

# Characterization of Sex Determination and Sex Differentiation Genes in *Latimeria*

Mariko Forconi<sup>1</sup>, Adriana Canapa<sup>1\*</sup>, Marco Barucca<sup>1</sup>, Maria A. Biscotti<sup>1</sup>, Teresa Capriglione<sup>2</sup>, Francesco Buonocore<sup>3</sup>, Anna M. Fausto<sup>3</sup>, Daisy M. Makapedua<sup>4</sup>, Alberto Pallavicini<sup>5</sup>, Marco Gerdol<sup>5</sup>, Gianluca De Moro<sup>5</sup>, Giuseppe Scapigliati<sup>3</sup>, Ettore Olmo<sup>1,9</sup>, Manfred Schartl<sup>6,9</sup>

1 Dipartimento di Scienze della Vita e dell'Ambiente, Università Politecnica delle Marche, Ancona, Italy, 2 Dipartimento di Biologia Strutturale e Funzionale, Università Federico II, Napoli, Italy, 3 Dipartimento per l'Innovazione nei Sistemi Biologici, Agroalimentari e Forestali, Università della Tuscia, Viterbo, Italy, 4 Faculty of Fisheries and Marine Science, University of Sam Ratulangi, Manado, Indonesia, 5 Dipartimento di Scienze della Vita, Università di Trieste, Trieste, Italy, 6 Physiological Chemistry, Biocenter, University of Wuerzburg, Wuerzburg, Germany

## **Abstract**

Genes involved in sex determination and differentiation have been identified in mice, humans, chickens, reptiles, amphibians and teleost fishes. However, little is known of their functional conservation, and it is unclear whether there is a common set of genes shared by all vertebrates. Coelacanths, basal Sarcopterygians and unique "living fossils", could help establish an inventory of the ancestral genes involved in these important developmental processes and provide insights into their components. In this study 33 genes from the genome of *Latimeria chalumnae* and from the liver and testis transcriptomes of *Latimeria menadoensis*, implicated in sex determination and differentiation, were identified and characterized and their expression levels measured. Interesting findings were obtained for *GSDF*, previously identified only in teleosts and now characterized for the first time in the sarcopterygian lineage; *FGF9*, which is not found in teleosts; and *DMRT1*, whose expression in adult gonads has recently been related to maintenance of sexual identity. The gene repertoire and testis-specific gene expression documented in coelacanths demonstrate a greater similarity to modern fishes and point to unexpected changes in the gene regulatory network governing sexual development.

Citation: Forconi M, Canapa A, Barucca M, Biscotti MA, Capriglione T, et al. (2013) Characterization of Sex Determination and Sex Differentiation Genes in *Latimeria*. PLoS ONE 8(4): e56006. doi:10.1371/journal.pone.0056006

Editor: Jennifer V. Schmidt, University of Illinois at Chicago, United States of America

Received September 21, 2012; Accepted January 3, 2013; Published April 25, 2013

**Copyright:** © 2013 Forconi et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. No current external funding sources for this study.

Competing Interests: The authors have declared that no competing interests exist.

- \* E-mail: a.canapa@univpm.it
- These authors contributed equally to this work.

### Introduction

Two major processes take place in sexual development: sex determination and sex differentiation. The former process determines whether the bipotential primordium will develop into a testis or an ovary; the latter takes place after sex determination and involves the actual development of testes or ovaries from the undifferentiated gonad [1]. Sex determination is considered as a default pathway or as suppression thereof and initiation of the opposite pathway; in contrast, sex differentiation seems to result from the antagonistic relationship among the genes influencing testis or ovary development [2,3]. Recently it has emerged that sex-specific mechanisms, which are critical to maintaining the male or female identity of the testis and ovary, also operate in adult mammalian gonads [4-6]. Other organs besides the gonads may also acquire elaborate male- and female-specific differences. In vertebrates—with the possible exception of birds [7]—such secondary sexual traits are generally believed to be instructed exclusively by the developing testis or ovary through sex steroids, whereas in invertebrates each somatic cell seems to have an inherent sexual identity [8]. Compared with eutherian mammals, sex steroids and the proteins involved in their metabolism and binding play an earlier role in the sex differentiation process of fish, amphibians, reptiles, birds, and marsupials [9–20].

In vertebrates sexual development is determined by two main factors: either the genetic makeup of the individual or the environment, through the influence of temperature during development, nutrients, pH, etc [21-23]. It has been demonstrated that in mammals the consecutive processes of sex determination, gonad differentiation and identity maintenance are brought about by a complex network of transcription factor interactions and signalling molecules; a master regulator upstream then directs the network towards male or female [24]. The male-determining gene in most mammals is the Y chromosome SRY gene, which however has only been detected in placental mammals [25]. In chickens (and possibly all birds) the master regulator of sexual development is Dmrt1; its homologues are dmrt1bY (or DMY) in the Japanese ricefish (medaka, Oryzias latipes) [26,27]; and DM-W in the frog Xenopus laevis [28]. In several fish species this function is served by gonadal soma-derived factor (GSDF) [29], anti-Müllerian hormone (AMH) [30], anti-Müllerian hormone receptor (AMHR2) [31], or other genes.

In contrast to the variety of upstream sex determinants, genome-wide studies and homology cloning in teleost fishes, amphibians, reptiles and birds have suggested that the downstream components of the network have a conserved function. This has

inspired the paradigm that in sex determination during evolution "masters change, slaves remain" [32–34]. However, it is unclear how far back in the evolutionary history this applies and in particular when and how the vertebrate sex regulation network evolved and whether the relevant genes represent an ancient, conserved mechanism or else they were repeatedly and independently recruited to the process.

The unique opportunity to examine high-quality RNA from the Indonesian coelacanth *Latimeria menadoensis* for transcriptome analysis of testis and liver tissue, and the availability of the whole genome sequence of the African coelacanth *Latimeria chalumnae*, enabled us to gain insights into a "living fossil" that is held to be among the nearest living relatives of tetrapods.

The genes involved in the regulatory network of sexual development described so far come mainly from mammalian studies and can be divided into functional groups as follows: 1) genes required for bipotential gonad development [Wilm's tumour suppressor-1 (WT1), steroidogenic factor-1 (SF-1), and GATA-binding protein 4 (GATA-4)]; 2) genes involved in male sex determination [double sex and mab-3 related transcription factor 1 (DMRT1), SRY-related box 9 (SOX9), dosage-sensitive sex-reversal-adrenal hypoplasia congenitalcritical region of X chromosome, gene 1 (DAX1), fibroblast growth factor 9 (FGF9), and desert hedgehog (DHH)]; 3) genes involved in male sex differentiation [AMH, AMHR2, and androgen receptor (AR)]; 4) genes involved in female sex determination [Wingless-type MMTV integration site family member 4 (WNT4), R-spondin-1 (RSPO-1), catenin  $\beta$ -1 (CTNNB1), forkhead box transcription factor L2 (FOXL2), and follistatin (FST)]; 5) genes involved in female sex differentiation [aromatase (also known as Cyp19A1 or P450arom), oestrogen receptor \alpha  $(ER\alpha)$ , and oestrogen receptor  $\beta$   $(ER\beta)$ ] (Figure 1).

These 22 genes and 11 other genes [DMRT3, DMRT6, GSDF, platelet-derived growth factors (PDGF)  $\alpha$  and  $\beta$  and their receptors (PDGFR $\alpha$ , PDGFR $\beta$ ), 11 $\beta$ -hydroxylase (CYP11B), and 5 $\alpha$ -reductase 1, 2, and 3 (SRD5A1, SRD5A2, SRD5A3)], whose involvement in sex development has been documented [35–47], were sought in the L. chalumnae genome and in the transcriptome of L. menadoensis;

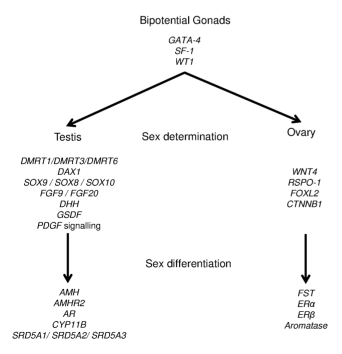


Figure 1. Genes involved in sexual development. doi:10.1371/journal.pone.0056006.g001

subsequently their expression levels were measured in the liver and testis of an adult specimen of *L. menadoensis*.

The coelacanth gene repertoire and expression profiles were much more similar to those of modern fish than to those of tetrapods, although they may also represent an intermediate condition; these data unexpectedly suggest that the major evolutionary changes accompanying the transition to terrestrial life were also involved in gonad development.

#### Methods

The genome of the African coelacanth L. chalumnae has recently been sequenced (project accession PRJNA56111) [48] and is available in the framework of the whole genome shotgun (WGS) sequencing project at http://www.ncbi.nlm.nih.gov and http:// www.ensembl.org. The transcriptome of its Indonesian congener, L. menadoensis, has been described by Pallavicini and colleagues [49] and Canapa and co-workers [50]. Briefly, an adult male specimen of L. menadoensis weighing 27 kg was caught in a shark net near Talise island, Indonesia [51]. Liver and testis were collected immediately after death and preserved in RNAlater (Applied Biosystems, Warrington, UK). A good quality RNA samples, extracted using Trizol Reagent (Ambion/Life Technologies, Carlsbad, CA) following the manufacturer's instructions (RNA integrity number was 7.0 for testis and 6.6 for liver), were used to generate cDNA libraries for transcriptome sequencing on the Illumina Genome Analyzer II platform (Illumina, San Diego, CA, USA). After filtering high-quality reads, removing reads containing primer/adaptor sequences, and trimming read length, the Illumina 100-bp paired-end reads were assembled on a 4-core server (72GB RAM). CLC Genomic Workbench 4.5.1 (CLC Bio, Katrinebjerg, Denmark) and Trinity [52] were used for de novo assembly of short reads. Contigs confirmed and improved by both methods were pooled in a high-quality set.

To identify the coelacanth homologues of the genes involved in sexual development, the corresponding *Xenopus tropicalis*, *Gallus gallus*, *Danio rerio* and *Homo sapiens* sequences were BLASTed on the *L. menadoensis* transcript dataset. The identity of each retrieved putative transcript was confirmed through NCBI BLAST by homology. BLASTx analyses allowed transcript completeness to be established (coding sequences, CDSs).

