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A supernumerary "B-sex" chromosome drives male sex determination in the Pachón cavefish, *Astyanax mexicanus*

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44 SUMMARY

45 Sex chromosomes are generally derived from a pair of classical type-A chromosomes, and relatively few alternative models have been proposed up to now^{1,2}. B chromosomes (Bs) are 46 47 supernumerary and dispensable chromosomes with non-Mendelian inheritance found in many plant and animal species^{3,4}, that have often been considered as selfish genetic elements that 48 behave as genome parasites^{5,6}. The observation that in some species Bs can be either restricted 49 or predominant in one sex $^{7-14}$ raised the interesting hypothesis that Bs could play a role in sex 50 determination¹⁵. The characterization of putative B master sex-determining (MSD) genes, 51 52 however, has not yet been provided to support this hypothesis. Here, in Astyanax mexicanus 53 cavefish originating from Pachón cave, we show that Bs are strongly male-predominant. Based 54 on a high-quality genome assembly of a B-carrying male, we characterized the Pachón cavefish B sequence and found that it contains two duplicated loci of the putative MSD gene growth 55 56 differentiation factor 6b (gdf6b). Supporting its role as an MSD gene, we found that the Pachón 57 cavefish gdf6b gene is expressed specifically in differentiating male gonads, and that its 58 knockout induces male-to-female sex reversal in B-carrying males. This demonstrates that 59 gdf6b is necessary for triggering male sex determination in Pachón cavefish. Altogether these 60 results bring multiple and independent lines of evidence supporting the conclusion that the 61 Pachón cavefish B is a "B-sex" chromosome that contains duplicated copies of the gdf6b gene 62 which can promote male sex determination in this species.

63

64 RESULTS AND DISCUSSION

65 Pachón cavefish B chromosomes are male-predominant B chromosomes

66 Supernumerary B chromosomes (Bs) are generally thought to arise from the duplication and assembly of A chromosome sequences^{16–19} and their relationship to sex chromosomes has often 67 been suspected and discussed¹⁵. Some hypotheses state that Bs are derived from sex 68 chromosomes or, alternatively, evolved to become sex chromosomes^{15,20-23}. Because Bs have 69 been described in *A. mexicanus*^{19,24,25}, we performed cytogenetic analyses in 17 males and 11 70 71 females of a laboratory population of Pachón cavefish (Pachón) to investigate whether Pachón 72 Bs could be sex-restricted. We found that Pachón Bs are euchromatic mitotically unstable microchromosomes (Figure 1A-B, Figure S1), that are present in one to three copies in most 73 74 male metaphases (mean number \pm SD of Bs per male metaphase = 1.08 \pm 0.41), contrasting with a barely detectable B occurrence in female metaphases (mean number \pm SD of Bs per 75 76 female metaphase = 0.05 ± 0.08 , Figure 1C and Data S1A). Chromosomal mapping by

77 fluorescence *in situ* hybridization (FISH) using probes generated from microdissected male Bs painted Bs in males and even the rare B in females, supporting that Pachón male-predominant 78 79 Bs and the low occurrence female Bs share a similar gDNA content (Figure 1D). In addition, weaker FISH signals were also detected on different terminal parts of some A chromosomes 80 81 (see small white arrows in Figure 1D), suggesting that Pachón Bs are made up of many duplicated fragments of A chromosomes¹⁶⁻¹⁹, and / or that they share repetitive DNAs with the 82 A chromosomes^{21,24,26}. Female or male sex-restricted or sex-predominant Bs have been 83 described in many fishes, like for instance in some cichlids^{12–14} and characiforms^{8,9,27,28}. In A. 84 *mexicanus*, Bs were recently described as being restricted only in some, but not all, males¹⁹, 85 suggesting population differences in the frequency and sex-linkage of Bs that have also been 86 reported in many species^{4–6,29}. In addition to metaphase spreads (Figure 1E-F), we also detected 87 Bs in pachytene chromosome spreads from testes of two Pachón males (N=52 and N=25). 88 Single Bs were found in most cells (44/52 and 23/25) and always as unpaired chromosomes 89 (Figure 1G-H). Hence, Pachón Bs are present in meiotic germ cells and they do not pair with 90 A chromosomes in line with what has been found in other species with Bs³⁰. The question of 91 92 whether Pachón Bs can pair to each other remains open because we detected no case of multiple 93 Bs in these pachytene chromosome spreads.

94 Characterization of the Pachón cavefish B chromosome sequence

95 Because the publicly available Pachón genome assembly was obtained from a female³¹, we sequenced the genome of a B+ Pachón male to assemble its B sequence. Bs are notoriously 96 97 difficult to assemble^{32,33}, due to their complex mosaic composition of A chromosome fragments¹⁶⁻¹⁹ and their high-repeat content^{17,19}, and most of the B sequence information is 98 from short-read sequencing of purified Bs^{17,18} or B+ versus B-devoid (B-) individuals^{17-19,34}. 99 100 To accurately assemble a high-quality Pachón B sequence, we used a combination of HiFi 101 PacBio and Oxford nanopore long-reads, 10X genomics Illumina short-linked reads, and a Hi-102 C chromosome contact map. The resulting whole-genome assembly (see assembly metrics in 103 Data S1B) contains 25 large scaffolds corresponding to the 25 Pachón A-chromosome pairs^{19,25} and 170 remaining unplaced scaffolds (2.12% of the total assembly size). 104

To identify the Pachón B sequence we used its male-predominant feature and a poolsequencing (pool-seq) approach to contrast whole-genome sequences of a genomic DNA (gDNA) pool of 91 phenotypic males versus a gDNA pool of 81 phenotypic females. By remapping these male and female pool-seq reads on our male Pachón genome assembly, we 109 identified a single 2.97 Mb unplaced scaffold (HiC Scaffold 28) that displays a clear male-110 biased read coverage profile (Figure 2A-A'). The sequence analysis of HiC Scaffold 28 111 revealed that it is made up from a complex mosaic of numerous duplicated fragments of A chromosomes (Figure 2B) including complete but also truncated duplicates of A chromosome 112 113 genes (Table S3). The B also displays a repeat content that is markedly different from A 114 chromosomes (Figure 2C-D, Figures S2A-B, Data S1D). Both its sex-biased profile and its 115 sequence characteristics indicate that HiC Scaffold 28 is the Pachón B. The contribution of 116 many A chromosome regions to the structure of Pachón Bs is in line with the recent findings 117 that A. mexicanus Bs contain a large number of transposable elements¹⁹. The high B proportion of satellite DNA (Figures S2 and Data S1D), was also reported in other species³⁵. Our 118 119 manually curated B gene annotation (Data S1C) identified 63 genes on the Pachón B. Of these, 120 20 show high-quality annotation over their full length, 11 are truncated compared to their 121 conserved homologs in other fish and one is a chimeric gene. Five genes show multiple copies and constitute almost one-third of the B gene content (Data S1C). These results contrast with 122 earlier studies which reported a much higher number of B genes in A. mexicanus¹⁹. These 123 124 annotation differences are likely due to indirect assessment of the B gene content based on short-read sequencing of very few B+ versus B- individuals¹⁹. This comparison clearly 125 126 illustrates the need for better, complete and high-quality B assemblies like we provided here for Pachón cavefish, to better understand the structure and gene content of Bs generally. 127

128 The Pachón cavefish B contains two copies of a putative master sex determining gene

129 Among the B genes with well-supported annotation evidence (Data S1C), we identified two 130 duplicated loci of the A-chromosome-3 growth differentiation factor 6b (gdf6b) gene (located at Chr03:863,919-866,170), that is the teleost ohnolog (teleost whole genome duplication^{36,37} 131 132 paralogous copy) of the *gdf6a* gene (Figure 3A). *Gdf6* genes belong to the TGF-β superfamily within which many master sex-determining (MSD) genes have been found, including TGF-B 133 receptors³⁸⁻⁴⁰ and ligands⁴¹⁻⁴⁴. Of note, *gdf6a* on the Y chromosome (*gdf6aY*) has been 134 135 characterized as the master sex-determining gene in the turquoise killifish, Nothobranchius *furzeri*⁴⁴. The Pachón B *gdf6b* genes (*B-gdf6b* = *B-gdf6b-1* and *B-gdf6b-2*) were thus retained 136 137 as potential candidate B MSD genes. Chromosome FISH hybridization with a gdf6b locus 138 probe revealed a gdf6b hybridization signal on the Pachón Bs, along with a single gdf6b 139 labelling on a single A chromosome pair, both in the male-predominant Bs and the low occurrence female Bs (Figure 1E-F, and inset in Figure 1E). 140

141 The two *B-gdf6b* loci are 99.6 % identical in the 21.6 kb region shared by the two genes, and 100% identical in their coding sequences (CDS), with *B-gdf6b-2* being derived from an internal 142 143 B duplication of the *B-gdf6b-1* locus (Figure S2C). Such internal B duplications / insertions 144 indicate that the origin of the B structure can be more complex than initially thought. 145 Comparison of these *B-gdf6b* loci with the overlapping sequence of their A chromosome 146 counterpart (A-gdf6b) revealed numerous differences in their proximal promoters and also their 147 intron that contains two *B-gdf6b* specific insertions (Figure S2D). However, differences within 148 the gdf6b CDS were limited to a T-to-C (A-gdf6b-to-B-gdf6b) synonymous substitution 149 (c.591T>C) and two nonsynonymous substitutions, i.e., a T-to-G (A-gdf6b-to-B-gdf6b) 150 transversion (c.180T>G) in exon 1 that switches a A-Gdf6b lysine into a B-Gdf6b asparagine 151 (p.Lys60Asn) and a G-to-A (A-gdf6b-to-B-gdf6b) transition (c.679G>A) in exon 2 that 152 switches the A-Gdf6b serine into a B-Gdf6b glycine (p.Ser227Gly) (Figure S2E). The 153 Lys60Asn does not impact a conserved amino-acid position of Gdf6b proteins, in contrast to 154 the Ser227Gly that impacts a glycine of the TGF- β /BMP propeptide domain that is conserved 155 in Pachón B-Gdf6b and in all vertebrate Gdf6 proteins, but not in Pachón A-Gdf6b (Figure 156 3B). It is interesting to note that this Ser227Gly modification suggests that the A-gdf6b acquired 157 this mutation after the *B-gdf6b* copy was duplicated on the B chromosome. This non-conserved 158 Ser227 engages in a central hydrogen bond network at a looptip region in the TGF-β/BMP propeptide domain (Figure 3C) that could affect the stability of this pro-domain. The 159 160 Gly227Ser exchange in Pachón A-Gdf6b leads to the gain of several hydrogen bonds that are 161 due to the side chain hydroxyl group of the Ser227 residue (Figure 3D-E). The gain of hydrogen 162 bonds can confer a higher folding stability of the proprotein complex and thereby might affect activation of Gdf6b as this requires release of the mature C-terminal domain from the 163 164 proprotein complex. The mature C-terminal growth factor domain, however, is likely to be 165 unaffected by this mutation. Whether these conformation differences between the A-Gdf6b and 166 B-Gdf6b proproteins could provide a potential functional explanation for a sex-determining 167 role of the male-predominant Pachón B-Gdf6b remains to be explored, but point mutations in 168 other MSD genes like *amhr2Y* in *Takifugu rubripes* and *amhY* in *Oreochromis niloticus*, have been described to be directly responsible for male sex determination^{39,41}. 169

