Monophyly of Heterandriini (Teleostei: Poeciliidae) revisited: 
a critical review of the data

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The systematics and taxonomy of poeciliid fishes (guppies and allies) remain poorly understood despite the relative importance of these species as model systems in the biological sciences. This study focuses on testing the monophyly of the nominal poeciliine tribe Heterandriini and the genus *Heterandria*, through examination of the morphological characters on which the current classification is based. These characters include aspects of body shape (morphometrics), scale and fin-ray counts (meristics), pigmentation, the cephalic laterosensory system, and osteological features of the neurocranium, oral jaws and suspensorium, branchial basket, pectoral girdle, and the gonopodium and its supports. A Maximum Parsimony analysis was conducted of 150 characters coded for 56 poeciliid and outgroup species, including 22 of 45 heterandriin species (from the accounted in Parenti & Rauchenberger, 1989), or seven of nine heterandriin species (from the accounted in Lucinda & Reis, 2005). Multistate characters were analyzed as both unordered and ordered, and iterative *a posteriori* weighting was used to improve tree resolution. Tree topologies obtained from these analyses support the monophyly of the Middle American species of “*Heterandria*,” which based on available phylogenetic information, are herein reassigned to the genus *Pseudoxiphophorus*. None of the characters used in previous studies to characterize the nominal taxon Heterandriini are found to be unambiguously diagnostic. Some of these characters are shared with species in other poeciliid tribes, and others are reversed within the Heterandriini. These results support the hypothesis that *Pseudoxiphophorus* is monophyletic, and that this clade is not the closest relative of *H. formosa* (the type species) from southeastern North America. Available morphological data are not sufficient to assess the phylogenetic relationships of *H. formosa* with respect to other members of the Heterandriini. The results further suggest that most tribe-level taxa of the Poeciliinae are not monophyletic, and that further work remains to resolve the evolutionary relationships of this group.

La sistemática y taxonomía de poeciliídos (gupys, platys, mollys y espadas) continúa siendo poco conocida, a pesar de su importancia como modelos experimentales en ciencias biológicas. Este estudio busca probar la monofilia de la tribu nominal Heterandriini (Poeciliinae) y del género *Heterandria*, mediante el estudio de los caracteres morfológicos en los que se basa la clasificación actual. Estos caracteres incluyen aspectos como la forma del cuerpo (morfometría), conteos de escalas y radios de las aletas (merísticos), pigmentación, sistema sensorial cefálico y características osteológicas del neurocráneo, mandíbula y sus suspersorios, canasta branquial, cintura escapular y del gonopodio y sus soportes. Se condujo un análisis de Maxima Parsimonia con 150 caracteres codificados para 56 poeciliídos y especies del grupo externo, incluyendo 22 de 45 especies de Heterandriini (según el conteo de Parenti & Rauchenberger, 1989), o siete de nueve especies (según el conteo de Lucinda & Reis, 2005). Los caracteres con estados múltiples fueron analizando de manera ordenada y no ordenada seguido de una ponderación iterativa *a posteriori* para mejorar la resolución de los árboles. Las topologías de los árboles obtenidos de estos análisis apoyan la monofilia de las especies mesoamericanas de “*Heterandria*”, que de acuerdo a la información filogenética disponible, son aquí reasignados al género *Pseudoxiphophorus*. Ninguno de los caracteres usados en estudios previos para caracterizar el taxón nominal Heterandriini fue encontrado de forma inequívoca diagnóstico. Algunos de estos caracteres son compartidos con especies de otras tribus de poeciliídos y otros han revertido entre Heterandriini. Estos resultados apoyan la hipótesis de que *Pseudoxiphophorus* es monofilético, y que este grupo no es grupo cercano de *H. formosa* (la especie tipo) del suroeste de Norte América. La información morfológica disponible no es suficiente para evaluar las relaciones de *H. formosa* con respecto a otros grupos de Heterandriini. Los resultados de este estudio sugieren que la mayoría de taxones a nivel de tribu de Poeciliinae no son monofiléticos, y que aún queda trabajo para resolver las relaciones evolucionarias de este grupo.

**Key words:** Fish diversity, *Heterandria*, Ichthyology, Osteology, *Pseudoxiphophorus*, Systematics.
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Introduction

The Poeciliidae (livebearers and relatives; sensu Ghedotti, 2000) is a clade of small-bodied (20–200 mm standard length) cyprinodontiform fishes with about 317 species currently recognized in 30 genera. Poeciliids are characterized by a suite of derived characters including the position of the pectoral fins high on side of the body, an anterior placement of the pelvic fins ventral to the pectoral fins, the presence of pleural ribs on the first several haemal arches, a bony cap on the ventral hypohyal lying over the anterior facet of the anterior ceratohyal, and supraorbital laterosensory pores with neuromasts embedded in fleshy grooves (Ghedotti, 2000). Despite the widespread use of poeciliid fishes in ecological and evolutionary studies (e.g., Rauchenberger, 1990; Meyer et al., 1994; Borrowsky et al., 1995; Marcus & McCune, 1999; Spencer et al., 1999; Spencer et al., 2000; Morris et al., 2001; Kallman et al., 2004; Gutierrez-Rodriguez et al., 2007a; 2007b; Reznick et al., 2007; Leborg & Firmin, 2008; Purcell et al., 2008; Martin et al., 2009; Pollux et al., 2009; Meredith et al., 2010; Pires et al., 2010; Albert & Johnson, 2011), the phylogenetic interrelationships among poeciliid species remain incompletely resolved.

The family Poeciliidae comprises three subfamilies that are restricted to fresh and brackish continental waters: Poeciliinae, Procatopodinae, and Aplocheilichthyinae (Ghedotti, 2000; Lucinda, 2003; Hrbek et al., 2007). The Poeciliinae includes at least 228 species distributed across much of the tropical and subtropical portions of the Americas, from the La Plata estuary of northern Argentina to southeastern United States, with species richness reaching a zenith in Middle America and the West Indies (Rosen & Bailey, 1963; Lucinda, 2003; Hrbek et al., 2007; Albert et al., 2011). The Procatopodinae includes at least 78 species in the humid tropical regions of South America (i.e. Fluviphylax) and Africa (Lucinda, 2003), and the Aplocheilichthyinae is represented by about 11 species of Aplocheilichthys from humid tropical regions of Africa (Huber, 1999).

All poeciliine species have internal fertilization achieved by means of an intromittent organ (gonopodium) in males, which is a modification of anal-fin rays 3–5. Poeciliine species bear their young as free-swimming juveniles (without a yolk sac), and exhibit either ovoviviparity (eggs hatch within the uterine cavity), or viviparity (maternal provisioning to the developing larvae while in the uterine cavity). The ovoviviparity condition is thought to be derived within the subfamily (Pollux et al., 2009). One poeciliine species, Tomeurus gracilis from northeastern South America, is facultatively ovoviviparous, releasing eggs or live young depending on environmental conditions (Breder & Rosen, 1966; Wourms, 1981). The Poeciliinae is also characterized by several salient morphological features in addition to the gonopodium; expansion of the fourth epibranchial, absence of exoccipital condyles, and neural arches of the first vertebra open (Rosen & Bailey, 1963; Parenti & Rauchenberger, 1989; Ghedotti, 2000).

The name Heterandriini was introduced by Hubbs (1924) for a group of poeciliins in which the third ray of the gonopodium lacks a pair of curved horn-like projections, and in which the fifth ray always is smooth on the posterior edge. This group originally included nine genera; Allogambusia (now Priapichthys), Alloheterandria (now Priapichthys and Gambusia), Darienichthys (now Priapichthys), Heterandria, Neoheterandria, Panamichthys (now Priapichthys), Priapichthys, Pseudopoecilia, and Pseudoxiphophorus (treated as a subgenus by Rosen, 1979). Hubbs (1926) published diagnoses to identify each of these genera. According to Hubbs (1926), species of the tribe Heterandriini evolved in Central America and northwestern (trans-Andean) South America.

Rosen & Bailey (1963) organized the Poeciliidae into five subfamilies, nine tribes, and 52 genera, primarily on the basis of similarities in the gonopodium and gonopodial suspensorium. They included six genera in the tribe Heterandriini; Heterandria, Neoheterandria, Phallichthys Poeciliopsis (which was divided into the subgenera Poeciliopsis and Alouphallus), and Priapichthys. The diagnostic characters of Heterandriini sensu Rosen & Bailey (1963) were also largely taken from the gonopodal complex; i.e., gonopodal length more than one-third standard length, presence of three or more gonapophyses, moderately or well-developed ligastyle, and two to four plate-like gonactinoids. However, these traits are shared with species of other nominal poeciliine tribes, and are not in fact unique to species of the Heterandriini.