The *L. menadoensis* sequences were then BLASTed against the WGS dataset of *L. chalumnae*, to identify the genomic scaffolds of the African coelacanth containing them. Species divergence was calculated with PAUP on the matching sequences as p-distance percentage; the Ka/Ks ratio was calculated with KaKs\_calculator [53] using the Nei and Gojobori method [54]. The synonymous distance was calculated using MEGA5 [55] by applying the uncorrected modified Nei and Gojobori method [56] to the concatenated CDSs, aligned with ClustalW2 (http://www.ebi.ac.uk/Tools/msa/clustalw2/; [57]).

The predicted transcripts of *L. chalumnae* were collected from ENSEMBL (http://www.ensembl.org/Latimeria\_chalumnae/Info/Index). The *GSDF* CDS was obtained manually by aligning *L. menadoensis* transcripts to the *L. chalumnae* genome; *FGF9*, not found in the transcriptome and not annotated in ENSEMBL, was obtained manually by BLASTing annotated amino acid sequences of other species to the *L. chalumnae* WGS. *GSDF* and *FGF9* putative transcripts were confirmed by homology through NCBI BLAST.

L. chalumnae and L. menadoensis transcripts were compared by ClustalW2 alignment; a graphical representation of each sequence pair is reported in Figures S1A and S1B.

Gene ontology (GO) terms involved in sex determination and sex differentiation (GO0007530 and GO0007548, respectively)

**Table 1.** Male sex-determining/differentiation gene inventory.

	Transcript in L. menadoensis			Gene location in L. chalumnae				Transcript prediction in L. chalumnae		
Gene	Accession	Length	CDS	Scaffold	N° exons³	Divergence <sup>4</sup>	Ka/Ks	ENSEMBL accession	Length	CDS
АМН	HF562302	1312 <sup>1</sup>	1312 <sup>2</sup>	JH126742	>5	0.046	0.000	ENSLACT00000009808	1689	1689 <sup>2</sup>
	HF562303	1039 <sup>1</sup>	670							
AMHR2	HF562304	693	693 <sup>2</sup>	JH126659	>9	0.289	0.343	ENSLACT00000020587	921	921 <sup>2</sup>
AR	HF562305	2590	2133 <sup>2</sup>	JH126641	>8	0.165	0.000	ENSLACT00000017177	2235	1239 <sup>2</sup>
CYP11B	-	-	-	JH127279	-	-	-	ENSLACT00000015536	1422	474 <sup>2</sup>
DAX1	HF562306	966	786	JH128268	2	0.207	0.309	ENSLACT00000007979	786	786
DHH	HF562307	926	926 <sup>2</sup>	JH126563	2	0.540	0.649	ENSLACT00000021749	1275	1275
DMRT1	HF562308	2244	998 <sup>2</sup>	JH127237	5	0.134	0.000	ENSLACT00000015034	798	798 <sup>2</sup>
DMRT3	-	-	-	JH127237	-	-	-	ENSLACT00000013757	1455	1455
DMRT6	HF562309	3121	957	JH130928	4	0.129	NA	ENSLACT00000003773	798	798 <sup>2</sup>
FGF9	-	-	-	JH128123	-	-	-	Manually identified		
FGF20	HF562310	370	370 <sup>2</sup>	JH127134	3	0.270	NA	ENSLACT00000014939	627	627
GATA-4	HF562311	1655	1200	JH128461	>6	0.064	NA	ENSLACT00000007000	1209	1209
GSDF	HF562312	1258	693	JH127632 <sup>5</sup>	>3	0.826	0.470	-	-	-
PDGFα	HF562313	968	594	JH126909	6	0.000	NA	ENSLACT00000025036	1892	594
PDGFβ	HF562314	664	483 <sup>2</sup>	JH128946	>4	0.000	NA	ENSLACT00000002931	630	630
PDGFRα	HF562315	823	823 <sup>2</sup>	JH128279	>7	0.243	NA	ENSLACT00000010417	3285	3285 <sup>2</sup>
PDGFRβ	HF562316	995 <sup>1</sup>	995 <sup>2</sup>	JH126585	>17	0.195	0.954	ENSLACT00000016664	3312	3312
	HF562317	1055 <sup>1</sup>	1055 <sup>2</sup>							
SF-1	HF562318	1686	1401	JH126572 <sup>6</sup>	7	0.000	NA	ENSLACT00000021404	591	591 <sup>2</sup>
SOX8	HF562319	735	735 <sup>2</sup>	JH126713	>3	0.000	NA	ENSLACT00000019016	1434	1410
SOX9	HF562320	3306	1428	JH126581	3	0.185	0.000	ENSLACT00000021484	1908	1428
SOX10	HF562321	1403	1353 <sup>2</sup>	JH127309	3	0.359	0.199	ENSLACT00000005034	4571	1356
SRD5A1	HF562322	3664	786	JH129903	5	0.164	NA	ENSLACT00000002047	6104	684
SRD5A2	HF562323	2711	765	JH126700	5	0.112	NA	ENSLACT00000025936	2918	765
SRD5A3	HF562324	1244	945	JH127256	5	0.000	NA	ENSLACT00000014423	2733	945
WT1	HF562325	2260	1257	JH126652	9	0.134	0.000	ENSLACT00000018732	1260	1257

<sup>1</sup>Fragmented contig. <sup>2</sup>Partial CDS. <sup>3</sup>Number of exons from the alignment of *L. menadoensis* transcripts to the *L. chalumnae* genome. Where the transcript carries only a partial CDS, the number of exon is partial. <sup>4</sup>Divergence between the two coelacanth sequences calculated as p-distance x100. <sup>5</sup>The *L. chalumnae GSDF* gene is split between scaffold JH127632 and contig AFYH01270444. <sup>6</sup> The *L. chalumnae SF-1* gene is split between scaffold JH126572 and contig AFYH01271535. doi:10.1371/journal.pone.0056006.t001

were selected and *L. menadoensis* orthologues to *D. rerio*, *X. tropicalis*, *G. gallus*, *Canis familiaris*, *Bos taurus*, *Sus scrofa*, *Mus musculus*, *Rattus norvegicus* and *H. sapiens* counterparts counted.

L. menadoensis liver and testis gene expression levels were calculated using the CLC Genomic Workbench 4.5.1 by mapping paired reads from the transcriptome on the assembled transcripts, and given as Fragments Per Kilobase of exon per Million sequenced fragments (FPKM). The lack of some transcripts in the assembled transcriptome may depend on poor gene expression, hence on the limited number of reads, which prevented assembly of a contig. In such cases ENSEMBL gene predictions were used to determine absence or low expression taking into account the predicted transcripts. The FPKM value is therefore still a function of transcript length rather than gene length. The FPKM value was calculated for DMRT3, FOXL2, aromatase, WNT4, and CTP11B on ENSEMBL transcript predictions as well as on the inferred sequence of L. chalumnae FGF9.

Besides genes expected to be involved in sexual development, the expression levels of some house-keeping genes, i.e. *phosphoglycerate kinase (PGK)*, *heat shock protein class B (HSPCB)*, and the

ribosomal proteins *RPS27*, *RPL19*, *RPL11*, *RPL32*, chosen according to Eisenberg and Levanon [58], were also evaluated.

Correct assignment to evolutionarily related gene groups was established by phylogenetic analysis. Sequences of SOXE, FGF9/16/20, and TGF- $\beta$  groups of other vertebrates were retrieved from the NCBI protein database and ENSEMBL. Multiple alignments were performed with ClustalW2 using default parameters. Phylogenetic trees were obtained using Bayesian Inference (BI) and Maximum Parsimony (MP) methods. BI analysis was performed with MrBayes 3.1.2 [59] by applying the amino acid model of Dayhoff et al. [60] to the SOXE and TGF- $\beta$  groups and the one by Jones et al. [61] to the FGF9/16/20 group. Parameters were set to 1,000,000 generations, sampling every 100; burn-in was set at 2,500 and stationarity was defined when the average standard deviation of split frequencies reached a value < 0.009.

MP analyses were performed with PAUP [62] by applying heuristic search with tree bisection-reconnection (TBR) branch swapping and random stepwise additions with 100 replications; 1,000 bootstrap replicates were calculated. Only minimal trees were retained. The outgroup, accession numbers, and constant,

parsimony informative, and parsimony non-informative sites are reported in the legend to each phylogenetic tree.

Conserved syntenic blocks were inferred from ENSEMBL annotation of putative CYP11B (Figure S2), DMRT1, FGF9, FGF16, and FGF20 flanking regions from some sequenced vertebrate genomes. Gene sizes and distances were calculated on the basis of the annotated coordinates of each element. Scaffolds containing FGF9 and flanking genes (EFHA1 and ZDHHC20) conserved in tetrapods were identified by homology through tBLASTn on L. chalumnae WGS data.

#### Results

GO analyses of 'sex determination' and 'sex differentiation' term annotations of the *L. menadoensis* transcriptome were conducted and the results compared to selected vertebrate genomes (Tables S1 and S2); 25 contigs were identified as orthologues of a GO0007530 (sex determination) annotation, and 297 contigs were orthologues of the GO0007544 (sex differentiation) annotation.

In this study we examined 33 genes with substantial evidence of involvement in sex determination and differentiation (Supplementary notes). CDSs were retrieved from the *L. chalumnae* genome and the *L. menadoensis* testis and liver transcriptomes (Tables 1 and 2) and their expression levels assessed. The putative orthology status of closely related genes was confirmed by tree topologies obtained by phylogenetic analysis. Furthermore the instances of microsynteny conservation described in other vertebrates for *DMRT1* [36] and *FGF9/16/20* [63,64] were analysed in the two coelacanths.

To establish whether the sequence information from *L. menadoensis* and *L. chalumnae* could be combined, their genetic distance was determined by comparing the transcripts of the former to the genomic sequences of the latter. The distance, calculated over all matching sequences, ranges between 0% and 0.826%, divergences being due mainly to mutations, insertions or deletions in untranslated regions (UTRs). Point mutations affecting the transcript coding region are predominantly synonymous (Tables 1 and 2). The synonymous distance calculated over the whole gene set was 0.0019 (standard error 0.0005). These findings showed that the data of the two species can be pooled and investigated together.

