

RESOURCE ARTICLE

An ancient truncated duplication of the anti-Müllerian hormone receptor type 2 gene is a potential conserved master sex determinant in the Pangasiidae catfish family

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Funding information

National Institutes of Health, Grant/Award Number: R01 OD011116 and R35 GM139635; Agence Nationale de la Recherche, Grant/Award Number: ANR-10-INBS-09, ANR-13-ISV7-0005 and ANR-16-CE12-0035

Handling Editor: Andrew DeWoody

Abstract

The evolution of sex determination (SD) in teleosts is amazingly dynamic, as reflected by the variety of different master sex-determining genes identified. Pangasiids are economically important catfishes in South Asian countries, but little is known about their SD system. Here, we generated novel genomic resources for 12 Pangasiids and characterized their SD system. Based on a *Pangasianodon hypophthalmus* chromosome-scale genome assembly, we identified an anti-Müllerian hormone receptor type II gene (*amhr2*) duplication, which was further characterized as being sex-linked in males

and expressed only in testes. These results point to a Y chromosome male-specific duplication (*amhr2by*) of the autosomal *amhr2a*. Sequence annotation revealed that the *P. hypophthalmus* *Amhr2by* is truncated in its N-terminal domain, lacking the cysteine-rich extracellular part of the receptor that is crucial for ligand binding, suggesting a potential route for its neofunctionalization. Reference-guided assembly of 11 additional Pangasiids, along with sex-linkage studies, revealed that this truncated *amhr2by* duplication is a male-specific conserved gene in Pangasiids. Reconstructions of the *amhr2* phylogeny suggested that *amhr2by* arose from an ancient duplication/insertion event at the root of the Siluroidei radiation that is dated to ~100 million years ago. Together these results bring multiple lines of evidence supporting that *amhr2by* is an ancient and conserved master sex-determining gene in Pangasiids, a finding that highlights the recurrent use of the transforming growth factor β pathway, which is often used for the recruitment of teleost master SD genes, and provides another empirical case towards further understanding of dynamics of SD systems.

KEYWORDS

amhr2, evolution, male genome assembly, pangasiid catfishes, sex determination

1 | INTRODUCTION

Catfishes (order Siluriformes) with ~4,000 species (Sullivan, Lundberg & Hardman, 2006) are economically and ecologically important fish worldwide. Among catfishes, the Pangasiid family (Pangasiidae) is recognized as a monophyletic group including four extant genera: *Helicophagus*, *Pangasianodon*, *Pangasius* and *Pteropangasius* (Pouyaud, Gustiano & Teugels, 2016). These species have a wide range of habitats both in fresh and brackish water across southern Asia, from Pakistan to Borneo (Roberts & Vidthayanon, 1991) and some species are listed in the IUCN red list (IUCN, 2021) as either endangered, such as the striped catfish *Pangasianodon hypophthalmus*, or even critically endangered, such as the Mekong giant catfish, *Pangasianodon gigas*, or the giant pangasius, *Pangasius sanitwongsei*. Many Pangasiids, because of their rapid growth rate, are also important aquaculture species, such as *Pangasius bocourti*, *Pangasius djambal* and *Pangasianodon hypophthalmus* (Lazard, Cacot, Slembrouck & Legendre, 2009). *P. hypophthalmus*, for example, has become a major aquaculture species extensively farmed in many Asian countries (Anka, Faruk, Hasan & Azad, 2014; Na-Nakorn & Moeikum, 2009; Phuong & Oanh, 2010; Singh & Lakra, 2012) and has even been recently introduced to Brazilian finfish aquaculture.

Sex determination (SD) mechanisms have not been investigated in detail in Pangasiid catfishes, but genetic sex-linked markers that could facilitate broodstock management for aquaculture or conservation purposes have been searched without success in both *P. hypophthalmus* and *P. gigas* (Sriphairoj, Na-Nakorn, Brunelli & Thorgaard, 2007). SD in vertebrates can rely on genetic (GSD for genetic SD), environmental (ESD for environmental SD) or a combination of both genetic and environmental factors (such as thermal effects on GSD = GSD+TE) (Baroiller, D'Cotta & Saillant, 2009; Kobayashi, Nagahama & Nakamura, 2013; Ospina-Alvarez

& Piferrer, 2008). In teleost fishes, SD has been found to be extremely variable with species expressing one or another of these three systems, GSD, ESD and GSD+TE, scattered across teleost fish phylogeny. In addition, teleosts exhibit a wide range of GSD systems, with both classical male (XX/XY) and female heterogamety (ZZ/ZW), but also more complex GSD systems relying on polygenic SD with or without multiple sex chromosomes (Devlin & Nagahama, 2002; Mank & Avise, 2009; Moore & Roberts, 2013). These transitions or turnovers of different GSD systems have been found in closely related species belonging to the same genus (Takehana, Hamaguchi & Sakaizumi, 2008) and even across populations of the same species (Kallman, 1973). A similar high turnover has also been found for master sex-determining (MSD) genes at the top of the genetic SD cascade (Matsuda et al., 2002; Myosho et al., 2012; Nanda et al., 2002; Q. Pan et al., 2016; Takehana et al., 2014). Many of these fish MSD genes belong to the "usual suspect" category (Herpin & Schartl, 2015) because they derive from key genes regulating the gonadal sex differentiation network. These "usual suspect" MSD genes currently belong to a few gene families, such as the *Dmrt* (Chen et al., 2014; Matsuda et al., 2002; Nanda et al., 2002), *Sox* (Takehana et al., 2014), steroid-pathway (Koyama et al., 2019; Purcell et al., 2018) and transforming growth factor beta (TGF β) families (Pan et al., 2021), which have been independently and recurrently used to generate new MSD genes. The greatest diversity of MSD genes is found within the TGF β family with the anti-Müllerian hormone, *amh* (Hattori et al., 2012; Li et al., 2015; Pan et al., 2019), the gonadal soma derived factor, *gsdf* (Myosho et al., 2012; Rondeau et al., 2013), or the growth/differentiation factor 6, *gdf6* (Imarazene et al., 2021; Reichwald et al., 2015) genes, but also TGF β type II and type I receptors with the anti-Müllerian hormone receptor type 2, *amhr2* (Feron et al., 2020; Kamiya et al., 2012), and the bone morphogenetic protein

receptor, type IBb, *bmpr1bb* (Rafati et al., 2020) genes. However, a few exceptions to the “usual suspects” rule have also been identified with, for instance, the conserved salmonid MSD *sdY* gene that evolved from an immunity-related gene (Bertho, Herpin, Schartl & Guiguen, 2021; Yano et al., 2012, 2013).

Based on a chromosome-scale high-quality genome assembly, and previously published whole-organ transcriptomic data (Pasquier et al., 2016) of *P. hypophthalmus*, we identified a male-specific duplication of *amhr2* (*amhr2by*) in that species. This potential Y chromosome-specific *amhr2by* encodes an N-terminal truncated protein that lacks the cysteine-rich extracellular part of the receptor, which is key for proper Amh ligand binding. Sex-linkage studies and genome sequencing of 11 additional Pangasiid species show that *amhr2by* is conserved as a male-specific gene in at least four Pangasiid species, stemming from a single ancient duplication/insertion event at the root of the Siluroidei suborder radiation that is dated to ~100 million years ago (Kappas, Vittas, Pantzartzi, Drosopoulou & Scouras, 2016). Together, these results bring multiple lines of evidence supporting the hypothesis that *amhr2by* is potentially an ancient and conserved MSD gene in Pangasiid catfishes and highlight the recurrent use of the TGF β pathway, which is often used for the recruitment of teleost MSD genes.

2 | MATERIAL AND METHODS

2.1 | Sample collection

For high-quality genome reference sequencing, a single *Pangasianodon hypophthalmus* male was sampled from captive broodstock populations originating from Indonesia and maintained in the experimental facilities of ISEM (Institut des Sciences de l'Evolution de Montpellier, France). High-molecular-weight (HMW) genomic DNA (gDNA) was extracted from a 0.5-ml blood sample stored in a TNES-urea lysis buffer (TNES-urea: 4 M urea; 10 mM Tris-HCl, pH 7.5; 125 mM NaCl; 10 mM EDTA; 1% SDS). HMW gDNA was then purified using a slightly modified phenol-chloroform extraction method (Q. Pan et al., 2019). For the chromosome contact map (Hi-C), 1.5 ml of blood was taken from the same animal and slowly cryopreserved with 15% dimethyl sulphoxide (DMSO) in a Mr. Frosty Freezing Container (Thermo Scientific) at -80°C . For sex-linkage analyses and short-read genome sequencing, fin clips were sampled from 12 Pangasiid species and stored in 90% ethanol. *Pangasius djambal* fin clips were sampled from captive broodstock populations originating from Indonesia and maintained in the experimental facilities of ISEM. *P. gigas* fin clips were sampled on broodstock populations kept for a restocking program in Thailand. *Pangasius bocourti* and *P. conchophilus* fin clips were sampled at market places in Vietnam. *Pangasius elongatus*, *P. siamensis*, *P. macronema*, *P. larnaudii*, *P. mekongensis* and *P. krempfi* were wild samples collected in Vietnam. *Pangasius sanitwongsei* fin clip samples were obtained through the aquaculture trade and their precise population of origin is unknown.

