Mimics (v. 11.01) software (see Figure 3A). A similar procedure was used for the red-legged seriema (Cariama cristata, FMNH 105635) and the White-tailed Eagle (Haliaeetus albicilla, from the University of Austin Digital Morphology Web site: http://www.digimorph.org). Cariama cristata was chosen for comparison because of its close phylogenetic relationships with phorusrhacids and Haliaeetus albicilla because of its predatory lifestyle.

Solid meshing was performed with Strand7 Finite Element software (v. 2.3) (Figure 3B). Models were assembled using 3D low-order four-noded tetrahedral “brick” elements (tet4). In the original surface mesh, maximum and minimum triangle edge lengths were kept at a 1:3 ratio (0.1 geometric error) to minimize discrepancies between triangle dimensions, which can lead to major discrepancies in brick element size in the final solid and introduce artifacts. Brick elements were assigned a single set of material properties as applied in previous studies [31,32], (Young’s modulus of Elasticity \[Y\] = 27.0 GPa; Poisson’s ratio = 0.4; density = 2.19005 T/mm\(^3\)).

Models were restrained to prevent free body motion. Point constraints (restricted to single nodes) can produce pronounced artifacts and inaccurate results [48]. For each loading we have applied more realistic constraints by positioning them within frameworks of rigid links at the occipital condyle as well as at the tip of the beak to more broadly distribute forces in accordance with previously published methods [32]. Statistical analyses were performed using a customized program written in R GUI (by K. Moreno).

Two kinds of load cases were arranged and solved: simulation of muscle forces generated by the jaw adductors and head flexors (intrinsic) and simulation of prey movement relative to the predator, or movement by the predator relative to the prey (extrinsic). To study biomechanical performance, we simulated dispatch (killing) forces: rostrocaudal (“pullback”), as well as dorsoventral and lateral shaking. To perform the simulations, we applied the bite force (intrinsic) obtained for each taxon as a load to the distal end of the premaxillary bone (i.e., the hook), with the occipital condyle fixed. Muscles themselves were not reconstructed. Relative mechanical performance was assessed on the basis of mean Von Mises stress for the skull, as well as visual output from the post-processing software.

It is important to note that in the absence of validation our results can only be considered in a comparative context, and, because the models are homogeneous, our approach addresses the influence of geometry, but not differences in the distribution of material properties. Thus, while our results provide insight into the influence of different geometries on the distribution of stress in a relative sense, such findings cannot be considered as absolute performance values [31,32].

The three taxa considered in this study span a considerable range of cranial dimensions and this raises the issue of scaling. Bite forces have effectively been scaled on the basis of regression and these same forces have been applied as extrinsic forces. In comparative linear-static analyses such as those performed here, scaling will influence the absolute magnitudes of recorded stresses, but not how they are distributed. For example, scaling will not affect interpretations of whether the cranium of Andalgalornis is relatively better or less well-adapted to resist lateral vs dorsoventrally directed forces (the ratios will remain the same). Thus, our interpretations are based on proportional rather than absolute differences in mean stress values and distributions.

**Supporting Information**

**Figure S1** Skull of Andalgalornis steulleti (FMNH P1435). Left lateral view (volume rendering of CT scan data) with slice planes (A-B) displaying the hollow beak cavity. Found at: doi:10.1371/journal.pone.0011856.s001 (3.96 MB TIF)

**Figure S2** Log bite force in birds plotted against log body mass. Passeriforms in blue (dark blue: Estrildidae, sky blue: Fringillidae), Rheidae in green, Accipitridae in red, Cathartidae in rose and Cariamidae in black. See Table S1 for raw data. Found at: doi:10.1371/journal.pone.0011856.s002 (3.39 MB TIF)

**Table S1** Bite force and body mass table. Found at: doi:10.1371/journal.pone.0011856.s003 (0.07 MB DOC)

**Text S1** Andalgalornis’ fossil. Found at: doi:10.1371/journal.pone.0011856.s004 (0.03 MB DOC)

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Terror Bird’s Feeding Behavior

Author Contributions
Conceived and designed the experiments: FJD CPT KM SW. Performed the experiments: FJD CPT. Analyzed the data: FJD KM SW. Contributed reagents/materials/analysis tools: LMW SW. Wrote the paper: FJD CPT KM LMW SW.

