SKELETAL ANATOMY, PHYLOGENETIC RELATIONSHIPS, AND PALEOECOLOGY OF THE EOCENE UROLOPHID STINGRAY ARECHIA CRASSICAUDATA (BLAINVILLE, 1818) FROM MONTE POSTALE (BOLCA LAGERSTÄTTE, ITALY)

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ABSTRACT—In this paper we re-examine the taxonomy and systematic position of the Eocene stingrays from Bolca Lagerstätte which are traditionally assigned to Urolophus crassicaudatus (Blainville). The analysis of their tooth morphology supports an assignment to the Eocene stingray genus Arechia Cappetta, a taxon known from isolated teeth from the Ypresian-Lutetian of northern and western Africa. The teeth of the Bolca specimens differ from the type species A. arambourgi in some characters (i.e., labial face with concave profile just below the crest, convex lower down; lingual face slightly more developed than the labial, with convex profile in its upper part and a concave profile in its mid-lower region) that justify the recognition of a second species within the genus, i.e., A. crassicaudata. This taxon also shows a unique combination of features (e.g., pectoral disc large and rhomboid; tail short, 44-57% of total length; ca. 238 vertebral centra; distal segment of propterygia located between mouth and antorbital cartilage; mesopterygium single, not fused to radials; 100-117 pectoral radials; 15-17 pelvic-fin radials; elongated caudal fin of apelosodic type) that supports its sister-group relationship with the living urolophids Urolophus and Trygonoptera. Arechia was a typical inhabitant of the near coastal and warm habitats that characterize the Monte Postale paleoenvironment.

SUPPLEMENTAL DATA—Supplemental materials are available for this article for free at www.tandfonline.com/UJVP

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INTRODUCTION

The Ypresian fossil Lagerstätte of Bolca, in northeastern Italy, is one of the few Paleogene deposits where fossils of cartilaginous fishes (Chondrichthyes) are represented by complete and articulated skeletal remains (Marramà et al., 2018a). In the last few years, several studies provided new insights into systematics, relationships, and paleobiology of sharks and rays from the Pescaiola and Monte Postale sites of Bolca (Fanti et al., 2016, 2019; Marramà et al., 2018a, 2018b, 2018c, 2019a, 2019b, 2019c, 2019d). The batoids, in particular, are rather abundant and represented by electric rays, guitarfishes, thornback rays, and stingrays, with the latter representing the most diverse lineage of chondrichthians of Bolca. One of the stingray taxa, described by Blainville (1818) as Trygonobatus crassicaudatus, was later assigned to the stingaree genus Urolophus Müller and Henle, 1837 by Jaekel (1894) in his comprehensive review of the elasmobranch fishes from Bolca. Since then, this taxonomic status has been widely accepted but a revisionary study of these fossils has not been provided up to now. According to Cappetta (2012), however, the size and morphology of the teeth of Urolophus crassicaudatus are quite different from those of the living Urolophus species and more similar to those of Arechia Cappetta, 1983, an extinct stingray genus described for isolated teeth from the Ypresian basin of Ouled Abdoun in Morocco (Cappetta, 1983), and occurring from the Ypresian to the Lutetian of northern and western Africa (Cappetta, 1987, 2012; Cappetta and Traverse, 1988; Noubhani and Cappetta, 1997; Tabuce et al., 2005; Sambou et al., 2017).

The taxonomic history of Urolophus crassicaudatus is very complex. After being poorly described and placed in the genus Trygonobatus by Blainville (1818), without figures or reference to any type specimen, the taxon was subsequently transferred without further description or illustrations to Trygon oblongus by Agassiz (1833-1844) and to Trygon brevicauda by Heckel (1851). A specimen housed in the Natural History Museum of Vienna (NHMW 1853.0027.0005) was later described as Urolophus princeps by Heckel (1853), whereas Molin (1861) named another specimen housed in the Museo di Geologia e Paleontologia dell’Università di Padova (MGP-PD 8875C/776c) as Tenaiera knerii. Subsequently, Jaekel (1894) concluded that all the taxa listed above should be regarded as junior synonyms of Trygonobatus crassicaudatus Blainville, 1818, and subsequently transferred the species to Urolophus. Unfortunately, Blainville (1818) never indicated
the type specimen and that indicated by Blot (1980) as the holotype of ‘Urolophus’ crassicaudatus in the collections of the Museum National d’Histoire Naturelle of Paris cannot be regarded as such. In fact, Eastman (1904, 1905) and more recently Marramà et al. (2019c) recognized that there is no stingray specimen in this institution that can be considered as the type reported by Blainville (1818). For this reason, the original holotype of Trygonobatus crassicaudatus must be considered lost and a neotype must be selected for this species.