2.2 | Chromosome-scale genome sequencing and assembly of *P. hypophthalmus*

2.2.1 | Oxford Nanopore sequencing

All library preparations and sequencing were performed using Oxford Nanopore Ligation Sequencing Kits SQK-LSK108 and SQK-LSK109 according to the manufacturer's instructions (Oxford Nanopore Technologies). For the SQK-LSK108 sequencing Kit, 90 μg of DNA was purified then sheared to 20-kb fragments using the megaruptor1 system (Diagenode). For each library, a DNA-damage repair step was performed on 5 μg of DNA. Then an end-repair-dA-tail step was performed for adapter ligation. Libraries were loaded onto nine R9.4.1 flowcells and sequenced on a GridION instrument at a concentration of 0.1 pmol for 48 h. For the SQK-LSK109 sequencing Kit, 10 μg of DNA was purified then sheared to 20-kb fragments using the megaruptor1 system (Diagenode). For this library, a one-step DNA-damage repair +end-repair-dA-tail procedure was performed on 2 μg of DNA. Adapters were then ligated to DNAs in the library. The library was loaded onto one R9.4.1 flowcell and sequenced on a GridION instrument at a concentration of 0.08 pmol for 48 h.

2.2.2 | 10X Genomics sequencing

The Chromium library was prepared according to 10x Genomics' protocols using the Genome Reagent Kit version 2. The library was prepared from 10 ng of HMW gDNA. Briefly, in the microfluidic Genome Chip, a library of Genome Gel Beads was combined with HMW template gDNA in master mix and partitioning oil to create Gel Bead-In-Emulsions (GEMs) in the Chromium apparatus. Each Gel Bead was then functionalized with millions of copies of a 10X barcoded primer. Dissolution of the Genome Gel Bead in the GEM released primers containing (i) an Illumina R1 sequence (Read 1 sequencing primer), (ii) a 16-bp 10x Barcode, and (iii) a 6-bp random primer sequence. The R1 sequence and the 10x barcode were added to the molecules during the GEM incubation. P5 and P7 primers, R2 sequence, and Sample Index were added during library construction. Ten cycles of PCR were applied to amplify the library. The library was sequenced on an Illumina HiSeq3000 using a paired-end format with read length of 150 bp with the Illumina HiSeq3000 sequencing kits.

2.2.3 | Hi-C sequencing

Hi-C library generation was carried out according to a protocol adapted from Rao et al. 2014 (Foissac et al., 2019). The blood sample was spun down, and the cell pellet was resuspended and fixed in 1% formaldehyde. Five million cells were processed for the Hi-C library. After overnight digestion with *HindIII* (NEB), DNA ends were labelled with Biotin-14-DCTP (Invitrogen) using the Klenow fragment (NEB) and religated. In total, 1.4 μg of DNA was sheared to an average size of

550 bp (Covaris). Biotinylated DNA fragments were pulled down using M280 Streptavidin Dynabeads (Invitrogen) and ligated to PE adaptors (Illumina). The Hi-C library was amplified using PE primers (Illumina) with 10 PCR amplification cycles. The library was sequenced using a HiSeq3000 (Illumina) in 150-bp paired-end format.

2.3 | Genome assembly

GridION data were trimmed using PORECHOP version 0.2.1 (<https://github.com/rrwick/Porechop>) and filtered using NANOFILT version 2.2.0 (De Coster, D'Hert, Schultz, Cruts & Van Broeckhoven, 2018) with the parameters `-l 3000` and `-q 7`. A *de novo* assembly was constructed with SMARTDENOVO (Ruan, 2015; Ruan, 2019), WTDDBG2 version 2.1 (Ruan & Li, 2020) and FLYE version 2.3.7 (Kolmogorov, Yuan, Lin & Pevzner, 2019), each with default parameters. The resulting assembly metrics were compared, and the draft assembly with the best metrics generated by SMARTDENOVO was kept and used as reference. This assembly was then further corrected using long reads. After mapping the trimmed and filtered GridION reads with MINIMAP2 version 2.11 (H. Li, 2018) with parameter `-x map-ont`, the assembly was polished using RACON (Vaser, Sović, Nagarajan & Šikić, 2017) version 1.3.1 with default parameters for three rounds. The assembly was then corrected using short reads. After mapping 10X short reads (with a 10× Genomics coverage of 116-fold the expected genome size) with LONG RANGER version 2.1.1, PILON (Walker et al., 2014) v1.22 was run with parameters `--fix bases,gaps --changes`. Again, three rounds of these short read polishing were performed. The final polished genome assembly was then scaffolded using Hi-C information. Reads were aligned to the draft genome using JUICER (Durand, Shamim, et al., 2016) with default parameters. A candidate assembly was then generated with a 3D *de novo* assembly (3D-DNA) pipeline (Dudchenko et al., 2017) with the `-r 0` parameter. The candidate assembly was manually reviewed using JUICEBOX (Durand, Robinson, et al., 2016) assembly tools. Gaps in this chromosome scaled assembly were filled using GAPCLOSER (https://github.com/CAFS-bioinformatics/LR_Gapcloser) version 1.1 with default parameters. Reads used to fill these gaps were GridION and PromethION reads filtered with NANOFILT and then corrected with CANU (Koren et al., 2017) version 1.6 using parameters `-correct genomeSize =753m -nanopore-raw`. The assembly was then corrected a final time using the short reads polishing pipeline.

2.3.1 | Genome analysis and protein-coding gene annotation

K-mer-based estimation of the genome size and heterozygosity were carried out with GENOMESCOPE version 2.0 (Ranallo-Benavidez, Jaron & Schatz, 2020; Vurtture et al., 2017). 10X reads were processed with JELLYFISH version 1.1.11 (Marçais & Kingsford, 2011) to count 21-mer with a maximum k-mer coverage of 10,000 and 1,000,000. BUSCO (Simão, Waterhouse, Ioannidis, Kriventseva & Zdobnov, 2015)

version 4.1.4 was run with parameters `-species zebrafish` and `-limit 10` on the single-copy orthologous gene library from the actinopterygii lineage. The first annotation step was to identify repetitive content using REPEATMASKER version 4.0.7 (<https://www.repeatmasker.org/>), DUST (Morgulis, Gertz, Schäffer & Agarwala, 2006) and TRF version 4.09 (Benson, 1999). A species-specific *de novo* repeat library was built with REPEATMODELER version 1.0.11 (<http://www.repeatmasker.org/RepeatModeler/>) and repeated regions were located using REPEATMASKER with the *de novo* and *Danio rerio* libraries. BEDTOOLS version 2.26.0 (Quinlan & Hall, 2010) was used to merge repeated regions identified with the three tools and to soft mask the genome. The MAKER3 genome annotation pipeline version 3.01.02-beta (Holt & Yandell, 2011) combined annotations and evidence from three approaches: similarity with fish proteins, assembled transcripts and *de novo* gene predictions. Protein sequences from 11 fish species (*Astyanax mexicanus*, *Danio rerio*, *Gadus morhua*, *Gasterosteus aculeatus*, *Lepisosteus oculatus*, *Oreochromis niloticus*, *Oryzias latipes*, *Poecilia formosa*, *Takifugu rubripes*, *Tetraodon nigroviridis*, *Xiphophorus maculatus*) found in Ensembl were aligned to the masked genome using EXONERATE version 2.4 (Slater & Birney, 2005). RNA-sequencing (RNA-seq) reads of *P. hypophthalmus* (NCBI BioProject PRJNA256973) from testis, ovary, embryo, brain, gills, heart, muscle, liver, head kidney, bones and intestine from the PhyloFish project (Pasquier et al., 2016) were used for genome annotation and aligned to the chromosomal assembly using STAR version 2.5.1b (Dobin et al., 2013) with `outWigType` and `outWigStrand` options to output signal wiggle files. CUFLINKS version 2.2.1 (Trapnell et al., 2010) was used to assemble the transcripts that were used as RNA-seq evidence. BRAKER version 2.0.4 (Hoff, Lange, Lomsadze, Borodovsky & Stanke, 2016) provided *de novo* gene models with wiggle files provided by STAR as hint files for GENEMARK (Hoff et al., 2016) and AUGUSTUS (Stanke et al., 2006) training. The best supported transcript for each gene was chosen using the Annotation Edit Distance (AED) quality metric (Eilbeck, Moore, Holt & Yandell, 2009).

2.3.2 | miRNA gene and mature miRNA annotation

Small RNA Illumina sequencing libraries were prepared using the NEXTflex Small RNA-Seq Kit version 3 (PerkinElmer) following the manufacturer's instructions and starting with the same total RNA extracts as for the Phylofish project (Pasquier et al., 2016). Total RNA was extracted using Trizol reagent (Euromedex) according to the manufacturer's instructions. Libraries were sequenced on an Illumina HiSeq 2500 sequencer and raw reads were preprocessed using CUTADAPT version 3.4 (Martin, 2011). All eight adult organ libraries (brain, gills, heart ventricle, skeletal muscle, intestine, liver, ovary and testis) were simultaneously analysed using Probst! (Thomas Desvignes, Batzel, Sydes, Eames & Postlethwait, 2019) selecting for read length of 17–25 nucleotides and with a minimum of five identical reads. Reads were then aligned to the species' reference genome using bbmapskimmer.sh version 37.85 of the BBMAP suite (<https://sourceforge.net/projects/bbmap/>). Gene and mature microRNA

(miRNA) annotations were performed as previously described (Thomas Desvignes et al., 2019) based on established miRNA gene orthologies among ray-finned fish species (Thomas Desvignes, Sydes, Montfort, Bobe & Postlethwait, 2021) and using previously published miRNA annotations in spotted gar, zebrafish, three-spined stickleback, Japanese medaka, shortfin molly and blackfin icefish as references (Braasch et al., 2016; Thomas Desvignes et al., 2019; Desvignes et al., 2021; Kelley et al., 2021; Kim et al., 2019). miRNA and isomiR nomenclature follow the rules established for zebrafish (Desvignes et al., 2015).

