

### Unraveling the genotype by environment interaction in a thermosensitive fish with a polygenic sex determination system

Benjamin Geffroy, Mathieu Besson, Núria Sánchez-Baizán, Frederic Clota, Alexander Goikoetxea, Bastien Sadoul, François Ruelle, Marie-Odile Blanc, Hugues Parrinello, Sophie Hermet, et al.

### ▶ To cite this version:

Benjamin Geffroy, Mathieu Besson, Núria Sánchez-Baizán, Frederic Clota, Alexander Goikoetxea, et al.. Unraveling the genotype by environment interaction in a thermosensitive fish with a polygenic sex determination system. Proceedings of the National Academy of Sciences of the United States of America, 2021, 118 (50), pp.e2112660118. 10.1073/pnas.2112660118. hal-03485506

### HAL Id: hal-03485506 https://hal.inrae.fr/hal-03485506v1

Submitted on 21 May 2024  $\,$ 

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Unraveling the genotype by environment interaction in a thermosensitive
fish with a polygenic sex determination system
Benjamin Geffroy <sup>a,1</sup> , Mathieu Besson <sup>b,c</sup> , Núria Sánchez-Baizán <sup>h</sup> , Frederic Clota <sup>a,c</sup> ,
Alexander Goikoetxea <sup>a</sup> , Bastien Sadoul <sup>a,d</sup> , François Ruelle <sup>e</sup> , Marie-Odile Blanc <sup>e</sup> , Hugues
Parrinello <sup>f</sup> , Sophie Hermet <sup>g</sup> , Eva Blondeau-Bidet <sup>g</sup> , Marine Pratlong <sup>f</sup> , Francesc Piferrer <sup>h</sup> ,
Marc Vandeputte <sup>a,c</sup> & François Allal <sup>a</sup>
<sup>a</sup> MARBEC Univ Montpellier, CNRS, Ifremer, IRD, Palavas-Les-Flots, France
<sup>b</sup> SYSAAF, Station LPGP/INRAE, Campus de Beaulieu, 35042 Rennes, France
<sup>c</sup> Université Paris-Saclay, INRAE, AgroParisTech, GABI, 78350 Jouy-en-Josas, France
<sup>d</sup> ESE, Ecology and Ecosystem Health, Institut Agro, INRAE, Rennes, France,
<sup>e</sup> Laboratoire Service d'Expérimentations Aquacoles, Ifremer, Palavas Les Flots, France
<sup>f</sup> MGX, BCM, Univ Montpellier, CNRS, INSERM, Montpellier, France.
<sup>g</sup> MARBEC Univ Montpellier, CNRS, Ifremer, IRD, Montpellier, France
<sup>h</sup> Institut de Ciències del Mar, Spanish National Research Council, Barcelona, Spain
<sup>1</sup> To whom correspondence may be addressed. Email: bgeffroy@ifremer.fr
Author contributions: B.G., F.P., M.V. and F.A. designed research; B.G., M.B., F.C., A.G.,
B.S., F.R., M-O.B., H.P., S.H. and E.B-B. performed research; B.G., M.B., E.B-B., F.P.,
M.P., N.S-B, and F.A. analyzed data; and B.G. wrote the paper.
Competing Interest Statement: The authors declare no conflict of interest.
Classification: Biological Sciences, Developmental biology

27 **Keywords:** sex determination, genomics, temperature, fish, epigenetic

#### 28 This PDF file includes:

29 Main Text30 Figures 1 to 6

31

32 Abstract

33 In most animals sex determination occurs at conception, when sex chromosomes are segregated following Mendelian laws. However, in multiple reptiles and fishes, this genetic 34 35 sex can be overridden by external factors after fertilization or birth. In some species, the 36 genetic sex may also be governed by multiple genes, further limiting our understanding of sex 37 determination in such species. We used the European sea bass (Dicentrarchus labrax) as a 38 model and combined genomic (using a SNPs chip) and transcriptomic (RNA-Sequencing) 39 approaches to thoroughly depict this polygenic sex determination system and its interaction 40 with temperature. We estimated genetic sex tendency (eGST), defined as the estimated 41 genetic liability to become a given sex under a liability threshold model for sex determination, 42 which accurately predicts the future phenotypic sex. We found evidence that energetic 43 pathways, concerning the regulation of lipids and glucose, are involved in sex determination 44 and could explain why females tend to exhibit higher energy levels and improved growth 45 compared to males. Besides, early exposure to high temperature upregulated sox3, followed 46 by sox9a in individuals with intermediate eGST, but not in individuals showing highly 47 female-biased eGST, providing the most parsimonious explanation for temperature-induced 48 masculinization. This gonadal state was maintained likely by DNA methylation and the 49 upregulation of several genes involved in histone modifications, including *jmjd1c*. Overall, 50 we describe for the first time a sex determination system resulting from continuous genetic 51 and environmental influences in an animal. Our results provide significant progress in our 52 understanding of the mechanisms underlying temperature-induced masculinization in fish.

53 Significance Statement

54

Traditionally, fish sex determination was considered to be governed by genetic or environmental factors. However, many teleost species defy this dichotomy. We combined genomic and transcriptomic approaches to characterize the temperature-dependent polygenic sex determination of European sea bass. We observed that the estimated genetic sex tendency (eGST) provides an accurate estimation of the phenotypic sex. Our data support the 60 hypothesis that sexually dimorphic growth is the consequence rather than the cause of sex 61 determination. We also showed that temperature-induced masculinization involves the 62 upregulation of *sox3* and *sox9a* for individuals in the middle of the eGST distribution. We 63 unprecedentedly show that sex determination system is influenced by continuous genetic and 64 environmental variation that results in variable proportions of males and females.

65

#### 66 Introduction

67

68 Sex determination is a central biological process with consequences relevant for natural 69 population dynamics and livestock production. A plethora of systems, from purely genetic sex 70 determination (GSD) to environmental sex determination (ESD), have been described in the 71 animal kingdom (1). Sex determination involves the interaction of pro-male and pro-female 72 genetic pathways in birds and mammals (1), but interestingly, those pathways are often 73 impacted by various environmental factors in reptiles and fish (2, 3). Undoubtedly, fishes 74 represent the taxon exhibiting the widest diversity of sex determination systems (4), in which 75 biotic (e.g. density) (5) or abiotic factors (e.g. pH and temperature) (6, 7) can interact with, or 76 even override, the genetic background of sex.

77 These external factors affecting the sex of individuals are then transduced at the physiological 78 level in different manners, depending on the species. Yet, two main routes have been 79 identified in fishes, one involving the stress-axis pathway (8) and another one involving 80 epigenetic mechanisms (differential methylation or histone modification) (9). Interestingly, 81 the latter seems more conserved in reptiles (10, 11) when compared to the former (12). 82 Temperature, the most studied environmental factor affecting fish sex, has been shown to 83 either increase cortisol production (the main stress hormone) with cascading effects on sex 84 (13) or to change methylation profiles in the promoters of key genes mostly involved in sex 85 differentiation (9).

86 Temperature-dependent sex determination (TSD) has been detected in various fish species 87 including the Atlantic silverside (Menidia menidia) (3), the Nile tilapia (Oreochromis 88 niloticus) (14), the olive flounder (Paralichthys olivaceus) (15), the African spiny catfish 89 (Clarias gariepinus) (16), the pejerrey (Odontesthes hatcheri) or the cobaltcap silverside 90 (Hypoatherina tsurugaen) (17). In these species, natural temperatures within the thermal 91 range of what fish usually encounter in the wild can impact sexual fate. Moreover, even in 92 species with a supposedly strong GSD, extreme water temperatures outside the natural 93 thermal range can sometimes override their sex determination pathway (18-20). In most of

94 the above-mentioned species, sex reversal (usually from female to male) induced by 95 temperature fluctuations is relatively easy to detect since the genetic sex can be identified 96 either at the gene (21-25) or at the chromosome level (26, 27), which enables the 97 investigation of the underlying physiological mechanisms of sex reversal on an individual 98 basis. Identifying cases of sex reversal becomes much more complicated when species exhibit 99 a polygenic sex determination system (28). In such instances, each individual presents a 100 specific combination of pro-male and pro-female genes involved in sex determination resulting in a genetic sex tendency (GST), defined as the genetic liability to become a male or 101 102 a female under a liability threshold model (29) for sex determination. This GST is by 103 definition continuous, as opposed to the dichotomic pattern found in species with a master 104 sex-determining gene, in which the presence or the absence of such gene governs sex at 105 conception.

106 The European sea bass (Dicentrarchus labrax) is a gonochoristic species that possesses a 107 GST whereby the genetic architecture (likely involving many genes) interacts with 108 temperature during a labile period where sex can be altered before the sexual fate of the gonad 109 is definetively fixed (30–32). The labile period encompasses the larval and the juvenile stages 110 (Fig. S1). Thus, as it occurs in many other fish species, exposure to relatively high 111 temperature (  $> 17^{\circ}$ C) during the larval stage promotes male differentiation (30–32). 112 However, long-term exposure to relatively low temperature (  $< 17^{\circ}$ C) before gonadal sex 113 differentiation is complete (*i.e.*, during the juvenile stage) can also trigger masculinization 114 (31, 33). In the European sea bass, future females are already bigger when compared to future 115 males (34), and it has been shown that sex-related differences in growth are established well 116 before the appearance of the first currently known molecular markers of sex (34). However, 117 whether being female induces enhanced growth rate or, conversely, if high early growth rate 118 promotes feminization need to be further studied.

119 The fact that this species possesses a polygenic sex determination system, where temperature 120 also influences sex determination, has complicated the studies aiming at deciphering the 121 underpinning mechanisms. Indeed, environmental effects have commonly been detected at the 122 group level (35–37) or after the labile period (36), and genetic effects are deduced from the 123 propensity of specific parents to produce a biased sex ratio (32, 38). While these earlier 124 studies have improved our knowledge of the potential mechanisms involved, they did not 125 allow the identification of the earliest molecular signs of environmental effects, when the 126 gonad is not yet differentiated, even at the molecular level.

