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The Development of Dermatonotus muelleri (Anura: Microhylidae: Gastrophryninae)

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ABSTRACT.—The monophyly of Microhylidae is supported by an overwhelming accumulation of synapomorphic larval features. Despite the distinctiveness of the microhylid tadpole, few studies have focused on larval development. Microhylid larval morphology is usually described and based on standard tables that imply that developmental events at equivalent stages of overall tadpole development are independent from species-specific patterns of developmental timing. Herein, we present additional developmental data based on external morphology and field data on larval growth for the gastrophrynine microhylid Dermatonotus muelleri. We describe internal morphological variation (e.g., skeletal and soft systems) during larval development. The results indicate that the onset of some metamorphic changes occurs earlier than those implied in current anuran developmental tables. This study provides baseline information for microhylid species that will allow comparisons of ontogenetic trajectories, heterochronic patterns influencing larval body plan, and the role of larval morphology on the adult microhylid body plan.

The understanding of anuran larval diversity has grown in the past 40 yr owing to studies of larval morphology that describe variation and provide baseline data relevant to interpreting anuran phylogenies, larval ecology, or both (e.g., Sokol, 1975; Altig and Johnston, 1989; Larson and de Sá, 1998; Haas, 2003). These studies focused on assessing variation among tadpoles from a wide spectrum of evolutionary clades independent of their specific larval developmental patterns, because such morphological descriptions of tadpoles were based on standard staging tables. Those most commonly used tables are Nieuwkoop and Faber (1956) on Xenopus laevis and Gosner (1960) on Inuiulus valliceps (Bufo valliceps Auctorum). The consequence of basing larval descriptions on standard tables is the assumption that developmental events (e.g., toe differentiation, forelimb emergence, and tail loss) are equivalent stages of tadpole development across all taxa and that these events are not subject to species-specific patterns of developmental timing. Studies assessing morphological variation of tadpole development provide detailed information for individual species (Hall and Larsen, 1998; Hall et al., 1997, 2002) or report the effects of heterochrony on larval growth, such as delayed loss of larval features and early differentiation of adult-like traits (e.g., Downie et al., 2004; Fabrez and Quinzio, 2008; Fabrez et al., 2009, 2010). These studies demonstrate the richness of information derived from the analyses of the entire larval development—including timing of development—to understand anuran diversity and the reciprocal influences between larval and adult body plans.

The clade Microhyliidae consists of 487 described species (Frost, 2011). New World microhylids are clustered in three clades; the largest is Gastrophryninae with 52 species and includes Dermatonotus muelleri. Microhylids are predominantly terrestrial (i.e., fossorial to semifossorial) and mostly inhabit tropical and subtropical lowland rain forests. The monophyly of Microhylidae is supported by adult morphology, molecular data, and a substantial accumulation of apomorphic larval features with several autapomorphic characters (Ford and Cannatella, 1993; Haas, 2003; Frost et al., 2006). Roelants et al. (2011) suggested that among the Neo-batrachia radiation, the Microhylidae stands out by its high evolutionary rate and low levels of homoplasy with other clades and that environmental or intrinsic factors that have limited larval evolution in other anurans played a minor role on the microhyloid radiation. Despite the distinctiveness of the microhylid tadpole, representing larval morphotype II (Orton, 1953, 1957), few studies had focused on larval development. The absence of developmental data for microhylid taxa is one of the limitations in interpreting their evolutionary history.

Herein, we describe morphological variation during larval development of the gastrophrynine microhylid Dermatonotus muelleri and include new characters and information derived from external morphology and field data on larval growth. Descriptions of developmental morphological variation in the lateral line system, skeletal development, and other aspects of soft morphology (e.g., tongue, digestive tract, gonads, and thyroid glands) are presented. The study provides baseline information for microhylid species that will 1) allow comparisons of ontogenetic trajectories within anurans, 2) address heterochronal patterns influencing the larval body plan, and 3) interpret the role of larval morphology on the adult microhylid body plan.

MATERIALS AND METHODS

Larval Dermatonotus muelleri (Boettger, 1885) were collected between November and April 2000–2007 in ephemeral ponds along Ruta Nacional 81 (23°10’S, 63°39’W to 23°14’S, 63°39’W) in Departamento San Martín, Provincia de Salta, Argentina. Specimens were fixed in 10% formalin in the field. Lots of specimens are deposited in the Herpetological Collection of the Museo de Ciencias Naturales (MCN), Universidad Nacional de Salta, Argentina, with the following collection data: MCN 1162 (25 November 2000), MCN 1163 (17 January 2000), MCN 672 (2 January 2001), MCN 1102 (11 November 2002), MCN 1016 (4 March 2004), MCN 946 (11 December 2004), MCN 960 (19 December 2004), MCN 1094 (29 December 2004), MCN 1080 (12 March 2005), MCN 1180 (13 April 2005), MCN 1066 (18 December 2005), MCN 1009 (28 December 2005), MCN 1357 (28 November 2007), 1358 (12 December 2007), MCN 1340 (27 December 2007), and MCN 1332 (5 January 2008), and six adult specimens that are deposited with lot MCN 997.

We selected 70 specimens from lots MCN 1332, 1357, 1358, and 1340 to include individuals in a larval series ranging from hind limb bud stages up to absence of the tail stage. Specimens of these lots were collected from the same pond and had started...
Table 1. Comparisons of key features between anuran larval stages by Gosner (1960) and larval stages of Dermatonotus muelleri (this study).

<table>
<thead>
<tr>
<th>Gosner larval stages</th>
<th>D. muelleri stages</th>
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<tr>
<td>26–30: Limb bud length from one-half diameter to twice diameter. Complete mouthparts.</td>
<td>26–30: Limb bud length from one-half diameter to twice diameter. Complete mouth, upper lip with well differentiated alae.</td>
</tr>
<tr>
<td>41: Maximum larval size. Breakdown of larval mouthparts.</td>
<td>41: Snout uplifted and nostrils protruded. Forelimbs protruding.</td>
</tr>
<tr>
<td>44: Angle of mouth between the midpoint and the posterior margin of eye. Caudal musculature greatly reduced.</td>
<td>44: Angle of mouth reaches the anterior border of eye. Spiracle scar is still present. Caudal fins and tail musculature well differentiated. Eyelids.</td>
</tr>
<tr>
<td>45: Angle of mouth at the posterior margin of eye. Tail reduced to stub.</td>
<td>45: Angle of mouth reaches the mid-level of the eye. Strong shortness of the tail, musculature and fins are indistinct. Vestige of spiracle has disappeared. Differentiation of the nictitans membrane.</td>
</tr>
<tr>
<td>46: Tail completely resorbed. Angle of mouth may be positioned caudal the posterior margin of eye.</td>
<td>46: Tail reduced to a dehiscent stub or absent. Angle of mouth between the mid-level and the posterior margin of eye.</td>
</tr>
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We examined transverse sections of tadpoles at stages 35 and 43 to analyze histological changes in the thyroid glands and operculum. Formalin-preserved tadpoles were dehydrated, embedded in paraffin, sectioned at 6 μm, and stained with hematoxylin and eosin following the protocol of Martoja and Martoja-Pierson (1970). Observations and photographs were made with Leica DM EP compound microscope equipped with digital camera.

