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**Integrative perspectives on bivalve sex
determination: A comparative and evolutionary
analysis across phylogeny**

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List of abbreviations

AASD	amino acid sequence divergence
CMS	cytoplasmatic male sterility
CUE	coupling of ubiquitin conjugation to endoplasmic reticulum degradation [domain]
DEAD/DEAH-box	Asp-Glu-Ala-Asp/Asp-Glu-Ala-His box
DGE	differential gene expression
dpf	days post fertilization
DM	<i>dsx</i> and <i>mab-3</i> [domain]
DMA	DM-associated [domain]
Dmrt	<i>dsx</i> and <i>mab-3</i> related transcription factor
Dmrt-1L	<i>Dmrt 1-like</i>
Dm-W	<i>dsx</i> and <i>mab-3</i> related gene <i>W</i>
Dmy	<i>dsx</i> and <i>mab-3</i> related gene <i>Y</i>
DSFG	Dmrt, Sox, and Fox gene
dsx	<i>doublesex</i>
DUI	doubly uniparental inheritance
ESD	environmental sex determination
FASW	filtered artificial sea water
FHA	forkhead-associated [domain]
Fox	forkhead box

GC	germ cell
GO	gene ontology
GRN	gene regulatory network
GSD	genetic sex determination
HCR	hybridization chain reaction
HeSC	heteromorphic sex chromosome
HMG	high mobility group [box domain]
HMM	hidden Markov model
HoSC	homomorphic sex chromosome
hpf	hours post fertilization
mab-3	<i>male abnormal-3</i>
MCL	Markov clustering algorithm
ML	maximum likelihood
mRNA-ISH	mRNA <i>in-situ</i> hybridization
Mya	million years ago
ORF	open reading frame
PBS	1× phosphate-buffered saline
PBS-Tw	1× PBS with 0.1 % Tween 20
PCA	principal component analyses
PFA	paraformaldehyde
PGC	primordial germ cell
pPGC	presumptive primordial germ cell
PSC	pluripotent stem cell
qRT-PCR	quantitative real-time polymerase chain reaction

RNAi RNA interference

RT room temperature

SC sex chromosome

SCO single-copy orthogroup

SD sex determination

SDf sex differentiation

SDG sex-determining gene

Sox *Sry*-related HMG-box

SRG sex-determination related gene

Sry *Sex-determining region of chromosome Y*

SSC-Tw 5× saline-sodium citrate with 0.1 % Tween 20

Sxl *Sex-lethal*

TBS 1× Tris-buffered saline

TBS-Tx 1× TBS with Triton X-100

tra *transformer*

TSD temperature-dependent [environmental] sex determination

Chapter 1

Introduction

1.1 The diversity of sexual processes in animals

The process of sex determination (SD) has been traditionally associated with the very first steps of gonad differentiation, where an initial trigger (or master switch) activates the molecular pathway that establishes the sexual identity of an organism. According to this view, two alternative types of SD have been traditionally recognised: the genetic sex determination (GSD) and the environmental sex determination (ESD), depending on whether the very first cues are of genetic or environmental origin. All the downstream events of gonad and morphological sex-specific development (i.e., after SD) have been instead appointed as sex differentiation (SDf), which consists of the entire set of morphogenetic, molecular, and physiological events leading to the full maturation of testes or ovaries and secondary sexual characters (**Uller and Helanterä, 2011; Bear and Monteiro, 2013; Beukeboom and Perrin, 2014**). GSD is found in many classical model systems—such as the mouse *Mus musculus*, the fruit fly *Drosophila melanogaster*, and the nematode *Caenorhabditis elegans*, but is by far the most prevalent SD system in animals, as it occurs in the majority of vertebrates and arthropods (**Bachtrog et al., 2014; Beukeboom and Perrin, 2014**). It encompasses a variety of sex-determining cues, ranging from single master genes (e.g., *Sex-determining region of chromosome Y (Sry)* in eutherians), to polygenic networks (as in the zebrafish *Danio rerio*) and chromosome countings (as in *D. melanogaster* and *C. elegans*). Conversely, ESD is more uncommon and is found mainly in reptiles, fishes, insects, crustaceans, annelids, nematodes, and rotifers (reviewed in **Korpelainen, 1990; Bachtrog et al., 2014**). It relies on several initial stimuli of environmental origin, such as light, food availability, and population density, though the most common process is the temperature-dependent SD

(TSD; **Bachtrög et al., 2014**; **Beukeboom and Perrin, 2014**). Currently, the molecular basis by which an environmental signal is transduced into the canalization of the male or female developmental pathway is unknown (**Bachtrög et al., 2014**; **Capel, 2017**).

Lately, a growing number of studies have challenged the traditionally binary views of both GSD/ESD and SD/SDf (**Bear and Monteiro, 2013**; **Bachtrög et al., 2014**; **Beukeboom and Perrin, 2014**; **Todd et al., 2016**; **Capel, 2017**). On the one hand, the characterisation of SD in new species has shown that GSD and ESD represent the ends of a continuum of mixed conditions, rather than two mutually exclusive phenomena. For example, in the red-eared slider turtle *Trachemys scripta*, a species with TSD, it has been shown that pairs of gonads cultured separately at the same pivotal temperature (i.e., the temperature producing 50 % of males and 50 % of females in offspring) tend to predominantly differentiate into one sex. Therefore, an underlying genetic/epigenetic mechanism controlling SD should exist when temperature effect is absent (**Mork et al., 2014**). In the Australian bearded-dragon *Pogona vitticeps*, some ZZ males were caught to sex-revert to fertile females in the wild after incubation at high temperatures, constituting a natural case of transition from GSD to ESD (in the form of TSD; **Holleley et al., 2015**). On the other hand, instead, the high evolutionary dynamics and the variable expression patterns of the genes involved in the processes of gonad commitment and development make the distinction between SD and SDf increasingly blurred (**Bear and Monteiro, 2013**; **Bachtrög et al., 2014**; **Beukeboom and Perrin, 2014**; **Todd et al., 2016**; **Capel, 2017**). In fact, considering that the SD of an organism may follow different routes, such as being triggered only in presumptive gonads—which then establish the sexual identity of the rest of the organism through hormone signalling (gonadal SD; a process traditionally associated to mammals), or occurring independently in every cell of the developing embryo (cell-autonomous SD; a process traditionally associated to fruit flies and nematodes), the tempo and modes of SDf may vary significantly among species and may not even necessarily depend on or originate from SD itself (**Bear and Monteiro, 2013**; **Capel, 2017**). Classic examples of the independence between SD and SDf are provided by gynandromorph animals (where both the male and the female phenotypes are found in the same organism in a bilaterally-distributed fashion), but also by the marsupial mammal tammar wallaby (*Macropus* [now *Notamacropus*] *eugenii*). In gynandromorph chickens (*Gallus gallus*), for example, it has been shown that the male half of the animal is made up mainly by ZZ cells, while the female half by ZW cells, and that each is selectively susceptible to either male or female gonad-secreted hormones, respectively.

This shows how in birds SD, which is initiated in the presumptive gonads, is independent from and occurs later than SDf, which is instead triggered by a cell-autonomous mechanism (**Zhao et al., 2010**). Similarly, in *N. eugenii* the somatic differentiation of the scrotum and the pouch precedes that of the gonads, indicating that their development (as part of SDf) is triggered by genes linked to the X chromosome, rather than by sex-differentiation inducing hormones as in other mammals (**Renfree et al., 1996**).

Considering this complex scenario, **Uller and Helanterä (2011)** proposed a unified and broad-scope definition for SD, that is, “the processes within an embryo leading to the formation of differentiated gonads as either testes or ovaries”, thus excluding any actual distinction between environmental/genetic initial triggers or the downstream effectors. However, it can be argued that this definition should be even more expanded to encompass not only the embryonic stage of the animal life cycle but also adulthood, since cases of sex reversals (sequential hermaphroditism) legitimately express proper SD processes also during post-embryonic life stages. For example, fishes represent a noteworthy example of how the establishment of sexual fate is not an irreversible process in an organism’s life, but instead a tradeoff trait, involving antagonistic gene regulatory networks (GRNs), that can be switched from one side to the other (**Todd et al., 2016; Capel, 2017**). Sex reversal in fishes is typically prompted by environmental signals, such as population density, sex ratio at spawning, and social factors, but also by the attainment of a threshold size and/or age (reviewed in **Todd et al., 2016**). Regardless of its biology and direction (i.e., from males to females [protandry], from females to males [protogyny], or both ways), sex reversal often results in the complete restructuring of gonads, as well as in remarkable changes in both somatic morphology and behaviour. Therefore, sequential hermaphroditism can be considered the expression of a genuine SD program inducing a second round of SDf in adult organisms (**Todd et al., 2016**).

Overall, decades of studies have revealed that SD is strikingly diverse among animals, even in closely related species. Therefore, the research effort is currently devoted to further characterising sex-determining processes in new species, as well as to understand how this fundamental aspect of animal development may sustain such a high plasticity among species (**Beukeboom and Perrin, 2014; Todd et al., 2016; Capel, 2017**).

1.2 Genetic sex determination and the evolution of sex-determination related genes

In its most intimate core, animal SD is the manifestation of complex GRNs where, according to **Wilkins (1995)**, the downstream actors appear to be nearly conserved across species, while the master top regulators (the commonly recognized sex determinants, such as the *Sry* in therians or the ratio between sex and autosomal chromosomes in *Drosophila*) are often the most variable part (**Matson and Zarkower, 2012; Mullon et al., 2012; Bachtrog et al., 2014; Beukeboom and Perrin, 2014**). Such a differential pattern of molecular evolution is considered to be the direct result of the mechanism by which a sex-determining cascade is assembled. Particularly, it has been proposed that a GRN may take on a role in SD through a retrograde growth, i.e., by progressively adding upper regulators in a bottom-up process (**Stothard and Pilgrim, 2003; Mullon et al., 2012; Capel, 2017**). This mechanism regards the SD cascade in *Drosophila* species (**Mullon et al., 2012**), *C. elegans* (**Stothard and Pilgrim, 2003**), and vertebrates, although the latter case has been questioned several times (reviewed in **Capel, 2017**). Remarkably, it appears that some gene families are more prone than others to be recruited in SD, as either primary sex-determining genes (SDGs) or in some key part of the cascade. For example, components of the *dsx* and *mab-3* related transcription factor (Dmrt) gene family have a main role as bottom effectors in the SD cascade of many animal species, as seen in *D. melanogaster* with *doublesex* (*dsx*) and in *C. elegans* with *male abnormal-3* (*mab-3*), but also in other invertebrates and the majority of vertebrates. In these cases, Dmrt genes dictate the sex-specific development in response to the primary SD decision (**Matson and Zarkower, 2012**). Nevertheless, paralogs of *Dmrt-1* have also repeatedly and independently taken on the role as SDGs in several vertebrate species, as in the medaka fish *Oryzias latipes* with *dsx* and *mab-3* related gene *Y* (*Dmy*), in the African clawed frog *Xenopus laevis* with *dsx* and *mab-3* related gene *W* (*Dm-W*), and in *G. gallus* with the Z-linked *Dmrt-1* (reviewed in **Matson and Zarkower, 2012; Mawaribuchi et al., 2019**). A similar but even more conserved sex-determining genetic axis is found in insects, where *transformer* (*tra*) directs the sex-specific splicing of *dsx* in almost every species investigated so far (**Verhulst et al., 2010; Bopp et al., 2014**). The GRN in which the *tra-dsx* module is placed, is instead more diversified and species-specific, as the top- and bottom-most parts are highly divergent, resulting in a SD cascade that can be represented by an hour-glass model (**Bopp et al., 2014**). Similarly, other highly-conserved genes involved in SD has been identified,

particularly as downstream effectors in vertebrates: these includes for example *Fox-L2* from the forkhead box (Fox) gene family and *Sox-9* from the *Sry*-related HMG-box (Sox) gene families, acting in the female- and male-specific cascades, respectively (**Capel, 2017**).

The significance of molecular evolution in shaping SDGs is also evident in the wider category of sex-determination related genes (SRGs), which includes all the genes that are responsible for the specification, development and maintenance of the sexual identity. For example, transcriptionally sex-biased genes often tend to evolve faster than unbiased genes at the level of protein sequences. In particular, male-biased genes generally show higher rate of sequence evolution in comparison to both female-biased and unbiased counterparts, as it has been repeatedly observed in well-studied organisms—such as fruit flies, nematodes, mice and primates (reviewed in **Parsch and Ellegren, 2013; Grath and Parsch, 2016**), but also in other emerging systems, such as the water flea *Daphnia pulex* (**Eads et al., 2007**), aphids (**Purandare et al., 2014**), and two wasp species of the genus *Nasonia* (**Wang et al., 2015**). That said, growing evidence is also showing cases in which female-biased genes have higher rates of sequence evolution than male-biased genes, such as in mosquitoes of the genus *Anopheles* (**Papa et al., 2017**), and European and Manila clams of the genus *Ruditapes* (**Ghiselli et al., 2018**). High rates of molecular evolution in SRGs is particularly evident in organisms with sex chromosomes (SCs)—both in XY/ZW and X0 systems, such as fruit flies, birds and mammals, where the so-called fast-X (or fast-Z) effect has been extensively reported (**Vicoso and Charlesworth, 2006; Mank et al., 2007; Meisel and Connallon, 2013**). In these species, accelerated sequence evolution is seen in general for genes residing on the X (or Z) chromosomes (i.e., the chromosomes determining the homogametic sex) with respect to genes of the autosomal chromosomes, and it could be explained by both adaptive and non-adaptive processes. In fact, the higher ratio of non-synonymous to synonymous mutations (dN/dS , or ω) may result from positive selection, driven either by natural or sexual selection (as in *Drosophila*), as well as form genetic drift (as in birds; **Vicoso and Charlesworth, 2006; Meisel and Connallon, 2013; Parsch and Ellegren, 2013; Grath and Parsch, 2016**).

1.3 Unravelling sex determination in bivalves

Bivalves are the second largest clade in molluscs, counting more than 23,000 species ([Catalogue of Life](#); accessed on 15/10/2024) distributed at all depths and in all marine environments, as well

as in some freshwater habitats. Thanks to their high diversity and biological peculiarities, they have been proposed as promising model organisms for investigating a wide array of biological, ecological and evolutionary issues (**Milani and Ghiselli, 2020; Ghiselli et al., 2021; Nicolini, Ghiselli, et al., 2023**). However, despite the socio-economic and scientific importance, the knowledge concerning the molecular basis of bivalve reproduction and SD is still quite limited (**Breton et al., 2018; Nicolini, Ghiselli, et al., 2023**). Clues from various works seem to suggest that both genetic and environmental factors are involved in SD, though the exact process by which sex is determined and gonad commitment is established is, currently, still unknown.

