Uniparental Genome Elimination in Australian Carp Gudgeons

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Abstract

Metazoans usually reproduce sexually, blending the unique identity of parental genomes for the next generation through functional crossing-over and recombination in meiosis. However, some metazoan lineages have evolved reproductive systems where offspring are either full (clonal) or partial (hemiclonal) genetic replicas. In the latter group, the process of uniparental genome elimination selectively eliminates either the maternal or paternal genome from germ cells, and only one parental genome is selected for transmission. Although fairly common in plants, hybridogenesis (i.e., clonal haploidization via chromosome elimination) remains a poorly understood process in animals. Here, we explore the proximal cytogenomic mechanisms of somatic and germ cell chromosomes in sexual and hybrid genotypes of Australian carp gudgeons (*Hypseleotris*) by tracing the fate of each set during mitosis (in somatic tissues) and meiosis (in gonads). Our comparative study of diploid hybrid and sexual individuals revealed visually functional gonads in male and female hybrid genotypes and generally high karyotype variability, although the number of chromosome arms remains constant. Our results delivered direct evidence for classic hybridogenesis as a reproductive mode in carp gudgeons. Two parental sets with integral structure in the hybrid soma (the F1 constitution) contrasted with uniparental chromosomal inheritance detected in gonads. The inheritance mode happens through premeiotic genome duplication of the parental genome to be transmitted, whereas the second parental genome is likely gradually eliminated already in juvenile individuals. The role of metacentric chromosomes in hybrid evolution is also discussed.

Key words: genome elimination, hemiclone, hybridogenesis, unisexual, gametogenesis, Hypseleotris.

Introduction

The great majority of metazoan organisms reproduce through sex in a two-stage process, commencing with meiosis in each parent and culminating in the fusion of one ploidy-reduced, genetically shuffled gamete from each parent into a single zygote (Crow and Kimura 1965; Kondrashov 1988). In sexual reproduction, both paternal and maternal genomes enter meiosis. In meiosis I, the parental chromosomes pair, and genetic material undergo regular DNA-recombination. The paired chromosomes are then segregated to the opposite poles of the meiotic spindle, whereas in meiosis II, sister chromatids disjoin (Petronczki et al. 2003; Suwa and Yamashita 2007). The resulting haploid gametes contain a unique cocktail of recombinant chromosomal DNA from the mother and father. In this way reproducing animals we call "sexual." Although the meiotic molecular machinery is highly conserved

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Significance

Most animals reproduce sexually, via biparental fusion of gametes resulting from the well-studied process of meiosis. However, only little is known about meiotic behavior of chromosomes and the extent of genetic recombination in animal groups that practice "asexual" reproduction. Here, we demonstrate that male and female hybrids of Australian carp gudgeons (*Hypseleotris*) are fertile and display uniparental chromosomal elimination, without genetic recombination, both as predicted by nuclear DNA markers. We show that genome elimination occurs premeiotically in juveniles, prior to gonad maturation. Asexual vertebrates are important animal models in research fields as diverse as speciation, toxicology, and unorthodox reproduction, for example, understanding how vertebrates clone their germ cells. These findings further enhance the potential of carp gudgeons in this regard.

among Metazoans (Bernstein and Bernstein 2010), gametogenesis has been repeatedly modified in a small proportion of organisms, giving rise to clonal, or asexual, taxa.

To transfer a whole intact genome from multicellular animal to next generation in a gamete, those taxa exploit a wide range of cytogenetic mechanisms ranging from processes completely without meiosis (apomixis) to those with modified meiosis (automixis) (Stenberg and Saura 2009; Lenormand et al. 2016). Hybridogenesis is a reproduction mode unique in its ability to transmit not a whole, but one-half of a somatic genome via genome elimination. The process of uniparental genome elimination selectively eliminates either the maternal or paternal genome from germ cells (Gardner and Ross 2014; Dedukh, Riumin, et al. 2020). During uniparental genome elimination, only one parental genome is thus selected for transmission; it then undergoes chromosome doubling and enters meiosis (Tunner and Heppich 1981; Heppich et al. 1982; Dedukh et al. 2015). In contrast to standard Mendelian inheritance, the resulting haploid gametes are clones (in the absence of mutation) of the gametes produced in the preceding generation (Doležálková-Kaštánková et al. 2018). Haploidization via chromosome elimination is a fairly common phenomenon in plants, with basic research leading to applications in accelerated plant breeding (Sanei et al. 2011; Comai 2014; Ishii et al. 2016). However, it remains a poorly understood process in animal ontogenesis.

Classic hybridogenesis, commonly also referred to as hemiclonal inheritance, has been demonstrated in a small range of organisms from various taxonomic groups, such as fishes from the genera *Poeciliopsis* (Schultz 1961), *Squalius* (Carmona et al. 1997), or *Hexagrammos* (Kimura-Kawaguchi et al. 2014; Munehara et al. 2016), frogs from the *Pelophylax esculentus* complex (Tunner 2009), and stick insects in the genus *Bacillus* (Mantovani and Scali 1992). In addition, it has also been inferred in several other groups, most notably *Hypseleotris* fishes (Bertozzi et al. 2000; Schmidt et al. 2011) but full confirmation ideally relies upon detailed chromosomal investigation.

