Research article

The B chromosomes of the African cichlid fish *Haplochromis obliquidens* harbour 18S rRNA gene copies

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Abstract

**Background:** Diverse plant and animal species have B chromosomes, also known as accessory, extra or supernumerary chromosomes. Despite being widely distributed among different taxa, the genomic nature and genetic behavior of B chromosomes are still poorly understood.

**Results:** In this study we describe the occurrence of B chromosomes in the African cichlid fish *Haplochromis obliquidens*. One or two large B chromosome(s) occurring in 39.6% of the analyzed individuals (both male and female) were identified. To better characterize the karyotype and assess the nature of the B chromosomes, fluorescence in situ hybridization (FISH) was performed using probes for telomeric DNA repeats, 18S and 5S rRNA genes, SATA centromeric satellites, and bacterial artificial chromosomes (BACs) enriched in repeated DNA sequences. The B chromosomes are enriched in repeated DNAs, especially non-active 18S rRNA gene-like sequences.

**Conclusion:** Our results suggest that the B chromosome could have originated from rDNA bearing subtelo/acrocentric A chromosomes through formation of an isochromosome, or by accumulation of repeated DNAs and rRNA gene-like sequences in a small proto-B chromosome derived from the A complement.

Background

Perciformes fishes are the largest order of vertebrates and include the family Cichlidae, which represents the most species-rich vertebrate family, with more than 3,000 species [1]. Cichlid fishes are native to tropical areas of the Americas, Africa and Asia and have undergone spectacular adaptive radiation in the lakes of East Africa (Victoria, Tanganyka, and Malawi) [2]. This group of fishes is an important model for evolutionary processes, mostly because their great phenotypic diversity has evolved in a short period of time, among lineages with similar genomes [3-5]. In addition, some species of Cichlidae, such as the tilapiines *Oreochromis*, *Tilapia* and *Sarotherodon*, are very important for aquaculture [6]. Because of their economic and evolutionary importance, cichlid fishes have been the focus of several genetic and genomic studies [7]. On the other hand, most cytogenetic studies in the group have been directed to *Oreochromis niloticus* and few data are available for other cichlid species. Cytogenetic data have shown that most cichlids from Central and South America have 48 chromosomes, while African species have 44 chromosomes (reviewed in [8]), with some characteristic chromosome pairs within each group.

Many animal and plant species have B chromosomes, also known as accessory, extra or supernumerary chromo-
somes. B chromosomes are additional and mostly dispensable chromosomes that are present in some individuals in some species. Most of these chromosomes probably arose from A chromosomes, but have followed their own evolutionary pathways [9]. Furthermore, B chromosomes have irregular and non-Mendelian modes of inheritance and are not believed to undergo recombination with any members of the basic A chromosome set [10]. They are mostly heterochromatic [11] and may have originated through chromosomal breakage and fusion of A chromosome(s) in a single species or by hybridization between species [12].

The occurrence of B chromosomes has been described for species of diverse fish groups. In general, the B chromosomes of fishes vary in number from one to eight [13-15]. The size of the B chromosomes can vary greatly, from macro-chromosomes as found in Astyanax sp. [16-20] and Alb Loaderus alb furnus [21,22], medium-sized chromosomes, as in Rhamdia sp. [23,24], small-sized chromosomes, as in Parachinicterus galleatus [25], or microchromosomes, as in Prochilodus sp. [13,15]. Among cichlids, B chromosomes have been described for only a few species of South American species. They were first described in the male germ cells of Gymnogeophagus balzani [26] and later found in Geophagus brasilienis, Cichlasoma paranaensis and Cren- dicichla niederleinii [27]. B microchromosomes have also been described for Cichla monocus, Cichla sp. and Cren- dicichla reticulata [28]. The presence of B chromosomes among the African cichlids has not previously been reported.

Despite being distributed in several fish taxa, the genomic nature and genetic behavior of B chromosomes are still poorly understood. In this study, we performed molecular cytogenetic analysis on the African cichlid fish H. obliqui- dens, in order to better understand the chromosomal organization and meiotic behavior of B chromosomes present in individuals of this species.

**Results**

**Basic cytogenetic analysis**

Approximately 30 metaphase spreads were analyzed per specimen to determine the diploid chromosome number and karyotype structure. H. obliqui- dens exhibited diploid chromosome numbers of 2n = 44 (12 m/sm and 32 st/a), without sex-related heteromorphism (Figure 1a). Two outstanding large pairs of chromosomes (pairs 1 and 7, Figure 1) are characteristic. One or two large metacentric B chromosomes (Figure 1b, c) were present in 38 out of 96 analyzed specimens (Table 1). In these specimens the B chromosomes were present in 100% of the analyzed cells.

