

The peculiar syrinx of *Rhea americana* (Greater Rhea, Palaeognathae)

MARIANA BEATRIZ JULIETA PICASSO¹ & JULIETA CARRIL^{2,3}

¹ División Paleontología Vertebrados, Museo de La Plata, Universidad Nacional de La Plata. Paseo del Bosque S/N, La Plata (1900), Buenos Aires, Argentina (Corresponding author); mpicasso(at)fcnym.unlp.edu.ar — ² Cátedra de Reproducción Animal, Instituto de Teriogenología, Facultad de Ciencias Veterinarias, Universidad Nacional de La Plata. Calle 60 y 118 S/N, La Plata (1900), Buenos Aires, Argentina — ³ Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET)

Accepted 01.xi.2013.

Published online at www.senckenberg.de/vertebrate-zoology on 18.xii.2013.

Abstract

This work studied the skeletal and muscular syringeal anatomy of Greater Rhea (*Rhea americana*) throughout postnatal ontogeny, by using muscle staining and differential coloring of cartilage and bone techniques. Anatomical syrinx dissections on four adults (one female and three males) and eight unsexed chicks, were made. The type of the syrinx was tracheobronchial and it was entirely cartilaginous in chicks and in the adult female but showed a partially cartilaginous and osseous pessulus in male adults. A pair of intrinsic muscles were found and the extrinsic muscles were represented by the muscles sternotrachealis and tracheolateralis, and a broad dorsal medial muscular band. The syrinx of Greater Rhea was notable for having a more complex morphology than other Paleognathae birds. Future studies on how *Rhea* produces vocalizations will allow the comparison with other birds, and contribute to the understanding of the evolution of sound-production mechanisms in birds.

Resumen

Se estudió la anatomía esquelética y muscular de la siringe del Ñandú Grande (*Rhea americana*) durante su ontogenia postnatal usando técnicas de tinción diferencial de tejido óseo y cartilaginoso. Se diseccionaron cuatro ejemplares adultos (una hembra y tres machos) y ocho juveniles sin sexar. La siringe fue del tipo tracheobronchial y fue completamente cartilaginosa en juveniles y en la hembra adulta mientras en los machos adultos presentó el pessulus parcialmente cartilaginoso. Un par de músculos intrínsecos fueron encontrados y la musculatura extrínseca estuvo representada por los músculos sternotrachealis y tracheolateralis y una ancha banda muscular de posición dorsomedial. La siringe de *Rhea americana* presentó una morfología más compleja que la descrita para otras aves Paleognatas. La realización de estudios focalizados en conocer cómo el Ñandú Grande produce sus vocalizaciones permitirá realizar comparaciones con otras aves y contribuirá al conocimiento de la evolución de la producción de sonidos en las aves.

Key words

Rheidae, tracheobronchial syrinx, syringeal muscles, pessulus.

Introduction

The Greater Rhea (*Rhea americana*) is the largest of South American birds, reaching 1.5 m in height and weighing 25 kg (FOLCH, 1992). *Rhea* is grouped with *Apertyx* (kiwi), *Casuaris* (cassowary), *Dromaius* (emu)

and *Struthio* (ostrich) in the Ratitae clade. This group together with Tinamidae (Tinamous) comprise the Palaeognathae. These birds are monophyletic and a basal group with respect to Neognathae (e.g. LIVEZEY & ZUSI, 2007;

BOURDON *et al.*, 2009; PHILLIPS *et al.*, 2010; JOHNSTON, 2011). Current studies on the Greater Rhea's anatomy are scarce (as well as in other Palaeognathae birds), despite their importance in the evolution and systematics of modern birds.