In this paper, we provide a detailed description of the skeletal anatomy of the fossil material from Bolca originally assigned to Urolophus crassicaudatus and re-evaluate its systematic position in the context of our current understanding of the stingray interrelationships.

**GEOLOGICAL SETTING**

The historical literature suggests that the specimens examined in this study were collected from the fossiliferous layers of the Monte Postale, which is located about 2 km north-east of the village of Bolca (Verona Province, northeastern Italy), in the eastern part of the Lessini Mountains, southern Alps, about 300 m from the better known Pesciara site. The Monte Postale sedimentary succession includes the Cretaceous Scaglia Rossa Formation up to Ypresian fossiliferous limestone; in the uppermost part, the latter contains abundant benthic foraminifera of the genus Alveolina, and marine and brackish mollusks, of almost the same age and similar sedimentological features as the Pesciara site, mostly comprising finely laminated micritic limestones with fish and plant remains. Papazzoni et al. (2017) investigated the stratigraphic relationships between the two fossiliferous deposits, suggesting that the uppermost productive sequence of Monte Postale should correlate with that of the Pesciara site, although the fossiliferous laminates of Pesciara appear to be slightly younger than those of Monte Postale. Based on large benthic foraminifera and calcareous nanoplankton, the uppermost strata of the Monte Postale site were assigned to Shallow Benthic Zone 11 by Papazzoni et al. (2017) and correspond to the late Cuisian (late Ypresian), around 50 Ma. Evidence of a coralgal rim, slightly younger than those of Monte Postale. Based on large benthic foraminifera and calcareous nanoplankton, the uppermost strata of the Monte Postale site were assigned to Shallow Benthic Zone 11 by Papazzoni et al. (2017) and correspond to the late Cuisian (late Ypresian), around 50 Ma. Evidence of a coralgal rim, lagoonal deposits, and a fore-reefsystem was hypothesized for the Monte Postale paleobiotope (Vescogni et al., 2016), and this interpret-
Trygon oblongus Agassiz; Agassiz, 1835a:297.
Trygon oblongus Agassiz; Agassiz, 1835b:14.
Trygon oblongus Agassiz; De Zigno, 1874:181.
Trygon brevicauda Heckel, 1851:324 (nomen nudum; no description or illustration).
Trygon brevicauda Heckel, 1851:324 (nomen nudum; no description or illustration).

Urolophus princeps Heckel, 1853:124 (nomen dubium; insufficient description; no illustration).
Urolophus princeps Heckel; Kner and Steindacher, 1863:32, pl. 6, fig. 2.
Urolophus princeps Heckel; De Zigno, 1874:182.
Taeniura knerii Molin, 1861:581 (nomen dubium; insufficient description; no illustration).
Urolophus princeps Heck.; Kner and Steindacher, 1863:32, pl. 6, fig. 2.

‘Urolophus’ sp.; Carvalho, Maisey, and Grande, 2004:11.
‘Urolophus’ crassicaudatus; Carpetta, 2012:425, fig. 416s.
‘Urolophus’ crassicaudatus (Blainville, 1818); Carnevale, Bannikov, Marrama, Tyler, and Zorzin, 2014:41.

Urolophus crassicaudatus de Blainville sp.; Jaekel, 1894:148, pl. 5.
Urolophus crassicaudatus (Blainville); Eastman, 1904:24.
Urolophus crassicaudatus (de Blainville); Eastman, 1905:9.
Urolophus crassicaudatus (de Blainville); D’Erasmo, 1922:22.
Urolophus crassicaudatus (de Blainville); Blot, 1980:345.
Urolophus crassicaudatus Eastman; Frickhinger, 1991:216.
‘Urolophus’ crassicaudatus; Carvalho, Maisey, and Grande, 2004:11.