Based on these *B-gdf6b* and *A-gdf6b* loci differences, we developed several B-specific PCR
genotyping tests on fin clips (Figure S3A). In our Pachón laboratory population, we found a
complete (100%) association between B-specific amplifications and the male phenotype in 723
males, with all the 787 tested females being negative (p-value of association with sex < 2.2e-

174 16). We also found the same complete association in wild-caught Pachón individuals (20 males and 20 females, recognized by external sex-specific traits without sacrifice; p-value of 175 176 association with sex = 1.87e-09) (Data S1F) showing that this male-predominant B is not the 177 result of a domestication effect. These results strengthen our cytological observations of a male-178 predominant B. The absence of B-specific amplifications in females despite the cytogenetic 179 detection of rare Bs in females is probably the result of a PCR sensitivity issue as increasing 180 the number of PCR cycles allows the detection of a faint PCR fragment in Pachón females 181 (Figure S3B).

182 Evidence supporting *gdf6b* as a potential master sex determining gene in Pachón cavefish

183 Sex-specific expression patterns during the sex differentiation period and alteration of gonadal 184 development upon knockout are key arguments for the evaluation of a candidate MSD gene. 185 Due to the high sequence identity of the *B-gdf6b* and *A-gdf6b* cDNAs, we were not able to specifically quantify the B-gdf6b expression. Quantification of the expression of gdf6b (B-186 187 gdf6b and/or A-gdf6b) showed that it has both a predominant expression in developing and 188 adult male gonads (as well as in male brain, intestine, kidney and swim-bladder; Figure S4A) 189 and a strong sexually dimorphic expression during early differentiation of B+ individuals (Figure 4A). Using *in situ* hybridization, the expression of *gdf6b* during the early differentiating 190 191 period was restricted to gonads of B+ individuals (at 15, 21, 30, and 60 days post-fertilization), 192 with no strict colocalization with the gonadal soma-derived factor gene (gsdf) (Figure 4B-C 193 and Figure S4B). Gsdf is a well-known gonad-restricted, somatic supporting cell lineage 194 marker⁴⁵⁻⁴⁷ that has also been described as one of the earliest Pachón gonadal sex differentiation marker genes⁴⁸. This result demonstrates that gdf6b mRNA (B-gdf6b and/or A-195 196 gdf6b) has an expression profile compatible with a male (B+) MSD function, being expressed 197 in the right place, i.e., only in the differentiating testis, and at the right time, i.e., during early 198 testicular differentiation.

To bring additional and functional evidence that *gdf6b* could act as an MSD gene, we generated *gdf6b* knockouts in Pachón cavefish using the genome-editing CRISPR-Cas9 system with two guide RNAs in order to remove a large part of the *gdf6b* CDS (Figure 4D). This large deletion (~470 bp) includes most of the TGF- β propeptide region and the beginning of the TGF- β like domain resulting in a truncated, likely non-functional, Gdf6b protein (Figure S2F). Among 200 first-generation microinjected individuals (which are mosaic for the genome-edited loci), eighteen B+ individuals (i.e., genotypic males) had a ~470 bp deletion in their *A*- and/or *B*-

206 gdf6b exon 2 (Figure S3C), and they were all sex-reversed into phenotypic females (Figure 207 4G). In contrast, all B+ males and B- females without the *gdf6b* deletion developed normal 208 testes (B+) or ovaries (B-) (Figure 4E-F). This result shows that gdf6b is necessary to trigger 209 Pachón testicular development in B+ individuals and brings further functional evidence that 210 gdf6b could be used as a male MSD gene in Pachón cavefish. However, because of the high 211 similarity of the *B-gdf6b* loci with the *A-gdf6b* locus, we have not been able to specifically 212 knockout the Pachón *B-gdf6b*. Further studies will bring more functional proof supporting the 213 role of the *B-gdf6b* genes in sex determination, including the specific *B-gdf6b* knockout in B+ 214 individuals and the overexpression of gdf6b by transgenesis in B- fish.

215 Conclusions

216 Altogether our results bring new pieces of evidence to support a role of some B chromosomes 217 in sex determination. The potential implication of Bs in sex determination had been suspected in many species including some fishes^{14,15}. Up to now, however, the characterization of 218 219 potential B MSD genes along with strong functional evidence has only been provided for the 220 bacterial-derived haploidizer gene in the jewel wasp, Nasonia vitripennis B chromosome (named PSR for paternal sex ratio chromosome)⁴⁹, although maleness in this haplo-diploid 221 organism is determined through elimination of the paternal A chromosome set⁵⁰. Here, 222 223 combining a variety of approaches we discovered that Pachón cavefish of the species A. 224 mexicanus carry male-predominant Bs that contain two copies of the gdf6b gene, which itself 225 behaves as an excellent MSD candidate gene. This indeed brings the interesting hypothesis that 226 these Pachón Bs could be considered as "B-sex" chromosomes. However, the question remains 227 open whether Pachón Bs are predominant in males because they are eliminated from female 228 tissues or whether they are by themselves necessary and sufficient to trigger maleness. Despite 229 being male-predominant, Pachón Bs are also found in some females albeit only in very few 230 metaphases (21.6 times less abundant than in males on average) and most often as a single B 231 copy. B frequencies have been described as being highly variable between species, sexes, individuals of the same population, and even in different cells of a single individual^{4–6,29}. This 232 233 variation is assumed to result from meiotic and/or mitotic instability of Bs that can be present only in some organs and absent from others^{4,29,51-53}. In the plant Aegilops speltoides, a 234 mechanism of programmed elimination of Bs occurs specifically in the roots⁵⁴. It results from 235 236 a B chromatid nondisjunction during mitosis, leading to the micronucleation of Bs and their subsequent degradation at early stages of the proto-root embryonic tissue differentiation⁵⁴. 237 238 Such a mechanism would potentially explain a specific B elimination in Pachón female organs,

as it has been hypothesized in another *Astyanax* species with male-restricted Bs¹¹. Further studies are now needed to better understand this sex-specific B drive mechanism, and if it reflects a cause or a consequence of sex determination in Pachón cavefish. Our results also lay a high-quality genome-based foundation in an important emerging fish model for studying the genomic evolution of Bs, including the micro- and macro-evolution of this B chromosome in

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line with the evolution of sex chromosomes.

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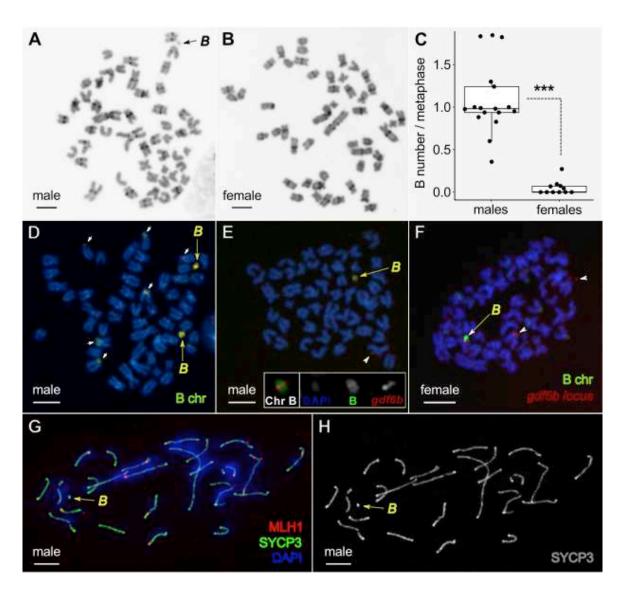
267 AUTHOR CONTRIBUTIONS

Conceptualization, S.R., A.H. and Y.G.; Methodology, B.I., S.R., A.H., A.S., and Y.G.; Formal
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 Writing –Original Draft, B.I., S.R., and Y.G.; Writing –Review & Editing, A.S., J.H.P.,
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275 DECLARATION OF INTERESTS

- 276 The authors declare no competing interests
- 277
- 278 FIGURE
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281 Figure 1: Karyological characterization of male-predominant supernumerary B chromosomes (Bs) in Pachón cave Astyanax mexicanus. A-B: Representative C-banding 282 283 patterns of single B+ male (A) and B- female (B) from Pachón cave. The Bs (black arrow) lack C-bands, suggesting that Pachón cavefish Bs are largely euchromatic. See also Figure S1. C: 284 285 Boxplots of the average number of Bs per metaphase in Pachón cavefish males and females. 286 Horizontal lines indicate the median, the box indicates the interquartile range (IQR), and the 287 whiskers the range of values that are within 1.5 x IQR. Statistical significance between males and females were tested with the Wilcoxon Rank Test (*** = P < 0.001). See also Data S1A. 288 289 D: Fluorescence in situ hybridization (FISH) of male Pachón cave mitotic metaphase labeled 290 with combined microdissected B probes. Yellow arrows point to the strong labelling of Bs and 291 the small white arrows to the lighter labelling of some different parts of A chromosomes. E-F: 292 FISH co-labelling of a 1B male (E) and a 1B female (F) metaphase with microdissected B 293 (green) and gdf6b-specific (red) probes. Bs are indicated by yellow arrows and the white 294 arrowheads point to pairs of A chromosome sister chromatids labelled by the *gdf6b* probe. Only 295 one A-chromosome gdf6b signal was detected in panel E. The two gdf6b signals (see inset in 296 E) that were often visible on male metaphases, cannot be interpreted as the two different B-297 gdf6b loci due to their genomic proximity (see Table S3) and the FISH resolution. G-H: 298 Synaptonemal complex (SC) analysis showing that Pachón cave Bs (yellow arrow) do not pair 299 with the 25 fully synapsed standard bivalents of A chromosomes. SCs were visualized by anti-300 SYCP3 antibody (green), the recombination sites were identified by anti-MLH1 antibody (red) 301 and chromosomes were counterstained by DAPI (blue). G: Merged image. H: SYCP3 302 visualization only. Scale bars = $5 \mu m$.