In the attempt to identify SRGs (including SDGs), and clarify whether a single genetic determinant or a parliamentary decision exist, several differential gene expression (DGE) analyses have been recently performed on a variety of species (e.g., **Milani et al., 2013; Teaniniuraite-moana et al., 2014; Zhang et al., 2014; Chen et al., 2017; Capt et al., 2018; Ghiselli et al., 2018; Shi et al., 2018**). Particularly, some of the genes that were found to be differentially expressed between gonads of different sex were systematically retrieved, such as those belonging to the Dmrt, Sox, and Fox gene (DSFG) families. To this regard, **Zhang et al. (2014)** proposed a working model for the sex-determining pathway of the Pacific oyster *Crassostrea gigas* in which: *Sox-H* promotes male gonad development by activating *Dmrt 1-like* (*Dmrt-1L*), and inhibiting *Fox-L2*; *Fox-L2*, when not inhibited by the pair *Sox-H/Dmrt-1L*, promotes the female gonad development. Additionally, *Fox-L2* has been appointed as the female SDG—following a ZW inheritance system, in *Patinopacten yessoensis* and *Chlamys farreri*, based on the analysis of read coverage and of the distribution of sexually dimorphic single-nucleotide polymorphisms (SNPs; **Han et al., 2022**). However, both the SD model in *C. gigas* and the role of *Fox-L2* as the female SDG in *P. yessoensis* and *C. farreri*, have never been fully tested from a functional point of view (e.g., through gene editing or knock-down), and thus remain only hypothesis. Overall, much of the recent research effort on bivalve SRGs (including DSFGs) has indeed been limited to their molecular cloning, differential transcription, and tissue localization (**Liang et al., 2019; Sun et al., 2022**), and few works have directly investigated the biological functions so far, mostly through post-transcriptional silencing of target mRNAs (RNA interference [RNAi]). For example, **Liang et al. (2019)** studied the role of *Sox2* in the spermatogenesis of the Zhikong scallop *C. farreri* and found that it likely regulates proliferation of spermatogonia and apoptosis of spermatocytes, since its knockdown resulted in the loss of male germ cells. **Wang et al. (2020)** proposed that in the female gonads of the freshwater

mussel *Hyriopsis cumingii*, *Fox-L2* might be related to the *Wnt/β-catenin* signalling pathway, which takes part in ovarian differentiation also in vertebrates. **Sun et al. (2022)**, instead, found instead that in *C. gigas*, *Fox-L2* and *Dmrt-1L* mRNA knockdown results in the size reduction of female and male mature gonads, respectively. The challenge in identifying SDGs, if they exist, is partly due also to the apparent lack of heteromorphic sex chromosomes (HeSCs) in all the bivalve species investigated to date (**Breton et al., 2018; Han et al., 2022**). In fact, any evidence of SCs has only been found in four scallop species (*Amusium japonicum*, *C. farreri*, *Placopecten magellanicus*, *P. yessoensis*), where they have been described as homomorphic sex chromosomes (HoSCs; **Han et al., 2022**). Though, considering that DSFGs generally work in a coordinated manner to regulate many developmental processes also in other animal species, including the SD cascade itself (see **Section 1.2**), it is reasonable to assume that they play similar roles also in bivalves.

Our understanding of the environmental influences on SD is possibly even more limited. Given that bivalves exhibit a wide array of reproductive strategies—ranging from strict gonochorism to sequential (either protandrous or protogynous) and simultaneous hermaphroditism, as well as the so-called ‘alternative’ and ‘rhythymical sexuality’ (reviewed in **Breton et al., 2018**), they represent an excellent model to investigate the mechanisms of ESD. Temperature, food availability, social factors, and xenobiotics all seem to influence SD, or at least to trigger sex reversal in several hermaphroditic species (mainly belonging to the Ostreida and Pectinida orders). As a matter of fact, ESD has been investigated only in adult individuals through sex-ratio studies, thus few or no experiments are available for the very first round of SD (i.e., that encompassing the first gonad specification cycle). The Pacific oyster *C. gigas*, along with other oyster species, is one of the most studied bivalves not only for GSD (as mentioned above), but also for ESD. It has been shown that the sex ratio of adults is influenced by the incubation temperature of immature spats: at 18 °C, the sex ratio is skewed towards females, while at 28 °C it favours males, with some simultaneous hermaphrodites also observed (**Santerre et al., 2013**). Considering that SD in *C. gigas* may be also under genetic control (**Santerre et al., 2013; Zhang et al., 2014**; reviewed in **Breton et al., 2018**), these observations contribute to the growing evidence that a mixture of different factors govern SD in the species. A similar hypothesis of a mixed SD system has been suggested also for other species, such as *Crassostrea corteziensis* (**Chávez-Villalba et al., 2008**), *Pinctada margaritifera* (**Teaniniuraitemoana et al., 2016**), and *Mytilus edulis* (**Dalpé et al., 2022**), although more insightful and thorough investigations are

needed (**Dalpé et al., 2022**).

Clearly, bivalves represent a dazzling example of how the traditional representation of sex as genetically- or environmentally-determined, as well as the distinction between SD and SDf, can no longer be assumed as strictly dichotomous. A multifactorial model, in which many genes and environmental cues act in concert to establish the sexual identity of the individual, seems to better explain the extreme diversity of bivalve SD systems (**Breton et al., 2018**). Nonetheless, much work still needs to be done, especially in the functional characterisation of the molecular ground plan. Functional assays employing RNAi and clustered regularly interspaced short palindromic repeats (CRISPR) and CRISPR-associated protein 9 (CRISPR-Cas9) techniques (e.g., **Wang et al., 2020; Sun et al., 2022; Wang et al., 2022**) are finally making their way into the study of bivalve biology and have been proved essential instruments also for the investigation of sex-related traits. However, very few works have made extensive use of the comparative and integrative approach in bivalve studies so far, which hampers the possibility to infer general patterns for such a vast and diverse class of organisms (**Milani and Ghiselli, 2020**).

Chapter 2

Project outlook and objectives

This PhD project focuses on understanding the evolutionary dynamics and molecular mechanisms underlying sex determination (SD) in bivalve molluscs. The research has leveraged a wide array of analytical tools—from comparative genomics, to transcriptomics, *in-situ* hybridization, and immunolocalization, in order to investigate sex-determination related genes (SRGs) across various species with an integrative and comparative approach. Particularly, special attention was given to the Dmrt, Sox, and Fox gene (DSFG) families, which are widely-recognised key actors in the SD process of the majority of animal species, including bivalves. Each major area of analysis in my research, together with its objectives, is presented in a dedicated chapter, resulting in three distinct sections.

Chapter 3, which consists of a perspective piece published in *Genome Biology and Evolution*, will examine bivalves as emerging model organisms in SD research, by reviewing their genomic and biological characteristics. Bivalve offers valuable insights into several topics, including (i) the transitions between environmental and genetic SD, (ii) the evolution of sex chromosomes (SCs), (iii) the tentatively interaction between mitochondrial inheritance and SD, and (iv) the evolutionary history of SRGs. Particularly, this chapter wants to emphasise the importance of establishing a comprehensive evolutionary genomics framework for studying SD across bivalve species.

Chapter 4 will explore the molecular evolution of some key SRGs. Using a broad genomic context that includes more than 40 annotated bivalve genomes and transcriptomes, this chapter aims to uncover how these genes have evolved and their potential roles in SD, by also adopting a cross-species validation assay. The analysis will focus on the evolution of the DSFG families, by using the tools of molecular evolution to assess whether some of them are tightly involved

in SD. Mammals and *Drosophila* spp. will be used as positive and negative control datasets, respectively, to validate the reliability of the approach.

Chapter 5 will focus on the expression patterns of three SRGs in the Mediterranean mussel *Mytilus galloprovincialis* during early developmental stages. Particularly, the spatial and temporal transcription patterns of *Dmrt 1-like* (*Dmrt-1L*), *Sox-H*, and *Fox-L2*—which have been identified as tightly linked to primary SD by analyses in **Chapter 4** and in previous works, will be investigated. By also including the analysis of the germline marker *Vasa/Vasa*, this chapter will provide novel insights into the mechanisms of SD and primordial germ cell (PGC) specification. Transcription patterns will be investigated through computational differential gene expression (DGE) analyses and mRNA *in-situ* hybridization chain reaction (HCR); the expression pattern of *Vasa/Vasa* will be investigated also through immunolocalization.

Overall, this PhD project aims to adopt a multi-layered and integrative approach that combines evolutionary genomics, gene expression analyses, and comparative biology to explore SD in bivalves. Bivalves represent a relatively underexplored group, and given the remarkable diversity of their SD processes, require a strong evolutionary perspective to decipher the mechanism. Here, the integration of genome-wide molecular evolution analysis with gene expression studies provides a novel framework for understanding how SRGs, such as those belonging to the DSFG families, contribute to SD and sexual differentiation. This work also benefits from cross-species comparisons, which places bivalve SD within a broader evolutionary context, allowing for the identification of commonalities and unique traits in sex-determining pathways across taxa. Moreover, by investigating the expression patterns of three SRGs during early development in *M. galloprovincialis*, this project addresses a critical gap in the understanding of how these genes may regulate the sexual process, as to date bivalve SD has been investigated mostly in adult life stages. Through a comprehensive and comparative methodology, the project promises to provide a first reference broad-scale evolutionary resource for bivalve SD, also pushing forward the boundaries of reproductive and evolutionary biology in non-model species.

Chapter 3

Bivalves as emerging model systems to study the mechanisms and evolution of sex determination: a genomic point of view

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Abstract. Bivalves are a diverse group of molluscs that have recently attained a central role in plenty of biological research fields, thanks to their peculiar life history traits. Here we propose that bivalves should be considered as emerging model systems also in sex-determination studies, since they would allow to investigate: (i) the transition between environmental and genetic sex determination, with respect to different reproductive backgrounds and sexual systems (from species with strict gonochorism to species with various forms of hermaphroditism); (ii) the genomic evolution of sex chromosomes, considering that no heteromorphic sex chromosomes are currently known and that homomorphic sex chromosomes have been identified just in few species of scallops; (iii) the putative role of mitochondria at some level of the sex determination signaling pathway, in a mechanism that may resemble the cytoplasmatic male sterility of plants; (iv) the evolutionary history of sex-determination related gene families with respect to other

animal groups. In particular, we think that this last topic may lay the foundations for expanding our understanding of bivalve sex determination, as our current knowledge is quite fragmented and limited to few species. As a matter of fact, tracing the phylogenetic history and diversity of sex-determination related gene families (such as the Dmrt, Sox and Fox genes) would allow to perform more targeted functional experiments and genomic analyses, but also fostering the possibility of establishing a solid comparative framework.

Significance. In this perspective, we provide an examination of the phylogenetic diversity of Dmrt genes, a sex-determination related gene family, to address the importance of bivalves in sex determination studies. By analyzing their taxonomic distribution and sequence diversity, we show how such a comparative study may set a common ground plan to settle down targeted functional experiments and essays. This kind of approach should be applied more extensively in future studies, especially when dealing with understudied organisms.

Bivalves are the second largest clade in molluscs, counting more than 18,000 species ([Catalogue of Life](#), accessed on 16/12/2022) distributed at all depths and in all marine environments, as well as in some freshwater habitats. Thanks to their high diversity and peculiar biological features, they have been proposed as promising model organisms for investigating a wide array of biological, ecological, and evolutionary issues, from mitochondrial biology and evolution to the physiological plasticity under fluctuating environmental conditions (**Milani and Ghiselli, 2020; Ghiselli et al., 2021**). In this context, bivalves may serve as a compelling model system to investigate the evolution and characteristics of sex determination (SD) as well, thanks to the diversity of their reproductive modes and genomic features. Nonetheless, this research field has been largely overlooked and many aspects of bivalve reproductive biology remain uncharacterized. In this perspective, we address the topic by first examining the relevant questions that bivalves may help to answer regarding processes and patterns of SD, and then providing a case study in the field of comparative genomics.

3.1 Open yet inspiring topics in bivalve sex determination

Despite the socio-economic and scientific importance of bivalves, the knowledge concerning the genetic and molecular bases of their SD system is quite limited and its study has been mostly neglected. Yet, bivalves may constitute a novel model system in SD studies that is as intriguing and valuable as other well-established models, such as vertebrates, insects and plants (**of Sex Consortium, 2014**), as they may provide complementary perspectives in many aspects of SD evolutionary studies. Topics such as (i) the transition between environmental and genetic SD, (ii) the evolution of sex chromosomes, (iii) the mito-nuclear interaction, and (iv) the evolution of SD related genes, can largely benefit from the integration with bivalve studies. But many others are likely to emerge as research in the field progresses.

3.1.1 Transitions between environmental and genetic sex determination

Clues from several works seem to suggest that both genetic and environmental factors are involved in bivalve SD, thus implying that a mixed system may exist (reviewed in **Breton et al., 2018**). The traditional dichotomy between environmental sex determination (ESD) and genetic sex determination (GSD) seems inapplicable in most bivalve species, where ESD and GSD rather represent the two ends of a continuum of mixed and plastic conditions. A weak

distinction between ESD and GSD is also found in amphibians, reptiles and teleost fish, three clades in which environment-dependent SD has been largely studied. Here, the interaction—or even the transition—between the two sexual systems have been reported in many species, suggesting that sex-determining mechanisms can be extraordinary plastic (**Bachtrog et al., 2014; Capel, 2017**). Adding a representative and diverse group of Lophotrochozoa (Protostomia) to those vertebrate taxa, can widely expand the comparative framework of the investigation, allowing to better understand the evolution of SD as a whole. In bivalves, ESD has been studied mostly in oysters, where hermaphroditic species show an effect of temperature on SD (reviewed in **Breton et al., 2018; Fig. 3.1**). Oysters may indeed constitute a prolific model to examine how the SD pathways are shaped in the presence of different initial triggers and highly dynamic reproductive backgrounds. In fact, various sexual systems can be found in oysters, such as (i) strictly gonochoric population, (ii) the coexistence of simultaneous hermaphroditic with strictly gonochoric individuals in the same population, (iii) the possibility of sex change according to environmental conditions, and (iv) the presence of both parasitic dwarf males and free-living males in the same species (**Collin, 2013**). Consequently, oysters may be extremely useful to understand how epigenetic control is involved in sex change, how gene regulatory networks can sustain the occurrence of different hermaphroditic conditions within gonochoric populations, and whether certain SD systems are more labile than others (**Abbott, 2011**).

3.1.2 Evolution of sex chromosomes

So far, heteromorphic sex chromosomes (HeSCs)—i.e., sex chromosomes showing strong morphological differentiation, have never been observed in bivalves (**Breton et al., 2018**), while the first evidence of homomorphic sex chromosomes (HoSCs)—i.e., sex chromosomes showing little or no differentiation, comes from a very recent study on several scallop species, where a non-homologous origin of the SD system has been proposed for different subfamilies (**Han et al., 2022; Fig. 3.1**). Theory predicts that, once originated, sex chromosomes (SCs) will eventually turn into HeSCs, because of the recombination arrest in the sex-determining region (**Bachtrog et al., 2014; Beukeboom and Perrin, 2014; Han et al., 2022**). Nonetheless, HoSCs are much more widespread in the animal kingdom than expected, sometimes also being of ancient age (**Bachtrog et al., 2014; Han et al., 2022**).

Species from the order Pectinida may thus be useful to investigate what determines the long-term maintenance of HoSCs and which genomic architectures and molecular dynamics

prevent HeSCs from evolving in bivalves. Additionally, they may be taken as model systems to investigate the origin of SCs in relation to the sexual systems and the route by which molecular pathways have been reprogrammed in the transition between different SD mechanisms (**Han et al., 2022**).

Researchers have been addressing this topic mainly in snakes, ratites and sturgeons (**Bachtrog et al., 2014; Han et al., 2022** and references therein). Though, scallops currently hold the oldest HoSC pairs, which are dated back to about 350 million years. The system is thus of great importance to investigate the role of sex-biased gene expression and selection forces in the long-term stability of SCs (**Han et al., 2022**), as well as the intertwining between SD systems.