Neotype — MCSNV IG.VR.27607, nearly complete, articulated skeleton, 1103.4 mm TL, 685.4 mm DW (Fig. 1).

Referred Material—MCSNV VII.B.82/83, a nearly complete articulated skeleton in part and counterpart, 1028.0 mm TL, 637.4 mm DW (Fig. 2A, B); MGP-PD 8875C/76C, a partially complete articulated skeleton lacking the tip of the tail, in part and counterpart, 567.1 mm DW; named by Molin (1861) as Taeniura knerii (Fig. 2C, D); MCSNV T.317/318, an incomplete, partially articulated skeleton, in part and counterpart, 955.5 mm TL, 530.0 mm DW, erroneously indicated by Cappetta (2012) as the holotype of ‘Urolophus’ crassicaudatus (Fig. 3A, B); MCSNV VII.B.84/85, a nearly complete articulated skeleton, in part and counterpart, 949.3 mm TL, 604.1 mm DW (Fig. 3C); NHMW 1853.0027.0005, an incomplete skeleton, with pieces of slab erroneously assembled together; described by Heckel (1853) as Urolophus princeps (Fig. 3D).

Type Locality and Horizon — Monte Postale site, Bolca Konservat-Lagerstätte, Italy; early Eocene, late Ypresian, middle Cuisian, SBZ 11, Alveolina dainelli Zone; 50.7–48.9 Ma (see Papazzoni et al., 2017).
Amended Diagnosis—A species of *Arechia* that differs from *A. arambourgi* in the following dental characters: crown with two transverse cutting crests separated by a depression; in profile view, the labial crown face is concave just below the main crest, but convex further basally; lingual crown face slightly more developed than the labial one, with convex profile in its upper part and concave profile in its mid-lower region; edge of the lingual visor slightly more convex than the labial one. Furthermore, *Arechia* can be diagnosed by the following skeletal features: pectoral disc rhomboidal, slightly longer than wide (about 1.1 times); tail short, 44–52% TL; basihyal not fragmented and separated from the first hypobranchials; approximately 238 vertebral centra; distal segment of the propterygium located between mouth and antorbital cartilage; mesopterygium single, with external margins straight and not fused to radials; 100–117 pectoral radials; 15–17 pelvic-fin radials; aplesodic caudal fin of 20–22% TL.

Description

The description of the cranial and postcranial morphology of *Arechia crassicaudata* (Blainville, 1818) is based on six articulated skeletons characterized by different degrees of completeness (Figs. 1–3). Although the overall outline and body morphology of the specimens is still recognizable, most of the cranial and girdle elements are disarticulated, fragmented, and displaced from their original position, due to the inadequate preservation that commonly characterizes the specimens from the Monte Postale site (Marràmà et al., 2016). Counts and measurements are listed in Supplemental Data 1 (Table S1). The specimens are of similar size, with the largest one being characterized by 69 cm disc width and 110 cm in total length. The disc is rhomboidal, not wing-like, with the maximum width in the anterior third of disc length. The disc length is slightly longer than the disc width (about 1.1 times), with the disc being 56–64% of the total length. The tail is short, thicker in its proximal part, measuring ca. 44–52% of total length.

**Neurocranium**—The neurocranium is most clearly preserved in MGP-PD 8875C/76C. It is anteroposteriorly elongated, with the greatest width at the level of the nasal capsules (Fig. 4). The rostral cartilage is absent as is the case in all adult stingrays (e.g., Miyake et al., 1992). The nasal capsules appear broad, ovoid in shape and laterally expanded with a rounded and
biconvex anterior margin. Small preorbital processes protrude from the posterolateral aspect of the nasal capsules (Fig. 4). The supraorbital process is difficult to detect. The postorbital processes are long, shelf-like, and anterolaterally directed. The otic capsules provide articulation surfaces for the proximal portion of the hyomandibulae. Although the specimen MGP-PD 8875C/76C mostly exposes its ventral side, it is possible to observe the outline of the dorsal fontanelle on the neurocranium (Fig. 4). The dorsal fontanelle is triangular in outline, anteroposteriorly elongated, and covers about 50–60% of the neurocranial length. However, the fontanelle does not exhibit any median constriction, which usually represents a remnant of the epiphyseal bar that separates an anterior precerebral from a posterior frontoparietal fontanelle in some stingrays (Miyake, 1988; Carvalho et al., 2004). Antorbital cartilages are not preserved in any specimen.