Active NORs were identified in the short arms of five chromosomes in the A complement, but were absent in the B chromosomes (Figure 2a). Constitutive heterochromatin was detected in the pericentromeric regions of most A chromosomes, in the short arm of the chromosome pair 1, as well as on the entire length of B chromosomes (Figure 2b, c).

**Cytogenetic mapping of repeated DNAs**

The 5S rDNA probe generated in situ hybridization signals in the centromeric area of 15 acrocentric chromosomes, including the chromosome pairs 1 and 7, not in the B chromosomes (Figure 2d). The 18S rRNA gene probe identified six sites in the short arms of small st/a chromosomes of the A complement. In addition, 18S rDNA clusters were observed in both telomeric regions and pericentromeric areas of the B chromosomes (Figure 2e).

Chromosomal mapping with the clone BAC-C4E09 from the O. niloticus genomic library, which contains LINE-like retrotransposons and satellite DNAs [29], labeled the short arms of two pairs of st/a chromosomes of the A complement (one of them being the largest chromosome pair), and evidenced weak signals along the whole extension and a strong cluster in the pericentromeric region of the B chromosomes (Figure 2f). Chromosomal mapping
with the clone BAC-C5E01, which also contains repetitive elements of the O. niloticus genome, evidenced strong signals in the short arms of some st/a chromosomes, in almost the whole extent of the largest chromosome of the A complement, and in the centromeric region of the B chromosomes (Figure 2g).

Chromosomal mapping with the SATA-satellite DNA isolated from O. niloticus labeled the centromeric region of all A chromosomes, but showed no clearly visible signal in the centromeric region of the B chromosomes (Figure 2h), although a faint signal was detected in the centromeric region of few metaphase B chromosomes (Figure 2h).

Chromosome mapping with the telomeric (GGGTTA)\textsubscript{n} probe labeled all the telomeres, including the telomeres of B chromosomes, and no interstitial signals were observed in either A and B chromosomes (Figure 2i).

**Meiotic analysis**

Meiotic chromosomes of mature male H. obliquidens with one or two B chromosomes were analyzed. Approximately 10 meiotic plates were analyzed per specimen to determine the haploid chromosome number and meiotic configuration. Chromosomes were Giemsa-stained, C-banded and in situ hybridized with the 18S rRNA gene probe (Figure 3). As expected in diakinoses, 22 bivalents resembling the 44 A chromosomes and one or two univalent B chromosome(s) were observed (Figure 3a). Although it was difficult to identify the B chromosome(s) among the A elements in the Giemsa-stained meiotic spreads, they were clearly distinguished by C-banding because they appeared totally heterochromatic (Figure 3b). The hybridization of 18S rDNA in diakinetic meiotic spreads was faintly detected in the centromeric region of the B chromosomes (Figure 3a).

**Table 1: B chromosome distribution in the analyzed H. obliquidens stock.**

| Number of B chromosomes | Total sample | Male | Female | Not sexed | % of total sample analyzed |
|-------------------------|-------------|------|--------|-----------|----------------------------|
| 0                       | 58          | 31   | 17     | 10        | 60.41%                     |
| 1                       | 30          | 10   | 13     | 07        | 31.25%                     |
| 2                       | 08          | 05   | 03     | -         | 8.33%                      |

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**Accumulation of repetitive DNAs in B chromosomes**

Heterochromatic B chromosomes as observed in H. obliquidens are common among fish species [13,30,31]. However, some cases of euchromatic B chromosomes have also been described in fish, such as Moenkhausia sanctaefilomenae [32] and Characidium cf. zebra [31]. The heterochromatin of the B chromosomes of H. obliquidens is enriched in repetitive DNAs, as evidenced by mapping with 18S rDNA, and the clones BAC-C4E09 and BAC-C5E01. These BACs contain repetitive DNAs with similarity to the LINE retrotransposons of the cichlids Haplochromis microlepis and O. niloticus, SATB satellite DNA of O. niloticus, and repeated segments of the Danio rerio genome [29]. On the other hand, the large number of 5S rDNA sites observed in H. obliquidens suggest that most centromeric heterochromatin contains repetitive 5S rRNA gene-like copies that are not clustered in the Bs. Repetitive satellite DNA originated from 5S rDNA was previously described in the fish Hoplias malabaricus [33]. The organization and evolution of tandem repetitive DNAs is governed by particular patterns of evolution such as unequal exchange, transposition, RNA-mediated transposition and gene conversion causing the origin of variant copies of rDNA repeats in the genome. These mechanisms could be responsible for the dispersion and accumulation of copies of 5S rDNA repeats in specific areas of the genome (reviewed in [33]). The genomic structure of 5S rDNA
repeats should be better investigated in the genome of *H. obliquidens* to clarify the origin and possible role of such dispersed pattern.