2.4 | Short-read sequencing and genome-guided assemblies of other Pangasiids

2.4.1 | Short-read sequencing

The *Pangasianodon gigas* and *Pangasius djambal* genomes were sequenced from male samples using an Illumina 2 × 250-bp format. DNA library construction was performed according to the manufacturer's instruction using the Truseq DNA nano library prep kit (Illumina). Briefly, gDNA was quantified using the HS dsDNA Assay kit on the Qubit (Invitrogen). In total, 200 ng of gDNA was sonicated on a Bioruptor (Diagenode). Sonicated gDNA was end repaired and size selected on magnetic beads aiming for fragments of an average size of 550 bp. Selected fragments were adenylated on their 3' ends before ligation of Illumina's indexed adapters. The library was amplified using eight PCR cycles and verified on a Fragment Analyzer using the HS NGS fragment kit (Agilent). The library was quantified by qPCR using the KAPA Library quantification kit (Roche, ref. KK4824) and sequenced on half a lane of HiSeq2500 in paired-end 2 × 250nt using the clustering and SBS rapid kit following the manufacturer's instructions. All other species were sequenced from males or unknown sex individuals (see Table S2), using an Illumina 2 × 150-bp strategy according to Illumina's protocols using the Illumina TruSeq Nano DNA HT Library Prep Kit. Briefly, DNA was fragmented by sonication, size selection was performed using SPB beads (kit beads) and adaptors were ligated to be sequenced. Library quality was assessed using an Advanced Analytical Fragment Analyzer and libraries were quantified by qPCR using the Kapa Library Quantification Kit. DNA-seq experiments were performed on one Illumina NovaSeq S4 lane using a paired-end read length of 2 × 150 bp with the Illumina NovaSeq6000 Reagent Kits.

2.4.2 | Assembly and annotation

The *Pangasianodon gigas* and *Pangasius djambal* genomes were assembled from 2 × 250-bp short reads using the DISCOVARDENOVO assembler (<https://github.com/bayolau/discovardenovo/>) with default parameters. For *Pangasius sanitwongsei*, *P. conchophilus*, *P. bocourti*, *P. larnaudii*, *P. mekongensis* and *P. krempfi*, 2 × 150-bp reads were assembled using SPADES version 3.11.1 (Bankevich et al., 2012) and

then purged using PURGE_DUPS (Guan et al., 2020). The *Pangasius elongatus*, *P. macronema* and *P. siamensis* 2 × 150-bp short reads were assembled with SPADES version 3.14.1 instead of version 3.11.1 because of a higher individual genome heterozygosity (>1%, calculated with GENOMESCOPE as described above), followed by a more stringent purge with REDUNDANS version 0.14a (Pryszcz & Gabaldón, 2016). All these species were then assembled into pseudochromosomes using a reference-guided strategy and the "query assembled as reference" function from DGENIES version 1.2.0 (Cabanettes & Klopp, 2018), and the GENO_Phyp_1.0 *P. hypophthalmus* assembly used as a reference. Genes from the NCBI annotation of GENO_Phyp_1.0 were then mapped to chromosome-scale assemblies using LIFTOFF (Shumate & Salzberg, 2021) except for *Pangasianodon gigas* and *Pangasius djambal* genomes that were annotated with the same in-house annotation pipeline described for the *P. hypophthalmus* genome.

2.5 | Species and gene phylogenies

Whole-genome species phylogeny analysis was carried out with protein gene annotation from our 12 Pangasidae species combined with protein sequences from *Ictalurus punctatus* (siluriformes) as a Pangasidae outgroup species. Outgroup species protein sequences were retrieved from Ensembl release 103 (Howe et al., 2021). Orthogroups were identified using ORTHOFINDER (Emms & Kelly, 2019), followed by multiple sequence alignment of concatenated one-to-one orthologues ($n = 8,151$) using MAFFT version 7.475 (Katoh & Standley, 2013). Species tree inference was performed via IQ-TREE 2 (Minh et al., 2020), the latter using a standard nonparametric bootstrap ($r = 100$).

Gene and protein phylogenetic reconstructions were performed on all *amhr2*/*Amhr2* homologous sequences from 28 catfish species along with *amhr2* sequences from *Astyanax mexicanus* (characiformes) and *Electrophorus electricus* (gymnotiformes) as siluriformes outgroups. Full-length coding sequences (CDS) were predicted based on their genomic and protein sequence annotation or retrieved from GenBank (see Table S2 and multi-fasta files of these sequences are publicly available at <https://doi.org/10.15454/M3HYAX>). To verify the tree topology of *amhr2*/*Amhr2* homologues, besides complete protein and cDNA sequences, we also constructed phylogenetic trees with only the first and second codons of the CDS (Lemey, 2009). All putative CDS and protein sequences were then aligned using MAFFT (version 7.450) (Katoh & Standley, 2013). Residue-wise confidence scores were computed with GUIDANCE 2 (Sela, Ashkenazy, Katoh & Pupko, 2015), and only well-aligned residues with confidence scores above 0.99 were retained. Phylogenetic relationships among the *amhr2* sequences were inferred with both maximum-likelihood implemented in IQ-TREE (version 1.6.7) (Minh et al., 2020), and Bayesian methods implemented in PHYLOBAYES (version 4.1) (Lartillot, Lepage & Blanquart, 2009). More precisely, alignment files from either full-length cDNA, third-codon-removed cDNA or full-length proteins were used for model selection and tree inference with IQ-TREE (version 1.6.7) (Minh et al., 2020) with 1,000 bootstraps

and the 1,000 SH-like approximate likelihood ratio test for robustness. The same alignment files were run in a Bayesian framework with PHYLOBAYES (version 4.1) (Lartillot et al., 2009) using the CAT-GTR model with default parameters, and two chains were run in parallel for ~2,000 cycles with the first 500 cycles discarded as burn-in until the average standard deviation of split frequencies remained ≤ 0.001 . The resulting phylogenies were visualized with FIGTREE (version 1.44).

2.6 | Selection analysis on *amhr2* sequences

Selection analysis was performed on the *amhr2* phylogeny using GODON (Davydov, Salamin & Robinson-Rechavi, 2019). Analyses were performed separately for (i) exons conserved in both *amhr2a* and *amhr2by* ("conserved exons") and (ii) the exon region found only in *amhr2a* ("first exons"). Three codon models were used: M8 (Yang, Nielsen, Goldman & Pedersen, 2000), M8 with codon gamma rate variation (Davydov et al., 2019) and the branch-site model (Zhang, Nielsen & Yang, 2005) (conserved exons only). For the branch-site model, the branch leading to the *amhr2by* clade was set as the foreground branch.

2.7 | Transcriptome analyses

To explore the expression profiles of *amhr2a*, *amhr2by* and other potential genes (i.e., *amh*, *dmrt1*, *gsdf*, *gdf6a*, *gdf6b* and *sox3*) known to be used as MSD genes in other species, reads from *P. hypophthalmus* adult organs and embryos (Pasquier et al., 2016) were mapped on the complete *P. hypophthalmus* reference transcriptome using BWA MEM version 0.7.17 (H. Li, 2013). Unique mapped reads were then filtered and a raw count matrix was generated with HTSEQ-COUNT (Anders, Pyl & Huber, 2015) and normalized using DESEQ2 (Love, Huber & Anders, 2014). Genes of interest were extracted from this complete transcriptome data set and missing values were replaced by a minimal value (0.1) in the normalized raw count matrix. Hierarchical classification was carried out after log transformation and gene median centring using the CLUSTER 3.0 software (de Hoon, Imoto, Nolan & Miyano, 2004) with an uncentred correlation similarity metric and an average linkage clustering method.

2.8 | Read-coverage analyses around the *amhr2a* and *amhr2by* loci in Pangasiids

To assess whether *amhr2by* is a potential Y-specific gene in species for which whole genome sequencing was only obtained from one sample, we computed the read coverage throughout the genome and extracted the read coverage information around the *amhr2a* and *amhr2by* loci. In *P. hypophthalmus*, ONT reads were mapped on its own genome assembly using MINIMAP version 2.11 (Li, 2018). In other Pangasiids, Illumina paired-end reads were mapped onto the *P.*

hypophthalmus genome assembly using BWA version 0.7.17 (Li, 2013), indexed using SAMTOOLS version 1.8 (Li et al., 2009) and sorted by PICARD SortSam. A pileup file was then generated using SAMTOOLS mpileup (H. Li et al., 2009) with per-base alignment quality disabled and (-B). Subsequently, a sync file containing the nucleotide composition for each position in the reference was created from the pileup file using POPOULATION MPILEUP2SYNC version 1.201 with a minimum quality of 20 (-min-qual 20) (Kofler, Pandey & Schlötterer, 2011). Read depth was then calculated in a 10-kb nonoverlapping window using PSASS (version 2.0.0, <https://doi.org/10.5281/zenodo.2615936>).