**Jaws**—The upper and lower jaws are massive and robust, extend laterally, and occupy almost the entire space between the propterygia (Figs. 4, 5). The antimeres are separated at the symphysis, and the occlusal width is greater than the diastema width. The palatoquadrate appears to be labiolingually compressed, slightly narrower than Meckel's cartilage, and relatively straight on its dorsal flange. The anterior processes of the Meckel's cartilage are difficult to detect. There is no evidence of the lateral projections of the lower jaws ('wing-like processes' of Carvalho et al., 2004) which are typical for myliobatids.

**Hyoid and Gill Arches**—The hyomandibulae appear slender and straight, not arched, and narrow at about their midlength (Figs. 4, 5). The hyomandibulae project anterolaterally, reaching the mesial wall of the propterygia just posterior to the posteroventral corner of the lower jaw. The distal end of the hyomandibulae articulates with the lower jaw through a stout and strong terminal portion in MCSNV IG.VR.27607, whereas their proximal portion at the articulation with the neurocranium is enlarged. The presence of the secondary hyomandibular cartilages that are characteristic of *Urolophus* are difficult to detect, due to poor preservation. The ventral gill arches of *A. crassicaudata* are poorly preserved. However, the outline of the central medial plate (derived from the fusion of the basibranchial copula and the basibranchial components) can be seen at least in MGP-PD 8875C-76C and MCSNV IG.VR.27607 (Figs. 4, 5). The medial
plate appears anteroposteriorly elongated. Its posterior distal tip seems to taper into small median projections. The basihyal appears unfragmented and is clearly separate from the first hypo-branchials, which are stout and robust in MCSNV IG.VR.27607 (Fig. 5). There are five pairs of ceratobranchials articulating with the lateral margins of the medial plate. The first appears fused to the pseudohyoid whereas the condition of the last two ceratobranchials is difficult to determine. The fifth ceratobran-chial pair articulates distally with the anterior margin of the scapulocoracoid. Filamentous branchial rays are associated with the ceratobranchials although the number of elements associated with each ceratobranchial is difficult to ascertain.

Synarcuals and Vertebral Column—The anterior (cervicothoracic) synarcual is difficult to discern whereas the outline of the posterior (thoracolumbar) synarcual cartilage can be detected (Figs. 4, 5). The thoracolumbar synarcual is triangular in shape, articulates anteriorly with the anterior synarcual, and tapers posteriorly with its posterior margin ending approximately mid-way between the scapulocoracoid and pelvic girdle. The vertebral column of *Arechia crassicaudata* consists of ca. 238 vertebral centra,
counted on the best preserved specimen MCSNV IG.VR.27607. However, it is difficult to diagnose the exact number of vertebræ because they are often damaged or lost. There are about 20 trunk (monospondyous) centra, counted from the first distinguishable centrum to the anterior margin of the puboischiadic bar. About 218 are diplospondylous (of which 148 are from the anterior margin of the puboischiadic bar to the anterior disc margin). The vertebral centra are small, subrectangular in shape and anteroposteriorly short. Some neural spines are visible in the examined material. Ribs are absent.