## Genes in male sexual development

Twenty-five genes involved in male sexual development were analysed in *Latimeria*: 3 genes containing a double sex and mab-3 (DM) domain (DMRT1, DMRT3, and DMRT6); 3 genes belonging to the SOXE subfamily (SOX8, SOX9, and SOX10) of SRY-related HMG box transcription factors; other transcription factors including WT1, DAX1, GATA-4, DHH, SF-1; the signalling molecules PDGF $\alpha$  and  $\beta$ , GSDF, AMH, FGF9, and FGF20; 4 receptors comprising AR, AMHR2, and PDGFR $\alpha$  and  $\beta$ ; and the steroidogenic enzymes SRD5A1, SRD5A2, SRD5A3 and CYP11B (Table 1).

ENSEMBL prediction recovered 23 out of 25 genes in the *L. chalumnae* genome annotation. The two missing sequences were inferred manually from the genome assembly: one, *FGF9*, was identified by comparison with orthologous sequences of other species, and the other, *GSDF*, by aligning an *L. menadoensis* transcript to WGS contigs of *L. chalumnae*. Fifteen of the 23 predicted transcripts of *L. chalumnae* carried complete CDS whereas 8 were partial. The manually inferred *L. chalumnae FGF9* covers the complete CDS, whereas the *L. chalumnae GSDF* homologue is incomplete (about 75% of the CDS).

The testis and liver transcriptomes of *L. menadoensis* contain 22 transcripts. Half of the contigs carried a complete CDS, the other half were partial or fragmented. Transcripts of 3 genes, *FGF9*, *CYP11B*, and *DMRT3*, were not found in liver and testis (Table 1). The male sex development sequences of *L. menadoensis* and *L. chalumnae* are compared in Figure S1A.

Since the expression of 13 male sex development transcripts in testis was<1 FPKM unit, they were considered as not being expressed above background. With the exception of AR, 11 genes (DMRT6, DMRT1, SOX9, SOX10, WT1, GSDF, AMH, SRD5A1, SRD5A3, DHH, and SF-1) were more expressed in testis than in liver, but only 3 (DMRT1, DMRT6, and SOX9) exhibited a differential expression with an FPKM difference>10. In liver 7 genes were expressed above background: SOX9, SRD5A1, AR, DAX1, PDGFa, GATA-4, and SRD5A2 (Figures 2A and 2B).

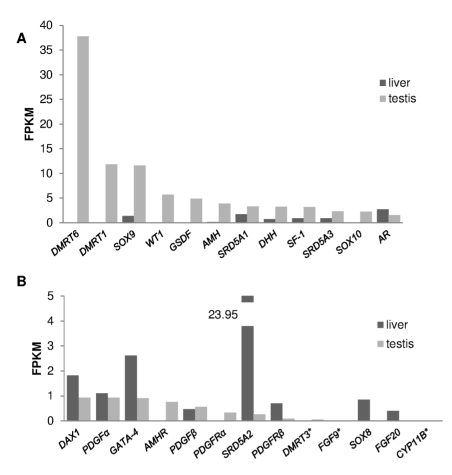
*DMRT6* was the most highly expressed transcript among the 25 male sex development genes analysed (37.79 FPKM in testis, no expression in liver) and one of the 2,000 most abundant transcripts among the 61,000 plus contigs measured in testis.

DMRT1, a major gene in male development, plays a key function in fish [65,66], chickens [67,68], and reptiles [69].

<b>Table 2.</b> Female	sex-determining	d/differentiation	gene inventory.

	L. menadoensis transcript			Gene location in L. chalumnae				Transcript prediction in L. chalumnae		
Gene	Accession	Length	CDS	Scaffold	N° exons³	Divergence <sup>4</sup>	Ka/Ks	ENSEMBL accession	Length	CDS
Aromatase	-	-	-	JH127307	-	-	-	ENSLACT00000010703	1329	1329 <sup>2</sup>
CTNNB1	HF562326	3325	2346	JH127054	15	0.702	0.000	ENSLACT00000017335	3458	2346
ERα	HF562327	2002	1541 <sup>2</sup>	JH129227 <sup>5</sup>	>8	0.352	NA	ENSLACT00000005056	396	396 <sup>2</sup>
ERβ	HF562328	3184	1689	JH126564	9	0.159	0.000	ENSLACT00000019235	2465	1689
FOXL2	-	-	-	JH127245	-	-	-	ENSLACT00000012991	915	915
FST	HF562329	2381	1044	JH127291	6	0.221	0.000	ENSLACT00000014112	2027	1032 <sup>2</sup>
RSPO-1	HF562330	474 <sup>1</sup>	425 <sup>2</sup>	JH126592	>5	0.626	NA	ENSLACT00000019383	747	747 <sup>2</sup>
	HF562331	485 <sup>1</sup>	269 <sup>2</sup>							
WNT4	-	-	-	JH126950	-	-	-	ENSLACT00000017139	1068	1068

<sup>1</sup>Fragmented contig. <sup>2</sup>Partial CDS. <sup>3</sup>Number of exons from the alignment of *L. menadoensis* transcripts to the *L. chalumnae* genome. Where the transcript carries only a partial CDS, the exon number is partial. <sup>4</sup>Divergence between the two coelacanth sequences calculated as p-distance x100. <sup>5</sup>The *ERα* gene in the *L. chalumnae* genome is split among scaffolds JH129227, JH129408, JH129637, and JH133026. doi:10.1371/journal.pone.0056006.t002



**Figure 2. Expression of male development genes.** Expression levels of male sex-determining/differentiation genes in *L. menadoensis* liver and testis transcriptomes. Values are expressed as FPKM (Fragments Per Kilobase of exon per Million sequenced fragments). A) genes highly expressed in testis; B) genes poorly expressed in testis. The expression levels of some housekeeping genes (not represented) were also analysed: *PGK* 96.95 (liver), 342.41 (testis); *RPS27a* 152.59 (liver), 128.43 (testis); *RPL19* 744.01 (liver), 64.89 (testis); *RPL11* 457.35 (liver), 282.59 (testis); *RPL32* 629.83 (liver), 373.75 (testis); *HSPCB* 507.99 (liver), 1213.75 (testis). Threshold value = 1. \* Expression level assessed on *L. chalumnae* orthologue. doi:10.1371/journal.pone.0056006.g002

Alignment of L. menadoensis transcripts to the L. chalumnae genome (Figure 3A) identified 5 exons which exceeded the ENSEMBL predicted transcript by 1,572 bp at the 3' end (Figure 3B). The DM domain is encoded in the first annotated exon. The long 3'UTR harbours a 320-bp region containing a low-copy interspersed repeat.

The size of the *DMRT1* gene in the *L. chalumnae* genome is>152 kb (Figure 3A), close to the 127 kb gene of *H. sapiens* (ENSEMBL annotation) but spanning a much longer range than the 3 kb gene of *Crocodylus palustris* [70], the 45 kb gene of *D. rerio* [71], and 53–58 kb gene of *G. gallus* ([72], ENSEMBL). Moreover the lack of a 5' UTR (Figure 3B), which in other fish is transcribed in the so-called exon 0 [73], both in sequences from the transcriptome and the ENSEMBL prediction, suggests the existence of another exon (which would further extend the genomic locus).

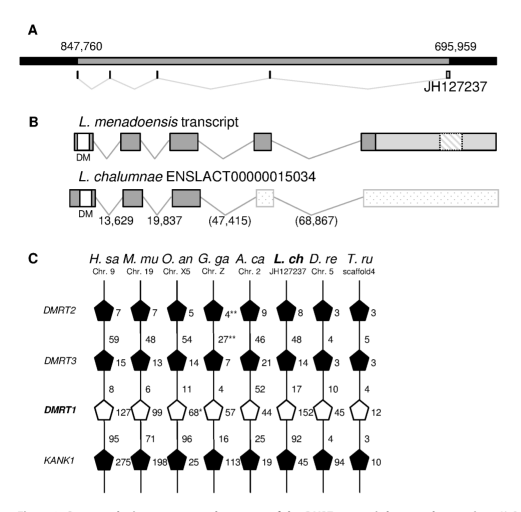
Brunner and colleagues [36] previously reported that the gene order around the *DMRT1* gene, involving two other DM domain genes, *DMRT2* and *DMRT3*, and the gene *KANK1* (*KIAA0172*), was strictly conserved. A similar micro-synteny conservation was also noted in the *L. chalumnae* genome when the genomic scaffold JH127237 (1,057,921 bp), from position 608,000 to 941,000, was compared to other vertebrate chromosomes (Figure 3C). Interestingly, this region is linked to the Z gonosome in *G. gallus* (where *DMRT1* is pivotal in male development) and to the X5 gonosome

in *Omithorhynchus anatinus*, whereas in other species of the actinopterygian and sarcopterygian lineages it is located on an autosome. To date it has been impossible to identify sex chromosomes in the *Latimeria* karyotype [74] or to relate the scaffold containing *DMRT1* to a definite chromosome.

*DMRT1* is the second most abundantly expressed gene in testis (11.84 FPKM units) and among the 10% most abundantly expressed transcripts (Figure 2A) of those analysed.

SOX9 is a transcription factor activating AMH; together with DMRT1 it inhibits WNT4 and FOXL2. In mammals it is activated by another SOX family protein, SRY, whereas in other vertebrates it is mainly regulated by SF-1 and DMRT1; together with SOX8 and SOX10 it belongs to SOX protein subgroup E. Phylogenetic analysis (Figure 4) of SOX E proteins from several vertebrates yielded a tree topology with 3 major clades corresponding to the 3 genes. In the SOX9 and SOX10 clades Latimeria sequences comprise a sister group of tetrapods, while the relationship of the Latimeria SOX8 was not clearly resolved given its phylogenetic position. SOX9 and SOX10 were more strongly expressed in testis than in liver (Figure 2A; FPKM: 11.60 and 1.38 for SOX9, FPKM: 2.25 and 0.04 for SOX10), whereas SOX8 expression was scanty in L. menadoensis liver (Figure 2B).