Larval and adult specimens of Elachistocleis bicolor (MCN 797, MCN 691, MCN 945, MCN 797, MCN 996, MCN 1010) and adult Phrynomantis bifasciatus (MCN 830), Gastrophrynecolus carolinensis (FML 3565), and Scinax acuminatus (MCN 800) were used for morphological comparisons. Museum abbreviations are as follows: FML, Instituto de Herpetología, Fundación Miguel Lillo, Tucumán, Argentina; and MCN, Museo de Ciencias Naturales, Universidad Nacional de Salta, Argentina.

Anatomical terminology is based on the amphibian anatomical ontology (www.amphibanat.org).

RESULTS

Larval Stages Based on External Morphology.—The development of D. muelleri is summarized and compared with standard Gosner’s stages (Table 1; Fig. 1).

Stages 26–37: These stages are based on hind-limb development and toe differentiation. Morphology of the developing limbs resembles that of Elachistocleis bicolor (Goldberg and Fabrezi, 2008). Throughout these stages, the color in life and in preservative of tadpoles is uniform, ranging from clear greenish to dark brownish in different individuals; the skin is opaque and only the tail fins are transparent. During this period, the upper labium of the terminal mouth grows and reaches the fully developed tadpole morphology, i.e., well-differentiated and quadrangular alae separated by a rounded notch; it completely...
Fig. 1. Changes in external morphology during larval development and metamorphosis in Dermatonotus muelleri. (A–I) Mouth transformations. (A) Stage 28, frontal view. The upper labium is divided by a medial notch into two quadrangular flaps overlapping the lower labium. (B) Stage 35, frontal view. Each flap presents rounded border and a fang-like latero-external projection. At this stage, the nostrils are perforated. (C) Stage 39, frontal view. The larval mouth becomes different. Upper labium is straight without the medial notch and flaps. Lower labium is now visible. Angle of the mouth is placed between the nostril and the eye. (D) Stage 42, frontal view. (E–F) Stage 43, frontal and lateral views respectively. (G) Stage 44, lateral view. Angle of the mouth reaches the anterior margin of the eye. (H) Stage 45, lateral view. Angle of the mouth is positioned at the midpoint of the eye. Upper eyelid has appeared. (I) Stage 46, lateral view. Angle of the mouth is located between the midpoint and the posterior margin of eye. The nictitating membrane is evident. (J–M) Tail changes. (J) Stage 28. Dorsal and ventral fins are symmetrical and tall. (K) Stage 42. Pigmentation is evident in fins. (L) Stage 44. Fins become lower. (M) Stage 45. Tail is an atrophied stub. (N) Stage 43, ventral view. There is a medial scar representing the vestigial spiracle. (O) Stage 44, ventral view. The body wall presents different patterns of pigmentation. Scale bar = 2 mm.
overlaps the lower jaw (Fig. 1A, B). The spiracle is posteromedial in position and opens slightly anterior and ventral to the vent tube; the vent tube is short, funnel-shaped, and attached to the ventral fin. External nostrils, eyelids, and nictitating membranes are absent. The tail conserves its shape during the period (Fig. 1J).

Stages 38 and 39: The metatarsal tubercle and subarticular tubercles on the toes develop in this period. The external nostrils open. The alae of the upper labium are oriented obliquely and have “fang-like” ventrolateral projections (Fig. 1C). The vent tube is anteriorly adjacent to the origin of the ventral fin. Dorsally, the tadpole is uniformly dark, whereas the abdomen is patterned with discontinuous, longitudinal, and dark stripes on a whitish background. The hind limbs are dark.

Stage 40: The vent tube is absent and the cloacal opening is developed. The reduced spiracle is located farther anterior on the ventral wall of the body than before. Anteriorly on the ventrolateral sides of the body, rounded and uniformly pigmented patches of skin mark the position where the forelimbs will emerge.

Stage 41: The tadpole’s snout begins to resemble that of the adult. It is slightly uplifted and acquires an overall pointed appearance; external nostrils are prominent and the mouth opening is ventral. In dorsal view, the forelimbs begin to protrude through the lateral body wall.

Stage 42: The forelimbs have emerged. The forelimb skin has a distinct line marking the union of the forelimb with the skin of the body wall. The upper labium of the larva has begun to disappear in the region of the lateral alae, but the “fang-like” projections are still visible (Fig. 1D); the mouth is now transversely oriented. The spiracular opening remains. The dorsal fin decreases in height, but the tail length does not change (Fig. 1K).

Stage 43: The mouth opening widens and the angle of the jaw lies between the nostril and the eye. The alae of the upper labium have almost disappeared (Fig. 1E, F), and the spiracle lies farther anterior, with the spiracular opening being posterior to the forelimbs (Fig. 1N).

Stage 44: The angles of the jaws are located at the level of the anterior border of the eyes at stage 44 (Fig. 1G). The spiracular opening is a longitudinal slit. The caudal fins are lower (Fig. 1L), and the eyelids are incipient (Fig. 1G).

Stage 45: The angles of the jaws lie at the midlevel of the eyes (Fig. 1H), and the spiracle is absent. The tail is substantially shorter, and the fins and musculature are atrophied (Fig. 1M). The nictitating membrane begins to differentiate (Fig. 1H).

Stage 46: The disappearance of larval structures at stage 46 marks the end of the metamorphosis. The angles of the jaws extend slightly posterior to the midlevel of the eyes (Fig. 1I), and the atrophied tail stub is short or absent. The ventral surface of the body has well-defined, colored areas, i.e., gular, pectoral, and ventral areas (Fig. 1O). The eyelids and nictitating membrane are not fully developed (Fig. 1I).

Size and Timing of Larval Development.—Larval somatic growth increases until stage 37 when the tail length is about equal to snout–vent length (SVL; Fig. 2) and the SVL is at its maximum, corresponding to the size of the metamorphosed individuals (≈14 mm). During metamorphosis, total length decreases owing to tail resorption that occurs abruptly in stage 45. The postmetamorphic size is approximately 3 times the size at metamorphosis, with adults being 40–50 mm. Development,
from fertilization to the end of metamorphosis, occurs in 25–30 days.

