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Independent Origin of XY and ZW Sex Determination Mechanisms in Mosquitofish Sister Species

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ABSTRACT Fish are known for the outstanding variety of their sex determination mechanisms and sex chromosome systems. The western (*Gambusia affinis*) and eastern mosquitofish (*G. holbrooki*) are sister species for which different sex determination mechanisms have been described: ZZ/ZW for *G. affinis* and XX/XY for *G. holbrooki*. Here, we carried out restriction-site associated DNA (RAD-) and pool sequencing (Pool-seq) to characterize the sex chromosomes of both species. We found that the ZW chromosomes of *G. affinis* females and the XY chromosomes of *G. holbrooki* males correspond to different linkage groups, and thus evolved independently from separate autosomes. In interspecific hybrids, the Y chromosome is dominant over the W chromosome, and X is dominant over Z. In *G. holbrooki*, we identified a candidate region for the Y-linked melanic pigmentation locus, a rare male phenotype that constitutes a potentially sexually antagonistic trait and is associated with other such characteristics, e.g., large body size and aggressive behavior. We developed a SNP-based marker in the Y-linked allele of *GIPC PDZ domain containing family member 1* (*gipc1*), which was linked to melanism in all tested *G. holbrooki* populations. This locus represents an example for a color locus that is located in close proximity to a putative sex determiner, and most likely substantially contributed to the evolution of the Y.

KEYWORDS Mosquitofish; sex determination; pool sequencing; *gipc1*; *sox9b*

TELEOST sex determination is perplexingly variable, ranging from genetic XX/XY, ZZ/ZW, and polygenic to environmentally regulated systems and hermaphroditism (Devlin and Nagahama 2002; Mank *et al.* 2006; Kikuchi and Hamaguchi 2013). The high diversity of teleosts, with over 26,000 species (Nelson *et al.* 2016) by far the largest clade of vertebrates, has been extensively studied at many levels of their biological organization, but surprisingly little is

known about the genes on the sex chromosomes that determine sex or fulfill sex-linked functions. In only a few teleosts, extensive gene expression analyses, positional cloning, and functional analyses led to the identification of the sex chromosome and the sex-determination locus. Master sex-determining genes of fish were found to have evolved either by allelic diversification on proto-sex chromosomes or by neofunctionalization of autosomal gene duplicates that moved to the prospective sex chromosomal regions (Herpin and Schartl 2015). So far, most identified fish sex determination genes fall into only two groups, which are implicated in the conserved genetic network regulating gonadal development. These genes are either members of the transforming growth factor- β signaling pathway, for instance, *anti-Müllerian hormone*, its *type II receptor*, and *gonadal soma derived factor*, or transcription factors, namely *dmrt1* and *sox3* (Matsuda *et al.* 2002; Nanda

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et al. 2002; Smith *et al.* 2009; Sutton *et al.* 2011; Kamiya *et al.* 2012; Myosho *et al.* 2012; Eshel *et al.* 2014; Takehana *et al.* 2014; Li *et al.* 2015). As an exception, the sex determination gene of salmonids is a duplicate of an immune regulatory gene that has no known function in sexual development (Yano *et al.* 2012, 2013). The region linked to sex determination can vary greatly in length and differentiation, ranging from no differentiation at all in fugu (*Takifugu rubripes*), where a single SNP in *amhr2* determines sex, to a 258-kb region in medaka (*Oryzias latipes*), and a region estimated to several megabases in the guppy *Poecilia wingei* and sticklebacks (*Gasterosteus aculeatus* and *Pungitius pungitius*) (Peichel *et al.* 2004; Kondo *et al.* 2006; Shikano *et al.* 2011; Kamiya *et al.* 2012; Nanda *et al.* 2014; White *et al.* 2015; Dixon *et al.* 2018; Natri *et al.* 2019). In most cases, information on the molecular differentiation of the sex chromosomes, the structure and function of sex-linked genes, and sex chromosome evolution is scarce. Only a few studies have compared the sex chromosomes of closely related teleosts, *e.g.*, of different species within the genus *Oryzias* and *Gasterosteus*, which highlight that, even within the same genus, different master sex-determining genes or sex-linked chromosomes were frequently recruited (Matsuda *et al.* 2002; Nanda *et al.* 2002; Kitano *et al.* 2009; Ross *et al.* 2009; Shikano *et al.* 2011; Myosho *et al.* 2012; Takehana *et al.* 2014).

Livebearing teleosts of the family Poeciliidae provide some well-studied, classical examples of genetic sex determination, which were already investigated in the 1920s (Schmidt 1920; Winge 1922, 1927). In this group, sex determination is often tightly linked to sex-specific coloration, as many color loci have been mapped in close proximity to the putative sex-determining loci (reviewed in Kottler and Schartl 2018). Many of these pigmentation loci are thought to be sexually antagonistic, or at least to be advantageous for only one of the sexes (reviewed in Kottler and Schartl 2018). Despite many efforts, neither a poeciliid master sex determiner nor a sexually antagonistic pigmentation gene has yet been identified on the molecular level.

The eastern (*Gambusia holbrooki*) and western (*Gambusia affinis*) mosquitofish are poeciliids native to the southeastern part of the USA and the northeastern part of Mexico. They were introduced for mosquito control in many freshwater habitats worldwide, where they threaten the native fauna (Hamer *et al.* 2002; Pyke 2008). Phylogenetic analyses demonstrated that *G. holbrooki* and *G. affinis* are sister species, with an estimated divergence time of ~2–7 MY (Lydeard *et al.* 1995; Lamatsch *et al.* 2015; Helmstetter *et al.* 2016). The natural distribution of the two species overlaps in an area extending from the Southeast of Mississippi (Mobile Bay) to the Northeast of Georgia, but hybridization between the two species is rare, suggesting the occurrence of reproductive barriers (Hubbs 1955; Black and Howell 1979; Wooten *et al.* 1988; Wooten and Lydeard 1990; Scribner and Avise 1993; Wilk and Horth 2016).

Within *G. holbrooki* populations, on average 1% of males present a spectacular melanic coloration phenotype that

consists of a conspicuous jet-black spotting pattern covering the entire body and fins (Figure 1A) (Horth and Travis 2002; Horth 2003, 2004). This pattern is generated by large, irregularly distributed macromelanocytes (Figure 1B) (nomenclature according to Schartl *et al.* 2016a; Regan 1961; Angus 1989; Horth 2006). Studies suggest that the melanic color pattern provides an advantage to the males carrying it, as they grow faster, have a larger gonopodium, and show more aggressive behavior toward females and competitors (Martin 1977; Horth 2003; Horth *et al.* 2010). Although melanic males are very conspicuous, they seem to be less prone to predation than the wildtype, silvery males (Figure 1C) (Horth 2004). Female preference is polymorphic, as female *G. holbrooki* that are derived from populations with melanic males prefer such males and vice versa (Bisazza and Pilastro 2000). The low numbers of melanic males seem to be stabilized by negative frequency-dependent selection (Horth and Travis 2002; Horth 2003). Melanism in *G. holbrooki* is a candidate for a sexually antagonistic phenotype (Martin 1977; Horth 2003; Horth *et al.* 2010), but further studies are required to determine whether the melanic coloration is neutral or would indeed be disadvantageous for females in the sense of a truly sexually antagonistic gene. As a potentially sexually antagonistic color locus, the melanic locus might have contributed to the evolution of the *G. holbrooki* Y chromosome and its identification is therefore of prime interest (Kottler and Schartl 2018).

Cytogenetic studies revealed that *G. holbrooki* lacks heteromorphic sex chromosomes (Black and Howell 1979). However, as the melanic macromelanocyte pattern is transmitted from fathers to sons, it has been generalized that all *G. holbrooki* most likely have an XX/XY sex determination system, in which the male-advantageous melanic locus became linked to the sex-determining locus on the Y chromosome (Angus 1989; Horth 2006). In stark contrast, *G. affinis* has conspicuous, morphologically differentiated sex chromosomes (Black and Howell 1979). Female *G. affinis* have a very large W chromosome, suggesting that the sister species of *G. holbrooki* evolved a ZZ/ZW sex determination mechanism (Black and Howell 1979). *G. holbrooki* is so far the only mosquitofish in which an XX/XY sex determination system has been postulated, while a distinct W chromosome was found not only in *G. affinis*, but also in *G. nobilis*, *G. gaigei*, and *G. hurtadoi* females (Campos and Hubbs 1971; Black and Howell 1979; Lydeard *et al.* 1995). The presence of different sex chromosomal systems makes *G. holbrooki* and *G. affinis* excellent model organisms for studying the evolution of sex chromosomes, Y-linked pigmentation, and other sex-linked traits.

We performed restriction-site associated DNA (RAD) and pool sequencing approaches to identify the sex linkage groups of *G. holbrooki* and *G. affinis*, revealing independent origins of the XY and ZW systems. We identified a unique allele of *GIPC PDZ domain containing family member 1* (*gipc1*) on the *G. holbrooki* Y chromosome (*gipc1Y*), which is linked to the potentially sexually antagonistic melanic phenotype. We also

present a detailed description of the melanic phenotype, which is of utmost importance in order to identify reasonable candidate genes that might underlie this phenotype in a gene set. With our newly developed SNP-based marker for the Y chromosome, we demonstrate that the Y chromosome of *G. holbrooki* is dominant over the W chromosome of *G. affinis* and the X is dominant over the Z. Our data give insights into the genetics and genomics of the complex evolution of different sex chromosome systems in closely related species.

