

Citation: Favarato RM, Ribeiro LB, Campos A, Porto JIR, Nakayama CM, Ota RP, et al. (2021) Comparative cytogenetics of Serrasalmidae (Teleostei: Characiformes): The relationship between chromosomal evolution and molecular phylogenies. PLoS ONE 16(10): e0258003. https://doi.org/10.1371/journal.pone.0258003

Editor: Paolo Ruggeri, Natural History Museum of London, UNITED KINGDOM

Received: June 11, 2021

Accepted: September 15, 2021

Published: October 7, 2021

Copyright: © 2021 Favarato et al. This is an open access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Data Availability Statement: All relevant data are within the manuscript and its Supporting

Information files.

Funding: This study was supported by the Brazilian agencies, Fundação de Amparo à Pesquisa do Estado do Amazonas (FAPEAM) FAPEAM/
SEPLANCTI/ Government of the State of Amazonas - Edital PAPAC 005/2019, Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES) - CAPES Pro-Amazon and Conselho

RESEARCH ARTICLE

Comparative cytogenetics of Serrasalmidae (Teleostei: Characiformes): The relationship between chromosomal evolution and molecular phylogenies

Ramon Marin Favarato 11*, Leila Braga Ribeiro², Alber Campos¹, Jorge Ivan Rebelo Porto³, Celeste Mutuko Nakayama^{3†}, Rafaela Priscila Ota 14*, Eliana Feldberg 15*, alber Campos¹, Eliana Feldberg 15*, alber Campos¹, Eliana Feldberg 15*, alber Campos¹, blana Feldberg 15*, alber Campos¹, alber Campos¹, blana Feldberg 15*, alber 15*, alb

- 1 Programa de Pós-Graduação em Genética, Conservação e Biologia Evolutiva, Instituto Nacional de Pesquisas da Amazônia, Petrópolis, Manaus, Amazonas, Brazil, 2 Centro de Ciências da Saúde, Universidade Federal de Roraima, Avenida Capitão Ene Garcêz, Boa Vista, RR, Brazil, 3 Coordenação de Pesquisas em Biodiversidade, Instituto Nacional de Pesquisas da Amazônia, Petrópolis, Manaus, Amazonas, Brazil, 4 Departamento de Biologia Estrutural e Funcional, Instituto de Biociências, Universidade Estadual Paulista "Júlio de Mesquita Filho", Botucatu, São Paulo, Brazil
- † Deceased.
- * ramonfavarato@gmail.com

Abstract

Serrasalmidae has high morphological and chromosomal diversity. Based on molecular hypotheses, the family is currently divided into two subfamilies, Colossomatinae and Serrasalminae, with Serrasalminae composed of two tribes: Myleini (comprising most of pacus species) and Serrasalmini (represented by Metynnis, Catoprion, and remaining piranha's genera). This study aimed to analyze species of the tribes Myleini (Myloplus asterias, M. lobatus, M. rubripinnis, M. schomburgki, and Tometes camunani) and Serrasalmini (Metynnis cuiaba, M. hypsauchen, and M. longipinnis) using classical and molecular cytogenetic techniques in order to understand the chromosomal evolution of the family. The four species of the genus Myloplus and T. camunani presented 2n = 58 chromosomes, while the species of Metynnis presented 2n = 62 chromosomes. The distribution of heterochromatin occurred predominantly in pericentromeric regions in all species. Tometes camunani and Myloplus spp. presented only one site with 5S rDNA. Multiple markers of 18S rDNA were observed in T. camunani, M. asterias, M. lobatus, M. rubripinnis, and M. schomburgkii. For Metynnis, however, synteny of the 18S and 5S rDNA was observed in the three species, in addition to an additional 5S marker in M. longipinnis. These data, when superimposed on the phylogeny of the family, suggest a tendency to increase the diploid chromosome number from 54 to 62 chromosomes, which occurred in a nonlinear manner and is the result of several chromosomal rearrangements. In addition, the different karyotype formulas and locations of ribosomal sequences can be used as cytotaxonomic markers and assist in the identification of species.

Nacional de Desenvolvimento Científico e Tecnológico (CNPq). The Center for Studies of Adaptation to Environmental Changes in the Amazon was founded by FAPEAM and CNPq (INCT ADAPTA II, FAPEAM/CNPq 573976/2008-2). EF was the recipient of a fellowship from CNPq (Grant #301886/2019-9) and RPO is founded by Capes (grant #12002011001P7). The National Amazon Research Institute (INPA)/Postgraduate Studies in Genetics, Conservation and Evolutionary Biology (INPA/GCBEv), given institutional support for this study. There was no additional external funding received for this study.

Competing interests: The authors have declared that no competing interests exist.

Introduction

Serrasalmidae is a family of fishes endemic to the Neotropical region, which are distributed mainly in the Amazon, Orinoco and Paraná-Paraguay basins [1, 2], with lower representativeness in the São Francisco River basin and introductions reported in coastal basins [3]. Despite the occurrence of some species in environments such as rapids, these fishes typically inhabit lakes and floodplains [4]. It is the fourth most diverse family within Characiformes, with 101 valid species, distributed in 16 extant genera [5–7], with more than 70 species occurring in the Amazon basin [8]. These fishes have different feeding habits, and can be frugivorous, herbivorous, piscivorous, and lepidophagous (consumer of scales) [9–11].

Family monophyly is supported by numerous morphological synapomorphies, such as the presence of a ventral keel, composed of spines derived from modified abdominal scales, an anteriorly oriented predorsal spine, only absent in *Colossoma* Eigenmann & Kennedy 1903, *Mylossoma* Eigenmann & Kennedy 1903, and *Piaractus* Eigenmann 1903, and by the presence of interlocking teeth in premaxilla and dentary [12, 13]. Serrasalmidae has also been recovered as monophyletic in molecular phylogenies proposals [14–17].

The evolutionary relationships within the family have been the subject of several studies [15, 18–20]. Commonly the species have been recovered and grouped into three large clades: (i) the first, composed by genera lacking the predorsal spine; (ii) the second, comprising *Acnodon* Eigenmann 1903, *Mylesinus* Valenciennes 1850, *Myleus* Müller & Troschel 1844, *Myloplus* Gill 1896, *Ossubtus* Jégu 1992, *Tometes* Valenciennes 1850, and *Utiaritichthys* Miranda Ribeiro 1937; and (iii) the third, represented by *Catoprion* Müller & Troschel 1844, *Metynnis* Cope 1878, *Pristobrycon* Eigenmann 1915, *Pygocentrus* Müller & Troschel 1844, *Pygopristis* Müller & Troschel 1844, and *Serrasalmus* Lacepède 1803 [15, 19]. Kolmann et al. [20] named these clades as subfamilies Colossomatinae, Myleinae, and Serrasalminae, respectively.

However, the hypothesis of the intrafamilial relationships within Serrasalmidae using of Ultraconserved Elements (UCEs), included all genera and the greatest number of species, and proposed a slightly different classification [6]. The family is presently composed by two major clades: (i) Colossomatinae, represented by species lacking a predorsal spine, and (ii) Serrasalminae, with species having a predorsal spine, and divided into Myleini (pacus) and Serrasalmini (*Metynnis* plus piranhas). Despite these advances, the monophyly of some genera was rejected, as *Myleus*, *Myloplus*, *Pristobrycon*, and *Tometes* [6, 20].