Anuran larval development is characterized by three functional developmental periods: 1) differentiation, development, and disappearance of unique larval features; 2) remodeling of larval into adult structures during metamorphosis; and 3) differentiation and development of adult structures.

**Disappearance of Larval Traits.**—Among the larval structures that disappear are the oral structures, vent tube, spiracle, and tail; these structures have been described in other studies (e.g., Gosner, 1960; Duellman and Trueb, 1986; Fabrezi, 2011). Herein, we focus on the lateral line system.

Tadpoles throughout stages 26–39 have a lateral line system consisting of nine pairs of lateral lines—three pairs of orbital lines, three pairs of mandibular lines, and three pairs of trunk lines (Fig. 3). The orbital lines are the preorbital (pre), infraorbital (io), and postorbital (po). Preorbital and infraorbital lines are continuous with each other behind the eye and extend anteriorly dorsal and ventral to the eye respectively (Fig. 3B). The postorbital line lies posterior to the confluence of the preorbital and infraorbital lines and is anteriorly concave. Mandibular lines comprise the oral (or), longitudinal oral (lor), and angular (an) lines that are visible on lateral and ventral views (Fig. 3B, C). The oral line runs between the angle of the mouth forming a ventral semicircle. The longitudinal oral lines extend from the angle of the mouth, posterior to the oral line, and run ventrally and posteriorly. The angular lines originate below the posterior margins of the eyes and descend between the longitudinal oral lines and the ventral lines. The angular lines converge ventrally without contacting with each other and terminate anterior to the ventral line and the spiracle (Fig. 3C).

The trunk lines are the dorsal (d), medial (m), and ventral (v). The dorsal line originates behind the postorbital line (internally at the level of the posterior margin of the otic capsule) and extends upward along the trunk and the anterior third of the dorsal fin (Fig. 3A, B). The medial line diverges from the dorsal line and runs along the trunk and the hypaxial musculature of the tail for about half the length of the tail. Laterally, the ventral line resembles a horizontal S. The anterior arm of the S descends and runs posteriorly to converge at midlength of the spiracle with its contralateral member (Fig. 3C).

The lateral lines resemble dashed lines under stereomicroscopy. Each dash consists of two to six neuromast organs. SEM images reveal that each neuromast organ is circular, with its perimeter defined by an outer layer of mantle cells surrounding a central area that consists of the sensory cells surrounded by microvilli of the supporting cells (Fig. 4A). The neuromasts are as much as 20 μm across and are located in shallow depressions of the epidermis with a variable number of sensory cells (three to six, Fig. 4B).

The orientation of the dashes in a lateral line varies. For example, in the oral, longitudinal oral, and medial lines, the main axis of the dashes are perpendicular to the orientation of the line, whereas in other lines, the axes are parallel to the orientation of the line. There is also variation in the orientation of the dashes along some lines; for example, along the angular and ventral lines, the dashes gradually shift from a perpendicular to parallel orientation.

The lateral line system is considerably reduced by stage 40. The preorbital, infraorbital, oral, and longitudinal oral lines are shorter, and the neuromasts of the dorsal and medial lines disappear along the trunk. The angular and ventral lines are anteriorly displaced concomitant with the displacement of the spiracle. The ventral line is disrupted by the appearance of lateral patches of skin marking the development of the forelimbs. During stage 42, the neuromasts of the preorbital line are restricted to the area between the nostrils. In addition, the postorbital line is shortened and the infraorbital, oral, and longitudinal oral lines have completely disappeared; the neuromasts of the angular and ventral lines are smaller. By stage 43, the preorbital, postorbital, angular, and ventral lines are absent, but the neuromasts of the medial line along the caudal musculature are still present. By stage 45, all neuromasts have disappeared.

**Remodeling of Larval Traits during Metamorphosis.**—Several tissues, organs, and organ systems that differentiate in early larval stages subsequently are remodelled to become adult structures. Herein, we describe the sequence of developmental events and the timing of changes in the digestive tract, chondrocranium, and hyobrachial skeleton.

Tadpoles in stages 26–39 characteristically have a long intestine that is nearly uniform in diameter with a thin and almost transparent wall. The foregut is located on the right side of the abdominal cavity and forms a U-loop around an elongated pancreas. Anteriorly the hindgut turns and forms a double spiral placed on the left side of the abdominal cavity. In contrast, the posterior hindgut is coiled into a double spiral that is medial in position, with the most inner segment opening into
the cloaca (Fig. 5A). The pancreas and an incipient liver lie to the right and overlap the foregut. Lungs are inflated and lie dorsolateral to the liver and intestine.

At stage 40, thin, transparent layers of tissue cover the abdominal cavity where a few longitudinal muscular fibers insinuate the development of the rectus abdominis muscle. The liver consists of three large lobes occupying the anterior portion of the abdominal cavity, and a large gall bladder lies among the lobes on the caudal portion of the liver. The foregut is dilated; it has lost its U shape and become straight, thicker, and distinct from the hindgut that has decreased in diameter. The anterior intestinal coils lack an organized pattern of turns; the posterior hindgut is dilated and located in the midline (Fig. 5B).
Between stages 41 and 43, four pairs of clustered muscle fibers differentiate in the ventral wall of the body to initiate the adult arrangement of the adult rectus abdominis muscle. Overall, the walls of the foregut thicken and the hindgut shortens.

At stage 44, the foregut has a differentiated and expanded stomach morphologically defined by anterior and posterior sphincters that separate it from the oesophagus and the intestines, respectively. The anterior hindgut is curved but does not spiral as seen in the posterior hindgut (Fig. 5C). By stage 45, the distal portion of the hindgut is larger in diameter and forms the rectum, and the rectum has a dilated caecum (Fig. 5D).

The chondrocranium and hyobrachial skeleton of *D. muelleri* tadpoles in stages 31–37 (Fig. 6A) were described by Lavilla (1992) and Vera Candioti (2007). Herein, we describe developmental changes in the chondrocranium starting at stage 38.