3.1.3 Mito-nuclear interactions

An additional pivotal topic in bivalve biology, tentatively connected to SD, regards the doubly uniparental inheritance (DUI) of mitochondria, a process in which two highly divergent mitochondrial genomes are transmitted uniparentally through the maternal and paternal lineages, respectively through eggs and sperm. This process, which has been reported in more than a hundred bivalve species from five different orders (**Fig. 3.1; Gusman et al., 2016; Capt et al., 2020**), has been proposed to interact with the major nuclear pathways that primarily establish the sexual identity, in a way that can resemble the cytoplasmatic male sterility (CMS) of plants (**Ghiselli et al., 2013; Breton et al., 2022**). In CMS, specific mitochondrial chimeric open reading frames (ORFs) cause the pollen to be sterile, while certain nuclear loci act in counterbalance to restore male fertility when occurring in the same individual. This Red-Queen scenario, in which balancing selection shapes the evolution of both CMS and restorer-of-fertility genes and keeps the two sexes viable, has been also hypothesized to be acting on bivalve DUI species (**Ghiselli et al., 2013; Xu, Iannello, et al., 2022**), where additional and effectively-transcribed ORFs have been observed in both the male-inherited and female-inherited mitochondrial lineages (**Milani et al., 2013, 2014**).

Clearly, if a functional interplay between DUI and SD in bivalves is proven, this will provide new research questions regarding not only bivalve biology itself but also broader evolutionary topics (e.g., are there any converging trait between DUI and CMS systems? What is the degree of plasticity of such mitochondria-related SD systems? Are mitochondria-related SD systems more widespread in eukaryotes than currently thought?).

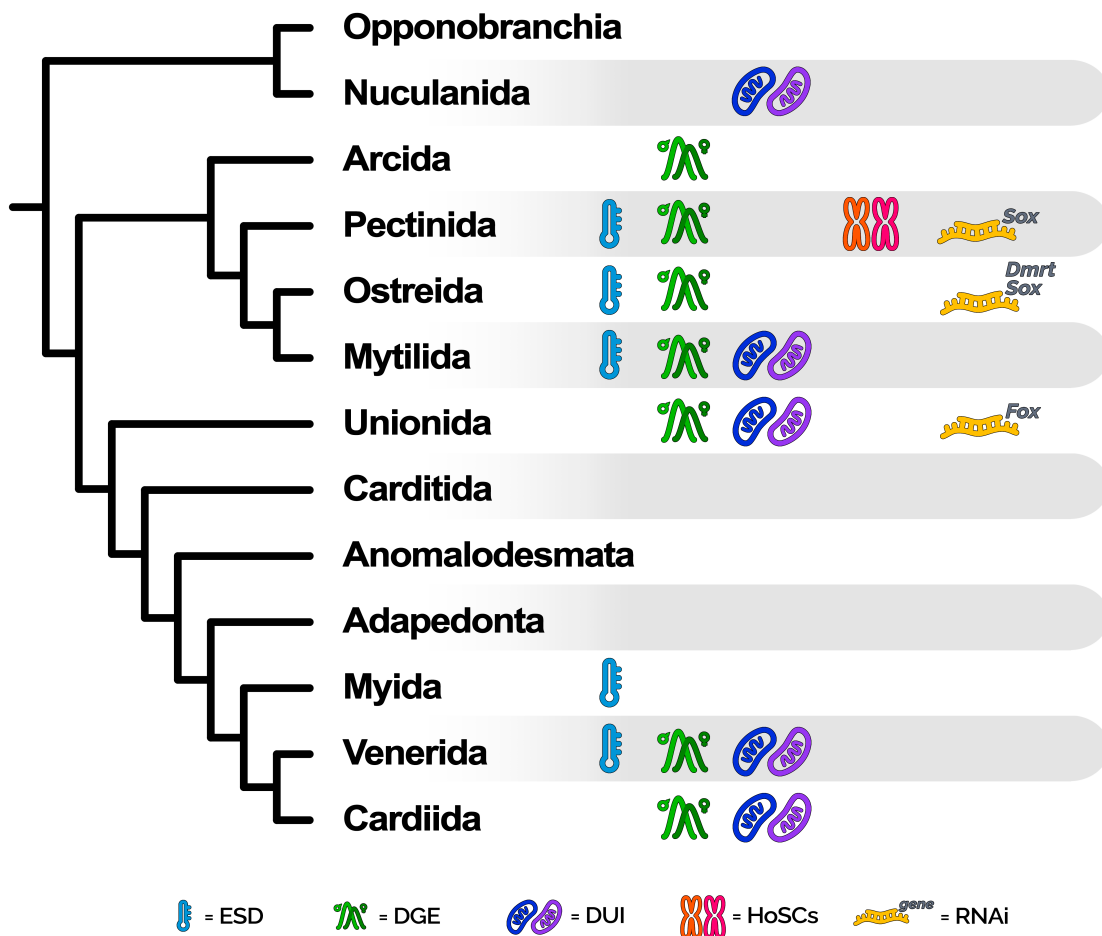


Figure 3.1 – Graphical summary of the available knowledge and experiments concerning the genetic basis of SD in bivalves, at the level of major taxonomic orders (as reported in WoRMS; accessed before or on 14/03/2023). For each bivalve clade it is reported: (i) the availability of records of ESD; (ii) the availability of differential gene expression (DGE) experiments specifically intended to investigate sex-biased or sex-specific genes; (iii) whether the DUI of mitochondria has been reported in at least one species; (iv) whether HoSCs have been identified in at least one species; (v) the availability of RNA interference (RNAi) experiments for genes belonging to the Dmrt, Sox, and Fox gene (DSFG) families. The phylogenetic tree on the left has been drawn on the basis of the most widely accepted topology for bivalves, according to analyses based on nuclear markers and morphological data. The tips of the tree correspond to major bivalve orders, except for Opponobranchia and Anomalodesmata, which represent higher-level taxonomic ranks. References for the availability of data and experiments can be found throughout the main text.

3.1.4 Evolution of sex-determination related genes

Considering this intricate scenario of SD mechanisms and the wide diversity of bivalves, in the last years many differential transcription analyses have been performed on several species with the attempt to identify the most probable sex-determination related genes (SRGs; e.g.,

Milani et al., 2013; Zhang et al., 2014; Chen et al., 2017; Capt et al., 2018; Shi et al., 2018; Fig. 3.1). Interestingly, certain genes consistently emerged across different bivalve species as being substantially more transcribed in one sex (sex-biased) or exclusively transcribed in one sex (sex-specific), suggesting their potential involvement in the SD pathway. These genes mainly belong to the *dsx* and *mab-3* related transcription factor (Dmrt), *Sry*-related HMG-box (Sox), and forkhead box (Fox) families, which play a role in various developmental processes (including the SD cascade) in most animals (Marshall Graves and Peichel, 2010; Bachtrog et al., 2014; Beukeboom and Perrin, 2014). Members of these three gene families are also included in the working model for the SD regulatory network proposed for the Pacific oyster *Crassostrea gigas* by Zhang et al. (2014), in which: *CgSoxH* (which belong to the Sox family) promotes male gonad development by activating *CgDsx* (also known as *Dmrt 1-like* [*Dmrt-1L*]; which belong to the Dmrt family) and inhibiting *CgFoxL2* (which belong to the Fox family); *CgFoxL2*, when not inhibited by the pair *CgSoxH/CgDsx*, promotes female gonad development. Similarly, Han et al. (2022) appointed *FoxL2* as a putative SD gene in the two scallop species *Patinopacten yessoensis* and *Chlamys farreri*. If their pivotal role in SD of bivalves is confirmed, an evolutionary genomic analysis may help in better understanding why members of the above-mentioned gene families appear particularly prone to be recruited in the SD cascade also in distantly related species, as it is observed for *Dmrt1* and *Sox3* homologs in vertebrates (Marshall Graves and Peichel, 2010; Bachtrog et al., 2014; and the following section). Furthermore, considering the occurrence of mixed SD systems in bivalves, DSFGs may provide new perspectives on the influence of different environmental cues on the molecular evolution of animal SRGs. However, to date, experiments have been limited to molecular cloning, differential transcription, and tissue localization of such genes (Liang et al., 2019; Sun et al., 2022), while only a few have directly investigated their biological functions in bivalves, for example through post-transcriptional silencing of target mRNAs (RNAi; Fig. 3.1; e.g., Liang et al., 2019; Wang et al., 2020; Sun et al., 2022).

Overall, DSFGs are highly interesting targets to be investigated in the framework of bivalve SD and have indeed obtained much more attention than the study of SCs or the role of environmental cues. However, much work is still to be done in order to understand their function in the SD signaling pathway and their evolutionary history.

3.2 The case of the Dmrt gene family in bivalves

Among the SRG candidates identified in bivalves, Dmrt genes (named after *doublesex* [*dsx*] from *Drosophila melanogaster* and *male abnormal-3* [*mab-3*] from *Caenorhabditis elegans*) are of particular interest. As a matter of fact, in vertebrates, besides their role in placode neurogenesis and somite patterning (reviewed in **Mawaribuchi et al., 2019**), Dmrt genes are also involved in the development of male gonads and the maintenance of the testicular function (**Sun et al., 2022**). Their role in the specification and organization of male sexual characters seems indeed to be common across Metazoa, suggesting that a similar function may have been already present in the Bilateria common ancestor (**Kopp, 2012; Beukeboom and Perrin, 2014**).

The first attempts to dig inside the phylogenetic history and diversity of bivalve Dmrt genes have been provided by **Li et al. (2018)** and **Evensen et al. (2022)**: besides retrieving all the canonical genes (i.e., *Dmrt2*, *Dmrt3* and *Dmrt4/5*), their inferences brought to light a monophyletic Dmrt group (named *Dmrt-1L*) which appears to be private to molluscs and present in several bivalve species. The *Dmrt-1L* monophyletic group is confirmed also when expanding the analysis by mining genomes from a wider range of bivalve taxa (**Tab. 3.1; Fig. 3.2A**), suggesting that *Dmrt-1L* genes are widespread in bivalves and were likely present in their common ancestor (**Evensen et al., 2022**). In particular, *Dmrt-1L* genes can be successfully retrieved in species of the orders Mytilida, Ostreida, Pectinida, Unionida, and from *Scapharca broughtonii* (Arcida), while the opposite holds for Venerida, *Sinonovacula constricta* (Adapedonta), and *Dreissena* spp. (Myida; **Fig. 3.2B**). Clearly, the absence of *Dmrt-1L* genes demands further investigations, as it may derive from errors in genome assembly and annotations.

The present analysis also supports a higher amino acid sequence divergence of the *Dmrt-1L* orthology group with respect to the other Dmrt orthology groups (**Fig. 3.2C**), which may be explained by a higher rate of sequence evolution related to their sex-biased expression in certain species (**Zhang et al., 2014; Shi et al., 2015; Li, Zhang, et al., 2018; Evensen et al., 2022**). This is consistent with what has been already observed for the SRGs *Dmrt1* and *dsx* in vertebrates and *Drosophila*, respectively (e.g., **Bewick et al., 2011; Baral et al., 2019**). In fact, sex-biased genes (including SRGs) often tend to evolve faster than unbiased genes at the level of protein sequences, either when considering male-biased (reviewed in **Parsch and Ellegren, 2013; Grath and Parsch, 2016**) or female-biased genes (e.g., **Papa et al., 2017; Ghiselli et al., 2018**). Another possible explanation for the higher amino acid divergence of *Dmrt-1L* genes

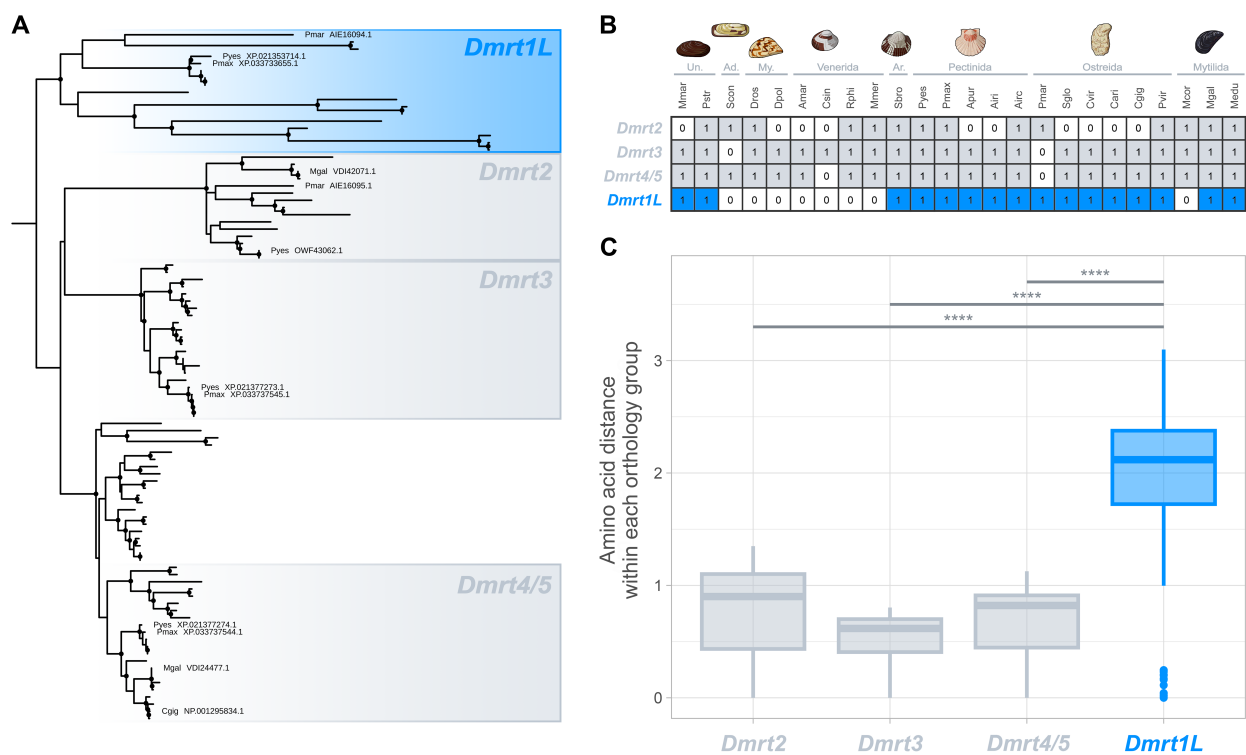


Figure 3.2 – Phylogenetic tree (A) and taxonomic distribution (B) of Dmrt genes in bivalves, and comparison of amino acid pairwise distances within *Dmrt-1L* and the other Dmrts (C). (A) Dmrt orthologs from bivalve genome assemblies were obtained with HMMsearch (HMMER toolkit; Eddy, 2011) with the Pfam HMM profile of the DM domain (PF00751). Amino acid alignment was obtained with MAFFT-DASH (Rozewicki et al., 2019), and manually inspected to remove poorly aligning sequences, and trimmed with trimAI (gap threshold of 60 %; Capella-Gutiérrez et al., 2009). The phylogenetic analysis was carried out using IQ-TREE 2 (Minh et al., 2020) with default parameters. Nodes with bootstrap values greater than 84 are marked with filled black circles. The tree was rooted according to Evensen et al., 2022. Dmrt genes analysed by Evensen et al., 2022 were used as reference to annotate the various orthology groups, and accession numbers are reported in the tree. The phylogenetic tree with all annotated tips and nodes can be accessed on supplementary material online. (B) Taxonomic distribution of identified Dmrt genes in bivalve genomes. Orders as reported in WoRMS (accessed before or on 14/03/2023) and in Fig. 3.1 are specified. (C) Pairwise amino acid distances were computed for amino acid sequences within each Dmrt orthology group identified in the tree, with the R package ‘phangorn’ (Schliep, 2011) under the JTT substitution model. After checking for normality with the Shapiro-Wilk test ($W = 0.88544$, $p < 2.2 \times 10^{-16}$) and for group effect with the Kruskal-Wallis test ($p < 2.2 \times 10^{-16}$), the pairwise Wilcoxon rank-sum test was used to compare the distributions of pairwise amino acid distances of *Dmrt-1L* and the other Dmrts. Horizontal bars mark the significative results with $p < 2.2 \times 10^{-16}$ (****; Bonferroni correction for multiple test was applied). The list of genome assemblies used for these analyses and species identifiers can be found in Tab. 3.1. Un.: Unionida; Ad.: Adapedonta; My.: Myida; Ar.: Arcida.

may lie on their expression breadth, that is, genes with a narrow tissue-specific expression tend to evolve faster than more ubiquitous genes (**Parsch and Ellegren, 2013; Xu, Martelossi, et al., 2022**). As a matter of fact, *Dmrt-1L* genes have been found to be significantly more transcribed in the gonadic tissue (particularly in testes) in *P. yessoensis* (**Li, Zhang, et al., 2018**) and *C. gigas* (**Yue et al., 2021**).