127 Here, we took advantage of the recently developed 57K SNP chip in the European sea bass 128 (39) to determine the estimated genetic sex tendency (eGST) of individuals, exposed to either 129 high or low temperatures. A prediction equation for eGST can be obtained by combining 130 multilocus SNP genotypes and sex phenotypes in a training population, and then this equation 131 can be used to estimate the eGST of fish for which only the SNP genotype is available, in a 132 genomic evaluation framework (40). We predicted that the sex of individuals at both extremes 133 of the eGST distributions would not be impacted by temperature, while those exhibiting 134 intermediate values would be sensitive to temperature. To test this hypothesis, individually-135 genotyped fish were sampled at four key time points, during the labile period (SI Appendix, 136 Fig. S1). We then used RNA-Sequencing (RNA-Seq) approaches, both at the whole-body and 137 at the gonadal level, during gonadal sex differentiation. Based on the transcriptomic analysis 138 of individually genotyped fish, we combined gene expression data with sex prediction 139 through eGST data, enabling the investigation of the pathways involved in sex determination 140 of the European sea bass at an early stage. Furthermore, our experimental design allowed us 141 to test the overlooked hypothesis that masculinization by elevated temperature may result 142 from sex-specific mortality of females rather than from induced female-to-male sex reversal, 143 which could not be tested previously, as mortality in larval temperature treatments occurs before any phenotypic or molecular difference between sexes is visible. 144

145

146 **Results** 

# 147 Sex ratio analysis of control and high-temperature-exposed 1-year-old fish validates the 148 eGST model

149 In individuals reared at 21 °C (high temperature, HT, n=493) from 8 to 390 days post hatching 150 (dph), the sex ratio was highly biased towards males ( $75 \pm 8.5\%$  males). On the contrary, those fish that were kept at 16 °C (low temperature, LT, n = 537) from 2 to 59 dph before 151 152 being switched to 21 °C had a more balanced sex ratio ( $46.5 \pm 1.4\%$  males), confirming a 153 strong effect of early exposure to high temperature on the sex of European sea bass (z-value = 154 -6.3, p-value < 0.001). When combining genomic relationships with the phenotypic sex in a 155 single trait threshold model, where low and high temperatures were considered a fixed effect, sex was found to be highly heritable, with a heritability estimate of  $h^2 = 0.56 \pm 0.06$ . In the 156 multi-trait analysis, where sex in each temperature treatment was considered a specific trait, 157 both sex at LT (sex LT) and at HT (sex HT) were also found highly heritable:  $h_{sex LT}^2 = 0.65$ 158  $\pm$  0.06 and h<sup>2</sup><sub>sex HT</sub> = 0.51  $\pm$  0.08, with a strong genetic correlation between both temperature-159

160 specific sex traits,  $r_G = 0.91 \pm 0.09$ . From a leave-one-out cross-validation approach, where 161 we predicted the sex of an individual based on a genomic prediction equation established 162 without providing information on its phenotype, the genomic prediction successfully 163 classified animals in 74.5% of the cases for the fish reared at LT, and 72.5% for the fish at 164 HT, based on the area under the curve values of the receiver operator characteristic (ROC) 165 curves (SI Appendix, Fig. S2). Importantly, we did not detect any skew in the distribution of 166 eGST over time (randomly sampled at the four time points, SI Appendix, Fig. S1) when comparing one temperature to the other (SI Appendix, Fig. S3), emphasizing that the skew in 167 168 sex ratio observed at high temperatures is due to sex reversal, rather than to genotype-specific 169 mortality. Based on the observed sex ratio of 1-year-old fish, we predicted that about 25% of 170 the whole population had sex reversed at HT, and that this would likely concern those 171 individuals with an eGST lower than 0.5 (i.e. "weak" genetic females exhibiting male sex 172 differentiation at high temperature see SI Appendix, Fig. S10).

173 A genome wide association study (GWAS) identified five genomic regions explaining more 174 than 2% of the total genetic variance of the GST (SI Appendix, Fig. S4), which we considered as putative quantitative trait loci (QTL). The region with the highest association with sex was 175 176 in LG7, between positions 6.3 and 6.8Mb (QTL LG7) with a 8% variance explained. Two other regions in LG19 in the region 6.6-7.3Mb (QTL LG19a) and in the region 17.0-17.9Mb 177 178 (QTL LG19b) explained 3.2% and 2.7% of the genetic variance, respectively. In LG13, the 179 region 24.0-24.6Mb (OTL LG13) explained 2.3% of the genetic variance. A last region in 180 LG1A (25.7-26.3Mb – QTL LG1A) explained 2.1% of the genetic variance for GST.

181

### 182 Sox genes and genes linked to histone modification processes are affected by high 183 temperature in fish with intermediate eGST values at the "flexion" stage

184 Whole-body RNA-Seq was performed at the "flexion" stage (25-40 dph), coinciding with the 185 early stages of the labile period for sex determination (SI Appendix, Fig. S1). At this stage, 186 only one germ cell per future gonad is observable in transversal fish sections (41). Ten 187 individuals (five per treatment) were selected for their intermediate, though positive eGST. 188 The DESeq2 analysis highlighted 341 differentially expressed genes (p-value < 0.05) between 189 HT and LT individuals. The GOs response to steroid hormone and cellular response to steroid 190 hormone stimulus, were among the biological processes upregulated at HT (SI Appendix, Fig. 191 S5). Specifically, we found that two sox (Sry-related HMG box) genes classically involved 192 in sex determination and differentiation, sox3 and sox9b, were upregulated at HT (Fig. 1A, 193 B). Seven genes were linked to histone modification processes, two of them upregulated at

194 HT (*ncoa6* and *lpin1*), and five upregulated at LT: *ube2a*, *zbtb7b*, *setd5*, *suz12*, and *auts2*.

195 Fig. 1. Both A) sox3 and B) sox9b were differentially expressed following the DeSeq2 analysis between neutral individuals (0 < eGST < 0.5) at the "flexion" stage kept at high 196 197 temperature (in red) and low temperature (in blue). The number of RNA-Seq transcripts of C) 198 sox3 and D) sox9a differed according to the temperature between groups of eGST < -0.5, -0.5199 < eGST < 0, 0 < eGST < 0.5 and eGST > 0.5 in fish sampled at the "all fins" stage following 200 the DeSeq2 analysis. The RNA-Seq analysis at the all fins stage revealed an overall negative 201 and significant correlation between E) sox3 and eGST and F) sox9a and eGST, as well as a 202 temperature effect. Abbreviations: \*\*\*= p-value < 0.001; \*\* = p-value < 0.01, \* = p-value < 203 0.05; ns= not significant.

204

## Sox genes and genes linked to energy regulation correlate with eGST at the "all fins" stage

207 Whole-body RNA-Seq was performed at the "all fins" stage (53-78 dph period) on 68 individuals (29 LT and 39 HT). This period coincindes with rapid primordial germ cell 208 209 proliferation (41). Among the 17303 genes that respected our inclusion criteria (> 30 reads per 210 gene), we detected 584 genes for which expression was correlated with the eGST (eGST p-211 value < 0.05; eGST x T (°C) > 0.5). Twelve and two genes, respectively (*SI Appendix*, Table 212 S1), were part of the GOs sex differentiation and sex determination, among which sox9a and 213 sox3 (p-value = 0.052) (Fig. 1E, F). For both these genes, we also detected a strong 214 temperature effect on transcript number (p < 0.001), with a higher number of transcripts at HT 215 compared to LT (Fig. 1C, D).

216 Sixteen genes were involved in the GO lipid biosynthetic process and 19 in the GO regulation 217 of growth (SI Appendix, Table S1). The gene encoding the growth hormone (gh) was one of 218 these genes, and was positively and significantly correlated with the eGST. Three other genes 219 (prkca, gfilb and eva2) that are in close vicinity of the three previously detected QTL 220 (OTL LG7, OTL LG19b and OTL LG1A) exhibited a significant correlation with the eGST, 221 though their expression was independent of their SNP genotype of the QTLs (AA, AB or BB). 222 The "response to glucose" was among the biological processes presenting a positive 223 correlation with the eGST, and thus more expressed in females (SI Appendix, Fig. S6). Eleven 224 genes involved in the GO histone modification were also significantly correlated to the eGST 225 (SI Appendix, Table S1). Interestingly, the genes from the GO "histone H3-K27 methylation" 226 and "histone H3-K4 methylation" were negatively correlated with the eGST (thus more 227 expressed in males; SI Appendix, Fig. S6).

228 Using a more stringent significance threshold (p-value < 0.001), four genes (*spry1*, *egfr*, *dpp4*, 229 and *dzip1*) were correlated to the eGST. Only the gene encoding the Daz interacting protein 1 230 (dzip1), involved in spermatogenesis, showed a clear dimorphic expression higher for 231 individuals with negative eGST. The three other genes have a role in growth rate (Epidermal 232 growth factor receptor, Egfr; Sprouty rtk signaling antagonist 1, Spry1) and glucose (The 233 dipeptidyl-peptidase IV, Dpp4) regulation (based on their gene ontology). The first axis of a 234 PCA, representing these four genes (Fig. 2A), was highly correlated to the eGST (Fig. 2B). With a quadratic model, only seven genes (SI Appendix, Table S1) showed both an overall 235 236 linear relationship with eGST (p < 0.01) and significant interaction with temperature, revealed 237 by a temperature-specific quadratic component (p-value < 0.05). Three of these genes were 238 involved in epigenetic processes: sgsm2, entpd2, and map3k3.

239

240 Fig. 2. A) Principal Component Analysis (PCA) of four genes (*dzip1*, *dpp4*, *egfr* and *spry1*) 241 having a highly significant (p-value < 0.001) and linear correlation with the sex tendency (eGST) and detected from the RNA-Seq analysis of whole individuals at the "all fins" stage. 242 243 B) The first component axis strongly correlated to the eGST. C) The energy content (joules.mg-1 of tissue) of fish sampled at the "all fins" stage correlated positively with eGST, 244 so that genetic females displayed slightly higher energy content than males. Fish kept at low 245 246 temperatures also displayed higher energy content than those kept at high temperature. Individuals are represented with a color gradient, from maroon to vellow, representing their 247 eGST. Circles represent fish kept at high temperature (HT = 21 °C; n = 39); and triangles 248 249 those kept at low temperature (LT = 16 °C; n = 29). Abbreviations: ns, not significant.