The syrinx is the organ responsible for producing vocalizations in birds and knowledge of its anatomy in Paleognathous birds is scarce and often outdated. Since the early 19th century, studies on the anatomy of the syrinx have focused mainly on Neognathae birds (e.g. MÜLLER, 1847; HUXLEY, 1872; CHAMBERLAIN *et al.*, 1968; AMES, 1971; WARNER, 1971, 1972; GABAN-LIMA & HÖFLING, 2006; KABAK *et al.*, 2007; MILLER *et al.*, 2008). On the other hand, FORBES (1881) was the first to study the macroscopic anatomy of this organ on Ratites. Subsequently, WUNDERLICH (1884), BEDDARD (1898) and PYCRAFT (1900) corroborated these anatomical descriptions and pointed some minor anatomical variations (e.g. conformation and numbers of tracheal rings) but did not provide further details. The anatomy of the Palaeognathae syrinx would not be studied again until BEEBES (1925) works with the Variegated Tinamous (*Crypturellus*), and later on, a few others in the ostrich (*Struthio*; YILDIZ *et al.*, 2003) and in Darwin's Nothura (*Nothura*; GARITANO-ZAVALA, 2009). BEAVER (1978) was the only researcher since the early 20th century, to perform a brief and superficial anatomical description of the syrinx of chicks of the Greater Rhea.

The works of FORBES (1881), WUNDERLICH (1884), BEDDARD (1898) and PYCRAFT (1900) state that the syrinx of the Greater Rhea has intrinsic muscles and a well-developed tympanum unlike that of other Ratites. Also, WUNDERLICH (1884) was the first to mention the cartilaginous nature of the pessulus. However, due to methodological limitations at the time, these studies did not provide detailed anatomical descriptions, nor age or sex of the individuals studied. Also, the illustrations only show two macroscopical syrinx drawings in dorsal and ventral aspects.

Studies on the comparative morphology of the syrinx contribute to understand the mechanisms of sound production (e.g. GOLLER & LARSEN, 1997; LARSEN & GOLLER, 1999, 2002) and have been used to establish phylogenetic relationships in systematic studies (e.g. AMES, 1971; LIVEZEY, 1986; PRUM, 1992; GRIFFITHS, 1994; GABAN-LIMA & HÖFLING, 2006; ZIMMER *et al.*, 2008; MANDIWANA *et al.*, 2011). The aim of this study was to describe the anatomy of the syrinx in *Rhea americana* throughout its ontogeny, by using muscle staining (BOCK & SHEAR, 1972) and differential colouring of cartilage and bone (CANNEL, 1988). This approach to the study of the anatomy of the syrinx of a Ratite bird would be useful to carry out further studies on its comparative and functional anatomy.

Materials and Methods

Anatomical dissections were performed on 12 specimens of *Rhea americana* at several ages: four adults (1 female and 3 males of two years old), four 3-month-old unsexed chicks, and four 1-month-old unsexed chicks. The birds were obtained from various commercial farms located in Buenos Aires province (Argentina) and reared in accordance with Argentinean regulations for Greater Rhea farming. Birds were sacrificed by cervical dislocation or electrical stunning, and their syringes were first observed in situ and then they were carefully removed. They were fixed in 10 % formaldehyde for 48 hours and then preserved in 70 % ethanol. An iodine solution that selectively stains muscular tissue with a reddish brown color (BOCK & SHEAR, 1972) was used to observe the muscles in preserved syringes. Lastly, the syringes were stained using a standard differential coloring technique for cartilage and bone (CANNELS, 1988), where cartilage tissues are stained blue (alcian blue) and ossified tissues are stained red (alizarin red). The anatomical nomenclature followed throughout corresponds to that proposed by BAUMEL *et al.* (1993). Photographs were taken with a Nikon D-40 digital camera.