Similar to previous stages, the neurocranium is broad; its length is ≈78% of its greatest width at level of the otic region. Cartilaginous structures roofing the cavum cranii are absent, and it is delimited by the tectum synoticum (posterior), the tectum nasi (anterior), and the orbital cartilages (lateral). The ethmoidal region and the olfactory tracts are short. The otic capsules are elongated (≈25% longer than wide) and oriented with their long axis slightly divergent from the sagittal plane. The septum nasi is broad and positioned medially and transversely to the ethmoidal plate (Fig. 6A). The ethmoidal plate extends anteriorly beyond the septum nasi to the level of the processes quadro-ethmoidalis, diverging into short and distally expanded cornua trabeculae. At the same time, the tectum nasi is developed and it is posteriorly continuous with the laminae orbitonasales. The palatoquadrate has a short and robust pars articularis quadrati, a broad commissura quadratocranialis anterior, a broad and low processus muscularis ratocranialis anterior, a broad and low processus muscularis (oriented slightly outward), and a relatively narrow and straight arcus subocularis that terminates in a stout and outwardly projecting cartilaginous bar, the pars posterior. The palatoquadrate attaches to the braincase by three cartilaginous bridges. (1) The broad commissura quadratocranialis anterior (diverging anteriorly in approximately a 45° angle with the main chondrocranial axis) joins the anterior portion of the palatoquadrate to the trabecular plate. (2) The processus ascendens diverges toward the braincase from the arcus subocularis at the level of the pars posterior and links the palatoquadrate with the orbital cartilage. (3) The larval processus oticus is ventrally continuous with the pars posterior and dorsally with the processus anterolateralis of the crista parotica. On the ventral surface of the pars posterior there is a narrow posterior process extending backward. Behind the larval processus oticus and extending from the otic capsule, a broad and thin cartilage extends ventrally and is continuous with the posterior edge of the pars posterior; this lamina is extensively perforated. The suprarostral is a single laminar cartilage with a medial concavity, which provides the skeletal support for the upper lip alae. The distal ends of the cornua trabeculae are expanded and straight and each half of the suprarostral is extensively fused with the anterior margin of each cornu. The lower jaw consists of a single infraorostral that is a slender element forming an inverted U-shaped structure, with lateral cartilaginous expansions representing the Meckelian diverticula (Fig. 6F, G) and a pair of lateral Meckel’s cartilages. Meckel’s cartilages are transverse bars with acuminate and pointed medial tips that are contiguous to the infraorostral; the outer ends are broad, robust, and articulate with the pars articularis of the palatoquadrate.

Changes in the ethmoidal region begin at stage 39 (Fig. 6B). The septum nasi is now slender. The paired anterior nasal walls grow laterally to the anterior end of the septum nasi. At this stage, the anterior nasal walls remain independent from the trabecular plate. Furthermore, the posterior nasal wall lifts up from the commissura quadratocranialis anterior.

By stage 40, the septum nasi has grown anteriorly reaching the divergence of the trabecular horns. The alary cartilages are visible as small, slender, elements located in front of the septum nasi on the trabecular plate. The alary cartilage is continuous with the anteriorly placed inferior prenasal cartilage and laterally with the developing laminae superior and inferior crista intermedia. The septum nasi projects forward as a short median prenasal process. The developing oblique cartilages are L-shaped, with the longer and transversely positioned arm being continuous with the septum nasi. The planum triangulare has been differentiated.

At stage 41, the erosion of the cornua trabeculae has begun and the cartilages of the nasal capsule form a complex structure. The incipient crista subnasalis develops ventral to the alary cartilages. The anterior and posterior nasal walls and the planum triangulare rise up bordering the commissura quadratocranialis anterior. The processus muscularis of palatoquadrate and the posterior cartilaginous connections between the palatoquadrate and the braincase have begun to erode. The suprarostral is still present, but resorption has started and progresses from medial to lateral direction.

By stage 42 (Fig. 6C), the cornua trabeculae are absent. Erosion of the suprarostral has advanced resulting in two lateral free pieces. The snout of the tadpole now is truncate in profile and composed of the frontal alary cartilages and the processus prenasalis. The oblique cartilages, as well as the laminae superior and inferior crista intermedia, are vertical in position. Ventrally, the crista subnasalis is massive and its posterior process is well defined. The nasal walls have grown and the planum triangulare has a processus maxillary anterior. The tectum nasi has expanded anteriorly and laterally covering the septum nasi along its length. The commissura quadratocranialis anterior is slender. Meckel’s cartilages and infrarostral have started to undergo morphological reshaping and fusion. The infrarostral forms a transverse bar with a distinct cartilaginous medial union. Lateral and ventral to the “mandibular synphysis” the pair of Meckelian diverticula became massive (Fig. 6H). Meckel’s cartilages are longer and oriented obliquely.

The medial and anterior cartilages of the nasal capsule have grown into a well-developed complex by stage 43. The commissura quadratocranialis anterior seems to be fused to the planum triangulare, forming a new anterior attachment of the palatoquadrate to the neurocranium. Erosion of the arcus subocularis has advanced anteriorly. The processus articularis is slender and rounded. The lower jaw lengths concomitantly with the processus pterygoideus that represents the posterior growth of most lateral portion of the commissura quadratocranialis anterior.

By stage 44 (Fig. 6D), the cartilaginous snout nearly has its adult form. The processus pterygoideus is curved, short, and continuous with the pars articularis of palatoquadrate that is now expanded, rectangular, and well differentiated. The cartilaginous pars articularis is perpendicularly oriented to the processus pterygoideus, and its anteroventral tip is modified to articulate with the Meckel’s cartilage. Lengthening of the Meckel’s cartilage continues throughout this stage.
Fig. 6. Cartilage remodeling during advanced larval stages. (A–E) Dorsal views of the chondrocranium. Scale bar = 0.5 mm. (A) Stage 35, the chondrocranium similar to that described for stage 38. (B) Stage 41. Differentiation of nasal walls and nasal capsules cartilages is underway. Erosion of suprarostral and cornua trabeculae is advanced. (C) Stage 42. Vestiges of the suprarostral alae appear as free cartilages. The cartilaginous skeleton of the nasal capsule is formed. Posterior attachments of the palatoquadrate to the neurocranium are eroded and resorption of the subocularis arch is evident. (D) Stage 44. The palatoquadrate has changed abruptly. The processus pterygoideus is well differentiated and it is the new cartilaginous union of the upper jaw to the skull. The pars articularis of the palatoquadrate migrated backwards concomitant with the growth of the processus
At stage 45 (Fig. 6E), the nares have a terminal position in dorsal view. The snout projects forward and beyond the lower jaw. The pars articularis of palatoquadrate is reduced and attaches by ligaments to the otic capsule; a poorly developed medial processus basalis is present. The cartilaginous lower jaw has distinct boundary between Meckel's cartilage and infrarostral, a well-developed mandibular symphysis, and the prominent Meckelian diverticula (Fig. 6I).