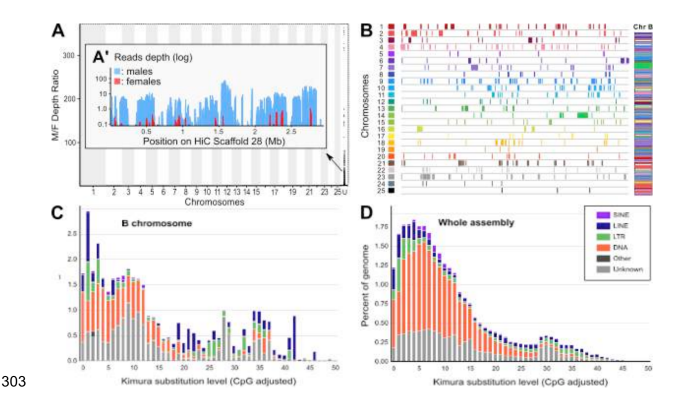
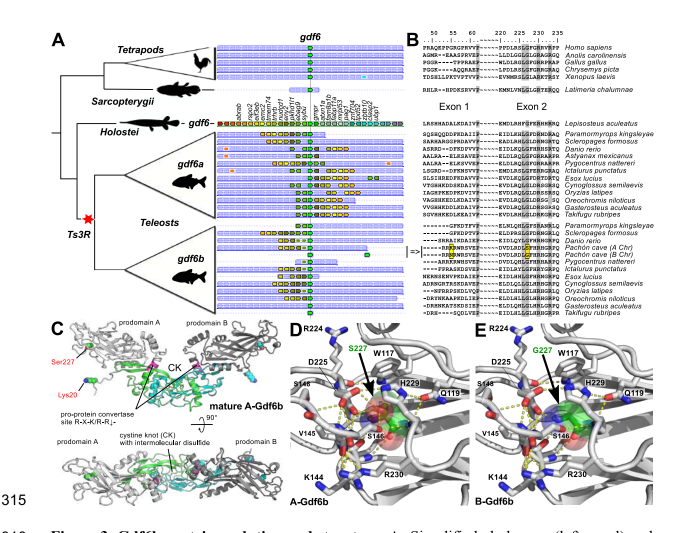
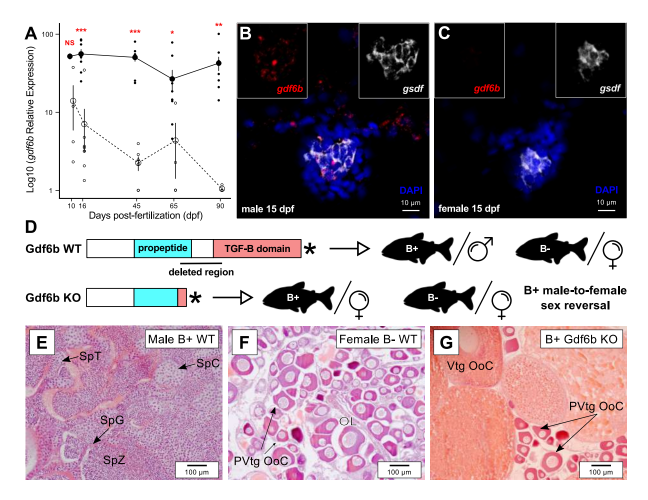


Figure 2: Genomic characterization of Pachón cavefish B chromosome (B). A: Read depth 304 ratio of male and female Pachón genomic pools showing a strong coverage bias in a single 305 306 scaffold, Hi scaffold 28 (enlarged in A' inset showing male and female read coverage). B: 307 Karyoplots of the A chromosome regions duplicated on the Pachón cavefish B (Chr B) showing 308 that the Pachón B is made from a complex mosaic of duplicated A chromosomal fragments. C-309 D: Comparison of the repeat landscapes of the Pachón B (C) and whole-genome including the 310 B (D), showing that the Pachón B has a very different repeat element (color code provided in 311 inset of panel C) content compared to A chromosomes. Short interspersed repeated sequences 312 (SINEs), long interspersed nuclear elements (LINEs), long terminal repeats (LTRs), DNA 313 repeat elements (DNAs), and terminal inverted repeat sequences (TIRs). See also Figure S2A-314 B and Data S1D for additional details.



316 Figure 3: Gdf6b protein evolution and structure. A: Simplified phylogeny (left panel) and synteny (middle panel) relationships of the gdf6a and gdf6b genes, with the additional B-gdf6b 317 Pachón cavefish duplication, showing that they are duplicated paralogues stemming from the 318 teleost whole genome duplication (Ts3R). Species names are given on the right side of panel 319 B. B: Corresponding multiple alignments of Gdf6 protein-coding sequences around the A-320 321 Gdf6b lysine to B-Gdf6b asparagine switch in Exon 1 (p.Lys60Asn) and the A-Gdf6b serine 322 to B-Gdf6b glycine switch in Exon 2 (p.Ser227Gly). C: Ribbon plot homology model of A-Gdf6b proprotein dimer (bottom panel view is rotated by 90° around the x-axis). The 323 prodomains are shown in light and dark grey, the furin processing site (R274-R-K-R-R278) is 324 325 indicated in magenta, the activity-containing mature C-terminal domain is shown in green and 326 cyan. The two residues differing between A-Gdf6b and B-Gdf6b are presented as spheres with their carbon atoms colored in green and cyan. D-E: Comparative magnifications of the 327 328 structure of A-Gdf6b (D) and B-Gdf6b (E) around the p.Ser227Gly switch. Amino acid 329 residues interacting with Ser227 or Gly227 are shown as sticks, and hydrogen bonds as yellow 330 stippled lines. As shown, the side chain hydroxyl group of Ser227 (shown with carbon atoms colored in green and transparent spheres highlighting the van der Waals spheres of the atoms) 331

engages in several hydrogen bonds with surrounding residues, e.g. Ser146, Asp225 thereby
stabilizing the tertiary and secondary structure in this region. Upon exchange of Ser227 with a
glycine as in B-Gdf6b these hydrogen bonds are lost thereby potentially destabilizing this
region in the prodomain.



336

Figure 4: Gene expression and functional evidence supporting a role of gdf6b as a 337 338 potential master sex-determining gene in Pachón cavefish. A: Expression profiles of gdf6b 339 in male and female trunks during early development from 10 to 90 days post-fertilization (dpf, males: solid line; females: dashed line) showing a significant over-expression in males 340 compared to females starting from 16 dpf. Results are presented as log_{10} mean \pm standard 341 342 errors; black and white dots represent the individual values of relative expression in males and 343 females, respectively. Statistical significance between males and females was tested with the 344 Wilcoxon Rank Sum Test (Wilcoxon-Mann-Whitney Test) and only significant differences are shown (*** = P < 0.01; ** = P < 0.01; * = P < 0.05). **B-C:** Gonadal expression of *gdf6b* (in 345 346 red) and the Sertoli and granulosa supporting cell marker gsdf (in white) in male (**B**) and female (C) differentiating gonads at 15 dpf showing that *gdf6b* is specifically expressed in male gonads 347 with no strict colocalization with gsdf in male. See Figure S6B for additional stages of 348

development. Nuclei were stained with DAPI (in blue). Scale bar = $10 \mu m$. **D:** Schematic 349 representation of the wild-type (WT) and knock-out (KO) Gdf6b proteins and the resulting 350 351 phenotypes of B+ males and B- females. E-F: Representative gonadal histology of WT males 352 (E), WT females (F), and Gdf6b KO B+ males showing that Gdf6b KO induces male-to-female 353 sex reversal (G) with ovaries containing vitellogenic (Vtg Ooc) and previtellogenic oocytes 354 (PVtg Ooc), like WT ovaries (F), contrasting with the testis in WT males (E). Ol: Ovarian lamellae. SpC: Spermatocytes; SpG; Spermatogonia; SpT: Spermatids; SpZ: Spermatozoa. 355 356 Scale bar = $100 \,\mu\text{m}$. See also Figure S4.

357

358 STAR METHODS

359

360 **RESOURCE AVAILABILITY**

361 Lead contact

Further information and requests for resources and reagents should be directed to and will befulfilled by the lead contact, Yann Guiguen (yann.guiguen@inrae.fr).

364

365 Materials availability

To request Pachón fish lines or constructs created in this study, please contact the lead contact.

368 Data and code availability

Raw sequences and the whole genome assembly of Pachón cavefish have been deposited in the National Center for Biotechnology Information DDBJ/ENA/GenBank databases under the BioProject PRJNA734455. This accession number is listed in the key resources table. This study did not generate new unique code. Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

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376 EXPERIMENTAL MODEL AND SUBJECT DETAILS

377 Cavefish breeding and sampling

Laboratory stocks of *A. mexicanus* Pachón cavefish were obtained in 2004 from the Jeffery laboratory at the University of Maryland, College Park, MD. Fish were raised as previously described. Fertilized eggs were provided by CNRS cavefish experimental facilities (Gif sur Yvette, France) and maintained at 24°C until the hatching stage occurring around 24 ± 2 hours 382 post-fertilization (hpf)⁵⁵. Subsequently, larvae were transferred and raised in the Fish Physiology and Genomics laboratory experimental facilities (LPGP, INRAE, Rennes, France) 383 384 under standard photoperiod (12 h light / 12 h dark) and at two different temperatures: 21 ± 1 °C and 28 ± 1 °C. Animals were fed twice a day, firstly with live artemia (Ocean Nutrition) 385 386 until 15 days post-fertilization (dpf), then with a commercial diet (BioMar) until adult stage. 387 For animal dissections and organ sampling, fish were euthanized with a lethal dose of tricaine 388 methanesulfonate (MS 222, 400 mg/l), supplemented by 150 mg/l of sodium bicarbonate. 389 Phenotypic sex of individuals was determined at 4 months and more, either by macroscopical 390 examination of the gonads when they were enough differentiated, or by histology when gonads were not totally differentiated⁴⁸. Caudal fin clips were collected from all individuals and stored 391 in ethanol 90% at 4°C before genomic DNA (gDNA) extraction. For the chromosome contact 392 393 map (Hi-C), 80 µl of blood was sampled from three males using a syringe rinsed with EDTA 394 2%. The fresh blood was slowly frozen in a Freezing Container (Mr. Frosty, Nalgene®) after 395 addition of 15% of dimethyl sulfoxide (DMSO). Karyotypic analyses were carried out in 17 396 males and 11 females of Pachón cave Astyanax mexicanus. Fin samples (a narrow strip of the 397 tail fin) were taken from the live specimens anesthetized by MS-222 (Merck KGaA, Darmstadt, 398 Germany), while for direct preparations (chromosomes from kidneys and gonads), fishes were 399 euthanized first using 2-phenoxyethanol (Sigma-Aldrich, St. Louis, MO, USA).