Results

Skeletal elements of the adult syrinx

The syrinx of Greater Rhea was of the tracheobronchial type and it was composed of cartilages, namely (cart.) tracheosyringaeals (T) and cart. bronchosyringaeals (B). The former were broad and conformed a well-developed tympanum (figs. 1: T1–T4 & 2a, b) together with the first cart. bronchial (figs. 1: B1 & 2a, b). In dorsal view, cartilages T2, T3 and T4 got to fuse in the middle region of tympanum together with the pessulus and the cart. B1 (Fig. 1 & 2a), whereas cart. B1 was free in dorsal aspect (fig. 2a). Unlike other cart. tracheosyringaeals, T4 was noticeably convex (fig. 1 & 2a). The cart. bronchosyringaeals (figs. 1: B2–B6 & 2a, b) were “C-shaped” and the cart. B1 showed a slightly concave shape, which added to the convexity of the T4, delimited a wide space occupied by the membrana tympaniformis lateralis (fig. 1; see below). The remaining cart. bronchosyringaeals had similar form and held the ligamentum bronchiale mediale (fig. 1).

The pessulus was a thin bar, fused with the tympanum (figs. 2a, b & 3). In females, the pessulus was entirely cartilaginous (figs. 2a, b). On the other hand, in males, it was cartilaginous on its dorsal half, whereas it was osseous

on its ventral half, forming an osseous plate (figs. 2c, d). The membrana tympaniformis lateralis (fig. 1) was located between the last cart. tracheosyringeal (T4) and the first cart. bronchosyringeal (B1). It was partially covered by the pair of intrinsic muscles (fig. 1; see below). The membrana tympaniformis medialis (figs. 1 & 3a) was suspended between the free ends of the cart. bronchosyringales B1 and B2, and extended, making contact with the pessulus (fig. 3a). This membrane formed a pair of intrusions into the lumen of the syrinx (fig. 3b). Finally, the ligamentum interbronchiale connected the left and right bronchii (fig. 1) and between this ligament and the pessulus the foramen interbronchiale could be observed (fig. 1).

The syrinx of chicks

In chicks, the syringes were cartilaginous and presented similar anatomy regardless of age (figs. 2e, f). Unlike adults, the cart. bronchosyringeal B1 was not fused with the cart. tracheosyringeal T4 in its ventral aspect (figs. 2e, f). Other differences were noted in the first cart. tracheosyringales: the fusion (or joint zones) between cart. T1–T3 was variable among the chicks considered in this study, as well as the presence of a partial bifurcation (divergence zones) of these cartilages (figs. 2e, f).

Syrinx musculature

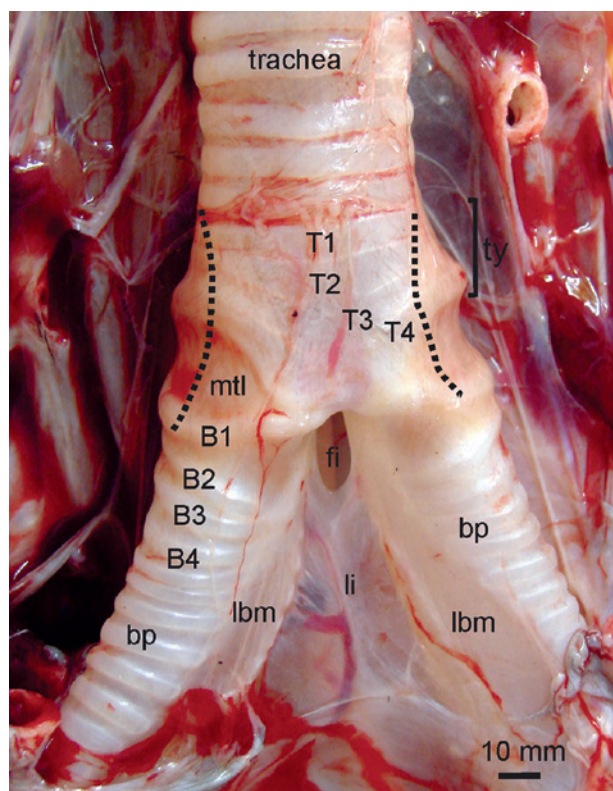
The musculature of the syrinx consisted of extrinsic and intrinsic muscles. The extrinsic musculature was represented by paired muscles (mm.), namely sternotrachealis and tracheolateralis (figs. 4a–c), and by a broad muscular band located on the mid-dorsal region of the trachea (figs. 4b, e). The muscle (m.) sternotrachealis was thin and delicate; it originated in the internal surface of the sternum and inserted on each side of the trachea, cranially to the tympanum (fig. 4a). The m. tracheolateralis (figs. 4b,c), was located on the lateral surface of the trachea; it was thin and weak, originating on the lateral surface of the larynx and inserting near to the cranial attachment of the intrinsic muscles (fig. 4c). The broad muscular band (figs. 4b, e) was thin and firmly attached to the dorsal surface of the trachea from near the larynx to the tympanum (at the level of the cart. T1 & T2). A single pair of intrinsic muscles were observed (figs. 4a, d). They consisted on wide and thin muscular bands with an oblique path extending from the dorsal region of the cart. tracheales (immediately preceding the tympanum) to the ventral surface of the first three cart. bronchosyringales. In unpreserved specimens, all muscles were thin and delicate and presented a pink pale color (figs. 4a & 1). No differences were found in muscle arrangement through ages.