2.9 | Primer design

P. hypophthalmus amhr2a and *amhr2by* genes were aligned with BIOEDIT version 7.0.5.3 and specific primers were designed based on this alignment to select highly divergent positions for each paralogue (Figure S1). Selected primer sequences were forward: 5'-GGAGTCTATAAACCCGTGGTAGC-3', and reverse: 5'-CTATGTCACGCTGAACCTCCAGTGT-3' (expected amplicon size: 153 bp) for the *amhr2by* gene and forward: 5'-GGAGTCTATAAGCCAGCGGTGGCT-3', and reverse: 5'-CTATGCCAGAATAACCCCTGCAATGC-3' (expected amplicon size: 142 bp) for the *amhr2a* gene.

2.10 | DNA extraction for PCR sex genotyping

For PCR genotyping of *amhr2a* and *amhr2by*, DNA from fin clips was extracted using a Chelex-based extraction method. Briefly, a piece of fin clip from each sample was placed into a PCR tube, and then 150 μ l 5% Chelex and 20 μ l 1 mg ml⁻¹ proteinase K were added to each tube. Tubes were then vortexed and quickly spun down. After that, samples were incubated for 2 h at 56°C followed by boiling for 10 min at 99°C. DNA was then centrifuged at 7,500 g for 5 min and diluted to 1:2 with double distilled water. Genotyping PCRs were run in 12.5 μ l with 1.25 μ l JumpStart PCR buffer 10X, 0.125 μ l 25 mM dNTP, 0.25 μ l 10 μ M forward and reverse primers, 8.5 μ l ddH₂O and 2 μ l DNA. PCR cycling conditions were: 95°C for 3 min as initial denaturation, then 35 cycles for amplification with denaturation at 95°C for 30 s, annealing at 52°C for 30 s and extension at 72°C for 30 s, and finally a further extension at 72°C for 30 s and hold at 4°C.

3 | RESULTS

3.1 | A high-quality chromosome-scale genome assembly of *P. hypophthalmus*

A high-quality reference genome of a male *Pangasianodon hypophthalmus* was sequenced using a combination of 10 \times Linked-Reads, Oxford Nanopore long reads and a chromosome contact map (Hi-C). Its genome size based on the kmer linked-reads distribution

was estimated to ~810 Mb including, respectively 65% and 35% of unique and repeated sequences. The level of heterozygosity of this *P. hypophthalmus* genome was estimated at ~1.2%. The integration of all sequencing data provided a genome assembly size of 760 Mb (93% of the kmer estimated size), containing 612 contigs, a contig and scaffold N50 of respectively 9.7 and 26.4 Mb (Table 1; Table S1), and 99.2% of all sequences anchored onto 30 chromosomes after Hi-C integration (see assembly metrics and comparison with other genome assemblies in Table 1 and Table S1). Combining *de novo* gene predictions, homology to teleost proteins and evidence from transcripts, 21,234 protein-coding genes were annotated with the following BUSCO (Simão et al., 2015) scores: Complete: 91.5% (Single: 90.2%, Duplicated: 1.3%), Fragmented: 2.0%, Missing: 6.5%, in our male *P. hypophthalmus* reference genome using our in-house genome annotation protocol. Because our *P. hypophthalmus* genome assembly has been derived by NCBI to produce a Reference Sequence (RefSeq) record (GCF_009078355.1) and was annotated by the NCBI Eukaryotic Genome Annotation Pipeline, the NCBI annotation will be used hereafter as reference in the following text. In addition to protein-coding genes, 323 miRNAs and 389 mature miRNAs were annotated using Illumina small-RNA sequencing data from a panel of eight organs. Gene and mature miRNA annotations as well as analysed expression patterns are publicly available on FishmiRNA (<http://www.fishmirna.org/>) (Thomas Desvignes et al., 2022). This genome-wide miRNA annotation represents the first exhaustive miRNA annotation available for a Pangasiid species.

3.2 | Characterization of a male-specific *amhr2* duplication in *P. hypophthalmus*

Because many teleost MSD genes evolved from the duplication of an autosomal “usual suspect” gene (see Introduction for references), we first searched for potential duplicates of *dmrt1*, *amh*, *amhr2*, *sox3*, *gsdf* and *gdf6* genes in the *P. hypophthalmus* genome assemblies using TBLASTN searches with the following bait proteins from the *Ictalurus punctatus* genome assembly (XP_017308041.1,

XP_017333187.1, XP_017331275.1, XP_017330512.1, XP_017347335.1, XP_017308337.1 and XP_017309564.1). We found no gene duplication for *dmrt1*, *amh*, *sox3*, *gsdf* or *gdf6* (*gdf6a* and *gdf6b*), but two *amhr2* homologues were found in the two male *P. hypophthalmus* assemblies (i.e., our own GENO_Phyp_1.0 assembly and the VN_pangasius assembly that is already published and publicly available; Kim et al., 2018), while only one *amhr2* gene was detected in the publicly available female *P. hypophthalmus* ASM1680104v1 assembly (Gao et al., 2021). In the GENO_Phyp_1.0 *P. hypophthalmus* assembly, these two *amhr2* homologues, LOC113540131 (annotated as bone morphogenetic protein receptor type-2-like) and LOC113533735 (annotated as anti-Müllerian hormone type-2 receptor-like), are located respectively on chromosome 4 (Chr04:32,081,919–32,105,291) and 10 (Chr10: 26,334,822–26,348,340). Annotation of *amhr2* genes as *bmpr2-like* (LOC113540131) has been already described (Adolfi, Nakajima, Nóbrega & Schartl, 2019) and is probably quite frequent in automatic genome annotations because these two genes belong to the same type II receptor family. The single *amhr2* locus found in the female ASM1680104v1 assembly (in ASM1680104v1 Chr04) is on chromosome 4 and shares 99% identity over 13.5 kb (100% overlap) with LOC113533735, and 87% identity on only 3% overlapping regions with LOC113540131. Specific primers were designed based on an alignment of the *P. hypophthalmus* LOC113540131 and LOC113533735 genes in order to select highly divergent positions for each paralogue. Using these primers, we genotyped *P. hypophthalmus* males ($N = 12$) and females ($N = 11$) and found that LOC113540131 is significantly linked with maleness ($p = 7.12e^{-05}$) with a single positive outlier among 11 phenotypic females (see Table 2). In contrast, LOC113533735 was detected in all males and females (Figure 1). These genotyping results, along with the absence of LOC113540131 in the female ASM1680104v1 assembly, strongly support the hypothesis that LOC113540131 is a Y-specific male-specific, gene. We thus called the LOC113540131 gene *amhr2by*, as the male-specific Y chromosome paralogue of the autosomal LOC113533735 gene named *amhr2a*.

TABLE 1 Comparison of our *Pangasianodon hypophthalmus* reference genome assembly metrics (our study) with the other *P. hypophthalmus* available assemblies

Assembly	GCA_003671635.1	Our study	GCA_016801045.1
Release date	05/04/2018	10/22/2019	14/10/2020
Sex of the sequenced individual	Male	Male	Female
Total sequence length (Mb)	715.8	758.9	742.5
Total ungapped length (Mb)	696.5	758.8	742.3
Number of contigs	2,334	612	808
Contig N50 (Mb)	0.06	9.7	3.48
Contig L50	3,254	18	63
Total number of chromosomes	N.A.	30	30
Number of component sequences (WGS or clone)	568	150	402

Abbreviation: N.A., not applicable.

3.3 | Comparison of *P. hypophthalmus amhr2by* and *amhr2a* and their inferred proteins

Overall, the predicted structure of the autosomal *P. hypophthalmus amhr2a* and the canonical vertebrate *Amhr2* are similar with the same number of introns and exons. The MISTA (Frazer, Pachter, Poliakov, Rubin & Dubchak, 2004) alignments of *P. hypophthalmus amhr2a* and *amhr2by* genes along with their CDS (Figure 2a) show that these two genes display some sequence identity only within their shared exons, with no significant homology detected in their intronic, 3' untranslated region (UTR) and 5'UTR sequences (Figure 2a). In addition, the *amhr2by* gene is lacking the first two exons of *amhr2a*, and the third *amhr2by* exon is also truncated. The *amhr2by* and *amhr2a* CDS (Figure S1) share 78.78% identity on 1,164 bp of overlapping sequences (78% of the *amhr2a* CDS that is 1,455 bp long). Correspondingly, the two deduced proteins share 70.32% identity over 380 overlapping amino acids, and *Amhr2by* lacks 112 amino acids at its N-terminal extremity corresponding to two first exons and part of exon 3 of *Amhr2a*. (Figure 2b,c). Hence, the *P. hypophthalmus Amhr2by* translates as an N-terminal-truncated type II receptor lacking its whole extracellular domain mediating ligand binding, while overall the remainder of the other functional domains (transmembrane and serine-threonine kinase domain) remain similar between *Amhr2a* and *Amhr2by* (Figure 2b,c).

3.4 | Expression of *amhr2by* and *amhr2a* in *P. hypophthalmus* adult tissues

Using *P. hypophthalmus* RNA-seq from the PhyloFish database (Pasquier et al., 2016), we examined the organ expression of *amhr2a* and *amhr2by* along with a series of SD genes previously identified in other teleosts, namely *amh*, *dmt1*, *gsdf*, *gdf6a*, *gdf6b* and *sox3*. Among these genes, *amh*, *dmt1* and *gsdf* display predominant expression in the adult testis and/or ovary with a much lower expression in the eight additional somatic organs examined or in embryos (Figure 3a). The two *amhr2* genes also have a gonadal-predominant expression pattern with *amhr2a* being expressed in both ovary and testis and *amhr2by* being strictly expressed in the testis as expected for a Y chromosome SD gene (Figure 3b). The two *gdf6* paralogues (*gdf6a*, *gdf6b*) and *sox3* have no expression or a low expression in gonads and are more expressed in embryos for *sox3* and *gdf6a* or in bones and brain for *sox3*.