250

# The juvenile gonadal transcriptome faithfully reflects the underlying eGST independently of temperature influences

253 RNA-Seq was performed on total RNA extracted from the gonads of 42 individuals (21 HT 254 and 21 LT) sampled at the juvenile stage (117-124 dph), before the first signs of 255 morphological sex differentiation (SI Appendix, Fig. S1). Among the 15724 genes that 256 respected our inclusion criteria, 1297 showed a significant (p-value < 0.01) linear correlation, 257 either positive or negative, between their expression level and the eGST, independently of the 258 initial temperature treatment (HT vs LT). Among those genes, nineteen and six genes (SI 259 Appendix, Table S3) were within the gene ontologies (GOs) of sex differentiation and sex 260 determination, respectively, including *cyp19a1a* (gonadal aromatase), *foxl2* (forkhead box l2), 261 *dmrt1* (doublesex and mab-3 related transcription factor 1), *gsdf* (gonadal soma derived 262 factor), amh (anti-Müllerian hormone), sox9a (sry-related HMG box 9a), and insr (insulin 263 receptor). Those genes, well described to be involved in sexual development, allowed to 264 distinguish two groups on the first axis of the Principal Component Analysis (PCA): the 265 differentiating males as opposed to the differentiating females (Fig. 3A). As expected, the 266 correlation was positive for genes involved in ovarian development and negative for those 267 involved in testis development (*SI Appendix*, Fig. 3B). This was confirmed by genes involved 268 in the GO steroids metabolic process, namely *hsd17b1*, *cyp26a*, and *3β-hsd* (*SI Appendix*, Fig. 269 S7).

270 Fig. 3. A) Principal component analysis (PCA) of 7 genes involved in sex determination and 271 differentiation. Data are from the RNA-Seq analysis of the gonads of fish at the juvenile stage (n = 42). The PC1 separated the sex horizontally and explained 88.4% of the variance. The 272 273 PC2 separated the variables vertically and explained 4.1% of the variance. The contribution of the variables (genes) are represented by the arrows. B) Significant (p < 0.01) linear correlation 274 between the estimated genetic sex tendency (eGST) and both insr and sox9a (relative number 275 276 of transcripts on the y axis). For five genes, *cvp19a1a*, *foxl2*, *dmrt1*, *amh*, *gsdf*, and the PC1 277 axis, a dichotomic distribution was observed and modelled with a "quasibinomial" function. Circles represent fish kept at high temperature (HT =  $21 \degree$ C; n = 21); and triangles those kept 278 279 at low temperature (LT =  $16 \degree C$ ; n = 21). Individuals are represented with a color gradient, from maroon to yellow, representing their lower or higher eGST. Abbreviations: \*\*\*= p-280 281 value < 0.001; \*\* = p-value < 0.01.

282

283 Overall, this allowed ascertaining the high relevance of the GST estimated with the Gibbs 284 model (eGST), especially for individuals at both extremes of the distribution, independently 285 of the temperature. Our results were further validated at the group level (eGST > 0 = genetic286 females vs eGST < 0 = genetic males) with DESeq2 on the GO of sex determination (Fig. 287 S8). Fifty-two genes involved in the GO histone modification were also significantly 288 correlated to the eGST (SI Appendix, Table S2), which was confirmed with the "without a 289 priori approach" showing that genes involved in histone methylation and acetylation were 290 also up- or downregulated in differentiating gonads (SI Appendix, Fig. S9). Interestingly, 291 other epigenetic processes such as those involved in the miRNA production, were negatively 292 correlated with the eGST, and thus positively with maleness (SI Appendix, Fig. S9). With the 293 quadratic model used for detecting changes linked to the temperature in the middle of the 294 eGST distribution, only seven genes (thop1, paxip1, sik3, jmjd1c, bcor, wiz, and auts2) showed both an overall linear correlation with eGST (p < 0.01) and a significant interaction 295 296 with temperature for the quadratic term (p-value < 0.05). Four of these genes are involved in 297 epigenetic processes: *jmjd1c*, *bcor*, *wiz*, and *auts2*. The expression of these four genes 298 increased in individuals with an eGST in the middle of the distribution and that were reared at 299 HT, which are the ones with a weak genetic sex determination that are expected to be more influenced by the environment (Fig. 4). 300

301

Fig. 4. Quadratic correlation between the estimated genetic sex tendency (eGST) and genes involved in histone modification, detected from the RNA-Seq analysis of the gonads of fish at the juvenile stage. The four genes exhibit a significant (\*\*= p-value < 0.01) linear correlation with the eGST, plus a significant (\*= p-value < 0.05) interaction with the temperature for the quadratic term (T  $^{\circ}C^{2}$ ). Red and blue points represent respectively fish kept at high temperature (HT = 21  $^{\circ}C$ ; n = 21) or low temperature (LT = 16  $^{\circ}C$ ; n = 21).

308

#### 309 DNA methylation levels of 1-year-old fish gonads

310 Reduced Representation Bisulfite Sequencing (RRBS) was conducted at the 1-year-old fish 311 stage using gonadal tissue from 65 males and 42 females. The statistical analysis of methylation data showed several differentially methylated cytosines (DMCs) between fish 312 313 reared at LT and HT in sox3 and sox9a genes (Fig. 5). For sox3 there was a decrease of 314 methylation levels at HT in both males (P = 0.01844), and females (P = 0.0001636). In males, 315 this gene showed seven hypomethylated DMCs in the first exon, close (< 200 bp) from the 316 transcription start site (TSS; Fig. 5A). In the females, the same positions were 317 hypomethylated in the first exon, with a total of up to 15 DMCs detected, among which two 318 of them, found around 600 bp from the TSS, were hypermethlated (Fig. 5B). The methylation 319 levels of sox9a showed an increase at high temperature in males (p-value = 0.01069), but no 320 significant difference in females (p-value = 0.1866) between LT and HT (Fig. 5 C-F). In 321 males, there were three hypermethylated DMCs towards the end of the gene body (Fig. 5C). 322 However, three out of the five DMCs identified in this region were hypomethylated in 323 females at HT (Fig. 5D).

324

Fig. 5. Boxplots of DNA methylation levels of sox3 (A, B) and sox9a (C, D) in of 1-year-old 325 326 fish testes (maroon) and ovaries (yellow), respectively. Individual DMCs identified within the 327 gene region (left side), and average methylation levels of the gene body  $\pm 2000$  bp (right side). The black line within the box indicates the median of the distribution, and the lower and 328 329 upper hinges display the distribution of values between the first and third quartiles. The upper whisker extends to the maximum value (1.5 \* interquartile range (IQR), and the lower 330 331 whisker extends to the minimum value (1.5 \* IQR). Individual DMCs are defined as CpGs 332 with methylation differences > 15% and q-value < 0.01, while significant differences between average data were assessed with the *t*-test. Abbreviations: \*\* = p-value < 0.01; ns, not 333 334 significant. Circles represent fish kept at high temperature (HT =  $21 \degree$ C; n = 65); and triangles 335 those kept at low temperature (LT =  $16 \degree C$ ; n = 42).

336

#### 337 Gonadal histology

The sampling at the juvenile stage (117-124 dph; n = 10 fish per temperature) confirmed that gonads were still not morphologically differentiated. Nevertheless, some oocytes were sparsely observable, but it was impossible to conclude with confidence on the actual phenotypic sex of individuals based on histological analyses alone. Furthermore, the number of oocytes was not correlated to the eGST.

#### 343 Relationship between energy content, body size and eGST

The energy content (joules.mg<sup>-1</sup>) was not significantly correlated to the eGST at any stage, 344 though it almost reached significance (p = 0.054) at the "all fins" stage, with genetic females 345 346 tending to have higher values than males regardless of their size (Fig. 2C). Individuals from 347 LT presented significantly (p-value < 0.001) higher energy content than those from HT at the 348 "all fins" stages (Fig. 2C), likely because LT fish were older. The size of fish was not 349 correlated to the eGST (length: t-value = 1.56, p-value = 0.13; wet weight: t-value = 1.3, p-350 value = 0.2), while there was an effect of temperature (length: t-value = 4.6, p-value < 0.001; 351 wet weight: t-value = 4.75, p-value < 0.001).

#### 352 **Discussion**

353 Our analysis combining genomic and transcriptomic data allowed to shed new light on the 354 mechanisms involved in temperature-induced masculinization of the European sea bass. The 355 results confirm the high heritability of the GST. Furthermore, the high genetic correlation 356 between sex LT and sex HT suggested low genotype-by-temperature interaction. In other 357 words, the ranking of animals based on their eGST remains very similar at least across the 358 two tested thermal environments. Hence, the effect of larval rearing temperature on sex was 359 mostly additive. This appears to contradict previous results where such interaction occurred 360 (31, 38). However, these previous results were obtained using between-family variation, while 361 we used a genomic relationship matrix for the present study, which is expected to accurately 362 estimate the true genetic parameters (42). Note that the low genotype-by-environment 363 interaction for GST between the two temperature treatments accounts for what happens 364 globally. But it does not impede local GxE interaction occurring at the genes level or the existence of GxE interactions in other populations and/or under other environmental 365 366 circumstances. The prediction equation for GST, which allowed us to estimate the eGST of 367 genotyped individuals, was established on phenotypically sexed individuals at one year of 368 age, and predicted their sex with a 72.5-74.5 % success. The relevance of eGST was further

369 confirmed by transcriptomic analysis of gonads at 117 and 124 dph, i.e. when the first signs of 370 molecular differentiation can be identified (43). Indeed, we detected a strong correlation 371 between key genes involved in sex differentiation in the gonad and the eGST at this juvenile 372 stage. This good match between phenotypic and genetic sex allowed us to determine, for the 373 first time, the eGST of one to three-month-old individuals, when the temperature is known to 374 act on the sex of European sea bass (30). Five putative OTLs, explaining a low but significant 375 part of the variance of GST, were identified in four different chromosomes, (LG7, LG19, 376 LG13 and LG1). Faggion et al. (44) already identified QTL\_LG7 in Northern Atlantic and 377 Mediterranean populations of European sea bass and the two LG19 QTLs in Mediterranean 378 populations only (origin of the present population). The minor QTLs found in LG13 and 379 LG1A were, however, specific of this study. None of the QTLs previously found in LG6, 380 LG11 and LG18-21 (45) were detected in the present study. Overall, our results are consistent 381 with those of previous studies, pinpointing a GST strongly driven by polygenic variation (~ 382 90% of the variance) with a low contribution of minor QTLs (~10%).