Discussion

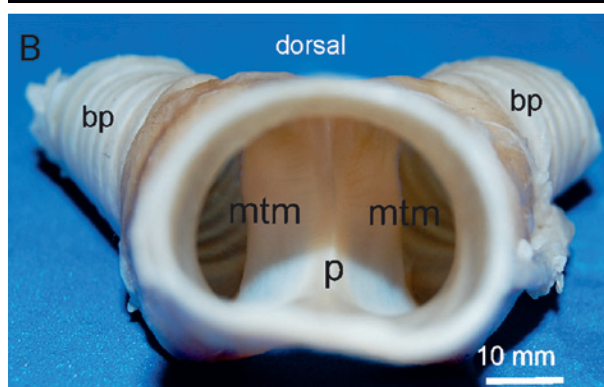
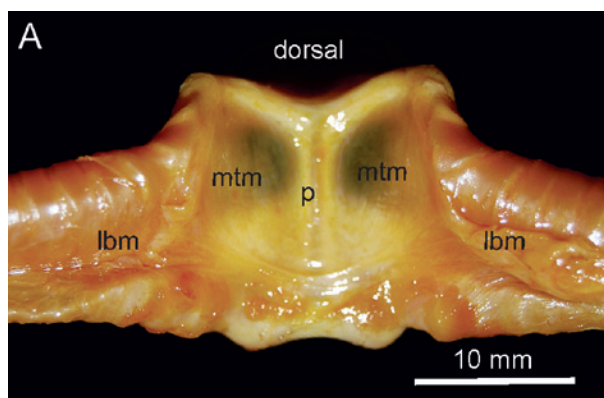
The syrinx of the Greater Rhea corresponds to the tracheobronchial type, as in the remaining Palaeognathae birds (FORBES, 1881; YILDIZ *et al.*, 2003; GARITANO-ZAVALA, 2009) and most Neognathae birds (BEDDARD, 1898; KING, 1989; BAUMEL *et al.*, 1993). But, the Greater Rhea syrinx was notable for having a well-developed *tympanum*, and for the presence of a pair of intrinsic muscles. In the rest of the Paleognathae birds, there are no intrinsic muscles and the tympanum has often been described as simple due to the presence of a lower degree of fusion between the components and the presence of a pessulus of connective tissue (FORBES, 1881; WUNDERLICH, 1884; YILDIZ *et al.*, 2003; GARITANO-ZAVALA, 2009). FORBES (1881) described the presence of a “vocal cord” inside the syrinx of *Rhea americana* (p. 240). It is possible that this author called “vocal cord” the two intrusions of the membranae tympaniformes mediales into the lumen of the syrinx (see fig. 3a). These intrusions are noticeable when the syrinx is fixed and preserved, whereas in, unpreserved specimens this is not as evident. In regard to the extrinsic muscles, the results of this study showed some differences with respect to information given by previous authors (i.e.: FORBES, 1881, WUNDERLICH, 1884; BEDDARD, 1898; PYCRAFT, 1900). Initially, these authors described the presence of a single pair of extrinsic muscles which were identified as the “lateral tracheal muscle” (FORBES, 1881 p. 240), without giving further details on their origin and extension. In our work, we identified the two typical pairs of extrinsic muscles of the syrinx (mm. sternotracheales and tracheolaterales). When comparing it with other Palaeognathae birds, FORBES (1881) found that the ostrich syrinx had no intrinsic or extrinsic muscles, but WUNDERLICH (1884), PYCRAFT (1900) and YILDIZ *et al.* (2003) described the presence of the m. sternotrachealis. Regarding the remain genera (*Casuaris*, *Dromaius* and *Apteryx*), these authors described the presence of the two typical pairs of extrinsic muscles. These disparities indicate that variations in the musculature (e.g. presence or absence of a muscle) is a common trait in birds (BERGER, 1956; BERMAN *et al.*, 1990).