Materials and Methods

Experimental animals/fish strains

Fish of the genus *Gambusia* were maintained under standard conditions as described for the related genus *Xiphophorus* by (Kallman (1975)). Nine *G. holbrooki* strains (six of which produced melanic individuals) and one *G. affinis* strain were included in our analyses: *G. affinis* (WLC1313), geographic origin: Pena blanca Lake, Arizona 31°24'01"N 111°05'20"W. *G. holbrooki* (WLC1475), geographic origin: Tamiami trail, Everglades, Florida. *G. holbrooki* (WLC1253), geographic origin: Fort Lauderdale, Florida. *G. holbrooki* (WLC2105), geographic origin: Everglades City, Florida. *G. holbrooki* (WLC2514), geographic origin: Florida City, Florida 25°23'54.2"N 80°34'20.2"W. *G. holbrooki* (WLC6439), geographic origin: Astouin, Camargue, France. *G. holbrooki* (WLC5589), substrain established from WLC2514. *G. holbrooki* (WLC6475), substrain established from WLC2105. *G. holbrooki* (WLC4598), substrain established from WLC1475. *G. holbrooki* (WLC6636), established from WLC4598. The observed sex ratio in our strains was 1:1.

Ethanol fixed individuals of wild caught *G. holbrooki* from Florida were kindly supplied by Ingo Schlupp, University of Oklahoma (Norman, OK). from his collection: ISO8042 (geographic origin: Kissimmee), ISO8043 (geographic origin: Satellite Beach), ISO8046 (geographic origin: Lake Eustis), and ISO8048 (geographic origin: Bulow Creek).

Males of *G. holbrooki* strains WLC1475, 1253, 2105, and 2514 are melanic, but occasionally nonmelanic males occur. In WLC1475, only a single nonmelanic male besides melanic males and nonmelanic females was recorded since its establishment in 1992. In the other strains, a higher frequency was noted. Nonmelanic males were removed promptly to keep the melanic strains monomorphic. Males of WLC6439 are nonmelanic (wildtype-pigmented). In WLC4598 and WLC6636, melanic pigmentation and sex determination are uncoupled, as both males and females are melanic, although nonmelanic fish occur sometimes. WLC5589 and WLC6475 are F₁ fish derived from a cross between a wildtype, nonmelanic female and a nonmelanic male of strain WLC2514 and WLC2105, respectively, resulting in exclusively nonmelanic offspring. The melanic pigmentation of *G. holbrooki* has been reported to be either sensitive or insensitive to temperature (Angus 1989; Horth 2006). We transferred one WLC2514 and

three WLC2105 nonmelanic males to a tank with a water temperature of 21°, a temperature sufficiently low to trigger melanic pigment development in temperature-sensitive strains (Angus 1989). None of them developed any melanic pigmentation after 3 months (data not shown), indicating that our melanic strains are not sensitive to temperature.

All animals were kept and sampled in accordance with the applicable EU and national German legislation governing animal experimentation. In particular, all experimental protocols were approved through an authorization (568/300-1870/13) of the Veterinary Office of the District Government of Lower Franconia, Germany, in accordance with the German Animal Protection Law (TierSchG).

Crosses

All crosses were performed with virgin females by unassisted matings. Pregnant females were kept isolated in 1 liter aquaria with high density plastic wool and controlled daily after the onset of illumination to avoid cannibalism. Offspring were removed and raised separately. Phenotypic sex was determined after puberty. No increased mortality compared to the parental strains was observed.

Imaging of fish and pigment cells

Pictures of whole fish were taken with a Nikon S5 Pro camera with an AF-S DX NIKKOR 18-105 mm 1:3.5-5.6G ED VR lens or with a VH-Z20W zoom lens connected to a VHX-S90BE and VHX-2000D with the OP-87429 polarization illumination attachment. Details of pigment cells were imaged with the VHX-2000D. To document the pigment cell development of fish after birth, the fish were immobilized and photographed with a Leica DFC450 C camera connected to a Nikon SMZ1000 dissecting microscope. The contrast, brightness, and background of some images were adjusted with GIMP 2.10.6.

Transmission electron microscopy

For transmission electron microscopy imaging, small pieces of the caudal peduncles of one melanic and one wildtype WLC2514 male were excised and fixed in a buffered solution containing 2.5% glutaraldehyde (w/v), 0.05 M KCl, and 2.5 mM MgCl₂ in 0.05 M cacodylate buffer (pH 7.2) at 4° overnight. The pieces were washed five times with 50 mM cacodylate buffer pH 7.2 for 10 min at 4° and then fixed with 2% (w/v) osmium tetroxide for 3 hr on ice. Afterward, the samples were washed five times with water at room temperature and stained in 0.5% (w/v) aqueous uranyl acetate solution at 4° overnight. The samples were then washed five times with water at room temperature and dehydrated (50%, 70%, 90% ethanol for 30 min at 4°, then three times 100% ethanol for 20 min at room temperature). They were washed two times with propylene oxide for 30 min and embedded in Epon/propylene oxide (1:1) overnight at room temperature. The final embedding was carried out with Epon and polymerized at 60° for at least 48 hr. Ultrathin sections (60–100 nm) along the longitudinal axis were made with a Leica EM UC7

microtome. The samples were inspected with a JEOL JEM-2100 transmission electron microscope at 200 kV; images were taken with TVIPS TemCAM F416 4k × 4k. The skin layers were named according to Kottler *et al.* (2014).

Adrenaline treatment

To investigate whether the melanocytes of melanic fish react to adrenaline by aggregating their melanosomes, fish were anesthetized and killed by decapitation. Subsequently, the trunk was kept in tank water containing 2.4 mg/ml epinephrine and moved gently. After 10 min, the pigment cells were photographed with the VHX-2000D. WLC2514 and WLC6439 fish were used.

Cytogenetics

Mitotic chromosome preparations of WLC2514 and WLC1313 fish were made directly from pooled spleen, gills, and cephalic kidney cells after exposing the fish for several hours to a 0.02% (w/v) solution of colchicine followed by hypotonic treatment [0.46% (w/v) KCl] and fixation with a mixture of methanol and acetic acid (3:1). Slides were stained with 5% Giemsa solution. C-banding using the standard BSG (barium hydroxide/saline/Giemsa) technique (Sumner 1972) was used to visualize the constitutive heterochromatin. DAPI and mithramycin staining of metaphase chromosomes were performed according to Schmid *et al.* (2010). To analyze meiotic chromosomes of hybrids, seminiferous tubules were exposed to hypotonic treatment followed by fixation in methanol-acetic acid (3:1) mixture. The meiotic cells from the fixed pieces of seminiferous tubules were mechanically released and allowed to spread on a hot plate in 50% acetic acid. After evaporation of acetic acid, slides were Giemsa stained and subjected to microscopic analysis. WLC2514, WLC1313, and WLC1475 fish were used.

DNA extraction

For RAD and pool sequencing, DNA from fish organs was extracted with phenol/chloroform. Tissue was lysed in 1 ml extraction buffer (0.1 M EDTA pH 8, 0.2% SDS, 0.2 M NaCl) containing 200 µg/ml proteinase K for 3 hr at 80°. Then, 500 µl phenol was added to each sample. After mixing and incubation at room temperature for 10 min, 500 µl chloroform:isoamyl alcohol (24:1) was added. The samples were incubated with occasional inversion at room temperature for 10 min and then centrifuged for 10 min at 5° (5000 × g). The upper layer was transferred to a new tube and one volume chloroform:isoamyl alcohol (24:1) was added. With occasional inverting, the samples were kept at room temperature for 10 min, and subsequently centrifuged for 10 min at 5° (5000 × g). The supernatant was transferred to a glass vial on ice and 2.5 times the sample volume of cold 100% ethanol was carefully added. The DNA was spooled with a glass rod and dissolved in TE buffer, pH 8. The DNA concentration was measured with a Qubit fluorometer (dsDNA BR Assay Kit, Invitrogen, Carlsbad, CA).

For PCR, DNA was extracted from trunk tissue of ethanol-fixed fish according to Reid *et al.* (2008). Lysis was carried out at 56° overnight.

RAD sequencing

For *G. affinis* RAD sequencing, DNA from 31 females and 30 males (WLC1313) was used. From *G. holbrooki*, 26 females and 30 melanic males of WLC2514 were included.

RAD-tag libraries were built from extracted DNA using Sbf1 as a single restriction enzyme, following the standard protocol described in Amores *et al.* (2011). All resulting libraries were sequenced on one lane of Illumina HiSeq 2500 in single end 150 bp mode. Reads were demultiplexed and quality-controlled using the `process_radtags.pl` script from the Stacks software version 1.44 (Catchen *et al.* 2011), with the following parameters: discard reads with low quality scores (-q), remove any reads with an uncalled base (-c), and rescue barcodes and RAD-tags (-r).