383 These QTLs however participate to the accuracy of the model, which appeared to be strong 384 (100%) for individuals at both extremes of the eGST distribution. However, some mismatches 385 were detected in the middle of the distribution, with some individuals with low negative 386 eGST value that likely were phenotypic females (15% at HT and 29% at LT) based on the 387 dichotomic expression of cyp19a1a, foxl2, dmrt1, gsdf and amh; and some individuals with 388 positive eGST values that were likely phenotypic males (25% at HT and 14% at LT). The 389 proportion of fish that are supposedly genetic females, but that exhibit a male phenotype 390 (25%), could well be explained by precocious and relatively long-term exposure to HT (31, 391 46, 47). It also corresponds well to the supposed percentage of masculinized genetic females 392 observed at the end of the experiment: 25%, a figure within the range of masculinization 393 typically observed in European sea bass exposed to HT (48). The mismatch occurring at low 394 temperature (14%) could also be due to the masculinization of fish kept too long at this 395 temperature, since there is what could be regarded as a second period of sensitivity to adverse 396 environmental conditions, including prolonged exposure to low temperature, in European sea 397 bass (31, 37, 46). However, temperature itself might not explain the pattern detected for 398 individuals that are supposedly genetic males. This could come from the fact that some errors 399 occurred in the estimation of eGST (as detected in one-year-old fish, with the leave-one-out 400 approach), which is expected with a polygenic trait that typically has incomplete penetrance, 401 when heritability is lower than 1, which is the case here (49, 50). It could also be that 402 phenological events linked to gonad development are involved. In the European sea bass, the 403 sex is considered "fixed" once animals reach a size of 8 to 10 cm according to some studies 404 (41) but at a size of 4 to 6 cm according to others (34). It is thus possible that individuals in 405 the middle of the distribution can still develop a phenotypic sex not predicted by their eGST, 406 according to the polygenic nature of sex determination in this species, and that their observed 407 transcript values of sex-related genes remain transitory at this stage/age (7.2 cm and 4.5 g). In 408 this sense, the histological analysis did not permit to unambiguously identify males and 409 females at the early juvenile stage, but some intra-testicular oocytes were detected in some 410 individuals, showing that they probably had undergone sex reversal (51). In European eels 411 (Anguilla anguilla), individuals presenting intra-testicular oocytes showed higher levels of 412 cyp19a1a than males (52). It is thus possible that the presence of both tissues in the same 413 gonad drives the mismatch observed between transcript values of sex-related genes and eGST 414 for intermediate individuals. A last plausible explanation involves the sampling design at this 415 stage for transcriptomic analysis. To ensure having sufficient tissue quantity (> 100 ng total 416 RNA), we sampled the biggest fish. Since early growth rate is known to impact sex, it is 417 possible that some of the genetic males (eGST < 0) developed as phenotypic females at this 418 stage.

419 Interestingly, the goodness of the linear fit between genetic and phenotypic sex was 420 confirmed by the gene expression of *sox9a* and *insr*, even when considering individuals in the 421 middle of the eGST distribution and at both temperatures. These two genes are within the GO 422 of sex determination and play a key role in male sex differentiation (1, 53-55). None of them 423 were pinpointed as essential for sex differentiation in previous studies on European sea bass (35, 43, 56). However, both genes are known to be overexpressed at high temperatures in 424 425 TSD reptile species during sex determination (57–59). The fact that both genes present a very 426 linear correlation (as opposed to the dichotomic expression of cyp19a1a, foxl2, dmrt1, gsdf, 427 and *amh*) with the eGST suggests that they are involved early in the process of sex 428 determination, while the other sex-related genes just transduce the future state of the gonad 429 (male or female). This was confirmed at the "all fins" stage with sox9a exhibiting a negative 430 correlation with the eGST, with higher expression at high temperatures for neutral fish. This 431 gene was also shown to play a key role in the ovary-to-testis transition in the zebrafish (60), 432 where it was strongly expressed in pre-Sertoli cells prior to oocyte apoptosis and 433 degeneration. The DNA methylation levels of sox9a increased with high temperatures in 434 testes of 1-year-old fish. This suggests that HT could affect sox9a gene expression already at 435 the "flexion" stage and maintain its state through to adulthood in males. However, the higher 436 expression of this gene found at HT for neutral eGST fish and the hypermethylation detected 437 at HT in males would not match the standard association of hypermethylation with 438 downreguation of gene repression (61). This could be explained by the fact that the DMCs 439 were identified towards the end of the gene body, while it is the methylation level of the first 440 intron, and to a lesser extent, the first exon, which was shown to play an important role and 441 inverse association with gene expression regardless of tissue and species (62).

442 Another gene that was also reported as a marker of germ cells and supporting cells that 443 preferentially develop into testes in zebrafish juvenile ovary-to-testis transformation (60) was 444 *dzip1*, which is concordant with our results where *dzip1* shows a clear dimorphic expression 445 at the "all fins" stage, predominantly at HT. Three other genes were strongly correlated with 446 the eGST at this stage, namely *egfr*, *dpp4*, and *spry1*. The specific function of these genes has 447 not been described in fish and much of our knowledge comes from murine models and 448 humans. Sprouty1 (Spry1 gene product) is involved in fat and bone development (63). Spry1 449 gene knockout in mice adipocytes results in decreased bone mass and increased body fat (63), 450 while adipose tissue-specific expression of the Spryl gene in mice protects against fat 451 accumulation and bone loss (64). EGFR is involved in protein kinase activity and the 452 inhibition of such protein was shown to improve glucose tolerance and favor insulin action 453 (65). Interestingly, its expression was shown to be regulated by spryl (66). DPP4 is also 454 involved in glucose homeostasis and targeted inactivation of this gene in mice yielded 455 individuals with enhanced insulin secretion and improved glucose tolerance (67). All those 456 studies advocate for genetic males having less capacity to produce fat and display appropriate 457 glucose levels. Although this could be viewed as only a conjunction of facts, this hypothesis 458 is enforced by the tendency of genetic females (individuals with high eGST values) to display 459 higher energy content in their tissue (in joules/mg), when compared to males (individuals with 460 low eGST values) at the "all fins" stage. At this stage (53-78 dph), no correlation was found 461 between size and eGST, and the earliest sexual size dimorphism was found at 103 dph in 462 another experiment (68). Overall, all these results support the hypothesis that early growth 463 differences are the consequence rather than the cause of sex differentiation. In that scheme, 464 the polygenic sex determination of the European sea bass provides information that is later 465 transduced at the phenotypic level, as exemplified by the positive correlation between eGST 466 and the *gh* gene at the "all fins" stage.

467 Regarding the role of temperature, at both the "flexion" and the "all fins" stages we detected 468 enhanced expression of *sox3* at a high temperature compared to a low temperature, while this 469 was not detected for genetic females (eGST > 0.5) at the "all fins" stage following 470 transcriptomic analysis. Sox3 is the evolutionary precursor of Sry (sex determining region Y) 471 in mammals (69) and a major master-sex determination gene in three medaka fish species 472 (70). Interestingly, ectopic expression of Sox3 in the developing XX gonads resulted in the 473 complete sex reversal of XX females to males in mouse (Mus musculus) (71), and loss-of-474 function of sox3 caused sex reversal of XY males in the Indian rice fish (Oryzias dancena) 475 (72). The expression of sox3 also gradually increased during the protogynous sex change 476 (female-to-male) of the hermaphroditic fish (*Epinephelus coioides*) (73). This gene is 477 essential in the upregulation of downstream key-related genes for testicular differentiation, 478 gsdf in the Indian rice fish (72) and sox9 in mouse (71). Here we also detected that 479 upregulation of sox3 appeared chronologically before the upregulation of sox9a suggesting 480 similar mechanisms. Sox3 was strongly hypomethylated at HT in both males and females. 481 These methylation levels were very close (< 200 bp) from the transcription start site and 482 within the first exon. The methylation and gene expression data together suggest that HT 483 could affect sox3 expression at the flexion stage through DNA hypomethylation-mediated, 484 unlocking of gene expression and continuing this state to adults through mitosis. The 485 hypomethylation levels found in this gene at 1-year-old gonads and the higher levels of 486 expression of this gene at HT found at earlier stages match the standard negative correlation 487 between DNA methylation and gene expression (61). It is highly difficult to assess whether 488 the observed methylation changes are the cause or the consequence. However, DNA 489 methylation is known to contribute to the acquisition and maintenance of cell identity, making 490 it possible that changes in DNA methylation during sex differentiation contribute to stabilize 491 the gene expression program of each sex. Regarding the exact function of sox9a and sox3, only a proper experiment involving the knockout of these genes would allow to fully 492 493 understand their role in temperature-induced masculinization.