Several subsequent phylogenetic studies based on morphological (Rodriguez, 1997; Ghedotti, 2000; Lucinda & Reis, 2005) and molecular (Hrbek et al., 2007) data suggest that some of the tribes proposed by Rosen & Bailey (1963) are not monophyletic. These papers also show that some genera not previously placed in the Heterandriini (i.e., Allofar, Brachyrhaphis, Pseudopoecilia, Neoheterandria) are more closely related to Heterandria formosa Girard (1859), the type species of the genus, than to the Middle American species of Heterandria which were previously assigned to Pseudoxiphophorus (Heckel, 1848). Despite these concerns, the pre-cladistic classification of Rosen & Bailey (1963) remains the most recent comprehensive morphological revision and taxonomic authority for Poeciliidae (Lucinda, 2003; Hrbek et al., 2007). The Poeciliidae of Rosen and Bailey (1963) is equivalent to the Poeciliinae of Parenti (1981).

This study focuses on the systematics of the nominal poeciliine taxon Heterandriini (sensu Lucinda & Reis, 2005) and the type genus Heterandria, using osteological and other morphological characters (morphometrics, meristics, pigmentation, and cephalic sensory system). As currently recognized, the Heterandriini includes five genera (Priapichthys, Neoheterandria, Heterandria, Poeciliopsis, and Phallichthys) with more than 60 species (Parenti & Rauchenberger, 1989; Ghedotti, 2000). The goal of this study is to test the monophyly of the taxa Heterandriini and Heterandria by evaluating all the characters used historically in the higher classification of the group within a formal phylogenetic context.
Material and Methods

A total of 56 species represented by 126 museum lots with more than 5,000 specimens were examined (Appendix 1). Museum abbreviations follow Leviton et al. (1985), and can be found at Fricke & Eschmeyer (2011) except USAC (Museo de Historia Natural de Escuela de Biologia, Universidad de San Carlos de Guatemala). The current definition of the Poeciliinae follows Parenti & Rauchenberger (1989), which is equivalent to the Poeciliidae of Rosen & Bailey (1963). Terminals were selected according to the list of species and previous taxonomic information for this group (see Rosen, 1979; Parenti, 1981; Greenfield, 1985; Rodriguez, 1997; Costa, 1998; Ghedotti, 1998, 2000; Lucinda & Reis, 2005; Hrbek et al., 2007) and available material in museums.

External characters, morphometrics, and meristics were examined from museum specimens of 66 species preserved in alcohol. Meristics and morphometrics followed the procedure outlined by Miller (1948) for cyprinodont fishes. Counts of scales in the lateral series follow Parenti (1981). All meristics and morphometrics were taken form the left side of specimens when possible using an Olympus SZX-12 dissecting microscope with 6-inch LCD digital caliper (± 0.1 mm error).

Clearing and staining for bones and cartilages followed the procedure of Taylor & van Dyke (1985) with some modifications to account for the small size of the specimens. Oxygen peroxide and xylene were not used to avoid decalcification. Specimens were transferred directly from ethyl alcohol to Alcian Blue to maximize uptake of the hydrophobic stain. The visceral cartilages were removed after staining with Alcian Blue because the small cartilages around the orbit are more apparent and the damage to the specimens is reduced (Reis, pers. comm., 2009). Prior to digestion in 30% trypsin solution (in sodium borate [NaBO]) specimens were treated one hour in 30% NaBO.

Dissections of cleared-and-stained fishes were made using microdissection tools following the method outlined by Weitzman (1974), and Ghedotti’s (2000) modification of removing the branchial basket before the suspensorium. Bones were disarticulated to functional groups (e.g., neurocranium, suspensorium, pectoral girdle), or into individual elements (e.g., maxilla). Specimens from a total of 56 species were dissected and coded for phylogenetic analysis. Outlines and standardized features of individual bones or functional groups were traced from lateral and medial aspects with the aid of a camera lucida mounted on an Olympus SZX-12 dissecting microscope. Images were digitized using an HP Scanjet 5470c (2400dpi, 48-bit color) scanner and edited using Adobe Photoshop 7.0.

Descriptions of 150 characters and their alternative states are provided with characters organized by functional group (Appendix 2). Character states were polarized using conditions observed in outgroup taxa. Question marks were used to indicate when a character state could not be coded due to a lack of available specimens, or when the coding was non-sensical; e.g., state of the ligastyle (a free ossified haemal spine dorsal to the gonopodial suspensorium and close to the vertebral column) when a gonopodium is not present in the species. Whenever possible characters were coded for states observed in multiple specimens (Albert & Crampton, 2005). Due to a lack of male specimens of Quintana and Fluviphylax gonopodial characters from these taxa were coded from literature. In most species, specimens of both sexes were examined, but in sexually dimorphic species only states observed in males were coded.


Microsoft Excel and Mesquite v. 2.72 ( Maddison & Maddison, 2009) software packages were used to assemble the data matrix of 56 taxa and 150 morphological characters. Characters were polarized using states in multiple outgroups, and character-state distributions were optimized in MacClade v. 4.0 PPC (Maddison & Maddison, 1993). Poeciliid outgroups were selected from the results of previous studies, and included representative Anablepidae (Jenynsia eirmostigma, Anableps dowi), Profundulidae ( Profundulus guatemalensis), and Aplocheilichthyinae ( Fluviphylax pygmaeus); (Parenti & Rauchenberger, 1989; Ghedotti, 2000; Lucinda & Reis, 2005; Lucinda et al., 2006). Poeciliid taxa included 52 species in 21 genera, representing 22 of 45 species of Heterandriini sensu Parenti & Rauchenberger (1989), which is equivalent to seven of nine species of Heterandriini sensu Lucinda & Reis (2005).

A Maximum Parsimony (MP) analysis was conducted to find tree topologies that most economically summarized the data matrix; e.g., maximized similarities as synapomorphies and minimized them as homoplasies (Ane & Sanderson, 2005; Campo et al., 2007; Albert, 2009). A heuristic search with the Tree-Bisection-Reconnection (TBR) algorithm was performed using PAUP* v. 4.0 b10 (Swofford, 1993, 2002). Multistate characters were analyzed as both unordered and ordered. Polytomies were resolved using a posteriori weighting on the rescaled consistency index (Farris, 1989; Albert, 2001; Kluge, 2001; Bryant, 2003; Kuklami & Moret, 2005). Bremer support (decay analysis) was used to assess node support (Bremer, 1988). Bremer support values were calculated using the reversed-constraint method (Sorensen, 1999) implemented in NONA v.2 (Goloboff, 1999a, 1999b; Goloboff et al., 2008) and Winclada v. 1.00.08 (Nixon, 2002) using the bremer.run script.

Results

These phylogenetic results are shown with multistate characters analyzed ordered and unordered, and used iterative weighting on the RC to resolve polytomies. The strict consensus tree of 12 MP trees based on 150 unordered characters is given

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A single MP tree was obtained from analysis of this same dataset with all multistate characters treated as unordered and followed by \textit{a posteriori} weighting based on the RC (Fig. 2; 1444 steps, CI = 0.20, RI = 0.47, RC = 0.09). A single MP tree with a slightly different topology (Fig. 3) was obtained when the multistate characters were treated as ordered (1446 steps, CI = 0.20, RI = 0.47, RC = 0.09). The strict consensus tree with all characters weighted equally, and with all multistate characters treated as unordered, is more-well resolved for the tribes Cnesterodontini, Scolichthyini, Gambusiini, and Alfarini (Fig. 1).

Bremer decay analysis showed that \textit{Poeciliopsis} is well supported even though there was incongruence and lack of resolution within that clade. According to this study, several clades recognized in previous literature are well supported. These clades are: 1. \textit{Gambusia} + \textit{Belonesox} (Lucinda & Reis, 2005); 2. \textit{Gambusia} + \textit{Belonesox} + \textit{Brachyrhaphis} (Mojica \textit{et al.}, 1997); 3. \textit{Scolichthys} + \textit{Neoheterandra} (Lucinda & Reis, 2005); 3. \textit{Limia} + \textit{Xiphophorus} (Rodriguez, 1997); 4. \textit{Limia} + \textit{Xiphophorus} (Rodriguez, 1997); 5. \textit{Pamphorichthys} + \textit{Micropoecilia} (Lucinda & Reis, 2005) (see Fig. 1).

This study also suggests that the characters used in previous morphological analyses of Poeciliinae do not support the monophyly of \textit{Heterandriini}. The monophyly of \textit{Heterandriini} sensu Lucinda & Reis (2005; see Table 3), including only species of the subgenus \textit{Heterandria} and \textit{Pseudoxiphophorus} was not supported by the results of this study. \textit{Heterandria formosa} (the type species) was not found most closely related to \textit{Pseudoxiphophorus} (from Mesoamerica) in any of the topologies recovered (Figs. 1-3). We therefore recommend restricting \textit{Heterandria} to \textit{H. formosa} (following Regan, 1913), and regard \textit{Pseudoxiphophorus} as the valid name for the Mesoamerican clade (following Hubbs, 1924).