Demultiplexed reads were analyzed with the RADSex pipeline (version 0.2.0, Zenodo, doi: 10.5281/zenodo.2616396). The underlying principle of RADSex is to split the demultiplexed reads into groups based on exact sequence match. Therefore, markers created with RADSex are nonpolymorphic, and reads that would be grouped into the same polymorphic marker using standard analyses software are split into multiple markers with RADSex. As a consequence, RADSex markers are only influenced by a single parameter (*i.e.*, minimum depth to consider a marker present in an individual), and markers can be compared between individuals using a straightforward presence/absence analysis. These properties enable easier cross-species comparison than with heavily parameter-driven software like Stacks. To analyze the RAD-Seq reads with RADSex, a table containing the depth of each marker in each individual from the dataset was first generated using the `process` command with default parameters. Then, the distribution of markers between males and females was computed using the `distrib` command with the minimum depth to consider a marker set to 10 (-min-cov 10), and the resulting distribution was plotted with the R package RADSex-vis. Sequences of markers significantly associated with sex were obtained with the `distrib` command also using a minimum depth of 10 (-min-cov 10). Markers were aligned to the poeciliid reference genome of the platyfish (*Xiphophorus maculatus*) [NCBI accession number: GCA_002775205.2; improved version of the genome described in Schartl *et al.* (2013)] with the `map` command using a minimum depth of 10 (-min-cov 10) and manhattan plots were generated with RADSex-vis. The manhattan plots showed $-\log_{10}(P_{\text{association with sex}})$ from a Chi-squared test on the number of males and number of females in which a marker is present for *G. affinis*/*G. holbrooki* markers aligned to the genome of *X. maculatus*. The *Xiphophorus maculatus* genome was used, as *Xiphophorus* is the closest relative of *Gambusia*, for which chromosome information is available. *Xiphophorus* and *Gambusia* have the same number of chromosomes.

Pool sequencing

For Pool-seq library preparation, individual gDNAs were pooled in equimolar ratios by sex (Supplemental Material, Table S1). Pool-seq libraries were prepared following Illumina's protocols using the Illumina TruSeq Nano DNA HT Library Prep Kit (Illumina, San Diego). Briefly, 200 ng of each gDNA pool (pools of males and females) was fragmented to 550 bp by sonication with a M220 Focused-ultrasonicator (Covaris, Woburn, MA). Size selection was performed using SPB beads and the 3' ends of blunt fragments were adenylated. Then, adaptors and indexes were ligated and the construction was amplified with Illumina-specific primers. Library quality was assessed using a Fragment Analyzer (Advanced Analytical Technologies, Ames) and libraries were quantified by qPCR using the Kapa Library Quantification Kit (Roche, Basel, Switzerland). Sequencing was performed on a NovaSeq S4 Lane (Illumina, San Diego, CA) using a paired-end read length of 2×150 nt with Illumina NovaSeq Reagent Kits. Sequencing produced 121 and 136 million paired reads for the male and female pooled libraries in *G. holbrooki*, respectively, and 94.7 and 94.2 million paired reads for the male and female pooled libraries in *G. affinis*, respectively.

For each species, reads from the male and female pools were aligned to the genomes of *Xiphophorus maculatus* [NCBI accession number: GCA_002775205.2; improved version of the genome described in Scharl *et al.* (2013)] and *G. affinis* (NCBI accession number: NHOQ01000000; Hoffberg *et al.* 2018) using BWA mem with default parameters (version 0.7.17; Li 2013). The resulting BAM files were sorted with Samtools *sort* (version 1.8; Li *et al.* 2009) with default parameters and PCR duplicates were removed with Picard tools MarkDuplicates (version 2.18.2) with the parameter REMOVE_DUPLICATES = true. A pileup file combining male and female pools was generated from the resulting BAM files using Samtools *mpileup* with minimum base quality set to 0 (-Q 0), and then converted to a sync file using popoolation2 (version 1201; Kofler *et al.* 2011) with minimum mapping quality set to 20 (-min-qual 20). This sync file was analyzed with PSASS (version 2.0.0, Zenodo, doi: 10.5281/zenodo.2615936, available at <https://github.com/RomainFeron/PSASS>), with parameters -range-het 0.1, -range-hom 0.01, -window-size 100000, and -output-resolution 100,000, to compute three metrics in 100-kb nonoverlapping windows over the genome: (1) F_{ST} between males and females over the window, (2) number of sex-specific SNPs, defined as SNPs with a frequency between 0.4 and 0.6 in one sex and a frequency higher than 0.99 in the other sex, and (3) reads depth for each sex. We filtered for a minimum depth of 10. We also carried out the same analysis with a 10 kb window size. Results were plotted with the R package PSASS-vis (version 2.0.0, available at <https://github.com/RomainFeron/PSASS-vis>).

Genes within the regions of interest were determined using the platyfish and *G. affinis* reference genomes. In platyfish, the genes in the regions of interest were determined with the NCBI Genome Data Viewer (<https://www.ncbi.nlm.nih.gov/>

[genome/gdv/](https://www.ncbi.nlm.nih.gov/)). In *G. affinis*, the regions of interest were extracted from the reference genome and aligned against the NCBI nr/nt databank using the discontinuous megablast algorithm (Altschul *et al.* 1990) to identify potential candidate genes.

Genome assembly of a melanic *G. holbrooki* male

A melanic WLC2514 *G. holbrooki* male genome was assembled from 250 bp long Illumina paired-end reads using DiscoverDeNovo (reference <https://software.broadinstitute.org/software/discover/blog/>) with standard parameters.

Sequence alignments

DNA and protein alignments of candidate genes were carried out with Clustal Omega (1.2.4) (Sievers *et al.* 2011). GenBank accession numbers: XP_007572354.1 (Amazon molly (*Poecilia formosa*) *Gipc1*); XP_007560319.1 (Amazon molly *Sox9a*); XP_007556425.2 (Amazon molly *Sox9b*); XP_025001585.1 (chicken (*Gallus gallus*) *Gipc1*); BAA25296.1 (chicken *Sox9*); XP_008436174.1 (guppy (*Poecilia reticulata*) *Sox9a*); NP_001284373.1 (guppy *Sox9b*); NP_005707.1 (human (*Homo sapiens*) *Gipc1*); CAA86598.1 (human *Sox9*); XP_023813268.1 (medaka (*Oryzias latipes*) *Gipc1*); AAX62152.1 (medaka *Sox9a*); AAX62151.1 (medaka *Sox9b*); XP_014866717.1 (Atlantic molly (*Poecilia mexicana*) *Gipc1*); NP_061241.1 (mouse (*Mus musculus*) *Gipc1*); NP_035578.3 (mouse *Sox9*); XP_003442153.1 (Nile tilapia (*Oreochromis niloticus*) *Gipc1*); XP_005448042.1 (Nile tilapia *Sox9a*); XP_003450167.1 (Nile tilapia *Sox9b*); XP_005801543.1 (platyfish (*Xiphophorus maculatus*) *Gipc1*); XP_005807407.1 (platyfish *Sox9a*); XP_005801494.1 (platyfish *Sox9b*). Accession numbers of *G. affinis* sequences: *sox9b* ORF, NHOQ01002125.1:61568-61995,62336-62583,62831-63594; *sox9b* protein sequence, PWA19450.1; *gipc1* ORF, NHOQ01002357.1:1316372-1316665,1316740-1316925,1317062-1317242,1318142-1318254,1318339-1318420,1318497-1318640,1318740-1318819,1319648-1319671 [shortened because probably not correctly assembled (does not end with stop codon)]; *gipc1* protein sequence, PWA17858.1 (shortened in accordance with ORF sequence).

Phylogenetic analysis

The phylogenetic tree was built with MEGA7.0.26 (Kumar *et al.* 2018) according to Hall (2013). The *sox9a* and *sox9b* ORF sequences were aligned with MUSCLE using the default settings. The Maximum Likelihood tree was calculated using the Tamura-Nei model with the partial deletion option (Tamura and Nei 1993). Initial trees for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood approach, and then selecting the topology with superior log likelihood value. A discrete Gamma distribution was used to model evolutionary rate differences among sites [five categories (+G, parameter = 0.3546)]. Fewer than 5% alignment gaps, missing data, and ambiguous bases were allowed at any position; 1000 replicates of bootstrapping were carried out.

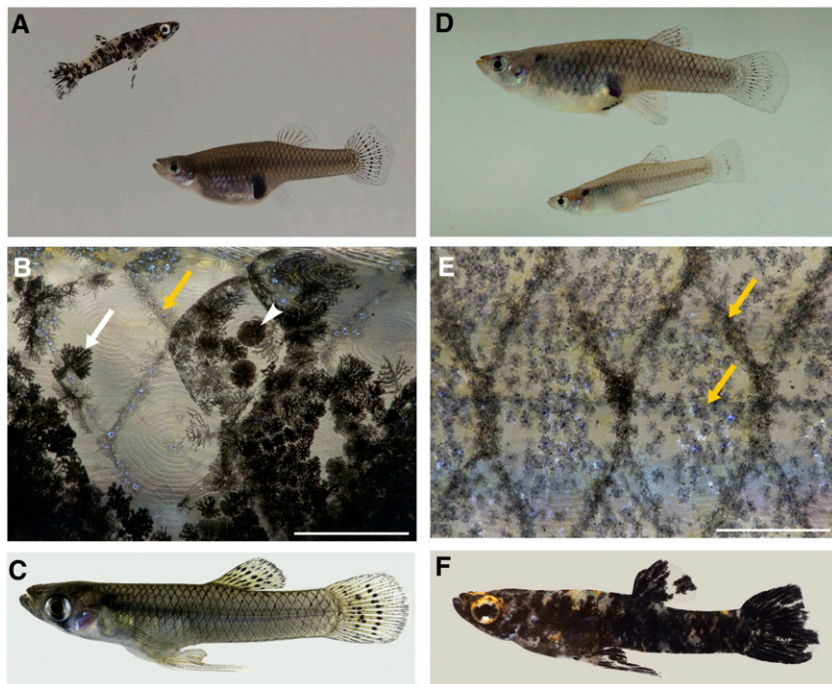


Figure 1 Phenotype of *G. holbrooki* and *G. affinis*. (A) Melanic *G. holbrooki* male (top) and wildtype female (bottom) (WLC2514). (B) Detail of the caudal peduncle of a melanic *G. holbrooki* male (WLC2514). Large dendritic- (white arrow) and corolla- (white arrowhead) shaped macromelanocytes and smaller melanocytes of the background pattern (yellow arrow) are visible. (C) Wildtype-pigmented *G. holbrooki* WLC6439 male. (D) *G. affinis* female (top) and male (bottom) (WLC1313). (E) Detail of the caudal peduncle of a wildtype-pigmented *G. holbrooki* male (WLC6439); the yellow arrows point to the melanocytes of the background pattern. (F) *G. holbrooki* male (WLC1475) with pronounced xanthophore-derived pigmentation on the head. All pictures taken under incident light conditions. Bar, (B), (E) 500 μ m. Length of males of both species is \sim 2–3 and of females 4–6 cm.