Advancements in molecular cytogenomics offer a spectrum of molecular tools allowing for new empirical insights into uniparental genome elimination. Research in various animal models has identified that DNA elimination mostly occurs before the initiation of meiosis (Cimino 1972: Ogielska 1994; Scali et al. 2003; Schön et al. 2009; Stenberg and Saura 2009; Dedukh et al. 2015, 2017; Dedukh, Riumin, et al. 2020). As shown in Pelophylax water frogs, DNA elimination seems to be a gradual process of individual chromosomes enclosed within micronuclei (Chmielewska et al. 2018; Dedukh, Riumin, et al. 2020). To allow the noneliminated chromosome set (the one to be transmitted) to enter effectively into meiosis, doubling of a haploid parental chromosome set must occur through 1) premeiotic genome endoreplication (cell cycle without mitosis), or 2) endomitosis (mitosis without chromosome segregation) (Dedukh, Riumin, et al. 2020). However, studies of insects and triploid vertebrates have shown that elimination of the uniparental chromosomal set may take place during meiosis as well (Zhang et al. 1998; Nabais et al. 2012; Gardner and Ross 2014). The inclusion of a wider concept of hybridogenesis into uniparental genome transmissions extends the number of mechanistic processes, from which genome elimination may be absent (Doležálková et al. 2016; Lavanchy and Schwander 2019). Therefore, to understand these processes, comprehensive and case-specific studies are needed to demonstrate any discordance in genomic content between a zygote, soma, and germ cells, and to identify pathways of elimination and transmission of genomes.

Carp gudgeons (*Hypseleotris*, Eleotridae) are a genus of small fishes with a widespread distribution across the Indo-Pacific, including moderate diversification in Australian freshwater environments (Thacker and Unmack 2005). Eastern Australia contains a species complex consisting of two described species (*Hypseleotris klunzingeri* and *Hypseleotris galii*), and several undescribed species (Hoese et al. 1980; Unmack 2000). In addition to four sexually reproducing species, Bertozzi et al. (2000) described the co-occurrence of three apparent F₁ hybrid forms derived from three distinctive taxa revealed in the lower Murray River and first suggested these were unisexual carp gudgeons. Subsequently, Schmidt et al. (2011) used microsatellite markers and proposed the occurrence of male and female hybrid lineages and sexual species. More

recently, Unmack et al. (2019) discovered one of the missing sexual parental species using SNPs. However, despite these population analyses and the demonstrated hemiclonal nature of several lineages (Unmack et al. 2019), it remains unclear what enables the persistence of this reproductive system from a cellular perspective.

In this article, we explore the proximal cytogenomic mechanisms maintaining the carp gudgeon's hybrid genotypes as F₁'s. We analyzed somatic and germ cell chromosomes, as well as gonadal microanatomy in both hybrid and sexual individuals. Using genomic in situ hybridization (GISH) with species-specific probes, we identified parental chromosomal sets in various hybrids and traced the destiny of each set during mitosis (in somatic tissues) and meiosis (in gonads). We specifically tested whether 1) chromosomal behavior supports a hypothesis of uniparental genome elimination and whether 2) reproduction mode of hybrids is linked with cytological principles of classic hybridogenesis.

Materials and Methods

Study Species

For a clarity, here we use the term "sexual," or "sexual species," for taxa of male and female individuals that use the regular meiotic (sexual) cycle. We also use the term "hybrid" for male and/or female individuals having parental chromosomal sets in their soma from the extant sexual species, and displaying uniparental chromosomal elimination in their reproduction cycle. Five sexual species have been recognized in eastern Australia: two, Hypseleotris compressa and H. klunzingeri frequently co-occur with the sexual/unisexual complex, but there is no record of them being involved in any hybridization. The sexual/unisexual complex consists of three sexual species, H. galii, H. sp. Midgley's, and H. sp. Bald, which have traditionally been identified in previous papers (Bertozzi et al. 2000; Schmidt et al. 2011; Unmack et al. 2019) by the codes HA, HB, and HX, respectively, which we use from this point forward in this article. Each of the three known interspecific diploid hybrid genotypes have the F1 genomic combinations designated as HA×HB, HA×HX, and HB×HX. With the exception of HX, which has an extremely restricted distribution (Unmack et al. 2019), the other species and hybrids in the sexual/unisexual complex are widespread across the Murray–Darling Basin (1,059,000 km²), in addition HB and HB×HX are present in the Bulloo River $(75,610 \text{ km}^2)$ and Cooper Creek catchments (298,000 km²) and HA×HB is present in coastal rivers from at least the Clarence River north to Waterpark Creek (110,000 km²). There is a strong sex bias in some hybrids, with HA×HB being strongly male biased (Schmidt et al. 2011 recorded 80 males, 2 females plus 5 indeterminate), HA×HX was strongly female biased (Schmidt et al. 2011 recorded 7 males, 152 females, and 9 indeterminate). While the sex ratio varies, $HB \times HX$ typically has both sexes present at most sites where they are found (Unmack et al. unpublished data).