Similarly, studies using the gene cytochrome c oxidase subunit I (COI) revealed that the diversity within Serrasalmidae is still underestimated. In the Brazilian Amazon, various lineages may represent new species, especially concerning *Myloplus* and *Serrasalmus* [21]. Two new species were recently described using the same marker, *Catoprion absconditus* Mateussi, Melo & Oliveira 2020 and *Myloplus nigrolineatus* Ota, Machado, Andrade, Collins, Farias & Hrbek 2020 [6, 7]. In the basins of the Paraná-Paraguay and Tocantins rivers, a species complex was detected in *Serrasalmus maculatus* Kner 1858 [22] and five lineages were recognized in *Pygocentrus nattereri* Kner 1858 in different river basins [23]. This underestimation of diversity is a consequence of the difficulty in identifying many species and their intra- and interspecific limits, due to the variation in body shape, sexual dimorphism, ontogeny [1, 4, 7, 24], and water color in Amazon basin [7, 25]. In addition, we highlight the scarcity of recent taxonomic revisions, with identification keys, of species-rich genera, such as *Metynnis*, *Myloplus*, and *Serrasalmus*.

From cytogenetic point of view, the greatest diversity is primarily related to diploid chromosome number (2n) and intra-and interspecific variations in the karyotype formula. In *Ser-rasalmus*, for example, three karyomorphs of *S. rhombeus* Linnaeus 1766 occurring in syntopy were found, which varied both in 2n and in the karyotype formula [26, 27]. Different

karyomorphs were also observed in specimens of *S. maculatus* (described as *S. spilopleura* Kner 1858) from the Paraná-Paraguay basin [28] and the Amazon basin [29, 30].

Significant advances in cytogenetic studies in Serrasalmidae utilizing different approaches also occurred, ranging from chromosomal characterization of species, such as *Pygocentrus cariba* Humboldt and Valenciennes 1821 [31] and *Myleus micans* Lütken 1875 [32], to the use of cytogenetic markers to identify hybrids between *Colossoma* and *Piaractus* [33, 34]. Recently, the presence of a B chromosome restricted to females was described to *Metynnis lippincottianus* (Cope 1870) [35].

The 2n in the family varies from 54 chromosomes in *Colossoma* [36, 37], *Mylossoma* [38], and *Piaractus* [33, 38] to 62—in *Metynnis* [35, 39, 40]. Intermediate 2n, such as 58 and 60 chromosomes, were already reported in *Pygocentrus* and *Serrasalmus* [26, 28, 29, 37, 41–43]. However, little is known about this number within Myleini and other species of *Metynnis* (the first genus to diverge within Serrasalmini). Given the chromosomal diversity observed in Serrasalmidae, this study aimed: (i) to analyze cytogenetically species of Myleini and *Metynnis*, and (ii) using modern molecular phylogenies as a framework, propose a general pattern of chromosomal evolution within Serrasalmidae.

Material and methods

In the present study, we analyzed a total of 39 specimens from eight species and three genera (<u>Table 1</u>). The sampling of specimens was authorized by Instituto Brasileiro do Meio Ambiente e Recursos Naturais Renováveis (IBAMA, License No. 28095–1) and the experimental procedure was approved by the Comitê de Ética na Utilização de Animais (CEUA) at Instituto Nacional de Pesquisas da Amazônia (INPA) (Approval No. 027/2017).

Cell suspensions were obtained from renal tissue, according to the protocol of [44]. Cbanding was based on the [45] protocol, with minor modifications and stained with propidium iodide [46]. The extraction of total DNA was from muscle tissue, using the Wizard® Genomic DNA Purification Kit (Promega), according to the manufacturer's guidelines. The extracted DNA was quantified in 1.5% agarose gel with NanoVueTM Plus (GE Healthcare). The sequences of 18S rDNA, 5S rDNA, and telomeric sequence were amplified by polymerase chain reaction (PCR), using the following primers: 18Sf (5'-CCG CTT TGG TGA CTC TTG AT-3') and 18Sr (5'-CCG AGGACC TCA CTA AAC CA-3') [47]; 5Sa (5'-TAC GCC CGA TCT CGT CCG ATC-3') and 5Sb (5' - CAGGCT GGT ATG GCC GTA AGC-3') [48]; and (TTAGGG)₅ and (CCCTAA)₅ [49]. Fluorescent in situ hybridization (FISH) was performed according to the protocol described by [50], but with modifications. The slides were denatured in 70% formamide/2xSSC at 70°C, and the spreads were dehydrated in an increasing ethanol series (70, 85 and 100%), for 5 min at each concentration. Subsequently, 20 μL of the hybridization mixture (100 ng of each probe, 50% deionized formamide, 20xSSC, and 10% dextran sulphate) was dropped onto each slide, and the mixture was hybridized at 37°C for 24 h in a moist chamber containing distilled water. The chromosomes were counterstained with DAPI (1.2 µg/mL) and mounted in antifade solution (Vector, Burlingame, CA, USA). The PCR products of the 18S rDNA gene and telomeric sequence were labelled by nick translation with digoxigenin 11-dUTP (Dig-Nick Translation mix; Roche) and 5S rDNA was labelled with biotin-14-dATP (Biotin Nick Translation mix; Roche), following the manufacturer's instructions. The detection of the hybridization signals was performed with anti-digoxigenin -rhodamine (Roche Applied Science) for the 18S rDNA probes and the telomeric sequence, and with streptavidin (Sigma-Aldrich) for the 5S rDNA probes. Subsequently, the chromosomes were counterstained with DAPI, analyzed in an Olympus BX51 fluorescence microscope and classified according to [51]. The 5S and 18S rDNA sequences were obtained from the eight species present in Table 1, with addition to published data from [35].

Table 1. Species and number of individuals analyzed, with their respective sampling locations and voucher number.

Species	Number of individuals Males Females	Location	Voucher
Metynnis cuiaba Pavanelli, Ota & Petry 2009	0 1	Negro River (Anavilhanas Archipelago), AM 2°37'28.5"S, 60° 58'16.8"W	INPA-ICT 59049
Metynnis hypsauchen (Müller & Troschel 1844)	3 6	Uatumã River (Balbina Hydroelectric Dam), AM 1°55'02.2"S 59°28'23.7"W	INPA-ICT 59050
Metynnis longipinnis Zarske & Géry 2008	1 2	Negro River (Anavilhanas Archipelago), AM 2°37'28.5"S, 60° 58'16.8"W	INPA-ICT 59051
Myloplus asterias* (Müller & Troschel 1844)	1 1	Apeu floodplain, Castanhal (PA)	INPA-ICT 59052
Myloplus lobatus (Valenciennes 1850)	3 0	Catalão Lake, AM 2°33'28.4"S, 60°46'29.7"W	INPA-ICT 59056
Myloplus rubripinnis* (Müller & Troschel 1844)	4 4	Guamá River, Belém (PA)	INPA-ICT 59053
Myloplus schomburgkii* (Jardine 1841)	9 3	Xingu River, Altamira (PA)	INPA-ICT 59054
Tometes camunani* Andrade, Giarrizzo & Jégu 2013	1 0	Belém (PA)	INPA-ICT 59055

AM = Amazonas state; PA = Pará state. / (*) Lacking precise geographic coordinates.

https://doi.org/10.1371/journal.pone.0258003.t001

For analysis of the evolution of the chromosome number in a phylogenetic context (adapted from [6]), data obtained from the eight species analyzed in this study were added to karyotype information already available in the literature *i.e.* [26, 29, 32, 33, 35, 37, 38, 42, 43].