The tree was rooted on mouse *sox9*. GenBank accession numbers: AY870394.1 (medaka *sox9a*), AY870393.1 (medaka *sox9b*), XM_005807350.2 (platyfish *sox9a*), XM_005801437.2 (platyfish *sox9b*), XM_008437952.2 (guppy *sox9a*), NM_001297444.1 (guppy *sox9b*), XM_007560257.2 (Amazon molly *sox9a*), XM_007556363.2 (Amazon molly *sox9b*), XM_005447985.2 (Nile tilapia *sox9a*), XM_003450119.4 (Nile tilapia *sox9b*), NM_011448.4 (mouse *sox9*).

Preparation of cDNA

To prepare cDNA of different *Gambusia* strains, total RNA was extracted from skin tissues using TRIzol (Invitrogen) according to the manufacturer's instructions. After DNase I (Thermo Fisher Scientific, Waltham, MA) treatment, cDNA was prepared with the RevertAid First Strand cDNA Synthesis Kit (Thermo Fisher Scientific) following the manufacturer's instructions. First strand cDNA was directly used for PCR.

Gipc1 and *sox9b* PCRs and sequencing

To amplify *gipc1* and *sox9b* from cDNA of different *Gambusia* strains, 5' and 3' UTR primers were used (*gipc1*: forward 5'-CTTCCTAGAGCGGGTCCAC-3', reverse 5'-AGTGTTAACACCGCACAGG-3'; *sox9b*: forward 5'-TTCAGTGCAGCAGATAGTC-3', reverse 5'-AGTAGTCTGGGTGGATGTGTTG-3'). Q5 High-Fidelity DNA Polymerase (New England BioLabs, Ipswich, MA) was used for amplification, which was carried out according to the manufacturer's instructions with an annealing temperature of 62° and an elongation time of 1 min. The PCR products were visualized by gel electrophoresis, purified with the GenElute PCR Clean-Up Kit (Sigma-Aldrich, St. Louis, MO), and sequenced by a sequencing provider (Eurofins, Ebersberg, Germany). For cloning, a TA-overhang was added to the PCR products using a *Taq*

DNA polymerase. Subsequently, the PCR products were cloned into pGEM-T Easy vector (Promega, Madison, WI) following the manufacturer's instructions. Plasmid DNA was extracted with the GenElute HP Plasmid Miniprep Kit (Sigma-Aldrich) and sequenced by Eurofins. Sequences were inspected with SnapGene Viewer Version 4.2.11.

For genotyping the genomic DNA of 71 fish from different strains (Table S2), SNP Pol (high discrimination of single nucleotides) DNA polymerase (Genaxxon, Ulm, Germany) was used with primers specific for *gipc1Y* (forward 5'-GCTCCAGTTCGGGCTCAGC-3', reverse 5'-AGCTCGTCCACGGTGGCGA-3') and *sox9bY* (forward 5'-ACCACCCTTGGTGGAGCAGC-3', reverse 5'-GAAGAGGGGCTGTAGTGCTGTAA-3'). The PCRs were carried out according to the manufacturer's instructions with an annealing temperature of 62° and an elongation time of 30 s. PCR products were inspected by gel electrophoresis.

SNP Pol DNA polymerase and gel electrophoresis were used as well to distinguish between *G. holbrooki sox9bX* (primers: forward 5'-GCAGCGAAGTCATCTCCAACATC-3', reverse 5'-GCTCTCCTGCTCCACCAAGGT-3'), *sox9bY* (primers: forward 5'-GCTGCGAAGTCATCTCCAACATC-3', reverse 5'-GCTCTGCTGCTCCACCAAGGG-3'), and *G. affinis sox9b* (primers: forward 5'-AGCAGCGAAGTCATCTCCAACATT-3', reverse 5'-GCTCTCCTGCTCCACCAAGGC-3'; based on a resequencing of *sox9b* of *G. affinis* strain WLC1313).

PCR using DNA of F₁ hybrids to determine sex chromosome genotypes was carried out using the *sox9bY* primers [forward 5'-GCTGCGAAGTCATCTCCAACATC-3', reverse 5'-GCTCTGCTGCTCCACCAAGGG-3'; SNP Pol DNA polymerase PCR protocol (Genaxxon, Ulm, Germany)] and the *amtW* primers [Gaf88 (Lamatsch *et al.* 2015); standard *Taq* DNA polymerase PCR protocol].

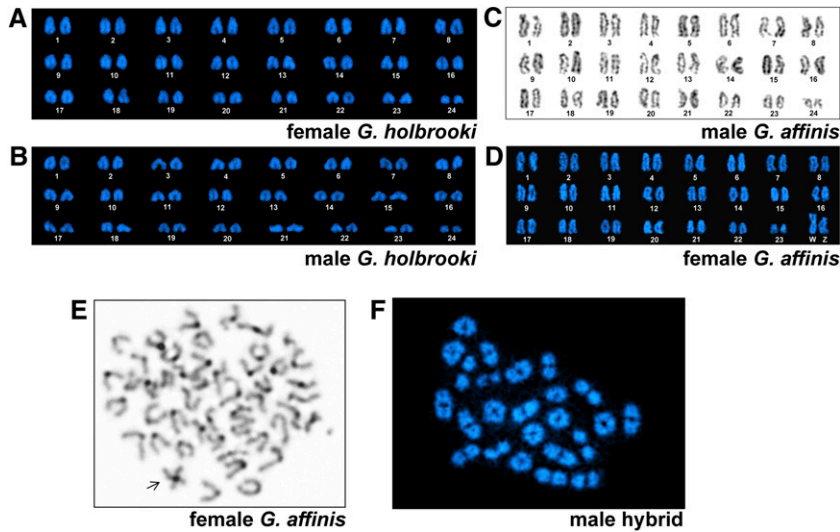


Figure 2 Cytogenetic analyses in the eastern (*Gambusia holbrooki*) and western (*G. affinis*) mosquitofish. (A) DAPI-stained female karyotype of *G. holbrooki* (WLC2514). (B) DAPI-stained male karyotype of *G. holbrooki* (WLC2514). No distinct heteromorphic sex chromosomes can be identified in the karyotype. (C) Giemsa-stained karyotype of male *G. affinis* (WLC1313). (D) DAPI-stained karyotype of female *G. affinis* (WLC1313) displaying distinguishable heteromorphic sex chromosomes (ZW). (E) C-banding of female metaphase of *G. affinis* (WLC1313) showing a large W chromosome (arrow). (F) DAPI stained metaphase I from a male F₁ hybrid of a *G. affinis* female (WLC1313) and a melanistic *G. holbrooki* male (WLC1475) revealing the expected number of paired bivalents. The chromosomes in (A–D) are arranged based on their size.

Data availability

The authors state that all data necessary for confirming the conclusions presented in the article are represented fully within the article. Figure S1 presents the results of the RAD-sequencing analysis. Figure S2 shows the analysis of *G. affinis* and *G. holbrooki* pooled-sequencing reads aligned to the genome of *X. maculatus* using 10 kb windows. Figure S3 represents a molecular phylogeny of fish *sox9*. Figure S4 are alignments of male *G. holbrooki* and *G. affinis* *gipc1* and *sox9b* alleles and protein sequences. Figure S5 are multiple species alignments of *Gipc1* and *Sox9* protein sequences. Figure S6 contains PCR results demonstrating that *sox9bY* was generated by allelic diversification. Figure S7 contains pictures of melanistic and nonmelanistic *G. holbrooki*. Table S1 gives information on the fish used for pool sequencing. Table S2 presents the results of the allele-specific PCRs. Table S3 contains the annotated loci within the 100 kb windows with the highest density of sex-specific SNPs. Table S4 contains the sex chromosome genotypes of F₁ fish (pedigree WLC20074) from a cross between a *G. affinis* female (WLC1313) and a *G. holbrooki* male (WLC2514) determined by PCR. Table S5 gives information on the number of offspring of nonmelanistic females and nonmelanistic males derived from melanistic strains.

RAD-seq and pool sequencing data are available at Sequence Read Archive, grouped under the same Bioproject (accession number: PRJNA563257). The genome assembly of a melanistic *G. holbrooki* male is available at NCBI, accession number VOPI00000000. *Gipc1* and *sox9b* cDNA sequences are available at GenBank (accession numbers: MK721001-MK721004). Supplemental material available at figshare: <https://doi.org/10.25386/genetics.7957376>.