The transformation of the larval chondrocranium is complete at stage 46. The skull is short and the braincase is wider posteriorly than anteriorly. Cartilaginous elements roofing the cavum cranii did not develop. The occipital condyles flank the foramen magnum and are widely separated. The lower has a medial cartilaginous symphysis (Fig. 6).

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The larval hyobranchial skeleton is characterized by 1) slender, transverse ceratohyalia with well-defined anterior, medial, lateral, and posterior processes; 2) a broad pars reuniens; 3) an elongated copula with a long processus urobranchialis; 4) small, unfused hypobranchialia; and 5) four ceratobranchialia with numerous and elaborated rays. The ovoid branchial baskets extend from the level of the solum nasi to the level of Presacral Vertebra IV.

Metamorphic changes begin at stage 40 (Fig. 7A) with the rapid resorption of the ceratobranchialia in a posterior-to-
anterior direction, and from Ceratobranchial IV to I. The hypobranchialia fuse medially and have a pair of posterior projections delimiting the laryngeal sinus.

During stage 41 (Fig. 7B), the anterior, lateral, and posterior processes of each ceratohyal have begun to erode. A medial chondrification—identified here as Cartilage Z—develops ventral to the hypobranchial plate and extends forward from the laryngeal sinus. The developing hyoglossus muscle attaches to the posterior margin of Cartilage Z.

At stage 43 (Fig. 7C), the hyobranchial skeleton still has identifiable larval ceratohyalia, pars reuniens, copula, and hypobranchialia; however, these elements form a single, contiguous, cartilaginous complex. Each robust ceratohyal still retains the rounded processus hyoquadrum. The pars reuniens and copula are approximately twice their original lengths. The hypobranchial region has lateral projections with remains of the eroding Ceratobranchial I that will differentiate into the posterolateral processes. On the hypobranchial plate, the posterior processes have reshaped into well-differentiated posteromedial processes.

During stage 44 (Fig. 7D), the ceratohyalia become slender and have an oblique position; the hyoglossal sinus is now well defined.

At stage 45 (Fig. 7E), each ceratohyal still possess rounded tip representing the vestige of the larval processus hyoquadrum. The anterolateral process of the adult hyoid apparatus forms from new cartilages lateral to the copula. The posteromedial processes have well-differentiated distal epiphyses.

Stage 46 marks the end of the metamorphic changes in the hyobranchial apparatus (Fig. 7F). The processus hyoquadrum has disappeared and the ceratohyal is uniformly slender. New cartilages give rise to the posterolateral processes on the hyoid plate. Cartilage Z is incompletely incorporated into the edge of the hyoid plate between the posterolateral processes. At this stage, the cartilaginous hyoid apparatus acquires the adult configuration and appearance.

The posteromedial processes ossifies postmetamorphically. Cartilage Z is incorporated into the ventral edge of the hyoid, ossifies as a spurlike process between the ossifications of the posteromedial processes, and projects ventrally. In adults, a medial and anterior mineralization develops on the hyoid plate that corresponds to a mineralized larval copula.

**Development of Postmetamorphic Traits.**—Some anuran morphological features traits have little or no function in larvae, but they are necessary components of the adult body plan. Among those that we consider herein are the gonads, tongue, thyroid glands, limbs, bony skull, and plectral apparatus.

At stage 26, the gonads are undifferentiated and resemble whitish, elongated, cylindrical, and paired cords diverging caudally; the left cord is larger than the right (Fig. 5F). In stage 38, the undifferentiated organs are thin, laterally compressed cords. During metamorphosis, at stage 43, the gonads remain undifferentiated (Fig. 5G).

The tongue (Fig. 5H–K) is first evident in stage 36 as an anterior anlage on the buccal floor. The anlage grows, conserving its initial cordiform shape until stage 42, when the tongue lengthens posteriorly and develops a pointed end that becomes rounded in stage 44. At the end of metamorphosis, the tongue is ovoid and occupies the entire buccal floor. The paired hyoglossus muscles begin to differentiate in the tongue in stage 40; each muscle grows posteriorly and the number of muscle fibers increases. By stage 42, the tendon of the hyoglossus muscle fibers attaches to the developing Cartilage Z on the

hyobranchial apparatus. The muscles lengthen and by stage 46 they are firmly attached to the protuberance formed by Cartilage Z.

Beginning at stage 36, the gland, follicle, and colloid volumes gradually increase and the epithelial cells become higher. There is no significant or abrupt variation of the thyroid glands during subsequent metamorphic stages (Figs. 5L–O, 8A–D).

Primary cartilages of limbs are differentiated by stage 34. The carpus is composed of seven cartilaginous elements (ulnare-intermedium, radiale, Distal Carpal 5+/4+3, Distal Carpal 2, and Element Y). The tarsus comprises six distinct cartilages distal to the enlarged proximal tarsals (fibulare-tibiale) (Distal Tarsals 3, 2, and 1, the proximal prehalical element in the dorsal plane, and two primary cartilages that form Element Y in the ventral plane). Fusion of Distal Tarsals 3 and 2, as well as the formation of Element Y (by fusion of the two primary cartilages) occur at the end of stage 35.

The humerus is the first bone to ossify in the forelimb at stage 33, closely followed by the radius and ulna that are not fused yet. The ossification of the hind limb begins at stage 34, slightly after the ossification in the forelimb. Ossification of autopodia begins before all the cartilaginous phalanges have differentiated. Before the beginning of metamorphosis, at stage 40, all diaphyses of limbs have ossification centers. The ossification sequences of the hand and foot progress in proximal–distal and postaxial–preaxial directions. Ossification of the carpals, prepollex, distal tarsals, and prehallux occurs after metamorphosis.

At the end of stage 37, the endochondral ossification of the exoccipitals develops around the jugular foramen and extends toward the perilymphatic foramina.

By stage 38, the dermal paraplitoid has a broad cultriform process and short alae that cover the cranial floor and partially overlap the otic capsules. The endochondral ossification of the prootics begins on the inner wall of the otic capsule and expands anteriorly from the auditory foramina to surround the prootic and oculomotor foramina in the orbital cartilage.

In stage 39, each frontoparietal differentiates and grow medially; the frontopariets grow rapidly and expand broadly behind the orbit.

At stage 40, the nasals appear over the developing cartilages of the nasal capsules. The septomaxillae, maxillae, and premaxillae are well differentiated by stage 41. The septomaxilla is a convex, laminar ossification attached to the developing oblique and alaris cartilages. Each premaxilla is a triangular bone with the ventral surface resting on the cornu trabecula. The maxilla ossifies anterior to the planum triangularis.