400 All animal protocols were carried out in strict accordance with the French and European 401 legislations (French decree 2013-118 and directive 2010-63-UE) applied for ethical use and 402 care of laboratory animals used for scientific purposes. Sylvie Retaux and CNRS institutional 403 authorizations for maintaining and handling A. mexicanus in experimental procedures were 91-404 116 and 91272105, respectively. For karyotypic analysis, all handling of fish individuals 405 followed European standards in agreement with §17 of the Act No. 246/1992 to prevent fish 406 suffering. The procedures involving fish were supervised by the Institutional Animal Care and 407 Use Committee of the Institute of Animal Physiology and Genetics CAS, v.v.i., the 408 supervisor's permit number CZ 02361 certified and issued by the Ministry of Agriculture of 409 the Czech Republic. Several sampling campaigns were carried out in the field in Mexico 410 between 2013 and 2019, resulting in a collection of tail fin clips from wild-caught individuals 411 from the Pachón cave. In the field, the phenotypic sex of animals was determined by checking 412 the presence or absence of denticles on the anal fins as described previously⁵⁶, a small fin clip 413 was gently taken and fish were rapidly returned to their natural pond. In addition, pictures of 414 each individual sampled were taken to confirm the phenotypic sex, back in the laboratory,

based on the morphological criteria described previously⁵⁶. The permits for field sampling
(02241/13, 02438/16, 05389/17 and 1893/19) were delivered by the Mexican authorities
(Mexican Secretaría del Medio 16 Ambiente y Recursos Naturales) to Sylvie Rétaux and
Patricia Ornelas-Garcia (UNAM, Mexico).

419

420 METHOD DETAILS

421 DNA extraction

422 For fish genotyping, gDNA was extracted from fin clips stored in 90% ethanol, after lysis in 423 5% chelex⁵⁷ and 10 mg Proteinase K at 55 °C for 2 h, followed by 10 min at 99 °C. Following 424 extraction, samples were centrifuged and the supernatant containing the gDNA was transferred 425 in clean tubes and stored at -20 °C. For pool-sequencing and TaqMan assay, gDNA was 426 extracted using NucleoSpin Kits for Tissue (Macherey-Nagel, Duren, Germany) according to 427 the supplier's recommendations. For long-read male genome sequencing, high molecular 428 weight (HMW) gDNA was extracted from a mature testis grounded in liquid nitrogen and lysed 429 in TNES-Urea buffer (TNES-Urea: 4M urea; 10 mM Tris-HCl, pH 7.5; 120 mM NaCl; 10 mM 430 EDTA; 5% SDS) for two weeks at room temperature. For HMW gDNA extraction the TNES-431 Urea solution was supplemented with Proteinase K at a final concentration of 150 µg/ml and 432 incubated at 37 °C overnight. HMW gDNA was extracted with a modified phenol-chloroform protocol as previously described⁴³. The gDNA concentrations for both pool-seq and genome 433 434 sequencing were quantified with Qubit3 fluorometer (Invitrogen, Carlsbad, CA) and HMW 435 gDNA quality and purity were assessed using spectrophotometry, fluorometry, and capillary 436 electrophoresis.

437 Primers and probe design

All primers used in this study including PCR genotyping, qPCR gene expression and cDNA
cloning were designed using Primer3web software⁵⁸ version 4.1.0 and are listed in Data S1E.

440 B polymerase chain reaction (PCR) genotyping

Genetic sex of Pachón cave individuals was determined by PCR tests using the fact that maleshave a B-predominant chromosome with three different sets of primers (see Table S6 for primer

- sequences) based on differences between the *A-gdf6b* and *B-gdf6b* loci. Three sets of primers
- (P) were designed (see Figure S3A) with two primer sets designed to amplify specifically the

445 two *B-gdf6b* copies based either on a single base variation between the A/B gdf6b CDS at position 679 bp (P1-P2), or based on primers located on both sides of the A/B breakpoints 446 447 downstream of the *B-gdf6b* gene (P3-P4). The third set of primers (P5-P6) was designed on 448 gaps/indels variations between A-gdf6b and the two B-gdf6b genes in the proximal promoter 449 of gdf6b genes. Another set of primers (P9-P10) was designed as a PCR positive control with 450 primers located on both sides of the A/B breakpoints downstream of the A-gdf6b gene. Primer 451 sets P1-P2 and P3-P4 produce a single PCR fragment only in males (B+), primer set P5-P6 452 amplifies two bands in males (all B+) and only a single band in females (all B-), and primer set 453 P9-P10 amplifies a single band in all individuals. For PCR reactions with the P1-P2 primer set, 454 HiDi Taq DNA polymerase (myPOLS Biotec) was used for detecting a single nucleotide 455 variation. PCRs were performed in a total volume of 12.5 µl containing 0.2 µM of each primer, 456 a final concentration of 20 ng/µl gDNA, 200 µM of dNTPs mix, 1X of HiDi buffer (10X), and 457 2.5 U per reaction of HiDi DNA polymerase. Cycling conditions were as follows: 95 °C for 2 min, then 35 cycles of (95 °C for 15 seconds (secs) + 60 °C for 10 sec + 72 °C for 30 sec), and 458 459 72 °C for 5 min. For PCR reactions with the P3-P4 and P9-P10 primer sets, PCR reactions were 460 performed in a final volume of 50 µl containing 0.5 µM of each primer, a final concentration 461 of 20 ng/µl gDNA, 10 µM dNTPs mix, 1X of 10X AccuPrime[™] buffer, and 0.5 µl per reaction 462 of AccuPrime[™] Taq DNA polymerase. Cycling conditions were as follows: 94 °C for 2 min, then 35 cycles of (94 °C for 30 sec + 58 °C for 30 sec + 68 °C for 1 min and 30 sec). For PCR 463 464 reactions with the P5-P6 primer set, PCR reactions were performed in a final volume of 25 µl 465 containing 0.5 µM of each primer, a final concentration of 20 ng/µl gDNA, 10 µM dNTPs 466 mixture, 1X of Jumpstart[™] buffer (10X), and 0.5 µl per reaction of Jumpstart[™] Taq DNA 467 polymerase. Cycling conditions were as follows: 95 °C for 2 min, then 35 cycles of (95 °C for $1 \min + 60 \degree C$ for $30 \sec + 72 \degree C$ for $1 \min$), and $72 \degree C$ for $5 \min$. 468

469 10× Genomics sequencing

470 10X Chromium Library was prepared according to 10X Genomics protocols using the Genome 471 Reagent Kits v2. Optimal performance has been characterized on input gDNA with a mean 472 length greater than 50 kb (~144Kb). GEM reactions were performed on 0.625 ng of genomic 473 DNA, and DNA molecules were partitioned and amplified into droplets to introduce 16-bp 474 partition barcodes. GEM reactions were thermally cycled (30 °C for 3 h and 65 °C for 10 min; 475 held at 4 °C) and after amplification, the droplets were fractured. P5 and P7 primers, read 2, 476 and sample index were added during library construction. The library was amplified using 10 477 cycles of PCR and the DNA was subsequently size selected to 450 bp by performing a double purification on AMPure Xp beads. Library quality was assessed using a Fragment Analyzer
and quantified by qPCR using the Kapa Library Quantification Kit. Sequencing has been
performed on an Illumina HiSeq3000 using a paired-end read length of 2x150 bp with the
Illumina HiSeq3000 sequencing kits.

482 Oxford nanopore genome sequencing

483 High molecular weight gDNA purification steps were performed using AMPure XP beads 484 (Beckman Coulter). Library preparation and sequencing were performed using Oxford 485 Nanopore (Oxford Nanopore Technologies) Ligation Sequencing Kit SQK-LSK109 according 486 to manufacturer's instructions "1D gDNA selecting for long reads (SQK-LSK109)". Five µg of DNA was purified then sheared at 20 kb using the megaruptor1 system (diagenode). A one-487 488 step DNA damage repair + END-repair + dA tail of double-stranded DNA fragments was 489 performed on 2 µg of sample. Then adapters were ligated to the library. Library was loaded 490 onto 1 R9.4.1 flowcell and sequenced on a PromethION instrument at 0.02 pM within 72 h.

491 PacBio HiFi genome sequencing

492 Library preparation and sequencing were performed according to the manufacturer's 493 instructions "Procedure & Checklist Preparing HiFi SMRTbell Libraries using SMRTbell 494 Express Template Prep Kit 2.0". Fifteen µg of DNA were purified and then sheared at 15 kb 495 using the Megaruptor3 system (Diagenode). Using SMRTbell Express Template prep kit 2.0, 496 a Single strand overhangs removal and then a DNA and END damage repair steps were 497 performed on 10 µg of the sample. Then blunt hairpin adapters were ligated to the library. The 498 library was treated with an exonuclease cocktail to digest unligated DNA fragments. A size 499 selection step using a 12 kb cutoff was performed on the BluePippin Size Selection system 500 (Sage Science) with "0.75% DF Marker S1 3-10 kb Improved Recovery" protocol. Using 501 Binding kit 2.0 kit and sequencing kit 2.0, the primer V2 annealed and polymerase 2.0 bounded 502 library was sequenced by diffusion loading onto 2 SMRTcells on Sequel2 instrument at 50 pM 503 with a 2 h pre-extension and a 30 h movie.