In mammals FGF9 has an important function in male development, creating a positive feedback cycle with *SOX9* and inhibiting the WNT4 pathway in testis [75]. It has not yet been



**Figure 3. Conserved micro-synteny and structure of the** *DMRT1* **genomic locus and transcripts.** A) Genomic representation of *DMRT1* on scaffold JH127237 of *L. chalumnae*. Grey box corresponds to gene. Small boxes and V signs represent the intron/exon map. B) Transcript representation of *DMRT1* in *L. menadoensis* and *L. chalumnae*. Boxes: exons; V signs: introns; white box: DM domain; light grey box: 3'UTR; dashed box: putative transposable element contained in the 3'UTR. Dotted boxes represent missing exons in the ENSEMBL transcript prediction. C) Microsyntenic conservation of genomic blocks containing the *DMRT1* gene. White pentagons represent *DMRT1* genes. The pentagon tip points to the relative gene orientation. Numbers near the pentagons stand for gene size expressed as kb, numbers on lines represent intergene distance expressed as kb. ENSEMBL data: *H. sa* (*Homo sapiens*), *M. mu* (*Mus musculus*), *O. an* (*Ornithorhynchus anatinus*), *G. ga* (*Gallus gallus*), *A. ca* (*Anolis carolinensis*), *L. ch* (*Latimeria chalumnae*), *D. re* (*Danio rerio*), *T. ru* (*Takifugu rubripes*). *L. chalumnae DMRT1* position was clarified using the *L. menadoensis* transcript, by integrating the *L. chalumnae* ENSLACT00000015034 coordinates. \*In *O. anatinus DMRT1* gene size was defined by comparison with other species. \*Values obtained in *G. gallus* from the annotation of NC\_006127.3 accession. doi:10.1371/journal.pone.0056006.q003

detected in teleosts and seems to be replaced by FGF20b [63,64] in sexual development. Interestingly, we found an FGF9-like sequence in L. chalumnae. To confirm the orthology relationships of the putative Latimeria FGF9, FGF16, and FGF20, sequence comparisons were performed and the conserved synteny arrangements of the flanking regions investigated (Figure 5). In tetrapods the two blocks harbouring FGF9 or FGF20 are characterized by an EFHA and a ZDHHC gene upstream the FGF genes. Extensive gene-deserted regions are found downstream FGF9, 16 and 20. In teleosts (where FGF9 is absent) the other genes forming the microsyntenic cluster are distributed on different chromosomes. In L. chalumnae the FGF9 cluster is split between two scaffolds whose colocalization on the same chromosome cannot as yet be confirmed. However, the proximity of a putative EFHA1 coding fragment upstream the 5' end of FGF9 suggests that the Latimeria FGF9 follows the tetrapod pattern.

Phylogenetic analysis of the FGF9/16/20 group (Figure 6) uncovered three major clades corresponding to the 3 genes. The

exact position of *L. chalumnae* FGF20 sequence is unresolved; like the *X. laevis* orthologue it is paraphyletic to teleosts and tetrapods. As expected, the coelacanth FGF16 sequence is basal to the tetrapods. However, the position of the *Latimeria* FGF9, albeit firmly nested within the FGF9 tetrapod clade, does not reflect its phylogenetic position in the taxonomic group.

Unexpectedly, neither FGF9 nor  $FGF2\theta$  expression was found in L. menadoensis testis.

GSDF, a recently described gene that appears to be critically involved in the development of male teleosts [29,39,40,45], has not been found in tetrapods and no sarcopterygian homologue has yet been described. However, BLAST analysis of teleost GSDF in the L. menadoensis transcript database suggested a putative GSDF gene, whose identity was confirmed by BLASTx analysis. Despite low similarity values (29% identity, 49% positive matching with Oncorhynchus mykiss GSDF NP\_001118051.1, and 28% identity and 50% positive matching with O. latipes GSDF NP\_001171213.1), BI and MP analyses reliably assigned the

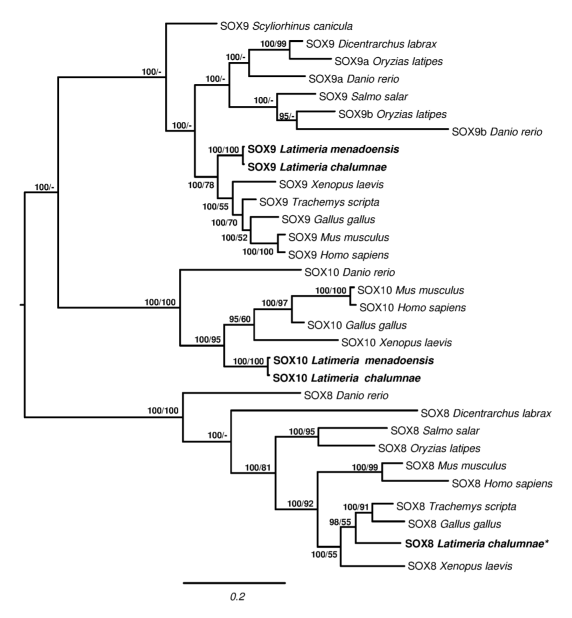
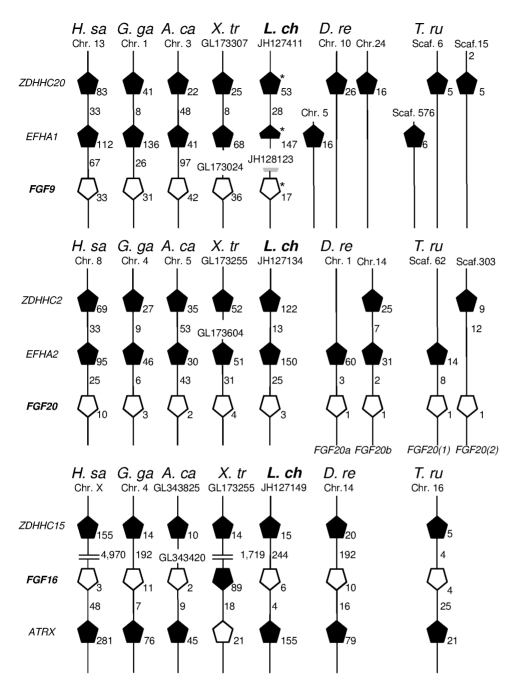


Figure 4. Phylogenetic tree of SOX8, SOX9, and SOX10. Phylogenetic analyses of vertebrate SOXE amino acid sequences. Midpoint rooting. Total characters: 592, constant: 164, parsimony non-informative: 77, parsimony informative: 351. Numbers close to nodes represent posterior probability in Bayesian Inference/bootstrap percentage in Maximum Parsimony. *Danio rerio* (SOX8: AAX73357.1; SOX9a: NP\_571718.1; SOX9b: NP\_571719.1; SOX10: AAK84872.1); *Dicentrarchus labrax* (SOX8: CBN81184.1; SOX9: CBN81190.1); *Gallus gallus* (SOX8: AAF73917.1; SOX9: BAA25296.1; SOX10: AAD38050.2); *Homo sapiens* (SOX8: AAH7797.1; SOX9: CAA86598.1; SOX10: CAG30470.1); *Latimeria chalumnae* (SOX8: ENSLACP00000018883; SOX9: ENSLACP00000021343; SOX10: ENSLACP00000004990); *Latimeria menadoensis* (SOX9, SOX10: this study); *Mus musculus* (SOX8: AAF35837.1; SOX9: AAH23953.1; SOX10: NP\_035567.1); *Oryzias latipes* (SOX8: NP\_001158342.1; SOX9a: AAX62152.1; SOX9b: AAX62151.1); *Salmo salar* (SOX8: ABC24688.1; SOX9: ACN10975.1); *Scyliorhinus canicula* (SOX9: ABY71239.1); *Trachemys scripta* (SOX8: AAP59791.1; SOX9: ACG70782.1; SOX10: ENSLACP00000004990); *Xenopus laevis* (SOX8: AAI69525.1; SOX9: NP\_001084276; SOX10: NP\_001082358.1). \*Only a partial *SOX8* sequence, perfectly matching the ENSEMBL prediction of the *L. chalumnae SOX8* gene, was retrieved in the transcriptome assembly of *L. menadoensis*. doi:10.1371/journal.pone.0056006.g004

sequence to the teleost GSDF clade (Figure 7). Besides GSDF the phylogenetic analysis included two other proteins of the TGF-β family, AMH and inhibin-α, selected for their close relationships to GSDF [39]. A multiple amino acid alignment of the conserved TGF-β domain of the 3 genes disclosed that the *L. menadoensis* GSDF is a sister group of teleost GSDFs, with a posterior probability of 100 in BI analysis and a bootstrap value of 97 in the MP tree (Figure 8). The lack of a glycine, a diagnostic amino acid not found in the GSDF protein [45], in a cysteine knot further confirms the inclusion of the *L. menadoensis* sequence in the GSDF

group, the first homologue to be described in the sarcopterygian lineage.

BLAST analysis of *L. menadoensis GSDF* on the *L. chalumnae* genome allowed identification of a genomic counterpart that was found partly on contig AFYH01270444 and partly on scaffold JH127632, with an intervening gap of 171 bp. The *L. menadoensis GSDF* is strongly expressed in testis but is not expressed in liver (Figure 2A).