Understanding the role and molecular interactions of *Dmrt-1L* genes in bivalve SD and gonad development would greatly enhance the possibility of outlining the evolutionary causes and consequences of their high amino acid divergence (**Fig. 3.2C**), for example by linking the molecular evolution to the degree of pleiotropy. However, most of our knowledge on *Dmrt-1L* biology is currently limited to the temporal and tissue localization of transcripts in a few species of bivalves (e.g., **Li, Zhang, et al., 2018; Yue et al., 2021**). In fact—apart from the work by **Sun et al. (2022)**, which confirmed the role of *Dmrt-1L* in the gonad development of *C. gigas* through non-invasive RNAi and found that the knocked-down phenotype results in size reduction of male gonads—no other experiments intended to elucidate the function of *Dmrt-1L* genes in bivalves have been carried out so far (**Fig. 3.1**). This clearly hinders any possible integration between molecular data with functional assays. If the role of *Dmrt-1L* as major sex determinants was confirmed, bivalves would become an intriguing clade in which investigate why, in Metazoa, certain genes (namely, the Dmrt gene family) appear particularly prone to being recruited at the top of the SD cascade. To date, this phenomenon has been widely examined in vertebrates, where *Dmrt1* genes have independently gained a primary role in male SD in fish, amphibians, and birds, and are considered candidate sex-determining genes also in monotreme mammals (**Marshall Graves and Peichel, 2010; Beukeboom and Perrin, 2014; Mawaribuchi et al., 2019**). Bivalves may provide an alternative evolutionary scenario to study the selective forces and molecular modifications that support Dmrt genes in repeatedly taking over the SD process. In fact, since *Dmrt-1L* genes seem to be restricted to molluscs (**Fig. 3.2A**), it would be intriguing to clarify if the putative involvement in the SD cascade of extant bivalve species is the result of shared ancestry or convergent evolution, which would establish a study system for the evolution of Dmrt genes parallel to that of vertebrates (see **Capel, 2017**).

Obviously, *Dmrt-1L* should not be expected to be the sole sex-determining gene (SDG). In fact, *Fox-L2* has already been appointed as the female SDG in *P. yessoensis* and *C. farreri* (**Han et al., 2022**). Consequently, we should expect that other primary genetic determinants

exist, consistently with the extremely high species diversity of the clade. Thus, bivalves may additionally serve as a valuable model system to study how genes from different families take over the SD cascade and are shaped by selection.

3.3 Conclusions: bivalves as new models in the study of sex determination

SD is undoubtedly a fascinating biological and evolutionary topic as much as it is challenging to investigate. Our understanding of the causes and consequences of the SD mechanism diversity strongly relies on the study of different systems and non-model model organisms (**Bachtrög et al., 2014; Milani and Ghiselli, 2020**), which provide the foundation for depicting a comprehensive evolutionary and comparative framework in which new and coherent research perspectives can be grounded.

In recent years, bivalves have been achieving growing importance in many fields of biology, from ecology to genomics, and from environmental biomonitoring to mitochondrial studies (**Milani and Ghiselli, 2020; Ghiselli et al., 2021**), but they can be a valuable model to address also SD studies. The diversity of their life history traits provides indeed a challenging, yet extremely fascinating framework, to put the SD processes into an evolutionary context.

Bivalves can help us explain how ESD and GSD interplay with each other in response to the environmental conditions, as a mixed system of both has been proposed to act in the establishment of bivalve sexual identity (reviewed in **Breton et al., 2018**). Moreover, the occurrence of the many existing variants of hermaphroditism and gonochorism even in closely related species, or within the same population, strongly suggests that the basic SD pathway (whether genetic, environmental, or mixed) should be plastic enough to sustain the existence of individuals of both sexes, thus providing the opportunity to study how SD gene regulatory networks are shaped and selected throughout evolution and how epigenetic regulation may influence SD. The unique DUI system further poses an undeniable challenge in SD studies since it may represent an SD-linked mechanism which relies on the non-nuclear portion of the genome and may unfold many new research paths (**Milani and Ghiselli, 2020; Ghiselli et al., 2021**). Nonetheless, much of the research effort on bivalve SD has been devolved to specific groups of socio-economic importance, such as Mytilida, Ostreida, Pectinida, and Unionida, while the

other lineages of the bivalve phylogeny have been neglected (**Fig. 3.1**). Our understanding of the SD processes of bivalves is thus restricted and is mainly lacking a broad comparative framework in which to draw comprehensive evolutionary inferences.

Genes from the DSFG families, which are involved in SD also in other Metazoa, may be considered excellent genomic targets to study the processes and patterns of molecular evolution in sex-biased genes, as well as of the recurrent recruitment of genes in the SD cascade. Also, identifying the major genetic regulators of SD in bivalves would burst the functional study of the interaction between ESD and GSD, by providing genetic targets that can be manipulated through RNAi and/or genome editing techniques to understand the role of environmental cues in SD. In the same way, knowing the main genetic actors of SD would allow researcher to identify SCs not only on the basis of *in-silico* techniques (such as k-mer based or single nucleotide polymorphism [SNP] methods) but also by less-expensive wet lab protocols (such as fluorescence mRNA *in-situ* hybridization [mRNA-ISH] on metaphase chromosome plates). Furthermore, it would help to understand whether and how the mitochondrial additional ORFs of DUI species interact with the SD system, by performing thorough gene expression essays.

In conclusion, we strongly urge researchers to invest more resources in the integrative study of bivalve SD to unravel the many underlying mechanisms and expand our understanding of this biological process. Given our limited knowledge in the field, one of the first routes that should be undertaken may rely on the comparative study of SRGs of bivalves from a genomic perspective, as this kind of data is nowadays growing at a rate faster than ever. Establishing such a genomic ground plan for understudied organisms will in fact allow researchers to develop evolutionary-aware experiments with better selected genetic targets.

Table 3.1 – List of bivalve genomes from which Dmrt genes have been extracted. For each species, the accepted name and the most-common synonym (in parentheses) are reported. NCBI accession numbers are provided, when available, as well as BUSCO scores of the predicted proteomes against the ‘metazoa_odb10’ dataset (Manni et al., 2021).

Species	ID	Order	Assembly level	BUSCO score	Reference	NCBI Acc. No.
<i>Anadara (Scapharca) broughtonii</i>	Sbro	Arcida	Chromosome	C:91.2% [S:85.6%,D:5.6%] F:2.6% M:6.2%	Bai et al., 2019	–
<i>Sinonovacula constricta</i>	Scon	Adapedonta	Chromosome	C:92.5% [S:80.4%,D:12.1%] F:3.4% M:4.1%	Ran et al., 2019	GCA_007844125.1
<i>Dreissena polymorpha</i>	Dpol	Myida	Chromosome	C:86.9% [S:75.1%,D:11.8%] F:6.4% M:6.7%	McCartney et al., 2022	GCA_020536995.1
<i>Dreissena rostriformis</i>	Dros	Myida	Scaffold	C:75.2% [S:73.2%,D:2.0%] F:15.2% M:9.6%	Calcino et al., 2019	GCA_007657795.1
<i>Mytilus unguiculatus (coruscus)</i>	Mcor	Mytilida	Chromosome	C:80.0% [S:79.1%,D:0.9%] F:7.7% M:12.3%	Yang et al., 2021	GCA_017311375.1
<i>Mytilus edulis</i>	Medu	Mytilida	Scaffold	C:83.7% [S:64.5%,D:19.2%] F:5.2% M:11.1%	Corrochano-Fraile et al., 2022	GCA_905397895.1
<i>Mytilus galloprovincialis</i>	Mgal	Mytilida	Scaffold	C:80.3% [S:47.5%,D:32.8%] F:8.8% M:10.9%	Gerdol et al., 2020	GCA_900618805.1

Tab. 3.1 continued from previous page

Species	ID	Order	Assembly level	BUSCO score	Reference	NCBI Acc. No.
<i>Perna viridis</i>	Pvir	Mytilida	Scaffold	C:99.4% [S:99.0%,D:0.4%] F:0.2% M:0.4%	Inoue et al., 2021	GCA_018327765.1
<i>Magallana (Crassostrea) ariakensis</i>	Cari	Ostreida	Chromosome	C:94.6% [S:90.9%,D:3.7%] F:0.9% M:4.5%	Li et al., 2021	GCA_020567875.1
<i>Magallana (Crassostrea) gigas</i>	Cgig	Ostreida	Chromosome	C:98.5% [S:67.6%,D:30.9%] F:0.3% M:1.2%	Penaloza et al., 2021	GCF_902806645.1
<i>Crassostrea virginica</i>	Cvir	Ostreida	Chromosome	C:98.1% [S:58.6%,D:39.5%] F:0.3% M:1.6%	Gómez-Chiarri et al., 2015	GCF_002022765.2
<i>Saccostrea glomerata</i>	Sglo	Ostreida	Scaffold	C:88.9% [S:85.3%,D:3.6%] F:5.1% M:6.0%	Powell et al., 2018	GCA_003671525.1
<i>Argopecten irradians concentricus</i>	Airc	Pectinida	Scaffold	C:94.8% [S:93.9%,D:0.9%] F:3.7% M:1.5%	Liu et al., 2020	GCA_004382765.1
<i>Argopecten irradians irradians</i>	Airi	Pectinida	Scaffold	C:94.8% [S:93.9%,D:0.9%] F:3.7% M:1.5%	Liu et al., 2020	GCA_004382745.1
<i>Argopecten purpuratus</i>	Apur	Pectinida	Scaffold	C:89.2% [S:88.5%,D:0.7%] F:5.0% M:5.8%	Liu et al., 2020	–

Tab. 3.1 continued from previous page

Species	ID	Order	Assembly level	BUSCO score	Reference	NCBI Acc. No.
<i>Pecten maximus</i>	Pmax	Pectinida	Chromosome	C:98.5% [S:74.7%,D:23.8%] F:0.4% M:1.1%	Kenny et al., 2020	GCF_902652985.1
<i>Mizuhopecten (Patinopecten) yessoensis</i>	Pyes	Pectinida	Scaffold	C:98.6% [S:75.2%,D:23.4%] F:0.4% M:1.0%	Wang, Zhang, et al., 2017	GCF_002113885.1
<i>Margaritifera margaritifera</i>	Mmar	Unionida	Scaffold	C:92.6% S:82.3%,D:10.3%] F:3.2% M:4.2%	Gomes-dos-Santos et al., 2021	GCA_015947965.1
<i>Potamius streckersoni</i>	Pstr	Unionida	Scaffold	C:74.7% [S:73.8%,D:0.9%] F:7.0% M:18.3%	Smith, 2021	GCA_016746295.1
<i>Calyptogena (Archivesica) marissinica</i>	Amar	Venerida	Chromosome	C:82.0% [S:80.0%,D:2.0%] F:6.1% M:11.9%	Ip et al., 2021	GCA_014843695.1
<i>Cyclina sinensis</i>	Csin	Venerida	Scaffold	C:94.0% [S:83.8%,D:10.2%] F:1.9% M:4.1%	Wei et al., 2020	GCA_012932295.1
<i>Mercenaria mercenaria</i>	Mmer	Venerida	Chromosome	C:95.4% [S:70.9%,D:24.5%] F:0.5% M:4.1%	Song et al., 2021	GCF_014805675.1
<i>Ruditapes philippinarum</i>	Rphi	Venerida	Chromosome	C:83.4% [S:74.5%,D:8.9%] F:8.8% M:7.8%	Xu, Martelossi, et al., 2022	GCA_026571515.1

3.4 Acknowledgments

The authors are extremely thankful to Sofía Blanco González from the University of Vigo for her willingness to engage in discussions and for genuinely sharing her opinion on this work.

3.5 Data Availability

Analyzed data and R scripts used to generate plots can be accessed in supplementary material online deposited at the following GitHub repository: [filonico/bivalve_sex_perspective](https://github.com/filonico/bivalve_sex_perspective).

Chapter 4

Identification of putative sex-determination related genes in bivalves through comparative molecular evolutionary analyses

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Manuscript in preparation.

4.1 Introduction

In sexually reproducing organisms, the modes of sex determination (SD), i.e., the process by which the male or female identity of an organism (or of the gonadic tissue) is established, is highly diverse, ranging from strictly genetic systems to environmentally-dependent processes (**Haag and Doty, 2005; Uller and Helanterä, 2011; Bachtrog et al., 2014; Beukeboom and Perrin, 2014**). Characterising the molecular basis of SD is crucial for understanding not only reproductive biology but also the evolutionary pressures shaping these systems (**Wilkins, 1995; Ellegren and Parsch, 2007; Grath and Parsch, 2016; Nicolini, Ghiselli, et al., 2023**), as sex-determination related genes (SRGs), including primary sex-determining genes (SDGs), are those responsible for the phenotypic differences of males and females, thanks to their sex-biased expression and interactions (**Ellegren and Parsch, 2007; Beukeboom and Perrin, 2014; Grath and Parsch, 2016**). One key aspect of SRGs is that they often exhibit accelerated rates of sequence evolution, due to their involvement in sex-related traits and reproduction. This represents the effects of sexual and/or adaptive selection, which act in sex-biased genes and produce high-divergent proteins at the interspecific level (**Civetta and Singh, 1998; Ellegren and Parsch, 2007; Meisel, 2011; Grath and Parsch, 2016**). Rapid sequence evolution is known for *Sex-determining region of chromosome Y (Sry)* of eutherians (**Pamilo and O'Neill, 1997; Mawaribuchi et al., 2012**), *dsx* and *mab-3* related gene *W (Dm-W)* of the African clawed frog *Xenopus laevis*, and *dsx* and *mab-3* related gene *Y (Dmy)* of the medaka fish *Oryzias latipes* (**Mawaribuchi et al., 2012**), all of which are master SDGs, that is, genes whose expression is primarily responsible for the establishment of the sexual fate of the organism. Evolution under episodic diversifying selection has been detected also in *Drosophila* for genes involved in the SD cascade (e.g., *Sex-lethal [Sxl]*, *transformer [tra]*, and *doublesex [dsx]*), in correspondence with its establishment in the genus common ancestor (**Mullon et al., 2012; Baral et al., 2019**); though, rapid sequence evolution seems to not be concerning extant amino acid sequences (**Haerty et al., 2007; Baral et al., 2019**), as they are globally evolving under purifying selection, especially in their catalytic domain (**Mullon et al., 2012; Baral et al., 2019**). Concerning the *dsx* genes, higher rates of nucleotide and amino acid sequence evolution can be however observed for male-specific regions, if compared to female-specific and oligomerization regions (**Baral et al., 2019**).