504 *Hi-C sequencing*

Hi-C data was generated using the Arima-HiC kit (Ref. 510008), according to the
manufacturer's protocols using 10 µl of blood as starting material, the Truseq DNA PCR-Free
kit and Truseq DNA UD Indexes (Illumina, ref. 20015962, ref. 20020590), and the KAPA
library Amplification kit (Roche, ref. KK2620). Hi-C library was sequenced in paired-end
2x150 bp mode on Novaseq6000 (Illumina), using half a lane of a SP flow cell (ref. 20027464).

- 510 Image analyses and base calling were performed using the Illumina NovaSeq Control Software
- and Real-Time Analysis component (v3.4.4). Demultiplexing was performed using Illumina's
- 512 conversion software (bcl2fastq v2.20). The quality of the raw data and potential contaminants
- 513 was assessed using FastQC $(v0.14.0)^{59}$ from the Babraham Institute and the Illumina software
- 514 SAV (Sequencing Analysis Viewer).

515 *Genome assembly*

Pacbio HiFi reads were assembled with hifiasm⁶⁰ version 0.9 using standard parameters. The 516 517 genome assembly fasta file was extracted from the principal gfa assembly graph file using an 518 awk command line. This assembly was then scaffolded using Hi-C and 10X as a source of 519 linking information. 10X reads were aligned using Long Ranger v2.1.1 (10x Genomics). Hi-C reads were aligned to the draft genome using Juicer⁶¹ with default parameters. A candidate 520 521 assembly was then generated with 3D de novo assembly (3D-DNA) pipeline⁶² with the -r 0 and --polisher-input-size 100000 parameters. Finally, the candidate assembly was manually 522 reviewed using the Juicebox assembly tools⁶³. Due to the specific structure of the Pachón cave 523 B chromosome, both Hi-C and 10X signals show some uncertainties in the order and 524 525 orientation of the contigs. To improve the quality of the B chromosome assembly, ONT reads were then aligned to the final version of the genome using minimap 2^{64} v2.11 with -x map-ont 526 527 parameter. Both reads spanning contig junctions and reads showing supplementary alignments 528 linking contigs belonging to the B chromosome were analyzed to resolve these ambiguities.

529 *Genome annotation*

530 The cavefish whole genome assembly was annotated using a pipeline adapted from previous 531 studies^{65,66}. In brief, RepeatModeler, RepeatProteinMask, and RepeatMasker (open-4.0.7, 532 http://www.repeatmasker.org/) were first used to scan the genome and mask out repeats. Then 533 protein-coding genes were annotated by collecting gene evidence from homology alignment, 534 RNA-seq mapping, and *ab initio* prediction. For homology alignment, 464,144 protein sequences collected from NCBI were aligned to the assembly using Genewise⁶⁷ and Exonerate 535 536 respectively. RNA-seq data were mapped to the assembly in two independent parallel steps. First Hisat⁶⁸ was used to align RNA-seq reads and then StringTie⁶⁹ was used to predict the gene 537 models; in the other step reads were first assembled into transcript sequences using Trinity⁷⁰ 538 and then PASA⁷¹ was used to map the transcripts to the assembly and model the gene structures. 539 For ab initio prediction and final integrating, Augutus⁷² was first trained using the high-quality 540 541 gene models and then ran in a hint-guide model.

543 **B** chromosome annotation

544 First, repeats were identified and masked from the B chromosome using RepeatModeler, 545 RepeatProteinMask, and RepeatMasker (open-4.0.7, http://www.repeatmasker.org/). To 546 annotate protein-coding genes, we collected all protein sequences of A. mexicanus annotated 547 by NCBI (Genome ID: 13073) and Ensembl (release-104), and aligned them onto the repeatmasked B sequence using GeneWise^{67,73} and Exonerate respectively. For each query, the best 548 549 hit was kept. To determine the best gene model when multiple ones compete for a splice site, 550 we introduced RNA-seq data to evaluate the quality of these homology gene models. Hence RNA-seq data of A. mexicanus from the previous study⁷⁴ were aligned to B using HISAT⁶⁸ and 551 552 parsed using StringTie⁶⁹. RNA-score of each homology gene model was then calculated as the 553 match-extend of splice sites to that of StringTie prediction. When multiple homology gene 554 models compete for a splice site, those with lower RNA-score were discarded. In cases when 555 some genes failed to be identified using homology alignment, we also implemented an *ab initio* 556 gene prediction using Augustus⁷², where all the homology and transcriptome evidence were 557 used as hints. The predicted results were included into the final gene set if 1) the splice sites 558 are not occupied and 2) the splice sites match 100% to that of StringTie predictions (RNA-559 score =100). To further evaluate the quality of the final gene set, we blasted their protein 560 sequences to **SWISSPROT** (https://www.uniprot.org/) and NR 561 (https://www.ncbi.nlm.nih.gov/), and took the alignment to the best hit to check how much of 562 the query and subject was aligned, respectively (query coverage & subject coverage). Genes 563 with query and subject coverage both >90% were considered as being of good quality.

564 To characterize the A chromosome content of the B chromosome, sequences of the B chromosome assembly were aligned to the sequences of the 25 A chromosomes with 565 566 minimap 2^{64} (v2.11) and the best match of each contig fragment was retained. Overlapping 567 matches were manually filtered considering match lengths and similarities (cigarline and edit distance) in order to build the best non-overlapping matching list. Karyoplots were then plotted 568 using the R package karyoploteR⁷⁵ (https://bernatgel.github.io/karyoploter tutorial/). The 569 median, minimum, and maximum sizes of the 628 B best matches on A chromosomes were 570 571 respectively 1,087 bp, 44 bp and 41,908 bp.

572 Male and female Pool-sequencing

573 DNA was collected from 91 phenotypic Pachón cave males and 81 phenotypic Pachón cave 574 females and was pooled as male and female pools separately. Before pooling, the DNA 575 concentration was normalized in order to obtain an equal amount of each individual genome in

576 the final pool. Pool-sequencing libraries were prepared using the Illumina TruSeq Nano DNA HT Library Prep Kit (Illumina, San Diego, CA) according to the manufacturer's protocol. After 577 578 the fragmentation of each gDNA pool (200 ng/pool) by sonication using an M220 Focused-579 ultrasonicator (COVARIS), the size selection was performed using SPB beads retaining 580 fragments of 550 bp. Following the 3' ends of blunt fragments mono-adenylation and the 581 ligation to specific paired-end adaptors, the amplification of the construction was performed 582 using Illumina-specific primers. Library quality was verified with a Fragment Analyzer 583 (Advanced Analytical Technologies) and then quantified by qPCR using the Kapa Library 584 Quantification Kit (Roche Diagnostics Corp, Indianapolis, IN). The enriched male and female 585 pool libraries were then sequenced using a paired-end multiplexed sequencing mode on a 586 NovaSeq S4 lane (Illumina, San Diego, CA), combining the two pools on the same lane and 587 producing 2×150 bp with Illumina NovaSeq Reagent Kits according to the manufacturer's 588 instructions. Sequencing produced 288 million paired reads and 267 million paired reads for 589 the male and female pool libraries, respectively.

590 *Pool-sequencing analysis*

Characterization of genomic regions enriched for sex-biased signals between Pachón cave 591 592 males and females, consisting of coverage and Single Nucleotide Variations (SNVs) was performed as described previously^{38,43}. Pachón cave *A. mexicanus* paired-end reads from male 593 594 and female pool-seq pools were mapped onto our own Pachón cave genome assembly using 595 BWA mem version 0.7.17⁷⁶. The resulting BAM files were sorted and the duplicate reads due 596 to PCR amplification during library preparation were removed using Picard tools version 597 2.18.2 (http://broadinstitute.github.io/picard) with default parameters. Then, for each pool and 598 each genomic position, a file containing the nucleotide composition was generated using samtools mpileup⁷⁷ version 1.8, and popoolation2⁷⁸ mpileup2sync version 1201. This file was 599 600 then analyzed with custom software (PSASS version 2.0.0; doi: 10.5281/zenodo.2615936) to 601 compute: (a) the position and density of sex-specific SNVs, defined as SNVs heterozygous in 602 one sex but homozygous in the other, and (b) the average read depths for male and female pools 603 along the genome to look for regions present in one sex but absent in the other (i.e., sex-specific 604 insertions). All PSASS analyses were run with default parameters except for the range of 605 frequency for a sex-linked SNV in the homogametic sex, --range-hom, that was set to 0.01 606 instead of 0.05, and the size of the sliding window, --window-size, that was set at 50,000 607 instead of 100,000.

608 Chromosome conventional cytogenetics

609 Mitotic or meiotic chromosome spreads were obtained either from regenerating caudal fin tissue as previously described⁷⁹, with slight modifications⁸⁰ and altered time of fin regeneration 610 (one week), or by direct preparation from the cephalic kidney and gonads⁸¹. In the latter, the 611 612 quality of chromosomal spreading was enhanced by a previously described dropping method⁸². 613 Chromosomes were stained with 5% Giemsa solution (pH 6.8) (Merck, Darmstadt, Germany) 614 for conventional cytogenetic analyses, or left unstained for other methods. For FISH, slides 615 were dehydrated in an ethanol series (70%, 80% and 96%, 3 min each) and stored at -20 °C 616 before analysis. Constitutive heterochromatin was visualized by C-banding⁸³, with 617 chromosomes being counterstained by 4',6-diamidino-2-phenolindole (DAPI), 1.5 µg/ml in 618 antifade (Cambio, Cambridge, United Kingdom).