Studied Material

We examined 33 individuals from 10 localities across eastern Australia (supplementary table S1, Supplementary Material online). We analyzed representatives of four sexual species: HA, HB, HX, and H. klunzingeri (HK) which was included as an outgroup as it is the sister species to the sexual/unisexual complex (Unmack et al. 2019). Two types of putative hybridogenetic hybrids were examined: HA×HB and HB×HX. A subset of 17 individuals was used for karyotype analyses and identification of somatic versus gonadal differences. The gonadal structure of 16 individuals was examined through confocal microscopy. All wild samples were obtained under state fisheries permits, and research was conducted with approval from the University of Canberra Ethics Committee (CEAE.15-05). Each individual was anesthetized with an overdose of clove oil. Complete information about the number of individuals, sex, localities, and methods employed is provided in supplementary table S1, Supplementary Material online.

Genotype Identification—DArT Sequencing

Genotypes of most individuals were confirmed via SNPs generated using DArTseq (DArT Pty Ltd.), a variation of the double-digest RAD technique as described by Kilian et al. (2012). The R-package dartR 1.8.3 (Gruber et al. 2018) was used for filtering the data, generating PCA plots and for exporting data for phylogenetic analysis to enable species identification. More information is provided in Supplementary Methods.

DNA Flow Cytometry

The genome size of cell populations from the testes was estimated by measurement of the cell nuclei using a BD FACSAria II flow cytometer on a subset of 17 individuals (same individuals as used for karyotype analyses; table 1 and supplementary table S1, Supplementary Material online). Testes and muscle tissues were fixed in 70% ethanol prior to measurements. Testes and muscle tissues were minced in 0.1% Triton X100, 10 µ/ml DAPI and 15 mM MgCl₂ to release nuclei from cells. These nuclei suspensions were incubated at +4°C overnight. After incubation of nuclei suspensions for 4–6 h (at +4°C), they were analyzed by BD FACSAria flow cytometer. At least 10,000 events were measured. BD FACSDiva software (6.1.3) was used to process the obtained data. Suspension of nuclei released from muscle cells was used as an internal control.

#	₽	Genotype	Locality	2n i	n Somatic	Cells	2n	in Germ C	ells	Genom	ne Sets Distinguished Based o	on GISH
				Zn	m/sm	st/a	2n	m/sm	st/a	Somatic	Cells	Germ Cells
. .	HA×HB_1	$HA \times HB$	Angas	46	2	44	4	4	4	B genome = 2 m/sm + 20st/a	A genome = 24 st/a	BB genome = 4 m/sm + 40st/a
5	$HA \times HB_2$	$HA \times HB$	Angas	46	2	44	4	4	40	B genome = 2 m/sm + 20st/a	A genome = 24 st/a	BB genome = 4 m/sm + 40st/a
m.	$HA \times HB_{-}3$	$HA \times HB$	Angas	46	2	44	4	4	40	B genome = 2 m/sm + 20st/a	A genome = 24 st/a	BB genome = 4 m/sm + 40st/a
4	$HA \times HB_{-}4$	$HA \times HB$	Angas	46	2	44	4	4	40	B genome = 2 m/sm + 20st/a	A genome = 24 st/a	BB genome = 4 m/sm + 40st/a
ы.	HA×HB_5	$HA \times HB$	Mudgeeraba	45	m	42	42	9	36	B genome = 3 m/sm + 18st/a	A genome = 24 st/a	BB genome = 6 m/sm + 36st/a
<u>.</u>	HA×HB_6	$HA \times HB$	Byfield	47	-	46	46	2	4	B genome = 1 m/sm + 22st/a	A genome = 24 a	BB genome = 2 m/sm + 44st/a
7.	$HB \times HX_1$	HB×HX	Gwydir	47	-	46	46	2	4	B genome = 24st/a	X genome = 1 m/sm + 22a	XX genome = 2 m/sm + 44st/a
œ.	HB×HX_2	HB×HX	Faithful	48	0	48	48	0	48	B genome = 24st/a	X genome = 24 st/a	XX genome = 48st/a
6.	HB×HX_3	HB×HX	Faithful	48	0	48	48	0	48	B genome = 24st/a	X genome = 24 st/a	XX genome = 48st/a
10.	HB×HX_4	HB×HX	Gwydir	46	2	44	46	2	4	B genome = 1 m/sm + 22st/a	X = 1 m/sm + 22st/a	XX genome = 2 m/sm + 44st/a
11.	HB_1	Ħ	Gwydir	47	-	46	47	-	46	No specif. signal	No specif. signal	No specif. signal
12.	HB_2	뛰	Gwydir	48	0	48	48	0	48	I		I
13.	HA_1	ЧA	Byfield	48	0	48	48	0	48	I	I	I
14.	HK_1	¥	Yabba	48	0	48	48	0	48	I		I
15.	HK_2	¥	Yabba	48	0	48	48	0	48	I	I	I
16.	HK_3	¥	Yabba	48	0	48	48	0	48			I
17.	HX_1	¥	Urumwalla	48	0	48	48	0	48	Ι	I	I

genome, haploid genome of HX; X genome, haploid genome of HX; m/sm, meta-submetacentric; st/a, subtelocentric-acrocentric chromosomes; transmitted genomes are highlighted with bold font.