3.5 | Sex-linkage of *amhr2by* in Pangasiids

To explore the evolution of *amhr2by* in Pangasiids, we obtained gDNA samples from 11 additional Pangasiid species with at least some specimens being phenotypically sexed for four of these species (Table S2). Samples from fish that were phenotypically sexed (i.e., *Pangasianodon gigas*, *Pangasius djambal*, *Pangasius conchophilus*

and *Pangasius bocourti*) were PCR-genotyped to explore the potential conservation of *amhr2by* male sex-linkage in Pangasiids. In three of these species, *amhr2by* was found to be significantly associated with male phenotype ($p < 8.528e^{-04}$) (Table 2; Figure S2), and in *Pangasianodon gigas*, the association was not significant ($p = .3865$) due to the combination of low sample size (three males and three females) and the presence of one female outlier (Table 2; Figure S2). To complement this genotyping information, one male individual of *Pangasianodon gigas*, *Pangasius djambal*, *P. conchophilus* and *P. bocourti* and one individual of unknown sex for *Pangasius elongatus*, *P. siamensis*, *P. sanitwongsei*, *P. macronema*, *P. larnaudii*, *P. mekongensis* and *P. krempfi* were sequenced using Illumina short-read strategies. These genomic short-read sequences were assembled and anchored using a reference-guided strategy (Lischer & Shimizu, 2017) on the *P. hypophthalmus* chromosome assembly, and the NCBI gene annotation of GENO_Phyph_1.0 was lifted over to these assemblies (see genome and annotation metrics in Table S1). The *amhr2a* genes were extracted from all these guided assemblies, and *amhr2by* homologues were extracted from the four male assemblies, that is *Pangasianodon gigas*, *Pangasius djambal*, *P. conchophilus*, and *P. bocourti* as well as from the unknown sex assemblies of *Pangasius sanitwongsei*, and *P. krempfi*. To better explore sex-linkage in species for which we only sequenced a single individual, read coverage was explored around the *amhr2a* and *amhr2by* loci using the genome *P. hypophthalmus* as reference (Figure S3). Under the hypothesis that *amhr2by* is also a male-specific Y chromosomal gene in additional Pangasiids, we expected half coverage around *amhr2by* in males (hemizygous in XY) and an average read coverage around the autosomal *amhr2a*. In agreement with that hypothesis, half coverage was found around the *amhr2by* locus for all species in which *amhr2by* was identified, namely the male individuals of *Pangasianodon hypophthalmus* and *P. gigas* and *Pangasius djambal*, *P. conchophilus* and *P. bocourti* and individuals of unknown sex in *Pangasius sanitwongsei* and *P. krempfi*. This result supports hemizygoty of *amhr2by* in these species as expected for a Y chromosomal gene. In other species (i.e., *Pangasius elongatus*, *P. siamensis*, *P. macronema*, *P. larnaudii*, and *P. mekongensis*), no conclusion can be drawn because the absence of *amhr2by* in these individuals could be because they are XX females without a Y chromosome and an *amhr2by* gene, or these species may have lost *amhr2by* as a Y chromosome gene.

3.6 | Evolution of *amhr2* in Siluriformes

These whole-genome annotations were combined with protein sequences from channel catfish, *Ictalurus punctatus* (Siluriformes, Ictaluridae) used as a Pangasiid outgroup, and 8,151 groups of one-to-one orthologues were used after concatenation to construct a whole-genome species tree inference (Figure 4). In addition, all Pangasiid *amhr2* sequences deduced from our genomic resources were used for phylogenetic analyses with other available catfish *amhr2* genes (Table S2), along with *amhr2* from a gymnotiform

(*Electrophorus electricus*) and a characiform (*Astyanax mexicanus*) as the closest species outgroups to the order Siluriformes. The topologies of all trees, that is using maximum-likelihood and Bayesian methods on proteins, CDS and CDS with third codons removed (see Materials and Methods), were all congruent in showing that most of the *amhr2* from the sub-order Siluroidei (Sullivan et al., 2006) cluster with the Pangasiid *amhr2a*, and that outside the Pangasiid family, only a single species (*Pimelodus maculatus*, Pimelodidae) has an *amhr2* duplication clustering with the *amhr2by* sequences (Figure 5; Figure S4). Within the Siluriformes, a single *amhr2* in *Corydoras* sp. (Callichthyidae, Loricarioidei) roots the *amhr2a* and *amhr2b* duplications (Figure 5; Figure S4), suggesting that *amhr2b* (*Pimelodus maculatus*) and *amhr2by* (Pangasiids) arose from an ancient duplication/insertion event at the root of the Siluroidei radiation that is dated to ~100 million years ago (Kappas et al., 2016). We also searched for selection acting on the Pangasiid *amhr2* sequences, but detected no statistically significant signal of positive selection (Table 3) for either all exons conserved in both *amhr2a* and *amhr2by* ("conserved exons") or for the exon region found only in *amhr2a* ("first exons").

TABLE 2 Sex-linkage of *amhr2by* in five different Pangasiid species. Associations between *amhr2by* specific PCR amplifications and sex phenotypes are provided for both males and females (number of positive individuals for *amhr2by*/total number of individuals) along with the *p* value of association with sex that was calculated for each species based on the Pearson's chi-square test with Yates' continuity correction

Species	Males	Females	<i>p</i> value
<i>Pangasianodon hypophthalmus</i>	12/12	1/11	7.12e-05
<i>Pangasianodon gigas</i>	3/3	1/3	.3865 ^a
<i>Pangasius bocourti</i>	12/12	1/20	8.411e-07
<i>Pangasius conchophilus</i>	22/22	0/10	1.559e-07
<i>Pangasius djambal</i>	6/6	0/9	8.528e-04

^aNon-significant association with sex.

4 | DISCUSSION

The Pangasiid family contains both important aquaculture species (Lazard et al., 2009) and key ecological catfish species (Eva et al., 2016) in many south Asian countries. Here, we present a reference genome for striped catfish, *Pangasianodon hypophthalmus*, and provide an additional high-quality genomic resource combining long-read sequencing and a chromosomal assembly for this species. This *de novo* genome (GENO_Phyp_1.0, GCA_009078355.1) was assembled into 30 large scaffolds that probably correspond to the 30 chromosomes reported previously in cytological studies (Sreeputhorn et al., 2017). This assembly also improves the metrics of the previously publicly available male assembly VN_pangasius (GCA_003671635.1) that was not anchored on chromosomes (O. T. P. Kim et al., 2018), and is comparable in terms of assembly metrics to the newest female ASM1680104v1 (GCA_016801045.1) chromosome-anchored assembly (Z. Gao et al., 2021). Furthermore, the *P. hypophthalmus* miRNA annotation represents the first exhaustive miRNA annotation available for a Pangasiid species, and completes among Siluriformes the two annotations available for catfishes in black bullhead, *Ameiurus melas* (Thomas Desvignes et al., 2022), and channel catfish, *Ictalurus punctatus* (Kozomara, Birgaoanu & Griffiths-Jones, 2019). This novel resource will enable studies of Pangasiid and more broadly of Siluriformes miRNA roles. In addition to this *P. hypophthalmus* chromosome-anchored assembly, we also provided short-read genome sequencing for 11 additional Pangasiid species belonging to the genera *Pangasianodon* (one additional species) and *Pangasius* (10 additional species). These short-read assemblies have been anchored and annotated on our reference *P. hypophthalmus* genome assembly and now present a large public set of genomic resources for the Pangasiid family.

Phylogenetic relationships within Siluriformes are still debated with no consensus for clear placement of some families within this order (Kappas et al., 2016; Sullivan et al., 2006). At a broader scale, however, it is generally accepted that the sub-order Loricarioidei (defined also as a super-family) containing the armored catfish families (Callichthyids and Loricariids) is the earliest-diverging

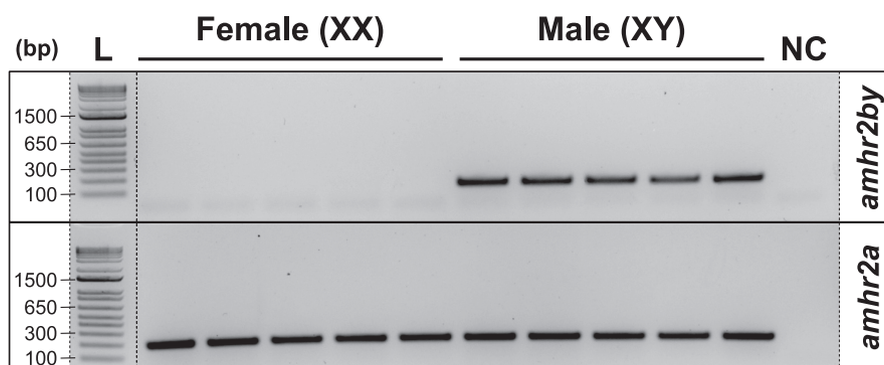


FIGURE 1 Sex genotyping in *Pangasianodon hypophthalmus*. The *amhr2a* sequence (bottom panel) is PCR amplified in both male and female samples, while the *amhr2by* sequence (upper panel) is only amplified in male samples, indicating that *amhr2by* is male-specific, that is Y-chromosome linked. L = 1-kb Plus DNA Ladder (Invitrogen). (bp) = DNA ladder fragment sizes in nucleotide base pairs. NC = negative control (water)