619 gdf6b probe synthesis for FISH mapping

620 gDNA was extracted using NucleoSpin Kits for Tissue (Macherey-Nagel, Duren, Germany) as 621 described above. A gdf6b fragment comprising the two exons, the intron and 2,260 bp of the 622 proximal promoter (with a total size of 4,368 bp) was amplified by PCR in a total volume of 623 50 µl. The mixture contained 0.5 µM of each primer, a final concentration of 20 ng/µl gDNA, 624 1X of 10X AccuPrime[™] PCR Buffer II, 1U/reaction of AccuPrime[™] Tag DNA Polymerase, High Fidelity (Thermofisher), was adjusted to 50 µl with autoclaved and distilled water. 625 Cycling conditions were as follows: 94 °C for 45 sec, then 35 cycles of (94 °C for 15 sec + 64 626 627 °C for 30 sec + 68 °C for 5 min and 30 sec), and 68 °C for 5 min. The resulting PCR product was cloned into TOPO TA cloning Kit XL (Thermofisher) and after sequence verification it 628 629 was purified using NucleoSpin plasmid DNA purification kit (Machery-Nagel, Düren, 630 Germany) according to the supplier's indications. This Pachón cave gdf6b cloned DNA fragment was labeled by nick translation with Cy3-dUTP using Cy3 NT Labeling Kit (Jena 631 632 Bioscience, Jena, Germany). The optimal fragment size of the probe (approx. 200-500 bp) was achieved after 30 min of incubation at 15 °C. 633

634 Chromosome microdissection and FISH mapping

Twelve copies of B chromosome from *A. mexicanus* male individual (male 10) and twelve copies encompassing two B chromosomes (per cell) from a male individual (male 9) were manually microdissected as previously described⁸⁴ under an inverted microscope (Zeiss Axiovert 135) using a sterile glass needle attached to a mechanical micromanipulator (Zeiss). The chromosomes were subsequently amplified by degenerate oligonucleotide primed-PCR 640 (DOP-PCR) following previously described protocols⁸⁵. One µl of the resulting amplification product was used as a template DNA for a labeling DOP-PCR reaction, with Spectrum Orange-641 642 dUTP and Spectrum Green-dUTP, for Male 9 (2B) and Male 10 (1B), respectively (both Vysis, Downers Grove, USA). The amplification was done in 30 cycles, following previously 643 644 described protocols⁸⁶. Depending on the experimental scheme, the final probe mixture contained i) both painting probes (200 ng each) or ii) a single painting probe (200 ng) and a 645 646 labeled 4,368 bp long fragment containing *gdf6b* gene and its promoter (300 ng; see below). To block the shared repetitive sequences, the probe also contained 4-5 µg of unlabelled 647 648 competitive DNA prepared from female gDNA (on male preparations) or male gDNA (on female preparations). Male and female gDNAs were isolated from liver and spleen using 649 650 MagAttract HMW DNA kit (Qiagen) and Cot-1 DNA (i.e., fraction of gDNA enriched with 651 highly and moderately repetitive sequences) was then generated from them according to previously described protocols⁸⁷. The complete B probe mixture was dissolved in the final 652 653 volume 20 μ l (in case of two painting probes) or 14 μ l (in case of one painting probe and a 654 gdf6b gene probe) of hybridization mixture (50% formamide and 10% dextran sulfate in $2\times$ SSC). 655

656

FISH and whole-chromosome painting

657 The FISH (Fluorescence in situ hybridization on chromosomes) experiments were done using a combination of two previously published protocols^{80,88}, with slight modifications. Briefly, 658 659 the aging of slides took place overnight at 37 °C and then 60 min at 60 °C, followed by 660 treatments with RNase A (200 µg/ml in 2× SSC, 60–90 min, 37 °C) (Sigma-Aldrich) and then 661 pepsin (50 µg/ml in 10 mM HCl, 3 min, 37 °C). Subsequently, the slides were incubated in 1% 662 formaldehyde in PBS (10 min) to stabilize the chromatin structure. Denaturation of 663 chromosomes was done in 75% formamide in 2× SSC (pH 7.0) (Sigma-Aldrich) at 72 °C, for 664 3 min. The hybridization mixture was denatured for 8 min (86 °C) and then pre-hybridized at 665 37 °C for 45 min to outcompete the repetitive fraction. After application of the probe cocktail 666 on the slide, the hybridization took place in a moist chamber at 37 °C for 72 h. Subsequently, 667 non-specific hybridization was removed by post-hybridization washes: two times in 1× SSC 668 (pH 7.0) (65 °C, 5 min each) and once in 4× SSC in 0.01% Tween 20 (42°C, 5 min). Slides 669 were then washed in PBS (1 min), passed through an ethanol series, and mounted in antifade 670 containing 1.5 µg/ml DAPI (Cambio, Cambridge, United Kingdom).

671 Synaptonemal complex immunostaining

Pachytene chromosome spreads from two Pachón cave A. mexicanus males (male 2 and 4) 672 were prepared from testes following the protocol for *Danio rerio*^{89,90} with some modifications. 673 674 Briefly, dissected testes were suspended in 200-600 μ l (based on the testes size and cell density) 675 of cold PBS. Cell suspensions were applied onto poly-l-lysine slides (ThermoFisher), with 1:30 676 (v/v) dilution in hypotonic solution (PBS: H₂O, 1:2 v/v). After 20 min at room temperature 677 (RT), slides were fixed with freshly prepared cold 2% formaldehyde (pH 8.0 - 8.5) for 3 min 678 at RT. Slides were then washed three times in 0.1% Tween-20 (pH 8.0-8.5), 1 min each, and 679 left to dry (1 h). Afterwards, immunofluorescence analysis of synaptonemal complexes took 680 place, using antibodies against the proteins SYCP3 (lateral elements of synaptonemal 681 complexes) and MLH1 (mismatch repair protein; marker for visualization of recombination 682 sites). The primary antibodies - rabbit anti-SYCP3 (1:300; Abcam, Cambridge, UK) and 683 mouse anti-MLH1 (1:50, Abcam) – were diluted (v/v) in 3% BSA (bovine serum albumin) in 684 0.05% Triton X-100/ PBS. After application onto the slides, the incubation was carried out overnight in a humid chamber at 37°C. Next day, slides were washed three times in 0.1% 685 686 Tween-20 in PBS, 10 min each and secondary antibodies, diluted (v/v) in 3% BSA (bovine 687 serum albumin) in 0.05% Triton X-100/ PBS, were applied. Specifically, we used goat anti-688 rabbit Alexa 488 (1:300; Abcam) and goat anti-mouse Alexa555 (1:100; Abcam), and the slides 689 were incubated for 3 h at 37 °C. Then, after washing in 0,1% Tween-20 in PBS (10 min) and 690 brief washing in 0.01% Tween-20 in distilled H₂O, slides were mounted in antifade containing 691 DAPI, as described above.

692 Microscopy and image analysis

693 At least 50 metaphase spreads per individual were analyzed to confirm the diploid chromosome 694 number (2n), karyotype structure, and FISH results. Giemsa-stained preparations were analyzed under Axio Imager Z2 microscope (Zeiss, Oberkochen, Germany), equipped with an 695 696 automatic Metafer-MSearch scanning platform. Photographs of the chromosomes were 697 captured under 100× objective using CoolCube 1 b/w digital camera (MetaSystems, 698 Altlussheim, Germany). The karyotypes were arranged using Ikaros software (MetaSystems, 699 Altlussheim, Germany). Chromosomes were classified according to their centromere 700 positions⁹¹, modified as metacentric (m), submetacentric (sm), subtelocentric (st), or acrocentric (a). FISH preparations were inspected using an Olympus BX53 epifluorescence 701 702 microscope (Olympus, Tokyo, Japan), equipped with an appropriate fluorescence filter set. 703 Black-and-white images were captured under 100× objective for each fluorescent dye with a 704 cooled DP30BW CCD camera (Olympus) using Olympus Acquisition Software. The digital

- images were then pseudo-colored (blue for DAPI, red for Cy3, green for FITC) and merged in
- 706 DP Manager (Olympus). Composed images were then optimized and arranged using Adobe
- 707 Photoshop CS6.

708 Phylogeny and synteny of Gdf6 proteins

Phylogeny and synteny relationships of *Gdf6* genes were inferred from a Genomicus instance⁹²
in which synteny and phylogeny have been reconciled with the Scorpios pipeline⁹³ on 4
tetrapod species, one sarcopterygii species, one holostei species and 13 teleosts (see Figure 3
for species names).

713 Three-dimensional protein modelling

714 Three-dimensional models for the proprotein of A. mexicanus A-Gdf6b and B-Gdf6b were 715 obtained by homology modeling. The amino acid sequences of full-length A-Gdf6b and B-Gdf6b were submitted to automated homology modeling using the hm build macro of the 716 717 software package YASARA⁹⁴. Modeling includes alignment of the two target sequences 718 against sequences from Uniprot via PSI-Blast to build a position-specific scoring function 719 matrix/profile, which is then used to search the PDB structure data bank for suitable modeling 720 targets. Five templates were identified with sufficiently high scores, i.e. PDB entries 4YCG 721 (proprotein complex of GDF2 (alternative naming BMP9))⁹⁵, 5HLY (proprotein complex of 722 Activin A)⁹⁶, 6Z3J (the mature C-terminal growth factor domain of GDF5 in complex with repulsive guidance molecule B)⁹⁷ as well as 6Z3M (which is like 6Z3J but includes neogenin 723 in complex with GDF5 and the repulsive guidance molecule B)⁹⁷, and 3QB4 (which is only the 724 C-terminal growth factor domain of GDF5)⁹⁸. Several initial models were built on the basis of 725 726 these template structures, missing sequence elements were modeled in an automated procedure 727 through YASARA on the basis of an indexed PDB structure database. By this scheme 13 728 models were built, three only covered the C-terminal growth factor domain comprising residues 729 293 to 398, while 10 models covered the proprotein complex consisting of residues 3 to 398 730 (models on the basis of the 5HLY entry) and of residues 66 to 398 (models on the basis of 731 template 4YCG). These models were individually refined by a short molecular dynamics 732 simulation in explicit water to optimize hydrogen bonding and protein packing. Due to the 733 overall low Z-score of the individual models, YASARA used the various models to form a 734 hybrid homology model combining the elements with the highest-scoring factors into a single 735 3D model. This hybrid model was then used for further analysis.

736 Expression analysis by Real- Time PCR

For gene expression studies, mRNA transcripts levels were quantified during gonadal development from 10 dpf to 90 dpf, male and female gametogenesis stages, and finally in 10 adult organs including gonads as described previously⁴⁸. All samples were frozen in liquid nitrogen and stored at -80 °C until RNA extraction. Total RNA extraction from gonads, trunks, and adult organs, followed by cDNA synthesis, and expression analysis by RT-PCR were carried out as previously described⁴⁸. Specific primers were designed for *gdf6b* in the most divergent sequence regions between the two paralogous *gdf6a* and *gdf6b* genes.