At stage 42 (Fig. 9A–C), the squamosals is a tiny, horizontal ossification attached to the base of the eroded processus muscularis of palatoquadrate. In the lower jaw, two ossifications develop anteriorly on Meckel's cartilages; the dentary differentiates along the labial surface and the angulosplenial on the lingual surface.

Subsequently, jaw and suspensorium bones (dentaries, angulosplenials, maxillae, and squamosals) and mandibular cartilages grow posteriorly. The ventral ramus of the squamosal gradually becomes vertical, along with the palatoquadrate and pars articularis; short rami (i.e., zygomatic and otic) develop and differentiate from the ventral ramus (Fig. 9D–L).

At stage 45, the quadratojugals and pterygoids are differentiated. The pterygoid, is attached to the posterior end of the processus pterygoideus of the palatoquadrate and remains small until the end of the metamorphosis, whereas the
quadratojugal grows anteriorly to articulate with the maxilla (Fig. 9M–O).

Ossification of cranial bones begins during larval development; however, the most extensive ossification of the skull takes place during postmetamorphic growth (Fig. 10A, B). Frontoparietals form a wide roof to the cavum cranii but they do not fuse medially. The large nasals completely cover the olfactory capsules and the alary cartilages and the processus prenasalis project forward over the anterior margins of the nasals. The superior and inferior prenasal cartilages are hidden by the premaxillae. The processus prenasalis, solum nasi, and septum nasi are mineralized. In ventral view, the ossification of the sphenethmoid extends to the premaxillary–maxillary articulation; neopalatines are absent (Fig. 10B). The sphenethmoid is not visible in dorsal view and forms an incomplete ring around the anterior end of the braincase; the two halves do not contact ventrally. The otic capsules are well ossified and the prootics are indistinguishably fused with the exoccipitals. The occipital condyles are widely separated. The ossification of the prootic extends anteriorly into the pila antotica; the prootic and oculomotor foramina are completely surrounded by this ossification. The mineralized crista parotica is wide, flat, and stout. Ossification of the braincase is extensive and contrasts with the reduced ossification of the jaws. Premaxillae and maxillae are edentulous. Each premaxilla has a rectangular pars palatine with a reduced processus palatinus. The pars alaris of

Fig. 8. (A–D) Histological changes in the thyroid glands. (A–B) Stage 35, the glands are organized into follicles, each containing a central lumen, and some lumina are beginning to accumulate colloid. The follicles are lined by a single layer of cuboidal epithelial cells. (C–D) Stage 43, the thyroids present enlarged follicles containing colloid and vacuoles within the colloid. (F–G) Transverse sections of otic capsule. (F) Stage 35, showing the incipient operculum in the fenestra ovalis. (G) Stage 43, the operculum is differentiated as well as the muscle opercularis. Abbreviations: o, operculum; oc, otic capsule; om, opercularis muscle; and th, thyroid gland.
FIG. 9. Sequence of ossification. Scale bar = 1 mm. Left column is dorsal view, central column is ventral view, and right column is lateral view. (A–C) Stage 42. Ossification of nasals (n), frontoparietals (fp), septomaxillae (smx), exoccipitals (exo), and prootics (pro) are advanced, and differentiation of premaxillae (pmx), maxillae (mx), squamosals (sq), dentaries (d), and angulosplenials (as) progresses. (D–F) Stage 43. (G–H) Stage 44. The squamosal turns and has a vertical position. Short zygomatic and otic ramii are differentiated. (I–L) Stage 45. The quadratojugals and pterygoids have appeared. (M–O) Stage 46. The skull at the end of larval development is incompletely ossified. The pterygoid is a tiny ossification adjacent to the internal face of squamosal and attached to the posterior end of the cartilaginous pterygoid processus of the palatoquadrate.
the premaxilla is short and vertical; it is visible only in frontal view. The pars facialis of the maxilla is low, without a differentiated processus nasalis, and the pars palatina is reduced. The quadratojugal is tiny and touches, but does not overlap, the posterior tip of the maxilla. The ventral ramus of the squamosal is short and vertical, and the otic and zygomatic rami are poorly differentiated. The ossification of the squamosal does not contact the otic capsule. The anterior ramus of the pterygoid lingually invests the pterygoid process throughout the posterior half of the maxilla, whereas the medial ramus is short and does not reach the otic capsule. The posterior ramus of the pterygoid terminates at the cartilaginous pars articularis of the palatoquadrate. The cartilaginous attachments of the upper jaw to the skull do not stain with Alcian Blue. The vomers are small, thin, and triangular “scales” located along the anteromedial margins of the choanae. On the lower jaw (Fig...
shaped. The tympanic membrane is absent. The pars externa plectri is small. The tympanic annulus is sickle-oriented obliquely towards the squamosal. The cartilaginous is slender and long, visible in dorsal and ventral views, and develops during postmetamorphic stages (Fig. 10H). The stapes ovalis. The developing myoblasts of the opercularis muscle continuous with the cartilaginous dorsal margin of the fenestra ovalis show connective tissue and a more uniform, extracellular matrix at the perichondral sections show chondrocytes and large lacunae placed medially of the membrane covering the fenestra ovalis show connective tissue with few, parallel, elastic fibers and small fibroblasts surrounding the anlage of the operculum (Fig. 8E). The anlage is composed of a small oval group of round, basophilic cells with spherical nuclei; the cells are immersed in a matrix lacking fibers and lacunae. The organization of this connective membrane is looser toward the periphery. In whole-mounted specimens, the operculum gradually becomes Alcian Blue positive; it grows to fill the fenestra ovalis and is well defined at stage 40. At stage 43, the well-differentiated operculum consists of condensed hyaline cartilage surrounded by a thin layer of connective tissue (i.e., the remains of the original membrane). Histological sections show chondrocytes and large lacunae placed medially and a more uniform, extracellular matrix at the perichondral limits (Fig. 8F). The perichondrium of the operculum is continuous with the cartilaginous dorsal margin of the fenestra ovalis. The developing myoblasts of the opercularis muscle appear directly attached to the external, ventrolateral surface of the operculum.

The components of the stapes and the tympanic annulus develop during postmetamorphic stages (Fig. 10H). The stapes is slender and long, visible in dorsal and ventral views, and oriented obliquely towards the squamosal. The cartilaginous pars externa plectri is small. The tympanic annulus is sickle-shaped. The tympanic membrane is absent.