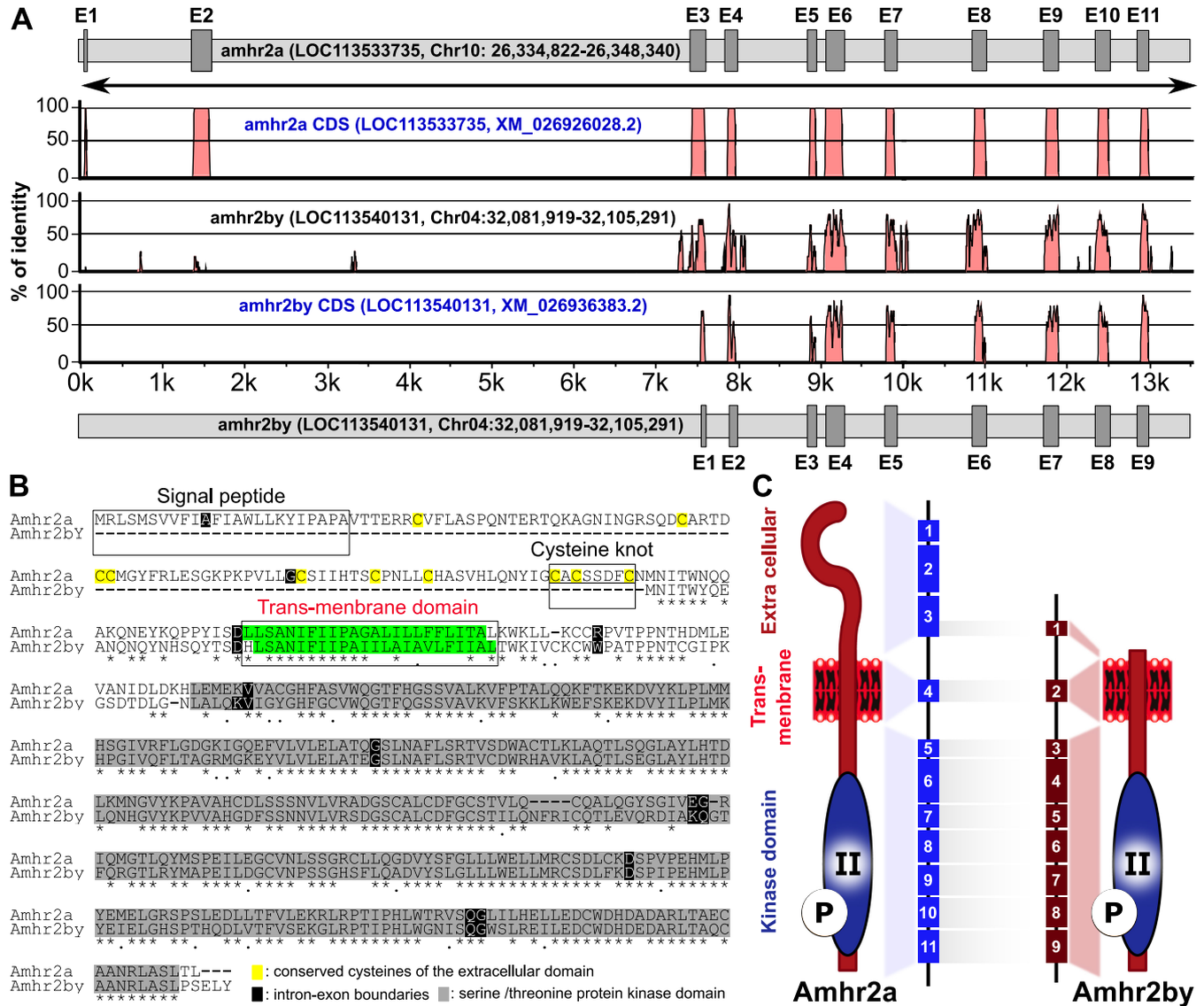


FIGURE 2 Structure of *amhr2a* and *amhr2by* and deduced proteins in *Pangasianodon hypophthalmus*. (a) Identity plot of the alignment of the autosomal *amhr2a* with the Y-linked *amhr2by* sequences. Exons (E) of both *amhr2* genes are depicted with grey boxes. (b) CLUSTAL W alignment of Amhr2a and Amhr2by proteins. Identical amino acids are shaded in grey and conserved cysteines in the extracellular domain of Amhr2a are highlighted in yellow. The different domains (signal peptide, cysteine knot and transmembrane domain) of the receptors are boxed. Intron-exon boundaries are boxed in black for both receptors. (c) Schematic representation of *P. hypophthalmus* autosomal Amhr2a and Y-linked Amhr2bY proteins showing the architecture of Amh receptors and the correspondence between exons of Amhr2a and Amhr2by, highlighting the absence of the entire extracellular domain in the truncated Amhr2bY

Siluriformes clade with the Diplomystoidei sub-order being the sister group to the remaining Siluroidei sub-order (Kappas et al., 2016; Sullivan et al., 2006). Pangasiids belong to the Siluroidei sub-order and have been characterized as the sister group to either Ictaluridae and Cranoglanididae (Kappas et al., 2016) or Schilbidae (Villela et al., 2017). Their phylogeny has been explored using both mitochondrial and nuclear markers (Karinthanyakit & Jondeung, 2012; Pouyaud et al., 2016). Here, using a phylogenomic approach (Delsuc, Brinkmann & Philippe, 2005), we were able to determine the precise phylogenetic relationships among the 12 Pangasiid species for which we produced genome sequencing. Our results confirmed the basal position of the genus *Pangasianodon* as already described

(Karinthanyakit & Jondeung, 2012; Na-Nakorn et al., 2006; Pouyaud et al., 2016) and, although we did not sequence any *Helicophagus* or *Pseudolais* species, the results allowed us to resolve the taxonomic positions of several *Pangasius* species (Karinthanyakit & Jondeung, 2012).

The molecular basis of genetic SD has been explored in only a few catfishes, with reports on the identification of male sex-specific sequences supporting an XX/XV SD system in *Pseudobagrus ussuriensis* (Z.-J. Pan, Li, Zhou, Qiang & Gui, 2015) and *Pelteobagrus (Tachysurus) fulvidraco* (Dan, Mei, Wang & Gui, 2013; Wang, Mao, Chen, Liu & Gui, 2009) from the family Bagridae, and in *Clarias gariepinus* from the family Clariidae (Kovács, Egedi, Bártfai &

FIGURE 3 Expression of some sex determination candidate genes in adult organs of *Pangasianodon hypophthalmus*. (a) Hierarchical clustering heatmap analysis of some sex determination genes previously identified in other teleosts, namely *amh*, *dmrt1*, *gsdf*, *gdf6a*, *gdf6b* and *sox3* in different organs and embryos of *P. hypophthalmus*. Each coloured cell corresponds to a relative expression value (see key on the left). (b) Normalized read counts of *amhr2a* and *amhr2by* in whole organs and embryos *P. hypophthalmus* transcriptomes

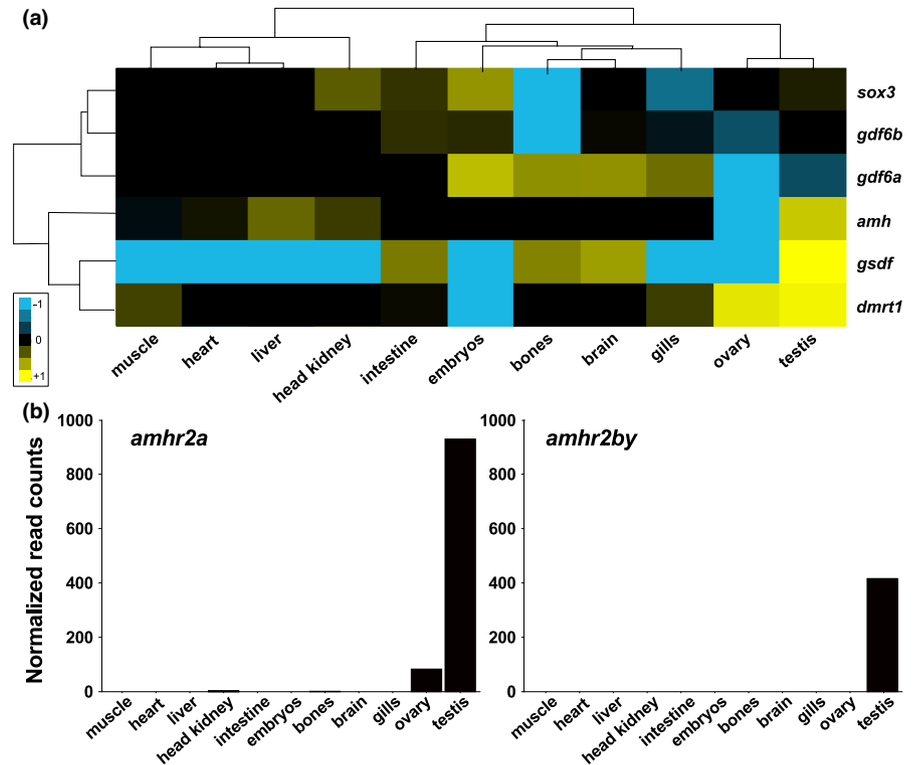
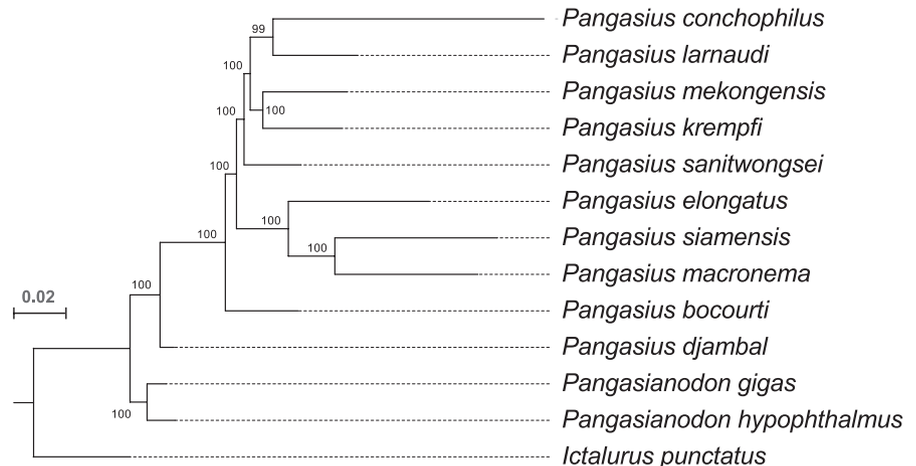