Results

The four species of the genus *Myloplus* and *T. camunani* have 2n = 58 chromosomes, while the species of *Metynnis* 2n = 62 chromosomes (Figs $\underline{1}$ and $\underline{2}$). The karyotype formulas and/or fundamental number (FN) are species-specific (Table 2).

The distribution of constitutive heterochromatin occurred mainly in the pericentromeric regions of all chromosomes of all analyzed species, in addition to the presence of blocks in terminal portions of some chromosomes (Figs 1 and 2). Some species also had entire heterochromatic short arms in some pairs, such as pair 12 of *T. camunani* (Fig 1B), pairs 7, 13, 14, and the proximal region of pairs 19 and 27 of *My. asterias* (Fig 1D), pairs 2, 5, 18, and 28 of *My. schomburgkii* (Fig 1F), pairs 19, 20, 23, 26, and 29 of *My. lobatus* (Fig 1H), pair 11 of *My. rubripinnis* (Fig 1J), pairs 21 and 29 *Me. cuiaba* (Fig 2D), and the pairs 1, 2, 14, 17, 21 of *Me. hypsauchen* (Fig 2F). Also, heteromorphism of heterochromatic block size was observed in pair 2 of *My. lobatus* (Fig 1H) and pair 24 of *Me. cuiaba* (Fig 2D).

The mapping of the 5S rDNA sequences showed only one pair with signal in the pericentromeric portion for the *Tometes* and *Myloplus* species, which were the pair 2 of *T. camunani* (Fig 3), 14 of *My. asterias* (Fig 3), and *My. rubripinnis* (Fig 3), 19 from *My. schomburgkii* (Fig 3) and 5 from *My. lobatus* (Fig 3). In relation to the 18S rDNA, the signals were found in three chromosomal pairs in four species: *T. camunani*, terminal portion, short arm, pairs 22, 25, and 27 (Fig 3), *My. asterias*, interstitial region of the short arm of pair 13, terminal region of the long arm of par 19, and pericentromeric portion of par 25 (Fig 3), *My. lobatus*, interstitial region of the short arm of pairs 2, 8, 22 (Fig 3), and *My. rubripinnis*, terminal region in pairs 20, 22, 24 (Fig 3). However, in *My. schomburgkii*, signals were seen in the interstitial region of the short arms of pairs 2, 3, 5, and 21 (Fig 3).

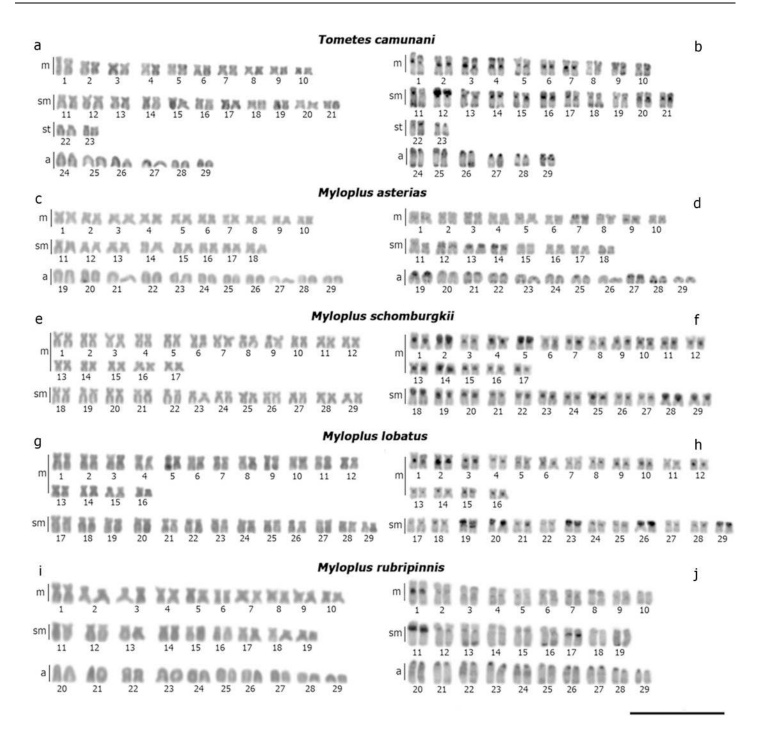


Fig 1. Karyotypes of the species with 2n = 58, stained with Giemsa (left column) and with C band (right column): (a, b) *Tometes camunani*; (c, d) *Myloplus asterias*; (e, f) *My. schomburgkii*; (g, h) *My. lobatus*; (i, j) *My. rubripinnis*. Scale bar = $10 \mu m$.

For *Metynnis*, the three species have synteny of the 18S and 5S rDNA in the interstitial portion of the long arms of the pairs 3 in *Me. longipinnis*, 29 in *Me. cuiaba*, and 10 in *Me. hypsauchen* (Fig 4). In *Me. longipinnis*, in addition to the aforementioned 5S marker, we detected another one in terminal portion of the short arm in pair 31 (Fig 4). Regarding the sites of the 18S rDNA, they were also observed in interstitial portion of the long arms of pair 23 in *Me*.

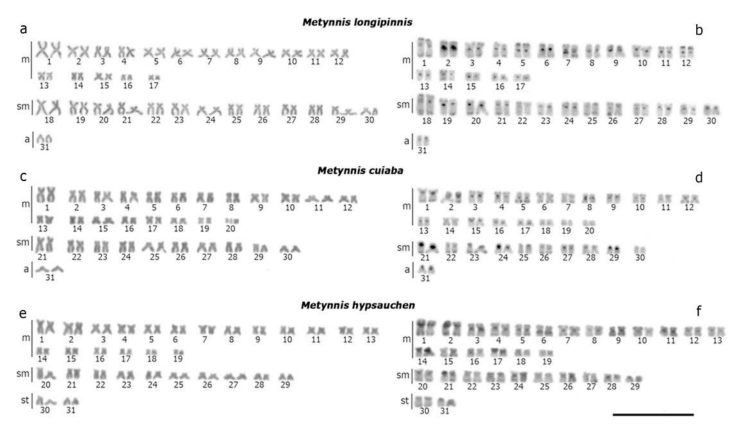


Fig 2. Karyotypes of Metynnis longipinnis (a, b); Me. cuiaba (c, d); Me. hypsauchen (e, f), stained with Giemsa (a, c, e) and C band (b, d, f). Scale bar = 10 μm.

longipinnis and *Me. hypsauchen* (Fig 4) and in the pericentromeric region in pair 8 in *Me. cuiaba* (Fig 4). Telomeric sequences were only detected in the terminal portions of the chromosomes in all species (S1 Fig).