FIGURE 6. A single tooth extracted from the neotype of Arechia crassicaudata (Blainville, 1818), MCSNV IG.VR.27607, in A, linguo-occlusal, B, lingual, C, labio-basal, D, basal, E, labial, and F, profile views. Scale bar equals 1 mm.
1983. The dentition exhibits a gradient monognathic heterodonty, with lateral teeth becoming smaller and enlarged labiolingually. The single tooth extracted from MCSNV IG.VR.27607 is about 2 mm in crown width, broader than long with a slightly convex labial contour and a very strongly convex lingual outline (Fig. 6). The crown is high and not cuspidate. The enameloid surface is completely smooth. The edge of the labial visor is rather broad and convex. The edge of the lingual visor is convex, with a central notch. There is a high, sharp, and lingually displaced transverse cutting crest that does not reach the lateral angles, which are blunt. A second transverse cutting crest occurs labially but it is lower and blunter than the lingual one, resembling the condition observed in *Urolophus, Trygonoptera, Uraboris*, and *Hiinantura* (Herman et al., 2000). Although not specifically mentioned, it seems that this character can be recognized also in the type species *A. arambourgi* (see Cappetta, 1983, 2012). At the labial surface, a weak depression can be observed in between the two transverse cutting crests. In profile view, the labial face has a concave profile just below the crest that becomes convex further basally. The lingual face is slightly more developed than the labial one; it has a convex profile in its upper part and a concave profile in its mid-lower region. The root is lower than the crown and slightly expanded mesiodistally. There are two root lobes whose basal face is subtriangular, slightly convex, and feebly joins the labial face of the oblique and straight root. The root lobes are divided by a broad and deep furrow with a large central foramen and one or two paracentral foramina. Other foramina open also on the labial face of the root, just below its junction with the crown.

**Squamation and Stings**—The skin of modern urolophid genera *Urolophus* and *Trygonoptera* is completely devoid of dermal denticles and thorns, whereas *Spindolophus* possesses some sparse denticles and thorns (Last et al., 2016). All the examined specimens of *Arechia* mostly lack dermal denticles and thorns. However, some of them show very small and scattered structures having star shape on the disc margin and tail that might be interpreted as dermal denticles or remains of the original pigmentation (Supplemental Data 1, Fig. S2A, B). However it is hard to draw firm conclusions given the poor preservation of the specimens. A single, elongate, and dorsoventrally flattened serrated caudal sting occurs in the majority of the specimens (Supplemental Data 1, Fig. S2C–F). The sting origin is located at about mid way down the length of the tail, posterior to the pelvic fins and just anterior to the caudal-fin origin, at ca. 70–77% of the total body length. The best-preserved stings are about 9% of the body length. Four to six irregular grooves run parallel to the main axis on the dorsal side of the sting. The serrations on the sting are small, hook shaped, and directed transversely to the caudal axis of the sting. The number of serrations per side is difficult to determine due to the lack of the distal tip of the stings. No specific characters of the sting, useful to discriminate *Arechia* from other urolophids, can be recognized.

**Stomach Contents**—Two specimens of *A. crassicaudata* show abdominal gut content consisting of partially digested bony fishes which are completely preserved in the abdominal cavity between the pectoral and pelvic girdles, on one side of the vertebral column, in a position comparable to that occupied by the gut-intestine tract in extant stingrays (Supplemental Data 1, Fig. S3). The stomach content of MCSNV VII.B.84/85 consists of a single vertebral column with associated arches, and disarticulated dorsal and anal fins. Conversely, the bones of the head and tail of the bony fish appear scattered around the vertebral column, showing little evidence of digestion thereby suggesting that the consumption occurred shortly before the death of the stingray. Despite the degree of disarticulation of the skeleton, the overall morphology of the body axis of the fish as well as the presence of considerably large dorsal-fin spines showing canals running parallel to the main spine axis are consistent with those of the extinct squirrelfish *Eoholocentrum macrocephalum* (Blainville, 1818), family Holocentridae (see Sorbini and Tirapelle, 1974).

The neotype of *A. crassicaudata*, MCSNV IG.VR.27607, shows two or three small fishes inside the abdomen. Due to the considerable degree of disarticulation, it was not possible to establish the precise taxonomic identity of the bony fishes, although the general morphology of the vertebral column appears to be consistent with those of small percomorphs.

A third specimen, MGP-PD 8875C/76C, shows a possible cololite, or intestinal fossil content still in situ in the abdominal cavity (Supplemental Data 1, Fig. S3C). The cololite is relatively short and uncoiled with a whitish coloration. It is not possible to distinguish what kind of food the cololite is composed of, with the exception of a small chela of a decapod crustacean.