744 **RNAScope in situ hybridization of gdf6b**

745 RNA in situ hybridization (ISH) assays have been carried out using the RNAscope® 746 technology (ACD BiotechneTM) on 7 µm cross-sections of 15, 21, 30 and 60 dpf Pachon 747 cavefish fixed in paraformaldehyde 4% overnight at 4 °C and embedded in paraffin after serial 748 dehydration in increasing methanol solutions. Specific probes for Pachon cavefish gdf6b and 749 gsdf were synthesized by ACD BiotechneTM. Sections were collected on Super frost+ slides, 750 heated at 60 °C for 1 h and dewaxed 2 x 5 min in xylene followed by 2x 2 min in Ethanol 100% 751 at RT. Fluorescent ISH was carried out with the Multiplex Fluorescent Reagent Kit v2 (ACD 752 BiotechneTM, ref: 323100) according to manufacturer's protocol. Following hybridization, 753 nuclei were labelled with DAPI (4',6-diamidino-2-phenylindole) staining and slides were 754 mounted with ProLongTM Gold Antifade Mountant (InvitrogenTM) and observed with a Leica 755 TCS SP8 laser scanning confocal microscope.

756 Knockout of Pachón A. mexicanus gdf6b

757 Pachón cave A. mexicanus inactivated for gdf6b were generated using the CRISPR/Cas9 method. Guide RNAs (sgRNAs) targeting two sites located in exon 2 of gdf6b were designed 758 759 using ZiFiT software⁹⁹ (http://zifit.partners.org/ZiFiT/Disclaimer.aspx). DR274 vector (Addgene #42250) containing the guide RNA universal sequence was first linearized with 760 761 Bsa1, electrophoresed in a 2% agarose gel and purified. PCR amplifications were then 762 performed using linearized DR274 as a template and two primers for each sgRNA. Forward 763 primers containing sgRNAs target sequences (#site 1 and #site 2) (bolded and underlined) 764 between the T7 promoter sequence in the 5' end and the conserved tracrRNA domain sequence 765 were as follows. Forward primer (#site 1): 5'-GAAATTAATACGACTCACTATAGGGAGTCTGAAACCGGTTCTGGTTTTAGAGCTA 766 767 GAAATAGCAAG-3'. Forward primer (#site 2): 5'-768

769 AGAAATAGCAAG-3'. Universal Reverse primer: 5'-770 AAAAGCACCGACTCGGTGCCACT-3'. Subsequently, residual plasmid was digested with 771 Dpn1 (renewed once) at 37°C for 3 hours. The final product was purified and used as a DNA template for transcription. The sgRNAs were transcribed using the MAXIscript[™] T7 772 773 Transcription Kit (Ambion) according to the manufacturer's instructions. The sgRNAs were 774 precipitated in 200 µl of isopropanol solution at -20°C, centrifuged and the supernatant was 775 removed. The precipitated sgRNAs were resuspended in RNase-free water. The sgRNAs were 776 co-injected with Cas9 protein. Synthesized RNAs were then injected into 1-cell stage A. 777 mexicanus Pachón cave embryos at the following concentrations: 72 ng/µl for each sgRNA and 778 216 ng/µl for the Cas9 protein (kindly provided by Tacgene, MNHN, Paris). Genotyping was 779 performed on gDNA from caudal fin-clips of adult fishes. CRISPR-positive fish were screened 780 for mutations using a set of PCR primers (P7-P8) (Figure S3A, Data S1E) flanking the sgRNAs 781 target sites leading to a ~470 bp deletion on the exon 2 of the *gdf6b* genes (Figure S10). The genetic sex of the mutants was determined by specific primers (P5-P6) on the gdf6b promoter 782 783 with gap/indel variation between A-gdf6b and B-gdf6b (see STAR methods above and Figure 784 S3A).

785 Histology

Gonads were fixed in Bouin's fixative solution for 48 h and then dehydrated serially in aqueous
70% and 95% ethanol, ethanol/butanol (5:95), and butanol. Tissues were embedded in paraffin
blocks that were cutted serially into 5 µm sections, and were stained with hematoxylin-eosinsafran (HES) (Microm Microtech, Brignais, France).

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791 QUANTIFICATION AND STATISTICAL ANALYSIS

792 Statistical analyses

For the sex genotyping marker based on the heterozygous and specific site of the B chromosome on the exon 2 (position 679 bp of the *B-gdf6b* CDS), the significance of the correlation between this polymorphism and the male phenotypic sex was tested with the Pearson's Chi-squared test with Yates' continuity correction. For gene expression, normality of data residuals, homogeneity of variances and homoscedasticity were verified before performing parametric or non-parametric tests. Consequently, statistical analyses were carried out only with non-parametric tests using RStudio (Open Source version) considering the level

- 800 of significance at p < 0.05. For comparisons between two groups, we used Wilcoxon signed rank
- 801 test. All data are shown as Mean \pm Standard Error of the Mean (SEM).

803 LEGEND OF SUPPLEMENTARY DATAFILE

804 Data S1: Supplementary information on metaphase numbers (A), genome assembly 805 metrics and annotation (B, C, D), primer names (E) and sex linkage of B-gdf6b (F) in 806 Pachón cavefish. Related to STAR Methods and Figures 1 and 2. A) Description of Data S1A. Number of metaphases (NM) containing B chromosomes (Bs) in males and females of 807 808 Pachón cave *A. mexicanus*. % M = percentage of metaphases. Only very few male metaphases 809 did not have a B. In contrast, Bs were not detected in most females (7/11), and when present in 810 females (4/11), they were detected only in very few metaphases and most often as a unique B 811 copy. B) Description of Data S1B. Genome assembly metrics of the Pachón cavefish A. 812 mexicanus male assembly (HiFi PacBio) and comparison with previous publicly available 813 surface (Astyanax mexicanus-2.0) and Pachón cave (Astyanax mexicanus-1.02) genome assemblies. C) Description of Data S1C. Gene annotation of Pachón cavefish B chromosome. 814 815 D) Description of Data S1D. Transposable elements in the Pachón cavefish genome. E) 816 Description of Data S1E. Primer names, sequences, target genes, and their corresponding 817 experiments. F) Description of Data S1F. Association between B-gdf6b specific 818 amplifications and sex phenotypes with different *B-gdf6b* primer sets in a laboratory stock and 819 in wild-caught Pachón cavefish. P-value of B-gdf6b association with sex is based on the 820 Pearson's Chi-squared test with Yates' continuity correction.

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Supplementary figures

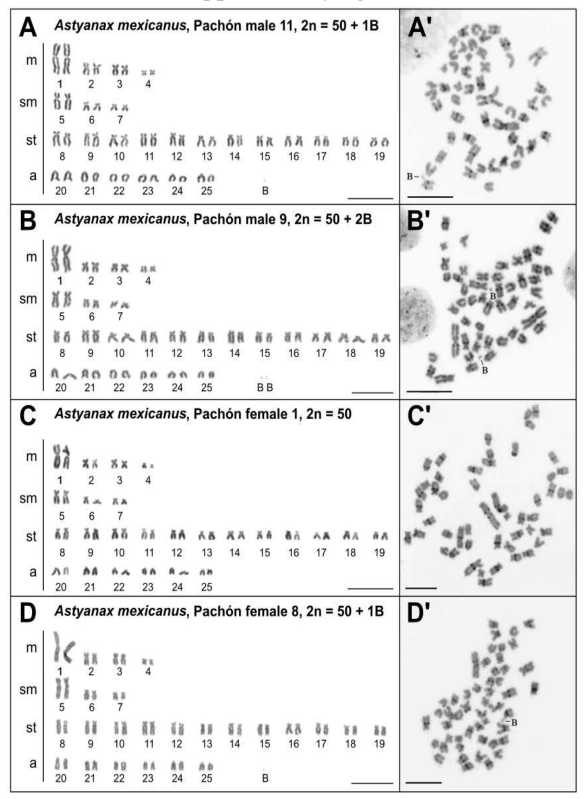
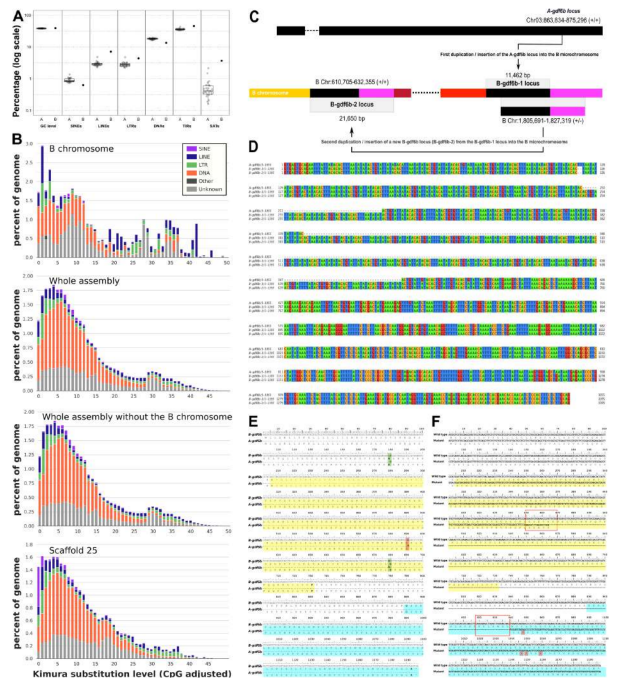


Figure S1. Karyotypes and corresponding C-banded mitotic metaphases of Pachón cave
 Astyanax mexicanus, with different male and female B chromosome (B) constitution.
 Related to Figure 1. Representative male and female Pachón cave karyotypes arranged from
 Giemsa-stained mitotic chromosomes (panels A-D) and their corresponding C-banding

1106 patterns (panels A'-D'). B numbers were found to be variable among individuals (from 0 to 3 1107 Bs) with all males having a single (panels A-A') or multiple Bs (panels B-B') in most of their 1108 metaphases, most females having no B (panels C-C'), and only a few females having rare B 1109 positive metaphases (panels D-D') Notice also the lack of C-bands on Bs suggesting that these 1110 Pachón cavefish male-predominant Bs are largely euchromatic. Scale bar = 10 μ m. Male and 1111 female numbers referred to Data S1A.

