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# 1 Evolution of sex hormone binding globulins reveals early

# 2 gene duplication at the root of vertebrates.

3 Yann Guiguen<sup>1</sup>, Jeremy Pasquier<sup>1</sup>, Alexis Fostier<sup>1</sup>, Julien Bobe<sup>1\*</sup>

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- 5 <sup>1</sup> INRAE, LPGP, 35000 Rennes, France
- 6 Correspondence to: Julien Bobe, julien.bobe@inrae.fr

#### 7

# 8 Highlights:9

- Phylogeny, synteny and expression analyses shed new light on Shbg evolution in vertebrates.
- Shbg diversity originates from a duplication event at the root of vertebrate evolution.
- This duplication was followed by many independent losses of *Shbg* paralogs in vertebrates.
- *Shbg* paralogs have acquired different tissue expression patterns.

#### 17 Abstract

Sex hormone-binding globulin (Shbg) is an important vertebrate blood carrier protein 18 19 synthetized in the liver and involved in the transport and local regulation of sex steroids in 20 target tissues. A novel shbg gene (shbgb) with a predominant ovarian expression was 21 recently characterized. Being initially found only in salmonids, this shbgb was originally 22 thought to result from the Salmonid-specific whole genome duplication. Using updated 23 transcriptomic and genomic resources we identified Shbab orthologs in non-salmonid 24 teleosts (European eel, arowana), holosteans (spotted gar, bowfin), polypteriformes 25 (reedfish), agnatha (sea lamprey) and in amphibians, and found that the classical Shbg gene 26 (Shbga) displays a predominant hepatic expression whereas Shbgb has a predominant 27 gonadal expression. Together, these results indicate that these two Shab genes most likely 28 originate from a whole genome duplication event at the root of vertebrate evolution, followed 29 by numerous and independent losses and by tissue expression specialization of Shbga and 30 Shbgb paralogs.

31

32 Keywords: evolution, Sex hormone-binding globulin, teleosts, vertebrates, gene expression33

# 34 **1 Introduction**

35 Sex hormone-binding globulin (Shbg) is mainly known as a blood-protein carrier involved in 36 the transport of sex steroids in the plasma and in the regulation of their bioavailability to 37 target organs. Shbg proteins present sequence similarities with the other LamG domain-38 containing proteins growth arrest-specific 6 (Gas6) and protein S alpha (Pros1) (Joseph, 39 1997; Joseph and Baker, 1992). By transporting and regulating androgens and estrogens 40 access to the gonads, Shbg plays important roles in vertebrates reproduction (Hammond, 2011). The Shbg protein was originally identified in the beta-globulin fraction of the human 41 42 serum (Rosner et al., 1969) and has previously been known as Androgen-binding protein

43 (Abp). Shbg genes and Shbg proteins have been characterized in a variety of tetrapod 44 species with the notable exception of birds (Westphal, 1986; Wingfield et al., 1984). In 45 aquatic vertebrates, Shbg was originally found in the plasma of an elasmobranch, the skate 46 (Raja radiata) (Freeman and Idler, 1969), and of a teleost, the rainbow trout (Oncorhynchus 47 *mykiss*) (Fostier and Breton, 1975). Since that moment on, Shbg have been subsequently 48 identified and studied in many fish species (see for review (Bobe et al., 2010)). Two different 49 shbg genes i.e., shbga and shbgb, have been characterized in teleosts with shbga being the 50 ortholog of the mammalian Shbg, which has been conserved from chondrichthyes to 51 tetrapods and shbgb that has only been reported up to now in salmonids. In contrast to 52 shbga that is mainly expressed in the liver (Bobe et al., 2008), the shbgb transcript was 53 mainly found in the ovary, suggesting a local mediation of the sex steroids effects by the 54 Shbgb protein (Bobe et al., 2008). These two salmonid Shbg proteins share very low identity 55 percentages, with for instance 26% identity between Shbga and Shgbb at the amino acid 56 level in the rainbow trout (Bobe et al., 2010, 2008). In comparison to other vertebrates and 57 teleost fishes, the salmonid ancestor has experienced an additional (4R) whole genome 58 duplication known as the salmonid specific whole genome duplication, or SaGD (Berthelot et 59 al., 2014). For this reason, and because no shbgb gene had ever been reported in a non-60 salmonid species, it was first hypothesized that shbga and shbgb were ohnologous genes 61 resulting from the SaGD (Bobe et al., 2008). This hypothesis was subsequently challenged in 62 another study and the possibility of an ancient duplication followed by a lineage-specific 63 retention in salmonids was suggested (Miguel-Queralt et al., 2009).