FIGURE 4 Whole-genome-based phylogenetic tree of all sequenced Pangasiid species. Maximum-likelihood phylogeny of 12 Pangasiidae species with *Ictalurus punctatus* (siluriformes) as a Pangasiidae outgroup, based on alignment of concatenated protein sequences. Branch length scale corresponds to 0.02 amino acid substitutions per site. Support values at each node are proportions of 100 standard nonparametric bootstrap replicates



Orbán, 2000). In the Ictalurid channel catfish, *Ictalurus punctatus*, based on whole genome sequencing of a YY individual and genome-wide analyses, an isoform of the breast cancer anti-resistance 1 (*bcar1*) gene has been characterized as the male MSD gene (Bao et al., 2019). In Pangasiids, genetic sex-markers have been searched without success in *P. hypophthalmus* and *P. gigas* (Sriphairoj et al., 2007). In our study, based on chromosome-scale genome assemblies of many Pangasiid species, transcriptomic data (Pasquier et al., 2016) and sex-linkage analyses, we identified a male-specific duplication of the *amhr2* (*amhr2by*) gene as a potentially conserved male MSD gene in that fish family. The role of *Amhr2* as an MSD gene has been functionally characterized in the tiger pufferfish, *Takifugu rubripes* and ayu, *Plecoglossus altivelis* (Kamiya et al., 2012; Nakamoto et al., 2021), and strongly suggested by sex-linkage information in common seadragon, *Phyllopteryx taeniolatus*, alligator pipefish, *Syngnathoides biaculeatus* (Qu et al., 2021), other species

of pufferfishes (Duan et al., 2021; Gao et al., 2020; Kamiya et al., 2012) and yellow perch, *Perca flavescens* (Feron et al., 2020). In addition, the anti-Müllerian hormone, Amh, which is the cognate ligand of Amhr2, has also been demonstrated or suggested as an MSD gene in a few fish species (Hattori et al., 2012; Li et al., 2015; Pan et al., 2019, 2021; Song et al., 2021). Our results thus provide a new example of the repeated and independent recruitment of Amh and TGF β pathway members in fish genetic SD (Pan et al., 2021). Although formal proof that this *amhr2by* gene is a conserved MSD gene in Pangasiids will require additional gene expression analyses and functional demonstrations, our results have application as a useful marker for sex control in many Pangasiid species in aquaculture. Sex dimorphic growth is often one of the main reasons for breeding all-male or all-female populations for aquaculture purposes. This could be applied for instance in *Pangasius djambal*, a species in which females have a faster growth rate above 3 kg,

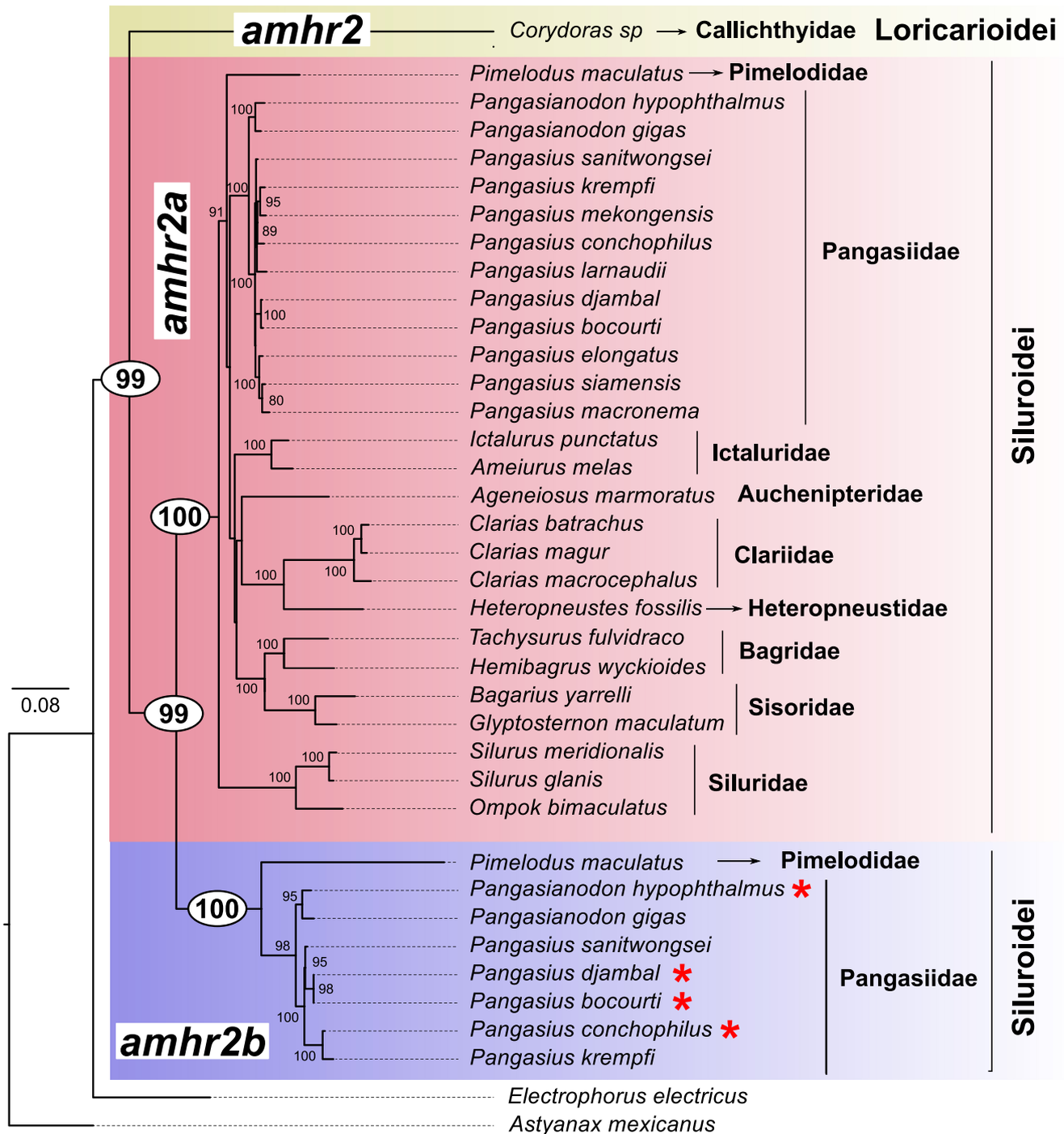


FIGURE 5 Phylogeny of *amhr2* in catfishes reveals an ancient *amhr2a/amhr2b* duplication in Siluroidei. Maximum-likelihood phylogeny of *amhr2* coding sequences (see Figure S4 for other phylogenetic approaches) from 28 catfish species with *amhr2* coding sequences from *Astyanax mexicanus* (Characiformes) and *Electrophorus electricus* (Gymnotiformes) as Siluriformes outgroups. Family and suborders are given for all catfish species on the right panel of the figure. The *amhr2b* cluster including the *amhr2by* of Pangasiids is shaded in purple, the *amhr2a* cluster shaded in red and the *Corydoras sp amhr2* preduplication is shaded in yellow. The branch length scale representing the number of substitutions per site is given at the root of the Siluriformes tree. Bootstrap values are given only for values over 80 and are inserted in a white circle at key nodes for the Siluroidei *amhr2* duplication. Species for which the association between the genome presence of *amhr2by* and male phenotype was significant are shown with a red asterisk (see Table 2 for additional details)

probably linked to the early maturation of males (Legendre et al., 2000). In addition, our results will also allow better management of breeders used for restocking in the large and endangered Mekong giant catfish, *Pangasianodon gigas*, because maturation takes as long

as 16–20 years in this species (Sripairoj et al., 2007). However, even if we found highly significant male sex-linkages of *amhr2by* in four pangasiid species, we also identified a few *amhr2by* positive females (three over 53 females genotyped in five species). This

TABLE 3 Positive selection analyses reveal no significant signal of positive selection on Pangasiid *amhr2*. *p*-values were computed using a chi-square distribution with 1 degree of freedom. None of the *p*-values passed a Bonferroni-corrected limit of significance: $.05/3 = .0167$. DlnL = difference in log-likelihood between models with and without positive selection; likelihood ratio test statistic

Model	Conserved exons		First exons	
	DlnL	<i>p</i> -value	DlnL	<i>p</i> -value
M8 gamma	0.0000000	.5	3.37482	3.309990e-02
Branch-site gamma	0.2976536	.2926786	—	—

discrepancy from an expected complete sex-linkage for a potential MSD gene could be due to phenotyping errors, as none of the individuals we used were sexed using gonadal histology, which would have prevented any misidentification. However, phenotype/genotype discrepancies could be also the result of sex-reversals that are sometimes detected in wild-type populations of some other fish species. However, these sex-reversals, which are often described as the result of environmental effects on SD (Baroiller et al., 2009; Kobayashi et al., 2013; Ospina-Alvarez & Piferrer, 2008), are mostly observed as female-to-male reversal, in contrast to what we observed here in Pangasiids. More accurate sex-linkage analyses would be now required to explore if such *amhr2by*-positive females are incorrectly sexed individuals or a reality in wild-type populations.