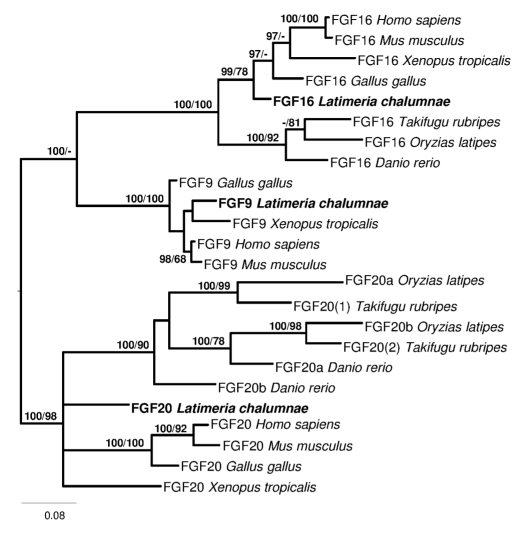
**Figure 5. Analysis of micro-syntenic conservation in** *FGF9, FGF16* **and** *FGF20* **blocks.** Micro-syntenic conservation of genomic regions containing the *FGF9, FGF20* and *FGF16* genes. White pentagons represent *FGF* genes. The pentagon tip points to the relative gene orientation. The grey mark on the top third of the figure indicates a *EFHA1* putative sequence of *Latimeria chalumnae*. Numbers near pentagons stand for gene size expressed as kb, numbers on lines represent intergene distance expressed as kb. ENSEMBL data: *H. sa* (*Homo sapiens*), *G. ga* (*Gallus gallus*), *A. ca* (*Anolis carolinensis*), *X. tr* (*Xenopus tropicalis*), *L. ch* (*Latimeria chalumnae*), *D. re* (*Danio rerio*), *T. ru* (*Takifugu rubripes*). Syntenic blocks for *FGF20* in *L. chalumnae* and *X. tropicalis*, and *FGF16* in *A. carolinensis* are split between two different scaffolds. The *ZDHHC15* genes belonging to the syntenic block of *FGF16* in *H. sapiens* and *X. tropicalis* lie on the same chromosome or scaffold, but are far removed from the genomic locus of *FGF16* and *ATRX*. \*Genes missing in the ENSEMBL prediction.

doi:10.1371/journal.pone.0056006.g005

## Genes in female sexual development

Eight female determining/differentiation genes were examined in the two coelacanths (Table 2): 3 genes belonging to the WNT signalling pathway (WNT4, RSPO-1, and CTNNB1), a transcription factor (FOXL2), two oestrogen receptors (ER $\alpha$  and ER $\beta$ ), a steroidogenic enzyme (aromatase), and an activin-binding protein (FST).

ENSEMBL prediction recovered all 8 gene sequences in the *L. chalumnae* genome. Four transcripts ( $ER\beta$ , CTNNB1, WNT4, and FOXL2) have a complete CDS; only two codons are missing at the 5' end of FST; RSPO-1 and aromatase are partial, whereas  $ER\alpha$ , subdivided into 4 different scaffolds in the WGS, could be only partially identified.



**Figure 6. Phylogenetic tree of FGF9, FGF16, and FGF20.** Phylogenetic analysis of amino acid sequences of the vertebrate FGF9/16/20. Midpoint rooting. Total characters: 237, constant: 91, parsimony non-informative: 35, parsimony informative: 111. Numbers close to nodes represent posterior probability in Bayesian Inference/bootstrap percentage in Maximum Parsimony. *Danio rerio* (FGF16: ENSDART00000061928; FGF20a: NP\_001032180.1; FGF20b: NP\_01034261.1); *Gallus gallus* (FGF9: NP\_989730.1; FGF16; NP\_001038115.1; FGF20: XP\_426335.2); *Homo sapiens* (FGF9: NP\_002001.1; FGF16: NP\_003859.1; FGF20: NP\_062825.1); *Latimeria chalumnae* (FGF9: manually inferred from JH128123; FGF16: ENSLACT00000011509; FGF20: ENSLACT000000114939); *Mus musculus* (FGF9: ADL60500.1; FGF16: BAB16405.1; FGF20: NP\_085113.2); *Oryzias latipes* (FGF16: ENSORLT00000007651; FGF20a: ENSORLT000000012578; FGF20b: ENSORLT000000025767); *Takifugu rubripes* (FGF16: ENSXETT000000021181; FGF20(1): ENSTRUT00000008788; FGF20(2): ENSTRUT000000039390); *Xenopus tropicalis* (FGF9: XP\_002938621.1; FGF16: ENSXETT00000009790; FGF20: NP\_001137399.1). *Latimeria menadoensis* is missing in this analysis because *FGF9* and *FGF20* are poorly or not expressed in the transcriptomes. doi:10.1371/journal.pone.0056006.g006

Analysis of the *L. menadoensis* transcriptome yielded 3 complete CDS sequences (CTNNB1,  $ER\beta$ , and FST) and 2 fragmented CDSs (RSPO-1 and  $ER\alpha$ ), whereas 3 transcripts were missing (FOXL2, WNT4, and aromatase).

The female sex development sequences of L. menadoensis and L. chalumnae are compared in Figure S1B. Their values in L. menadoensis testis and liver are shown in Figure 9. As expected, WNT4, FOXL2 and aromatase — held to be responsible for female development and pathway maintenance — were not expressed in testis. CTNNB1, FST, and  $ER\beta$  were a strongly expressed in liver (56.08, 27.33, 12.93 FPKM, respectively); the expression of FST and CTNNB1 was expected, because their expression is ubiquitous [76]. Finally,  $ER\beta$  liver expression in the L. menadoensis specimen, a male individual, was unexpected.

## Discussion

In this study a set of 33 genes held to be critically involved in sexual development were isolated and characterized for the first time in coelacanths. Comparison of the gene sequences of the two *Latimeria* species confirmed the very slow rate of gene evolution that has recently been documented in *HOX* genes [77], although the latter genes are known to evolve particularly slowly. The 33 genes examined belong to a range of different families, thus providing valuable information.

Interpretation of our data is of course limited by the fact that they come from a single adult individual. However, given the importance of this living fossil in understanding tetrapod and fish evolution, and the exceptional opportunity provided by the availability of high-quality RNA from a specimen of an endangered species, we nonetheless cautiously draw some conclusions.

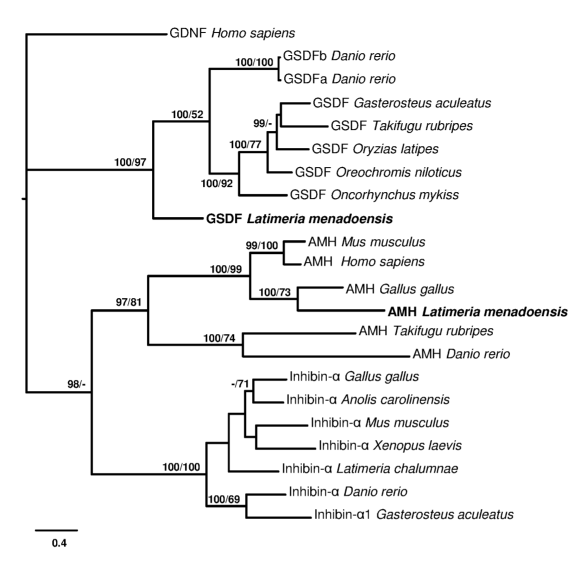


Figure 7. Phylogenetic tree of GSDF, AMH, and inhibin-α. Phylogenetic analysis of amino acid sequences of vertebrate GSDF, inhibin-α and AMH. Total characters: 849, constant: 84, parsimony non-informative: 225, parsimony informative: 540. Outgroup: human glial-derived nerve growth factor (GDNF). Numbers close to nodes represent posterior probabilities in Bayesian Inference/bootstrap percentage in Maximum Parsimony. *Anolis carolinensis* (inhibin-α: ENSACAT00000014331); *Danio rerio* (GSDFa: AEL99890.1; GSDFb: AEL99889.1; AMH: NP\_001007780.1; inhibin-α: ENSDART0000057348); *Gallus gallus* (AMH: NP\_990361.1; inhibin-α: NP\_001026428.1); *Gasterosteus aculeatus* (GSDF: ENSGACT00000021595; inhibin-α: ENSGACT0000018909); *Homo sapiens* (AMH AAC25614.1; GDNF: NP\_000505.1); *Latimeria chalumnae* (inhibin-α: ENSLACT0000017535); *Latimeria menadoensis* (GSDF, AMH this study); *Mus musculus* (AMH: AAI50478.1; inhibin-α: AAH56627.1); *Oreochromis niloticus* (GSDF: BAJ78985.1); *Oryzias latipes* (GSDF: NP\_001171213.1); *Oncorhynchus mykiss* (GSDF: ABF48201.1); *Takifugu rubripes* (GSDF: ENSTRUT00000036269; AMH: ENSTRUT00000045919); *Xenopus laevis* (inhibin-α: NP\_001106349.1). The reliability of *L. menadoensis* CDSs is supported by the same sequence resulting from application of two different assembly procedures. doi:10.1371/journal.pone.0056006.q007

Ka/Ks analysis indicated that no gene in the set studied here is under positive selection in coelacanths. A totally unexpected finding was the very high *DMRT6* expression in testis, which was actually the most abundant male-specific transcript. To date the gene had only been found in amniotes and is not annotated in *Xenopus* and all fish genomes. This phylogenetic pattern could be explained by its being a newly arisen paralogue of the *DMRT* family at the base of amniote vertebrates. Detection of a bona fide *DMRT6* homologue in *Latimeria* points to a much earlier origin of the gene and supports a possible origin from the 1R/2R whole genome duplication events that occurred in ancestral vertebrates [78] and a subsequent, repeated loss in the teleost fish and amphibian lineages and even in basal chordates. Since information on *DMRT6* expression is quite scanty the interpretation of some of these data is merely speculative. In mouse embryo it is expressed in

the developing brain but not in the gonads [43]. In the human microarray database (https://www.genevestigator.com) it is highly expressed exclusively in ovary and testis, whereas studies of mouse organs have disclosed that only erythroblasts and oocytes show elevated expression. Whatever its original function, it is reasonable to assume that *DMRT6* was taken over by other members of the gene family, and that it has ceased to be required in those lineages where it is no longer extant. Its persistence in *Latimeria* may indicate an important function in male (and possibly female) development which, according to current knowledge, was then at least partially conserved in amniotes. Our findings suggest its being a putative novel gene in the gonad regulatory network.