The chromosome number within the main clades of Serrasalmidae was relatively stable. In Colossomatinae all species have 2n = 54, in Myleini: *Mylesinus*, *Myleus*, *Myloplus*, and *Tometes* possess 2n = 58 chromosomes, and in Serrasalmini: the species of *Serrasalmus* and *Pygocentrus* have 2n = 60 chromosomes (Fig 5).

	respective diploid and fundamental numbers, karyotype formula, and 18S and 5S rDNA position.
--	--

Species	2n	FN	KF	188	58
Metynnis cuiaba	62	122	34m+26sm+2a	8p; 29p	29p
Metynnis longipinnis	62	122	40m+20sm+2a	3p; 23q	3p; 31p
Metynnis hypsauchen	62	120	38m+20sm+4a	10p; 23q	10p; 23q
Myloplus asterias	58	94	20m+16sm+22a	13p; 19q; 25q	14q
Myloplus schomburgkii	58	116	34m+24sm	2p; 3p; 5p; 21p	19q
Myloplus lobatus	58	116	32m+26sm	2p; 8p; 22p	5q
Myloplus rubripinnis	58	96	20m+18sm+20a	20p; 22p; 24p	14q
Tometes camunani	58	104	20m+22sm+4st+12a	22p; 25p; 27p	2q

(2n = diploid number; FN = fundamental number; KF = karyotype formula); m = metacentric; sm = submetacentric; st = subtelocentric; a = acrocentric; p = short arm; q = long arm).

https://doi.org/10.1371/journal.pone.0258003.t002

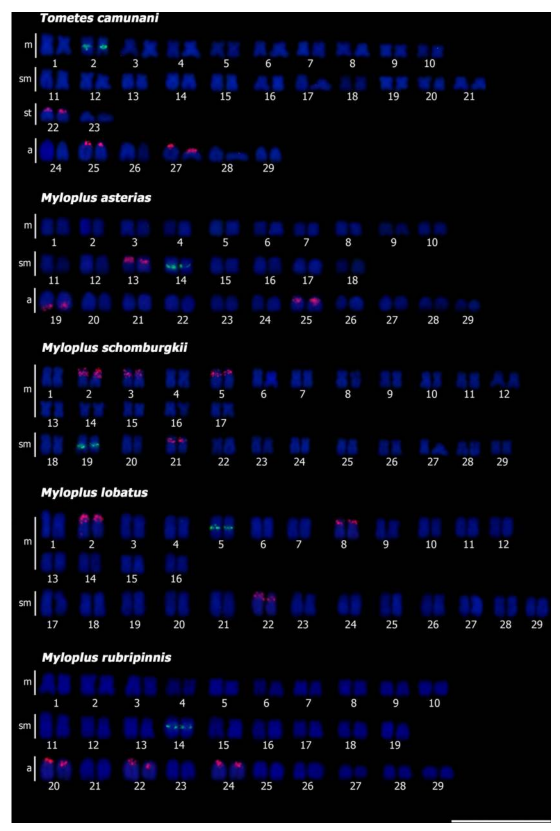


Fig 3. Karyotype of the five species analyzed by Double-FISH with 18S rDNA (red) and 5S rDNA (green) and counterstained with DAPI: (a) *Tometes camunani*; (b) *Myloplus asterias*; (c) *My. schomburgkii*; (d) *My. lobatus*; (e) *My. rubripinnis*. Scale bar = $10 \mu m$.

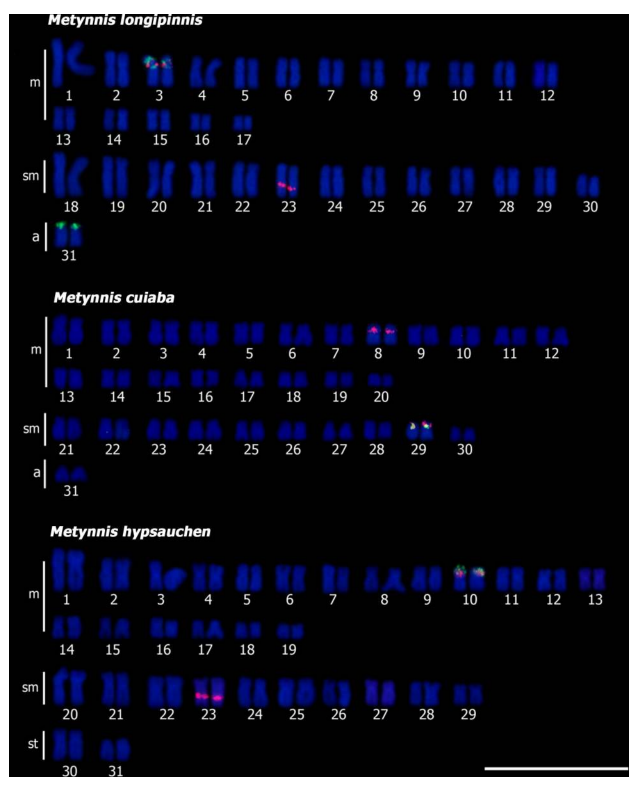


Fig 4. Double-FISH in *Metynnis*: (a) *Me. longipinnis*; (b) *Me. cuiaba*; (c) *Me. hypsauchen*, 18S rDNA (red); 5S rDNA (green) and counterstained with DAPI. Scale bar = $10 \mu m$.

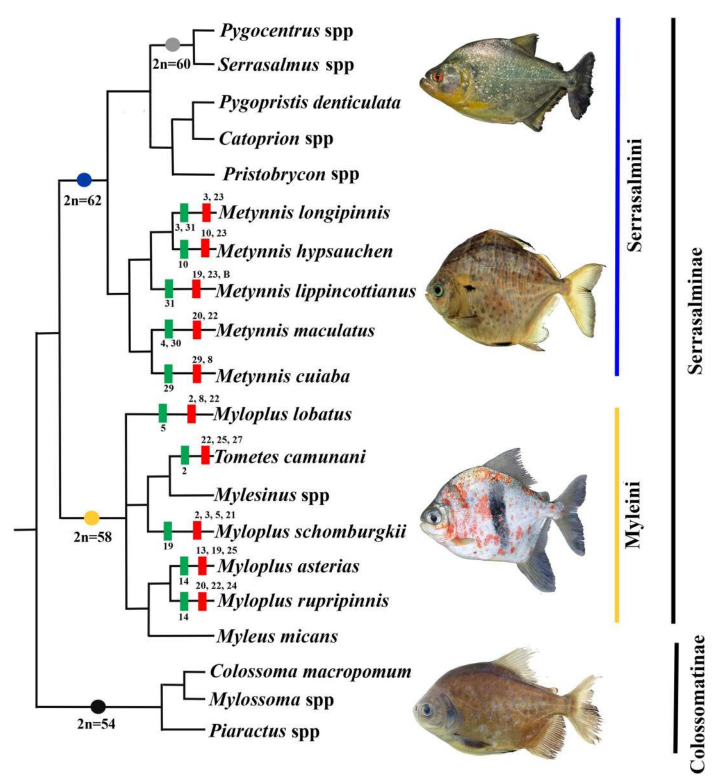


Fig 5. Cladogram adapted from Mateussi *et al.* [6], indicating the variation of the diploid number between the species and genera with available cytogenetic information, with the probable ancestral diploid number of the clades indicated by circles of different colors (2n = 54 black, 2n = 58 yellow, 2n = 60 gray, 2n = 62 blue). Data for *Pygocentrus* spp. and *Serrasalmus* spp. [26, 29, 37, 42, 43], and for *Metynnis* [35]. Data for Colossomatinae [33, 38]; and for *Myleus micans* [32]. Numbers indicate chromosomal pairs with 18S rDNA (red) and 5S rDNA (green).

Discussion

The present study contributes to the knowledge about Serrasalmidae cytogenetics, mainly concerning Myleini, including for the first time species of the genera *Myloplus* and *Tometes*; and within Serrasalmini expanding to five the number of *Metynnis* species with chromosomal information. *Metynnis* was the first genus to diverge within Serrasalmini [6] and is essential to understand the evolution of remaining members of the tribe. The karyotype formulas and/or fundamental number were species-specific for the eight species, which can represent informative data to be used in combination with molecular markers to better understand the phylogenetic relationship in intergeneric level, especially concerning paraphyletic genera as *Myleus*, *Myloplus*, *Tometes*, and *Pristobrycon*.