Results

Sex chromosomes are not heterochromatic in *G. affinis*

A previous study described a large W chromosome in the karyotype of female *G. affinis*, but found no heteromorphic

sex chromosomes in *G. holbrooki* (Black and Howell 1979). To confirm this for our study material, we carried out cytogenetic analyses of female and male *G. affinis* and *G. holbrooki*. Both species have $2n = 48$ chromosomes (Figure 2) (Black and Howell 1979). In accordance with this earlier study, no heteromorphic sex chromosomes were discernible in *G. holbrooki* male and female chromosome spreads, while female *G. affinis* had a large W chromosome (Figure 2, A–D). The W chromosome is the largest and only biarmed chromosome of the karyotype (Figure 2D). Despite the considerable morphological differentiation in size, C-banding of female metaphases revealed that the W is not heterochromatic except for its centromeric region (Figure 2E). This indicates that evolution of the W did not develop through canonical heterochromatinization.

The Y and W chromosome correspond to different linkage groups

The chromosomes of F₁ hybrids between *G. affinis* females and *G. holbrooki* males form 24 bivalents during meiosis, which suggests that the genomes of the two species are highly homologous (Figure 2F). To identify the sex linkage groups of *G. holbrooki* and *G. affinis*, we first analyzed the RAD-sequencing data from each species using RADSex.

In *G. affinis*, 433 markers were significantly associated with phenotypic sex ($P < 0.05$, Chi-squared test with Bonferroni correction), 397 of which (92%) were present in all females and absent from all males, confirming that this species has a ZZ/ZW sex determination system (Figure S1A). In *G. holbrooki*, 309 markers were significantly associated with phenotypic sex ($P < 0.05$, Chi-squared test with Bonferroni correction), 195 of which (63%) were present in all males and absent from all females, confirming that this species has an XX/XY sex determination system (Figure S1B). When aligning the significant RADSex markers from *G. affinis* to the genome of the platyfish, which is closely related to *Gambusia* and has the same number of chromosomes (Walter *et al.* 2004; Hrbek *et al.* 2007), 337 markers (78%)

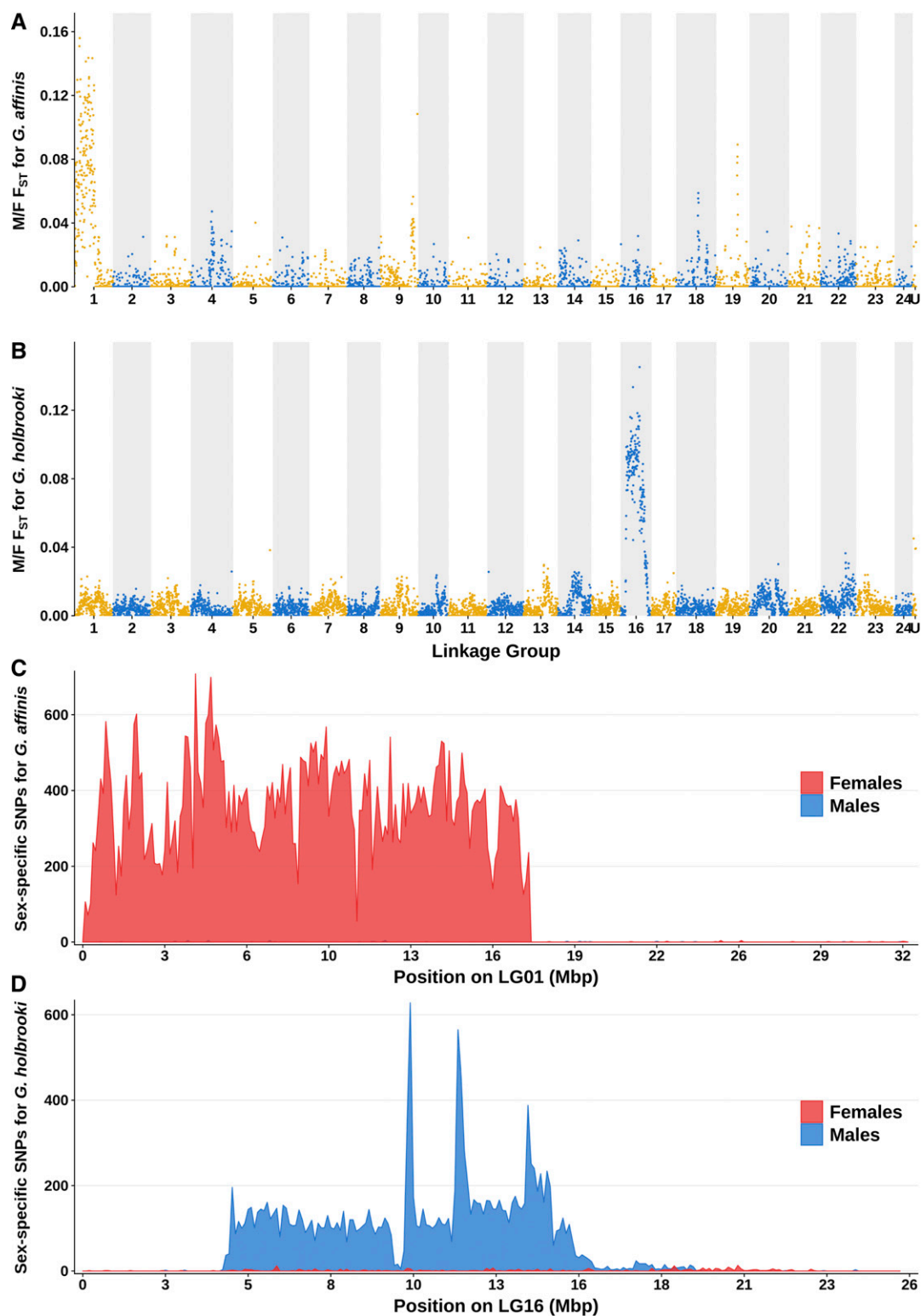


Figure 3 Analysis of *G. affinis* and *G. holbrooki* pooled-sequencing reads aligned to the genome of *X. maculatus*. Male/female F_{ST} was computed in nonoverlapping 100 kb windows from *G. affinis* (A) and *G. holbrooki* (B) pooled-sequencing reads aligned to *X. maculatus* genome. Odd linkage groups (LGs) are colored in yellow with light background, and even LGs are colored blue with dark background. For *G. affinis*, a large region at the first half of LG1 shows a high male/female F_{ST} compared to the rest of the genome, while for *G. holbrooki*, a large region in the center of LG16 has the highest male/female F_{ST} . These results indicate that around half of LG1 of *X. maculatus* is homologous to the sex chromosome of *G. affinis*, while half of LG16 of *X. maculatus* is homologous to the sex chromosome of *G. holbrooki*. The number of female-specific SNPs (FSS, in red) and male-specific SNPs (MSS, in blue) was similarly computed, and results are shown for LG1 for *G. affinis* (C) and LG16 for *G. holbrooki* (D). For *G. affinis*, a region spanning from 0 to 17.7 Mb on LG1 was strongly and homogeneously enriched in FSS compared to the rest of the genome, while for *G. holbrooki*, the region enriched in MSS was located between 4.31 and 16.1 Mb on LG16, and showed three peaks of MSS density ~ 10.2 , 11.9, and 14 Mb.

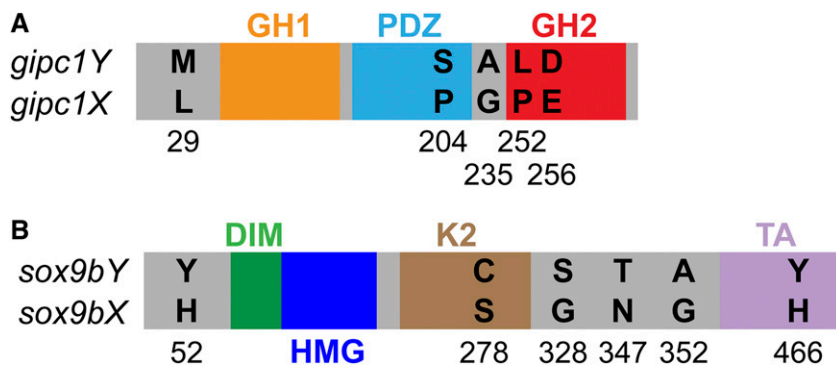


Figure 4 Differences between the X- and Y-encoded Gipc1 and Sox9b proteins found in a melanistic WLC2514 male. (A) Amino acid differences between Gipc1Y and Gipc1X; domains according to (Kato 2013). (B) Amino acid differences between Sox9bY and Sox9bX; domains according to Jo *et al.* (2014). A PQA (proline-, glutamine- and alanine-rich) domain is not present in mosquitofish Sox9b. DIM, dimerization domain; GH1, GIPC homology one domain; GH2, GIPC homology two domain; HMG, high mobility group domain; K2, transactivation domain; PDZ, postsynaptic density-95/Disks large/zona occludens-1 domain; TA, transactivation domain. Protein and domains are not drawn to scale.

were aligned to LG1 (NC_036443.1) (Figure S1C), suggesting that this linkage group is homologous to the sex chromosome in the genome of *G. affinis*. The platyfish genome was used, as the current *G. affinis* genome does not contain chromosome information. The same analysis for *G. holbrooki* revealed 249 markers (81%) significantly associated with sex were aligned to LG16 of *X. maculatus* (NC_036458.1) (Figure S1D), suggesting that LG16 is homologous to the sex chromosome in the genome of *G. holbrooki*. Overall, these results show that, despite being sister species, *G. affinis* and *G. holbrooki* have different sex determining systems and different sex chromosomes.