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Figure S2: Genomic repeated elements, *gdf6b* evolution and *gdf6b* sequence alignments.
Related to Figure 2. A. Comparison of the GC content and repeated elements in Pachón cavefish B chromosome (B) versus Pachón cavefish A chromosomes. Percentage (in log scale) of GC content (GC level), short interspersed repeated sequences (SINEs), long interspersed nuclear elements (LINEs), long terminal repeats (LTRs), DNA repeat elements (DNAs),

1121 terminal inverted repeat sequences (TIRs), and satellite DNA (SATs) in the 25 A Pachón chromosomes (boxplots, A) versus Pachón B (black dots, B). B. Comparison of the repeat 1122 landscapes of the Pachón B chromosome (B), the whole assembly with (whole assembly) and 1123 1124 without B and of Scaffold 25. Color code for repeat elements is provided in the top right inset of the figure. B repeat landscape is different from the whole assembly with or without the B 1125 chromosome and to the repeat landscape of scaffold 25. The B tends to have a higher content 1126 1127 of interspersed repeats, mainly due to the expansion of long interspersed nuclear elements 1128 (LINEs). Short interspersed repeated sequences (SINEs), long terminal repeats (LTRs), DNA repeat elements (DNAs), and terminal inverted repeat sequences (TIRs). See Data S1D for 1129 1130 additional details. C. The *B-gdf6b* loci on the Pachón cave *A. mexicanus* B chromosome (B) 1131 stemmed from two successive A and B duplications. Schematic representation of the duplication / insertion history of the two *B-gdf6b* loci on the Pachón cave B (HiC scaffold 28) 1132 1133 with a two-steps duplication hypothesis scheme suggesting that the *B-gdf6b-1* locus originated from an initial duplication of a 11.4 kb region surrounding the *A-gdf6b* locus and inserted in 1134 1135 the B, followed by a second independent duplication of a 21.6 kb region surrounding the B-1136 gdf6b-1 locus that was inserted also in the B. This second independent duplication is probably 1137 very recent as the two *B-gdf6b* loci are 99.6 % identical in the 21.6 kb region shared by the two genes. Locations of the duplicated regions are given with respect to the 5'-3' orientation of the 1138 gdf6b cDNA. The colors of the schematic B fragments depict their different A chromosome 1139 origin. D: Multiple sequence alignment of Pachón cavefish intron 1 of A-gdf6b with B-gdf6b-1140 1141 1 and *B-gdf6b-2*. Sequences were extracted from the whole genome Pachón cavefish assembly 1142 with the following coordinates: For A-gdf6b = HiC scaffold 3:864791:865845:-, For B-gdf6b-1143 HiC scaffold 28:617028:618422:+ and for B-gdf6b-2 1 1144 HiC scaffold 28:1832027:1833421:-. The percentage of identity between the A- and B-gdf6b is respectively 1.7 % for intron 1 (18 differences in 1054 bp after gaps and indels collapsed to 1145 1146 1 bp) compared to 0.58 % for their proximal promoter (7 differences in 1200 bp after gaps and indels collapsed to 1 bp; proximal promoter defined from the ATG to the 3'end of the adjacent 1147 gene, ~ 1400 bp). This could suggest different evolutionary constraints between these two 1148 different gdf6b regions. E. Coding sequence and protein alignments of A-gdf6b and B-gdf6b. 1149 Sequences were aligned with CLUSTALW^{S1}. As the gdf6b-B1 and gdf6b-B2 are 100% 1150 identical from their ATG to STOP codons the alignment only shows differences between B-1151 1152 gdf6b (B-gdf6b-1 and B-gdf6b-2) and A-gdf6b coding sequences (CDS). The three nucleotide variations between the *B-gdf6b* and *A-gdf6b* CDS are boxed (green for nonsynonymous sites 1153 and red for synonymous sites) at positions 180 bp, 591 bp and 679 bp positions of the CDS. 1154 1155 Regions highlighted in yellow and blue indicated respectively the Pfam "TGF-β propeptide" and "TGF-β-like" domains^{S2}. ".": Identical nucleotides; "*": stop codon. F. Alignment of 1156 nucleotide (B-gdf6b) and translated protein sequences (B-Gdf6b) in wild type and one 1157 1158 representative example of a mutant male showing a 470 bp deletion in its *B-gdf6b* gene in F0 fish in the exon 2 encoding for the Pfam "TGF- β propeptide" (yellow) and the "TGF- β -like" 1159 (blue) domains^{S2}. The resulting fish displayed a frame-shifted and truncated Gdf6b-B protein 1160 with premature stop codon (red). The positions of the guide RNAs selected to inactivate gdf6b 1161 in Pachón cavefish are boxed in red. Similar 470 bp deletions have been also within the A-1162 gdf6b gene in other fish. 1163 1164

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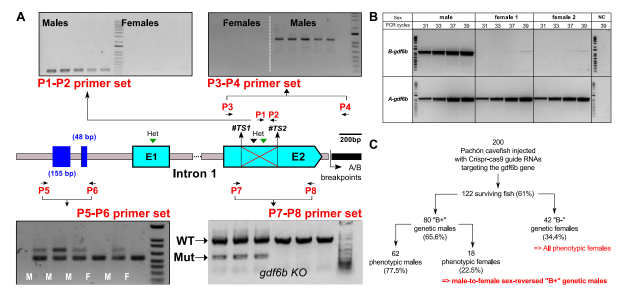


Figure S3: PCR genotyping of the Pachón cavefish male-predominant B chromosome and 1168 the gdf6b knockout mutant individuals. Related to Figure 4 and STAR Methods. A. 1169 1170 Absence / presence of the Pachón cavefish B chromosome was detected with different primer 1171 pairs based on differences between the A-gdf6b and B-gdf6b loci. Three sets of primers (P) were designed with two primer sets designed to amplify specifically the *B-gdf6b* copy based 1172 either on a single base variation between the A/B gdf6b CDS at position 679 bp (P1-P2 primer 1173 1174 set), or based on primers located on both sides of the A/B breakpoints downstream of the Bgdf6b gene (P3-P4 primer set). The third set of primers (P5-P6 primer set) was designed on 1175 gaps/indels variations between A-gdf6b and B-gdf6b in the proximal promoter of gdf6b genes. 1176 1177 Primer sets P1-P2 and P3-P4 produce a single PCR fragment only in B+ individuals, and primer set P5-P6 amplifies two bands in B+ individuals and only a single band in B- individuals. Gene 1178 knockout (KO) was performed by genome editing using the CRISPR-cas9 method with two 1179 guide RNAs target sites (#TS1 and #TS2) designed in order to target gdf6b exon 2 (E2). KO 1180 individuals were genotyped using primers (P7-P8 primer set) flanking the 2 target sites that 1181 induced a 470 bp deletion in *gdf6b* mutant individuals (Mut) compared to the wild type (WT) 1182 1183 gdf6b sequence. B. Increasing PCR cycle numbers allows the detection of a faint PCR fragment 1184 specific to the B chromosome in Pachón cavefish females. PCRs were carried out in one male and two female Pachón cavefish with the protocol described for the P3-P4 primer set and with 1185 1186 increasing PCR cycle numbers. The B chromosome was detected using primers specifically 1187 designed to amplify the B-gdf6b loci (primer set P3-P4) and a control was incorporated with 1188 primers specifically designed to amplify the A-gdf6b locus (primer set P9-P10). NC: PCR negative control. C. Numbers of gdf6b knockout generated by CRISPR-Cas9 method including 1189 1190 the number of fish injected, and the number and percentage of sex-reversed males obtained (in bold red type). Out of the 200 micro-injected eggs (at 1 cell stage), 122 adult fishes were 1191 1192 obtained including 80 genetic males and 42 genetic females. Among the 80 genetic males, we 1193 found 62 phenotypic males (77.5%), and 18 phenotypic females (22.5%) displaying a 470 bp 1194 deletion on the exon 2 of their A-gdf6b and/or B-gdf6b gene. 1195

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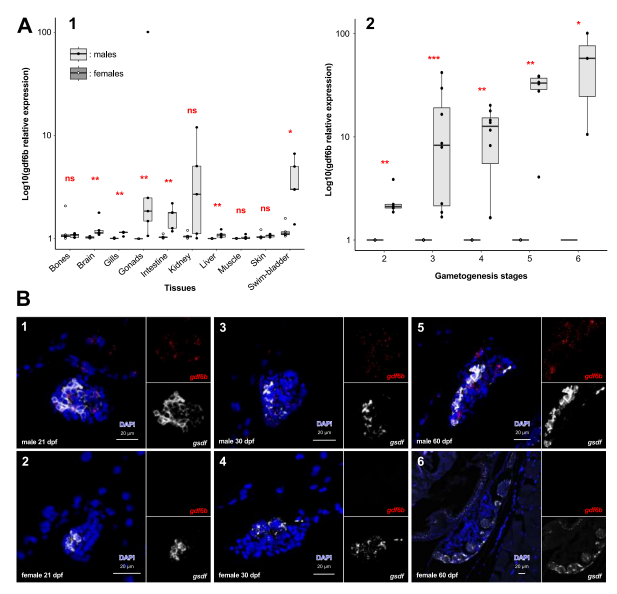


Figure S4. Expression patterns of gdf6b in adult Pachón cave Astvanax mexicanus. 1199 Related to Figure 4. A. Expression profiles of gdf6b (A-gdf6b and B-gdf6b) in different adult 1200 1201 tissues (A1) in males (light grey) and females (dark grey) and during male (light grey) and 1202 female (dark grey) gametogenesis (A2). Gametogenesis stages 2 to 6 were defined as previously described^{S3}. Results are presented as boxplots with individual expression values 1203 1204 displayed as dots, the expression median as a line, and the box displaying the first and third quartiles of expression. Statistical significance between males and females were tested with the 1205 1206 Wilcoxon Rank Sum Test (Wilcoxon-Mann-Whitney Test) and significant differences are *** = P < 0.01; ** = P < 0.01; * = P < 0.05. **B.** Gonadal expression of *gdf6b* (in red) and the Sertoli 1207 1208 and granulosa supporting cell marker gsdf (in white) in male (B1, B3 and B5) and female (B2, B4 and B6) differentiating gonads. At all stages i.e., 21, 30- and 60-days post-fertilization 1209 (dpf) gdf6b is specifically expressed in male gonads with no strict colocalization with gsdf in 1210 1211 males. Nuclei were stained with DAPI (in blue). Scale bar = $20 \,\mu m$.

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1215 SUPPLEMENTAL REFERENCES

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