The evolutionary history of *Shbg* genes in vertebrates thus remained unclear and deserved further investigations. Using the increasing amount of genomic and transcriptomic data available for many vertebrate species we revisited the evolutionary history of *Shbg* genes. The transcriptomes of 24 actinopterygian species (including 22 teleosts) and vertebrate genomes were included in the analysis, which led to the identification of previously noncharacterized *Shbgb* genes in several non-salmonid vertebrate lineages. Using phylogenomic analyses, we identified several *Shbgb* orthologs in a variety of non-salmonid

71 vertebrate species, including teleosts, non-teleost actinopterygians, amphibians and one 72 agnatha. Combined with synteny reconstruction analysis, we demonstrated that Shbg 73 diversity results from a duplication event much older than the SaGD. To gain new information 74 on the functional evolution of shbg genes, we also used quantitative PCR and next 75 generation sequencing approaches, to characterize the expression profiles of shbga and 76 shbgb transcripts for several actinopterygian species. This showed that the paralogous shbg 77 genes have acquired different expression profiles with shbgb having a predominant gonadal 78 expression contrasting with a predominant liver expression of *shbga*.

# 79 2 Material and Methods

#### 80 2.1 Genomic and transcriptomic databases

81 The genomes of the following species, human, Homo sapiens; tropical clawed frog, Xenopus 82 tropicalis; coelacanth, Latimeria chalumnae; spotted gar, Lepisosteus oculatus and zebrafish, 83 Danio rerio explored usina the Ensembl were genome browser 84 (http://www.ensembl.org/index.html). The rainbow trout (Oncorhynchus mykiss) genomic 85 database was searched using the Genoscope trout genome browser 86 (http://www.genoscope.cns.fr/trout/). The European eel (Anguilla Anguilla) genomic database 87 was investigated using the European eel assembly available at ZF-Genomics 88 (http://www.zfgenomics.org/sub/eel). Transcriptomes of holostean and teleostean species 89 were investigated using the PhyloFish project resource (Pasquier et al., 2016) available at 90 http://phylofish.sigenae.org. The protein sequences of Human SHBG, Xenopus Shbg, 91 zebrafish Shbg, and rainbow trout Shbga and Shbgb were used as queries to identify 92 homologs of Shbga and Shbgb in the different genomic and transcriptomic databases 93 investigated. A similar methodology was used for Gas6 and Pros1 proteins that were 94 relevant to study due to their phylogenetical proximity and structural similarity.

#### 95 2.2 Phylogenetic and synteny analyses

Amino-acid sequences of 126 predicted Shbg (a, b), Gas6 and Pros1 proteins were first 96 97 aligned using ClustalW (Thompson et al., 1994), then alignments were manually adjusted, to improve the quality of the multiple sequence alignments. The JTT (Jones, Taylor and 98 99 Thornton) protein substitution matrix of the resulting alignment was determined using 100 ProtTest software (Darriba et al., 2011). Phylogenetic analysis of the proteins presenting 101 LamG domains (i.e. Shbga, Shbgb, Gas6 and Pros1) was performed using the neighbour 102 joining (NJM) method (MEGA 5.1 software), with 1000 bootstrap replicates (Tamura et al., 103 2011). Trees were edited online with iTOL (Letunic and Bork, 2016) and exported as 104 Scalable Vector Graphics.