494 Regarding other epigenetic mechanisms, genes from the "histone modifications" GO term 495 were always differentially expressed in all our analyses, confirming their implication in the 496 sex-specific response to temperature (75). However, the specific upregulation of DNA 497 methyltransferases (DNMTs), which are key in the regulation of DNA methylation, was never 498 detected. This was unexpected owing to the previous demonstration of specific methylation of 499 promoters of both cyp19a1 and dmrt1 in males and females of European sea bass, 500 respectively (9, 36). Indeed, epigenetic reprogramming mediated by changes in sexually 501 dimorphic DNA methylation has been suggested to be a key mechanism in the determination 502 of sexual fate in sexually isogenic species (76, 77). Further, we found that key Jumonji family 503 (Jmj) gene *jmjd1c*, involved in histone demethylation at the H3K9 site, was upregulated in 504 male gonads compared to female gonads. Additionally, *jmjd1c* expression was highly

505 impacted by temperature in individuals in the middle of the eGST distribution, with 506 upregulation in the gonads of HT-exposed individuals and downregulation in those exposed to 507 LT. Interestingly, this signal was still dectectable in gonads of LT fish even after the LT 508 treatment had ended (e.g. fish from the LT treatment were at 21 °C from day 60). It has been 509 hypothesized that temperature may reset epigenetic marks thus redirecting sexual fate (1), 510 supporting our observed results. In certain reptilian species, some specific and related histone 511 demethylases (JMJD3, JARID2, and KDM6B), were shown to be crucial in the shaping of the 512 gonadal phenotype during temperature-induced sex-determination (10, 78). Furthermore, 513 although the results here presented are not sufficient to infer the functionality of genes bcor, 514 wiz, and *auts2*, to the best of our knowledge this is the first study to suggest that these genes 515 could be involved in the transduction of temperature signals influencing sexual fate in 516 teleosts, which warrants further examination.

517 To conclude, we found evidence that sex reveral rather than genotype-specific mortality was 518 the cause of some mismatches between the sex predicted by the eGST and the actual 519 phenotypic sex. We did not find any evidence that stress-axis activation was involved in 520 masculinization. Rather, our data supports the involvement of conserved sex-related 521 pathways, epigenetic and energetic processes as key to understand temperature-induced 522 masculinization in the European sea bass. We propose a model where the GST of individuals 523 drives the specific expression of genes involved in lipid and glucose metabolism, 524 independently of temperature (Fig. 6). In that scheme, individuals presenting higher energy 525 content (transduced by higher transcript levels of genes involved in lipids and glucose 526 production) would become females and those with lower energy would become males (Fig. 527 6). This may explain the early sexually dimorphic differences in growth usually observed in European sea bass, and why domestication (and selection for growth) leads to an increased 528 529 proportion of females (79). Once the sex is fixed, a classical cascade of genes involved in sex 530 differentiation is activated, starting by sox9a. However, if fish are exposed very early to high 531 temperatures, this first triggers an upregulation of sox3 that is then followed by an 532 upregulation of sox9a, determining the sex of individuals in the middle of the eGST 533 distribution (Fig. 6). This signal is then maintained, likely thanks to epigenetic processes 534 (DNA methylation and histone modifications, e.g. through *jmjd1c*), which leads to individuals 535 with intermediate eGST values developing as males.

536 From an adaptive and evolutionary point of view, the conditions favoring the emergence of 537 ESD over GSD have been extensively discussed (28, 80). But species where both strategies 538 coexist, with additive effects, provide new challenges for evolutionary scientists. 539 Understanding how external factors can override genetic information, affecting an essential 540 trait such as phenotypic sex is indeed of major importance in a global warming context. 541 Furthermore, the insights provided by this study on European sea bass can help to illuminate 542 other systems where temperature, stress or other envornmental cues can override a GSD 543 system, as reported in vertebrates, from fish to mammals (81). The present study, therefore, 544 helps pave the way for our understanding of such mechanisms: the genetic architecture likely provides information linked to the regulation of energy and growth, constituting a strong 545 546 example in support of the hypothesis linking metabolism and sex determination (82). Any 547 environmental cues that might affect this relationship (e.g. temperature influencing 548 metabolism) will trigger a specific expression of a cascade of genes that will, in turn, affect 549 the phenotypic sex of sensitive genotypes.

550

551 Fig. 6. Summary of the polygenic sex determination system of the European sea bass. In the 552 top panel (temperature-independent sexual phenotype), all fish have their sexual genotype (estimated genetic sex tendency, eGST) within a Gaussian like distribution, with those that 553 554 present a male tendency (maroon) in the left extreme and those that present a female tendency 555 (yellow) in the right extreme. The genes involved in lipid and glucose (in dashed square) regulation are correlated to the eGST, which likely explain the difference in energy content 556 557 between future males and future females at the "all fins" stage (around 60 dph). This "energetic" information likely drives sex determination, starting with the overexpression of 558 559 sox9a. At 120 dph, the gonad is undergoing molecular sex differentiation involving classical 560 sex pathways. Those in the middle of the distribution can still change sex. Now, if European 561 sea bass are kept at high T (°C), these results in the overexpression of sox3 and sox9b for individuals with a relatively "low" female eGST at 30 dph and that is conserved at 60 dph, 562 563 followed by an increase in the expression of sox9a. This sexual phenotype is then likely maintained thanks to the overexpression of key genes involved in histone modification at 564 565 high, but not low, temperatures for individuals in the middle of the eGST distribution. 566 Twenty-five percent of the population is then likely masculinized at HT following these processes. 567

568

#### 569 Material and Methods

- 570
- 571 SI Appendix provides a detailed description of the materials and methods used in this study.
- 572

#### 573 Fish production and rearing

574

575 The fish population used was the result of a mating design including eight males and one 576 female from a West Mediterranean Sea strain of European sea bass, performed by artificial

577 fertilization (22/03/2017). Fertilized eggs were incubated at 14 °C until 48 hours post-

578 fertilization. Eggs were then evenly dispatched in six tanks of 500 L each, and the 579 temperature was gradually increased from 14 °C to 16 °C within one day. Following hatching, 580 fish density was of 50 larvae per liter. Larvae were then exposed to 21 °C (HT) in triplicates 581 or kept at 16 °C (LT) in triplicates, as described in (83). HT treatment consisted in gradually 582 increasing temperature to reach 21 °C, from 3 dph to 8 dph. From 10 dph onwards, fish were 583 fed Artemia nauplii for 40 days, then weaned on a commercial sea bass diet (Pro Start and Pro 584 Wean, BioMar, Nersac, France). For the LT treatment, the temperature was also increased 585 from day 59 (1 °C/day) to day 64, to reach 21 °C. Fish were reared at the Ifremer Plateforme 586 Expérimentale d'Aquaculture (Palavas-les-Flots, France), accredited to use and breed laboratory animals (n°C341926), and the project was approved by the Animal Care 587 588 Committee # 36 COMETHEA under project authorization number APAFIS 19676.

589

#### 590 Sampling

591

592 Fish were evenly sampled in the triplicates of each temperature treatment, at four different 593 developmental stages. Since fish growth is favored at high temperature, the development of 594 fish generally greatly differs between thermal treatments. Hence, to enable data comparison 595 between individuals kept at different temperatures, the first three samplings were carried out 596 at the same sum of degree-day (base 10 °C,  $DD_{10 \ \circ C}$ ), a procedure previously used to allow 597 standardized measurement of growth in fishes (84). Thus, larvae were not collected at the 598 same date, but at the same DD<sub>10 °C</sub>: 77 DD<sub>10 °C</sub>, 242 DD<sub>10 °C</sub>, and 550 DD<sub>10 °C</sub> (see Table S4 599 for details regarding stage, age and size). The fourth and last sampling of juveniles, aiming at 600 obtaining developing gonads, was, however, based on size rather than age, since growth 601 difference between treatments declined with age (Table S4, Fig. S1). For this reason, gonads 602 were sampled on juveniles at 117 dph (HT) or 124 dph (LT). At the end of the experiment, 603 after one year (at 390 dph), all fish were euthanized using benzocaine (150 mg/L), measured, 604 weighed, and sexed by in situ gonad examination. All samples were genotyped with an SNP chip (see below), but only subsamples of those performed at 242 DD<sub>10 °C</sub>, 550 DD<sub>10 °C</sub>, and 605 606 117-124 dph were used for molecular analysis (SI Appendix, Table S4).

607

#### 608 Genotyping

609

610 We genotyped three generations of European sea bass using the Thermofisher DlabChip 611 European sea bass array of 57k SNP markers (Griot et al. 2021). Generation 0 (G0) includes 612 the parents of the 8 sires. Generation 1 (G1) corresponds to the parents (the dam and the 8 613 sires). Generation 2 includes the 2030 offspring, composed of the larvae sampled at 614 77 DD<sub>10 °C</sub> (n = 192), at 242 DD<sub>10 °C</sub> (n = 300) and at 550 DD<sub>10 °C</sub> (n = 280), the juveniles 615 collected at 117 dph (HT, n = 70) and 124 dph (LT, n = 70), and fish sexed at 390 dph 616 (n = 1118). Fish were sampled at random at each stage, and the number of fish per family was 617 known a posteriori following genotyping and parentage assignment (SI Appendix, Table S4). 618 Complete details of the genotyping analysis can be found in SI Appendix, Materials and 619 Methods.

620

## 621 Heritability, estimate genetic sex tendency prediction and genome wide association scan622

623 We predicted the sex using a liability threshold model, which has been used in a large variety 624 of settings, often with diseases (29, 85), but also for sex determination (28, 32). With this 625 model the binary phenotype is the realization of an underlying continuous phenotype, the 626 liability trait, here called phenotypic sex tendency (PST). When PST exceeds a given threshold, animals differentiate as females, while they differentiate as males when PST 627 628 remains below the threshold (SI Appendix, Fig. S10). The phenotypic sex tendency is itself 629 the addition of a genetic and an environmental sex tendency. In a polygenic sex determination 630 system, the PST is considered a quantitative trait, influenced by many genes with small 631 effects plus environmental effects (28). The genetic sex tendency (GST), which cannot be 632 measured directly, is the genetic part of this PST, positive for individuals more likely to 633 develop as females in a neutral environment and negative for individuals more likely to 634 develop as males (SI Appendix, Fig. S10). Variance components and heritability were 635 estimated for phenotypic sex tendency (PST), the underlying liability of the binary sex, with a 636 threshold model using THRGIBBS1F90 (86). Complete details for the establishment of 637 eGST, QTL presence and heritability assessment can be found in SI Appendix, Materials and 638 Methods.

