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# Do Timing and Pattern of Myogenesis Correlate with Life History Mode in Anurans?

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## Recommended Citation

Smetanick, Matthew T., et al. "Do Timing and Pattern of Myogenesis Correlate with Life History Mode in Anurans?" [Journal of Herpetology](https://www.jstor.org/journal/jherpetology), vol. 34, no. 4, [Society for the Study of Amphibians and Reptiles,](https://www.jstor.org/publisher/ssar) 2000, pp. 637–42, https://doi.org/10.2307/1565288.

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Do Timing and Pattern of Myogenesis Correlate with Life History Mode in Anurans? Author(s): Matthew T. Smetanick, Rafael O. de Sá and Gary P. Radice Source: Journal of Herpetology, Vol. 34, No. 4 (Dec., 2000), pp. 637-642 Published by: Society for the Study of Amphibians and Reptiles Stable URL: https://www.jstor.org/stable/1565288 Accessed: 01-11-2021 12:42 UTC

# **REFERENCES**

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 cozumela. They accepted the idea that parthenogens cozumela. They accepted the idea that parthenogens derived from separate hybridizations should be rec- cozumela com ognized as different species, but they did not consider  $\longrightarrow$ , AND it as an exclusive criterion. They agreed with Echelle ation among (1990) that postformational mutational events of suf- (1990) that postformational mutational events of suf ficient magnitude can serve as points of origin for new progenitor s parthenogenetic species (Taylor and Cooley, 1995a, b). We agree with this position. The karyotypic and mor- WALKER, J. M. 19 phological differences and the geographical barrier phological differences and the geographical barrier between *C. cozumela* and *C. maslini* appear to be suf-<br>
and the set of Chemido ficient elements to give species recognition to C. mas-<br>*in the set of the species* recognition to C. mas $lini$ , thereby reflecting the substantial evolutionary di-  $\blacksquare$  WRIGHT,  $\blacksquare$  W. vergence between these two groups of parthenoge-<br>
genus Cnemic netic lizards. parthenogenetic species (Taylor and Cooley, 1995a, b). We agree with this position. The karyotypic and mor- WALKER

Acknowledgments.-To Oswaldo Hernández Gallegos and Felipe Rodriguez Romero for field assistance, to Carmen Loyola and Felipe Villegas for technical assis tance, to DGAPA (Project No. IN210594) and PADEP (Project No. 003007) for financial support to conduct this study, to Mark Paulissen for comments and help in translation, especially to Orlando Cuellar for holis tic support and discussions, and to R. Seigel, J. Walker, and an anonymous reviewer for comments and sug gestions.

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Accepted: 26 July 2000.

Journal of Herpetology, Vol. 34, No. 4, pp. 637-642, 2000 Copyright 2000 Society for the Study of Amphibians and Reptiles

### Do Timing and Pattern of Myogenesis Correlate with Life History Mode in Anurans?

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 The timing and pattern of myogenesis varies among anurans that have been studied and the different pat terns may provide useful phylogenetic information. Specific myogenic markers have been described (Muntz, 1975; Kielbowna, 1981; Boudjelida & Muntz, 1987; Radice et al., 1989) and they can provide infor mation on evolutionary changes for closely related lin eages within a clade. For example, we previously com pared first appearance of a muscle-specific protein, first twitch of axial muscle, onset of multinucleation within axial myotome, and first heartbeat in two pipid genera (Smetanick et al., 1999). We found that al though the timing of myogenesis differed, the se quence of events was the same for these two pipids. The similarities we saw in the two pipids could be due their common lineage, or alternatively, be a result of sharing a life history mode. For example, appear ance of muscle twitch prior to multinucleation could be an adaptation for rapid development, an advantage in frogs with free-swimming tadpoles. If so, it might occur in other lineages with free-swimming tadpoles regardless of phylogenetic distance.