We observed the presence of large heterochromatic blocks in the pericentromeric and terminal regions of some chromosomes in Myleini and in analyzed species of *Metynnis*. This pattern was already reported for other species of Myleini [33, 38] and Serrasalmini [26, 29, 35, 42]. The presence of fully heterochromatic short arms, as in *Myloplus schomburgkii* and *My. lobatus*, was also observed in *Serrasalmus compressus* Jégu, Leão & Santos 1991 and *S. elongatus* Kner 1858 [42] and is possibly related to interspecific variations in *Myloplus*. These heterochromatic blocks and arms may indicate chromosomal rearrangements such as the Robertsonian or non-R rearrangements, which can cause changes in 2n or not [52].

In the species analyzed in this study, some heterochromatic blocks are associated with ribosomal sequences mapped. In Serrasalmidae, this association was previously described in different species of Serrasalmus [37, 43], in Colossoma macropomum Cuvier 1816 and Piaractus mesopotamicus Holmberg 1887 [38], and in Myleus micans (Lütken 1875) [32]. In the case of piranhas, as Serrasalmus rhombeus, this association directly influenced the differentiation among karyomorphs [26], where the chromosomes, mainly subtelocentric/acrocentric, with 18S ribosomal cistrons, are C-band positive. These chromosomes are apparently those that underwent rearrangements, causing variations in the 2n and karyotype formulas, revealing that heterochromatin may be generating points that are susceptible to chromosomal breaks and contributing to the karyotype evolution of the group [26, 37].

The location pattern of the 18S and 5S rDNA sites registered in *Myloplus* and *Tometes* species resembles that already described in the family, with single 5S and multiple 18S labeling [37, 43]. To date, all analyzed *Serrasalmus* species also presented this 5S rDNA localization pattern. This pattern of ribosomal sequences can be used as a taxonomic marker, since both sites provide unique markers for each species analyzed. In addition, 5S rDNA was already reported as a relevant cytogenetic marker within the family, in which all species of *Serrasalmus* analyzed had this site in the interstitial region of pair 7 [37, 43]. Therefore, we recommend 5S rDNA to be used in integrative taxonomy approach of the family, along with DNA barcoding that is already being greatly employed in the past years [e.g. 6, 7, 53]. Despite the advances in species description of Serrasalmidae, 14 new taxa in the last decade [5], none of them included a *Serrasalmus* discovery, even with the several evidences of new species using COI [e.g. 21, 22].

On the other hand, the synteny between the 18S and 5S sequences, observed in the *Metynnis* species, had not yet been reported, and can be considered an unprecedented characteristic for the family. In addition to the syntenic pair, one additional chromosome pair presented 18S rDNA sites, coinciding with the number of markers for *Me. maculatus* (Kner 1858) and *Me. lippincottianus* [35], two very similar looking congeners, but not closely related according to molecular phylogenies [e.g. 6, 20]. The co-location of these cistrons in the three species seems to indicate that this condition is maintained and being propagated within *Metynnis*, suggesting some adaptive advantage for maintaining this organization in the genera, as suggested for some genera of Julidini (Perciformes) [54].

The data from the rDNA show an apparent conservation and organization within the clades/genera of the family. The 5S rDNA sequence is observed in pericentromeric portions in two pairs in *C. macropomum*, *Mylossoma* spp., and *Piaractus mesopotamicus* (Colossomatinae) [38], while in *Myleus micans*, *Myloplus*, and *Tometes* (Myleini), *Pygocentrus* and *Serrasalmus* (Serrasalmini) only one meta/submetacentric pair is a marker of this ribosomal site [32, 35, 37, 43]. On the other hand, for 18S rDNA an increase in the number of sites of the basal clade towards the derived clade can be observed. In Myleini, the species have 1 to 3 pairs of 18S bearers [33, 38, this study]. While in Serrasalmini all species of *Serrasalmus* and *Pygocentrus* have at least five pairs carrying these sequences, with a clear predominance in acrocentric chromosomes [37, 43].

In general, there is a similarity in the number of sites of the ribosomal sequences in each clade, which suggests that there is conservation of the chromosomal structure in Serrasalmidae. Although the mapping of telomeric sequences did not show any rearrangement, the variations in relation to the number and location of rDNA sequences between species, may indicate that these sequences have an evolutionary independence [55], between or within the genera of Serrasalmidae. As for example, in *Metynnis* with the presence of synteny in the three analyzed species, and with the homeology of the 5S rDNA in pair 7 of the *Serrasalmus* species [37, 43].

This conservation is also observed regarding the diploid chromosome number. The comparison to the phylogenetic relationship proposals of Mateussi et al. [6] and Kolman et al. [20], we observe that in Colossomatinae, all species have 2n = 54 [33, 38]. Within Serrasalminae, diploid chromosome number increased, the tribe Myleini (or Myleinae *sensu* Kolman et al. [20]) presented 2n = 58 (*Myleus*, *Myloplus*, and *Tometes*). In Serrasalmini (or Serrasalminae *sensu* Kolman et al. [20]), *Metynnis* species have 2n = 62 chromosomes, while, *Serrasalmus* and *Pygocentrus* have 2n = 60 chromosomes [26, 29, 37, 42, 43]. Previous hypotheses proposed that 2n = 54 chromosomes is the ancestral number of the family, with a tendency to increase the 2n number from 54 to 62 chromosomes [26, 29, 35, 38, 42]. The increase in diploid number would have occurred through chromosomal fission, since in Colossomatinae all chromosomes have two arms (meta and submetacentric type) while in the derived Serrasalmini there are some pairs of chromosomes with only one arm (acrocentric) [37, 38].

The increase in chromosome number in the tribe Myleini, along with a greater amount of chromosomes of subtelocentric/acrocentric types, is a condition that was already reported in different fish families. In Curimatidae, for example, most analyzed species have 2n conserved, equal to 54 meta/submetacentric chromosomes [52, 56-59]. It is interesting to note that the conservation of the diploid number (2n = 54) [17] can be considered a synapomorphy of Curimatoidea [60-64]. The only families of this clade, for which 2n diverged from 54 are Serrasalmidae [26, 32, 33, 37, 38, 67] and Curimatidae (e.g. Potamorhina latior, P. altamazonica and P. squamoralevis) [26, 33, 38, 56, 65-67], however they have 2n = 54 present in the species of the clades that first diverged.

In spite of the indication that the chromosomal fissions, coupled with the emergence of acrocentric chromosomes, are associated with an increase in the diploid number in Serrasal-midae, this change did not occur in a linear path from Myleini to Serrasalmini, since Myloplus asterias, My. rubrippinis, and T. camunani have 2n = 58, and several acrocentric chromosomes, while in Metynnis, the highest chromosomal number (2n = 62) is observed, and a larger number of the meta- and submetacentric, and fewer acrocentric chromosomes. This suggests that, in addition to fission, other rearrangements, such as fusions, translocations and pericentric inversions, were involved in the evolution of these species and modified the 2n and karyotype formulas among the clades. These rearrangements occurred in a dynamic and complex way, independently in the different clades, since each of them has unique characteristics, as the synteny present in Metynnis and the homeology of the 58 rDNA pair in Serrasalmus.