The high *DMRT1* expression found in *Latimeria* testis and its lack of expression in liver is in line with its expression pattern and important role in testis development and in maintenance of the



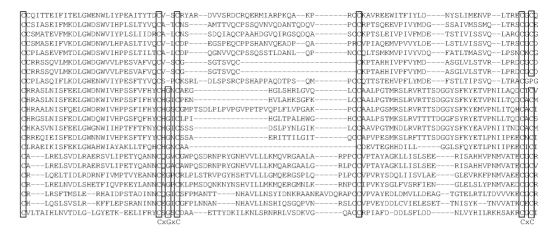
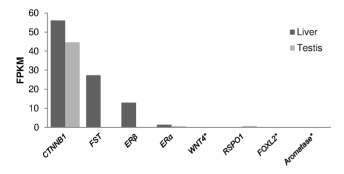


Figure 8. Multiple alignment of the TGF-β domain in GSDF, AMH, and inhibin-α. Conserved amino acids of the cysteine knot are boxed. Anolis carolinensis (inhibin-α: ENSACAT00000014331); Danio rerio (GSDFa: AEL99890.1, GSDFb: AEL99889.1; AMH: NP\_001007780.1; inhibin-α: ENSDART0000057348); Gallus gallus (AMH: NP\_990361.1; inhibin-α: NP\_001026428.1); Gasterosteus aculeatus (GSDF: ENSGACT00000012595; inhibin-α: ENSGACT0000018909); Homo sapiens (AMH: AAC25614.1; GDNF: NP\_000505.1); Latimeria chalumnae (inhibin-α: ENSLACT00000017535); Latimeria menadoensis (this study); Mus musculus (AMH: AAI50478.1; inhibin-α: AAI56627.1); Oreochromis niloticus (GSDF: BAJ78985.1); Oryzias latipes (GSDF: NP\_001171213.1); Oncorhynchus mykiss (GSDF: ABF48201.1); Takifugu rubripes (GSDF: ENSTRUT00000036269; AMH: ENSTRUT00000045919); Xenopus laevis (inhibin-α: NP\_001106349.1). The reliability of L. menadoensis CDSs is supported by the same sequence resulting from application of two different assembly procedures.

doi:10.1371/journal.pone.0056006.g008

male gonad identity documented in vertebrates, from fish to mammals [65,66]. In teleost fish adult testis *DMRT1* is found in germ cells, in somatic cell types or both [65]. Unfortunately, RNA-Seq transcriptome data provide no information on the cell type expressing *DMRT1* in coelacanth testis. In medaka a duplicated version of *DMRT1* on the Y chromosome, designated *dmrt1bY*, is the master male sex determining gene [26,27]. Its major function appears to suppress germ cell proliferation at the critical sex-determining stage in males [79]. In adult testes it is dramatically downregulated [80], and its high expression suggests that only the autosomal *DMRT1* (*dmrt1a* in medaka) may function in mature testis. As in all the teleosts studied so far, a single *DMRT1* copy was found in *L. menadoensis*, suggesting that in coelacanths it may not serve a major role in primary sex determination, but may do so in testis differentiation and adult testis function.



**Figure 9. Expression of female development genes.** Expression of female-determining/differentiation genes in *L. menadoensis* liver and testis transcriptomes. Expression levels are reported as FPKM (Fragments Per Kilobase of exon per Million sequenced fragments). The expression levels of some housekeeping genes were also analyses. The expression levels of some housekeeping genes were also analyses *PLS* 96.95 (liver), 342.41 (testis); *RPS27a* 152.59 (liver), 128.43 (testis); *RPL32* 744.01 (liver) 64.89 (testis); *RPL11* 457.35 (liver), 282.59 (testis); *RPL32* 629.83 (liver), 373.75 (testis); *HSPCB* 507.99 (liver), 1213.75 (testis). Threshold value = 1. \* Expression level assessed on *L. chalumnae* orthologue.

doi:10.1371/journal.pone.0056006.g009

Unlike all the other vertebrates studied [38,43,81–83], *DMRT3* is not expressed in *L. menadoensis* male gonad.

The  $TGF-\beta$  family member GSDF is an important gene in teleost fish gonad development and displays much higher expression in testis than in ovary [39,40]. A duplicate of GSDF may actually have become the master male sex determination gene in Oryzas luzonensis [29]; there is strong evidence that in O. latipes the master male sex determining gene dmrt1bY upregulates GSDF and that upregulation correlates with early testis differentiation [45]. No GSDF homologue has yet been identified outside teleosts. Identification in our study of a bona fide GSDF sequence in Latimeria and its high expression in testis (which also points at its functional conservation) suggests that the gene arose already at the base of the fish lineage, but was later lost during tetrapod evolution. GSDF thus appears to be an ancestral male sexdetermining gene. In the absence of functional data on GSDF function in fish, it remains unclear whether during tetrapod testis development another TGF-\beta family member may have taken over the function it exerted in teleosts and coelacanths.

The high expression in the *L. menadoensis* testis transcriptome of *SOX9*, *SOX10*, *WT1*, *AMH*, *DHH*, *SF-1* and *SDR5A1* and *3* (at least compared to liver), the low expression of *AMHR2*, and the absence of the female factors *FST*, *RSPO-1*, *WNT4*, *FOXL2*, aromatase, and oestrogen receptor transcripts are in line with their expression patterns documented in many vertebrate species and with their proposed function in sexual development.

In particular, the AMH/AMH-receptor system is of interest for *Latimeria* sexual development. In mammals and most likely in all tetrapods AMH induces Müllerian duct regression. Teleosts do not have Müllerian ducts, whereas lungfish and *Latimeria* possess oviducts that are homologous to those of tetrapods [84]. Despite the absence of Müllerian ducts, AMH/AMH-receptor system has an important function in the manifestation of gonadal sex in teleosts, because in medaka AMH signalling is crucial in regulating germ cell proliferation in early gonad differentiation [85]. Given that *AMH* and *AMH-receptor* are expressed in *L. menadoensis* adult testis, the AMH signalling system is present and probably active

there, as in adult teleosts [86,87,88], whereas in mouse testis the system is downregulated before sexual maturity [89].

Several of the 33 genes tested, all of which are involved in sex determination and differentiation in other organisms, were found to be abundantly expressed in the liver transcriptome. The high CTNNB1 levels were expected, due to the ubiquitous function of this signal transducer of the WNT pathway. High FST expression agrees with its expression in all vertebrates and with the finding that it is required for liver cell growth homeostasis in mice [90]. This non-gonadal function of the gene may be conserved in coelacanths. Similarly the transcription factor GATA-4, besides a role in gene regulation in testis development [91], is also involved in the control of a number of liver genes, explaining why transcripts of the coelacanth homologue were found in both tissues. In contrast to coelacanths, where  $5\alpha$ -reductase 2 is highly expressed in liver, the  $5\alpha$ -reductase 1 isoform is differentially regulated by androgens and glucocorticoids in rat liver, resulting in high expression in this tissue, while  $5\alpha$ -reductase 2 is preferentially expressed in gonads [92]. This may indicate lineage-specific subfunctionalization of the isozymes during evolution.

The absence of SOX8 expression in Latimeria testis was unexpected. In other vertebrates, including teleost fish, it is readily detected in this organ, and in mammals it has been assigned an important function in the FGF9/SOX9 interaction loop to maintain Sertoli cell identity by acting redundantly to SOX9 [6,93]. Such back-up function does not seem to be required in Latimeria testis maintenance, or may have been lost in the extant coelacanth lineage. In medaka SOX9 is required for germ cell proliferation and survival, but not for testis determination [94]. Together with the other L. menadoensis findings this may indicate that the sex-determining function was acquired later in tetrapod lineage, after the split of teleost and coelacanth lineages.

Intriguing data were found for FGF9 and 20, which together with FGF16 constitute a gene subfamily of paracrine FGFs. The critical role of FGF9 in mammalian testis development is well established and appears to be conserved in all tetrapods. On the other hand, the gene is not found in any teleost genome ([63,64], ENSEMBL), unlike FGF16 and 20 (the latter being duplicated due to the teleost genome duplication). In the amphioxus an FGF gene is basal to the three FGFs in tetrapods [95]. FGF9 could thus be a later duplicate of either FGF16 or 20, and its role in testis development could be interpreted as an innovation arising in tetrapods. However, identification of FGF9 in Latimeria supports an origin during the 1R/2R whole genome duplication events that took place in ancestral chordates and its loss in the lineage leading to teleosts. In the teleost Oreochromis niloticus (Tilapia) FGF20b and FGF16 are both expressed in ovary, whereas only FGF16 is (poorly) expressed in testis [64]. Together with the complete absence of FGF9, FGF20 and FGF16 expression in L. menadoensis liver and testis, this indicates that the function of FGF signalling in testis, in particular the central role of FGF9, was acquired later in tetrapod evolution.

Surprisingly, the  $ER\beta$  gene was expressed in the liver of the male coelacanth. A previous study of the same individual had disclosed expression of the vitellogenin genes vtgABI, II and III [50]. Vitellogenins are yolk proteins physiologically expressed in the liver of reproductive females upon induction by oestrogens. Thus expression of vitellogenins and oestrogen receptor indicates the presence of oestrogens in this male specimen. They could derive from environment pollutants, as reported in a number of specimens from polluted waters; however this individual lived in Bunaken Marine Park in submarine caves at a depth of 100 to 200 m, i.e. in a relatively protected environment. Alternatively,  $ER\beta$  expression could be the result of a pathological condition, of a

hormone imbalance due to ageing, or of a physiological feature of coelacanths.

## **Conclusions**

Analysis of the coelacanth testis transcriptome, reported here for the first time, disclosed important new information on which genes involved in sexual development and testis differentiation in other organisms are present and expressed in this living fossil and on the evolution of this process in vertebrates. Interestingly, some genes that are generally considered critical for testis maintenance in all vertebrates, like SOX8 or a fibroblast growth factor gene from the FGF9/16/20 subfamily, do not play this role in Latimeria. This finding and the high GSDF expression found in the coelacanths make their transcript profile more similar to that of modern fish. In summary, the coelacanth testis transcriptome is expected to contribute further important information to reconstruct the ancestral tetrapod situation and indicates that evolutionary innovations for sexual development occurred already during the transition from water to land.

# **Supporting Information**

Figure S1 Sequence pair comparison of male sex development genes. Sequence pair comparison of male sexdetermining/differentiation transcripts from the L. menadoensis transcriptome and L. chalumnae ENSEMBL predictions. Boxes represent CDSs. Lines represent UTRs. Dashed boxes represent a missing part in the CDS. Green lines/boxes represent an inaccurate gene prediction or a mismatch between L. chalumnae and L. menadoensis sequences. Scale dimension are preserved. B) Sequence pair comparison of female sex development genes. Sequence pair comparison of female sex-determining/differentiation transcripts from the L. menadoensis transcriptome and L. chalumnae ENSEMBL predictions. Boxes represent CDSs. Lines represent UTRs. Dashed boxes represent a missing part in the CDS. Green lines/boxes represent an inaccurate gene prediction or a mismatch between L. chalumnae and L. menadoensis sequences. Scale dimension are preserved. (PDF)

Figure S2 Micro-syntenic conservation of CYP11B. Micro-syntenic conservation of genomic regions containing CYP11B genes. Black pentagons represent CYP11B genes. The pentagon tip points to the relative gene orientation. ENSEMBL data: H. sa (Homo sapiens), M. mu (Mus musculus), B. ta (Bos taurus), L. ch (Latimeria chalumnae), D. re (Danio rerio), T. ru (Takifugu rubripes). (PDF)

Table S1 Gene Ontology analysis of the "sex determination" term.