References
1. Tambussi CP, Noriega JI (1996) Summary of the Avian Fossil Record from Southern South America. In: Arratia G, ed. Contributions of the southern South American vertebrate paleontology, Müncher Geowissenschaftliche Abhandlungen pp 245–264.
2. Tambussi C, Ubilla M, Perea D (1999) The youngest large carnassial bird (Phorusrhacidae, Phorusrhacinae) from South America (Pliocene-Early Pleistocene). J Vert Paleontol 19: 404–408.
3. Andrews C (1909) On the extinct birds of Patagonia. Trans Zool Soc London 15: 55–86.
4. Livezey BC (1990) A phylogenetic analysis of the Gruiformes (Aves) based on morphological characters, with an emphasis on the rails (Rallidae). Trans R Soc London 353: 2077–2151.
5. Alcarenza HMF, Holling E (2003) Systematic revision of the Phorusrhacidae (Aves: Raillioformes). Papeis Avaluos de Zoologia 43 (4): 55–91.
6. Gasc JP (2002) A new specimen of Salamilia robusta (Aves: Gruiformes: Salmilidae n. fam.) from the Middle Eocene of Messel. Palaeontologische Zeitschrift 76 (2): 305–316.
7. Kruglova L (1931) Contribucion al conocimiento de las aves fósiles de la época araucana-contrariana. Physiol 10: 304–315.
8. Chiappe LM, Berthel S (2006) Skull morphology of giant terror birds. Nature 439: 929.
9. Wroe S, Deroose J, Claes NL, McHenry C (2008) Bite forces and evolutionary adaptations to big biting mammals and the prediction of predatory behaviour in fossil taxa. Proc Biol Sci 275: 619–625.
10. Christiansen P, Wroe S (2007) Bite forces and evolutionary adaptations to feeding ecology in carnivores. Ecology 88 (2): 347–358.
11. Bluem J, Rauhut ORJ (1993) Anthropologie. In: Blumel J, King A, Brezale Evans H, Vanden Berge J, eds Handbook of avian anatomy: Nomina Anatomica Avium Publications of the Nott Archontological Club pp 133–187.
12. Zusi RL (1993) Patterns of diversity in the Avian Skull. In: Hanken J, Hall BK, eds. The Skull Volume 2: Patterns of structural and systematic diversity University of Chicago Press pp 391–437.
13. McDowell S (1948) The bony palate of birds. Part I the Palaeognathae. Auk 65 (4): 520–539.
14. Gussakovskiy WS, Bout RG (2005) Cranial in kinesis in palaegnathus birds. J Exp Biol 208: 3400–3419.
15. Gussakovskiy WS, Vosselman MG, Bout RG (2001) Three-dimensional kinematics of skeletal elements in avian prokinetic and rhynchokinetic skulls determined by roentgen stereophotogrammetry. J Exp Biol 204: 1735–1744.
16. Broom MJ, Rauhut ORJ (2005) Using finite-element analysis to suture morphology: a case study with large carnivorous dinosaurs. Anat Rec 281A: 349–365.
17. Moazen M, Curtis N, O’Higgins P, Evans SE, et al. (2009) Assessment of the role of sutures in a lizard skull a computer modelling study. Proc R Soc B 276: 29–46.
18. Bour T, Zee GA (2001) The role of cranial in kinesis in birds. Comp Biochem Physiol A 131(1): 197–205.
19. Evans H, Vanden Berge J, eds. Handbook of avian anatomy: Nomina Anatomica Avium Publications of the Nott Archontological Club pp 133–187.
20. Schooners J (1963) Some aspects of the cranial morphology of Colius indicus. Annale Universite van Stellenbosch 30, 7 (7): 213–246.
21. Holling E, Gasc JP (1984) Biomécanique du crâne et du bec chez Ramphastidae, Aves, GénoèBes morphologies morphéques. Jahrbuch 130 (2): 235–262.
22. Sims SW (1955) The morphology of the head of the hawks (Cocytius cocytus). Bull Brit Museum, Zool 2 (13): 371–393.
23. van der Meij M, Bout RG (2004) Scaling of jaw muscle size and maximal bite force in tinamous. J Exp Biol 207: 2745–2753.
24. van der Meij M, Bout RG (2006) Seed husking time and maximal bite forces in tinamous. J Exp Biol 209: 3329–3335.
25. Herrell A, Podos J, Huber SK, Hendry AP (2005a) Bite performance and morphology in a population of Darwin’s finches: implications for the evolution of beak shape. Funct Ecol 19: 43–48.