GBE

Table 1

Chromosome Preparation

Metaphase chromosomes were prepared according to Bertollo et al. (2015) with slight modifications. Briefly, fish were injected with 0.1% colchicine solution (1 ml/100 g of body weight) 45 min before being sacrificed using an overdose of anesthetic. The kidneys, gills, part of spleen, and guts were dissected in 0.075 M KCl at room temperature. The cell suspension free of tissue fragments was hypotonized for 30 min in 0.075 M KCl, fixed in freshly prepared fixative (methanol:acetic acid 3:1, v/v), washed twice in fixative and spread onto microscope slides. For inspection of chromosomal composition and structure in germ cells metaphases and germ cells meiotic metaphases I, suspensions from testes were prepared, using the same protocol, with hypotonization prolonged to 45 min. The same protocol cannot be used for females as they have a low number of dividing cells as well as large yolky oocytes, preventing examination of meiosis using classical cytogenetic methods.

Cytogenetic Analyses

Mitotic metaphase chromosomal preparations from all individuals were stained with 5% Giemsa solution for 10 min to confirm ploidy and morphology of chromosomes. To confirm the genome composition in hybrid individuals and to detect possible genome elimination, GISH was performed on chromosomes obtained from both somatic and gonadal tissue of 12 putative hybridogenetic hybrids. Probes used in GISH experiments were prepared from whole genomic DNA (gDNA) of the three parental sexual HA, HB, and HX. gDNA was extracted from muscles using the DNeasy Blood and Tissue Kit (Qiagen, Hilden, Germany) according to the manufacturer's instructions. gDNA samples were labeled via nick translation using Fluorescein Nick Translation Labeling Kit (Jena Bioscience, Jena, Germany) and Cy3 Nick Translation Labeling Kit (Jena Bioscience) following the protocol supplied by the manufacturer. The best results were obtained after 45 min of nick translation until labeled DNA fragments were 200-500 bp long. Species-specific hybridization probes combined gDNA of sexual species (HB with either HA or HX) to perform GISH experiments on chromosomes of hybrid individuals (table 1). Salmon sperm was used as a blocking reagent for repetitive DNA. The hybridization and detection procedure were carried out under conditions described by Majtánová et al. (2016). The chromosomes were counterstained with Vectashield/DAPI (1.5 mg/ml) (Vector, Burlingame, CA).

Microscopy and Image Analyses

Chromosomal preparations were examined by a Zeiss Axioplan epifluorescence microscope equipped with a CCD camera and a ZEISS Axio Imager.Z2 epifluorescence microscope (Zeiss, Oberkochen, Germany). Images of metaphase chromosomes were recorded with a CoolCube 1 camera (MetaSystems, Altlussheim, Germany). The IKAROS and ISIS imaging programs (Metasystems) were used to analyses grayscale images. The captured digital images from GISH experiments were pseudocolored (red for Anti-Digoxigenin-Rhodamin, green for Invitrogen FITC-Streptavidin) and superimposed using Adobe Photoshop software (CS5).

3D Immunofluorescence Staining

To compare gonadal morphology between sexual and hybrid individuals as well as to identify the ontogenetic stage when genome elimination occurs, we examined gonads of 16 individuals (supplementary table S1, Supplementary Material online) under a laser scanning confocal microscope. Juveniles (1–2 months, 10 mm long) and adult fish were sacrificed using an anesthetic overdose. The caudal muscle tissue of each individual was collected in 80% ethanol for genotyping using DArT sequencing. The main body (for juveniles) or part of gonads (in case of adults) was fixed in 2% paraformaldehyde for 10 h and then transferred to $1 \times$ PBS with 0.02% NaN₃ for long term storage. Prior to immunofluorescence staining, gonads were placed in 1% solution of Triton X100 in $1 \times$ PBS and incubated for 4–5 h at room temperature. Afterward, tissues were washed in 1× PBS at room temperature and incubated for 1-2 h in a 1% blocking solution (Roche) prepared with $1 \times PBS$. Germ cells were visualized by rabbit polyclonal antibodies against Vasa protein (DDX4 antibody [C1C3], GeneTex, Irvine, CA). Incubation with primary antibodies was carried out at room temperature overnight, followed by washing in $1 \times PBS$ with 0.01% Tween (ICN Biomedical Inc.). Secondary antibodies conjugated with Alexa-594-conjugated goat anti-rabbit IgG (H + L) (Thermo Fisher Scientific, Waltham, MA) were added according to the manufacturer's instructions and incubated for 12 h at room temperature. Tissues were then washed in $1 \times PBS$ with 0.01% Tween (ICN Biomedical Inc.) and counterstained with DAPI (1 μ g/ μ l) (Sigma) in 1 × PBS at room temperature for overnight.