(PDF)

Table S2 Gene Ontology analysis of the "sex differentiation" term.

(PDF)

# Acknowledgments

FM, CA, BMA, BM, and OE are affiliated to Istituto Nazionale Biosistemi e Biostrutture (INBB).

### **Author Contributions**

Conceived and designed the experiments: MF AC EO MS. Performed the experiments: MF AC MB MAB AP MG GDM EO. Analyzed the data: MF AC MB MAB TC AP MG GDM EO MS. Contributed reagents/

materials/analysis tools: MF AC MB MAB FB AMF AP MG GDM DMM GS EO MS. Wrote the paper: MF AC MB MAB EO MS.

#### References

- Hayes TB (1998) Sex determination and primary sex differentiation in amphibian: genetic and developmental mechanisms. J Exp Zool 281: 373–399.
- Graves JAM, Peichel CL (2010) Are homologies in vertebrate sex determination due to shared ancestry or to limited options? Genome Biol 11: 205.
- McClelland K, Bowles J, Koopman P (2012) Male sex determination: insights into molecular mechanisms. Asian J Androl 14: 164–171.
- Uhlenhaut NH, Jakob S, Anlag K, Eisenberger T, Sekido R, et al. (2009) Somatic sex reprogramming of adult ovaries to testes by FOXL2 ablation. Cell 139: 1130–1142.
- Matson CK, Murphy MW, Sarver AL, Griswold MD, Bardwell VJ, et al. (2011) *DMRT1* prevents female reprogramming in the postnatal mammalian testis. Nature 476: 101–104.
- Herpin A, Schartl M (2011) Sex determination: switch and suppress. Curr Biol 21: R656–9.
- Clinton M, Zhao D, Nandi S, McBride D (2012) Evidence for avian cell autonomous sex identity (CASI) and implications for the sex-determination process? Chromosome Res 20: 177–190.
- Zarkower D (2006) Somatic sex determination (February 10, 2006), in WormBook. Edited by The C. elegans Research Community.doi/10.1895/ wormbook.1.84.1, http://www.wormbook.org.
- Abinawanto, Shimada K, Yoshida K, Saito N (1996) Effects of aromatase inhibitor on sex differentiation and levels of P450<sub>17 alpha</sub> and P450<sub>arom</sub> messenger ribonucleic acid of gonads in chicken embryos. Gen Comp Endocrinol 102: 241–246.
- Akazome Y, Mori T (1999) Evidence of sex reversal in the gonads of chicken embryos after oestrogen treatment as detected by expression of lutropin receptor. J Reprod Fertil 115: 9–14.
- Baker PJ, Moore HD, Burgess AM, Mittwoch U (1993) Gonadal sex differentiation in embryos and neonates of the marsupial, Monodelphis domestica: arrest of testis development in postterm embryos. J Anat 182: 267–273.
- 12. Coveney D, Shaw G, Renfree MB (2001) Estrogen-induced gonadal sex reversal in the tammar wallaby. Biol Reprod 65: 613–621.
- Fadem BH (2000) Perinatal exposure to estradiol masculinizes aspects of sexually dimorphic behavior and morphology in gray short-tailed opossums (Monodelphis domestica). Horm Behav 37: 79–85.
- Guiguen Y, Baroiller JF, Ricordel MJ, Iseki K, Mcmeel OM, et al. (1999) Involvement of estrogens in the process of sex differentiation in two fish species: the rainbow trout (*Oncorhynchus mykiss*) and a tilapia (*Oreochromis niloticus*). Mol Reprod Dev 54: 154–162.
- Kobayashi T, Kajiura-Kobayashi H, Nagahama Y (2003) Induction of XY sex reversal by estrogen involves altered gene expression in a teleost, tilapia. Cytogenet Genome Res 101: 289–294.
- Mackenzie CA, Berrill M, Metcalfe C, Pauli BD (2003) Gonadal differentiation in frogs exposed to estrogenic and antiestrogenic compounds. Environ Toxicol Chem 22: 2466–2475.
- Mittwoch U (1998) Phenotypic manifestations during the development of the dominant and default gonads in mammals and birds. J Exp Zool 281: 466–471.
- Pieau C, Dorizzi M (2004) Oestrogens and temperature-dependent sex determination in reptiles: all is in the gonads. J Endocrinol 181: 367–377.
- Renfree MB, Coveney D, Shaw G (2001) The influence of estrogen on the developing male marsupial. Reprod Fertil Dev 13: 231–240.
- Shaw G, Renfree MB, Short RV, WS O (1988) Experimental manipulation of sexual differentiation in wallaby pouch young treated with exogenous steroids. Development 104: 689–701.
- Ramsey M, Crews D (2009) Steroid signaling and temperature-dependent sex determination. Reviewing the evidence for early action of estrogen during ovarian determination in turtles. Semin Cell Dev Biol 20: 283–292.
- Schartl M (2004) Sex chromosome evolution in non-mammalian vertebrates. Curr Opin Genet Dev 14: 634

  –641.
- 23. Uller T, Helanterä H (2011) From the origin of sex-determining factors to the evolution of sex-determining systems. Q Rev Biol 86: 163–180.
- 24. Biason-Lauber A (2010) Control of sex development. Best Pract Res Clin Endocrinol Metab 24: 163–186.
- Kashimada K, Koopman P (2010) Sty: the master switch in mammalian sex determination. Development 137: 3921–3930.
- Matsuda M, Nagahama Y, Shinomiya A, Sato T, Matsuda C, et al. (2002) DMY
  is a Y-specific DM-domain gene required for male development in the medaka
  fish. Nature 417: 559–563.
- Nanda I, Kondo M, Hornung U, Asakawa S, Winkler C, et al. (2002) A duplicated copy of *DMRT1* in the sex-determining region of the Y chromosome of the medaka, *Oryzias latipes*. Proc Natl Acad Sci USA 99: 11778–11783.
- Yoshimoto S, Okada E, Umemoto H, Tamura K, Uno Y, et al. (2008) A W-linked DM-domain gene, DM-W, participates in primary ovary development in Xenopus laevis. Proc Natl Acad Sci USA 105: 2469–2474.
- Myosho T, Otake H, Masuyama H, Matsuda M, Kuroki Y, et al. (2012) Tracing the emergence of a novel sex-determining gene in medaka, *Oryzias luzonensis*. Genetics 191: 163–170.

- Hattori RS, Murai Y, Oura M, Masuda S, Majhi SK, et al. (2012) A Y-linked anti-Müllerian hormone duplication takes over a critical role in sex determination. Proc Natl Acad Sci USA 109: 2955–2959.
- Kamiya T, Kai W, Tasumi S, Oka A, Matsunaga T, et al. (2012) A Trans-Species Missense SNP in Amhr2 Is Associated with Sex Determination in the Tiger Pufferfish, Takifugu rubripes (Fugu). PLoS Genet 8: e1002798.
- Angelopoulou R, Lavranos G, Manolakou P (2012) Sex determination strategies in 2012: towards a common regulatory model? Reprod Biol Endocrinol 10: 13.
- Graves JAM (1995) The evolution of mammalian sex chromosomes and the origin of sex-determining genes. Philos Trans R Soc Lond B Biol Sci 350: 305– 311.
- Graves JAM (2008) Weird animal genomes and the evolution of vertebrate sex and sex chromosomes. Annu Rev Genet 42: 565–586.
- Borg B (1994) Androgens in teleost fishes. Comp Biochem Physiol Part C 109: 219–245.
- Brunner B, Hornung U, Shan Z, Nanda I, Kondo M, et al. (2001) Genomic organization and expression of the doublesex-related gene cluster in vertebrates and detection of putative regulatory regions for *DMRT1*. Genomics 77: 8–17.
- Chardard D, Kuntz S, Chesnel A, Flament S (2003) Effects of androgens on sex differentiation of the urodele *Pleurodeles waltl*. J Exp Zoolog A Comp Exp Biol 296: 46–55.
- El-Mogharbel N, Wakefield M, Deakin JE, Tsend-Ayush E, Grützner F, et al. (2007) DMRT gene cluster analysis in the platypus: new insights into genomic organization and regulatory regions. Genomics 89: 10–21.
- Gautier A, Le Gac F, Lareyre JJ (2011) The gsdf gene locus harbors evolutionary conserved and clustered genes preferentially expressed in fish previtellogenic oocytes. Gene 472: 7–17.
- Gautier A, Sohm F, Joly JS, Le Gac F, Lareyre JJ (2011) The proximal promoter region of the zebrafish gsdf gene is sufficient to mimic the spatio-temporal expression pattern of the endogenous gene in Sertoli and granulosa cells. Biol Reprod 85: 1240–1251.
- Gnessi L, Emidi A, Jannini EA, Carosa E, Maroders M, et al. (1995) Testicular development involves the spatiotemporal control of PDGFs and PDGF receptors gene expression and action. J Cell Biol 131: 1105–1121.
- Gnessi L, Basciani S, Mariani S, Arizzi M, Spera G, et al. (2000) Leydig cell loss and spermatogenic arrest in platelet-derived growth factor (PDGF)-A-deficient mice. J Cell Biol 149: 1019–1025.
- Kim Š, Kettlewell JR, Anderson RC, Bardwell VJ, Zarkower D (2003) Sexually dimorphic expression of multiple doublesex-related genes in the embryonic mouse gonad. Gene Expr Patterns 3: 77–82.
- O'Donnell L, Stanton PG, Wreford NG, Robertson DM, McLachlan RI (1996) Inhibition of 5-alpha-reductase activity impairs the testosterone-dependent restoration of spermiogenesis in adult rats. Endocrinology 137: 2703–2710.
- 45. Shibata Y, Paul-Prasanth B, Suzuki A, Usami T, Nakamoto M, et al. (2010) Expression of gonadal soma derived factor (GSDF) is spatially and temporally correlated with early testicular differentiation in medaka. Gene Expr Patterns 10: 283–289
- Smith CA, McClive PJ, Hudson Q, Sinclair AH (2005) Male-specific cell migration into the developing gonad is a conserved process involving PDGF signaling. Dev Biol 284: 337–350.
- Zaccanti F, Petrini S, Rubatta ML, Stagni AM, Giorgi PP (1994) Accelerated female differentiation of the gonad by inhibition of steroidogenesis in amphibia. Comp Biochem Physiol A 107: 171–179.
- Amemiya CT, Alföldi J, Lee AP, Fan S, Brinkmann H, et al. (2013) The African coelacanth genome provides insights into the tetrapod evolution. Nature (in press).
- Pallavicini A, Canapa A, Barucca M, Alföldi J, Biscotti MA, et al. (2013) Analysis
  of the transcriptome of the Indonesian coelacanth *Latimeria menadoensis*.
  (submitted).
- Canapa A, Olmo E, Forconi M, Pallavicini A, Makapedua MD, et al. (2012) Composition and phylogenetic analysis of vitellogenin coding sequences in the Indonesian coelacanth *Latimeria menadoensis*. J Exp Zool B Mol Dev Evol 318: 404–416.
- Makapedua DM, Barucca M, Forconi M, Antonucci N, Bizzaro D, et al. (2011)
   Genome size, GC percentage and 5 mC level in the Indonesian coelacanth Latimeria menadoensis. Mar Genomics 4: 167–172.
- Grabherr MG, Haas BJ, Yassour M, Levin JZ, Thompson DA, et al. (2011) Fulllength transcriptome assembly from RNA-Seq data without a reference genome. Nat Biotechnol 29: 644–652.
- Zhang Z, Li J, Zhao XQ, Wang J, Wong GK, et al. (2006) KaKs Calculator: Calculating Ka and Ks through model selection and model averaging. Geno Prot Bioinfo 4: 259–263.
- Nei M, Gojobori T (1986): Simple methods for estimating the numbers of synonymous and nonsynonymous nucleotide substitutions. Mol Biol Evol 3: 418–426.
- Tamura K, Peterson D, Peterson N, Stecher G, Nei M, et al. (2011) MEGA5: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. Mol Biol Evol 28: 2731–2739.