Therefore, the chromosomal macrostructure of the Serrasalmidae species is conserved within the main clades, with higher variation in Serrasalmini. This fact makes the family a very interesting group to study, because the different karyotype formulas and locations of ribosomal sequences, recorded in some species can be used as cytotaxonomic markers and assist in the identification of species, given the difficulty and taxonomic uncertainties that still persist in Serrasalmidae, despite all these advances. Furthermore, the diversity of chromosomal markers highlights the importance of integrating cytogenetic studies with systematic studies, whether they are morphological or molecular. The expansion of both chromosome studies and the number of localities sampled would contribute further to confirm the evolutionary process that occurred in Serrasalmidae and also to corroborate the diversity of species in the different clades.

Supporting information

S1 Fig. Metaphases of eight species of Serrasalmidae analyzed by FISH using a telomeric probe and counterstained with DAPI. (a) T. camunani; (b) M. asterias; (c) M. schomburgkii; (d) M. lobatus; (e) M. rubripinnis; (f) M. longipinnis; (g) M. altidorsalis; (h) M. hypsauchen. Scale bar = $10 \mu m$. (TIF)

Author Contributions

Conceptualization: Ramon Marin Favarato, Jorge Ivan Rebelo Porto, Celeste Mutuko Nakayama, Rafaela Priscila Ota, Eliana Feldberg.

Data curation: Ramon Marin Favarato, Rafaela Priscila Ota.

Formal analysis: Ramon Marin Favarato. Funding acquisition: Eliana Feldberg.

Investigation: Ramon Marin Favarato, Rafaela Priscila Ota.

Methodology: Ramon Marin Favarato, Alber Campos, Eliana Feldberg.

Project administration: Eliana Feldberg.

Resources: Eliana Feldberg.

Supervision: Ramon Marin Favarato, Rafaela Priscila Ota, Eliana Feldberg.

Visualization: Ramon Marin Favarato, Leila Braga Ribeiro, Rafaela Priscila Ota, Eliana Feldberg.

Writing - original draft: Ramon Marin Favarato, Leila Braga Ribeiro, Eliana Feldberg.

Writing - review & editing: Rafaela Priscila Ota, Eliana Feldberg.

References

- Jégu M. Subfamily Serrasalminae (Pacus and piranhas). In: Reis RE, Kullander SO, Ferraris CJ Jr, editors. Check list of the freshwater fishes of South and Central America. Porto Alegre, Brasil: Editora da Pontifícia Universidade Católica, 2003. pp. 182–196.
- Nico LG, Jégu M, Andrade MC. Family Serrasalmidae—Piranhas and Pacus. In: van der Sleen P, Albert JS, editors. Field guide to the fishes of the Amazon, Orinoco, and Guianas (Princeton Field Guides). Princeton: Princeton University Press, 2018. pp. 172–96.

- Britski HA, Sato Y, Rosa ABS. Manual de Identificação de Peixes da Região de Três Marias (Com Chaves de Identificação para os Peixes da Bacia do São Francisco). 3rd ed. Brasília, Brazil: Câmara Dos Deputados/CODEVASF; 1988.
- Ota RP, Röpke CP, Zuanon JAS, Jégu M. Serrasalmidae. In: Queiroz LJ, Torrente-Vilara G, Ohara WM, Pires TH, Zuanon JAS, Dória C, editors. Peixes do rio Madeira, volume II, a ictiofauna do rio Madeira, Dialeto Latin American Documentary. 1st ed. São Paulo, Brasil: 2013 pp. 15-47.
- Fricke R, Eschmeyer WN, Fong JD. Eschmeyer's Catalog of Fishes: Species by Family/Subfamily. 2020. Available from: Electronic version accessed 7.16.2020. (http://researcharchive.calacademy.org/ research/ichthyology/catalog/SpeciesByFamily.asp).
- Mateussi NTB, Melo BF, Ota RP, Roxo FF, Ochoad LE, Foresti F. Phylogenomics of the Neotropical fish family Serrasalmidae with a novel intrafamilial classification (Teleostei: Characiformes). Mol Phylogenetics Evol 2020; 153. https://doi.org/10.1016/j.ympev.2020.106945 PMID: 32861778
- Ota RP, Machado VN, Andrade MC, Collins RA, Farias IP, Hrbek T. Integrative taxonomy reveals a new species of pacu (Characiformes: Serrasalmidae: Myloplus) from the Brazilian Amazon. Neotrop Ichthyol 2020; 18(1):e190112. https://doi.org/10.1590/1982-0224-20190112.
- Dagosta FCP, De Pinna M. The fishes of the Amazon: distribution and biogeographical patterns, with a comprehensive list of species. Bull. Am. Mus. Nat. Hist. 2019; 431:1–163. https://doi.org/10.1206/ 0003-0090.431.1.1.
- Goulding M. The fishes and the forest exploration in Amazonian natural history. Berkeley: University of California press; 1980.
- 10. Sazima I, Machado FA. Underwater observations of piranhas in western Brazil. Environ Biol Fishes 1990; 28:17-31. https://doi.org/10.1007/BF00751026
- Correa SB. Winemiller KO. Lopez-Fernandez H. Galetti PM Jr. Evolutionary perspectives on seed consumption and dispersal by fishes. BioScience 2007; 57:748-756. https://doi.org/10.1641/B570907
- Machado-Allison A. Estudios sobre la sistemática de la subfamília Serrasalminae (Teleostei, Characidae). Parte II. Discussion sobre la condicion monofiletica de la subfamília. Acta Biol Venez 1983; 11 (4):145-195.
- Kolmann MA, Cohen KE, Bemis KE, Summers AP, Irish FJ, Hernandez LP. Tooth and consequences: Heterodonty and dental replacement in piranhas and pacus (Serrasalmidae). Evolution & Development 2019; e12306. https://doi.org/10.1111/ede.12306 PMID: 31449734
- Calcagnotto D, Schaefer SA, De Salle R. Relationships among Characiformes fishes inferred from analysis of nuclear and mitochondrial gene sequences. Mol Phylogenet Evol 2005; 36:135-153. https://doi. org/10.1016/j.ympev.2005.01.004 PMID: 15904862
- Thompson AW, Betancur-R R, López-Fernández H, Ortí G. A time-calibrated, multi-locus phylogeny of piranhas and pacus (Characiformes: Serrasalmidae) and a comparison of species tree methods. Mol Phylogenet Evol 2014; 81:242-257. https://doi.org/10.1016/j.ympev.2014.06.018 PMID: 25261120
- Arcila D, Ortí G, Vari RP, Armbruster JW, Stiassny MLJ, Ko KD, et al. Genome- wide interrogation advances resolution of recalcitrant groups in the tree of life. Nat Ecol Evol 2017; 1(2): 0020. https://doi. org/10.1038/s41559-016-0020 PMID: 28812610
- Betancur-R R, Arcila D, Vari RP, Hughes LC, Oliveira C, Sabaj MH, et al. Phylogenomic incongruence, hypothesis testing, and taxonomic sampling: The monophyly of characiform fishes. Evolution 2018; 73 (2):329-345. https://doi.org/10.1111/evo.13649 PMID: 30426469
- Ortí G, Petry P, Porto JIR, Jégu M, Meyer A. Patterns of nucleotide change in mitochondrial ribosomal RNA genes and the phylogeny of piranhas. J Mol Evol 1996; 42:169-182. https://doi.org/10.1007/ BF02198843 PMID: 8919869
- Ortí G, Sivasundari A, Dietz K, Jégu M. Phylogeny of the Serrasalmidae (Characiformes) based on mitochondrial DNA sequences. Genet Mol Biol 2008; 31:343-351. http://doi.org/10.1590/S1415-
- Kolmann MA, Hughes LC, Hernandez LP, Arcila D, Betancur-R R, Sabai MH, et al. Phylogenomics of piranhas and pacus (Serrasalmidae) uncovers how dietary convergence and parallelism obfuscate traditional morphological taxonomy. Systemat Biol 2021; 70(3):576-592. https://doi.org/10.1093/sysbio/ syaa065 PMID: 32785670
- Machado VN, Collins RA, Ota RP, Andrade MC, Farias IP, Hrbek T. One thousand DNA barcodes of piranhas and pacus reveal geographic structure and unrecognised diversity in the Amazon. Sci Rep 2018; 8. https://doi.org/10.1038/s41598-018-26550-x PMID: 29849152
- Bignotto TS, Maniglia TC, Gomes VN, Oliveira IJ, Agostinho CS, Prioli SMAP. Genetic evidence for a species complex within the piranha Serrasalmus maculatus (Characiformes, Serrasalmidae) from three Neotropical river basins based on mitochondrial DNA sequences. Genet Mol Biol 2020; 43:e20190131. https://doi.org/10.1590/1678-4685-GMB-2018-0131 PMID: 31454404