Confocal Laser Scanning Microscopy

Tissues were placed in a drop of DAPI with Vectashield (1.5 mg/ml) (Vector, Burlingame, CA), solution and mounted with cover slides and examined under Leica TCS SP5 confocal microscope based on the inverted microscope Leica DMI 6000 CS (Leica Microsystems, Germany). Specimens were analyzed using HC PL APO $20\times$, $40\times$, and $63\times$ objective. Diode and helium-neon lasers were used to excite the fluorescent dyes DAPI and fluorochrome Alexa-594, respectively. The images were captured using LAS AF and processed in LAS AF Lite software (Leica Microsystems, Germany).



Fig. 1.—Karyotype differences between somatic cells and germ cells in hybrid individuals HA×HB (*Hypseleotris galii* × *H*. sp. Midgley's). Giemsa-stained karyotypes obtained from somatic cells (first column), germ cells (second and third column). m/sm, meta-submetacentric; st/a, subtelocentric–acrocentric chromosomes. In meiotic metaphase I, we observed bivalents forming circles (arrows). The numbers of such bivalents correspond to pairs of metacentric chromosomes observed in mitosis of germ cells after the expected elimination of one genome and duplication of the second one (i.e., hybridogenetic reproduction). Bars equal 10 μ m.

Results

Genotype Identification—DArT Sequencing

All results obtained from DArT sequencing are provided in the Supplementary Materials. Genotyping confirmed the identifications of all individuals examined in this study as provided in table 1, supplementary table S1 and supplementary figures S2–S5, Supplementary Material online. We were unable to sequence juvenile individuals from Albury and Wagga Wagga, but these populations only consisted of HB and HB×HX adults at the time when the juveniles were collected.

Karyotype Variability among Analyzed Individuals

We observed variability in numbers and morphology of chromosomes obtained from somatic tissue among the studied individuals. Whereas some individuals displayed a karyotype 2n = 48, composed of solely acrocentric chromosomes, others exhibited chromosomal counts ranging from 45 to 47 with one to three metacentric chromosomes. All individuals, regardless of the chromosomal number, displayed constant chromosome arm numbers, NF = 48. Detailed information about karyotype composition for each individual is provided in table 1.

Karyotype Differences of Mitotic Spreads between Somatic and Gonadal Tissues in Hybrids

In hybrid individuals, we observed within-individual differences in chromosome classification and numbers. These characters can be used to compare metaphases obtained from somatic tissues (mixed kidneys, gills, and livers) and gonadal tissue (testes) (fig. 1), because the absence of chromosomes characterizing a specific chromosomal set in one tissue may indicate their programmed loss. Differences in the chromosome numbers suggest the possible premeiotic elimination of one genome and subsequent duplication of the other in these individuals (fig. 1). Detailed information about differences of chromosome numbers between somatic and germ cells for each individual are provided in table 1. In addition to the mitotic metaphase spreads obtained from gonads, we observed also other stages of meiosis, that is, metaphase I. These spreads consist of bivalents. The numbers of such bivalents correspond to pairs of chromosomes observed in mitosis of germ cells after the expected elimination of one genome



Fig. 2.—Genomic in situ hybridization (GISH) in somatic and germ cells of hybrid individuals. Both haploid parental chromosomal sets were clearly distinguishable in metaphases obtained from somatic tissue after GISH. In metaphases obtained from gonadal tissue, only one parental genome was detected. Red dye represents *H*. sp. Midgley's (HB) gDNA; green dye represents *Hypseleotris galii* (HA) gDNA or *H*. sp. Bald (HX) gDNA. To visualize the proper morphology of chromosomes, Giemsa stained metaphase spreads are presented. Chromosomes are arranged in a decreasing size order, metacentric/ submetacentric chromosomes are marked with asterisks. Bars equal 10 µm.

and duplication of the second one (i.e., hybridogenetic reproduction). The pairs of metacentric chromosomes (if presented), formed circle-shaped bivalent (fig. 1).

Chromosomal Evidence of Hybridogenesis via Genome Elimination

GISH was performed on 12 hybrid and one sexual individual used as a control (table 1 and supplementary table S1, Supplementary Material online) to identify parental chromosomal sets. Both haploid parental chromosomal sets were clearly distinguishable in metaphases obtained from somatic tissue in all hybrids (fig. 2, left panel). The metaphase chromosomes showed no visible intergenomic exchanges between the parental sets, supporting a scenario of the genomic integrity and F_1 hybrid state on a whole-chromosomal level. In metaphases obtained from gonadal

tissue (germ cells), only one parental genome was detected after GISH staining (fig. 2, right panel). Based on the identification of haploid parental chromosomal sets in somatic cells, we were able to distinguish which genome is presented in gonads. In all HA×HB hybrids, we only detected the parental HB genome (corresponding to parental species H. sp. Midgley's) in gonadal metaphases (fig. 2, table 1). That means that parental genome HA (corresponding to H. galii) was eliminated, followed by duplication of HB genome (table 1). Based on these observations we conclude that such individuals uniparentally transmit only the HB genome into their gametes. In HB×HX hybrids, we observed that parental HB genome was eliminated. Nevertheless, in two cases (i.e., IDs: HB×HX_3 and HB×HX_4; fig. 2), when the parental haploid sets consist of the morphologically same karyotypes we cannot clearly conclude which genome is propagated without using more specific cytogenetic markers.