Lastly, concerning the medium-dorsal muscular band present in the Greater Rhea, FORBES (1881) called it “fibrous band” and mentioned its presence also in the Cassowary. After a review of the available bibliography, we could not find a similar muscle in other birds.

The presence of intrinsic muscles in the syrinx of the Greater Rhea is interesting. They are a distinctive feature of Passerine birds (especially the Oscines songbirds), that have at least four pairs that contribute to the control of a wide variety of vocalizations (SUTHERS, 2001; LARSEN & GOLLER, 2002). Some groups of non-Passeriformes birds also have intrinsic muscles, although in lesser numbers, namely hummingbirds (Trochiliformes, MÜLLER, 1847; GAUNT, 1983), parrots (Psittaciformes, GAUNT & GAUNT, 1985; GABAN-LIMA & HÖFLING, 2006) and the oilbird



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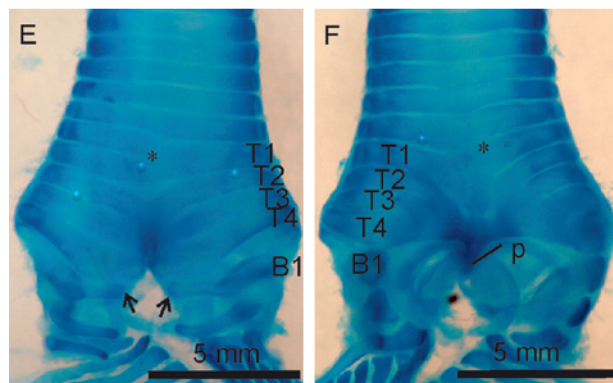
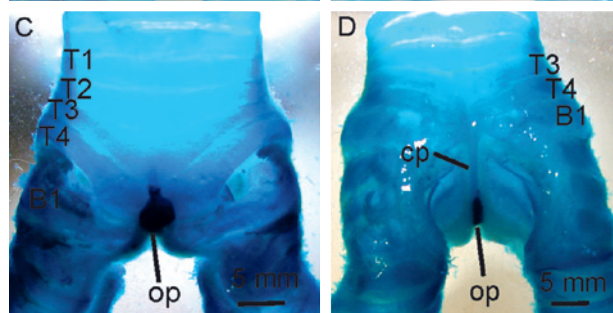
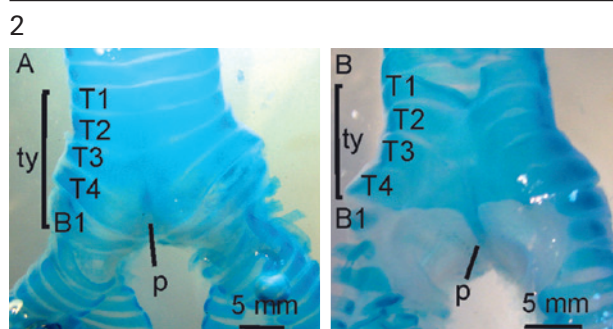