 Herein, we analyze myogenesis among seven spe cies in four anuran families. Species selected for this study also differ in their life histories and general modes of reproduction. We sought to determine whether the timing or sequence of myogenesis, or both, correlated with phylogenetic group or with life history mode.

 We grouped species in three categories based on reproductive mode. The non-direct developers, i.e.,

 TABLE 1. Gosner (1960) stages correlated with Nieuwkoop and Faber (1975) (NF) and Townsend and Stewart (1985) (TS) stages. \* Dashes indicate an in ability to directly correlate a NF or TS stage with a particular Gosner stage. \*\* Gosner correlations of stag es 26-27 to NF stages 46-47 from Just et al 1981.

NF stage	Gosner stage	TS stage
18	15	3
$19 - 21$	16	3
$22 - 23$	17	4
24	18	4
$25 - 32$	∸	
$33 - 34$	19	5
$35 - 39$		5
40	20	5
41	$21 - 23$	6
42	24	6
$43 - 45$	25	6
$46***$	26	6
$47**$	27	6

riymenochirus boettgeri, Kana sylvatica, K. utricularia,<br>Xenopus laevis, and X. tropicalis, lay large clutches of at 2  $\mu$ m using glass knives, transferred to a slide, and<br>eggs in lentic water. Embryos hatch into free-swi *Xenopus iaevis,* and *X. tropicalis,* lay large clutches of<br>eggs in lentic water. Embryos hatch into free-swim-<br>ming tadpoles before developing into juvenile frogs.<br>*Xenopus* and *Hymenochirus* are pipids, but differ in<br>W eggs in ientic water. Embryos hatch into tree-swim-<br>ming tadpoles before developing into juvenile frogs.<br>*Xenopus* and *Hymenochirus* are pipids, but differ in<br>their diet. *Xenopus* tadpoles are filter feeders whereas<br>thei their diet. Senopus below the eveloping into juvenile rrogs.<br>
Xenopus and Hymenochirus are pipids, but differ in<br>
their diet. Xenopus tadpoles are filter feeders whereas<br>
Hymenochirus tadpoles are carnivorous and eat mainl Actiopus and Trymenochirus are pipids, but differ mediator of the initial presence of muscle<br>their diet *are operation* and eat mainly and eat mainly protein. All embryos were preserved in Dent fixative<br>copepods, mollusc l dellary the control of the methods are there is whereas<br> *Hymenochirus* tadpoles are carnivorous and eat mainly<br> *copepods, mollusc larvae, and in the laboratory, brine*<br> *copepods, mollusc larvae, and in the laboratory, b Hymenochirus* tadpoles are carnivorous and eat mainly<br>copepods, mollusc larvae, and in the laboratory, brine<br>shrimp. Rana tadpoles were chosen as detritivors or<br>grazers, grazing on algae by scratching rocks with<br>tibody 12 copepods, mollusc larvae, and in the laboratory, brine<br>shrimp. Rana tadpoles were chosen as detritivors or<br>grazers, grazing on algae by scratching rocks with<br>their horny beaks, and, because they are commonly ognizes a spec shrimp. Kana tadpoles were chosen as detritivors or<br>grazers, grazing on algae by scratching rocks with<br>their horny beaks, and, because they are commonly<br>studied, were readily available. The direct developer, cle (Kintner grazers, grazing on aigae by scratching rocks with<br>their horny beaks, and, because they are commonly by a specific antigen in amphibian skeletal mus-<br>studied, were readily available. The direct developer, cle (Kintner and ency of the set of the discussed and the set of the disculied, were readily available. The direct developer, the tained from the Developmental Studies Hybri studied, were readily available. The direct developer,<br>Eleutherodactylus coqui, lays arboreal eggs that lack the tained from the Developmental Studies Hybridoma<br>intermediate free-swimming tadpole stage. Instead, Bank, Univ Eleutherodactylus coqui, lays arboreal eggs that lack the<br>intermediate free-swimming tadpole stage. Instead,<br>embryos are positioned on top of a yolk sac from<br>which they obtain nourishment throughout develop-<br>Wit (biotin-av merineum enters winning taupole stage. Instead,<br>explored to the vector of a yolk sac from<br>which they obtain nourishment throughout develop-<br>which they obtain nourishment throughout develop-<br>ment and hatch directly into fro embryos are positioned on top or a yolk sac from<br>which they obtain nourishment throughout develop-<br>ment and hatch directly into froglets. The "interme-<br>ment and hatch directly into froglets. The "interme-<br>diate developer," which diey obtain hourismment unoughout develop-<br>ment and hatded embryos were cleared with<br>diate developer," Agalychnis callidryas, has a reproduc-<br>tive mode intermediate between those previously de-<br>Whole embryos and hist ment and nature directly into froglets. The intermeduce one part benzyl alcohol: two parts benzyl benzoate.<br>diate developer," Agalychnis callidryas, has a reproduction part benzyl alcohol: two parts benzyl benzoate.<br>tive m diate developer," Agalychnis callidryas, has a reproductional conterpart betizy in altohol: two parts benzyl benzoate.<br>tive mode intermediate between those previously de-<br>scribed. Eggs are arboreal, and intracapsular devel tive mode intermediate between those previously de-<br>scribed. Eggs are arboreal, and intracapsular devel-<br>opment is extended beyond the embryo stage into<br>opment is extended beyond the embryo stage into<br>what are, morphologic scribed. Eggs are arboreal, and intracapsular devel-<br>opment is extended beyond the embryo stage into Correlation of Developmental Tables.—Our analysis<br>what are, morphologically, early tadpole stages. Upon compares the stag opment is extended beyond the embryo stage into<br>what are, morphologically, early tadpole stages. Upon<br>hatching, they drop into the water below and become skeletal muscle myogenic markers appear. Unfortu-<br>free-swimming tadp what are, interprotogramy, early tadpole stages. Upon<br>hatching, they drop into the water below and become<br>free-swimming tadpoles. Nourishment during intra-<br>nately, at least three different tables of normal devel-<br>capsular natching, they drop into the water below and become<br>free-swimming tadpoles. Nourishment during intra-<br>capsular development is provided by yolk reserves, opment are commonly used to gauge developmental<br>whereas the free-swim