639

#### 640 **Transcriptomics**

641

The transcriptomic approach aimed to detect genes whose expression would be correlated to the eGST. Since we expected to have fewer females at HT vs LT, we collected more individuals in the HT than in the LT treatment to ensure having a sufficient number of females to analyze. Overall, 70 larvae, 40 from the HT group and 30 from the LT group, were

randomly collected at the "flexion" (242 DD<sub>10 °C</sub>,) and the "all fins" stages (550 DD<sub>10 °C</sub>,), 646 647 euthanized (benzocaine 150 mg/L) and snap-frozen in liquid nitrogen. At these stages, the 648 RNA extraction was performed on entire individuals because of the impossibility to neatly 649 dissect the gonads of such small individuals. This was justified from two points of view: i) it 650 has been previously shown that the chance of underestimating gene expression of sex-related 651 genes using body trunks was negligible in sea bass (87) and ii) we aimed to have a whole 652 picture of the physiological processes associated with TSD. At the juvenile stage (117-124 653 dpf), the gonads of 35 euthanized (benzocaine 150 mg/L) individuals from each temperature 654 treatment were collected and snap-frozen in liquid nitrogen. At this stage, the biggest 655 individuals were collected to ensure the sampling/collection of enough gonadal tissue for 656 transcriptomic analysis. All samples were stored at -80°C until RNA extraction. Complete 657 details for RNA extraction, RNA-seq and RNA-seq data analysis can be found in SI 658 Appendix, Materials and Methods.

659

#### 660 Histology

661

At the juvenile stage, 20 gonads (10HT and 10 LT) were fixed in Bouin's fluid for 6–8 h, rinsed in clear water for 1 h and stocked in a 70% alcohol solution. Each gonad was stained with eosin and then placed in agarose (to improve detection) before being dehydrated, and embedded in paraffin. Sections of 5–6 mm thickness were stained with Trichrome de Masson, Haematoxylin Groat, Fuschine Ponceau and Aniline blue using an automated device.

667

#### 668 Elemental analysis

669

670 Elemental analysis was performed to estimate the energy content of fish with a known eGe. 671 Elemental composition was determined using a microVario Elemental analyzer (Elementar), 672 allowing to obtain the percentage of carbon, hydrogen, nitrogen, sulphur and oxygen 673 (CHNSO) per milligram of dry weight. Seventy sampled fish were weighted and measured 674 before being euthanized (benzocaine 150 mg/L) and kept at -20°C. Then, fish at each stages 675 ("flexion", "all fins" and juveniles) were lyophilized simultaneously for 24 hours. Dried fish 676 were then individually homogenized using ball mills. About 1 mg was used for CHNS 677 analyses. Another 1 mg of the same samples were used for oxygen analysis. The analysis was 678 performed by the laboratory of Physical measures (https://lmp.edu.umontpellier.fr/elem) of 679 Montpellier (certified AFNOR, Iso 9001). Individual elemental composition was then

- transformed to a relative energetic content (cal.mg-1 dry weight) using the Given (1986)
- 681 formula, as recommended by the analyser program:
- 682
- 683 Relative energetic content (cal.mg-1) = ((78.3C + 339.1H 33O + 22S) + 152)/1000
- 684 Where C, H, O, and S refer to the percentages of carbon, hydrogen, oxygen, and sulphur in a
- sample. It was then divided by 4.184 to obtain the energetic content in joules.mg-1.
- 686

#### 687 **RRBS library preparation and analysis**

688

689 Genomic DNA was isolated from ~25 mg of frozen gonadal tissue from the 1-year-old fish 690 with the Qiagen Blood and Cell Culture kit (cat. no. 13323). Genome-wide profiling of DNA 691 methylation levels was performed by RRBS using the Premium RRBS Kit (cat. no. 692 C02030033; Diagenode) according to the manufacturer's instructions. Complete details for 693 RRBS sequencing and preliminary analysis can be found in *SI Appendix*, Materials and 694 Methods.

695

#### 696 Data analysis

697

698 Final sex ratio was analysed using a binomial mixed model, with "tank" added as a random 699 factor and "initial temperature treatment" as a fixed factor. For the three transcriptomes 700 datasets of fish at the "flexion" stage, "all fins" stage, and juvenile stage, we adopted a 701 traditional approach based on group comparison, where differentially expressed genes are 702 detected using the Bioconductor (88) package DESeq2 v.1.18.1 (89) in R (90) and data 703 normalized using the default method of DESeq2. Regarding the 10 transcriptomes at the 704 "flexion" stage, group comparisons were performed between LT (n=5) and HT (n=5) fish 705 with a positive, but not extreme eGST (i.e. 0 < eGST < 0.5). Regarding the 68 transcriptomes 706 at the "all fins" stage, individuals were grouped based on their eGST, within each initial 707 temperature treatment (note that they were collected at the same T (°C)). Those at each 708 extreme of the eGST distribution were considered as extremes genetic males (eM, eGST < -709 (0.5) or extreme genetic females (eF, eGST > 0.5), respectively, and those in the middle of the 710 distribution were considered neutrals males (nM, -0.5 < eGST < 0) or neutrals females (nF, 0 711 < eGST < 0.5). Group comparisons between temperature treatments with DESeq2 were thus 712 performed between eight groups, eM-HT (n=6), nM-HT (n=10), nF-HT (n=12), eF-HT 713 (n=11), eM-LT (n=8), nM-LT (n=6), nF-LT (n=13), eF-LT (n=3). For juveniles' gonads, we

considered only two groups, those with an eGST > 0 (genetic females, n=26) and those with an eGST < 0 (genetic males, n=17) to detect differentially expressed genes according to the genetic sex. For all comparisons, genes with an adjusted p-value less than 5% (according to the FDR method from Benjamini-Hochberg) were declared differentially expressed.

718 For the two transcriptomic datasets with the highest number of samples (68 and 43), we also 719 considered another approach using our linear predictor, the eGST, as a fixed value in linear 720 models. Using normalized (DESeq2) and filtered data (keeping those with more than 30 reads 721 in average), we ran a loop in R (R version 4.0.4) to detect all genes that are significantly 722 correlated to the eGST in a linear and a quadratic model where the temperature treatment (HT 723 or LT) is added as an interaction term with the eGST (T x eGST). For the linear model, we 724 considered only the genes without significant interaction with temperature. Once detected, we 725 also ran a linear model with a quasibinomial distribution (link= Logit) for those interesting 726 genes displaying a dichotomic pattern of expression. For the quadratic model, we considered 727 either the genes with a significant correlation with eGST plus a significant quadratic term for 728 the interaction (I(eGST<sup>2</sup>) x T) or those where the quadratic term (I(eGST<sup>2</sup>)) plus the 729 interaction between the temperature treatment and the quadratic term (I(eGST^2) x T) are 730 significant. One outlier (LT) was removed since it was at the extreme for eGST with unusual 731 intermediate values for all sex-related genes, though keeping this outlier resulted in similar 732 outcomes for those genes (SI Appendix, Fig. S11). Linear models were also used to assess the 733 effect of eGST on energy content (joules.mg-1) and the size of the fishes.

734 For all transcriptomic analyses, the GO dataset of the mouse was used as it gives a more 735 complete overview of the genes involved in each pathway. We adopted two strategies: 1) with 736 a priori, where GOs were selected based on our hypotheses (SI Appendix, Materials and 737 Methods) and 2) without a priori, where GOs were automatically detected from a functional 738 enrichment analysis. For this second approach, we used a functional enrichment analysis. The 739 clusterProfiler version 3.16.1 (91), an R package, was used to analyze function profiles of 740 genes to identify major biological functions of genes. GO terms were predicted based on 741 differentially expressed genes (from the group comparisons and linear models), including 742 biological process and cellular component categories. Enrichment analysis was performed and 743 a p-value < 0.05 was considered to indicate a statistically significant difference. PCA was 744 performed using gene expression levels and illustrated using the ggplot2 package. The same 745 package was used for representing raw values of genes for linear, quadratic regressions and 746 selected genes from group comparisons. The "lme4" package was used for mixed models. 747 Heatmaps of standardized expressions (mean subtraction followed standard deviation

- division) were created using the pheatmap package. Methylkit and Genomic Ranges v1.44.0
- (92) packages were used to filter percent methylation data for the target regions: gene body  $\pm$
- 750 2000 bp. DMCs were defined as cytosines with a methylation difference of  $\geq 15\%$  between
- temperature treatments and a significance threshold of q-value < 0.01.
- 752

#### 753 Acknowledgments

754

The study was supported by the European Maritime and Fisheries Fund (3S, Seabass Sex and Stress, grant number 4320175237) allocated to BG. Production of the fish benefited from AQUAEXCEL<sup>2020</sup> TNA grant "Transsexbass" to FP and NSB. Research at the FP lab supported by Spanish Ministry of Science grant no. PID2019-108888RB-I00. We thank Pierre Lopez (MARBEC) for kindly drawing fish at the different stages of development, Thomas Régnier for providing background on CHNSO analysis, Sandrine Skiba for discussion of the

- results and Mako Pegart for samplings.
- 762

#### 763 **References**

764

B. Capel, Vertebrate sex determination: evolutionary plasticity of a fundamental
switch. *Nat. Rev. Genet.* (2017) https://doi.org/10.1038/nrg.2017.60.

M. Charnier, [Action of temperature on the sex ratio in the Agama agama (Agamidae,
Lacertilia) embryo]. *C. R. Seances Soc. Biol. Fil.* 160, 620–622 (1966).

769 3. D. O. Conover, B. E. Kynard, Environmental Sex Determination: Interaction of
770 Temperature and Genotype in a Fish. *Science* 213, 577–579 (1981).

R. H. Devlin, Y. Nagahama, Sex determination and sex differentiation in fish: an
overview of genetic, physiological, and environmental influences. *Aquaculture* 208, 191–364
(2002).

5. B. Geffroy, A. Bardonnet, Sex differentiation and sex determination in eels:
consequences for management. *Fish Fish.* 17, 375–398 (2016).