The presence of repeated sequences with similarities to retrotransposons in a B chromosome has been previously reported for the fish *Astyanax scabripinis* [34], as well as for the B chromosome of the parasitoid wasp *Nasonia vitripennis* [35,36]. A retrotransposon also appears to be involved in the transposition of chloroplast DNA into the repeated element Bd49 of the B chromosomes of *Brachycome dichromosomatica* [37]. The repeated elements are...
thought to accumulate in the heterochromatic regions of chromosomes, which are characterized by lower gene density and reduced recombination [38]. The absence of recombination of B chromosomes of *H. obliquidens*, as shown by the univalent conformation that they assume during meiosis, may facilitate the accumulation of repeated DNAs. Repetitive DNAs were long considered to be junk DNA because they had no clearly identified function [39,40]. However, further studies of their accumulation in specific genomic areas, causing chromosome rearrangements through breakages, deletions, inversions and amplifications [41,42], have shown they can operate at the chromosomal level in the speciation process. These recent findings have suggested that junk DNA may be one of the major drivers of genome evolution [43]. Consequently, the study of repetitive DNA in B chromosomes could yield new insights into the evolutionary mechanisms involved with the origin and evolution of particular chromosome elements.

**Presence of 18S rRNA genes copies in B chromosomes**

Another remarkable characteristic described here is the presence of 18S rRNA gene copies in both telomeric regions and centromeric areas of B chromosomes. The presence of rDNA sequences in the B chromosome suggests its possible origin from A chromosomal elements that harbor clusters of 18S rRNA genes. It is believed that extra rDNA copies could initially increase transcription rates and can present an advantage to the bearers of the B chromosome (reviewed in [12]). On the other hand, the 18S rDNA sites in the B chromosomes of *H. obliquidens* appeared not stained by silver nitrate, suggesting that they are not actually being transcribed, as silver has a strong affinity for the proteins that are assembled during nucleolus formation [44]. The presence of non-transcribed rRNA genes was described in the B chromosomes of the black rat *Rattus rattus* [45]. In this case, the B chromosomes seem to have arisen from A chromosomes that harbour rRNA genes and was subsequently invaded by repetitive sequences with the dispersion of rRNA genes throughout the B chromosome [45]. On the other hand, active rRNA genes were observed in the telomeric regions of B chromosome of the rodents *Akodon montensis* and *Oryzomys angouya* [46]. In *H. obliquidens* the rRNA genes accumulated in the centromeric and terminal regions of the B chromosomes and the repeated DNAs are dispersed throughout the entire B chromosomes with enrichment in the centromeric area. Considering that no silver signal was detected on the Bs of any cell of *H. obliquidens* we can hypothesize that the rRNA genes in the B chromosome are permanently inactive representing variant copies (pseudogenes) of the true rRNA genes.

Previous studies have found evidence of dispersed or clustered elements similar to 45S rDNA in several eukaryotic genomes. These elements have mainly been characterized as noncoding, small-unit tandem repeats of variable copy number. Such elements have been identified in various eukaryotic species, including yeast [47], animals [48], and plants [49]. Moreover, some authors have hypothesized that ribosomal cistrons could undergo active transposition followed by tandem amplification. This transposition event could be facilitated by recombination between preexisting repetitive DNAs in the constitutive heterochromatins and/or rDNA clusters. Repeated DNAs seem to be associated with ribosomal DNAs in *A. scabripinnis*, facilitating recombination between terminal segments in the same chromosome pair or between terminal portions from non-homologous chromosomes [34,50]. The results presented here for *H. obliquidens* and previously observed for other vertebrates [45,46] show rRNA gene-like copies may have originated from the true rRNA genes and further accumulated in the B chromosomes through the same mechanism that drives the evolution of repeated DNAs. These mechanisms could have been involved in the distribution of rDNA copies in B chromosomes of *H. obliquidens*. Moreover, the regions of chromosomes with repetitive DNAs, including 18S rDNA copies, are among the last regions of the genome to segregate during cell division, as previously reported in yeast [51,52], and may therefore be involved in the non-disjunction and accumulation of the B chromosome in *H. obliquidens* cells.
Possible origin and evolution of the B chromosomes

The meiotic behavior of the B chromosomes show they do not associate with any of the A chromosomes and do not form true bivalents in the cell when two Bs are present during prophase I of meiosis. The two Bs appeared slightly associated in metaphasic images of a few cells, probably as consequence of the similarity in their repetitive DNA content. The chromosomes appeared to associate themselves through the rDNA sequences in their terminal regions. Indeed, this slight association was rarely seen in the majority of analyzed meiosis plates, and it could be favored due to the accumulation of the 18S rDNA repeats.