cussed below and a summary given in Table 1. The mation, first neartbeat, gill circulation and gill expantitilized Agalychnis eggs. Adult Eleutherodactylus were is summarized in Table 1.<br>tilized Agalychnis eggs. Adult Eleu collected in the wild in Puerto Rico and bred in the laboratory (non-hormone induced). Agalychnis and Eleutherodactylus embryos were raised at 22 C in ster ilized culture dishes on sterile filter paper (moistened

 with 10% HEPES-buffered Steinberg solution, Peng, 1991). Hymenochirus adults were purchased from Blue Lobster Farm, Madeira, CA; embryos were raised from clutches obtained in laboratory aquaria (non-hormone induced breeding). Eggs were removed and trans ferred to sterilized disposable culture dishes, and em bryos raised at 22-24 C. Rana sylvatica eggs and em bryos were collected at Fort AP Hill, VA. Fertilized eggs of Rana utricularia were purchased from Charles D. Sullivan Co. (Nashville, TN). Both Rana species were raised in bowls of dechlorinated tap water at 22- 24 C. Xenopus embryos were obtained from hormon ally-induced breeding. Clutches were raised in 10% Steinberg solution at 22-24 C in sterile dishes. Exper iments were conducted in accordance with approved Institutional Animal Care and Use guidelines (98-2).