- Mateussi NTB, Melo BF, Foresti F, Oliveira C. Molecular data reveal multiple lineages in piranhas of the 23. genus Pygocentrus (Teleostei, Characiformes). Genes 2019; 10:371. https://doi.org/10.3390/ genes10050371.
- Zarske A, Géry J. Revision der neotropischen Gattung Metynnis Cope, 1878. Beschreibung zweier neuer Arten und zum Status von Metynnis goeldii Eigenmann, 1903 (Teleostei: Characiformes: Serrasalmidae). Vertebr Zool 2008; 58:173-196.
- Ota RP, Py-Daniel LHR, Jégu M. A new Silver Dollar species of *Metynnis* Cope, 1878 (Characiformes: Serrasalmidae) from Northwestern Brazil and Southern Venezuela. Neotropical Ichthyology (Online) 2016; 14:12-30. http://doi.org/10.1590/1982-0224-20160023
- Nakayama CM, Jégu M, Porto JIR, Feldberg E. Karyological evidence for a cryptic species of piranha within Serrasalmus rhombeus group (Characidae, Serrasalminae) in the Amazon. Copeia 2001; 3:866-869. http://doi.org/10.1643/0045-8511(2001)001[0866:KEFACS]2.0.CO;2
- Teixeira AS, Nakayama CM, Porto JIR, Feldberg E. Esterase- D and chromosome patterns in Central Amazon piranha (Serrasalmus rhombeus Linnaeus, 1766) from lake Catalão. Genet Mol Biol 2006; 29 (3):498-502. http://doi.org/10.1590/S1415-47572006000300018.
- Cestari MM, Galetti PM Jr. Chromosome studies of Serrasalmus spilopleura (Characidae, Serrasalminae) from the Paraná-Paraguay River: evolutionary and cytotaxonomic considerations. Copeia 1992a; 1:108-112.
- Nakayama CM. Porto JIR. Feldberg E. Ocorrência de dois citótipos em Serrasalmus spilopleura Kner. 1958 (Characiformes, Serrasalmidae) na região de confluência dos rios Negro e Solimões, Amazonas, Brasil. Acta Amazonica 2000; 30(1):149-154.
- Centofante L, Porto JIR, Feldberg E. Chromosomal polymorphism in Serrasalmus spilopleura Kner, 30. 1858 (Characidae, Serrasalminae) from central Amazon basin. Caryologia 2002; 55:37-45. http://doi. org/10.1080/00087114.2002.10589256.
- Gaviria JI, Nirchio M, Granado A, Estrada A. Karyotype and nucleolar organizer regions of Pygocentrus cariba (Serrasalminae) from Caicara Del Orinoco, Venezuela. Interciencia 2005; 30(1): 44-47.
- Moreira-Peres AW, Bertollo LAC, Moreira Filho O. Karyotypic characterization of Myleus micans (Lütken, 1875) (Pisces, Characidae, Serrasalminae). Caryologia 2006; 59(2):125-130. http://doi.org/10. 1080/00087114.2006.10797907.
- Nirchio M, Fenocchio AS, Swarça AC, Pérez JE, Granado A, Estrada A, et al. Cytogenetic characterization of hybrids offspring between Colossoma macropomum (CUVIER, 1818) and Piaractus brachypomus (CUVIER, 1817) from Caicara del Orinoco, Venezuela. Caryologia 2003; 56(4):405-411. http:// doi.org/10.1080/00087114.2003.10589351.
- Ribeiro LB, Moraes Neto A, Artoni RF, Matoso DA, Feldberg E. Chromosomal mapping of repetitive sequences (Rex3, Rex6, and rDNA genes) in hybrids between Colossoma macropomum (Cuvier, 1818) and Piaractus mesopotamicus (Holmberg, 1887). Zebrafish 2017; 2:155-160. http://doi.org/10. 1089/zeb.2016.1378.
- Favarato RM, Ribeiro LB, Ota RP, Nakayama CM, Feldberg E. Cytogenetic characterization of two 35. Metynnis species (Characiformes, Serrasalmidae) reveals B chromosomes restricted to the females. Cytogenet Genome Res 2019; 158(1):38-45. https://doi.org/10.1159/000499954 PMID: 31079097
- Almeida-Toledo LF, Foresti F, Toledo-Filho SA, Bernardino G, Ferrari VA, Alcantara RCF. Cytogentic 36. studies in Colossoma mitrei, C. macropomum and their interspecific hybrid. In: Tiews K editors, Selection, Hybridization and Genetic Engineering in Aquaculture 1987. pp. 189-195.
- Nakayama CM, Feldberg E, Bertollo LAC. Karyotype differentiation and cytotaxonomic considerations 37. in species of Serrasalmidae (Characiformes) from the Amazon basin. Neotrop Ichthyol 2012; 10(1):53-58. http://doi.org/10.1590/S1679-62252012000100005.
- 38. Ribeiro LB, Matoso DA, Feldberg E. Chromosome mapping of repetitive sequences in four Serrasalmidae species (Characiformes). Genet Mol Biol 2014; 37:46-53. https://doi.org/10.1590/s1415-47572014000100009 PMID: 24688290
- Arefiev VA. Chromosome sets of four characid fish species (Teleostei, Characidae). Zool Zh 1989; 68 39. (5): 82-91.
- Baroni S, Lopes CE, Almeida-Toledo LF. Cytogenetic characterization of Metynnis maculatus (Teleos-40. tei, Characiformes): the description in Serrasalminae of a small B chromosome bearing inactive NORlike sequences. Caryologia 2009; 62:95-101. http://doi.org/10.1080/00087114.2004.10589674.
- Cestari MM, Galetti PM Jr. Chromosome evolution in the genus Serrasalmus and citotaxonomic considerations about Serrasalminae (Characidae, Pisces). Brazilian Journal of Genetics 1992b; 15(3):555-567.