To confirm the results from the RAD sequencing, and to identify differences between the sex chromosomes at a higher resolution, we generated pool sequencing data for a pool of at least 30 males and a pool of at least 30 females of each species (Table S1) and aligned the reads to the genome of the platyfish and the genome of *G. affinis*. The resulting BAM files were analyzed with PSASS to output male/female F_{ST} , number of sex-specific SNPs, and reads depth for each sex in 100 kb nonoverlapping windows over the genomes. In the following, we focus on the mapping results against the *X. maculatus* genome, as no chromosome information was available for the *G. affinis* genome.

Male/female F_{ST} was highest on platyfish LG1 for *G. affinis* (Figure 3A) and highest on platyfish LG16 for *G. holbrooki* (Figure 3B), confirming the results from the RAD sequencing analysis. The F_{ST} was computed for the respective entire 100 kb window. Using a 10 kb window size did yield similar results (Figure S2). In *G. affinis*, we identified 64,290 female-specific SNPs (FSS) and 827 male-specific SNPs (MSS). Female-specific SNPs are only found in females and male-specific SNPs only in males. Among the FSS, 63,323 (98%) were aligned to LG1 of *X. maculatus* in a region spanning from 0 to 17.7 Mb (Figure 3C), suggesting that this region is homologous to a highly differentiated region in the sex chromosome pair of *G. affinis*. In *G. holbrooki*, we identified 1581 FSS and 17,231 MSS, among which 15,956 (93%) were aligned to LG16 of *X. maculatus* in a region spanning from 4.31 to 16.1 Mb (Figure 3D), suggesting that this region is homologous to a highly differentiated region in the sex chromosome pair of *G. holbrooki*. While the distribution of FSS in

G. affinis was homogeneous in the region on LG1 (Figure 3C), there were three peaks of MSS in *G. holbrooki* located within the region of LG16 ~10.2, 11.9, and 14 Mb, suggesting that the regions homologous to these locations have a higher male/female differentiation on the sex chromosomes of *G. holbrooki* (Figure 3D). The FSS and MSS that did not align to LG1 and LG16, respectively, were not clustered on any other linkage group.

Taken together, these results show a ZZ/ZW sex-determining system with a 17.7 Mb sex-differentiated region homologous to the first half of LG1 of *X. maculatus* in *G. affinis*, and an XX/XY sex-determining system with an 11.8 Mb region homologous to the middle part of LG16 of *X. maculatus* in *G. holbrooki*.

A Y-specific allele of GIPC PDZ domain containing family member 1 (*gipc1*) is linked to *G. holbrooki* melanistic pigmentation

To identify candidate genes for *Gambusia* sex determination and pigmentation based on published literature on gene function, we investigated the annotated loci within the platyfish and *G. affinis* genomic areas with the highest density of female- or male-specific SNPs from our pool sequencing analysis. In *G. affinis*, the 100 kb window with the peak density of female-specific SNPs (708 SNPs when aligned to the platyfish genome and 1058 SNPs when aligned to the *G. affinis* reference genome assembly) contains two annotated miRNAs, a lncRNA, and *piggyBac* transposable element-derived protein 3-like (*pgbd3-like*) in platyfish and a lncRNA and *gata5-like* in *G. affinis* (Table S3). The distance between *pgbd3-* and *gata5-like* is ~87 kb in the platyfish genome. In the *G. affinis* genome, *pgbd3-* and *gata5-like* are not located on the same scaffold. In a previous study, a female-specific *G. affinis* marker in the 3'UTR of *aminomethyl transferase (amt)* was discovered (Lamatsch *et al.* 2015). Platyfish *amt* is located ~4.5 Mb downstream of *pgbd3-like* on chromosome 1. In the *G. affinis* genome, *amt* and *pgbd3-like* are located on separate scaffolds. Based on the available information, it is not possible to further narrow down a region that may harbor a female sex-determining gene.

In *G. holbrooki*, three distinct male-specific SNP peaks are clearly visible on LG16 (Figure 3D). The window containing

Table 1 SNPs at predicted heterozygous sites detected by direct sequencing in *gipc1* *G. holbrooki*

<i>Gipc1</i>	SNPs at predicted heterozygous sites ^a						
	85	610	702	704	755	768	
<i>gipc1X</i> (genome prediction)	T	C	G	G	C	G	
<i>gipc1Y</i> (genome prediction)	A	T	A	C	T	C	
predicted amino acid change X > Y	L > M	P > S	silent	G > A	P > L	E > D	
Strain	Sex	Phenotype					
WLC2514	male	melanic	T/A	C/T	G/A	G/C	G/C
WLC2105	male	melanic	T/A	C/T	G/A	G/C	G/C
WLC1475	male	melanic	T/A	C/T	G/A	G/C	G/C
WLC4598	female	melanic	T/A	C/T	G/A	G/C	G/C
WLC6439	male	wildtype	T	C	G	G	G
WLC2514	female	wildtype	T	C	G	G	G

^a Positions according to *G. holbrooki gipc1* ORF sequences (Figure S4A); SNPs predicted by genome assembly (see Figure S4)

the highest peak (628 SNPs when aligned to the platyfish and 683 SNPs when aligned to the *G. affinis* reference genome) spans 10 loci in the platyfish and 8 in the *G. affinis* genome, among them *gipc1* (also named *synectin*, *neuropilin-1-interacting protein*, or *GAIP interacting protein*, *C terminus*) (Table S3). *Gipc1* interacts with other proteins by its peptide-binding PDZ domain and is involved in vesicular trafficking, thereby affecting, for instance, blood vessel and nervous system formation (De Vries *et al.* 1998; Cai and Reed 1999; Chittenden *et al.* 2006; Naccache *et al.* 2006; Varsano *et al.* 2006; Valdembrri *et al.* 2009; Sanchez *et al.* 2011; Katoh 2013). *Gipc1* might also play a role in the sorting of tyrosine-related protein 1 (Typr1) to the melanosomes, as it binds the cytoplasmic tail of this protein (Liu *et al.* 2001). Typr1 is required for pigment synthesis and melanosome formation in fish (Braasch *et al.* 2009; Krauss *et al.* 2014).

The second highest peak (565 and 546 SNPs, respectively) contains only one locus, the transcription factor *SRY-related HMG box9-like (sox9-like)*, in both platyfish and *G. affinis* (Table S3). Phylogenetic analysis revealed that this gene corresponds to the teleost paralog *sox9b* (Figure S3). *Sox9*, the main target of the mammalian sex-determining gene *SRY*, is crucial for testis formation, making this gene an excellent candidate for *G. holbrooki* sex determination (Wagner *et al.* 1994; Sekido and Lovell-Badge 2008; Capel 2017; Gonen *et al.* 2018). *Sox9* is also expressed in neural crest cells and required for the development of neural crest cell derivatives like bone and cartilage cells (Wagner *et al.* 1994; Bi *et al.* 1999; Yan *et al.* 2005). Moreover, it has been shown that *Sox9* plays a role in melanocyte differentiation and melanoma formation (Passeron *et al.* 2007; Shakhova *et al.* 2015). Within the third peak, three loci were found, none of which, to our knowledge, has been implicated in sex determination or pigmentation so far (Table S3).

As *gipc1* and *sox9b* are potential candidate genes for pigmentation and sex determination, we further investigated both genes, first utilizing our preliminary genome assembly of a melanic *G. holbrooki* WLC2514 male. In the genome assembly, two alleles of both genes were present. The two *gipc1* alleles differ in six positions within the open reading

frame (ORF), translating into five potential amino acid changes, one of which is located in the PDZ and two in the GH2 domain (Figure 4A and Figure S4, A and B). The *sox9b* alleles have 10 variable positions in the ORF, leading to six nonsynonymous amino acid changes (Figure 4B and Figure S4, C and D). The transactivation domains K2 and TA each contain one nonconservative change (Figure 4B and Figure S4D). Multiple species alignments revealed which of the alleles are most likely Y-linked, as the putative Y-linked proteins contain rare or unique amino acid changes (Figure S5). The amino acids at position 204, 235, 252, and 256 are fully conserved in all *Gipc1* protein sequences from fish to human except for *Gipc1Y* (Figure S5A). In *Sox9bY*, the H > Y at position 52 and 466 are unusual, as the histidine at position 52 is conserved in all *Sox9b* sequences and the one at position 466 is present in all of the investigated fish *Sox9a/b* sequences, as well as in human, mouse, and chicken *Sox9* (Figure S5B). Moreover, position 278 is a serine in almost all of the sequences except for Nile tilapia *Sox9b* and *G. holbrooki Sox9bY* (Figure S5B). A glycine is conserved in all fish *Sox9a* and *Sox9b* sequences at position 352, but not in *Sox9bY* (Figure S5B).

To confirm these results, we designed primers in the 5' and 3' untranslated regions of both genes based on our genome assembly. We then amplified *gipc1* and *sox9b* from cDNA prepared from skin tissue of six individuals from different *G. holbrooki* strains and directly sequenced the PCR products. This allowed us to investigate whether the fish were heterozygous at the sites predicted in our genome assembly.

We found that the predicted six heterozygous sites between *gipc1X* and *gipc1Y* were present in all melanic fish, suggesting that *gipc1Y* might be closely linked to the melanic locus (Table 1). They were not linked to being male, as the heterozygous sites could not be found in the wildtype WLC6439 male sequence (Table 1).