- Zhang J, Rosenberg HF, Nei M (1998) Positive Darwinian selection after gene duplication in primate ribonuclease genes. Proc Natl Acad Sci USA 95: 3708– 3713.
- Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, et al. (2007)
   ClustalW and ClustalX version 2 (2007). Bioinformatics 23: 2947–2948.
- Eisenberg E, Levanon EY (2003) Human housekeeping genes are compact. Trends Genet 19: 362–365.
- Huelsenbeck JP, Ronquist F, Nielsen R, Bollback JP (2001) Bayesian inference of phylogeny and its impact on evolutionary biology. Science 294: 2310–2314.
- Dayhoff M, Schwartz R, Orcutt B (1978) A model of evolutionary change in protein. Atlas Protein Seq Struct 5: 345–352.
- Jones DT, Taylor WR, Thornton JM (1992) The rapid generation of mutation data matrices from protein sequences. CABIOS 8: 275–282.
- Swofford DL (2002) PAUP\*. Phylogenetic analysis using parsimony (\* and other methods) version 4. Sunderland, MA: Sinauer Associates.
- 63. Itoh N, Konishi M (2007) The zebrafish FGF family. Zebrafish 4: 179-186.
- Sun YL, Zeng S, Ye K, Yang C, Li MH, et al. (2012) Involvement of FGF9/16/ 20 subfamily in female germ cell development of the Nile tilapia, *Oreochromis niloticus*. Fish Physiol Biochem 38: 1427–1439.
- Herpin A, Schartl M (2011) Dmtl genes at the crossroads: a widespread and central class of sexual development factors in fish, FEBS I 278: 1010–1019.
- Matson CK, Zarkower D (2012) Sex and the singular DM domain: insights into sexual regulation, evolution and plasticity. Nat Rev Genet 13: 163–174.
- Chue J, Smith CA (2011) Sex determination and sexual differentiation in the avian model. FEBS J 278, 1027–1034.
- Raymond CS, Shamu CE, Shen MM, Seifert KJ, Hirsch B, et al. (1998) Evidence for evolutionary conservation of sex-determining genes. Nature 391: 691–695.
- Rhen T, Schroeder T (2010) Molecular mechanisms of sex determination in reptiles. Sex Dev 4: 16–28.
- Anand A, Patel M, Lalremruata A, Singh AP, Agrawal R, et al. (2008) Multiple alternative splicing of *DMRT1* during gonadogenesis in Indian mugger, a species exhibiting temperature-dependent sex determination. Gene 425: 56–63.
- Guo Y, Cheng H, Huang X, Gao S, Yu H, et al. (2005) Gene structure, multiple alternative splicing, and expression in gonads of zebrafish *DMRT1*. Biochem Biophys Res Commun 330: 950–957.
- Zhao Y, Lu H, Yu H, Cheng H, Zhou R (2007) Multiple alternative splicing in gonads of chicken DMRT1. Dev Genes Evol 217: 119–126.
- Shinomiya A, Otake H, Togashi K, Hamaguchi S, Sakaizumi M (2004) Field survey of sex-reversals in the medaka, *Oryzias latipes*: genotypic sexing of wild populations. Zoolog Sci 21: 613–619.
- Bogart JP, Balon EK, Bruton MN (1994) The chromosomes of the living coelacanth and their remarkable similarity to those of one of the most ancient frogs. J Hered 85: 322–325.
- Kim Y, Kobayashi A, Sekido R, DiNapoli L, Brennan J, et al. (2006) Fgf9 and Wnt4 Act as Antagonistic Signals to Regulate Mammalian Sex Determination. PLoS Biol 4: e187.
- Nakatani M, Takehara Y, Sugino H, Matsumoto M, Hashimoto O, et al. (2008)
   Transgenic expression of a myostatin inhibitor derived from follistatin increases skeletal muscle mass and ameliorates dystrophic pathology in mdx mice.

   FASEB J 22: 477–487.

- Higasa K, Nikaido M, Saito TL, Yoshimura J, Suzuki Y, et al. (2012) Extremely slow rate of evolution in the HOX cluster revealed by comparison between Tanzanian and Indonesian coelacanths. Gene 505: 324–332.
- Johnsen H, Andersen Ø (2012) Sex dimorphic expression of five dmrt genes identified in the Atlantic cod genome. The fish-specific dmrt2b diverged from dmrt2a before the fish whole-genome duplication. Gene 505: 221–232.
- Herpin A, Schindler D, Kraiss A, Hornung U, Winkler C, et al. (2007) Inhibition of primordial germ cell proliferation by the medaka male determining gene DmrtIbY. BMC Dev Biol 7: 99.
- Hornung U, Herpin A, Schartl M (2007) Expression of the male determining gene dmrtlbY and its autosomal coorthologue dmrtla in medaka. Sex Dev 1: 197–206
- 81. Bratus A, Slota E (2009) Comparative cytogenetic and molecular studies of *DM* domain genes in pig and cattle. Cytogenet Genome Res 126: 180–185.
- Smith CA, Hurley TM, McClive PJ, Sinclair AH (2002) Restricted expression of DMRT3 in chicken and mouse embryos. Gene Expr Patterns 2: 69–72.
- Winkler C, Hornung U, Kondo M, Neuner C, Duschl J, et al. (2004)
   Developmentally regulated and non-sex-specific expression of autosomal dmrt genes in embryos of the Medaka fish (Oryzias latipes). Mech Dev 121: 997–1005.
- 84. Kapoor BG, Khanna B (2004) Ichthyology Handbook. Berlin: Springer 487.
- Nakamura S, Watakabe I, Nishimura T, Picard JY, Toyoda A, et al. (2012) Hyperproliferation of mitotically active germ cells due to defective anti-Müllerian hormone signaling mediates sex reversal in medaka. Development 139: 2283–2287.
- Klüver N, Pfennig F, Pala I, Storch K, Schlieder M, et al. (2007) Differential expression of anti-Müllerian hormone (amh) and anti-Müllerian hormone receptor type II (amhrII) in the teleost medaka. Dev Dvn 236: 271–281.
- Pala I, Klüver N, Thorsteinsdóttir S, Schartl M, Coelho MM (2008) Expression pattern of anti-Müllerian hormone (amh) in the hybrid fish complex of Squalius albumoides. Gene 410: 249–258.
- Halm S, Rocha A, Miura T, Prat F, Zanuy S (2006) Anti-Müllerian hormone (AMH/AMH) in the European sea bass: its gene structure, regulatory elements, and the expression of alternatively-spliced isoforms. Gene 388: 148–158.
- Beau C, Rauch M, Joulin V, Jégou B, Guerrier D (2000) GATA-1 is a potential repressor of anti-Müllerian hormone expression during the establishment of puberty in the mouse. Mol Reprod Dev 56: 124–138.
- Ooe H, Chen Q, Kon J, Sasaki K, Miyoshi H, et al. (2012) Proliferation of rat small hepatocytes requires follistatin expression. J Cell Physiol 227: 2363–2370.
- Rey R, Lukas-Croisier C, Lasala C, Bedecarrás P (2003) AMH/MIS: what we know already about the gene, the protein and its regulation. Mol Cell Endocrinol 211: 21–31.
- El-Awady MK, El-Garf W, El-Houssieny L (2004) Steroid 5alpha reductase mRNA type 1 is differentially regulated by androgens and glucocorticoids in the rat liver. Endocr J 51: 37–46.
- Barrionuevo F, Georg I, Scherthan H, Lécureuil C, Guillou F, et al. (2009)
   Testis cord differentiation after the sex determination stage is independent of
   Sox9 but fails in the combined absence of Sox9 and Sox8. Dev Biol 327: 301–312.
- Nakamura S, Watakabe I, Nishimura T, Toyoda A, Taniguchi Y, et al. (2012)
   Analysis of Medaka sox9 Orthologue Reveals a Conserved Role in Germ Cell Maintenance. PLoS ONE 7: e29982.
- Bertrand S, Camasses A, Somorjai I, Belgacem MR, Chabrol O, et al. (2011)
   Amphioxus FGF signaling predicts the acquisition of vertebrate morphological traits. Proc Natl Acad Sci USA 108: 9160–9165.