**Systematic account**

**Genus Pseudoxiphophorus** Heckel, 1848

**Composition.** \textit{Pseudoxiphophorus anzueto}i (Rosen, 1979); \textit{P. bimaculatus} Heckel; \textit{P. cataractae} (Rosen, 1979); \textit{P. diremptus} (Rosen, 1979); \textit{P. jonesii} Gunther, \textit{P. litoperas} (Rosen, 1979), \textit{P. obliquus} (Rosen, 1979).

**Distribution.** From the Tamesi river, Atlantic versant of Mexico (22°13’N 97°54’W), to Marceligo Creek near Miranda (now Rosita), Tunky (now Tunki) river, Bambana-Prinzapolka basin, Atlantic versant of Nicaragua (13°52’N 84°20’W).

**Diagnosis.** Species of \textit{Pseudoxiphophorus} share the following synapomorphies: 1. Post-temporal unbranched (ch. 49-1); 2. Less than 9 dorsal-fin rays (ch. 63-0); 3. Two vertebrae with widened neural spines (ch. 66-2); 4. Ten anal-fin rays in females (ch. 78-1); 5. Subdistal spines on gonopodial ray 3 enlarged ventrally forming a slight elbow along the ventral profile of the ray (ch. 95-1); 6. Gonopodial ray 4p as long as ray 5a (ch. 99-1); 7. Two or more penultimate short segments on gonopodial ray 4a with a membranous envelope (ch. 100-1); 8. More than four subdistal short segments on gonopodial ray 4a (ch. 101-1); 9. Keel without axial projection on the posterior ventral surface of anal-fin ray 5 (ch. 105-0); 10. Dorsal-fin melanophores forming two or more rows of discrete spots and anal-fin melanophores extending distally from anal-fin base as streaks in interradial membrane (ch. 129-1); 11. Anal fin with small diffuse dots (ch. 130-2); 12. Presence of a basicaudal spot (ch. 131-1); 13. Basicaudal spot in adults partly or wholly above midlateral line and extending onto caudal-fin base (ch. 132-1); 14. Pigment of scale pockets forming a strong reticular network along the sides (ch. 134-1).

**Discussion**

**Comparisons with previous studies.** Some of the phylogenetic results reported above are consistent with the conclusions of previous studies of Poeciliinae. The phylogeny reported in the strict consensus tree (Fig. 1) supports the monophyly of the Cnesterodontini (\textit{Pseudoxiphophorus} sensu Lucinda & Reis, 2005), Gambusiini (\textit{Poecilia} sensu Rosen & Bailey, 1963), and Tomeurini (\textit{Poecilia} sensu Rosen & Bailey, 1963; Parenti & Rauchenberger, 1989; Lucinda & Reis, 2005). The monophyly of the Priapichthyini (\textit{Poecilia} sensu Lucinda & Reis, 2005) and Alfarini (\textit{Pseudoxiphophorus} sensu Ghedotti, 2000) were not well tested, as each was represented by species in a single genus.

Other results of this study are not consistent with the published taxonomy of Poeciliinae. The Cnesterodontini was not well resolved in the topologies obtained from weighting the unordered multistate characters by the RC (Fig. 2), or from analyzing the data with ordered multistate characters (Fig. 3). Those topologies recovered the Cnesterodontini within \textit{Poeciliopsis}. The Gambusiini is not monophyletic according to the topology obtained by analyzing the data as ordered-multistate characters (Fig. 3). According to that topology, \textit{Brachyrhaphis} is not most closely related to the clade \textit{Gambusia} + \textit{Belonesox}.

None of the phylogenies recovered in this study support the monophyly of the Poeciliini (\textit{Pseudoxiphophorus} anzuetoii sensu Ghedotti 2000; Lucinda & Reis 2005). In the strict consensus tree (Fig. 1), \textit{H. formosa} is sister to \textit{Quintana} and is closely related to \textit{Limia} and \textit{Xiphophorus} suggesting that these four genera should be included in Heterandriini. However, the clade of \textit{Quintana} and \textit{H. formosa} is not well supported by Bremer decay analysis. Further, male specimens of \textit{Quintana atrizona} were not available for this study, and the data matrix was coded based on published description of this monotypic genus (Hubbs, 1934; Rosen & Bailey, 1963; Lucinda & Reis, 2005). In the topology obtained from \textit{a posteriori} weighting of the unordered-multistate characters (Fig. 2), \textit{H. formosa} is most closely related to other genera of Poeciliini (\textit{Micropoecilia}, \textit{Pamphorichthys}, \textit{Limia}, and \textit{Poecilia}).

The topology obtained from analyzing the dataset as ordered-multistate characters (Fig. 3) supports \textit{H. formosa} as closely related to the Brachyrhaphini. To verify the phylogenetic relationship between \textit{Brachyrhaphis} and \textit{Heterandria} it will be necessary to examine all (or most) of
the 12 valid species of *Brachyrhaphis*. If *H. formosa* and *Brachyrhaphis* are not closely related, the whole Heterandriini could be restricted to *H. formosa* and *Brachyrhaphis* could be referred to as *Brachyrhaphini* *sensu* Lucinda & Reis (2005).

The strict consensus tree (Fig. 1) suggests that *Neo heterandria*, *Xenophallus*, and *Scolichthys* are closely related. If this relationship is true, *Neo heterandria* and *Xenophallus* might be included in the tribe *Scolichthyini* (rather than the tribe *Gambusiini* *sensu* Lucinda & Reis, 2005). This topology is supported by the analysis of unordered multistate characters with *a posteriori* weighting (Fig. 2), but not by the analysis of ordered-multistate characters (Fig. 3).

*Poeciliopsis* was found to be monophyletic in the strict consensus tree (Fig. 1), and is well supported by the Bremer decay analysis, as in Mateos *et al.* (2002). However, in the other two topologies arising from this study *Poeciliopsis* was not found to be monophyletic (Figs. 2, 3). Even though the relationships and monophyly of *Poeciliopsis* are uncertain in all of the analyses *Poeciliopsis* is not closely related to *H. formosa* and therefore should not be included in Heterandriini.

According to Lucinda & Reis (2005), *Phallicthys* should be included in the tribe *Girardinini*, with *Girardinus*, *Poeciliopsis*, *Neo heterandria*, and *Phalloptychus*. In the present study specimens of *Girardinus* and other species of *Neo heterandria* were not examined but *Poeciliopsis* and *Phallicthys* were not found to be closely related. The position of *Carlhubbsia* varied in this study, but it should be noted the species of *Carlhubbsia* used here (*C. stuarti*) differed from that (*C. kidderi*) used by Lucinda & Reis (2005).

Gonopodial characters have been proven effective in

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**Fig. 1.** Strict consensus tree of twelve MP trees with multistate characters analyzed as unordered. Each MP tree of 1,430 steps, $CI = 0.20$, $RI = 0.48$, $RC = 0.10$. Numbers indicate Bremer support values.
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Fig. 2. Single MP tree with multistate characters analyzed as unordered, and with a posteriori weighting using RC (1,444 steps, CI = 0.20, RI = 0.47, RC = 0.09).

Taxonomy of Heterandria and Heterandriini

The taxonomic history of Heterandria and Pseudoxiphophorus is complex. Girard (1859) recognized Limia formosa from the coastal plains of southeastern United States (South Carolina to Mexico) on the basis of (among other things) a stout body with low vertebral counts (17+13 = 30), small teeth in the outer row of each jaw, a relatively tall dorsal fin (having long rays) positioned over the anal fin, and a long alimentary canal (longer than body) with numerous convolutions. This species was subsequently designated as the type of Heterandria Agassiz (1853; see Rosen & Bailey, 1963; Rosen, 1979), as a distinct genus from the type species of Limia L. vittata (Guichenot, 1853), from Cuba.
Bleeker (1860) recognized as *Pseudoxiphophorus* a species previously described as *Xiphophorus bimaculatus* (Heckel, 1848) collected from the coastal plains of Vera Cruz, Mexico. *Pseudoxiphophorus* was described on the basis of having a relatively elongate body (with high vertebral counts; 18+14=32), a flat head, large conical teeth in the outer row of both jaws, many and small gill rakers, a low, long dorsal fin positioned in advance of the anal fin, and a short simple alimentary canal (less than length of body).

Regan (1913) treated *Pseudoxiphophorus* as a subgenus of *Heterandria* while Hubbs (1924) recognized *Pseudoxiphophorus* and *Heterandria* as distinct genera. Miller (1974) illustrated the gonopodia, gonopodial suspensorium, and color patterns of these two taxa, and also examined morphometric and meristic data. He concluded that the differences between the gonopodia are sufficient to recognize the two genera. Although Rosen & Bailey (1963) synonymized *Pseudoxiphophorus bimaculatus* and *Heterandria formosa* into the genus *Heterandria*, the description of new *Heterandria* species caused Rosen (1979) to recognize these taxa as two distinct subgenera. In his review of the genus *Heterandria* Rosen (1979) added more species to the taxon he referred to as the subgenus *Pseudoxiphophorus* referring to the assemblage of *Heterandria* species from Middle America, and limited the subgenus *Heterandria* to *H. formosa*. Rosen (1979) also provided a cladogram based on synapomorphies of the described species, area cladograms, and proposed the first identification key for the all species of the subgenus *Pseudoxiphophorus*.