In *sox9b*, no heterozygous sites fully associated with sex or melanic pigmentation were found (Table 2). The melanic WLC2514, WLC2105, and WLC1475 males shared the heterozygous sites found in the preliminary genome assembly of the melanic WLC2514 male, except for the silent SNP at

Table 2 SNPs at predicted heterozygous sites detected by direct sequencing in *sox9b* *G. holbrooki* ORFs

<i>Sox9b</i>	SNPs at predicted heterozygous sites ^a											
	154	364	513	675	832	982	1040	1055	1174	1396		
<i>sox9bX</i> (genome prediction)	C	C	C	G	A	G	A	G	C	C		
<i>sox9bY</i> (genome prediction)	T	T	T	T	T	A	C	C	T	T		
predicted amino acid change X > Y	H > Y	silent	silent	silent	S > C	G > S	N > T	G > A	silent	H > Y		
Strain	Sex	Phenotype										
WLC2514	male	melanic	C/T	C/T	C/T	G/T	A/T	G/A	A/C	G/C	C/T	C/T
WLC2105	male	melanic	C/T	C/T	T	G/T	A/T	G/A	A/C	G/C	C/T	C/T
WLC1475	male	melanic	C/T	C/T	T	G/T	A/T	G/A	A/C	G/C	C/T	C/T
WLC4598	female	melanic	C	C	T	G	A	G	A	G	C	C
WLC6439	male	wildtype	C	C	T	G	A	G	A	G	C	C
WLC2514	female	wildtype	C	C	C	G	A	G	A	G	C	C

^a Positions according to *G. holbrooki sox9b* ORF sequences (Figure S4C); SNPs predicted by genome assembly (see Figure S4).

position 513 (Table 2). The melanic WLC4598 female, the wildtype female from the Florida population (WLC2514), and the wildtype-pigmented male from the European population (WLC6439) did not have these heterozygous sites (Table 2). The sequences of the *gipc1* and *sox9b* alleles of the melanic WLC2514 male were confirmed by cloning the PCR products and sequencing (GenBank accession numbers: MK721001-MK721004).

Next, we designed primers specific for *gipc1Y* and *sox9bY* and carried out allele-specific PCRs using DNA from 71 fish from different strains as template (Table S2). *Gipc1Y* was identified in all melanic fish, while *sox9bY* was found in 25 of 28 investigated melanic individuals (Table S2). The three melanic fish lacking *sox9bY* were melanic females, a rare incidence in *G. holbrooki* (Angus 1989). *Gipc1Y* and *sox9bY* were not detected in any of the wildtype-pigmented fish (Table S2).

To find out if *sox9bY* is a Y-specific duplicate of *sox9bX*, we designed primers that distinguish between *G. holbrooki sox9bX*, *sox9bY*, and *G. affinis sox9b*. PCRs using DNA from F₁ individuals of a cross between a *G. affinis* female and a melanic *G. holbrooki* male as template revealed that melanic F₁ males have the *G. affinis sox9b* and *G. holbrooki sox9bY*, but not *sox9bX* (Figure S6). This suggests that *sox9bY* was generated by allelic diversification, not duplication, of *sox9b* (Figure S6). Such an analysis was not possible for *gipc1*, as no suitable primers could be designed.

The Y-linked macromelanocyte pattern of *G. holbrooki* is composed of a divergent pigment cell lineage

G. affinis females and males are grayish silvery, as are most *G. holbrooki* fish (Figure 1, A, C, and D). The uniform pigmentation is caused by melanocytes (Figure 1E), iridophores, and xanthophores, which are distributed over the whole body. In melanic *G. holbrooki*, macromelanocytes are superimposed on this background pattern (Figure 1B). Some macromelanocytes have a compact shape, while others are highly dendritic (Figure 1B). Compared with the background melanocytes, they are considerably darker, and, with an average diameter of 167 μm (up to 300 μm), almost twice as large. Transmission electron microscopy

imaging revealed that the macromelanocytes are located in the dermis and hypodermis, where they are frequently associated with iridophores and xanthophores (Figure S7A). Their melanosomes show the same structure as those of the normal melanocytes (Figure S7, A–D). The first macromelanocytes are usually already discernible 1 or 2 days after birth, and quickly increase in size (Figure S7, E–H). At older age, some melanic males develop pronounced yellow areas, which are usually located on the head and belly (Figure 1F).

To investigate whether macromelanocytes can aggregate their melanosomes like normal melanocytes, we treated melanic and nonmelanic (wildtype) fish with epinephrine (adrenaline). While the background melanocytes of wildtype and melanic males aggregated their melanosomes upon exposure to epinephrine, the macromelanocytes did not change their appearance (Figure 5). As epinephrine is a potent physiological substance to elicit melanosome contraction, we conclude that the macromelanocytes in contrast to the background melanocytes are compromised with respect to this function. This information is crucial in order to identify potential candidate loci underlying this phenotype, e.g., *gipc1*. *Gipc1* affects a protein involved in melanosome development (Liu *et al.* 2001).

G. holbrooki Y is dominant over *G. affinis* W and X is dominant over Z

To determine dominance relationships among sex chromosomes, we set up reciprocal crosses between *G. affinis* and *G. holbrooki*. All crosses resulted in viable and fertile offspring. The brood size of the interspecific crosses and backcross matings was comparable to the brood size of the purebred parental species.

In platyfish, the W chromosome is dominant over the Y chromosome, as WY fish develop as females (Kallman 1973; Volff and Schartl 2002). If, like in platyfish, the *G. affinis* W chromosome would be dominant over the *G. holbrooki* Y chromosome, we would expect 25% WY melanic females, 25% WX nonmelanic (wildtype) females, 25% ZY melanic males, and 25% ZX nonmelanic (wildtype) males in crosses between ZW *G. affinis* females and XY melanic *G. holbrooki*

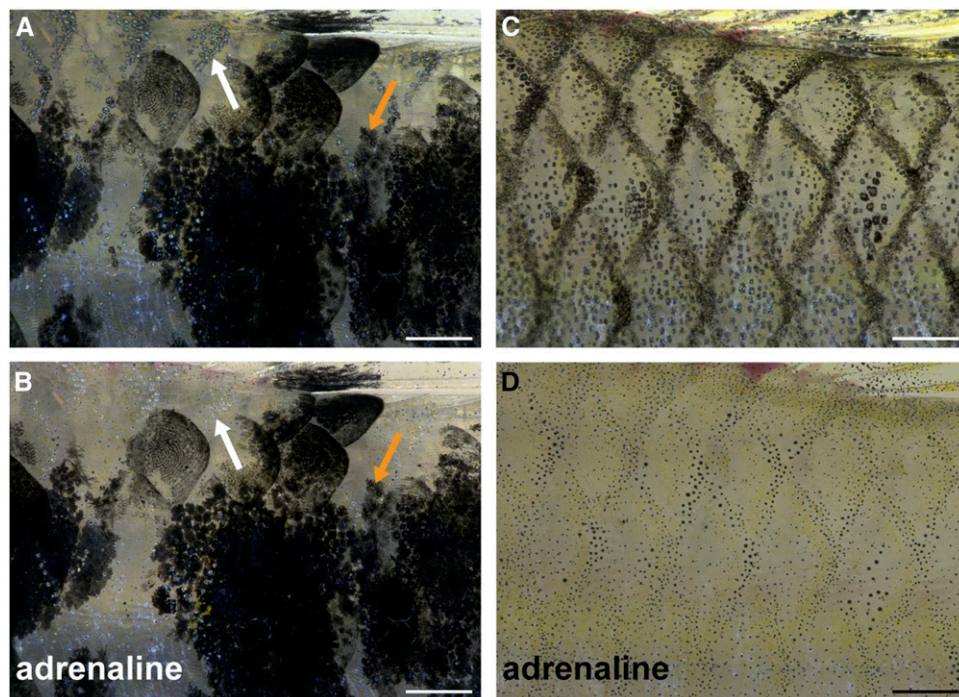


Figure 5 Effect of adrenaline on melanocytes and macromelanocytes of *G. holbrooki*. Skin from a melanic male (WLC2514) before (A) and after (B) treatment with adrenaline. The melanosomes of the background melanocytes (white arrows) aggregate, while those of the macromelanocytes do not (orange arrows). Wildtype, nonmelanic male (WLC6439) skin before (C) and after (D) treatment with adrenaline. The melanocytes aggregate their melanosomes. Pictures were taken under incident light conditions below the dorsal fin. Bar, 500 μ m.

males (note that the W and Y do not correspond to the same linkage group). Deviating from this expectation, only one melanic F₁ female and 280 melanic F₁ males were found in a total of 568 F₁ fish (Table 3). This suggests that almost all WY F₁ fish become males. Support for this comes from the cytogenetic analysis, which revealed that melanic F₁ males had two different karyotypes: some with a W chromosome and others without a morphologically recognizable sex chromosome (data not shown).

The nonmelanic F₁ of our crosses consisted of 236 females and 51 males (Table 3). In some of the crosses, no nonmelanic F₁ males were obtained at all (Table 3). This indicated that, at variable frequencies depending on the cross, many or most ZX fish become females. We further investigated this by crossing XX *G. holbrooki* females with ZZ *G. affinis* males. Their offspring consisted of 83% females despite the uniform ZX genotype and several broods consisted exclusively of females (Table 4).

To confirm that WY fish usually develop as males and ZX fish as females, we genotyped F₁ nonmelanic females and melanic males from one cross of *G. affinis* x *G. holbrooki* for the *G. affinis* W chromosome and the *G. holbrooki* Y chromosome using molecular markers. In this cross, all WY F₁ fish were males and all ZX F₁ fish developed as female (Table S4).