Fig. 3.—Comparison of gonadal microanatomy in sexual and hybrid individuals. Whole-mount immunofluorescent staining with antibodies against Vasa protein (red) identifying germ cells (G). DAPI is visualizing chromatin (cyan). (A) Sexual male *Hypseleotris klunzingeri* (ID: HK_3); (B) hybrid male HB×HX (*H.* sp. Midgley's × *H.* sp; ID: HB×HX_7); (C) sexual female HB (*H.* sp. Midgley's; ID: HB_3); (D) hybrid female *H.* sp. Midgley's × *H.* sp. Bald (ID: HB×HX_5). According to the morphology of gonads, several cell types can be determined: S, spermatids; P, cells in the pachytene stage of meiotic division; G, germ cells; D, diplotene cells of meiotic division. Bars equal 50 µm.

Gonadal Structure of Hybrid and Sexual Individuals

As hybrid males are absent or rare in most sexual/unisexual complexes, we analyzed their ability to produce sperm via DNA flow cytometry. We analyzed seven sexual and 10 hybrid adult males (supplementary table S1, Supplementary Material online). Both hybrid and sexual individuals possessed haploid (1C, corresponding to spermatids and spermatozoa), diploid (2C, corresponding to spermatogonia and somatic cells) and cells after DNA synthesis cell populations (4C, corresponding to primary spermatocytes) (supplementary fig. S1, Supplementary Material online). We also did not observe the accumulation of aneuploid cells (supplementary fig. S1, Supplementary Material online). Our results suggest that meiosis in hybrids likely does not affect regular spermatogenesis and that hybrid males are able to produce haploid sperm.

Additionally, to investigate whether gametogenesis operates normally in hybrid males and females, we performed the analysis of gonadal microanatomy using confocal scanning microscopy in seven adult individuals. We checked one hybrid male and two hybrid females as well as three sexual males and one sexual females. The gonadal morphology of adult hybrids of both sexes is similar to those of sexual individuals (fig. 3). In hybrid males, we detected large clusters of spermatids. Smaller clusters were represented by cells during the pachytene stage of meiosis. Individual gonocytes, as well as primordial germ cells (PGC), were identified by immunostaining of Vasa protein. Gonads of adult females clearly showed diplotene oocytes as well as individual gonocytes located on the periphery of the gonad (fig. 3). In combination with results from DNA flow cytometry, we found no obvious differences between sexual and hybrid individuals, suggesting that the fertility of hybrid males and females is not reduced when compared with their sexual relatives.

Genome Elimination Occurs in Juvenile Individuals

According to the analysis of chromosomal spreads from gonads of adult hybrid individuals and the absence of one parental set during meiosis, we looked for evidence whether genome elimination of one set followed by genome duplication of another set takes place prior to meiosis. In order to detect the process of genome elimination, we analyzed juvenile fish (before fully developed gonads, i.e., 1–2 months old).



Fig. 4.—Comparison of gonadal microanatomy in sexual and hybrid juvenile individuals. Whole-mount immunofluorescent staining with antibodies against Vasa protein (red) identifying germ cells. DAPI is visualizing chromatin (cyan). (A) Sexual individual H. sp. Midgley's (HB_6); (B) and (C) hybrid individuals H. sp. Midgley's \times H. sp. Bald (HB \times HX_8 and HB \times HX_9); arrows indicate micronuclei in the cytoplasm of germ cells. Bars equal 50 μ m.

In gonads of two sexual and four juvenile hybrids, gonial cells were identified with antibodies against the Vasa protein as large cells with multiple nucleoli and less intensive chromatin staining compared with somatic cells (fig. 4). At this developmental stage, meiotic cells were isolated or absent, and most cells were gonial and actively dividing as we observed multiple mitotic divisions. In all observed hybrid individuals, we detected micronuclei in the cytoplasm of germ cells (fig. 4B). Micronuclei were presented as a round chromatin positive body, usually with the more intense chromatin staining, suggesting possible heterochromatinization (fig. 4B and C). The number of micronuclei varied from one to seven per individual germ cell with an average of four micronuclei per cell. In sexual species, we have not detected any micronuclei in fish of the same age. Our results suggest that genome elimination has already occurred via micronuclei formation before meiosis commences in juvenile carp gudgeons.

Discussion

The First Hybridogenetic Animal from the Southern Hemisphere

Both sexual reproduction and uniparental genome elimination require fertilization, meiosis and formation of haploid gametes. However, only the uniparental genome elimination leads to segregation of the genomes nonrandomly, creating asymmetric genetic systems with uneven sex ratio as an evolutionary playground for peculiar phenotypes (Normark 2001; Austin et al. 2009; Ross et al. 2011). Our study has confirmed and delivered direct evidence for classic hybridogenesis as a reproductive mode for the unisexual Australian carp gudgeon hybrids (fig. 5), correctly predicted by Bertozzi et al. (2000) and later Schmidt et al. (2011). Apart from the well-known cases of unisexual reproduction, including hybridogenesis in the Northern Hemisphere, all the obligate unisexual animals from South America, Australia, and New Zealand appear to reproduce through parthenogenesis (i.e., "virgin birth"; reproduction without mating; Schön et al. 2009). Australian carp gudgeons add to the knowledge of the formation and global distribution of unisexual reproduction as the firstknown animals using hybridogenesis in the Southern Hemisphere.