The monophyly and taxonomic composition of *Heterandria* has subsequently remained poorly resolved. Whereas Radda
Heterandriini the face of increased taxon sampling. Despite more than a gonopodial characters have not been corroborated as species within these higher taxa. In particular, many now known to be present in different combinations among traditionally used to diagnose tribe and genus-level taxa are since Rosen & Bailey (1963), and most of the characters suggested that the Heterandriini should be limited to species of the genus Heterandria sensu Rosen & Bailey (1963).

More than twenty-seven species have been described since Rosen & Bailey (1963), and most of the characters traditionally used to diagnose tribe and genus-level taxa are now known to be present in different combinations among species within these higher taxa. In particular, many gonopodial characters have not been corroborated as phylogenetically informative at the tribe or generic-level in the face of increased taxon sampling. Despite more than a century of research, the monophyly and interrelationships of Heterandriini and Heterandria remain uncertain, and a formal taxonomic revision is necessary.

According to the present study, the monophyly of Heterandria sensu Rosen & Bailey (1963) is not supported by morphological data. However, by design, the character definitions used here were compiled from previous studies on groups other than Heterandriini (Rauchenberger, 1989; Rodriguez, 1997; Costa, 1998; Ghedotti, 1998, 2000; Lucinda & Reis, 2005). It is therefore not surprising that these characters do a relatively poor job of resolving phylogenetic relationships within the Heterandriini. However, the results of this study do show that Heterandria and Pseudoxiphophorus are not closely related, supporting the hypothesis of Regan (1913), Miller (1974), and Radding (1985) that Pseudoxiphophorus should be recognized as a generic-level taxon. The results of the present study also suggest that Heterandriini should be restricted to H. formosa and its closest relatives. The dataset arising from this study is not sufficient to address the phylogenetic relationships of H. formosa to other possible members of Heterandriini.

Langerhans et al. (2005) and Hrbek et al. (2007) hypothesized that gonopodial characters are phylogenetically plastic, having evolved multiple times in the Poeciliidae under similar genetic and environmental circumstances. A prediction of such plasticity is that species of Poeciliidae should exhibit many derived changes in the morphology of the gonopodium, a condition that has indeed been observed. Therefore, in light of the high amounts of homoplasy observed in gonopodial characters, accurate tribe-level taxonomic designations of species must await more complete phylogenetic studies at the species level.

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Monophyly of Heterandriini


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Appendix 1. Material examined included in the phylogenetic analyses.


Appendix 2. Material examined included in the outgroup analyses.

Appendix 3. Material examined included in the ingroup analyses.

**Appendix 2.** Descriptions of characters and alternative states. Characters are grouped by functional group as follows: jaws, dentary, neurocranium, suspensorium, branchial arches, pectoral fin and girdle, dorsal fin, vertebrae and ribs, caudal skeleton, anal fin and gonopodium, scales and pigmentation, external morphology, and cephalic sensory system.

**Oral Jaws**

1. **Shape of the tip of the distal arm of maxilla (Fig. 4):** (0) square, (1) round, (2) triangular.
   
   The distal arm of the maxilla present different shape at the tip, the character was considered square if it had a completely flat bottom, even if the tip was slender. If the shape of the tip was slightly or totally rounded, it was given character state 1. When the tip ended in a peak, even if it had round edges it was given character state 2.

   2. **Notch on the dorsal portion of maxilla (Costa, 1998, ch. 3):** (0) present, (1) absent.
      
      A dorsal notch was present in most of the species analyzed and was shared by most of the Poeciliinae and Profundulidae.

   3. **Width of the distal arm of maxilla (Costa 1998, ch. 6; Fig. 4):** (0) slender, (1) broad.

   4. **Length of the dorsal process of the maxilla compared with ventral process of the maxilla (Fig. 4):** (0) long, (1) same length, (2) short.
      
      This character was a modification of character 7 in Costa (1998), who proposed that the shape of the dorsal process of the maxilla was continuous or curved. In this study, the size of the dorsal process was compared with the size of the ventral process.

   5. **Length of premaxillary ascending process (Fig. 4):** (0) dorsal margin with a simple, undivided ascending process, (1) ascending process divided, posterior ramus longer, but less than twice length than anterior ramus, (2) ascending process divided, posterior and anterior rami of subequal length, (3) ascending process divided, posterior ramus more than twice length of anterior ramus.
      
      Lucinda & Reis (2005) used six character states to describe the shape of the ascending process of the premaxilla, including aspects of both shape and size; in this study each of these aspects of trait diversity is coded as a separate character (characters 4 and 5).

   6. **Shape of premaxillary ascending process (Parenti, 1981; Costa, 1998, ch. 18; Lucinda & Reis, 2005, ch. 12, figs. 5a-f; Fig. 4):** (0) triangular, (1) rounded, (2) triangular with rounded tip.
      
      Lucinda & Reis (2005) used six character states to describe the shape of the ascending process of the premaxilla, including aspects of both shape and size; in this study each of these aspects of trait diversity, is coded as a separate character (characters 4 and 5).
Fig. 4. Lateral view of premaxilla in: a) *Pseudoxiphophorus diremptus*, LSUMZ 14337; b) *Xiphophorus mayae*, LSUMZ 14336; c) *Carlhubbsia stuartii*, USAC 204; d) *Phallichthys amates*, CAS 587873; e) *Phallichthys fairweatheri*, USAC 1797; f) *Xiphophorus alvarezi*, USAC 1782; g) *Alfaro hubber*, USAC 850; h) *Profundulus labialis*, LSUMZ 14328; i) *P. guatemalensis*, USAC 769; maxilla in: j) *Pseudoxiphophorus diremptus*, LSUMZ 14337; k) *Poeciliopsis latidens*, CAS 54436; l) *Brachyrhaphis olomina*, TU 24997; m) *Xenophallus umbratilis*, TU 186285; n) *Poecilia rositae*, USAC 1822; o) *Poeciliopsis gracilis*, TU 84966; p) *Alfaro cultratus*, TU 24927; q) *Pamphorichthys araguaiensis*, MCP 40919; r) *Heterandria formosa*, TU 24002; s) *Limia melanogaster*, MCZ 258314; t) *Fluviphylax pygmaeus*, MCP 38830; u) *Profundulus guatemalensis*, USAC 769. Scale bars = 1 mm.
17. Anterior margin of frontals (Ghedotti, 1998, ch. 1; Lucinda & Reis, 2005): (0) straight (Ghedotti, 1998, figs. 2b-d), (1) extending anteriorly between the nasals (Ghedotti, 1998, fig. 2a).

Ghedotti (1998) reported extended frontals for Alfar cultratus. Lucinda & Reis (2005) reported that the frontal extends anteriorly between the nasals for Fluviphylax, Alfar, Priapichthys, Priapella (not included here), and Belonesox.

18. Parietals (Rosen, 1967; Parenti, 1981; Rauchenberger, 1989, fig. 1; Ghedotti, 2000, fig. 3; Lucinda & Reis, 2005, ch. 2): (0) long, (1) medium, (2) narrow or short, (3) absent.

Lucinda & Reis (2005) proposed three character states, reporting large parietals for Jenynsia, Alfar, Brachyrhaphis, Priapichthys, H. formosa, Gambusia, Belonesox, Neoheterandria, Scolichthys, Girardinus (not included in the study), and Xiphophorus. Ghedotti (2000) reported reduced parietals for Tomeurus, Phallicthys, Poecilia, Girardinus, and Phallichthys, and the absence of parietals in Cnesterodon, Poeciliopsis, and Phallotorynus. Lucinda & Reis (2005) reported this absence condition for Pseudopoecilia, Phallopichthys, Xenodexia, Pamphorichthys, Micropoecilia, and Phallotorynus. Parenti (1981) reported absence of parietals as synapomorphies among cyprinodontines and other species such as Fluviphylax.

19. Epiotic processes in adults (Rosen, 1967; Rauchenberger, 1989; Ghedotti, 1998, ch. 5; Ghedotti, 2000, ch. 4, fig. 3; Lucinda & Reis, 2005, ch. 3): (0) large (beyond first vertebra; Ghedotti, 1998, figs. 2a-b, d), (1) medium (reaches first vertebra), (2) small (does not reach first vertebra; Ghedotti, 1998, fig. 2c), (3) absent (Ghedotti, 1998).