Crosses suggest that female-to-male sex reversal occurs in *G. holbrooki*

All of our melanic *G. holbrooki* strains occasionally produce nonmelanic male offspring. Nonmelanic males derived from melanic fathers can be the result of a recombination or deletion event leading to the loss of the melanic locus from the Y or by a female-to-male sex reversal in XX fish. To test this, we

carried out crosses between such exceptional males and nonmelanic *G. holbrooki* females (Table S5). Variation in these crosses was high, as they resulted in 0–87.7% male offspring (average 29.2%). We genotyped 15 of these males as being negative for *gipc1Y* and *sox9bY* (Table S2). The generally lower percentage of male offspring and the consistency with which such nonmelanic males occur in our melanic strains suggests that at least some of these males are XX.

Discussion

Gambusia affinis and *G. holbrooki* provide the unique opportunity to study different sex chromosome systems in two sister species. The sex determination mechanism of the last common ancestor of *G. holbrooki* and *G. affinis* is not known. Both systems might have evolved in parallel or a transition might have occurred in the common ancestor, separating the lineages from each other. Higher differentiation of the W than the Y chromosome, at least with respect to size, does not indicate that the W is older than the Y. Large W chromosomes are also found in other nonamniotes with young sex chromosomes (Schartl *et al.* 2016b).

The finding from our interspecific crosses that WY hybrids usually become males demonstrates that the sex determination mechanism of *G. holbrooki* is dominant over the one of *G. affinis*. This indicates that the Y harbors a strong, dominant male determiner. Whether sex-determination in *G. affinis* is mediated by a dosage-sensitive locus on the Z similar to the situation in chicken (Smith *et al.* 2009), or a dominant female-sex determiner on the W, still needs to be determined.

Sex chromosome turnover driven by sexually antagonistic selection requires that the new sex determiner is genetically

Table 3 Segregation of offspring pigmentation and sex from crosses between *G. affinis* females and melanic *G. holbrooki* males

Parental ♀	Parental ♂	Pedigree #	F ₁ ♀ melanic	F ₁ ♀ wt	F ₁ ♂ melanic	F ₁ ♂ wt
<i>G. aff.</i> WLC1313	<i>G. hol.</i> WLC2105	2353	1	22	51	21
<i>G. aff.</i> WLC1313	<i>G. hol.</i> WLC1475	5964/l	0	74	118	30
<i>G. aff.</i> WLC1313	<i>G. hol.</i> WLC1475	5964/ll	0	51	44	0
<i>G. aff.</i> WLC1313	<i>G. hol.</i> WLC1475	6244	0	56	43	0
<i>G. aff.</i> WLC1313	<i>G. hol.</i> WLC1475	6245	0	33	24	0
		Σ	1	236	280	51
		Σ		568		

wt, nonmelanic wildtype-pigmented fish.

dominant over the old one, meaning that a new Y chromosome can only invade a ZW sex determination system if the Y is dominant over the W (van Doorn and Kirkpatrick 2007). Thus, we predict that the Y from *G. holbrooki* may invade *G. affinis* in the hybrid zone between the two species and the W will be eventually lost. This is supported by a recent study using microsatellite markers, which provided evidence for nonlinear gene flow within the hybrid zone of the two species (Wilk and Horth 2016). The distortion of the sex ratio caused by interspecific crosses might lead to a decreased fitness of hybrid populations and multiple introgression events could even lead to WW offspring that might not be viable. A previous study using microsatellite markers found that the heterozygosity of hybrid mosquitofish in the Mobile Bay area is low, and that the *G. holbrooki* phenotype is more abundant (Wilk and Horth 2016). This motivates future studies to further sample the region where the two species are in contact, to determine which sex chromosomes are present in the hybrid zone, and to carry out crosses to find out whether WW fish are viable. Hybrid speciation and introgression are prevalent in fish evolution and have been documented in particular in poeciliids (Meyer *et al.* 2006; Roberts *et al.* 2009; Cui *et al.* 2013; Kang *et al.* 2013).

Our study provides an important first step toward identifying the sex determination locus in *G. holbrooki* and *G. affinis* and the Y-linked pigmentation locus in *G. holbrooki* by pinpointing regions on the sex chromosomes that are especially highly differentiated. These regions were identified by pool sequencing, which defines such areas in the genome more precisely than RAD-seq and is therefore a promising method to investigate sex determination systems in nonmodel organisms. *G. affinis* and *G. holbrooki* confirm that highly differentiated sex chromosomes can be either heteromorphic (ZW) or homomorphic (XY). Like in *G. holbrooki*, the sex chromosome pair of medaka appears homomorphic, but, on the molecular

Table 4 Sex ratio of offspring between *G. holbrooki* females and *G. affinis* males

Parental ♀	Parental ♂	Pedigree #	Brood	F ₁ ♀	F ₁ ♂
<i>G. hol.</i> WLC1475	<i>G. aff.</i> WLC1313	5964r/ll	1	22	0
			2	25	0
			3	11	5
			4	28	6
			5	13	6
			6	8	12
			7	34	6
<i>G. hol.</i> WLC1475	<i>G. aff.</i> WLC1313	5964r/l	1	33	0
			Σ	174	35
			Σ	209	

level, the medaka Y is highly differentiated over a region of at least 3 Mb (Takeda 2008; Wilson *et al.* 2014). Even between populations, the accumulation of repetitive elements and pseudogenes on the sex chromosomes can vary greatly (Nanda *et al.* 2014).

As a potentially sexually antagonistic Y-linked locus, the melanic locus might have played a major role in the evolution of the *G. holbrooki* Y chromosome (Kottler and Schartl 2018). Studies suggest that the low percentage of melanic males in natural populations is maintained by negative frequency-dependent selection (Horth and Travis 2002; Horth 2003). Large numbers of melanic males seem to trigger increased male and female mortality rates (Horth and Travis 2002; Horth 2003). One possible explanation for this would be the aggressive behavior of melanic males, leading to, ultimately detrimental, stress (Horth and Travis 2002; Horth 2003). Our genotyping results suggest that *gipc1Y* either contributes to the formation of the melanic color pattern or is closely linked to the melanic locus. In zebrafish, a dominant mutation in *tyrp1a* leads to defects in eumelanin production and melanosome morphology, ultimately resulting in melanocyte death (Krauss *et al.* 2014). A similar phenotype is observed in *tyrp1a* and *tyrp1b* knockdown zebrafish (Braasch *et al.* 2009). *Gipc1* is involved in the intracellular transport of newly synthesized Tyrp1 protein, but Tyrp1 is probably only one of many *Gipc1* interaction partners within the melanocytes and the effect of the detected mutations, or of an overexpression of *Gipc1*, is difficult to predict (Liu *et al.* 2001).

Because of the repressed recombination on the Y, genotyping of more populations with melanic males is required to confirm the linkage between the melanic phenotype and *gipc1Y*. In general, the detection of an array of SNPs differing between the X and Y suggests that a large part of the Y is differentiated from the X, as it is the case in the *G. affinis* W and Z. This implies that the expression and/or coding sequences of many genes on the sex chromosomes are most likely altered. It is therefore possible that the melanic phenotype is caused by several additive Y-linked mutations that

became fixed on the Y due to the repressed recombination. For instance, *cAMP-dependent protein kinase catalytic subunit alpha*, which encodes a subunit of protein kinase A (PKA), is located close to *gipc1*. PKA regulates melanosome movement via cAMP regulation, and mutations at this locus leading to permanently high internal levels of cAMP could explain the insensitivity of the macromelanocytes to adrenaline (Fujii 2000; Rodionov *et al.* 2003). The temperature-sensitivity of the melanic phenotype in some *G. holbrooki* populations indicates that the expression of the causative locus, or loci, is affected and that melanocytes are transformed into macromelanocytes at lower temperatures (Angus 1989; Horth 2006). However, further studies are necessary to confirm that the temperature-sensitive melanic phenotype is caused by the same locus. The melanic locus, or a locus closely linked to it, affects xanthophores as well, as older *G. holbrooki* males frequently develop yellow areas on the head and trunk. In our strains, macromelanocytes were first visible shortly after birth, well before males could be distinguished from females by their secondary sexual characteristics. This demonstrates that, in contrast to, for instance, the famous male pigment patterns of the guppy, the melanic pattern is not under the control of high androgen levels that are only established after reaching sexual maturity (Kottler and Schartl 2018).

Occasionally, our melanic *G. holbrooki* strains produce nonmelanic males. While some of these males might have lost the melanic locus on the Y, the sex bias to female in our crosses suggests that many of these males are XX. As we currently only have a marker for the melanic locus (*gipc1Y*) but not for the Y chromosome in general, we could not investigate yet whether most of these males indeed lack a Y chromosome. Male development in the absence of the Y can be induced by autosomal modifiers that occur with varying frequency and penetrance within a population. This has been reported from other poeciliid fish of the related genus *Xiphophorus* (Kallman 1984; Schartl *et al.* 2011). Recurrence of XX males in crosses from female-to-male sex-reversed XX fishes due to autosomal modifiers was also observed in medaka (Nanda *et al.* 2003). Another possibility would be that the environment affects sex determination in *Gambusia*.

Sox9bY appeared as a reasonable sex determination candidate gene in our study, but was not linked to male sex in all of our fish, as wildtype-pigmented males lacked *sox9bY*. However, we currently do not know whether melanic and wildtype males share the same sex chromosome. If melanic males have a distinct Y chromosome, *sox9bY* would be an excellent candidate for their sex-determining locus.

Taken together, understanding how different sex determination mechanisms in closely related species evolved will shed light on one of the most fundamental and fascinating aspects of vertebrate genome evolution.

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