The presence of crustacean and fish remains as stomach/gut content of three out of six individuals of *Arechia*, suggests that the food preferences of this taxon were quite similar to those of extant urolophids, which feed mainly on polychaete worms, crustaceans, and small bony fishes (Last and Compagno, 1999; Last et al., 2016a).

**Phylogenetic Analysis**

The analysis conducted in TNT resulted in a single parsimony tree with a length of 215 steps, a consistency index (CI) of 0.64, and a retention index (RI) of 0.80 (Fig. 7), and a topology that is consistent with those depicted in previous studies on Bolca stingrays (e.g., Marramà et al., 2019c). The Myliobatiiformes are recovered as a monophyletic group (Bremer value 4) supported herein by ten characters: basihyal as a single element, but separated from the first hypobranchials (ch. 19[1]); presence of a median projection of the basibranchial medial plate (ch. 22[1]); presence of levator and depressor rostri muscles (ch. 66[1]), serrated tail stings (ch. 67[1]); thorns absent (ch. 69[1]); rostral cartilage vestigial or absent (ch. 73[1]); postbital process very broad and shelf-like (ch. 74[1]); jugal arch absent (ch. 75[1]); presence of ball and socket articulation between scapular process and synarcual (ch. 78[1]); presence of a thoracolumbar synarcual (ch. 79[1]). As in the molecular analysis of Ashciman et al. (2012b), the sixgill stingray *Hexatrygon* is inferred to be the sister taxon to all other stingrays, in contrast with molecular phylogenies interpreting this taxon as sister to *Gymnura* (Bertozzi et al., 2016) or to the urolophids (Naylor et al., 2012a, 2012b).

The remaining stingrays are dichotomously grouped into two large clades (*Myliobatoidea* and *Dasyatoidea*) whose nature can be possibly linked to the different body shape, calcifications of radials, swimming mode, and life style (Schaefer and Summers, 2005; Hall et al., 2018).

The monophyly of the clade comprising all benthic stingrays (i.e., *Dasyatoidea*) is weakly supported (Bremer value 1) by a single character, the spiracularis split into lateral and medial bundles, with the medial bundle inserting on the posterior surface of Meckel’s cartilage and the lateral bundle inserting onto the dorsal edge of the hyomandibula (ch. 88[1]). The dasytooids include taxa having a rhomboid or ovoid pectoral disc, and a ‘catenated’ calcification pattern of radials, which possibly reflect their undulatory swimming mode and benthic habits (Schaefer and Summers, 2005).

*Arechia* is herein inferred to be a basal member of the dasytooid stingrays, sister to the living urolophids *Urolophus* and *Trygonoptera*. This relationship is supported by a single ambiguous character: the presence of a second transverse keel on teeth (ch. 98[1]). *Urolophus* and *Trygonoptera* are united by a single character: an external margin of the mesopterygia which is fused to radial elements (ch. 29[2]). This character appears absent in *Arechia* in which the external margins of the
mesopterygia are straight and not fused to the radials. The presence of the enlarged optic nerve foramen (ch. 8[1]) supports the sister-group relationship between *Urolophus* and *Trygonoptera* in previous phylogenetic analyses (e.g., Carvalho et al., 2004; Marramà et al., 2019a, 2019b, 2019c). However, this character does not support the group in this analysis, possibly because its presence is difficult to determine in *Arechia*. The monophyly of the Urolophidae was also recovered by previous morphological (e.g., Carvalho et al., 2004; Claeson et al., 2010) and molecular analyses (e.g., Naylor et al., 2012b; Bertozzi et al., 2016), whereas the family appears to be paraphyletic according to Naylor et al. (2012a) with *Trygonoptera* being sister to *Hexatrygon*, and *Urolophus* sister to *Plesiobatis*. The systematic position of *Arechia* also is supported when the data matrix is analyzed with PAUP* 4.0, which yielded 18 equally parsimonious trees with a tree length of 216 steps, a CI of 0.63, and a RI of 0.80.