Ghedotti recognized three states for this character, in the present study was included another state, a medium size state, when the epiotic processes reach the first vertebra but do not go beyond it (referred to as “long” by Lucinda & Reis, 2005). According to Ghedotti (1998), this character was not a synapomorphy for Anablepidae because some species of that family lack epiotic processes. Ghedotti (1998) stated that the epiotic processes in Fluviphylax obscurum were absent, while in this study the specimen of F. pygmaeus examined possessed small epiotic processes (state 2). Lucinda & Reis (2005) stated that long epiotic processes were present in Jenynsia, Aplocheilichthys Phallichthys, Alfar, Poeciliopsis, Quintana, and Carlhubb'sia. According to Ghedotti & Reis (2005), Priapichthys, Girardinus, Poecilia, Neoheterandria, Scolichthys, Xiphophorus, and Limia have medium-sized epiotic processes, while short processes are present in Heterandria and Pseudopoecilia. Ghedotti (2000) reported the absence of epiotic processes in Tomeurus and Cnesterodon. Lucinda & Reis (2005) also reported them absent in Pamphorichthys and Micropoecilia.

20. Supraoccipital process (Lucinda & Reis, 2005, ch. 4): (0) bifid, outer portion large (Lucinda & Reis, 2005; fig. 4c), (1) bifid, outer portion diminutive (Lucinda & Reis, 2005; fig. 4b), (2) simple, not split (Lucinda & Reis, 2005; fig. 4a), (3) absent.
Fig. 5. Lateral view of retroarticular in: a) Phalloceros buckupi, SU 64230; b) Heterandria formosa, TU 24002; c) Brachyrhaphis olomina, TU 24997; d) Xenophallus umbratilis, TU 186285; e) Phalichthys fairweatheri, USAC 797; f) Carlhubbsia stuarti, USAC 204; g) Poeciliopsis turrobaensis, TU 24130; h) Brachyrhaphis terrabensis, AMNH 2503; i) Pseudoxiphophorus bimaculatus, TU 183396; j) Jenynsia eirmostigma, MCP 40527; k) Phalloceros caudimaculatus, MCP 40527; hyomandibula in: l) Heterandria formosa, TU 24002; m) Pseudoxiphophorus obliquus, LSUMZ 14339; n) Pseudoxiphophorus diremptus, LSUMZ 14337; o) Pseudoxiphophorus bimaculatus, TU 183396; p) Brachyrhaphis olomina, TU 24997; q) Jenynsia eirmostigma, MCP 40527; r) Anableps dowi, TU 39076; s) Profundulus guatemalensis, USAC 769; dentary in: t) Pseudoxiphophorus obliquus, LSUMZ 14339; u) Pseudoxiphophorus diremptus, LSUMZ 14337; v) Brachyrhaphis episcopi, SU 50303; w) Brachyrhaphis olomina, TU 24997; x) Pamphorichthys araguaensis, MCP 40919; y) Heterandria formosa, TU 24002; z) Profundulus guatemalensis, USAC 769. Scale bars = 1 mm.
In this study, this character was modified from Lucinda & Reis (2005) by adding state 3. Lucinda and Reis (2005) reported that *Girardinus* and *Pamphorichthys* have minute external bifid halves minute, while *Phalloceros* and *Cnesterodon* have a large external half.

21. Supraoccipital crest robust and long (Ghedotti, 1998): (0) absent, (1) present.

22. Ascending process of parasphenoid (Ghedotti, 1998, 2000, ch. 4, figs. 3b, f-g; Lucinda & Reis, 2005, ch. 20): (0) long (Ghedotti, 1998, fig. 4), (1) short.

Ghedotti (1998) stated that for most species of *Jenynsia* and *Fluviphylax* examined the parasphenoid was short. Only two states were examined in this study because *Xenodexia* was not available for loan.

23. Vomer (Ghedotti, 2000, ch. 6, figs. 3f-h): (0) present, (1) absent.

Vomer was present (state 0) in most of the species.

Sensory system

24. Extent and shape of dorsomedial lamella of the autopalatine (Costa, 1998, ch. 29, fig. 4): (0) long, (1) short, (2) absent.

25. Shape of cartilaginous head of autopalatine (Costa, 1998, ch. 31, fig. 4c): (0) narrow, (1) wide.

26. Extent of lateral lamella of the hyomandibula (Costa, 1998, ch. 35, fig. 4; Ghedotti, 1998, ch. 23, fig. 8; Ghedotti, 2000, ch. 29, figs. 6b-d; Fig. 5): (0) not expanded (1) expanded (Ghedotti, 1998, fig. 8c).

27. Shape of the medial process of the lachrymal (Ghedotti, 2000, ch. 25, fig. 4): (0) short and broad, (1) long and narrow.

28. Shape of the dorsoposterior region of lachrymal bordering the orbit (Ghedotti, 2000, ch. 26, figs. 4a-d): (0) dorsally indented, (1) without indentation, (2) short.

29. Dorsoposterior process of the autopalatine (Ghedotti, 2000, ch. 27, fig. 6): (0) long, (1) short, (2) absent.

30. Anterior margin of the dermalplatine (Ghedotti, 2000, ch. 27, fig. 6): (0) straight, (1) concave, (2) convex.

Branchial skeleton

31. Shape of dorsal process of urohyal (Costa, 1998, ch. 37, fig. 5): (0) straight, (1) bent.

32. Condyles on anterior portion of anterior ceratohyal (Costa, 1998, ch. 42; Parenti, 1981, fig. 28): (0) two, (1) one.

33. Anterior process of anterior ceratohyal extending ventrally, to ventral hypohyal (Ghedotti, 2000, fig. 8; Lucinda & Reis, 2005, ch. 24): (0) present, (1) absent.

According to Lucinda & Reis (2005), the absence of the anterior process was a synapomorphy for Poeciliinae except *Tomewurus*, *Brachyrhaphis*, and *Heterandria*.

34. (Lucinda & Reis, 2005, ch. 23): First and second branchiostegal rays united at the base: (0) absent, (1) present.

According to Lucinda & Reis (2005), *Scolichthys* and *Neo heterandria* present a synapomorphic condition that branchiostegal rays were united at the base.

35. Anterior margin of the first hypobranchial (Ghedotti, 2000, ch. 35, figs. 9a-b, Lucinda & Reis, 2005, ch. 29): (0) straight, (1) concave.

36. Posterior border of first hypobranchial with a pronounced concavity (Costa, 1996): (0) absent, (1) present.

37. Teeth on the second and third hypobranchials (Ghedotti, 2000, ch. 36, fig. 9): (0) absent, (1) present.

38. Cartilaginous heads on the third hypobranchial (Ghedotti, 2000, ch. 37, fig. 9): (0) two heads, (1) one, continuous, (2) none.

39. Teeth on the fourth ceratobranchial (Ghedotti, 2000, ch. 38, fig. 9; Lucinda & Reis, 2005, ch. 27): (0) absent, (1) present.

40. Teeth in the fifth ceratobranchial (Lucinda & Reis, 2005, ch. 28): (0) wide regular, (1) narrow irregular, (2) absent.

41. Third and fourth pharyngobranchial toothplates (Ghedotti, 2000, ch. 41, fig. 10b; Lucinda & Reis, 2005, ch. 26): (0) fused, (1) separated.

Pectoral fin and girdle

42. Number of pectoral-fin rays (Miller, 1979; Rosen, 1979; Parenti, 1981; Ghedotti, 1998, ch. 34): (0) 10 or less, (1) 11-12, (2) 13-14, (3) 15 or more.

43. Shape of supracleithrum (Costa, 1998, ch. 45, fig. 4a; Ghedotti, 2000, ch. 54, fig. 12; Fig. 6): (0) broad, (1) narrow, (2) absent.

44. Fusion of supracleithrum with post-temporal (Costa, 1998, ch. 76; Fig. 6): (0) absent, (1) present.

45. Position of pectoral-fin origin in relation to midline of the body (Ghedotti, 2000, ch. 51; Lucinda & Reis, 2005, ch. 32): (0) above, (1) below.

46. Ventral arm of post-temporal (Ghedotti, 2000, ch. 52, figs. 12a-c; Fig. 6): (0) ossified, (1) non-ossified.

47. Posterior extension of dorsal enclosure of cleithrum (Ghedotti, 2000, ch. 55, figs. 12a-d; Fig. 6): (0) slightly curved, (1) straight.
48. Broad postcleithrum: (0) absent, (1) present (Costa, 1998).

49. Dorsal proximal pectoral radials (Ghedotti, 2000, ch. 56, fig. 12; Figs. 6, 17): (0) fused, (1) free.

50. Posteroventral coracoid (Ghedotti, 2000, ch. 57, fig. 16; Fig. 6): (0) straight, (1) round.

51. Post-temporal (Rosen, 1967, fig. 4; Lucinda & Reis, 2005, ch. 31; Fig. 6): (0) bifid, (1) unbranched.

Pelvic fin and girdle

52. Number of pelvic-fin rays (Ghedotti, 2000, ch. 60; Lucinda & Reis, 2005, ch. 33): (0) 3-4, (1) 6-8.

53. Ventral lamella on central part of basipterygium in males (Ghedotti, 2000, ch. 6, figs. 13a-e): (0) absent, (1) present.