Uniformity in Fertility and Ploidy Level but High Variability in Karyotypes

Most unisexual animal taxa reproduce as all-female populations, with males being typically absent or sterile. Undeveloped gonads and inability to produce sperm was frequently found among hybrids from various genera, for example, Cobitis loaches (Juchno et al. 2017; Juchno and Boroń 2018; Jablonska et al. 2020), Misgurnus loaches (Itono et al. 2006), oribatid mites (Heethoff et al. 2009), Bacillus stick insects (Mantovani and Scali 1992; Mantovani et al. 1999), or Diadromus wasps (El Agoze et al. 1994). In all these examples, hybrid females were able to reproduce normally and did not exhibit any gonadal aberrations. Previous studies have shown that the hybrid male's sterility is caused by the inability to modify their gametogenesis in order to achieve clonality as females do. In this respect, males have problems with orthologous pairing in meiosis (Kuroda et al. 2019; Dedukh, Majtánová, et al. 2020; Spangenberg et al. 2017). Our analysis of gonadal microanatomy and flow cytometry did not indicate any aberration in male and female fertility, as both of them exhibited normal gonads with cells on various gametogenic stages.

In carp gudgeons, all observed hybrid males demonstrated the usual pairing of chromosomes during meiotic division (fig. 1). Thus, hybridogenetic reproduction does not restrain any gametogenic stages in males, and those can produce visually functional gametes (fig. 3). One comparative diploid system exists in Central Europe in water frogs, where the sexual species lives with the all-male hybrid lineages (Doležálková et al. 2016; Doležálková-Kaštánková et al. 2018). In these populations, male gametes are also produced hybridogenetically. Less surprisingly, ovarian microanatomy of carp gudgeons confirmed the functional gametogenesis for female hybrids, which produces oocytes (fig. 3). However, carp gudgeons represent an enigmatic model group, since



Fig. 5.—Schematic diagram of genome elimination in carp gudgeons. Diagram represents one of the case examples of this study.

the all-diploid ploidy level is linked with co-occurring hybrids of both sexes. Our flow cytometric and karyotype analyses showed no evidence of triploid individuals, which is in agreement with previous carp gudgeon studies (Bertozzi et al. 2000; Schmidt et al. 2011; Unmack et al. 2019). Indeed, sexual/unisexual complexes comprising strictly diploid hybrids are rare and were recently described in *Hexagrammos* fish populations (Suzuki et al. 2017). All other animal systems, in which both males and females can reproduce through mechanisms alternative to sexual reproduction, need the presence of polyploid individuals to be functional and stable (Alves et al. 2001; Scali et al. 2003; Stöck et al. 2012; Collares-Pereira et al. 2013; Dedukh et al. 2015; Zhang et al. 2015).

This study represents the first report of karyotype composition in sexual species and hybrid individuals in the fish genus *Hypseleotris*. We described intraspecific karyotype variability in sexual species and hybrids bearing their genomes. Surprisingly, our comparative analysis revealed a high level of karyotype variability within and between sexual species as well as hybrids (table 1). Karyotypes of these fishes included mostly a number of acrocentric chromosomes accompanied by metacentric chromosomes varying from zero to three across individuals. Variation in chromosome numbers has been frequently observed in hybrid fish taxa. However, the pattern was caused by leakage of individual chromosomes from sperm during gynogenesis, or due to aberrant cell divisions (Sola et al. 1992; Fontana et al. 2007; Zhang et al. 2015; Suzuki et al. 2017). Hybridization events may thus be one of the driving forces causing karyotype reorganization.

Despite the detected karyotype variability, the number of chromosome arms remains constant for all observed individuals, NF = 48. That suggests that presented metacentric chromosomes could arise via Robertsonian rearrangements involving the centric fusion of acrocentric chromosomes. Robertsonian translocations represent a relatively frequent phenomenon causing variability of chromosomal numbers within individuals of the same species. They were described in various fish species (Galetti et al. 2000; Morescalchi et al. 2011; Guyomard et al. 2012), and are considered to have generally little impact on meiotic chromosome pairing (Lanzone et al. 2007). Therefore, they might not represent a barrier to hybridization among individuals with different karyotypes (Lajus 2007).

Evidence for Intact Parental Chromosomal Sets and Uniparental Inheritance in Hybrids

Using GISH, we identified two clear groups of chromosomal constituents in hybrid soma. The origin of haploid sets to the parental species from which they derive was difficult to trace back only at some individuals due to a limitation of GISH markers and intraspecific chromosomal variation within sexuals (fig. 2, table 1). Nevertheless, chromosomes were divergent enough in DNA sequence variation to bind labeled DNA species-specifically and split them into haploid sets, and the method is still powerful enough to detect possible intergenomic exchanges between chromosomes as found in unisexual salamanders (Bi et al. 2007). Studied carp gudgeon hybrids had a rather integral structure of parental sets, typical for wellmaintained F₁ hybrid constitutions like other gynogenetic fishes or hybridogenetic water frogs (Zaleśna et al. 2011; Majtánová et al. 2016). Similarly, the integral character of a single parental (species-specific) set in germ cells of several hybrid individuals (fig. 2) suggests that parental sets might have been formed clonally and their reproduction was hemiclonal.