While SD has been extensively studied in model organisms, like mammals, insects, and

nematodes, comparatively little is known about the molecular ground plans in non-model organisms. A remarkable example of this is represented by bivalve molluscs, which exhibit a wide variety of reproductive strategies and sexual systems (**Breton et al., 2018**). Notwithstanding the considerable importance in the human socio-economic landscape (reviewed in **Haszprunar and Wanninger, 2012; Gomes-dos-Santos et al., 2020**), the study of SD mechanisms in bivalves has been hampered by the striking divergence among species (**Li et al., 2022**), and thus largely overlooked and limited to few case studies (**Breton et al., 2018; Nicolini, Ghiselli, et al., 2023**). So far, no master SDG has been unambiguously identified, and the only working hypothesis on the functioning of the SD gene regulatory network is available for the Pacific oyster *Crassostrea gigas* (now *Magallana gigas*; **Zhang et al., 2014**). Nonetheless, the field still lacks both a robust functional investigation and an evolutionary framework in which to place the current knowledge (**Nicolini, Ghiselli, et al., 2023**). As a matter of fact, major efforts have been dedicated to identify sex-biased genes through differential gene expression (DGE) analyses (e.g., **Milani et al., 2013; Teaniniuraitemoana et al., 2014; Zhang et al., 2014; Capt et al., 2018; Afonso et al., 2019**), but very few have leveraged cutting-edge techniques to investigate their actual role in SD and/or gonad differentiation and development (e.g., **Liang et al., 2019; Sun et al., 2022**).

Components of the Dmrt, Sox, and Fox gene (DSFG) families are notoriously known as key actors in several developmental processes across Metazoa (**Benayoun et al., 2011; Matson and Zarkower, 2012; Sarkar and Hochedlinger, 2013; Mawaribuchi et al., 2019**), including SD in certain clades: the aforementioned *Dm-W*, *Dmy*, and *dsx* all belong to the *dsx* and *mab-3* related transcription factor (Dmrt) gene family, while *Sry* belongs to the *Sry*-related HMG-box (Sox) gene family; *Fox-L2*, which takes part in most of the vertebrate SD processes as a downstream effector of the female pathway, belongs to the forkhead box (Fox) gene family. Members of the DSFGs have been identified as putative SRGs also in bivalves, thanks to both DGE analyses and mRNA *in-situ* hybridization (mRNA-ISH; e.g., **Naimi et al., 2009; Li, Zhang, et al., 2018; Liang et al., 2019; Yue et al., 2021**), suggesting that their role in morphological and sexual development is maintained also in the clade. However, the clear role of DSFGs has yet to be elucidated, probably as a consequence to the lack of (i) a systematic classification of the families and (ii) a comprehensive understanding of their evolutionary history.

In order to overcome such limitations, the present study aims to perform a thorough investigation of the DSFG families in bivalves, with the attempt to provide a high-quality resource

to be used as a reference for future studies. Through the analysis of more than 40 annotated bivalve genomes and transcriptomes, we aim (i) to describe the complete set and evolutionary history of DSFGs in bivalves by means of phylogenetic inferences, manual curation, and orthology prediction; furthermore, we aim (ii) to identify DSFGs potentially involved in bivalve SD by investigating their sequence evolution in a genome-wide context. As a matter of fact, our hypothesis is that, if any of the DSFGs is directly involved in SD (i.e., is a SDG), then we should expect it to be experiencing a higher rate of sequence evolution, as already found in previous studies (**Pamilo and O'Neill, 1997; Mawaribuchi et al., 2012**) and discussed earlier; this characteristic, in turn, would be reflected in a high diversity of the extant amino acid sequences across the bivalve clade. To assess the robustness and reliability of our approach, we additionally applied our pipeline to two non-bivalve datasets, composed of mammal and *Drosophila* species, respectively (hereon referred to as the ‘mammal dataset’ and the ‘fruit fly dataset’). By choosing two clades for which SD is well characterised, we wanted to compare our results with those obtained on taxa for which a deeper and more detailed knowledge is available. Particularly, mammals and *Drosophila* provide two different frameworks to study the patterns of molecular evolution in SDGs: the former is a system where SD is completely genetic (i.e., the development into a male or into a female is triggered by the up- or down-regulation of *Sry* in undifferentiated gonads, respectively), while the latter is a system where SD is chromosomal, thus lacks a master SDG (the sexual fate of the individual is determined by the ratio between autosomal and X chromosomes). Hence, they represent opposing control datasets to be compared to bivalves, as it is expected that a higher rate of sequence evolution concerns only master SDGs (i.e., the top regulatory part of the SD cascade), but not also the downstream genes (i.e., the bottom effectors). If our method is robust, we should thus expect that, (i) in the mammalian dataset *Sry* is detected as rapidly-evolving (**Pamilo and O'Neill, 1997; Mawaribuchi et al., 2012**), while (ii) in the fruit fly dataset no gene among those working within the sex-determining cascade (including *dsx*) is evolving at a higher pace (**Haerty et al., 2007; Mullon et al., 2012; Baral et al., 2019**). By testing the performance of the pipeline in mammals and fruit flies, we were able to assess the reliability of results in bivalves.

This work offers novel insights into the evolutionary dynamics of SRGs and contributes a valuable genomic resource for understanding SD in bivalves, one of the most ecologically and economically important groups of marine organisms. Particularly, here we provide the first extensive phylogenetic-based classification of DSFGs in bivalves, covering many species from

the major bivalve orders, along with a comprehensive investigation of their sequence evolution.

4.2 Materials and methods

4.2.1 Dataset of bivalve annotated genomes and transcriptomes

Annotated genome assemblies of bivalves were obtained from various publicly available resources, while reference genome assemblies for gastropods and cephalopods were downloaded from NCBI (**Tab. S1**). Isoforms were removed from genome annotations using a perl script from the AGAT toolkit (v0.8.0; **Dainat et al., 2022**). Concerning *Sinonovacula constricta* (Adapedonta), the nucleotide coding sequence fasta file was not available for download. To avoid excluding the species from our analyses, the file was generated in-house by mapping the annotated protein sequences on the reference genome using miniprot (v0.13-0; **Li, 2023**). Then, the corresponding nucleotide sequences were extracted using AGAT.

In order to provide an extensive identification of SRGs also for underrepresented bivalve orders (mainly belonging to the Heterodonta clade), 14 additional species represented by sequenced transcriptomes were included in the analyses. Assembled and annotated transcriptomes were obtained from **Piccinini et al., 2021** and **Iannello et al., 2023**. The resulting set of annotated genomes and transcriptomes (hereafter referred to as the ‘comprehensive set’) was checked for completeness using BUSCO with the Metazoa reference dataset (v5.2.2; **Manni et al., 2021**).

4.2.2 Identification and classification of Dmrt, Sox and Fox genes in bivalves

Members of DSFG families were retrieved in the comprehensive set with HMMsearch from the HMMER package (v3.3.2; **Eddy, 2011**). The signature catalytic domains of each family were used as queries. Specifically, hidden Markov model (HMM) profiles were built after the Pfam databases for the *dsx* and *mab-3* (DM) domain (PF00751), the high mobility group (HMG) box (PF00505) and the forkhead domain (PF00250) to retrieve members of the DSFG families, respectively. The e-value for both the per-target and the per-domain inclusion threshold was set to 10^{-5} .

Obtained hits were then annotated using (i) the PANTHER HMM standalone sequence scoring against the PANTHER library v18.0 and (ii) RPS-BLAST (v2.5.0+) against the Con-

served Domain Database (CDD; pre-compiled version, downloaded from <ftp.ncbi.nih.gov> on 09/11/23). In both cases, hits with an e-value of 10^{-5} were retained. Genes which were correctly annotated by both systems (on the basis of the PANTHER gene family and CDD domain identifiers; **Tab. S2**) were kept for subsequent analyses.

DSFGs from *Homo sapiens*, *Drosophila melanogaster*, and *Caenorhabditis elegans* (**Tab. S3**; hereafter referred to as ‘reference species’) were retrieved from NCBI and were used as reference genes for annotation (see below). Classification and nomenclature of each family was retrieved from: **Mawaribuchi et al. (2019)** for Dmrt genes; **Phochanukul and Russell (2010)** and **Sarkar and Hochedlinger (2013)** for Sox genes; **Mazet et al. (2003)** for Fox genes.

The alignments of mollusc and reference DSFGs were guided by the aforementioned Pfam HMM profiles and performed with Clustal Omega (v1.2.3; **Sievers et al., 2011**), then trimmed with trimAl (v1.4.rev15; **Capella-Gutiérrez et al., 2009**) with a gap threshold of 40 %. Resulting alignments were manually inspected to remove sequences with incomplete catalytic domains, then aligned and trimmed again as before. Phylogenetic trees were inferred using IQ-TREE (v2.1.4-beta COVID-edition; **Minh et al., 2020**) with automatic model selection (**Kalyaanamoorthy et al., 2017**), 1,000 bootstrap replicates and 5 independent runs. The phylogenetic tree of Dmrt genes was midpoint rooted, as no clear homology relationship has been found with other gene families or zinc-finger proteins so far (**Wexler et al., 2014**). Phylogenetic trees of Sox and Fox gene families were rooted using two fungi mating protein A (Mat-A) sequences (XP_62685912.1, CCD57795.1) and two Amoebozoa forkhead-like domains (XP_004368148.1, XP_004333268.1), respectively (**Nakagawa et al., 2013**; **Heenan et al., 2016**). The rooting was performed with Gotree (v0.4.5; **Lemoine and Gascuel, 2021**). To identify and annotate bivalve homology groups within each gene family, we employed a species overlap algorithm followed by a Markov clustering algorithm (MCL) weighted by node supports as implemented in Possvm (v1.2; **Grau-Bové and Sebé-Pedrós, 2021**). DSFGs from *H. sapiens*, *D. melanogaster*, and *C. elegans* were used as reference annotation.

In order to better establish the orthology relationships among ambiguous groups of Dmrt and Fox genes, we run a series of other phylogenetic reconstructions (see **Section 4.4**), by using the same pipeline as before. In the case of *Fox-Y* genes, we also employed Fox gene sequences from the sea urchin *Strongylocentrotus purpuratus*, as given by **Tu et al. (2006)**. All the phylogenetic trees were plotted using the R package ‘ggtree’ (**Yu et al., 2017**).

4.2.3 Sequence diversity of bivalve single-copy orthogroups

As a metrics to measure the sequence diversity of bivalve DSFGs, and test whether those putatively involved in SD show higher values than other genes, we employed the amino acid sequence divergence (AASD). As a matter of fact, this metric is fast and straightforward to obtain, as it only requires the amino acid alignment and the corresponding best-fit substitution mode.

To this purpose, we produced amino acid alignments of bivalve single-copy orthogroups (SCOs) and built the distribution of their median AASD. Specifically, we assembled a second dataset (hereafter referred to as the ‘reduced bivalve dataset’) which includes, for each bivalve genus, only the best genomes and transcriptomes in terms of either BUSCO scores (on the ‘metazoan_odb10’ dataset; **Manni et al., 2021**) or assembly statistics (**Tab. S1**), in order to reduce computational time. *Archivesica marissinica* (now *Calyptogena marissinica*) and *Saccostrea glomerata* were also removed, as their annotated coding sequences contain many stop codons, which prevent accurate amino acid guided alignments. Genes were clustered in orthologous groups using OrthoFinder (v2.5.5; **Emms and Kelly, 2019**) with DIAMOND ultra-sensitive and default parameters. Resulting orthogroups were splitted into SCOs using DISCO (v1.3.1; **Willson et al., 2022**), and orthogroups with at least 17 species (50 % of the species included in the bivalve reduced dataset) were retained. Amino acid and nucleotide sequences of SCOs were then aligned using Clustal Omega as implemented in TranslatorX (v1.1; **Abascal et al., 2010**), and jointly trimmed using trimAl with a gap threshold of 40 % and the removal of spurious sequences (`-resoverlap 50 -seqoverlap 50`). Eventually, orthogroups containing (i) internal stop codons, (ii) with less than 17 species left (50 % of the species included in the bivalve reduced dataset), or (iii) containing DSFGs were removed from downstream analyses. The best amino acid substitution model was inferred for each trimmed alignment using ModelFinder as implemented in IQTREE2 (model search was restricted to matrices accepted by the ‘phangorn’ R library; i.e., Blosum62, cpREV, Dayhoff, DCMut, FLU, HIVb, HIVw, JTT, JTTDCMut, LG, mtART, mtMAM, mtREV, mtZOA, rtREV, VT, WAG) and the corresponding pairwise amino acid distances were computed with the function ‘dist.ml’ from the ‘phangorn’ R package (**Schliep, 2011**). The same pipeline was also employed to obtain pairwise amino acid distances for each DSFG single-copy orthologous group. We decided to employ the pairwise amino acid distance instead of the tip-to-tip phylogenetic distance (which accounts for a more comprehensive evolutionary signal) in order to save computational time. However,

to check whether the two metrics were comparable to each other, we randomly selected 200 SCOs (including orthogroups from the DSFGs) and computed the maximum likelihood (ML) trees using IQTREE2, with ModelSelection restricted as before. Then, the tip-to-tip pairwise distances were obtained with the R package ‘adephylo’ (**Jombart and Dray, 2010**).

The distribution of amino acid distances was then built after the median values of pairwise distances of each SCO, and genes were categorised accordingly into three groups: Group 1, consisting of genes from the 1 % upper quantile of the distribution; Group 2, consisting of genes between the 1 and 5 % upper quantiles; and Group 3, consisting of all the remaining genes. Group 1 and Group 2 genes will be referred to as ‘highly-divergent genes’.

4.2.4 Mammals and *Drosophila* spp. as test datasets

To validate our approach for the study of bivalve SRG molecular evolution, we run the same analysis on two additional datasets, consisting of reference genomes of mammals and *Drosophila* species (**Tab. S4** and **S5**, respectively), whose sex-determining mechanisms are well studied and characterised. As a matter of fact, despite it is well known that SDGs tend to evolve faster than genes not involved in SD, the hypothesis has never been tested extensively across the entire phylogenetic diversity of a group: molecular evolution of SDGs and SRGs has mainly been tested on single/pairs of species or inside the boundaries of taxonomic genera (**Stothard and Pilgrim, 2003; Haerty et al., 2007; Mank et al., 2007; Mullon et al., 2012; Papa et al., 2017; Ghiselli et al., 2018**). For both mammals and fruit flies, annotated genomes were downloaded from NCBI using the command-line tool ‘datasets’, then processed using the same pipeline and scripts as before (**Fig. 4.1**).