54. Medial process of basipterygium in males (Ghedotti, 1998, ch. 39; Ghedotti, 2000, ch. 62): (0) separated (Ghedotti, 1998 fig. 14b), (1) overlap (Ghedotti, 1998, fig. 14a), (2) fused.

55. Position of pelvic girdle in adult males (Ghedotti, 2000, ch. 65; Lucinda & Reis, 2005, ch. 35): (0) same as females, (1) more anterior in males, (2) more posterior in males.

The character states proposed by Lucinda & Reis (2005) were reduced in this study.

56. Pelvic-fin insertion between the pleural ribs of vertebrae 3 to 6 (Costa, 1998, ch. 85): (0) absent, (1) present.

57. Pelvic-fin length in adult males (Lucinda & Reis, 2005, ch. 34): (0) short (extending up to anal-fin origin), (1) medium (extending beyond origin of anal fin but do not reaching end of anal fin), (2) long (reaching beyond end of anal fin).

58. Ray 3 of the pelvic fin expanded distally and with several branches (Rodriguez, 1997): (0) absent, (1) present.

59. Ray 2 of the pelvic fin with a distal triangular shape, comb-like, formed by spiny bony processes of these segments of the ray and covered by a fleshy membrane (Rodriguez, 1997; Fig. 8): (0) absent, (1) present.

60. Second ray of pelvic fin separated from rays 3-5 by a large gap (Rodriguez, 1997): (0) absent, (1) present.

61. Dorso-lateral process of the basipterygium in adult males (Lucinda & Reis, 2005, ch. 36, fig. 8): (0) short, (1) long.

62. Lateral keel of the basipterygium in adult males (Lucinda & Reis, 2005, ch. 38): (0) absent, (1) present.

Lucinda & Reis (2005) reported that this was a derived feature in Scolichthys, Pamphorichthys, Poeciliopsis, and Phalloptychus.

63. Shape of anterior tip of the basipterygium in adult males (Lucinda & Reis, 2005, ch. 37, figs. 8c-d, 9, 11): (0) triangular, (1) triangular with round tip (2) sinuous, (3) pointed, (4) round, (5) round with keel-tip.

The character states used in this study differ from Lucinda & Reis (2005).

64. Narrowing of the lateral surface of the basipterygium in males (Lucinda & Reis, 2005, ch. 39, fig. 9): (0) present, (1) absent.

Axial skeleton

65. Number of dorsal-fin rays in males (Rosen, 1979; Ghedotti, 2000, ch. 86; Lucinda & Reis, 2005, ch. 64): (0) 9 or fewer, (1) 10-11, (2) 12 or more.

66. Position of dorsal-fin origin in adult males (Ghedotti, 1998, ch. 58; Ghedotti, 2000, ch. 87; Lucinda & Reis, 2005, ch. 62): (0) anterior to vertebra 9, (1) between vertebrae 9 and...
10. (2) between vertebrae 10-11 or 11-12, (3) between vertebrae 12-13, (4) between vertebrae 13-14, (5) behind vertebra 14. This character was coded into more states than in Ghedotti (1998).

67. Number of vertebrae (Regan, 1913; Miller, 1979; Rosen, 1979; Parenti, 1981; Rauchenberger, 1989; Ghedotti, 1998, ch.31): (0) 25-26, (1) 27, (2) 28, (3) 29, (4) 30, (5) 31 or more.

Ghedotti (1998) coded this character for the family Anablepidae using only two character states, fewer than 45, and 45 or more. This character was coded into more states than in previous publications.

68. Total number of vertebrae with a widened neural spines: (0) none, (1) one, (2) two, (3) three, (4) four, (5) nine.

69. Distal tips of pleural ribs associated with sixth to tenth vertebrae (Rosen & Bailey, 1963, figs. 17, 26, 49; Rodriguez, 1997, ch. 1; Ghedotti, 2000, ch. 50): (0) slender and straight, (1) slender and curved, (2) thickened and curved.

70. Haemal arches and spines of vertebrae 13-17 in males (Lucinda & Reis, 2005, ch. 47): (0) rudimentary, not modified into gonapophyses, (1) modified into gonapophyses.

71. Size of caudal accessory cartilage between distal neural spines of preural vertebrae 3 and 4 (Costa, 1998; Ghedotti, 2000, figs. 17a-b; Fig. 7): (0) large, (1) small.

72. Shape of caudal accessory cartilage between distal neural spines of pleural vertebrae 3 and 4 (Fig. 7): (0) rounded, (1) ovoid, (2) squared, (3) n-shaped, (4) 7-shaped, or triangular.

73. Complete median horizontal gap between dorsal and ventral hypurals (Costa 1998): (0) absent, (1) present.

74. Hypural plate in adult males (Parenti, 1981; Ghedotti, 1998, ch. 59: Ghedotti, 2000, ch. 88, fig. 17; Lucinda & Reis, 2005, ch. 131; Fig. 7): (0) posterior margin completely separate or bipartite, (1) with a median aperture of 3/4 to 1/2 of its size, (2) with a median aperture less than half its size, (3) fused.

75. Number of caudal-fin rays in contact with hypural plate (Lucinda & Reis, 2005, ch. 132): (0) 7 or less, (1) 8-9, (2) 10-11, (3) 12-13, (4) 14-15.

76. Ossification state of hypural plates: (0) fully ossified, (1) only tips of posterior margin cartilaginous, (2) 1/3 cartilage, (3) 2/3 cartilage, (4) fully unossified.

Anal fin and Gonopodium

77. Male anal fin (Hubbs, 1924, Parenti, 1981, Ghedotti, 1998, ch. 41; Ghedotti, 2000, ch. 67, fig. 14; Lucinda & Reis, 2005, ch. 86): (0) unmodified, almost identical to female anal fin, (1) modified into an intromittent organ, fleshy tube supported by more than three anal-fin rays, (2) modified into a gonopodium supported by anal-fin rays three to five.

78. Sexual laterality in males (Ghedotti, 1998, ch. 68; Ghedotti, 2000, ch. 94): (0) absent, (1) present (Ghedotti, 1998, fig. 24).

This character refers to laterally directed papillae in relation to the midline.

79. Male anal-fin length (gonopodium length) (Ghedotti, 2000, ch. 68): (0) short, extending from vertebrae 9 to vertebrae 15, (1) medium, extending from vertebrae 16 to vertebrae 29, (2) long, extending further back from vertebrae 30.

This character was coded taking into account the number of vertebrae from the position of the gonopodium origin to the last vertebrae reached by the gonopodial tip.

80. Number of anal-fin rays in females (Ghedotti, 1998, ch. 40; Ghedotti, 2000, ch. 69; Lucinda & Reis, 2005, ch. 85): (0) 9 or less, (1) 10, (2) 11, (3) 12 or more.

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Fig. 7. Lateral view of caudal-fin skeleton in: a) Phalloceros caudimaculatus, MCP 4052; b) Heterandria formosa, TU 24002; c) Gambusia affinis, TU 114758; d) Pamphorichthys araguaiensis, MCP 40919; e) Brachyrhaphis episcopi, SU 50303; f) Pseudoxiphophorus obliquus, LSUMZ 14339; g) Pseudoxiphophorus bimaculatus, CAS 78795; h) Brachyrhaphis rhabdophora, TU 84483. Scale bars = 1 mm.
### 81. Ventral margin of the gonopodium (Rosen, 1979): (0) straight, (1) concave behind spines.

### 82. Gonopodium symmetry (Rosen & Bailey, 1959; Rosen 1979; Lucinda & Reis, 2005, ch. 87): (0) symmetric, (1) asymmetric.

### 83. Distal tip of anal-fin ray 3 in males (Rosen & Bailey, 1963, fig. 31; Ghedotti, 2000, ch. 72, fig. 14): (0) absent, (1) simple, (2) segmented, (3) with spinous cirri.

### 84. Fleshy palp on ventral surface of anal-fin ray 3 in males (Rosen & Bailey, 1963, fig. 25; Rodriguez, 1997, fig. 5; Ghedotti, 2000, ch. 73; Lucinda & Reis, 2005, ch. 88; Fig. 8): (0) absent, (1) present.

### 85. Pedicle at tip of anal-fin ray 3 (Rosen & Bailey, 1963, fig. 31; Lucinda & Reis, 2005, ch. 90; Figs. 8): (0) absent, (1) present.

### 86. Subdistal spines of anal-ray 3 curved forward (Rodriguez, 1997, Fig. 8): (0) absent, (1) present.

### 87. Anal-fin ray 3 with a large decurved hook distally and an ossified bladelike structure on the dorsal margin of the decurved hood (Rosen, 1979; Rodriguez, 1997; Fig. 8): (0) absent, (1) present.