Fig. 1. *In situ* ventral view of the syrinx of Greater Rhea, B1–B4: cartilages bronchosyringaealis, T1–T4: cartilages tracheosyringaeales, bp: bronchus primarius, fi: *foramen* interbronchiale, lbm: ligamentum bronchiale mediale, li: ligamentum interbronchiale, mtl: membrana tympaniformis lateralis; ty: tympanum. Dotted line indicate the pair of intrinsic muscles.

Fig. 2. Stained and clarified syrinx, (a,b): adult female in ventral and dorsal view respectively; (c,d): adult males in ventral and dorsal view respectively, note the differences between the female pessulus (p) (entirely cartilaginous) and the male pessulus (partially osseous and cartilaginous) (op/cp); (e,f): chick of one month old in ventral and dorsal view respectively, B1: cart. brochosyringaeal, T1–T4: cart. tracheosyringaeales, p: pessulus, ty: tympanum, arrows indicates the *cart.* B1 not fused with the *tympanum*, asterik indicate tracheal rings with partial bifurcations.

Fig. 3. Location and extension of the membrana tympaniformis medialis (mtm) and pessulus (p) in fixed syringes, (a) Caudal view of syrinx showing the pessulus and the membrana tympaniformis medialis (specimen of three months old); (b) Internal view of the syrinx showing the intrusions of the membrana tympaniformis medialis (adult specimen), lbm: ligamentum bronchiale mediale, bp: bronchus primarius

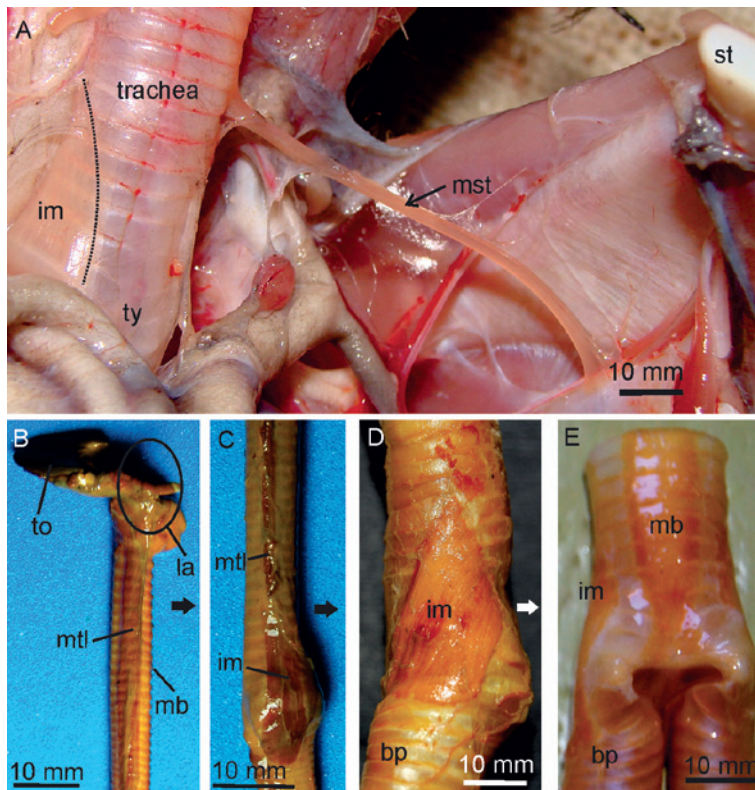


Fig. 4. Musculature of the syrinx. (a) In situ ventral view of the syrinx showing the m. sternotrachealis (mst) and the right intrinsic muscle (im) (specimen of three months old); (b) and (c) Proximal and caudal left view respectively of the syrinx showing the origin and insertion of m. tracheolateralis (mtl) (specimen of one month old); (d) Left lateral view showing the left intrinsic muscle (im) (specimen of three month old); (e) Dorsal view showing the medial muscular band (mb) (specimen of three month old), (b–e): specimens stained with iodine solution, bp: bronchus primarius, la: larynx, st: sternum, to: tongue, tr: trachea, ty: tympanum. Arrow in figures b and c indicates dorsal surface.