26. Herrell A, Podos J, Huber SK, Hendry AP (2005b) Evolution of bite force in Darwin’s finches: a key role for head width. J Evol Biol 18: 669–675.
27. Herrell A, Van Damme R, vanhooydonck B, De Vree F (2001) The implications of bite performance for diet in two species of lacedizard. Can J Zool 79: 662–670.
28. Herrell A, O’Reilly JC, Richmond AM (2002) Evolution of bite force in turtles. J Evol Biol 15: 1083–1094.
29. Jimenez J, Jakse F (1990) Historia natural del aguja Geomatias melanoleucus: una revisión. El Hornero 13: 97–110.
30. Absourachi A, Holling E, Renu N (2005) Walking kinematics parameters in some palaegnathous and neognathous neotropical birds. Ornithologia Neotropica 16: 471–479.
31. Wroe S, Moreno K, Clausen PD, McHenry CR (2007) High-resolution computer simulation of hominid cranial mechanics. The Anatomical Record 290: 1240–1255.
32. McHenry CR, Wroe S, Clausen PD, Moreno K, Cunningham E (2007) Super-modeled sabercat, predatory behaviour in Smilodon fatalis revealed by high-resolution 3-D computer simulation. PNAS 104: 16010–16015.
33. Preuchoff H, Witzel U (2002) Biomechanical investigations on the skulls of reptiles and mammals. Senckenbergiana lethaea 82 (1): 207–222.
34. Metzger K (2002) Cranial kinesis in lepidosaurs: skulls in motion. In: Aerts P, D’Aout K, Herrell A, Van Damme R, eds. Topics in Functional and Ecological Vertebrate Morphology. Scholten Publishing, Maastricht pp 15–46.
35. Moazen M, Curtis N, O’Higgins P, Evans SE, Fagan MJ (2009) Biomechanical assessment of evolutionary changes in the lepidosaurian skull. PNAS 106: 8273–8277.
36. Sinclair W, Forr M (1932) Aves of the Santa Cruz beds. In: Scott W, ed. Reports of the Princeton University expeditions to Patagonia (1896–1899) Princeton University. pp 157–191.
37. Degrange P (2008) M. adductor mandibulae externus of Andalgalornis steulleti (Aves, Phorusrhacidae): Reconstruction and biomechanics (in Spanish). III Congreso Latinoamericano de Palaeontologia de Vertebrados, Simposio Morfología y Adaptación en Aves Nuevas Herramientas y Conceptos: 76.
38. Wroe S, Huber DR, Lowry M, McHenry C, Moreno K, et al. (2008) Three-dimensional computer analysis of white shark jaw mechanics: how hard can a great white bite? J Zool 276: 336–342.
39. Sustaita D (2008) Musculoskeletal underpinnings to differences in killing behavior between North American Accipiters (Falconiformes: Accipitridae) and Falcons (Falconidae). J Morphol 269: 283–301.
40. Wroe S, McHenry C, Thomason J (2005) Bite club: comparative bite force in big biting mammals and the prediction of predatory behaviour in fossil taxa. Proc Biol Sci 272: 361–362.
41. Christiansen P, Wroe S (2007) Bite forces and evolutionary adaptations to feeding ecology in carnivores. Ecology 88 (2): 347–358.
42. Blair RE, Jones WW (2005) Terror birds on the run: A mechanical model to estimate its maximum running speed. Proc R Soc B 272: 1769–1773.
43. Tanaka CP (1997) Zoon morphology aspects of the phorusrhacid locomotion (Aves, Gruiformes) (in Spanish). Americaniana 34 (4): 541.
44. Patterson B, Kraglevich L (1960) Sistemática y nomenclatura de las aves fororracoides del Ploceno argentino. Publicación del Museo Municipal, Ciencias Naturales y Tradicionales de Mar del Plata 1: 1–51.
45. Campbell KE, Marcus L (1992) The relationship of head bone dimensions to body weight in birds. In: Campbell KE editors. Papers in avian paleontology honoring Pierce Brodkorb. Natural History Museum of Los Angeles County, Science Series 36: 395–412.
46. Rayfield EJ (2004) Cranial mechanics and feeding in Terror Bird’s Feeding Behavior. PLoS ONE 9 August 2010 | Volume 5 | Issue 8 | e11856