### 88. Shape of anal-fin ray 4a (Rosen, 1979): (0) curved up, (1) curved down, (2) straight, (3) curved left, (4) curved right.

### 89. Keel structure situated ventrally on anal-fin ray 4a (Rauchenberger, 1989; Fig. 8): (0) absent, (1) present.

### 90. Serrae on posterior margin of posterior branch of anal-fin ray 4 in adult males (Howell & Hubbs, 1936; Miller, 1974; Ghedotti, 2000, ch. 74, fig. 15): (0) absent, (1) present.

### 91. Anal-fin ray 4p with two series of distal serrae separated at the level of the hook of ray 5a in undifferentiated segments (Rosen, 1979; Rodriguez, 1997): (0) absent, (1) present.

### 92. Ventral projection of anal-fin ray 4 (Rauchenberger, 1989, figs. 20, 38; Lucinda & Reis, 2005, ch. 113): (0) ventral projection absent, (1) ventral projection curved toward ray 3, (2) ray 3 curved toward ray 4, (3) ventral projection twisted.

### 93. Depth of distal segments of anal-fin ray 4p serrae (Rodriguez, 1997, ch. 12, fig. 5; Lucinda & Reis, 2005, ch. 115): (0) segments absent, (1) wider than tall, (2) taller than wide.

### 94. Number of subdistal recurved spines on anal-fin ray 4p (Lucinda & Reis, 2005, ch. 116): (0) less than 9, (1) 10-12, (2) 13-14, (3) more than 15.

### 95. Elongate dorsal protuberance just behind recurved spines series of anal-fin ray 4p (Rosen, 1979, fig. 6; Lucinda & Reis, 2005, ch. 118): (0) absent, (1) present.

Elongated dorsal protuberance behind recurved spines of ray 4p (state 0) are absent in most species. An elongate dorsal protuberance behind recurved spines of ray 4p (state 1) is present in Alfaro cultratus and Micropoecilia most Poecilia (except P. butleri), H. formosa, Brachyrhaphis terrabensis, Phallichthys, Limia, Xiphophorus, Scolichthys, Cnesterodon, Phalloceros, Priapichthys, and Pseudoxiphophorus (except a reversal in P. obliquus P. anzuetoi and P. diremptus).

### 96. Pedicle of anal-fin ray 3 united to anal-fin ray 4 (Rosen & Bailey, 1963, fig. 31; Lucinda & Reis, 2005, ch. 90; Fig. 8): (0) absent, (1) present.

### 97. Subdistal spines on gonopodial ray three enlarged ventrally to form a slight elbow along the ventral profile of the ray (Rosen, 1979): (0) absent, (1) present.

### 98. Distal fourth of gonopodium with an increased ventral flexure of its ray and moderately to strongly concave along ventral profile behind enlarged spines of ray 3 (Rosen, 1979): (0) absent, (1) present.

### 99. Terminal segment of gonopodial ray 4a forming large decurved hook six times or more as long as the height from the base to the tip (Rosen, 1979): (0) absent, (1) present.

### 100. Large decurved terminal hook on gonopodial ray 4a L-shaped or in form of open K (Rosen, 1979): (0) absent, (1) present.

### 101. Gonopodial ray 4p as long as or longer than ray 5a excluding the latter from direct contact with ray 4a (Rosen, 1979): (0) absent, (1) present.

### 102. Penultimate short segments of gonopodial ray 4a withdrawn from base of terminal membranous envelope and two or more in number (Rosen, 1979): (0) absent, (1) present.

### 103. Subdistal short segments of gonopodial ray 4a more than four in number and bearing ventral peg-like processes (Rosen, 1979): (0) absent, (1) present.

### 104. Intermediate segments of gonopodial ray 5p enlarged, forming a semi-elliptic swelling (Lucinda & Reis, 2005, ch. 112): (0) absent, (1) present.

### 105. Terminus of anal-fin rays 5a, 5p, 4a, 4p (Rauchenberger, 1989, fig. 23; Lucinda & Reis, 2005, ch. 112): (0) directed dorsally, (1) straight, (2) directed ventrally, (3) Anal-fin ray 5 directed dorsally and ray 4 directed ventrally, (4) Ray 5 directed ventrally and ray 4 directed dorsally, (5) twisted.
Fig. 8. Lateral view of the tip of gonopodium in: a) Heterandria formosa, TU 24002; b) Pseudoxiphophorus cataractae, LSUMZ 14338; c) Pseudoxiphophorus obliquus, LSUMZ 14339; d) Pseudoxiphophorus litoperas, USAC 1813; e) Phalloceros caudimaculatus, MCP 40527; f) Cnesterodon decemmaculatus, MCP 35215; g) Xiphophorus mayae, LSUMZ 14336; h) Poecilia rositae, USAC 1822; i) Gambusia affinis, TU 114758; j) Phallocerichthys fairweatheri, USAC 1797; k) Brachyrhaphis rhabdophora, TU 84483; l) Pamphorichthys araguaiensis, MCP 40919; m) Alfaro cultratus, TU 24927. Scale bars = 1 mm.
106. Posteriorly directed claw on distal tip of anterior branch of anal-fin ray 5 (Ghedotti, 2000, ch. 76, fig. 15; Fig. 8): (0) absent, (1) present.

107. Keel on posterior ventral surface of anal-fin ray 5 (Rodriguez, 1997, fig. 5f; Lucinda & Reis 2005, ch. 119): (0) keel without axial projection, (1) keel projects anteriorly toward anal-fin ray 4p.

108. Anal-fin ray 5 decurved away from body axis, around end of ray 4 (Rodriguez, 1997, fig. 5f; Lucinda & Reis, 2005, ch. 120): (0) absent, (1) present.

109. Groove on dorsal margin of anal-fin ray 5 (Rodriguez 1997, ch. 2; Lucinda & Reis, 2005, ch. 121): (0) narrow, (1) broad.

110. Distal segment at tip of anal-fin ray 5a (Miller, 1974; Rauchenberger, 1989; fig. 20; Rosa & Costa, 1993; Ghedotti, 2000, fig. 15a; Lucinda & Reis, 2005, ch. 122; Fig. 8): (0) absent, (1) normal, (2) hooked, (3) triangular with spine, (4) with claw.

111. Dorsal expansion of anal-fin ray 5p (Rosa & Costa, 1993; Lucinda & Reis, 2005, ch. 124, fig. 28): (0) absent, (1) present.

112. Serrae on anal-fin ray 5p (Lucinda & Reis, 2005, ch. 125): (0) absent, (1) present.

113. Fusion between lower and upper branches of anal-fin ray 6a and 6p (Lucinda & Reis, 2005, ch. 126): (0) absent, (1) partial, (2) total.

114. Degree of fusion among more distal elements of branches of anal-fin ray 6 (Lucinda & Reis, 2005, ch. 127): (0) not fused, (1) partial, (2) total fused.

115. Distal portion of anal-fin ray 6 (Lucinda & Reis, 2005, ch. 128): (0) not expanded, (1) expanded.

116. Inclination of gonactinost complex (anal-fin radials 3-5 in males) relative to longitudinal axis of body (Ghedotti, 2000, ch. 79, figs. 14, 16; Lucinda & Reis, 2005, ch. 68): (0) absent, (1) backward, (2) forward, (3) perpendicular.

117. Small lateral process on base of fifth middle anal-fin radial (Rodriguez, 1997: Fig. 8): (0) absent, (1) present.

118. Gonactinost 1-5 (Rauchenberger, 1989, fig. 22; Fig. 9): (0) gonactinost 5 not ossified, (1) gonactinost 5 ossified and fused with gonactinosts 1-4 into a single laminar plate resembling spokes in a fan (2) gonactinost 5 ossified and not fused with gonactinosts 1-4, (3) gonactinosts 1-5 all ossified separately.

119. Gonactinost 8 (Rosen & Bailey, 1963, fig. 53; Lucinda & Reis, 2005, ch. 83; Fig. 9): (0) absent, (1) straight, (2) wing like.

120. Gonactinost 9 (Rosen & Bailey, 1963, fig. 53; Lucinda & Reis, 2005, ch.84; Fig. 9): (0) absent, (1) wing-like, (2) straight and smaller than other gonactinost, (3) straight and same size as other gonactinosts, (4) straight and half the size of other gonactinosts.

121. Distal portion of gonactinosts 3-4 completely fused: (0) absent, (1) present.

122. Ligastyle, a free haemal spine in adult males located dorsal to the gonopodial suspensorium and close to the vertebral column near the thirteenth vertebra (Ghedotti, 2000, ch. 44; Fig. 9): (0) cartilaginous, (1) ossified.