Among the other dasyatoid stingrays, *Plesiobatis*, the Eocene stingrays from the Green River Formation, the urotrygonids and *Lessiniaabatis* form successive sister taxa to all potamotrygonids and dasyatids. *Asterotrygon* and *Heliobatis* form a monophyletic clade that is weakly supported (Bremer value 1) by a single ambiguous character (caudal fin reduced to tail folds; ch. 34[1]) that seems to have been achieved independently from the dasyatids (Marramà et al., 2019b). Urotrygonid stingrays (*Urotrygon* and *Urobatis*) are monophyletic as detected in Aschliman et al. (2012a) and Bertozzi et al. (2016). The enigmatic extinct stingray *Lessiniaabatis* from Bolca is sister to a polytomy that includes potamotrygonids and dasyatids. Although *Styracura* is certainly closely related to the freshwater potamotrygonids as suggested by morphological, molecular, and chrono/geographic evidence (Lovejoy, 1996; Lovejoy et al., 1998; Carvalho et al., 2004, 2016; Aschliman et al., 2012b; Naylor et al., 2012a, 2012b; Bertozzi et al., 2016), our phylogeny did not recognize *Styracura* as a genuine member of the Potamotrygonidae, possibly because *Styracura* lacks some characters of the lateral line, and pectoral and pelvic girdle skeleton, which are typically found in freshwater potamotrygonids (Carvalho et al., 2016).

Finally, our analysis weakly supports (Bremer value 1) a clade that includes *Gymnura* as sister to all pelagic stingrays with six synapomorphies: short orbital region with more anteriorly placed supraorbital and postorbital processes (ch. 10[1]); lateral expansion of the radials in pectoral region (ch. 28[1]); caudal fin absent (ch. 34[2]); first segment of propterygium adjacent to anterior margin of antorbital cartilage or anterior to margin of nasal capsule (ch. 81[3]); 'crustal' calcification pattern of radials (ch. 99[0]); and wing-like pectoral disc, with fins greatly expanded.
portions and meristic counts (Supplemental Data 1, Table S2). In fact, more robust molecular analyses

DISCUSSION

Comparative Remarks

The Urolophidae, also known as stingarees or round stingrays, are a myliobatiform family represented by 28 living species arranged in three genera (Urolophus, Trygonoptera, and Spinilophus; Last et al., 2016). They are small to medium-sized stingrays (up to 90 cm TL) characterized by an oval, circular or rhomboid pectoral disc and a short tail with an elongate lobe-like caudal fin. A dorsal fin can be present in some species. The skin is completely devoid of denticles and thorns in Urolophus and Trygonoptera, but usually one or more serrated stings are present on the tail, well behind the pelvic fins (Last and Compagno, 1999, Last et al., 2016). The monophyly of the family Urolophidae appears to be supported in morphology-based phylogenetic analyses by the presence of mesopterygia fused to radial elements, an enlarged foramen for the optic nerve in the neurocranium, and a second transverse tooth keel (e.g., Carvalho et al., 2004; Claeson et al., 2010; Marramà et al., 2019a, 2019b, 2019c).

In our study, the presence of the thoracolumbar synarcual, a serrated sting, and the absence of rostral cartilage support the inclusion of Arechia within the batoid order Myliobatiformes. The presence of a rhomboidal pectoral disc, a short tail with an elongate lobe-like caudal fin, skin mostly devoid of denticles and thorns, and second transverse tooth keel support the inclusion of Arechia within the Urolophidae and its sister-group relationship with Urolophus plus Trygonoptera in our phylogeny. A combination of several characters argues against the placement of Arechia within the other stingray lineages. For example, the presence of a complete caudal fin excludes its assignment to the Dasypodidae, Potamotrygonidae, ‘myliobatids’, or to the Eocene stingrays from the Green River Formation, because these taxa have replaced the caudal fin with tail folds or a cartilaginous stiffening rod in the terminal part of the tail (e.g., Carvalho et al., 2004). The absence of some traits characterizing pelagic/benthopelagic stingrays (e.g., wing-like pectoral disc, crustal calcification of radials, crushing dental plates, bashyal absent) rules out a close relationship with Arechia. Although the overall body plan of Arechia and living urolophids is similar to that of the Urotrygonidae, the presence of a single bashyal in Arechia excludes a close relationship with Urobatis and Urotrygon (bashyal absent or fragmented; Carvalho et al., 2004). Furthermore, the unique tooth crown morphology and the absence of any tooth enameloid ornamentation in the Bolca specimens excludes their alignment with other Eocene stingray taxa such as Anurobatis, Coupatexia, Heterorotpedo, Hypolophodon, Iacquinhermania, Merabatis, Meridiana, and Uoleia (see Cappetta, 2012).