Discussion

The Microhylidae comprises 487 described species (Frost, 2011) that usually are medium- to small-sized species; however, there are several miniature and a few large taxa. Most species have free-swimming and feeding tadpoles, although nonfeeding tadpoles, direct development, or both have evolved independently in many lineages (Clarke, 1996; Glaw and Vences, 2007). Species with free-swimming and feeding tadpoles are characterized by “explosive” seasonal breeding in ephemeral ponds; some may breed multiple times during one season, and tadpoles of different ages are found simultaneously in the same pond. Most available descriptions report that microhylid larval development lasts from 20 to 60 days (Krügel and Richter, 1995), with larvae reaching a maximum total size of 40 mm and maximum SVLs at metamorphosis of no more than 16 mm (Cei, 1981; Donnelly et al., 1990; Chou and Lin, 1997; Vera Candioti, 2006; Glaw and Vences, 2007; Lehr et al., 2007).

Dermatonotus muelleri inhabits the semiarid environments of the Chaco Region in South America; its reproduction occurs explosively three or four times during the wet season on warm days with heavy rainfall. Like other microhylids (e.g., Elachistocleis bicolor), D. muelleri deposits eggs as a film on the surface of the water (Lavilla et al., 1995) and larval development corresponds to the typical biphasic life style of microhylids of tropical and subtropical habitats, conforming overall with the life-history patterns described above.

The end of the larval period in D. muelleri is marked by the absence of the tail and the beginning of the terrestrial life style (Fig. 11). Froglets are small (14 mm), and significant postmetamorphic growth is needed to reach the adult size (40–50 mm). Females are larger than males and gonad differentiation occurs during postmetamorphic stages; rates of body growth could be related to differentiation and maturation of ovaries and testes. Under natural conditions, the development (i.e., from fertilization to metamorphosis) of D. muelleri lasts between 22 and 26 days. The period between stages 41 and 46 corresponds to metamorphic climax and at a maximum, lasts 3 days. By stage 37, tadpoles have attained their metamorphic snout–vent size; this seems to distinguish larvae of D. muelleri from other Type-IV tadpoles in which the snout–vent size increases until the forelimbs emerge (Hall et al., 1997; Quinzio et al., 2006; Fabrezi, 2011). Larval somatic growth and metamorphic development seem to parallel each other in most anurans; thus, one might expect that any negative external factor (e.g., diet, temperature, competence, and predation) might affect the rate of development and modify somatic growth or vice versa (Wilbur and Collins, 1973). However, the early acquisition of maximum larval somatic growth (i.e., metamorphic size) in D. muelleri suggests that size would not be affected by later changes in larval developmental rate.

The loss of larval features during development in D. muelleri occurs between stages 41 and 46 (Fig. 11); structures are lost in the following sequence: 1) vent tube, 2) larval mouth, 3) lateral line system, 4) spiracle, and 5) tail. At the same time (i.e., stages 41–46), internal larval tissues are undergoing transformation. The synchrony of the onset and offset of transformations in digestive tract, jaws, and the hyoglossal apparatus is associated with the development of the adult feeding mechanism. The onset of ossification of the limbs and skull at stage 34 indicates these events are partially independent of other anatomical larval components (Fig. 11).

Tadpoles of D. muelleri have nine pairs of symmetrical lines of neuromasts. The distribution of these lines resembles the generalized pattern described for anurans except for the unusual lengths of ventral and angular lines that may be related to the medial position of the spiracle. The organization of the neuromasts within each lateral line, in irregular groups of five to seven organs, coincides with descriptions for other microhylids (Lannoo, 1987). Studies of the temporal and spatial sequences of the disappearance of the lateral line system reveal that the process starts simultaneously in receptor organs, ganglion cells, and central nuclei at metamorphic climax. The beginning of this process is correlated with high levels of circulating thyroid hormone and the expression of thyroid hormone receptors in the parts of the lateral line system mentioned (Wahnschaffe et al., 1987; Schlosser, 2002). These findings suggest that thyroid hormone triggers the disappearance of the lateral line system (Schlosser, 2002). In D. muelleri, all lateral lines begin to disappear almost simultaneously before the forelimbs emerge. The lateral-line system is lost while caudal fins and tail musculature are still well differentiated. This is in striking contrast to descriptions of other anurans (e.g., Rana temporaria, Hyperolius mitchelli, Wahnschaffe et al., 1987; Pseudis paradoxa, Fabrezi et al., 2009; ceratophryids, Quinzio, unpubl. data).

Limb development in D. muelleri has a spatial and temporal pattern of differentiation of primary cartilages and ossification
resembling that of other anurans (de Sá, 1988; de Sá and Trueb, 1991; Fabrezi and Alberch, 1996; Fabrezi and Goldberg, 2009). This suggests that developmental events related to adult anatomical structures are not closely linked to the development of species-specific unique larval structures. The endochondral ossifications of the diaphyses and some skull bones (e.g., exoccipitals and prootics) are the earliest to appear and continue development in postmetamorphic stages (Fig. 11). Dermal skull bones (e.g., those forming the floor and roof of the braincase and snout) are well differentiated at stage 41 (Fig. 11). However, dermal bones associated with the development of the jaw apparatus seem to be linked to changes in larval jaw cartilage. This pattern of ossification is in agreement with available reports on the ossification of most anurans (Trueb, 1985; de Sá and Trueb, 1991; among others).

Patterns of larval development and gonadal differentiation also are dissociated developmental events (Lopez, 1989; Ogielska and Kotusz, 2004). Differentiation of the gonads occurs before metamorphosis in species with long larval periods (e.g., *Pseudis* spp., Downie et al., 2009; Fabrezi et al., 2010) and after metamorphosis in species with short larval periods (e.g., *Scaphiopus couchii*, Buchholz and Hayes, 2005). *Dermatonotus muelleri* and other anurans with ecological similar breeding sites (e.g., *Scinax acuminatus*, *S. nasicus*, Fabrezi et al., 2010) have a moderately short larval period and delayed gonadal differentiation (Fig. 11).

Transformation of the digestive tract of *D. muelleri* begins before forelimb emergence at stage 41 with changes occurring in an anterior–posterior direction. The onset of other developmental changes related to adult feeding—but involving different cellular types, such as those of the skeleton, muscles, and nerves that will form the hyoglossal apparatus—occurs before stage 41 (Fig. 11).