Our results on Pangasiid SD also raise interesting questions on *Amhr2* structure and evolution. For instance, the N-terminal truncation of all the Pangasiid *Amhr2by* proteins is intriguing because this N-terminal part of the TGF β type II receptors encodes the complete extracellular ligand-binding domain that is known to be crucial for ligand binding specificity (Hart et al., 2021). N-terminal truncations of TGF β receptors acting as sex-determining genes have been already reported for *Amhr2* in yellow perch (Feron et al., 2020) and common seadragon (Qu et al., 2021), and for *Bmpr1b* in the Atlantic herring, *Clupea harengus* (Rafati et al., 2020). In the Atlantic herring, the N-terminal truncated *Bmpr1bby* protein lacks the canonical TGF β receptor extracellular domains, but has maintained its ability to propagate a specific intracellular signal through kinase activity and Smad protein phosphorylation (Rafati et al., 2020). Together, these studies suggest that some TGF β receptors truncated in their N-terminal extracellular ligand-binding domain can still trigger a biological response independent from any ligand activation. The fact that, convergently, many fish MSD genes encoding a TGF β receptor with a similar N-terminal truncation suggests that such a ligand-independent action is probably an important step that could have been selected independently in different lineages to allow an autonomous action of the MSD gene. A second interesting and unexpected result from our study is that the duplication of *amhr2* genes that gave birth to the Pangasiids *amhr2by* gene is potentially ancient and so is likely to still be present in additional catfish species outside the family Pangasidae. This result is well supported by the topologies of our *amhr2* phylogenetic gene trees that place the origin of this duplication at the root of the Siluroidei sub-order, which is dated around 100 million years ago (Kappas et al., 2016). We also found one example of an *amhr2b* that is retained in the *Pimelodus maculatus*

(family Pimelodidae) genome, although we do not know if this gene is also sex-linked in this species. Surprisingly, however, no other *amhr2* duplication has been reported yet in other catfish species. Gains and losses of MSD genes have been already described such as in Esociformes in which some species have completely lost the *amh* duplication (*amhby*) that is an MSD gene in other closely related species from the same family (Pan et al., 2019, 2021). Such complete gene losses can also be expected in catfishes, for instance in the channel catfish that relies on the *bcar1* gene as an MSD gene (Bao et al., 2019), with no remains of an *amhr2* gene duplication. This situation is also probably the case for additional catfish species in which we did not find any *amhr2* duplication in male genome assemblies, such as in the Ictaluridae, *Ameiurus melas*, the Clariidae, *Clarias magur*, and the Auchenipteridae, *Ageneiosus marmoratus*. However, if *amhr2b* is also male-specific as in Pangasiids, the question remains open for the additional catfish species where only female genome assemblies are currently available, such as in the families Sisoridae, Siluridae and Bagridae. A more extensive search for a potential duplication of *amhr2* genes in additional Siluroidei catfishes would be needed to better understand the fate of the *amhr2b* gene and whether it remains an MSD gene like in the Pangasiid family.

Together our results bring multiple lines of evidence supporting the hypothesis that the conserved Pangasiid *amhr2by* is a potential sex determining gene that stemmed from an ancient duplication common to all Siluroidei catfishes. Our results highlight the recurrent use of the TGF β pathway in teleost SD (Q. Pan et al., 2021) and the potential functional innovation through protein truncation. Furthermore, our results showcase the less considered long-term stability of SD genes in teleosts, a group that often receives attention for its dynamic evolution of SD systems.

ACKNOWLEDGEMENTS

This project was supported by funds from the “Agence Nationale de la Recherche” (PhyloSex project, grant No. ANR-13-ISV7-0005; GenoFish project, grant No. ANR-16-CE12-0035) and R35 GM139635 and R01 OD011116 from the National Institutes of Health, USA. The GeT-PlaGe, Gentyane and Montpellier GenomiX (MGX) facilities were supported by France Génomique National infrastructure, funded as part of the “Investissement d’avenir” programme managed by Agence Nationale pour la Recherche (grant No. ANR-10-INBS-09). M.R.R. was supported by the Swiss National Science Foundation (grant 173048). We are grateful to the Genotoul bioinformatics platform Toulouse Midi-Pyrenees (Bioinfo Genotoul) for providing computing and/or storage resources.

CONFLICT OF INTERESTS

All authors declare no competing interests.

AUTHOR CONTRIBUTIONS

Y.G. and J.H.P. designed the project. J.C.A., R.D., M.C., T.T.T.H., R.G., K.S., J.R. and F.L.A. collected the samples, E.J., M.W., C.I., A.C., C.R., O.B., S.V., C.L., C.P., E.B., V.G. and H.A. extracted the gDNA, made the genomic libraries and sequenced them. C.C., C.K., M.Z., M.W., Q.P. and Y.G. processed the genome assemblies and/or analysed the results. T.D., J.M. and J.B. processed and analysed the small RNA sequencing data for miRNA analysis. C.F.B., M.W., Q.P. and M.R.R. performed phylogenetic analyses. C.F.B. and M.R.R. performed the selection analysis. M.W., J.H.P., C.C., C.K., C.R., Q.P. and Y.G. wrote the manuscript with inputs from all other coauthors. J.H.P., C.D., J.B. and Y.G. supervised the project administration and raised funding. All the authors read and approved the final manuscript.

BENEFIT-SHARING STATEMENT

A research collaboration was developed with scientists from the countries providing genetic samples (K.S. in Thailand, G.R. in Indonesia, T.T.T.H. in Vietnam, and J.R. and F.L.A. in Brazil), all collaborators are included as co-authors, the results of research have been shared with the provider communities, and the research addresses a priority concern, in this case the conservation of organisms being studied. More broadly, our group is committed to international scientific partnerships, as well as institutional capacity building.

DATA AVAILABILITY STATEMENT

The Whole Genome Shotgun project of *Pangasianodon hypophthalmus* is available in the Sequence Read Archive (SRA), under BioProject reference [PRJNA547555](https://doi.org/10.1093/bioinformatics/btu638) with 10X genomics and Hi-C Illumina sequencing data are available in SRA under accession nos. [SRX6071341](https://doi.org/10.1093/bioinformatics/btu638) and [SRX6071345](https://doi.org/10.1093/bioinformatics/btu638) and Oxford Nanopore long reads data under SRA accession nos. [SRX6071342](https://doi.org/10.1093/bioinformatics/btu638) to [SRX6071344](https://doi.org/10.1093/bioinformatics/btu638) and [SRX6071346](https://doi.org/10.1093/bioinformatics/btu638) to [SRX6071355](https://doi.org/10.1093/bioinformatics/btu638). *P. hypophthalmus* small RNA sequences are available in SRA under Bioproject [PRJNA256963](https://doi.org/10.1093/bioinformatics/btu638). *Pangasianodon gigas* and *Pangasius djambal* genomes assembled with a *P. hypophthalmus* reference-guided strategy have been submitted to SRA under the respective BioProjects [PRJNA593917](https://doi.org/10.1093/bioinformatics/btu638) and [PRJNA605300](https://doi.org/10.1093/bioinformatics/btu638). All other Pangasiidae genomes assembled with a *P. hypophthalmus* reference-guided strategy without their genome annotations are available in SRA under BioProject [PRJNA795327](https://doi.org/10.1093/bioinformatics/btu638), and their genome assemblies plus their annotations are available in the omics dataverse (Open source research data repository) server with the following DOI (<https://doi.org/10.15454/M3HYAX>). *Pangasius siamensis* has been considered by NCBI curators as a synonym of *P. macronema* and its genome is then recorded in NCBI with *P. macronema* as a Biosample species name, with sample name PaSia (for *Pangasius siamensis*) under accession BioSample no. [SAMN24707637](https://doi.org/10.15454/M3HYAX).

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How to cite this article: Wen, M., Pan, Q., Jouanno, E., Montfort, J., Zahm, M., Cabau, C., Klopp, C., Iampietro, C., Roques, C., Bouchez, O., Castinel, A., Donnadiou, C., Parrinello, H., Poncet, C., Belmonte, E., Gautier, V., Avarre, J.-C., Dugue, R., Gustiano, R., ... Guiguen, Y. (2022). An ancient truncated duplication of the anti-Müllerian hormone receptor type 2 gene is a potential conserved master sex determinant in the Pangasiidae catfish family. *Molecular Ecology Resources*, 22, 2411–2428. <https://doi.org/10.1111/1755-0998.13620>