105 Synteny maps of the conserved genomic regions in human, Xenopus, coelacanth, spotted 106 gar and zebrafish were constructed based on information available within the Genomicus 107 (Muffato et al., 2010) v75.01 website (http://www.genomicus.biologie.ens.fr/genomicus-108 75.01/cgi-bin/search.pl). Synteny map of the conserved genomic regions in the rainbow trout 109 performed was using the Rainbow Trout Genomicus Server 110 (http://www.genomicus.biologie.ens.fr/genomicus-trout-01.01/cgi-bin/search.pl). The synteny 111 analyses of European eel conserved genomic regions were obtained performing TBLASTN 112 searches in the corresponding genomic database. For each studied gene, the protein 113 sequences of human and zebrafish were used as queries.

114 Multiple alignments plots of *shbgb* genes in salmonids were processed online 115 (http://genome.lbl.gov/vista/) with mVISTA (Dubchak and Ryaboy, 2006; Poliakov et al., 116 2014) using genomic *shbgb* sequences of rainbow trout, *Oncorhynchus mykiss*, Atlantic 117 salmon, *Salmon salar*, and Coho salmon, *Oncorhynchus kisutch*. Putative *shbgb* 118 pseudogenes were retrieved by TBLASTN searches on whole genome sequences using as 119 query the protein sequence of rainbow trout Shbgb.

120

#### 121 2.3 RNA-seq *shbga* and *shbgb* tissue expression in holosteans and teleosts.

122 RNA-seg and *de novo* assembly were performed for all studied species as previously 123 described (Berthelot et al., 2014; Braasch et al., 2016; Pasquier et al., 2016). In order to 124 study the expression patterns and levels of *shbg* transcripts for each actinopterygian species 125 with two shbg genes, we mapped RNA-seq reads on the corresponding shbg coding 126 sequence (CDS) using BWA-Bowtie (Langmead and Salzberg, 2012) with stringent mapping 127 parameters (maximum number of allowed mismatches -aln 2). Mapped reads were counted 128 using SAMtools (Li et al., 2009) idxstat command, with a minimum alignment quality value (-129 q 30) to discard ambiguous mapping reads. For each species, the numbers of mapped reads 130 were then normalized for each *shbg* gene across the eleven tissues using the reads per kilo 131 base per million mapped reads (RPKM) normalization. All RNA-seq data are available here: 132 (http://phylofish.sigenae.org/index.html)

133

#### 134 2.4 Quantitative PCR analysis (QPCR).

135 QPCR was performed using the RNA collections of the PhyloFish RNA-seg project as previously described (Braasch et al., 2016; Pasquier et al., 2016). Briefly, tissues were 136 137 sampled from the same female individual and testis from a male individual, when possible. In 138 some species and depending on the tissues, RNA samples from different individuals were 139 pooled to obtain sufficient amounts of RNA. Total RNA was extracted using Tri-Reagent 140 (Molecular Research Center, Cincinnati, OH, USA) according to the manufacturer's instructions. Reverse transcription (RT) was performed using 1 µg of RNA for each sample 141 142 with M-MLV reverse transcriptase and random hexamers (Promega, Madison, WI, USA). 143 Briefly, RNA and dNTPs were denatured for 6 min at 70°C, chilled on ice for 5 min before the 144 RT master mix was added. RT was performed at 37°C for 1 h and 15 min followed by a 15-145 min incubation step at 70°C. Control reactions were run without reverse transcriptase and used as negative control in the real-time PCR study. Quantitative RT-PCR (QPCR) 146 experiments were performed sing an Applied Biosystems StepOne Plus. RT products, 147

148 including control reactions, were diluted to 1/25, and 4 µl were used for each PCR. All QPCR 149 were performed in triplicates. QPCR was performed using a real-time PCR kit provided with 150 a Fast-SYBR Green fluorophore (Applied Biosystems) with 200 nM of each primer in order to 151 keep PCR efficiency between 80% and 100% for all target shbg genes. The relative abundance of target cDNA within a sample set was calculated from serially diluted cDNA 152 153 pool (standard curve) using Applied Biosystem StepOne V.2.0 software. After amplification, a 154 fusion curve was obtained to validate the amplification of a single PCR product. The fusion 155 curves obtained showed that each primer pair used was specific of a single *shbg* transcript. 156 The negative control reactions were used to estimate background level. Genes were 157 considered significantly expressed when measured level was significantly above background 158 at p<0.05 and within the range of the standard curve. For each studied tissue, cDNA 159 originating from three individual fish were pooled and subsequently used for real-time PCR. 160 Before further analysis, real-time PCR data were collected using the same detection 161 threshold for all studied genes. Data were subsequently normalized using the  $\Delta\Delta$ Ct method 162 to 18S transcript abundance in samples diluted to 1:2,000.