Carp gudgeons are a remarkable group for its nonrandom pattern in uniparental chromosomal inheritance. All of those hybrids having one to three metacentrics in the soma also had the metacentric chromosomes in gonadal tissue, and in twice the number. We provide evidence that genome duplication occurs premeiotically and results in the uniparental hybridogenetic reproduction as previously suggested based on allozyme and microsatellites data (Bertozzi et al. 2000; Schmidt et al. 2011). Moreover, based on our results, it seems there is a correlation between the presence of metacentric chromosome(s) and the genome being transmitted (fig. 5). Uzzell et al. (1980) proposed a hypothesis for water frogs that one parental genome may contain factors responsible for the induction of hybridogenetic gametogenesis and its preferential transmission. Despite the fact the exact mechanism of genome elimination in carp gudgeons remains unknown, and such information can only be obtained from future long-term breeding experiments, the observed preferential propagation of a genome bearing the metacentric chromosomes is documented for the first time and requires further study to shed light on the mechanisms of selective genome elimination. As such, the evolution of metacentric chromosomes from the ancestral acrocentric carp gudgeons karyotype (2n = 48) and the actual mosaic variation and distribution pattern in the genus remain enigmatic and deserves more detailed study.

Mechanism of Uniparental Genome Elimination in Carp Gudgeons

Our gonocyte analysis in adults did not detect any traces of DNA degradation or chromosomal lagging that usually accompany uniparental genome elimination in various organisms (Ishii et al. 2016; Chmielewska et al. 2018). It allowed us to infer that genome elimination does not occur in adult individuals. However, we detected micronuclei in the cytoplasm of germ cells in juvenile hybrids during stages of differentiation in which fish gonads contain somatic and germ cells. Germ cells that arise from PGC of an embryo actively proliferate, giving rise to primary oogonia and prespermatogonia (Van Winkoop et al. 1994). These ontogenetic stages usually correspond with the sexual differentiation of fish gonads. Micronuclei were shown to be connected with genome elimination in Pelophylax water frogs (Ogielska 1994; Chmielewska et al. 2018; Dedukh, Riumin, et al. 2020). These tetrapods gradually eliminate chromosomes through the accumulation of heterochromatin markers and degradation inside autophagosomes (Chmielewska et al. 2018; Dedukh, Riumin, et al. 2020). Such observations contrast with hybridogenetic Poeciliopsis females, which have variation in uniparental elimination modes. In this fish the whole paternal genome is eliminated during single oogonial division when attached to the unipolar spindle (Cimino 1972).

In carp gudgeons, we observed from one to seven micronuclei per cell, which is mechanistically closer to the gradual process of elimination rather than elimination all at once. The gradual genome elimination is a widespread pathway in eukaryotes, operating in plant hybrids, during programmed genome rearrangements in sea lampreys, or elimination of B chromosomes in insects and birds (Subrahmanyam and Kasha 1973; Gernand et al. 2006; Staiber 2006; Timoshevskiy et al. 2016; Torgasheva et al. 2019). The micronuclei are wellknown structures that appear as a result of chromosome missegregation and reflect chromosome instability in many kinds of cancer cells (He et al. 2019), or a variety of cells exposed to genotoxic agents (Sánchez et al. 2000). Thus, the presence of micronuclei observed in hybrid individuals from carp gudgeons may indicate the gradual elimination of one of the parental genomes during their hybridogenetic reproduction.

In this article, we have presented cytological mechanisms underlying uniparental genome elimination and hybridogenesis in the Australian carp gudgeons. We anticipate that carp gudgeons will provide a good model system to help unveil some fundamental biological phenomena. A comparison of karvotypes provides the first view of the preferential transmission of the genome bearing the metacentric chromosomes. Chromosomal remodeling resulting in diverse karyotype variation seems to be linked with local hybridization and asexual reproduction when compared with a conservative karyotype of pure sexual populations free of hybrids. Second, there are not many groups of animal hybrids in which both sexes have functional gonads maintaining their reproductive potential. Carp gudgeons may, therefore, provide insights into conditions both for hybrid fertility and sterility. Finally, the occurrence of hybridogenesis in the Southern Hemisphere suggests that not only the geographic parthenogenesis (sensu Kearney 2005) is distributed worldwide, and may tell us more about the geography and demography in the unisexual origins.

Supplementary Material

Supplementary data are available at *Genome Biology and Evolution* online.

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Author's Contribution

ZM, TE and PJU designed the study and codrafted MS together with LC, DD and PR. MA and PJU collected material. ZM and DD performed cytogenetic analyses. DD performed immunofluorescent staining and confocal microscopy. ZM and DD performed flow cytometry. PJU performed genotype analyses with DArT sequencing. All co-authors contributed to the final text version.

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