4.2.5 GO-term enrichment

After having obtained the distributions of AASD in the three datasets (reduced bivalves, mammals, and fruit flies) and having sorted SCOs genes up into 3 groups (Group 1, Group 2, and Group 3), we performed a gene ontology (GO) enrichment analysis of genes from Group 1 and genes from Group 1 + Group 2. To do so, we firstly selected one gene per SCO, giving priority to few chosen species: (i) for bivalves, we selected genes from *Pecten maximus*, or alternatively from *C. gigas*, *Hyriopsis bialata* (now *Unio delphinus*), *Tridacna squamosa*, and *Solen grandis*; (ii) for mammals, we selected genes from *H. sapiens*, or alternatively from *Bubalus bubalis*, *Pan-*

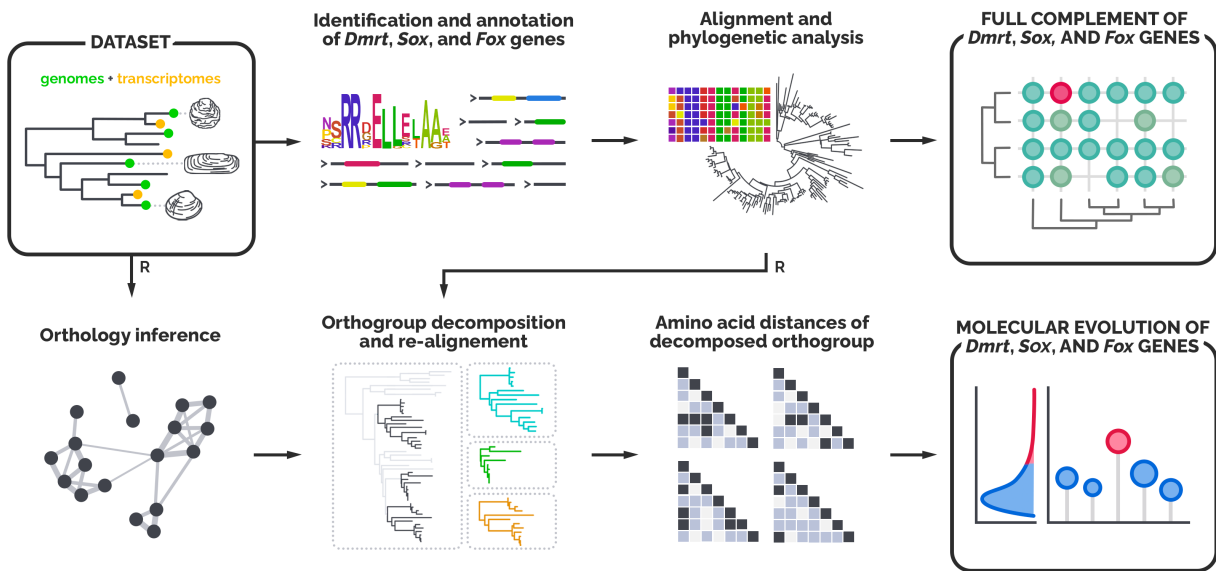


Figure 4.1 – Workflow of the analyses for the bivalve dataset. Starting from a set of both genomes and transcriptomes covering a great portion of bivalve taxonomic diversity, we first characterized the entire complement of DSFG genes (upper row). In particular, we used sequence annotation and phylogenetic tools to obtain reliable sequences and filter out any putative mis-assembled or mis-annotated sequence. Afterwards, we built a reduced set of transcriptomes and genomes (the reduced bivalve dataset, where we minimized the redundancy of congeneric species) from which to draw the molecular evolution patterns of orthologous genes (bottom row). In particular, after having obtained SCOs, we calculated the amino acid distances within each orthogroup and then built the distribution of median values. The same pipeline was also employed for the mammal and the fruit fly datasets, with just two minor differences: the starting dataset was composed of only genomes, and that the reduction step (R) was not necessary.

thera tigris, *Camelus dromedarius*, and *Monodelphis domestica*; (iii) for fruit flies, we selected genes from *D. melanogaster*, or alternatively from *Drosophila hydei*, *Drosophila pseudoobscura*, and *Drosophila suzukii*. By doing so, we ensured that each SCO was represented by one gene. Afterwards, we annotated the obtained datasets with the corresponding GO terms using the OMA browser (accessed on 18/09/2024; **Altenhoff et al., 2024**). The GO-term enrichment of Group 1 genes and Group 1 + Group 2 genes was performed with the R package ‘topGO’ with the Fisher exact test (**Alexa and Rahnenführer, 2009**).

4.3 Results

4.3.1 Genomic and transcriptomic datasets

The complete bivalve dataset consists of 29 bivalve genomes, 14 bivalve transcriptomes, and 7 outgroup genomes (5 gastropods and 2 *Octopus* spp.; **Tab. S1**). BUSCO statistics for complete single-copy genes spanned from the 64.9 % in *Modiolus modiolus* to the 99.4 % of *Perna viridis*, with a median value of 94.7 %. We were able to get at least one representative species for 11 different bivalve orders, covering a good proportion of the phylogenetic diversity of the clades Pteriomorpha, Palaeoheterodonta, and Imparidentia, and thus building the most extensive genomic and transcriptomic dataset for bivalve comparative analyses so far (**Tab. S1**). Unfortunately, no genomes or transcriptomes for Protobranchia, Archiheterodonta, and Anomalodesmata were available at the time of the project, thus we were not able to include any of those clades in our analysis. The reduced bivalve dataset (used for the orthology inference and the molecular evolution analysis; **Fig. 4.1**) consists instead of 36 genomes and transcriptomes (**Tab. S1**), and was built to retain just one species for each taxonomic genera.

The mammal dataset consists of 32 species and 1 outgroup (*Gallus gallus*, Aves; **Tab. S4**), and covers 12 major orders, while the fruit fly dataset consists of 17 species and 1 outgroup (*Anopheles gambiae*, Culicidae; **Tab. S5**), and covers 2 *Drosophila* subgenera (i.e., *Drosophila* and *Sophophora*). BUSCO statistics for complete single-copy genes were generally higher than those of bivalves, with a median of 98.3 % for mammals and of 99.8 % for fruit flies (**Tab. S4** and **S5**).

4.3.2 The Dmrt, Sox, and Fox complements in bivalves

Our annotation pipeline managed to successfully identify and annotate DSFGs in bivalves, as proved by the same analysis in mammals and fruit flies (see **Section 4.3.4**).

We retrieved four main orthology groups of Dmrt genes in bivalves (**Fig. 4.2** and **S1**; **Tab. S6**), three corresponding to the groups present in the Bilateria common ancestor (*Dmrt-2*, *Dmrt-3*, and *Dmrt-4/5*; **Mawaribuchi et al., 2019**), and one additional group with no unambiguous ortholog among reference genes, and thus putatively specific to molluscs (named *Dmrt 1-like* [*Dmrt-1L*], as per **Li, Zhang, et al., 2018**; **Evensen et al., 2022**). The majority of identified Dmrt genes are present in single-copy in each species, but *Dmrt-4/5s* show a group-

specific expansion in Palaeoheterodonta and Heterodonta, while *Dmrt-1L* is completely absent from Heterodonta. The degree of missing data for Dmrt genes in bivalves is about 35 %, with *Dmrt-2* having the highest (~56 %) and *Dmrt-4/5* the lowest (~7 %; **Tab. S7**). The coupling of ubiquitin conjugation to endoplasmic reticulum degradation (CUE)-like DM-associated (DMA) domain has been annotated in most of the *Dmrt-3* and *Dmrt-4/5* genes, while an additional DM domain has been annotated in *Dmrt-1L* genes in Mytilida and the gastropod *Pomacea canaliculata* (**Tab. S6**). Concerning Sox genes, we retrieved six main orthology groups, none of which is restricted to molluses or bivalves (**Fig. 4.2** and **S2**; **Tab. S6**). Five Sox groups (*Sox-B1/2*, *Sox-C*, *Sox-D*, *Sox-E*, and *Sox-F*) are those traditionally considered to be present in the Bilateria common ancestor (**Phochanukul and Russell, 2010**), while one has been identified outside mammals only recently (*Sox-H*, or *Sox-30*; **Han et al., 2010**). *Sox-B2* and *Sox-B1* have been grouped in the same clade, as in our phylogenetic reconstruction the former results in a paraphyletic group with the latter (**Fig. S2**), despite being traditionally recognised as a separate paralogy group in humans, fruit flies, and nematodes. The degree of missing data for Sox genes in bivalves is ~8 %, with *Sox-H* having the highest (~21 %) and *Sox-B1/2* and *Sox-C* both having no missing genes (**Tab. S7**). The Sox N-terminal signature domain was annotated for *Sox-E* genes (**Tab. S6**). Concerning Fox genes, we retrieved 27 main orthology groups (**Fig. 4.2** and **S3**; **Tab. S6**), two of which are specific to molluscs (*Fox-OG13/NA*, *Fox-OG16/NA*). Additionally, other potential mollusc-specific Fox groups have been identified, but these have been excluded from the final orthology analysis as they are present in less than half of bivalve species (see **Section 4.2**; **Tab. S6**). The two major Fox gene subgroups, Group I (monophyletic, specific to Metazoa; includes *Fox-A*, *Fox-B*, *Fox-C*, *Fox-D*, *Fox-E*, *Fox-F*, *Fox-G*, *Fox-H*, *Fox-L1*, *Fox-L2*, *Fox-Q2*) and Group II (paraphyletic, specific to Opisthokonta; includes *Fox-O*, *Fox-P*, *Fox-J2*, *Fox-J1*, *Fox-K*, *Fox-N2/3*, *Fox-N1/4*; **Larroux et al., 2008**), have been recovered, including the four Fox genes that were present in the Bilateria common ancestor (*Fox-C*, *Fox-F*, *Fox-L1*, and *Fox-Q1*; **Shimeld et al., 2010**). Two putative lineage-specific expansions have been recovered for *Fox-OG28/NA*, one regarding *Mytilus* spp. and one regarding the two Myida species (**Fig. 4.2**; **Fig. S3**). The degree of missing data for Fox genes in bivalves is about 22 %, with *Fox-H* having the highest (~42 %) and *Fox-J1* having no missing genes (**Tab. S7**). The forkhead-associated (FHA) domain was annotated for *Fox-K* genes, the *Fox-P* coiled-coil signature domain was annotated for *Fox-P* genes, while both the forkhead N- and C-terminal signature domains were annotated for *Fox-A* genes (**Tab. S6**). Regarding bivalve

species, the amount of missing data greatly differs between genomes and transcriptomes, with a mean of about 9 % and about 45 %, respectively. *Argopecten irradians concentricus*, *Mytilus coruscus* (now *unguiculatus*), and *P. maximus* have no missing data, while *Loripes orbiculatus* has the highest proportion (~64 %; **Tab. S7**).

4.3.3 Amino acid sequence divergence of Dmrt, Sox, and Fox genes in bivalves

In the reduced bivalve dataset, OrthoFinder collectively analysed >1.2 M genes distributed in 34 species. 89.4 % of these genes were placed in orthogroups, while 10.6 % were not. The number of retrieved SCOs is 5, which is drastically low but can be explained considering the mixed nature of the dataset, that is, including both genomes and transcriptomes with highly different BUSCO scores (**Tab. S1**). In order to be able to analyse a greater number of genes, we decomposed OrthoFinder orthogroups using DISCO and eventually obtained ~11 k SCOs with at least 50 % of the species. By running the same pipeline on DSFGs, we included in the AASD analysis 32 SCOs (**Fig. 4.2**) out of 33 initial Possvm-identified groups (*Fox-H* didn't meet the species occupancy threshold; **Fig. 4.3**).

From the distribution of median AASD, 112 genes were assigned to Group 1 (1 % upper quantile), 447 to Group 2 (5 % upper quantile), and 10.603 to Group 3. Most of the DSFGs (29/32) fell in Group 3 (**Fig. 4.3**), which means they have a median AASD comparable to the vast majority of other genes in bivalves (median level of the genomes). Just *Dmrt-1L*, *Sox-H*, and *Sox-F* showed higher divergences, and have been accordingly placed in Group 2. Overall, pairwise AASD proved to be a good approximation of the tip-to-tip distances ($R = 0.84, p < 2.2 \times 10^{-16}$, calculated on 200 randomly-selected trees; **Fig. 4.3C**), while it showed no influence from the alignment length ($R = 0.11$) or the number of represented species ($R = -0.23$; **Fig. 4.3D** and **4.3E**). Genes from Group 1 and Group 2 are strongly involved in cellular regulatory processes (such as those related to the metabolism of nucleic acids, proteins, and other macromolecules), but also in development and response to external stimuli, as shown by the GO-term enrichment analysis (**Tab. 4.1** and **S10**).

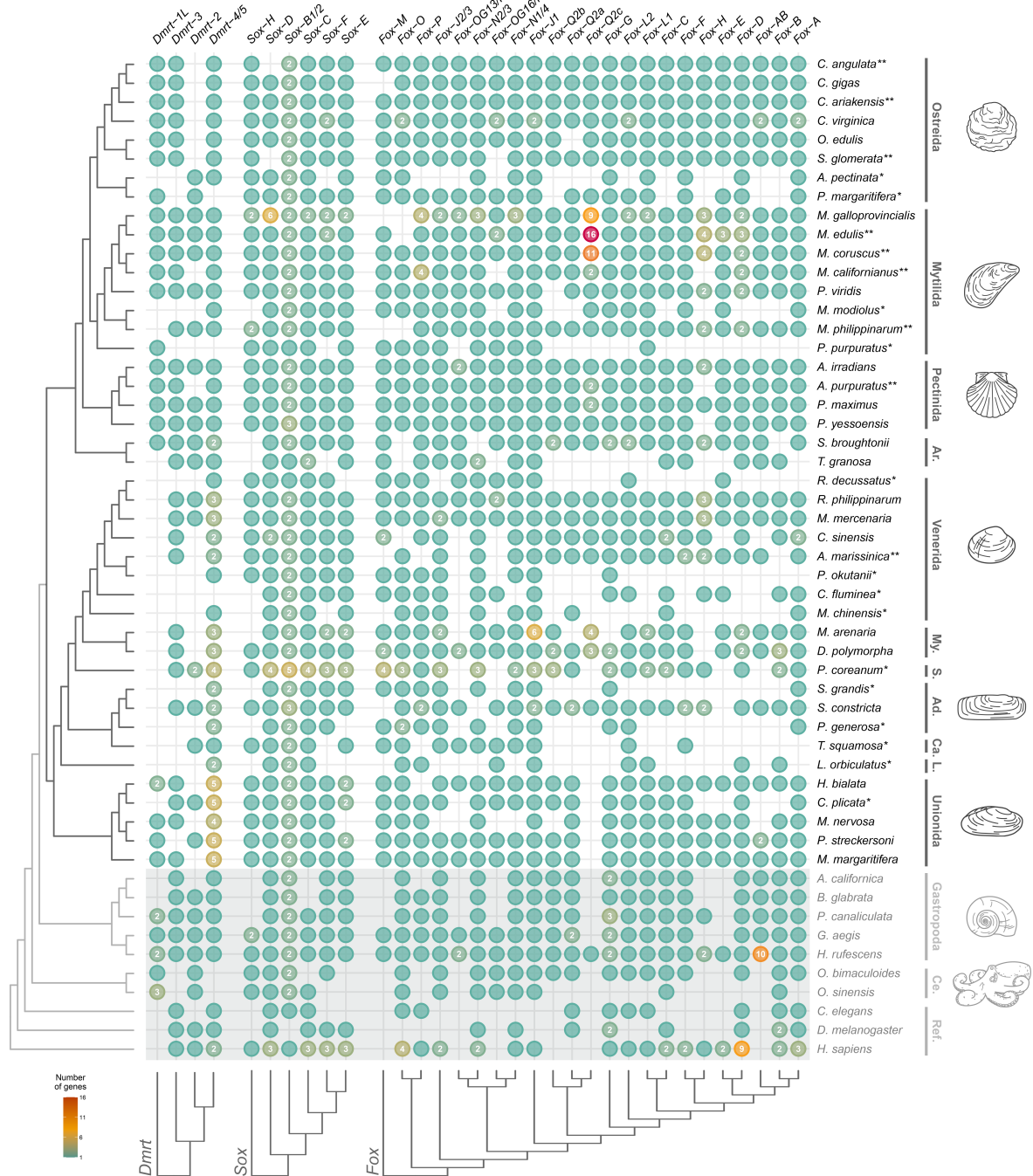


Figure 4.2 – DSFG complement in bivalves and their outgroups. Presence/absence of genes in various species are indicated by filled circles. Numbers inside each circle specify genes with 2 or more copies. The shaded area highlights non-bivalve species, belonging either to other molluscs or to the references. The phylogenetic tree of analyzed species, as inferred from literature, is shown on the left, while major taxonomic groups are reported on the right. Species represented by transcriptomic data are marked with an asterisk ('*'), and species not present in the reduced bivalve dataset are marked with two asterisks ('**'; see main text and **Fig. 4.1**); note that the two categories do not overlap. DSFG trees are shown on the bottom (full trees can be found in **Fig. S1–S3**). Full species names, along with all assembly and taxonomic information, can be found in **Tab. S1**. Ad.: Adapedonta; Ar.: Arcida; Ca.: Cardiida; Ce.: Cephalopoda; L.: Lucinida; My.: Myida; Ref.: reference genes; S.: Sphaeriida.