163

#### 164 2.5 Clustering analysis

Expression profiles originating from either QPCR and RNA-seq were represented using supervised clustering methods (Eisen et al., 1998). Hierarchical clustering was processed using centroïd linkage clustering, that uses the average value of all points in a cluster as a reference to calculate distance of other points, with Pearson's uncentered correlation as similarity metric on data that were normalized and median-centered using the Cluster program (Eisen et al., 1998). Results (colorized matrix) of hierarchical clustering analyses were visualized using the Java TreeView program (Saldanha, 2004).

### 173 **3 Results**

#### 174 3.1 Shbg gene evolution in vertebrates

175 In order to decipher phylogenetic relationships among Shbg sequences, a phylogenetic reconstruction of the evolution of Shbg was made based using the alignment of 126 176 177 vertebrate LamG domain-containing proteins. This phylogeny includes Shbg proteins, growth 178 arrest specific 6 proteins (Gas6) and Vitamin K-dependent protein S (Pros1) and the tree 179 was rooted using as outgroup the zebrafish Laminin subunit alpha 4 (lama4). This analysis 180 (Fig. 1) shows that these vertebrate LamG domains proteins cluster into two major clades 181 containing Shbg proteins on one side and the Gas6 and Pros1 proteins on the other side. 182 These Shbg and Gas6/Pros1 clades are both significantly supported with high bootstrap 183 values (i.e. 75% and 100%, respectively). The Shbg clade contains two sub-clades both 184 supported by significant bootstrap values. The Shbga cluster (100% bootstrap support), in 185 red, contains all classical vertebrate Shbg proteins from chondrichthyes to teleosts with the 186 notable exception of birds in which no Shbg proteins have been detected (see also Fig.S2). 187 The Shbgb cluster (93% bootstrap support), in blue, contains not only the salmonid Shbgb 188 proteins, but also other vertebrate sequences outside the salmonid family including 189 sequences from teleosts (European eel and silver arowana), non-teleost bony fishes 190 (reedfish, spotted gar and bowfin), some amphibians and an agnatha, i.e., the sea lamprey 191 (Fig.1 and Fig.S1). The tree topology indicates that Shbgb proteins are not specific to the 192 salmonids lineage and thus suggests a much more ancient origin of Shbgb genes in 193 vertebrates than previously hypothesized (Bobe et al., 2008).

To strengthen this phylogeny-based analysis of Shbg protein evolutionary history we carried out a synteny analysis in order to better support this hypothesis of an ancient origin *Shbga* and *Shbgb* genes. The synteny analysis first revealed that these two *Shbg* genes are located on two different syntenic chromosome regions in vertebrates (see Fig. 2A for the *Shbga* locus and Fig. 2B for the *Shbgb* locus). In contrast, *Gas6* and *Pros1* genes are located on the same syntenic chromosome region in most studied species, with the exception of

200 primates (Fig. 2C). In vertebrates, Shbga, Shbgb, and Gas6/Pros1 are also located in 201 regions containing other syntenic gene families. These neighboring genes are spread over 202 four syntenic regions (Fig. 2A-2D) like for instance for the Atp-related genes (Atp1b2, 203 Atp1b3, Atp4b) that are found in all four syntenic chromosome regions. In addition, Zbtb-204 related genes (Ztb4, Zbt38, Zbt33) and Lamp-related genes (Lamp1, Lamp2, Lamp3) are 205 present in three of the four different syntenic chromosome regions depending of the gene 206 family. Altogether, these results strongly suggest that the diversity of the gene family present 207 on these four syntenic chromosome regions probably results from early whole genome 208 duplication events (VG1 and/or VG2) that occurred at the root of vertebrate evolution with a 209 subsequent complex pattern of gene retention and gene losses.