in glycol methacrylate (JB4+ embedding kit, Polysci-<br>Hymenochirus boettgeri, Rana sylvatica, R. utricularia, ences, Inc). Axial muscle was sectioned longitudinally<br>Xenopus laevis, and X. tropicalis, lay large clutches of a Hymenochirus boettgeri, Rana sylvatica, R. utricularia, ences, Inc). Axial muscle was sectioned longitudinally<br>Xenopus laevis, and X. tropicalis, lay large clutches of at 2 µm using glass knives, transferred to a slide, an Observations of first twitch and immunohistochem ical staining were made using a Nikon stereomicro scope. First stimulated twitch of muscle was examined by poking live specimens with metal probes. Obser vations of multinucleation in axial myotomes were made using a Nikon Optiphot microscope. In order to observe muscle cell nuclei, specimens were embedded in glycol methacrylate (JB4+ embedding kit, Polysci ences, Inc). Axial muscle was sectioned longitudinally at 2  $\mu$ m using glass knives, transferred to a slide, and stained for 10 sec with 0.1% toluidine blue in 1% so dium tetraborate (Dawes, 1979).

> Whole-mount immunohistochemical staining was performed to identify the initial presence of muscle protein. All embryos were preserved in Dent fixative prior to staining. The procedure was adapted from Hanken et al. (1992, 1997) and used monoclonal an tibody 12/101, a muscle-specific antibody that rec ognizes a specific antigen in amphibian skeletal mus cle (Kintner and Brockes, 1984). Antibody was ob tained from the Developmental Studies Hybridoma Bank, University of Iowa. The 12/101 primary anti body was visualized using the Vectastain Universal Kit (biotin-avidin complex) and diaminobenzidine (DAB) substrate. Stained embryos were cleared with one part benzyl alcohol: two parts benzyl benzoate. Whole embryos and histological sections were pho tographed with Kodak Technical Pan film.

Examplement is provided by volk reserves. The commonly used to gauge developmental<br>whereas the free-swimming tadpoles are filter feeders. age in the species we studied. We therefore needed to<br>A total of 454 specimens were psular development is provided by york reserves,<br>hereas the free-swimming tadpoles are filter feeders. age in the species we studied. We therefore needed to<br>A total of 454 specimens were examined (27  $Aga$ -correlate the di A total of 454 specimens were examined (27 Aga-<br>lychnis callydrias, 36 Eleutherodactylus coqui, 86 Hymen-<br>ochirus boetteeri. 60 Rana sulvatica. 35 R. utricularia. 120 stages. To do this we accepted previous partial stage Fried the Cosnet set continued that the distribution of the cosnet (1960)<br>dehinis callydrias, 36 Eleutherodactylus coqui, 86 Hymen-<br>ochirus boettgeri, 60 Rana sylvatica, 35 R. utricularia, 120 stages. To do this we accept nychmis callyarias, 36 Eleutheroadcrylus coqui, 86 Hymen-<br>ochirus boettgeri, 60 Rana sylvatica, 35 R. utricularia, 120 stages. To do this we accepted previous partial stage<br>Xenopus laevis, and 90 X. tropicalis). Specimens behinds boeliger), ob Kanu Sylolinda, 35 K. alimbularia, 120<br> *Xenopas lacevis*, and 90 X. tropicalis). Specimens were correlations by Just et al. (1981) and Townsend and<br>
preserved in either Dent fixative (1 part dimethy *Xenopus laevis,* and 90 *X. tropicalis*). Specimens were correlations by just et al. (1981) and lownsend and<br>preserved in either Dent fixative (1 part dimethyl sulf-<br>oxide: 4 parts methanol; Dent et al., 1989) or 4% neu-g preserved in either Dent fixative (1 part dimethyl sulf-<br>oxide: 4 parts methanol; Dent et al., 1989) or 4% neu-<br>gaps during the stages of skeletal muscle myogenesis<br>tral-buffered formaldehyde. Species were staged fol-<br>by n oxide: 4 parts memanoi; Dent et al., 1989) or 4% neu-<br>tral-buffered formaldehyde. Species were staged fol-<br>lowing, or correlating with, the Gosner (1960) table of characters that we presume are independent of skele-<br>normal tral-buffered formaldehyde. Species were staged fol-<br>lowing, or correlating with, the Gosner (1960) table of characters that we presume are independent of skele-<br>normal development. Staging correlations are dis-<br>cussed bel lowing, or correlating with, the Gosner (1960) table of characters that we presume are independent of skelenormal development. Staging correlations are dis-<br>normal development. Staging correlations are dis-<br>cussed below an Final development. Staging correlations are dis-<br>seed below and a summary given in Table 1. The Nation, first heartbeat, gill circulation and gill expan-<br>The National Aquarium in Baltimore provided fer- sion, and limb deve Correlation of Developmental Tables.--Our analysis compares the stages during development at which skeletal muscle myogenic markers appear. Unfortu nately, at least three different tables of normal devel opment are commonly used to gauge developmental age in the species we studied. We therefore needed to correlate the different normal tables to a standard, which we arbitrarily chose to be the Gosner (1960) stages. To do this we accepted previous partial stage correlations by Just et al. (1981) and Townsend and Stuart (1985). We then extended these and filled in gaps during the stages of skeletal muscle myogenesis by normalizing to the appearance of developmental characters that we presume are independent of skele tal muscle development, including neural tube for mation, first heartbeat, gill circulation and gill expan sion, and limb development. Our complete correlation is summarized in Table 1.