Steatornis caripensis (Caprimulgiformes, SUTHERS & HECTOR, 1985). These birds do not have a large repertoire of vocalizations, but they can imitate sounds (e.g. parrots) and are capable of vocal learning (e.g. hummingbirds and parrots) (GAUNT, 1983; GAUNT & GAUNT, 1985; SUTHERS, 2001). Moreover, in the echolocating oilbird, the intrinsic muscles have evolved for the production of sonar clicks (SUTHERS & HECTOR, 1985). The role that intrinsic muscles perform in the functioning of the syrinx of the Greater Rhea is still unknown, however, is interesting to note that adults of *R. americana* produce sounds like “hisses” and during breeding season, male adults make a deep-toned “grunt” (termed booming call), that can be heard at great distances (RAIKOW, 1969; BRUNING, 1974; BEAVER, 1978; FOLCH, 1992; CODENOTTI & ALVAREZ, 2001; DAVIES, 2002). In contrast, BEAVER (1978) found that young birds have a wide repertoire of sounds (consisting of about five types of vocalizations), that impoverishes and disappears as the chicks grow. This author related this vocal modification with the more developed membranae tympaniformes mediales in chicks than in adults. Nonetheless, he did not perform measurements on this membrane to corroborate this statement and the published figures are unclear (see BEAVER, 1978 fig. 3, p. 387), without scale and taken in different views. In our study no macroscopic differences were found in the membranae tympaniformes mediales when comparing chicks to adults. We also believe that the degree of intrusion of the membrane could be a fixation artifact. Therefore, we conclude that the impoverishment of richness of sounds found by BEAVER (1978) should be studied from other perspectives.

The syrinx of the Greater Rhea was previously described as being completely cartilaginous (FORBES, 1881; WUNDERLICH, 1884; PYCRAFT, 1900), but the differential staining techniques incorporated in this study showed the presence of a pessulus in the male adult formed by osseous and cartilaginous tissues. A similar condition (yet of unknown significance) was found in the male of the Tufted Duck, *Aythya fuligula* (WARNER, 1971). Sexual dimorphism in syrinx anatomy has been found in other birds (MILLER *et al.*, 2008), varying from the presence of larger syringes in males (e.g. in the collared dove, *Streptopelia decaocto*, BALLINTIJN & CATE, 1997), to the presence of specialized structures such as the syringeal bulla in males of Anatidae (e.g. FRANK *et al.*, 2007; WARNER, 1971), to differences in structures like labia and cartilaginous rings (e.g. in the Zebra Finch, *Taeniopygia guttata*; RIEDE *et al.*, 2010). Also, differences in vocalizations between male and female could be associated with sexual dimorphism in syrinx anatomy (BALLINTIJN & CATE, 1997; MILLER *et al.*, 2008; RIEDE *et al.*, 2010; WARNER, 1971), but in the Greater Rhea this topic still remains to be explored.

In conclusion, the present study points to the complexity in the morphology of the syrinx of *Rhea americana* and the more attention it deserves. Future studies on air flow and air sac pressure, intrinsic and extrinsic muscles electromyography and endoscopic filming of the syrinx during the generation of sounds are needed to complement these findings. Such information will allow to compare if the mechanisms of sound production of Palaeognathae differs from those known for Neognathae birds, and eventually shed new light on the evolution of this feature in birds.

Acknowledgements

We would like to thank to F. J. Degrange for his help during the dissections, to J.R. Casciotta and J.A. Vanegas-Rios for their valuable suggestions in staining methods.

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