N. Ospina-Álvarez, F. Piferrer, Temperature-Dependent Sex Determination in Fish
Revisited: Prevalence, a Single Sex Ratio Response Pattern, and Possible Effects of Climate
Change. *PLoS ONE* 3, e2837 (2008).

779 7. U. Römer, W. Beisenherz, Environmental determination of sex in Apistogrammai
780 (Cichlidae) and two other freshwater fishes (Teleostei). *J. Fish Biol.* 48, 714–725 (1996).

781 8. R. S. Hattori, D. C. Castañeda-Cortés, L. F. Arias Padilla, P. H. Strobl-Mazzulla, J. I.
782 Fernandino, Activation of stress response axis as a key process in environment-induced sex

- plasticity in fish. *Cell. Mol. Life Sci.* (2020) https://doi.org/10.1007/s00018-020-03532-9 (July
  20, 2020).
- 785 9. F. Piferrer, *et al.*, The Model of the Conserved Epigenetic Regulation of Sex. *Front*.
  786 *Genet*. 10 (2019).
- 10. I. W. Deveson, *et al.*, Differential intron retention in Jumonji chromatin modifier
  genes is implicated in reptile temperature-dependent sex determination. *Sci. Adv.* **3**, e1700731
  (2017).
- A. Georges, C. E. Holleley, How does temperature determine sex? *Science* 360, 601–
  602 (2018).
- 12. B. Geffroy, M. Douhard, The Adaptive Sex in Stressful Environments. *Trends Ecol. Evol.* 34, 628–640 (2019).
- B. Geffroy, C. Wedekind, Effects of global warming on sex ratios in fishes. J. Fish
  Biol. 97, 596–606 (2020).
- J. F. Baroiller, H. D'Cotta, E. Bezault, S. Wessels, G. Hoerstgen-Schwark, Tilapia sex
  determination: Where temperature and genetics meet. *Comp. Biochem. Physiol. A. Mol. Integr. Physiol.* 153, 30–38 (2009).
- K. Tabata, Reduction of Female Proportion in Lower Growing Fish Separtated from
  Normal and Feminized Seedlings of Hirame *Paralichthys olivaceus*. *Fish. Sci.* 61, 199–201
  (1995).
- 802 16. S. Santi, *et al.*, Thermosensitivity of the sex differentiation process in the African
  803 catfish, Clarias gariepinus: Determination of the thermosensitive period. *Aquaculture* 455,
  804 73–80 (2016).
- R. S. Hattori, *et al.*, The Duplicated Y-specific amhy Gene Is Conserved and Linked to
  Maleness in Silversides of the Genus Odontesthes. *Genes* 10, 679 (2019).
- T. Kitano, Y. Hayashi, E. Shiraishi, Y. Kamei, Estrogen rescues masculinization of
  genetically female medaka by exposure to cortisol or high temperature. *Mol. Reprod. Dev.* 79,
  719–726 (2012).
- K. Valdivia, *et al.*, High Temperature Increases the Masculinization Rate of the AllFemale (XX) Rainbow Trout "Mal" Population. *PLOS ONE* 9, e113355 (2014).
- 812 20. H. Zhou, *et al.*, Temperature-control-induced masculinization in tiger puffer Takifugu
  813 rubripes. *J. Oceanol. Limnol.* 37, 1125–1135 (2019).
- R. S. Hattori, *et al.*, Cortisol-Induced Masculinization: Does Thermal Stress Affect
  Gonadal Fate in Pejerrey, a Teleost Fish with Temperature-Dependent Sex Determination? *PLoS ONE* 4, e6548 (2009).
- 817 22. Y. Hayashi, *et al.*, High temperature causes masculinization of genetically female
  818 medaka by elevation of cortisol. *Mol. Reprod. Dev.* 77, 679–686 (2010).
- T. Kamiya, *et al.*, A Trans-Species Missense SNP in Amhr2 Is Associated with Sex
  Determination in the Tiger Pufferfish, Takifugu rubripes (Fugu). *PLoS Genet.* 8 (2012).

- 821 24. K. Miyoshi, R. S. Hattori, C. A. Strüssmann, M. Yokota, Y. Yamamoto,
- Phenotypic/genotypic sex mismatches and temperature-dependent sex determination in a wild
  population of an Old World atherinid, the cobaltcap silverside Hypoatherina tsurugae. *Mol. Ecol.* 29, 2349–2358 (2020).
- A. Yano, *et al.*, The sexually dimorphic on the Y-chromosome gene (sdY) is a
  conserved male-specific Y-chromosome sequence in many salmonids. *Evol. Appl.* 6, 486–496
  (2013).
- 828 26. J. F. Baroiller, D. Chourrout, A. Fostier, B. Jalabert, Temperature and sex
  829 chromosomes govern sex ratios of the mouthbrooding Cichlid fish Oreochromis niloticus. *J.*830 *Exp. Zool.* 273, 216–223 (1995).
- X. Wang, *et al.*, High temperature causes masculinization of genetically female olive
  flounder (Paralichthys olivaceus) accompanied by primordial germ cell proliferation
  detention. *Aquaculture* 479, 808–816 (2017).
- 834 28. M. G. Bulmer, J. J. Bull, Models of Polygenic Sex Determination and Sex Ratio
  835 Control. *Evolution* 36, 13–26 (1982).
- M. L. A. Hujoel, S. Gazal, P.-R. Loh, N. Patterson, A. L. Price, Liability threshold
  modeling of case–control status and family history of disease increases association power. *Nat. Genet.* 52, 541–547 (2020).
- 839 30. F. Piferrer, M. Blázquez, L. Navarro, A. González, Genetic, endocrine, and
  840 environmental components of sex determination and differentiation in the European sea bass
  841 (Dicentrarchus labrax L.). *Gen. Comp. Endocrinol.* 142, 102–110 (2005).
- 842 31. E. Saillant, *et al.*, Temperature effects and genotype-temperature interactions on sex
  843 determination in the European sea bass (Dicentrarchus labrax L.). *J. Exp. Zool.* 292, 494–505
  844 (2002).
- 845 32. M. Vandeputte, M. Dupont-Nivet, H. Chavanne, B. Chatain, A Polygenic Hypothesis
  846 for Sex Determination in the European Sea Bass Dicentrarchus labrax. *Genetics* 176, 1049–
  847 1057 (2007).
- M. Vandeputte, *et al.*, Low temperature has opposite effects on sex determination in a
  marine fish at the larval/postlarval and juvenile stages. *Ecol. Evol.* **10**, 13825–13835 (2020).
- 850 34. N. Díaz, L. Ribas, F. Piferrer, The relationship between growth and sex differentiation
  851 in the European sea bass (Dicentrarchus labrax). *Aquaculture* 408–409, 191–202 (2013).
- 852 35. N. Díaz, F. Piferrer, Lasting effects of early exposure to temperature on the gonadal 853 transcriptome at the time of sex differentiation in the European sea bass, a fish with mixed 854 genetic and environmental sex determination. *BMC Genomics* **16**, 679 (2015).
- 855 36. L. Navarro-Martín, *et al.*, DNA Methylation of the Gonadal Aromatase (cyp19a)
  856 Promoter Is Involved in Temperature-Dependent Sex Ratio Shifts in the European Sea Bass.
  857 *PLoS Genet* 7, e1002447 (2011).
- 858 37. M. Vandeputte, *et al.*, Low temperature has opposite effects on sex determination in a
  859 marine fish at the larval/postlarval and juvenile stages. *Ecol. Evol.* n/a (2020).

- B60 38. D. Anastasiadi, M. Vandeputte, N. Sánchez-Baizán, F. Allal, F. Piferrer, Dynamic
  epimarks in sex-related genes predict gonad phenotype in the European sea bass, a fish with
  mixed genetic and environmental sex determination. *Epigenetics* 13, 988–1011 (2018).
- 863 39. R. Griot, *et al.*, Genome-wide association studies for resistance to viral nervous
  864 necrosis in three populations of European sea bass (Dicentrarchus labrax) using a novel 57k
  865 SNP array DlabChip. *Aquaculture* 530, 735930 (2021).
- 40. T. H. Meuwissen, B. J. Hayes, M. E. Goddard, Prediction of total genetic value using
  genome-wide dense marker maps. *Genetics* 157, 1819–1829 (2001).
- 868 41. C. Roblin, J. Bruslé, Ontogenèse gonadique et différenciation sexuelle du Loup
  869 Dicentrarchus labrax, en conditions d'élevage. *Reprod. Nutr. Dév.* 23, 115–127 (1983).
- 42. J. Ødegård, T. H. Meuwissen, Estimation of heritability from limited family data using
  genome-wide identity-by-descent sharing. *Genet. Sel. Evol.* 44, 16 (2012).
- 43. L. Ribas, *et al.*, Characterization of the European Sea Bass (Dicentrarchus labrax)
  Gonadal Transcriptome During Sexual Development. *Mar. Biotechnol.* 21, 359–373 (2019).
- 874 44. S. Faggion, M. Vandeputte, B. Chatain, P.-A. Gagnaire, F. Allal, Population-specific
  875 variations of the genetic architecture of sex determination in wild European sea bass
  876 Dicentrarchus labrax L. *Heredity* 122, 612–621 (2019).
- 45. C. Palaiokostas, *et al.*, A new SNP-based vision of the genetics of sex determination in
  European sea bass (Dicentrarchus labrax). *Genet. Sel. Evol.* 47, 68 (2015).
- 46. M. Bláquez, S. Zanuy, M. Carillo, F. Piferrer, Effects of rearing temperature on sex
  differentiation in the European sea bass (Dicentrarchus labrax L.). *J. Exp. Zool.* 281, 207–216
  (1998).
- 47. M. Pavlidis, *et al.*, Evidence of temperature-dependent sex determination in the
  European sea bass (Dicentrarchus labrax L.). *J. Exp. Zool.* 287, 225–232 (2000).
- 48. L. Navarro-Martín, M. Blázquez, J. Viñas, S. Joly, F. Piferrer, Balancing the effects of
  rearing at low temperature during early development on sex ratios, growth and maturation in
  the European sea bass (Dicentrarchus labrax).: Limitations and opportunities for the
  production of highly female-biased stocks. *Aquaculture* 296, 347–358 (2009).
- 888 49. C. A. Alper, Z. Awdeh, Incomplete penetrance of MHC susceptibility genes:
  889 prospective analysis of polygenic MHC-determined traits. *Tissue Antigens* 56, 199–206
  890 (2000).
- 50. D. N. Cooper, M. Krawczak, C. Polychronakos, C. Tyler-Smith, H. Kehrer-Sawatzki,
  Where genotype is not predictive of phenotype: towards an understanding of the molecular
  basis of reduced penetrance in human inherited disease. *Hum. Genet.* 132, 1077–1130 (2013).
- E. Saillant, *et al.*, Sexual differentiation and juvenile intersexuality in the European sea
  bass (Dicentrarchus labrax). *J. Zool.* 260, 53–63 (2003).
- 896 52. B. Geffroy, Y. Guiguen, A. Fostier, A. Bardonnet, New insights regarding gonad
  897 development in European eel: evidence for a direct ovarian differentiation. *Fish Physiol.*898 *Biochem.* **39**, 1129–1140 (2013).