 Townsend and Stewart (1985) provided correlations of their Eleutherodactylus coqui stages from Gosner Stages 1-18. We extended the correlation for TS Stages 5 to Gosner Stages 19/20 based on initial heart beat



 FIG. 1. Immunohistochemical detection of muscle specific antigen expression. Whole embyros were re acted with anti-muscle antibody 12/101. Binding was detected with a peroxidase-linked secondary antibody and detected with diaminobenzidine. In some embry os this procedure stained the central nervous system in addition to skeletal muscle. Arrows indicate areas of specific muscle staining. Scale bars, 0.5 mm. (A.) Lateral view of Rana utriucularia at Gosner stage 17. Anterior is to the left. Antigen is detected in anterior myotome but not posterior unsegmented mesoderm. The staining pattern of Rana sylvatica is similar (not shown) (B.) Xenopus tropicalis, Nieuwkoop and Faber stage 20 (dorsal view, left) and 22 (lateral view, right). Anterior is toward the top. Note that embryos of this species are much smaller than the other anurans shown. (C.) Ventro-lateral view of Eleutherodactylus co qui, Gosner stage 5. Staining is just detectable above background in anterior myotome. (D.) Agalychnis cal lidryas, Gosner stage 5. Arrows indicate myotome staining.

 and beginning of gill circulation. TS stage 7 is distin guished by the first appearance of foot paddles, which can be directly correlated with Gosner stage 31. Con sequently, Gosner stages 21-30 were correlated with TS Stage 6, which mainly involves elongation of the limbs (Townsend and Stewart, 1985).

12/101 Protein (Fig. 1 and Table 2).-The initial pres-

 ence of axial muscle protein was detected in somites of both Xenopus species by Stage 16. It was observed slightly later, at early Stage 17, in both Rana species, and at Stage 18/19, in Hymenochirus. The protein was detected at Stage 19/20 in Eleutherodactylus, whereas in Agalychnis it was found at Stage 17/18. We did not section the specimens to confirm that staining oc curred in muscle. However, the antibody is known to recognize a muscle specific protein (Kintner and  $b$ rockes, 1984) and we could easily visualize entire somites stained in the cleared specimens. Since the bulk of amphibians somites is myotome, with only small contributions from sclerotome and dermatome (see Keller, 1999, for review), we are confident that the staining we observed represents expression in myo blasts or muscle cells. Note also that our results for the stage of first expression depend on the sensitivity of whole mount immunocytology. More sensitive methods could detect earlier expression. Hence they should be considered the latest stages at which muscle protein first appears.