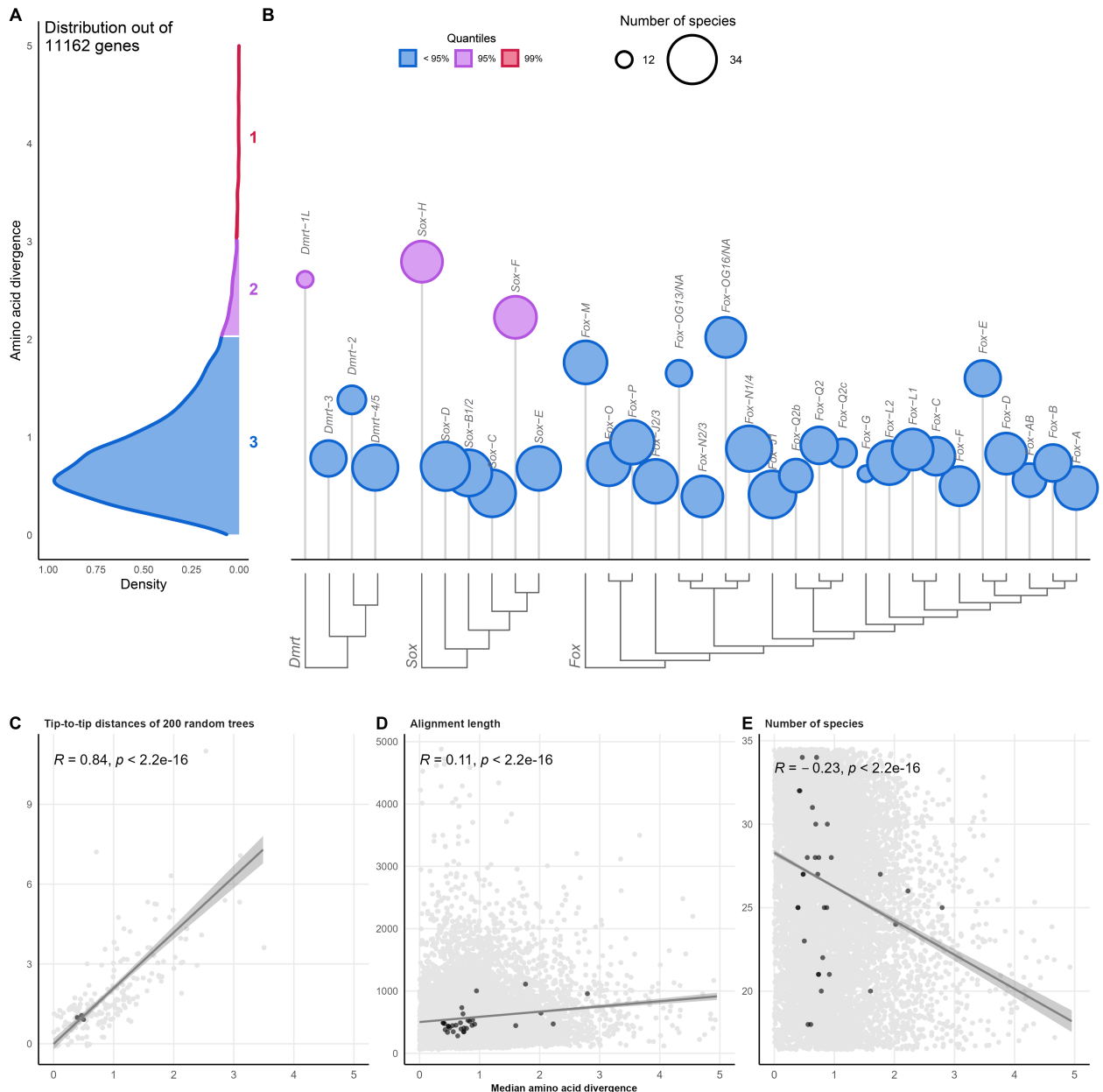


Figure 4.3 – Distribution of AASD of single-copy orthogroups in bivalves (A), including DSFGs (B), and their correlations with tip-to-tip distances (C), alignment lengths (D), and number of species (E). The distribution of AASD has been computed on the median values of pairwise distances of >11k SCOs from the reduced bivalve dataset (see main text and **Fig. 4.1**). Genes have been divided according to their median AASD value into three different groups, which are indicated by different colors and increasing numbers (Groups 1, 2, and 3). Circle heights of DSFGs show the median value of their AASD, while the size indicates the number of represented species. DSFG trees are shown on the bottom (full trees can be found in **Fig. S1** and **S3**). Darker points in C–E indicate DSFG SCOs. The correlation between the amino acid distance and the tip-to-tip distance has been computed on 200 randomly-selected orthogroups.

Table 4.1 – Top enriched GO terms for highly-divergent genes of bivalves, mammals, and *Drosophila*. The extended version of the table, which includes also the expected number of annotated genes per GO term and all the other enriched GO terms, can be accessed in [Tab. S10](#).

Dataset	GO.ID	Term	Annotated genes	Significant genes	Classic genes	Fisher
Bivalvia	GO:0060255	regulation of macromolecule metabolic process	737	59	0.0453	
	GO:0080090	regulation of primary metabolic process	673	53	0.0182	
	GO:0019219	regulation of nucleobase-containing compound metabolic process	541	41	0.0239	
	GO:0006351	DNA-templated transcription	571	39	0.0377	
	GO:0032774	RNA biosynthetic process	579	39	0.0449	
	GO:0051252	regulation of RNA metabolic process	517	37	0.0272	
	GO:0006355	regulation of DNA-templated transcription	490	35	0.0375	
	GO:2001141	regulation of RNA biosynthetic process	491	35	0.0384	
	GO:0006950	response to stress	370	33	0.0195	
	GO:0032502	developmental process	261	27	0.0445	
	GO:0006468	protein phosphorylation	345	23	0.0248	
	GO:0031325	positive regulation of cellular metabolic process	125	17	0.0080	
	GO:0010604	positive regulation of macromolecule metabolic process	151	17	0.0405	
	GO:0051172	negative regulation of nitrogen compound metabolic process	117	16	0.0081	
	GO:0051173	positive regulation of nitrogen compound metabolic process	137	15	0.0245	
	GO:0006310	DNA recombination	66	14	0.0009	
	GO:0048513	animal organ development	83	12	0.0409	
	GO:0010629	negative regulation of gene expression	78	11	0.0005	
	GO:0023051	regulation of signaling	133	11	0.0287	
	GO:0045934	negative regulation of nucleobase-containing compound metabolic process	64	11	0.0364	
	GO:0009605	response to external stimulus	90	11	0.0454	

Tab. 4.1 continued from previous page

Dataset	GO.ID	Term	Annotated genes		Significant genes	Classic Fisher
			63	11		
Bivalvia	GO:0044419	biological process involved in interspecies interaction between organisms	1,297	145	0.0006	
	GO:0006955	immune response	853	112	0.0207	
	GO:0098542	defense response to other organism	647	82	0.0000	
	GO:0045087	innate immune response	630	51	0.0466	
	GO:0001817	regulation of cytokine production	233	45	0.0000	
	GO:0042742	defense response to bacterium	642	45	0.0174	
	GO:0006954	inflammatory response	382	44	0.0000	
	GO:0019221	cytokine-mediated signaling pathway	342	44	0.0000	
	GO:0002250	adaptive immune response	402	41	0.0272	
	GO:0001819	positive regulation of cytokine production	308	37	0.0443	
	GO:0002697	regulation of immune effector process	432	35	0.0256	
	GO:0042110	T cell activation	257	34	0.0000	
Mammalia	GO:0051607	defense response to virus	491	32	0.0226	
	GO:0048232	male gamete generation	478	31	0.0280	
	GO:0007283	spermatogenesis	273	29	0.0129	
	GO:0070661	leukocyte proliferation	221	29	0.0483	
	GO:0002449	lymphocyte mediated immunity	212	25	0.0187	
	GO:0070663	regulation of leukocyte proliferation	300	24	0.0024	
	GO:0050727	regulation of inflammatory response	240	24	0.0124	
	GO:0031349	positive regulation of defense response	177	22	0.0034	
	GO:0002768	immune response-regulating cell surface receptor signalling pathway	66	17	0.0000	
	GO:0050829	defense response to Gram-negative bacterium	164	17	0.0001	
	GO:0071222	cellular response to lipopolysaccharide				

Tab. 4.1 continued from previous page

Dataset	GO.ID	Term	Annotated genes		Significant genes	Classic Fisher
			genes	genes		
Mammalia	GO:0010466	negative regulation of peptidase activity	163	16	0.0004	
	GO:0002429	immune response-activating cell surface receptor signalling pathway	164	16	0.0024	
	GO:1903555	regulation of tumor necrosis factor superfamily cytokine production	137	16	0.0124	
	GO:0071706	tumor necrosis factor superfamily cytokine production	137	16	0.0124	
	GO:0070665	positive regulation of leukocyte proliferation	132	16	0.0277	
	GO:0045089	positive regulation of innate immune response	113	16	0.0322	
	GO:0071356	cellular response to tumor necrosis factor	175	15	0.0022	
	GO:0002695	negative regulation of leukocyte activation	148	15	0.0115	
	GO:0002456	T cell mediated immunity	82	15	0.0161	
	GO:0002705	positive regulation of leukocyte mediated immunity	113	15	0.0184	
<i>Drosophila</i>	GO:0032680	regulation of tumor necrosis factor production	133	15	0.0326	
	GO:0032640	tumor necrosis factor production	133	15	0.0326	
	GO:0050866	negative regulation of cell activation	165	15	0.0405	
	GO:0000819	sister chromatid segregation	140	11	0.0293	
	GO:0070192	chromosome organization involved in meiotic cell cycle	54	9	0.0085	
	GO:0007131	reciprocal meiotic recombination	37	7	0.0007	
	GO:0007143	female meiotic nuclear division	54	6	0.0227	
	GO:0035967	cellular response to topologically incorrect protein	44	5	0.0333	
	GO:0035966	response to topologically incorrect protein	47	5	0.0427	
	GO:0007141	male meiosis I	13	4	0.0015	
	GO:0140543	positive regulation of piRNA transcription	3	3	0.0001	
	GO:0010526	retrotransposon silencing	8	3	0.0033	
	GO:0007130	synaptonemal complex assembly	10	3	0.0067	

Tab. 4.1 continued from previous page

Dataset	GO.ID	Term	Annotated genes		Significant genes	Classic Fisher
			genes	genes		
<i>Drosophila</i>	GO:0030719	P granule organization		11	3	0.0089
	GO:0071218	cellular response to misfolded protein		12	3	0.0115
	GO:0051788	response to misfolded protein		12	3	0.0115
	GO:0007135	meiosis II		15	3	0.0217
	GO:0034508	centromere complex assembly	19	3	0.0409	

4.3.4 Dmrt, Sox, and Fox genes, and amino acid sequence divergence in the test datasets

The DSFG datasets retrieved in mammals and fruit flies are far more complete than those in bivalves, and most of the already-recognised orthology groups have been identified.

In mammals, we retrieved 7 Dmrt orthology groups with about 3.1 % of missing data, 20 Sox orthology groups with about 8.1 % of missing data, and 42 Fox orthology groups with about 4.6 % of missing data (**Fig. S4A, S5, and S7; Tab. S8**). Of these, just *Sox-5* was not included in the subsequent AASD analysis, as it did not meet the 50 %-species occupancy threshold. OrthoFinder analysed about 650 M genes, and the number of SCOs used in the AASD analysis (thus resulting from the DISCO-based orthogroup decomposition pipeline) is >16 k (**Fig. 4.4A**). From the distribution of median AASD, 163 genes were assigned to Group 1, 649 to Group 2, and 15,355 to Group 3. Most of the DSFGs (66/68) fell in Group 3 (**Fig. 4.4B**), while *Sry* and *Fox-D4* showed higher divergences, and have been accordingly placed in Group 1 and Group 2, respectively. Highly-divergent genes show a strong enrichment in immune-related functions (such as innate and adaptive immune response, defence response to bacteria and viruses, lymphocyte methabolism, etc.), but also in reproductive processes (such as spermatogenesis; **Tab. 4.1** and **S10**).

Concerning *Drosophila*, we retrieved 4 Dmrt orthology groups with about 1.7 % of missing data, 7 Sox orthology groups with about 3.9 % of missing data, and 17 Fox genes with about 8.3 % of missing data (**Fig. S4B, S8, and S10; Tab. S9**). OrthoFinder analysed about 240 M, and the distribution of median AASD was built after >12k SCOS (**Fig. 4.4C**). 126 genes were assigned to Group 1, 501 to Group 2, and 11,880 to Group 3. All of the DSFGs have been used in the AASD analysis, but none of them have been placed in Group 1 or 2, that is, all the DSFGs in *Drosophila* have an AASD comparable to the median level of the genome (**Fig. 4.4D**). Genes of Group 1 and Group 2 show a GO-term enrichment in meiotic processes, such as chromosome/chromatid organisation, and retrotransposon silencing (**Tab. 4.1** and **S10**).