210 Using the recently released salmonid genome resources, we also re-investigated the 211 presence of additional copies of shbg genes in salmonids and confirmed that shbga and 212 shbab were both retained as single copies (Fig. 1) in rainbow trout. Atlantic salmon and coho 213 salmon suggesting that no functional duplicated copies were retained after the salmonid 214 whole genome duplication (SaGD). However, we also found an additional shbgb gene 215 conserved in these three salmonid species (Fig.3A), but with many stop codons in its deduced open reading frame (see example for rainbow trout in Fig.3B), suggesting that this 216 217 gene ( $\psi$  shbgb) was subsequently pseudogenized after the SaGD.

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#### 219 **3.2** Expression patterns of shbga and shbgb

In all investigated actinopterygians, *shbga* was found to be mainly expressed in the liver, supporting a conserved role for this blood-secreted Shbg (Fig.4). However, in the silver arowana a low *shbga* expression in the gonads is also detected in addition to the predominant liver expression (Fig.4A and Fig.4B). In contrast to *shbga*, expression of *shbgb* is not predominant in the liver in all investigated actinopterygians (Fig.4A and Fig.4B). In contrast, *shbgb* expression is predominantly detected in the gonads (ovary and/or testis) with the notable exception of the European eel.

### 227 4 Discussion

228 In this study, we aimed at investigating the diversity of the Shbg family in vertebrates and the 229 evolutionary history of Shbg genes. To date, despite the recent discovery of a second shbg gene (i.e. *shbgb*) in salmonids, the origin and diversity of *Shbg* genes in vertebrates has 230 231 remained controversial. Because salmonids experienced an additional whole genome 232 duplication (SaGD) approximately 100 Mya (Berthelot et al., 2014; Macqueen and Johnston, 233 2014) compared to other teleost fish and as *shbab* genes were initially found in salmonids 234 and, never reported at that time in any non-salmonid species, shbga and shbgb have been 235 first hypothesized to be the result of the SaGD (Bobe et al., 2010, 2008). However pairwise 236 comparison of Shbga and Shbgb reveals a surprisingly low sequence identity (around 25% at 237 the amino acid level) that was initially interpreted as Shbga and Shbgb being highly divergent 238 SaGD paralog. However, this paralogy relationship was not supported by the phylogeny 239 reconstruction (Bobe et al., 2008) and this discrepancy was thus explained as being the 240 result of a long-branch attraction artifact resulting from the dramatic divergence of these two 241 sequences (Bobe et al., 2010, 2008). Based on the cloning of another salmonid *shbgb* gene 242 in coho salmon, Oncorhynchus kisutch and the low sequence identity between salmonids 243 Shbga and Shbgb, other authors (Miguel-Queralt et al., 2009), hypothesized that the shbgb 244 gene could stemmed from a much more ancient duplication than the SaGD. In order to 245 decipher the evolutionary history of Shbg genes we re-analyzed the phylogenetic 246 relationships of Shbg genes, their local synteny context, and the evolution of the 247 phylogenetically and structurally closely related genes i.e., Gas6 and Pros1 that also contain 248 LamG domains and are often identified as potential members of the same family. The 249 identification of new Shbgb genes in vertebrates, the Shbg phylogenetic tree topology and 250 their local synteny relationships strongly suggest that Shbga and Shbgb genes result from a 251 whole genome duplication event that occurred very early at the root on the vertebrate 252 lineage. The presence of a single Shbga gene and a single Shbgb gene in amphibians, 253 holosteans, polypteriformes, agnatha and some teleost fishes, suggests that this Shbg