Stimulated Twitch (Table 2).-First stimulated twitch of axial muscle occurred by Stage 17 in X. laevis and slightly later in X. tropicalis, Stage 17/18. It was ob served at Stage 18 in Rana species, Stage 18/19 in Hy menochirus, Stage 19/20 in Eleutherodactylus, and at Stage 18/19 in Agalychnis.

Spontaneous Twitch (Table 2).-The first spontaneous twitch of axial muscle was observed at Stage 18/19 in Xenopus species, Stage 18 in Rana species, Stage 18/ 19 in Hymenochirus, between Stages 21-30 in Eleuth erodactylus, and at Stage 18/19 in Agalychnis.

Multinucleation (Table 2).-Multinucleated axial myo tome was initially detected at Stage 26/27 in X. tro picalis and at Stage 26 in X. laevis. It was observed at Stage 17 in Rana species. Similar to the other pipids, Hymenochirus multinucleation was present by Stage 26. Multinucleation was observed at Stage 19/20 in Eleutherodactylus and at Stage 18 in Agalychnis.

 Hymenochirus boettgeri, Xenopus laevis, and X. tropi calis were originally staged according to the Nieuwk oop and Faber (1975) normal table of development for X. laevis. Just et al. (1981) and Trueb and Hanken (1992) provided a correlation for NF stages 46-47 with Gosner stages 26-27. We determined the correlation between NF stages 18-45 and Gosner stages 15-25. NF stage 18 was correlated with Gosner Stage 15 based on neural groove formation. Neural tube devel opment linked NF Stages 19-21 with Gosner Stage 16. NF Stages 22-23 were correlated with Gosner Stage

<b>Species</b>	Gosner stage when first detected			
	12/101 protein expression	Stimulated twitch	Spontaneous twitch	Multinucleation
E. coqui	$19 - 20$	$19 - 20$	$21 - 30$	$19 - 20$
H. boettgeri	18/19	18/19	18/19	26
X. tropicalis	16	17/18	18/19	26/27
X. laevis	16	17	18/19	26
R. sylvatica	16/17	18	18	17
R. utricularia	17	18	18	17
A. callidryas	17–18	$18 - 19$	18/19	18

TABLE 2. Comparative stages of myogenesis of several anuran species.



 FIG. 2. Myofiber multinucleation in several anurans. A-D represent frontal sections through the myotome, oriented so that the medial surface (facing the notochord) is towards the bottom of the panel and the lateral aspect toward the top. Portions of two myotomes are shown in each panel. Individual myofibers extend the length of a myotome. Examples of multiple nuclei within a single myofiber are marked arrows. (A) Rana sylvatica, Gosner Stage 17; (B) Xenopus tropicalis Gosner St. 26/27; (C) Agalychnis callidryas Gosner St 18; (D) Eleutherodactylus coqui Gosner St.21-30. A-D were photographed at the same magnification. Bar, 50  $\mu$ m.

 17; these stages involve the early formation and ap pearance of the tail bud. NF stage 33/34 is character ized by first heartbeat, which is observed at Gosner Stage 19. We found no direct correlation between NF stages 25-32 and any given Gosner stage. Consequent ly, NF Stages 25-32 would correspond to the transi tional period between Gosner Stages 18 and 19. Gill circulation links NF Stage 40 and Gosner Stage 20; thus, NF Stages 35-39 correspond to the period be tween Gosner Stages 19 and 20. NF Stage 41, at which the gills become "broader and flatter" (Nieuwkoop and Faber, 1975) corresponds to Gosner Stages 21-23 which involve further development of the gills. NF Stage 42 correlates with Gosner Stage 24 based on the initial development of the operculum. Further oper culum development linked NF Stages 43-45 with Gos ner Stage 25 just before the initial presence of the hin dlimb bud. Initial limb bud development directly cor relates NF Stage 46 with Gosner stage 26.