899 J. Kent, S. C. Wheatley, J. E. Andrews, A. H. Sinclair, P. Koopman, A male-specific 53. 900 role for SOX9 in vertebrate sex determination. Development 122, 2813–2822 (1996). 901 54. S. Nef, et al., Testis determination requires insulin receptor family function in mice. 902 *Nature* **426**, 291–295 (2003). 903 55. I. Stévant, S. Nef, Genetic Control of Gonadal Sex Determination and Development. 904 *Trends Genet.* **35**, 346–358 (2019). 905 56. M. Blázquez, L. Navarro-Martín, F. Piferrer, Expression profiles of sex 906 differentiation-related genes during ontogenesis in the European sea bass acclimated to two 907 different temperatures. J. Exp. Zoolog. B Mol. Dev. Evol. 312B, 686–700 (2009). 908 57. N. Moreno-Mendoza, V. R. Harley, H. Merchant-Larios, Temperature Regulates 909 SOX9 Expression in Cultured Gonads of Lepidochelys olivacea, a Species with Temperature 910 Sex Determination. Dev. Biol. 229, 319-326 (2001). 911 S. Radhakrishnan, R. Literman, J. Neuwald, A. Severin, N. Valenzuela, 58. 912 Transcriptomic responses to environmental temperature by turtles with temperature-913 dependent and genotypic sex determination assessed by RNAseq inform the genetic 914 architecture of embryonic gonadal development. PLOS ONE 12, e0172044 (2017). 915 59. C. Shoemaker, M. Ramsey, J. Queen, D. Crews, Expression of Sox9, Mis, and Dmrt1 916 in the gonad of a species with temperature-dependent sex determination. Dev. Dyn. 236, 917 1055-1063 (2007). 918 D. Sun, et al., Sox9-related signaling controls zebrafish juvenile ovary-testis 60. 919 transformation. Cell Death Dis. 4, e930-e930 (2013). 920 L. D. Moore, T. Le, G. Fan, DNA Methylation and Its Basic Function. 61. 921 Neuropsychopharmacology 38, 23–38 (2013). 922 D. Anastasiadi, A. Esteve-Codina, F. Piferrer, Consistent inverse correlation between 62. 923 DNA methylation of the first intron and gene expression across tissues and species. 924 Epigenetics Chromatin 11, 37 (2018). 925 S. Urs, et al., Sprouty1 is a critical regulatory switch of mesenchymal stem cell lineage 63. 926 allocation. FASEB J. 24, 3264-3273 (2010). 927 S. Urs, T. Henderson, P. Le, C. J. Rosen, L. Liaw, Tissue-specific expression of 64. 928 Sprouty1 in mice protects against high-fat diet-induced fat accumulation, bone loss and 929 metabolic dysfunction. Br. J. Nutr. 108, 1025-1033 (2012). 930 P. O. Prada, et al., EGFR Tyrosine Kinase Inhibitor (PD153035) Improves Glucose 65. 931 Tolerance and Insulin Action in High-Fat Diet-Fed Mice. Diabetes 58, 2910–2919 (2009). 932 Z. Koledova, et al., SPRY1 regulates mammary epithelial morphogenesis by 66. 933 modulating EGFR-dependent stromal paracrine signaling and ECM remodeling. Proc. Natl. 934 Acad. Sci. 113, E5731-E5740 (2016). 935 67. D. Marguet, et al., Enhanced insulin secretion and improved glucose tolerance in mice 936 lacking CD26. Proc. Natl. Acad. Sci. 97, 6874-6879 (2000). 937 68. S. Faggion, et al., Sex dimorphism in European sea bass (Dicentrarchus labrax L.): 27

- New insights into sex-related growth patterns during very early life stages. *PLOS ONE* 16,
  e0239791 (2021).
- 69. A. Herpin, M. Schartl, Plasticity of gene-regulatory networks controlling sex
  determination: of masters, slaves, usual suspects, newcomers, and usurpators. *EMBO Rep.* 16,
  1260–1274 (2015).
- 70. T. Myosho, Y. Takehana, S. Hamaguchi, M. Sakaizumi, Turnover of Sex
  Chromosomes in Celebensis Group Medaka Fishes. *G3 Bethesda Md* 5, 2685–2691 (2015).
- 945 71. E. Sutton, *et al.*, Identification of *SOX3* as an XX male sex reversal gene in mice and
  946 humans. *J. Clin. Invest.* 121, 328–341 (2011).
- 947 72. Y. Takehana, *et al.*, Co-option of Sox3 as the male-determining factor on the Y
  948 chromosome in the fish Oryzias dancena. *Nat. Commun.* 5, 4157 (2014).
- 949 73. B. Yao, L. Zhou, Y. Wang, W. Xia, J.-F. Gui, Differential expression and dynamic
- changes of SOX3 during gametogenesis and sex reversal in protogynous hermaphroditic fish. *J. Exp. Zool. Part Ecol. Genet. Physiol.* **307A**, 207–219 (2007).
- 952 74. S. Nakamura, *et al.*, Analysis of Medaka sox9 Orthologue Reveals a Conserved Role
  953 in Germ Cell Maintenance. *PLOS ONE* 7, e29982 (2012).
- 954 75. F. Piferrer, Epigenetics of sex determination and gonadogenesis. *Dev. Dyn.* 242, 360–
  955 370 (2013).
- 956 76. N. J. Gemmell, E. V. Todd, A. Goikoetxea, O. Ortega-Recalde, T. A. Hore, "Chapter
  957 Three Natural sex change in fish" in *Current Topics in Developmental Biology*, Sex
  958 Determination in Vertebrates., B. Capel, Ed. (Academic Press, 2019), pp. 71–117.
- 959 77. A. Tsakogiannis, *et al.*, The Gene Toolkit Implicated in Functional Sex in Sparidae
  960 Hermaphrodites: Inferences From Comparative Transcriptomics. *Front. Genet.* 9 (2019).
- 961 78. C. Ge, *et al.*, The histone demethylase KDM6B regulates temperature-dependent sex
  962 determination in a turtle species. *Science* 360, 645–648 (2018).
- 963 79. B. Geffroy, *et al.*, Parental selection for growth and early-life low stocking density
  964 increase the female-to-male ratio in European sea bass. *Sci. Rep.* 11, 13620 (2021).
- 80. E. L. Charnov, J. Bull, When is sex environmentally determined? *Nature* 266, 828–
  830 (1977).
- 967 81. F. Piferrer, D. Anastasiadi, Do the Offspring of Sex Reversals Have Higher Sensitivity
  968 to Environmental Perturbations? *Sex. Dev.* 15, 134–147 (2021).
- 969 82. U. Mittwoch, The elusive action of sex-determining genes: mitochondria to the
  970 rescue? J. Theor. Biol. 228, 359–365 (2004).
- 83. A. Goikoetxea, *et al.*, Genetic pathways underpinning hormonal stress responses in
  fish exposed to short- and long-term warm ocean temperatures. *Ecol. Indic.* 120, 106937
  (2021).
- 84. K. A. Chezik, N. P. Lester, P. A. Venturelli, Fish growth and degree-days I: selecting a base temperature for a within-population study. *Can. J. Fish. Aquat. Sci.* **71**, 47–55 (2014).

- 85. O. Weissbrod, C. Lippert, D. Geiger, D. Heckerman, Accurate liability estimation
  improves power in ascertained case-control studies. *Nat. Methods* 12, 332–334 (2015).
- 86. S. Tsuruta, I. Misztal, THRGIBBS1F90 for estimation of variance components with
  threshold and linear models. *Proc. 8th World Congr. Genet. Appl. Livest. Prod. Belo Horiz. Minas Gerais Braz. 13-18 August 2006*, 27–31 (2006).
- 87. M. Blázquez, A. González, M. Papadaki, C. Mylonas, F. Piferrer, Sex-related changes
  in estrogen receptors and aromatase gene expression and enzymatic activity during early
  development and sex differentiation in the European sea bass (Dicentrarchus labrax). *Gen. Comp. Endocrinol.* 158, 95–101 (2008).
- 88. R. C. Gentleman, *et al.*, Bioconductor: open software development for computational
  biology and bioinformatics. *Genome Biol.* 5, R80 (2004).
- 987 89. M. I. Love, W. Huber, S. Anders, Moderated estimation of fold change and dispersion
  988 for RNA-seq data with DESeq2. *Genome Biol.* 15, 550 (2014).
- 989 90. R Core Team, R: A language and environment for statistical computing. *R Found*.
  990 *Stat. Comput. Vienna Austria* (2021).
- 991 91. G. Yu, L.-G. Wang, Y. Han, Q.-Y. He, clusterProfiler: an R Package for Comparing
  992 Biological Themes Among Gene Clusters. *OMICS J. Integr. Biol.* 16, 284–287 (2012).
- 993 92. M. Lawrence, *et al.*, Software for Computing and Annotating Genomic Ranges. *PLOS*994 *Comput. Biol.* 9, e1003118 (2013).

995