Finally, Arechia can be distinguished from other members of the family Urolophidae (Urolophus, Trygonoptera, and Spinilophus) based on size (110 vs. 90 cm) and by several different body proportions and meristic counts (Supplemental Data 1, Table S2).

Fossil Record and Paleoecology of Urolophids

The paleoecological role of the urolophids from the Monte Postale site of the Bolca Lagerstätte has been poorly investigated so far. Living representatives of the Urolophidae are temperate to tropical marine batoids mostly occurring on continental and insular shelves of the Indo-Australian Archipelago and northwestern Pacific, although some species can occur in estuaries or slopes up to 420 m depth (Last and Compagno, 1999; Last et al., 2016). They are usually slow-swimming bottom-dwellers, occurring on soft mud, and feeding on polychaete worms, crustaceans, and small benthic fishes (Last and Compagno, 1999; Last et al., 2016). Quantitative paleoecological and taphonomic analyses of the fish assemblage of Monte Postale site suggest that the fossiliferous sediments were deposited close to an emerged coastal area, possibly characterized by the presence of mangroves and seagrass, in a coral reef context (Marramà et al., 2016; Vescogni et al., 2016). From this perspective, the presence of Arechia inhabiting the warm shallow-water habitats of the Monte Postale paleobiotope is therefore consistent with the ecological and environmental preferences of modern urolophids.

The fossil record of urolophids is poor when compared with the other stingray lineages, possibly because their isolated teeth are often mis-assigned to the wastebasket genus Dasysatis. Beside Arechia crassicaudata other occurrences of the genus are reported from the Ypresian to Lutetian of northern and western Africa (Cappetta, 1983, 1987, 2012; Cappetta and Travers, 1988; Noubhani and Cappetta, 1997; Tabuce et al., 2005; Sambou et al., 2017, 2020), and possibly Mississippi, U.S.A., the Netherlands, and France (Bor, 1985; Cappetta, 1982). Urolophus was reported by Hasse (1882) in the Lutetian of Belgium, and by Noetling (1885) in the Lutetian of Russia, on the basis of vertebral centra. Isolated teeth of Urolophus from the Oligocene of Germany were described by Fresss (1991). However, all the occurrences of Urolophus need to be verified according to Cappetta (2012). Indeterminate urolophid fossil teeth were also reported from the Lutetian of Morocco (Tabuce et al., 2005). Teeth of Urolophus halleri were reported from Pliocene of California (Fitch, 1964) although this taxon is today regarded as a species of Urobatis (family Urotrygonidae).

Molecular analyses suggested that the clade containing Urolophus and Trygonoptera possibly diverged from gymnurids around 75 Ma ago (Aschliman et al., 2012b) or from plesiobatids around the K-Pg boundary (Bertozzi et al., 2016). Since Arechia possibly represents the oldest urolophid (50 Ma), the divergence time estimates of the family appear to be quite consistent with its stratigraphic occurrence in the fossil record, or at least, a small gap exists (15–25 Ma). Moreover, it is plausible to assume a Tethyan origin for the family Urolophidae based on the oldest fossils occurring in the Tethyan realm, as also suggested for other batoid lineages.

CONCLUSIONS

The systematic revision of the fossil stingrays from Bolca traditionally identified as Urolophus crassicaudata (Blainville, 1818) confirmed previous assumptions that these articulated fossil
specimens cannot be attributed to the genus *Urolophus*. Considering that the general dental characters of the specimens from Bolca are consistent with those of *Arechia* these individuals represent the first skeletal record of this Eocene stingray genus and the only articulated fossil remains of the Urolophiidae known so far.

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