 Having correlated normal stages we then compared the stage and sequence in which common skeletal muscle myogenic events occurred (Table 2). Then we  determined whether a particular pattern of myogen esis corresponded to a life. history mode. We selected Rana, with free-swimming and completely aquatic tadpoles, as the standard reproductive mode to com pare myogenic events among species because this ge nus has been widely studied and was readily avail able. The initial presence of muscle protein, as judged by whole-mount immunocytology, was detected at Gosner Stage 17 in Rana. The appearance of muscle protein in other non-direct developing larvae exam ined differed. It was detected earlier, Stage 16, in Xen opus laevis and X. tropicalis, whereas it was not found until later in Hymenochirus, Stage 18/19. In Eleuthero dactylus synthesis of muscle protein appears delayed and was first detected at least two stages later than in Rana. Multinucleation precedes the first twitch of axial muscle in Rana; however, Hymenochirus and Xenopus exhibit functional, but mononucleated, axial myotome at a point remarkably earlier than the onset of multin ucleation. In Agalychnis and Eleutherodactylus multinu cleation is also slightly delayed, but it occurs at about  the same time as the activation of axial muscle (first twitch).

 Overall, with the exception of Hymenochirus, the non-direct developing taxa have an earlier expression of muscle protein and muscle function than species representing other reproductive modes. The myogenic pattern of Agalychnis is delayed in its entirety relative to Rana, but it exhibits muscle protein expression, muscle function, and multinucleation earlier than in Eleutherodactylus. Initially, we ranked Agalychnis as an intermediate developer, based on its extended intra capsular development. The present analysis shows that the myogenic pattern of Agalychnis is intermediate between that of non-direct and direct developing an urans. Myogenesis in Agalychnis occurs faster than in Eleutherodactylus, but it is generally slower relative to Rana and pipids.

 The present study suggests that myogenic events vary with reproductive modes. Furthermore, it sug gests a progressive delay of myogenesis associated with a delay in hatching, i.e., extended intracapsular development. If this is correct, we would predict that myogenic events in other anuran taxa such as Dendro bates and centrolenids would resemble the pattern de scribed for Agalychnis whereas myogenic events in Co phixalus, Ceratobratrachus, and some Gastrotheca, (all di rect developers) would resemble the pattern of Eleuth erodactylus.

 Although life history may correlate with patterns of myogenesis, some myogenic events may be better un derstood in the light of evolutionary relationships. For example, among the anurans with free swimming tad pole, the pipid taxa studied showed delayed multin ucleation of axial muscle, delayed further than it is in Eleutherodactylus. However, within pipids Hymenochi rus differs from X. laevis and X. tropicalis in its delayed expression of muscle protein and first twitch. The de parture of Hymenochirus from the pattern of myogen esis found in other pipids is not surprising since tad poles of this taxon have been shown to differ in the characteristics and development of other musculoskel etal structures (e.g., Sokol, 1959, 1962, 1977; de Sá and Swart, 1999). Moreover, multinucleation occurs in pip ids after first twitch, whereas in the other species mul tinucleation precedes or coincides with muscle func tion. Hence part of the myogenic sequence is altered in pipids relative to other anurans studied thus far. Thus, multinucleation following function may be a synapomorphy for the pipidae. If so, then it should also occur in the genus Pipa. The genus Pipa will be particularly interesting to examine because there are both free swimming tadpole (e.g., P. carvalhoi) and direct developing  $(P. pipa)$  members of the genus. This could allow one to determine more directly whether myogenic patterns are related to life history rather than historical events.

Acknowledgments.-We are grateful to the Departa mento de Recursos Naturales y Ambientales, Puerto Rico (DRNA 96-186 to RDS), and to the Virginia De partment of Game and Inland Fisheries (8372 to JM) for providing collecting permits. The National Aquar ium in Baltimore generously provided specimens. Funding for this work was provided by NSF Grant BIR 9510228.

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Accepted: 27 July 2000.