Changes in the cartilaginous cranial skeleton and visceral arches in *D. muelleri* include the following coordinated events: 1) differentiation and development of the olfactory capsules concomitant to the resorption of the trabecular horns and suprarobranchials; 2) resorption of ceratobranchials and remodelling of ceratohyalia, pars reuniens, copula, and hypobranchialia to form the hyoid skeleton; and 3) lower jaw elongation concomitant to the resorption of the palatoquadrate-neurocranium attachments and differentiation of new cartilaginous connections. These changes are synchronous with each other and precede the development and/or changes that will occur in the musculature and bony skeleton. For example, olfactory cartilages appear before bones of the snout (nasals, vomers, and maxillae). Changes in the hyobranchial skeleton advance accompany tongue development and differentiation of associated hyoglossus and genioglossus muscles, and changes in the palatoquadrate and lower jaw begin just before transformations of the mandibular and hyoid muscles and the differentiation of the mandibular bones (maxillae, squamosals, pterygoids, and quadratojugals). Endochondral ossification of the braincase occurs without major remodelling of cartilages before metamorphic events. The dermal cranial ossification of *Hamptophryne boliviana* is delayed and occurs after metamorphosis (de Sá and Trueb, 1991); however, cartilage remodelling is similar to *D. muelleri*.
During the development of *Dermatonotus muelleri*, as well as that of *H. boliviana* (de Sá and Trueb, 1991), transformations of larval jaws reflect a low developmental rate in which the offset of the palatoquadrate/infrarotal-Meckel's cartilages lengthening at the end of metamorphosis exhibits the condition that most anurans reach at the middle of metamorphosis (stage 44) (Fabrezi, 2011). Paedomorphic features of cartilaginous jaws impact the development of the associated bones that are either reduced (e.g. vomers, quadratejuguale) or absent (e.g. neopalatines).

Microhydil skulls are characterized by the anterior location of the lower jaw articulation (Zweifel, 1985; de Sá and Trueb, 1991; Lehr and Trueb, 2007). Ossification of the squamosal and pterygoïd provides weak reinforcement to the cartilaginous suspensorium. The ventral ramus of the squamosal is nearly vertical and short, the zygomatic ramus is usually small but variable in its development, and the otic ramus is reduced or absent; furthermore, the squamosal does not contact the otic capsule. The pterygoïd also has short posterior and medial rami. In addition, quadratojuguale, vomers, and palatinis may be absent; if present, they are small or less differentiated. The position of the jaw articulation defines a small mouth gape that is characteristic of several adult microhylids.

Trueb and Alberch (1985) described the sequence of appearance of anuran skull bones; furthermore they proposed that paedomorphic cranial patterns could be related to the overall small size of some species given that loss of skull bones (e.g., neopalatines, vomers, quadratojuguale) is infrequent in large-sized species. Yeh (2002) explored anuran intraadinal variation in adult sizes, skull shapes, and skull bones and described some patterns that could be predictive for clades that exhibit size reduction. Despite limited knowledge of the developmental variation within Microhylidae, skull development, skull shape, and skull bone loss in *Dermatonotus muelleri* suggests that these features are a consequence of larval morphology and gradual metamorphosis of the cartilaginous upper and lower jaws (Wassersug and Hoff, 1982).

In *Dermatonotus muelleri* the anterior medial segment of the larval lower jaw consists of a single piece. Vera Candidioti (2006, 2007) described it as fused infrarotals and de Villiers (1930) demonstrated the absence of a discernible symphysis through histological sections of the jaw of *Phrynomantis bifasciatus*. Two infrarotals joined by a copula mandibularis were reported in *H. boliviana* (de Sá and Trueb, 1991). Whether New World microhylids have a single or paired infrarotals needs to be demonstrated with supporting histological evidence from early developmental stages. However, it is worthwhile to highlight that during the metamorphosis of *D. muelleri*, the contribution of the infrarotals to the formation of the cartilaginous skeleton of the adult lower jaw is limited to their most anterior segments that include the Meckelian diverticula (Fig. 6F–J). This fact is different from other anurans in which the metamorphic lengthening of the infrarotals contributes to form up to the 30% of the adult cartilaginous jaw (Fabrezi, 2011:fig. 6). At least in the microhylid *D. muelleri*, the anterior placement of the jaw articulation is both the result of the larval morphology of the infrarotals and the early offset of the metamorphosis of the jaw.

Another feature related with the skeleton configuration of microhylid frogs is the reduction of the tympanic system. *Dermatonotus* and other microhylids lack a tympanic membrane but have a well-developed plectral apparatus as well as the tympanic annulus that differentiates during postmetamorphic growth. The pars interna and media plectri form a slender, long, and well-ossified bar reaching the squamosal (Fig. 10C–F vs. Fig. 9G). The pars externa plectri is a small and oval cartilage surrounded by a small and incomplete tympanic annulus. Absence of the tympanic membrane and reduction in size of the tympanic annulus and pars externa plectri in *Dermatonotus* and other microhylids could be related with the paedomorphic characteristics of the palatoqquadrate (Smirnov and Vorobyeva, 1988). Interspecific comparisons of the plectral apparatus in microhylids reveal that it could be characterized by the reduction of the footplate (Fig. 10C–F).

In summary, the tadpole of *D. muelleri* corresponds to the Orton's morphotype II (Microhyloidea), and it is characterized by an accumulation of several apomorphic features, some of which are unique among anuran larvae (Haas, 2003). Larval development in *D. muelleri* (Fig. 11) reveals an interesting pattern in which remodeling of larval into adult jaws during metamorphosis has an early offset with consequences in the differentiation and development of the bony skull. The morphological changes observed in *D. muelleri* begin at larval stage 35; moreover, they have been related to the thyroid glands' function in anurans with free-swimming larvae and direct development (Etkin, 1936; Jennings and Hanken, 1998; Callery and Elinson, 2000). Standard developmental tables based on Type-IV larvae traditionally define the beginning of metamorphic climax by forelimb emergence (that also involves spiracle disappearance), loss of larval mouthparts, and onset of transformations in the gut and visceral arches (mandibular and hyobranchial skeleton) (Etkin, 1936; Gosner, 1960). Within this context, the developmental sequences in *D. muelleri*, as well as that of other anurans, such as *Pseudis* spp. (Fabrezi et al., 2009, 2010; Fabrezi, 2011), ceratophryines (Fabrezi and Quinzi, 2008; Fabrezi, 2011), and *Sphaenorhynchus bromelicola* (Bokermann, 1974), suggest that the onset of metamorphic changes varies and is not fixed as might be inferred from standard anuran developmental tables. However, the end of larval development is marked by the complete loss of the tail, an ontogenetic event in which changes in larval developmental modules undergo integration with those of the adult stages to converge in the unique adult body plan that seems to be independent with larval growth.

A more comprehensive taxonomic revision of larval morphology, larval development, and their variation among microhylids could provide help to understand the morphological evolution in the group.

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