254 duplication stems at least from the second round of vertebrate genome duplication (VG2). It 255 is however also possible that Shbga and Shbgb originate the first round of vertebrate 256 genome duplication (VG1) followed by the loss of one duplicate of each gene before VG2. 257 Following this early duplication, these two Shgb paralogs have evolved through many 258 different phylum-specific gene retentions and/or gene losses. Among them the case of birds 259 is interesting as they not only lost their Shbgb gene like reported here for many other 260 tetrapods, but also their Shbga gene that is found to be conserved in all other vertebrates. 261 This complete absence of Shbga in birds has been already reported and it was hypothesized 262 that this specific steroid hormone-binding transport would then be performed by a 263 corticosteroid-binding globulin (Wingfield et al., 1984). Similarly, no Shbg homologs were 264 detected by homology searches in Chondrichthyes (data not shown) but their complete 265 absence in this clade requires further in-depth analysis and additional genome information as 266 Shbg-like sex steroid binding capacities exist in the serum of the Thorny skate (Freeman and 267 Idler, 1969). In tetrapods, Shbgb was only found in Amphibians along with Shbga. Shbgb 268 was also found in holosteans (spotted gar and bowfin), polypteriformes (reedfish), agnatha (sea lamprey) and in a few teleost fish orders i.e., in Elopomorphs (European eel), 269 270 Osteoglossiforms (silver arowana) and Salmoniforms even though the protein is frequently 271 misannotated in GenBank (Fig.S1). Interestingly we did not find any retention of additional 272 whole genome [SaGD and the teleost specific duplication (TGD)] paralogs for both shbga 273 and shbgb gene with the exception of a pseudogenized SaGD shbgb paralog ( $\psi$  shbgb). This 274 indicates that these extra whole genome duplications did no impact the repertoire of shbg 275 genes with a maximum of one shbga and one shbgb functional copies in all investigated 276 teleost clades. This systematic and independent losses of additional shbga and shbgb 277 duplicated paralogs in teleosts may reflect an evolutionary constraint of maintaining a correct 278 gene and protein dosage as it has been suggested in other organisms (Conant et al., 2014; 279 Gout and Lynch, 2015).

In consistency with existing data in mammals, our expression data showed that *shbga* is predominantly expressed in the liver in the different teleost species studied here. This

confirms what has previously been reported in various teleost species including zebrafish
(*Danio rerio*) (Miguel-Queralt et al., 2004), rainbow trout (*Oncorhynchus mykiss*) (Bobe et al.,
2008), Coho salmon (*Oncorhynchus kisutch*) (Miguel-Queralt et al., 2009), pejerrey
(*Odontesthes bonariensis*) (González et al., 2017) and sea bass (*Dicentrarchus labrax*)
(Miguel-Queralt et al., 2007). In addition, this strong hepatic expression is also observed in
spotted gar (*Lepisosteus oculatus*) and bowfin (*Amia calva*) as shown by both RNA-seq and
QPCR data.

289 In contrast to *shbga*, data on the tissue distribution of *shbgb* remain scarce. The ovarian 290 predominant expression of shbgb was originally reported in rainbow trout, in which the 291 transcript could also be detected at lower levels in muscle and stomach (Bobe et al., 2008). 292 Semi quantitative data in Coho salmon confirmed the expression of *shbgb* in the ovary and 293 stomach and revealed its presence in gills (Miguel-Queralt et al., 2009). Here we show that 294 shbgb is also predominant expressed in the ovary in brown trout, silver arowana and 295 grayling. We also report a strong testicular expression of *shbqb* in the two holostean species, 296 spotted gar and bowfin, that appears to be lost in teleosts. In addition, the shbgb gene does 297 not exhibit any gonad predominant expression in European eel. Together, our data show that 298 shbga and shbgb have a very specific expression patterns with a predominant expression in 299 liver and gonads, respectively. This pattern appears to be conserved during evolution without 300 any significant change following whole genome duplications events (TGD and SaGD), with 301 the exception of European eel in which the gonad predominant expression of shbgb appears 302 to be lost. Finally, the strong testicular expression of *shbgb* revealed in bowfin and spotted 303 gar is not found in any teleost species suggesting a specific role of Shbgb in testicular 304 physiology in holostean species.