123. Ligastyle shape (Lucinda & Reis, 2005, ch. 46, fig. 13): (0) short, (1) long one-axis, (2) long triangular.

124. Number of well-developed gonapophyses (Rodriguez, 1997, ch. 9, fig. 4; Lucinda & Reis, 2005, ch. 48; Fig. 10): (0) rudimentary, (1) 1-2, (2) 3-4.
Fig. 10. Lateral view of gonapophyses in: a) *Brachyrhaphis olomina*, TU 24997; b) *Brachyrhaphis rhabdophora*, TU 84483; c) *Poeciliopsis pleurospilus*, USAC 838; d) *Brachyrhaphis episcopi*, SU 50303; e) *Xiphophorus alvarezi*, USAC 1782; f) *Brachyrhaphis terrabensis*, AMNH 20503; g) *Phallichthys fairweatheri*, USAC 1797; h) *Poecilia rosiatae*, USAC 1822; i) *Priapichthys annectens*, TU 25193; j) *Pseudoxiphophorus bimaculatus*, CAS 78795; k) *Poeciliopsis turrubarenensis*, TU 24130; l) *Pseudoxiphophorus obliquus*, LSUMZ 14339; m) *Poeciliopsis gracilis*, TU 84966; n) *Phalloceros caudimaculatus*, MCP 40527; o) *Pseudoxiphophorus litoperas*, USAC 1813; p) *Gambusia affinis*, TU 114758; q) *Heterandria formosa*, TU 24002; r) *Pamphorichthys araguaiensis*, MCP 40919; s) *Phalloptychus iheringi*, MCP 11054. Scale bars = 1 mm.
125. Position of functional gonapophyses (Lucinda & Reis, 2005, ch. 49): (0) between vertebrae 12-13 or 13-14, (1) between vertebrae 13-15, (2) between vertebrae 13-16, (3) between vertebrae 15-19 or 15-17.

126. First gonapophysis reduced to a support for adjacent gonapophyses located between vertebrae 12-13 (Rodriguez, 1997, fig. 4b; Lucinda & Reis, 2005, ch. 50): (0) absent, (1) present.

127. Hollister's foramen on first or second gonapophyses (Rodriguez, 1997, ch. 7, fig. 4b; Lucinda & Reis, 2005, ch. 51; Fig. 10): (0) absent, (1) present.

128. Shape of gonapophysis at vertebra 14 (Rosen & Bailey, 1963, figs. 24a, 26c-d, 29b, 46, 56a-d; Lucinda & Reis, 2005, ch. 52; Fig. 10): (0) same as other vertebrae, (1) rudimentary, (2) straight, (3) forming an acute angle, (4) curved.

Scales and pigmentation:
129. Number of lateral-line scales in male (Lucinda & Reis, 2005, ch. 125): (0) less than 23, (1) 24-25, (2) 26-27, (3) 28-29, (4) more than 30.

130. Dorsal-fin pigment (Rosen, 1979): (0) diffuse spots among rays, (1) diffuse spots only at the tips, (2) pigmented rays, (3) one spot at the base.

131. Dorsal-fin melanophores forming two or more rows of discrete spots and anal-fin melanophores (Rosen, 1979): (0) not extending distally from the anal-fin base as streaks in interradial membrane, (1) extending distally from the anal-fin base as streaks in interradial membrane.

132. Anal-fin pigmentation (Rosen, 1979): (0) small pigment spots at distal fin margin, (1) small pigment spots along the fin, (2) pigments at the base, (3) large pigment spots in the fin base.

133. Basicaudal spot (Rosen, 1979): (0) absent, (1) present.

134. Basicaudal spot in adults partly or wholly above lateral line and extending onto caudal-fin base (Rosen, 1979): (0) absent, (1) present.

135. Basicaudal spot large spherical dusky filling area between midlateral line and dorsal margin of caudal-fin peduncle (Rosen, 1979): (0) absent, (1) present.

136. Pigment of scale pockets forming a strong reticular network along the sides of the body (Rosen, 1979): (0) absent, (1) present.

137. Midlateral pigment stripe (Rosen, 1979; Ghedotti, 1998, ch. 62): (0) slight line, (1) diffuse, formed by small dots, (2) single large dot at mid body, (3) strong, formed by small dots, (4) strong formed by large dots, (5) strongly developed.

138. Crosshatching on the body (Rosen, 1979): (0) well-developed, (1) diffuse.

139. Elongate vertical bars on the lateral surface of the body (Lucinda & Reis, 2005, ch. 133): (0) absent, (1) present.

140. Sexual dimorphism in color pattern (Costa, 1998, ch. 106): (0) absent, (1) present.

Body proportions:
141. Adult sexual size dimorphism (Ghedotti, 2000, ch. 95): (0) no difference between sexes, (1) females larger, (2) males larger.

142. Body shape (Rosen, 1979; Ghedotti, 2000, ch. 98): (0) abrupt transition between abdomen and caudal peduncle, (1) no defined transition between abdomen and caudal peduncle.

143. Mouth terminal (Ghedotti, 1998): (0) absent, (1) present.

Cephalic sensory system
144. Posterior supraorbital canal (pores 2b, 3, 4a) (Costa, 1998, ch. 98; Ghedotti, 2000, ch. 90; Lucinda & Reis, 2005, ch. 5): (0) pore present, open, forming a depression, (1) pore present, closed (Gosline, 1949, plate II, figs. 1, 4), (2) pore absent or forming a shallow groove (Rosen & Mendelson, 1960, figs. 3B, J-M).

145. Lachrymal canal in adults (Ghedotti, 2000, ch. 91): (0) closed canal, forming pores, (1) open canal.

146. Preorbital canal (Rosen & Mendelson, 1960, fig. 2; Lucinda & Reis, 2005, ch. 9): (0) present open, (1) present closed, (2) absent.

147. Anterior section of posterior remnant of infraorbital system (pore 4b, 5, 6a) (Lucinda & Reis, 2005, ch. 6): (0) pores closed, not visible, (1) pores were absent, forming a groove (Rosen & Mendelson, 1960, figs. 3a-e, g-p), (2) pores were closed, forming a depression (Gosline, 1949, plate II, figs. 1, 4), (3) pores were closed forming a groove, (4) pores open forming a depression, (5) pores open, forming a groove.

148. Posterior section of posterior remnant of infraorbital system (canal 6b, 7) (Lucinda & Reis, 2005, ch. 7): (0) open (Rosen & Mendelson, 1960, figs. 2a-b), (1) closed (Gosline, 1949, plate II, fig. 4; Parenti, 1981, fig. 14a).

149. Preopercular canal (Lucinda & Reis, 2005, ch. 8): (0) absent (Rosen & Mendelson, 1960, figs. 2a-b), (1) present (Gosline, 1949, plate II, fig. 2).

150. Nasal canal (Parenti, 1981): (0) large, (1) reduced.
**Appendix 3.** Data matrix of 150 characters for 52 poeciliine and four outgroup taxa. “?” indicates a character not coded due to lack of specimens, structures, or inapplicable codings. 

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Monophyly of Heterandriini
Appendix 3 (cont.). Data matrix of 150 characters for 52 poeciliine and four outgroup taxa. “?” indicates a character not coded due to lack of specimens, structures, or inapplicable codings. P = 0&1; Q = 0&2; R = 1&3; S = 1&4; T = 0&1&2.
Appendix 3 (cont.). Data matrix of 150 characters for 52 poeciliine and four outgroup taxa. "?” indicates a character not coded due to lack of specimens, structures, or inapplicable codings. P = 0&1; Q = 0&2; R = 1&3; S = 1&4; T = 0&1&2.

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Profundulus guatemalensis

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Anableps dowi

Fluviphax pygmeaus

Tomeurus gracilis

Alfaro cultratus

Alfaro huberi

Belonosox belizanus

Brachyhaphis episcopi

Brachyhaphis olomina

Brachyhaphis rhabdophora

Brachyhaphis tarrabensis

Carlbubbasia stuarti

Cnesterodon decemmaculatus

Gambusia affinis

Heterandria anzueto

Heterandria bimaculatus

Heterandria cataractae

Heterandria diremptus

Heterandria forma

Heterandria ilotorpus

Heterandria obliquus

Limia dominicensis

Limia melanogaster

Microprocenia branneri

Monophyly of Heterandriini

Poecilia latipinna

Poecilia mexicana

Poecilia retropinna

Poecilia rositaen

Poecilia salvatoris

Poecilia sphenops

Poeciliopsis elongate

Poeciliopsis fasciata

Poeciliopsis gracilis

Poeciliopsis infans

Poeciliopsis latidens

Poeciliopsis lucida

Poeciliopsis julzi

Poeciliopsis pleurospilus

Poeciliopsis presidionis

Poeciliopsis prolificus

Poeciliopsis turrubarenensis

Poeciliopsis viriosa

Pripichthys annectens

Quintana strizouna

Scolichthys grceenwayi

Xiphophorus alvarezi

Xiphophorus mayae