The presence of repeated DNAs (including rRNA gene copies) in specific chromosomes of the A complement suggests that Bs could have arisen from genomic segments of such A elements. This proto-B chromosome(s) would then be freed from the selection pressures that act on the maintenance of standard chromosomes and could have increased of size due to the accumulation of repetitive DNAs in the absence of recombination.

Methods

Specimens, and chromosome preparation and banding

We analyzed ninety-six specimens of *H. obliquidens* (46 males, 33 females and 17 of undetermined sex) obtained from the aquarium trade in Botucatu, SP, Brazil (Table 1). The animal experiments were performed with the approval of the appropriate ethics committee of UNESP - São Paulo State University. Metaphase chromosomes were obtained from cells of the anterior kidney following in vivo treatment with colchicine at 0.025% (1 ml/100 g of body weight) according to the air-drying method [59]. Heterochromatin was identified by C-banding [60], and the nucleolus organizer regions (NORs) were visualized by silver nitrate staining [61]. Chromosomes were classified as meta/submetacentric (m/sm) and subtelo/acrocentric (st/a), and were organized by decreasing order of size in the karyotype. Meiotic cells from testes were obtained as described by Kligerman and Bloom [62].

Fluorescence in situ hybridization (FISH)

Fluorescence in situ hybridization (FISH) was performed to map repeated DNAs on the mitotic and meiotic chromosomes of *H. obliquidens*. Five DNA probes containing sequences of different classes of repeated DNA were used for chromosome hybridization. (i) 5S rDNA probe: complete repeat units of 5S rDNA of *H. obliquidens* were obtained by the polymerase chain reaction (PCR) with the primers 5SA (5’-TAC GGC CGA TCT CCT CCG ATC - 3’) and 5SB (5’ - CAG GCT GGT ATG GCC GTA AGC-3’) designed from the rainbow trout 5S rRNA sequence [63] and successfully applied for the amplification of 5S rDNA of other cichlids [64,65]. (ii) 18S rDNA probe: a segment of 1,400 base pairs (bp) of the 18S rRNA gene of *H. obliquidens* was obtained by PCR with the primers 18SF 5’CCG CIT TGG TGA CTC TTG AT and18SR 5’CCG AGG ACC TCA CTA AAC CA. The 18S primers were designed from the catfish *Ictalurus punctatus* (GenBank accession number AF021880) and have been successfully used to amplify 18S rRNA genes of different fish species [65,66]. (iii) SATE satellite: repeated satellite DNA isolated and cloned from *O. niloticus* [29]; (iv) Telomeric DNA sequences: in vitro synthesized oligomers of telomeric repeats (GGGTTA)n (TAACCC)n; (v) Clones BAC-C4E09 and BAC-C5E01: Bacterial artificial chromosomes containing several classes of repeated elements from the *O. niloticus* genome [29].
Probes were labeled by nick translation with biotin 14-dATP (Bionick labeling system-Invitrogen). After denaturation of chromosomal DNA in 70% formamide/2× SSC for 40 seconds at 70°C, hybridization mixtures containing 100 ng of denatured probe, 10 mg/ml dextran sulfate, 2× SSC and 50% formamide, in a final volume of 30 μl, were dropped on the slides and the hybridization was performed overnight at 37°C in a 2× SSC moist chamber. Post-hybridization washes were carried out at 45°C in 2× SSC/50% formamide for 15 min, followed by a second wash in 2× SSC for 15 min, and a final wash at room temperature in 4× SSC for 15 min. Detection of hybridized probes was carried out with 0.07% avidin FITC conjugate (Sigma) in C buffer (0.1 M NaHCO3/0.15 M NaCl) for 1 h, followed by two rounds of signal amplification using 2.5% anti-avidin biotin conjugate (Sigma) in blocking buffer (1.26% NaHCO3, 0.018% sodium citrate, 0.0386% Triton X-100 an 1% non-fat dried milk) for 30 min. Each treatment with anti-avidin biotin conjugate was followed by a treatment with avidin-FITC. The treatments with avidin-FITC and anti-avidin-biotin were conducted in a 2× SSC moist chamber at 37°C. After each amplification step, the slides were washed three times for 5 min each in blocking buffer at 42°C. Chromosomes were counterstained with propidium iodide diluted in antifade (Vectashield Mounting Medium, Vector). Hybridized chromosomes were visualized using an Olympus BX 61 microscope, and images were captured with a digital camera Olympus DP71 with the software Image-Pro MC 6.0.

**Authors’ contributions**

ABP carried out the cytogenetic analyses and drafted the manuscript. IAF helped in cytogenetic analysis and drafted the manuscript. CM designed and coordinated the study, and drafted and revised the manuscript. All authors read and approved the final manuscript.

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