305

The multiple independent losses of *Shbgb* across vertebrates, while *Shbga*, *Gas6* and *Pros1* have been conserved in almost all vertebrates, could reflect different adapative and reproductive strategies as Shbg have been shown to be important carrier proteins for the blood transport of sex steroids and for their delivery to target reproductive tissues

(Hammond, 2011). However, despite this discrepancy among species, the distinct roles of Shgba in hormone transport in the blood and of Shbgb in local hormone action in reproductive organs as well as the associated expression in liver and gonads, respectively, appears to be evolutionary conserved in species that have retained both genes despite a few intriguing species-specific exceptions.

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Shbgb

Figure 1: Phylogenetic reconstruction of the evolution of LamG domains proteins
(Shbg, Gas6 and Pros1) in vertebrates. Circular NJM phylogenetic tree of LamG domains
proteins including Shbga in red, Shbgb in blue, Gas6 in purple and Pros1 in green. The tree
is rooted using zebrafish Laminin Subunit Alpha 4 (lama4) and bootstrap values over 0.75
are shown with a black dot on each significant node.



441 Figure 2: Synteny maps of conserved genomic regions around LamG domains 442 proteins (i.e. Shbga, Shbgb, Gas6 and Pros1) in human, Xenopus, coelacanth, spotted 443 gar, Euroean eel, zebrafish and rainbow trout. Synteny maps are given for genomic 444 regions around Shbga (A), Shbgb (B), Gas6 and Pros1 locus (C) and for a fourth region containing homologs of neighbouring Shbga, Shbgb, Gas6 and Pros1 genes (D). Genes are 445 represented by blocks with an arrowed side indicating the gene orientation on chromosomes, 446 447 linkage group or scaffolds. Gene location on chromosomes (Chr for Human and zebrafish), 448 Linkage group (LG for spotted gar) and scaffolds (Ensembl reference or scaffold number) is 449 given in Mb below each gene block. Genes belonging to the same Chr, LG or scaffolds are 450 linked by a solid line.



Figure 3: Multiple alignments plots of shbgb genes in salmonids. (A). Percentage of sequence identity of shbgb genes in Atlantic salmon, Salmon salar, and Coho salmon, Oncorhynchus kisutch compared to the rainbow trout, Oncorhynchus mykiss, shbgb gene on linkage group (LG) 24. In addition to the functional shbgb genes that were found in all salmonid species investigated, all these species have an additional shbgb homolog containing multiple stop codons and thus considered as a pseudogene ( $\psi$  shbgb). (**B**). Rainbow trout Shbgb and  $\psi$  Shbgb protein alignment showing that the  $\psi$  Shbgb contains multiple stop codons (red asterisks) and a large deletion in exon 4 of the Shbgb protein. 



467 Figure 4: Tissue expression profiles of shhbga and shbgb genes. (A). Heatmap 468 (colorized matrix) of the hierarchical clustering of tissue RNA-Seq expression profiles (NGS 469 values) of *shbg* genes in different Holostean (spotted gar and bowfin) and teleost species. A 470 predominant expression of shbga is found in the liver contrasting with the predominant expression of shbgb in gonads (B). Heatmap (colorized matrix) of the hierarchical clustering 471 472 of tissue expression profiles of shbg genes analyzed by guantitative PCR (QPCR values) in 473 different Holostean (spotted gar and bowfin) and teleost species. Colorized matrixes highlight 474 the high expressing tissues in yellow, the low expressing tissues in blue and the median 475 expression in black (see color scale).

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480 Figure 5: Evolution of Shbg/shbg, Gas6/gas6, and Pros1/pros1 genes through successive whole genome duplications and independent gene losses. (A). Schematic 481 482 representation of the evolution of LamG domains proteins including, Shbga (red squares), Shbgb (blue squares), Gas6 (purple squares) and Pros1 (green squares) in some jawed 483 vertebrates' representative species. Whole genome duplications (black circles; VG1 and 484 VG2: vertebrate genome duplications 1 and 2, TGD: teleost genome duplication, SaGD: 485 486 salmonid genome duplication) are indicated at each duplication nodes. Gene losses are represented by square boxes with a cross inside. (B). Simplified representation of the 487 488 evolution of Shbg proteins after whole genome duplications showing the systematic losses of 489 one Shbga and Shbgb paralog after each duplication.