- Nakayama CM, Porto JIR, Feldberg E. A comparative cytogenetic study of five piranha species (Serrasalmus, Serrasalminae) from the Amazon basin. Genetica 2002; 114:231-236. https://doi.org/10.1023/ a:1016275505655 PMID: 12206361
- Nakayama CM, Feldberg E, Bertollo LAC. Mapping of ribosomal genes and chromosomal markers in three species of the genus Serrasalmus (Characidae, Serrasalminae) from the Amazon basin. Genet Mol Biol 2008; 31(4): 868-873. https://doi.org/10.1590/S1415-47572008005000018
- Gold JR, Li YC, Shipley NS, Powers PK. Improved methods for working with fish chromosomes with a review of metaphase chromosome banding. J Fish Biol 1990; 37:563-575. https://doi.org/10.1111/j. 1095-8649.1990.tb05889.x
- Sumner AT. A simple technique for demonstrating centromeric heterochromatin. Expe Cell Res 1972; 75:304-306. https://doi.org/10.1016/0014-4827(72)90558-7 PMID: 4117921
- Lui RL, Blanco DR, Moreira-Filho O, Margarido VP. Propidium iodide for making heterochromatin more evident in the C-banding technique. Biotech Histochem 2012; 87(7):433-8. https://doi.org/10.3109/ 10520295.2012.696700 PMID: 22747174
- Gross MC, Schneider CH, Valente GT, Martins C, Feldberg E. Variability of 18S rDNA locus among Symphysodon fishes: chromosomal rearrangements. J Fish Biol 2010; 76:1117–1127. https://doi.org/ 10.1111/j.1095-8649.2010.02550.x PMID: 20409165
- Martins C, Galetti PM Jr. Chromosomal localization of 5S rDNA genes in Leporinus fish (Anostomidae, Characiformes). Chromosome Res 1999; 7:363-367. https://doi.org/10.1023/a:1009216030316 PMID: 10515211
- 49. Ijdo JW, Wells RA, Baldini A, Reeders ST. Improved telomere detection using a telomere repeat probe (TTAGGG)n generated by PCR. Nucleic Acids Res 1991; 19:4780. https://doi.org/10.1093/nar/19.17. 4780 PMID: 1891373
- 50. Pinkel D, Straume T, Gray JW. Cytogenetic analysis using quantitative, high sensitivity, fluorescence hybridization. P Natl Acad Sci USA 1986; 83:2934-2938. https://doi.org/10.1073/pnas.83.9.2934
- Levan A, Fredga K, Sandberg AA. Nomenclature for centromeric position on chromosomes. Hereditas 51. 1964: 52:201-220
- Pinheiro VSDS, Carvalho NDM, do Carmo EJ, Schneider CH, Feldberg E, Gross MC. Karyoevolution in Potamorhina (Cope, 1878) (Ostariophysi, Curimatidae): Using repetitive DNA for the elucidation of genome organization. Zebrafish 2016; 13:118-131. https://doi.org/10.1089/zeb.2015.1187 PMID: 26840804
- Andrade MC, Machado VN, Jégu M, Farias IP, Giarrizzo T. A new species of Tometes Valenciennes 53. 1850 (Characiformes: Serrasalmidae) from Tocantins-Araquaia River Basin based on integrative analysis of molecular and morphological data. PLoS One 2017; 12(4):e0170053. https://doi.org/10.1371/ journal.pone.0170053 PMID: 28422969
- 54 Amorim KDJ, Cioffi MB, Bertollo LAC, Soares RX, de Souza AS, Costa GWWF, et al. Co-located 18S/ 5S rDNA arrays: an ancient and unusual chromosomal trait in Julidini species (Labridae, Perciformes). Comp Cytogenet 2016; 10:555–570. https://doi.org/10.3897/CompCytogen.v10i4.10227 PMID: 28123678
- Calado LL, Bertollo LAC, Costa GWWF, Molina WF. Cytogenetic studies of Atlantic mojarras (Perciformes—Gerreidae): chromosomal mapping of 5S and 18S ribosomal genes using double FISH. Aquac Res 2012; 44:829-835. https://doi.org/10.1111/j.1365-2109.2012.03089.x
- Feldberg E, Porto JIR, Bertollo LAC. Karyotype evolution in Curimatidae (Teleostei, Characiformes) of the Amazon region. I. Studies on the genera Curimata, Psectrogaster, Steindachnerina and Curimatella. Brazil J Genet 1992; 15:369-383.
- Venere PC, Souza IL, Silva LKS, dos Anios MB, de Oliveira RR, Galetti PM Jr, Recent chromosome 57. diversification in the evolutionary radiation of the freshwater fish family Curimatidae (Characiformes). J Fish Biol 2008; 72:1976–1989. http://doi:10.1111/j.1095-8649.2008.01814.x
- 58. Sampaio TR, Gravena W, Gouveia JG, Giuliano-Caetano L, Dias AL. B microchromosomes in the family Curimatidae (Characiformes): mitotic and meiotic behavior. Comp Cytogenet 2011; 5:301-313. https://doi.org/10.3897/CompCytogen.v5i4.1650 PMID: 24260637
- Sampaio TR, Pires LB, Venturelli NB, Usso MC, da Rosa R, Dias AL. Evolutionary trends in the family 59. Curimatidae (Characiformes): Inferences from chromosome banding. Comp Cytogenet 2016; 10:77-95. https://doi.org/10.3897/CompCytogen.v10i1.6316 PMID: 27186339
- Porto JIR, Feldberg E, Falcão JN, Nakayama CM. Cytogenetic studies in Hemiodidae (Ostariophysi, 60. Characiformes) fishes from the Central Amazon. Cytologia 1993; 58:397-402.
- Pastori MC. Roncati H. Aichino DR. Ledesma MA. Fenocchio AS. Chromosome characterization and cytotaxonomic considerations on Characidae, Acestrorhynchidae and Cynodontidae (Pisces,

- Characiformes) from the Paraná River (Argentina). Caryologia 2009; 1(32):69–74. http://doi.org/10.1080/00087114.2004.10589668.
- 62. Traldi JB, Vicari MR, Martinez J, Blanco DR, Lui RL, Moreira-Filho O. Chromosome Analyses of Apareiodon argenteus and Apareiodon davisi (Characiformes, Parodontidae): An Extensive Chromosomal Polymorphism of 45S and 5S Ribosomal DNAs. Zebrafish 2016; 13:19–25. https://doi.org/10.1089/zeb.2015.1124 PMID: 26625282
- 63. Barros LC, Galetti PM Jr., Feldberg E. Mapping 45S and 5S ribosomal genes in chromosomes of Anostomidae fish species (Ostariophysi, Characiformes) from different Amazonian water types. Hydrobiologia 2016; 789(1):77–89. https://doi.org/10.1007/s10750-015-2583-8
- **64.** Pinheiro-Figliuolo VS, Goll L, Ferreira VP, Feldberg E, Gross MC. First record on sex chromosomes in a species of the Family Cynodontidae: *Cynodon gibbus* (Agassiz, 1829). Cytogenet Genome Res 2020; 160:29–37. https://doi.org/10.1159/000505889 PMID: 32092757
- 65. Feldberg E, Porto JIR, Bertollo LAC, Nakayama CM. Karyotype evolution in Curimatidae (Teleostei, Characiformes) from the amazon region. II. Centric fissions in the genus *Potamorhina*. Genome 1993; 36:372–376. https://doi.org/10.1139/g93-051 PMID: 18469994
- **66.** Navarrete MC, Julio-Junior HF. Cytogenetic analysis of four curimatids from the Paraguay basin, Brazil (Pisces: Characiformes: Curimatidae). Cytologia 1997; 62:241–247.
- **67.** Nirchio M, Rossi AR, Foresti F, Oliveira C. Chromosome evolution in fishes: a new challenging proposal from Neotropical species. Neotrop Ichthyol 2014; 12:761–770.