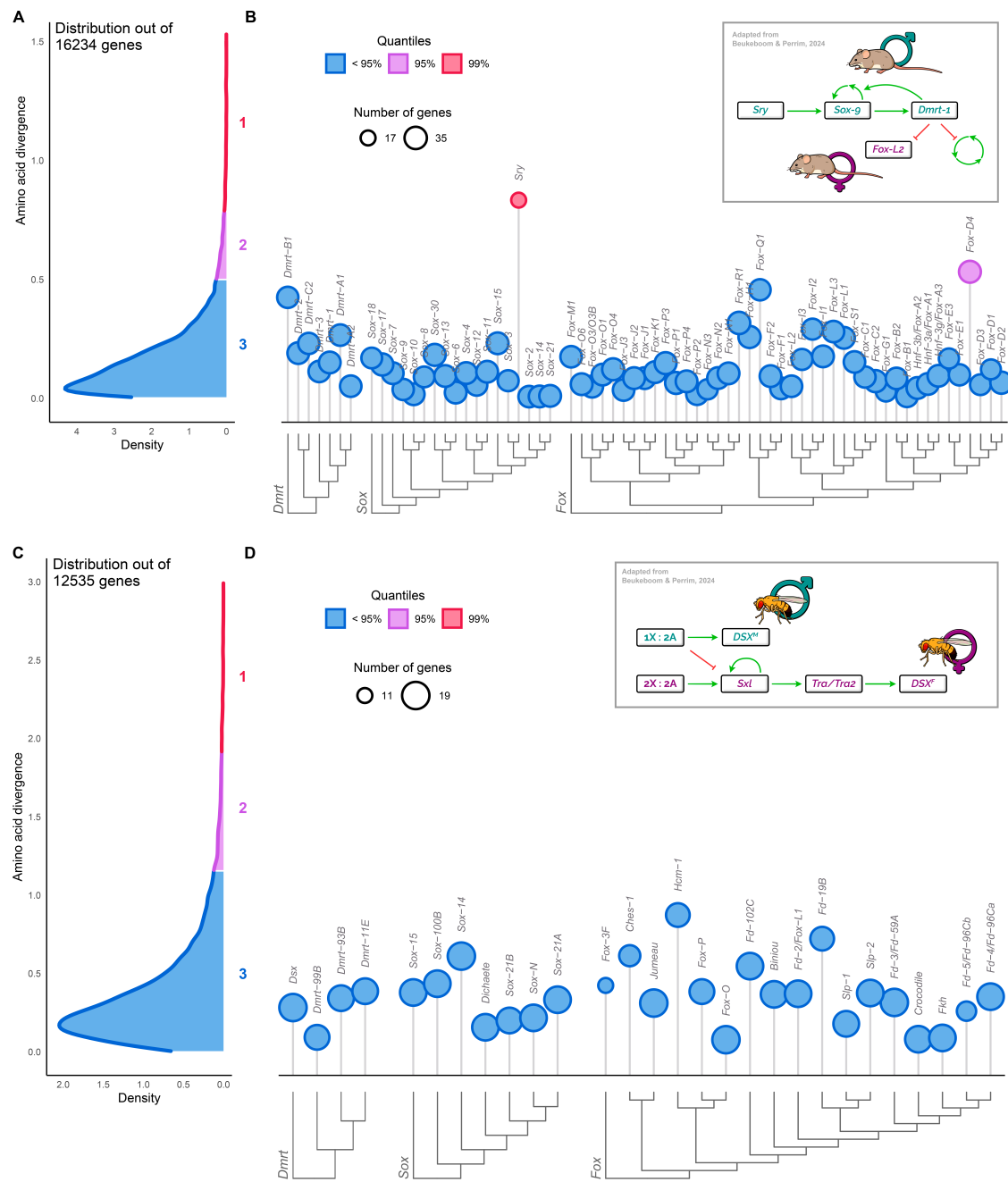


Figure 4.4 – Distribution of AASD of single-copy orthogroups in Mammalia (A) and *Drosophila* (C), including DSFG (B-D). The distributions of AASD in mammals and fruit flies have been computed on the median values of pairwise distances of over 16k and 12k SCOs, respectively. Genes have been divided according to their median AASD value into three different groups, which are indicated by different colors and increasing numbers (Groups 1, 2, and 3). Circle heights of DSFGs show the median value of their AASD, while the size indicates the number of represented species. DSFG trees are shown on the bottom (full trees can be found in Fig. S5–S7 for mammals and in Fig. S8–S10 for fruit flies). Insets: scheme of the sex-determination molecular pathways in *Mus musculus* and in *Drosophila melanogaster*, with shown the main genes involved (adapted from Beukeboom and Perrin, 2014). Green arrows indicate transcription activations, red arrows indicate transcription suppressions. X: sex chromosomes; A: autosomal chromosomes; *DSX^{M/F}*: *DSX* splicing variants present in males or females, respectively.

4.4 Discussion

4.4.1 A new manually-curated and phylogenetic-based reference dataset of Dmrt, Sox, and Fox genes in bivalves

The annotation and characterisation process of a gene family in a certain clade of organisms may harbour many overlooked challenges (**Vizueta Moraga et al., 2020**). For example, the presence of highly-conserved catalytic domains may hamper the correct identification of the components of a gene family because of insufficient phylogenetic signal, as it is the case for Hox and ParaHox genes and their homeobox motif (**Baldwin-Brown et al., 2018; Nicolini, Martelossi, et al., 2023**). Conversely, the components of dynamic gene families characterised by abrupt and sequential duplication events may be difficult to sort into separate groups. As a matter of fact, varying levels of sequence heterogeneity and gene copy numbers makes the inference of orthologous groups hard, as for certain clans of the P450 gene family (**Dermauw et al., 2020**). Regardless of the causes, having a solid and wide phylogenetic context in which to study gene duplications and losses, and orthology relationships, is crucial to overcome these difficulties. In the same way, manual curation and visual inspection of multiple sequence alignments, phylogenetic trees, and gene structures (in terms of domain composition, start and stop codons, and other feature representations) is helpful, despite being time-demanding and possibly low reproducible. In this study, we characterised the full complement of DSFGs in the vast class of bivalves, by leveraging sequence domain annotation, phylogenetics, and manual curation of the dataset. Our aim was to obtain the most reliable gene complements as possible, combined with a vast taxonomic dataset, a solid phylogenetic inference, an openly-available dataset of gene sequences, and a reproducible pipeline for the annotation of gene identity. By doing so, we want to provide a reliable resource for future studies of DSFGs, either focused on bivalves or generally in Metazoa.

Concerning the Dmrt gene family, we identified orthologs of the vertebrate *Dmrt-2*, *Dmrt-3*, and *Dmrt-4/5* (or *A1/A2*; **Fig. 4.2** and **S1**; **Tab. S6**), which are also expected to have been present in the Bilateria common ancestor (**Mawaribuchi et al., 2019**). **Wang et al. (2023)** found that *Dmrt-4/5* is duplicated in *Mercenaria mercenaria* and *Cyclina sinensis* (Venerida), and in *Dreissena polymorpha* (Myida), and we confirm this result by tracing back the duplication event to the split between Palaeoheterodonta (here represented by Unionida) and Heterodonta (here represented by Venerida, Myida, Sphaeriida, Adapedonta, Cardiida, and Lucinida; **Fig. 4.2**).

Furthermore, we confirm *Dmrt-1L* to be present in many bivalve species (mainly belonging to the Ostreida, Pectinida, Mytilida, and Unionida orders; **Fig. 4.2**), as well as in gastropods and *Octopus*. Though, our phylogenetic analysis did not retrieve any unambiguous orthology relationship among *Dmrt-1L* and either vertebrate *Dmrt-1* or *Drosophila dsx* genes, as instead it was proposed in previous works (Li, Zhang, et al., 2018; Evensen et al., 2022). As a matter of fact, the amino acid sequence of the *Dmrt-1L* DM domain does not recall that of any other Dmrt gene. Furthermore, it must be considered that various phylogenetic analyses have recovered both *Dmrt-1* and *dsx* genes to be restricted to vertebrates and arthropods, respectively (Wexler et al., 2014; Mawaribuchi et al., 2019; Panara et al., 2019), that is, they do not have any direct ortholog outside their relative clades. Thus, if *Dmrt-1L*, *dsx*, and *Dmrt-1* are true orthologs, their origin would need to be placed at least in the Bilateria common ancestor, which seems however to be not the case. All considered, we thus confirm that *Dmrt-1L* is not orthologous to *Dmrt-1* and *dsx* and is rather a mollusc-specific gene (Evensen et al., 2022). The monophyly of the group is not supported by the phylogenetic tree inferred with Dmrt genes from molluscs and the reference species (**Fig. S1**); though, it is recovered when analysing just genes from mollusc species (**Fig. S11**). To this regard, we speculate that in our analysis, the difficulty in obtaining the monophyly of *Dmrt-1L* genes may have arisen primarily because of the many *C. elegans*-restricted genes (**Tab. S3**), which are placed among the other bivalve genes (**Fig. S1**), but also because of the high AASD of *Dmrt-1L* genes (see **Section 4.4.2**), which hampers a straight-forward phylogenetic reconstruction. Furthermore, our broad-context analysis allowed us to identify some cases of incorrect gene identification in bivalves, which have arisen because of erroneous or ambiguous annotations in previous works, as a result of limited datasets or analyses. For example, the scallop-specific cluster of Dmrt genes retrieved by Wang et al. (2023) rather belongs to the *Dmrt-1L* group, while the classification of Dmrt genes in *Crassostrea* species provided by Zeng et al. (2024) needs to be revised following the one of this work (*Dmrt-1* genes *sensu*-Zhang are *Dmrt-4/5*; *Dmrt-2* genes *sensu*-Zhang are *Dmrt-3*; *Dmrt-3* genes *sensu*-Zhang are *Dmrt-1L*; hence, *Crassostrea* species do not have *Dmrt-2* genes).

For what concerns the Sox gene family, bivalves (or molluscs) do not show any major clade-restricted gene, as only the five Bilateria-specific Sox groups (*Sox-B1/2*, *Sox-C*, *Sox-D*, *Sox-E*, and *Sox-F*) and *Sox-H* have been identified (**Fig. 4.2** and **S2**; **Tab. S6**), in accordance with previous findings (Evensen et al., 2022; Wang and Nie, 2024; Yu et al., 2017). *Sox-B1/2* is

clearly made up of two subgroups (i.e., *Sox-B1* and *Sox-B2*), as expected, but their respective identity could not be unambiguously established, as *Sox-B1/2* genes of reference species do not form separate clusters (**Fig. S2**). Even when inferring the phylogenetic tree only of components of the *Sox-B1/2* group from molluscs and reference species, the identity can not be properly established (**Fig. S12**).

Compared to Dmrt and Sox genes, the Fox gene family appears as the most dynamic in terms of gene presence/absence, as already shown by other works (**Wu et al., 2020; Schomburg et al., 2022; Seudre et al., 2022**). Our phylogenetic analysis successfully recovered Group I and Group II of Fox genes (**Larroux et al., 2008**), which include the four Fox genes that were present in the Bilateria common ancestor (*Fox-C*, *Fox-F*, *Fox-L1*, and *Fox-Q1*; **Fig. 4.2** and **S3**; **Tab. S6**; **Shimeld et al., 2010**). To our knowledge, this is the first broad-taxonomic identification and classification of Fox genes in bivalves, as up to now they have been systematically characterised only in *C. gigas* (**Yang et al., 2014**), *Patinopacten yessoensis* (now *Mizuhopecten yessoensis*; **Wu et al., 2020**), and *Ruditapes philippinarum* (**Liu et al., 2024**). Firstly, our analysis confirms the absence in molluscs of *Fox-I*, *Fox-Q1*, *Fox-R*, *Fox-S* (**Fig. S3**), which are in fact thought to have emerged with the diversification of deuterostomes or vertebrates (**Yang et al., 2014; Wu et al., 2020; Schomburg et al., 2022; Seudre et al., 2022**). Furthermore, we have found many Fox groups that appeared as mollusc-specific and/or still-unnamed at a first analysis (*Fox-OG2/NA*, *Fox-OG39/NA*, *Fox-OG15/NA*, and *Fox-OG28-NA*; **Tab. S6**). However, a more in-depth investigation revealed a different scenario. *Fox-OG2/NA* appears close to the human *Fox-M* gene in the phylogenetic tree, but they do not form a monophyletic group (**Fig. S3**). However, by comparing *Fox-OG2/NA* sequences and phylogenetic tree with those analysed by **Yang et al. (2014)**, **Wu et al. (2020)**, **Schomburg et al. (2022)**, and **Seudre et al. (2022)**, it appears clear that this group of Fox genes is indeed *Fox-M*. However, our analysis has failed to retrieve a monophyletic relationship among bivalve and human *Fox-M* genes, even when inferring a tree with just *Fox-J2*, *Fox-M*, *Fox-O*, and *Fox-P* complements (**Fig. S13**), which belong to the same Fox group. Regarding the *Fox-OG39/NA* group, it does not have any homolog in reference species (**Fig. S3**) but is found to belong to the *Fox-AB* group by sequence comparison with previous works (**Yang et al., 2014; Wu et al., 2020; Seudre et al., 2022**). *Fox-AB* was formerly described only in the sea urchin *S. purpuratus* and the lancelet *Branchiostoma floridae* (**Tu et al., 2006; Yu et al., 2008**), but was later identified also in several Spiralia lineages, including molluscs (e.g., **Yang et al., 2014; Wu et al., 2020; Seudre et al., 2022**). A similar

situation concerns *Fox-OG15/NA* and *Fox-OG28/NA*, which again could not be named based on orthology relationships with the reference species genes (**Fig. S3**), but actually represent two lineage-specific expansions of the *Fox-Q2* group (named *Fox-Q2b* and *Fox-Q2c*), as already appointed in previous studies (**Yang et al., 2014; Wu et al., 2020**). This observation fits within the wider context of the *Fox-Q2* group expansion in Bilateria and, particularly, in Spiralia, that led to remarkable differences in their gene copy numbers across various clades (**Seudre et al., 2022**). Two additional Fox genes have been previously identified in bivalves, and were named *Sox-Y* and *Sox-Z* (**Yang et al., 2014; Wu et al., 2020**). In our analysis, these Fox groups were at first identified as *Fox-OG13/NA* and *Fox-OG16/NA*, after sequence comparison of Fox genes from *C. gigas* and *P. yessoensis*. On one hand, *Fox-Y* was firstly identified in *S. purpuratus* (**Tu et al., 2006**) and only recently in a few bivalve species (**Yang et al., 2014; Wu et al., 2020**). However, when analysing bivalve and *S. purpuratus* Fox genes, we failed in retrieving such a clear orthology relationship, as *S. purpuratus* *Fox-Y* does not fall within the phylogenetic range of bivalve *Fox-OG13/NA*, which contains the supposed *Fox-Y* orthologs (**Fig. S14**). Also, the forkhead domains of *Fox-OG13/NA* genes were annotated as ‘forkhead domain P’ (**Tab. S6**). On the other hand, *Fox-Z* was firstly identified in bivalves and in several other protostomes, thanks to a phylogenetic work including the brachiopod *Lingula unguis*, the annelid *Capitella teleta*, the scorpion *Centruroides sculpturatus*, and the centipede *Strigamia maritima* (**Wu et al., 2020**). However, later works have not recovered this Fox gene group, even when analysing annelids (**Seudre et al., 2022**) and panarthropods (**Schomburg et al., 2022**) in a more dedicated effort. In this case, the forkhead domains were annotated as either a generic ‘forkhead domain’ or a ‘forkhead domain Q2’ (**Tab. S6**). All considered, we argue that bivalves possess two additional Fox groups (here *Fox-OG13/NA* and *Fox-OG16/NA*; **Fig. 4.2** and **S3**; **Tab. S6**) which are shared with other mollusc species, as revealed also by other authors. However, given the discordant results of the phylogenetic hypothesis and domain annotation, we think that a more thorough investigation on their orthology relationships with Fox genes from other Metazoa is needed, and thus we chose to not employ their former names *Fox-Y* and *Fox-Z*.

Besides the DSFG groups discussed so far, it must be also considered that many orphan genes have been identified (**Fig. S1** and **S3**; **Tab. S6**). For example, **Wu et al. (2020)** identified a duplication event of *Fox-H* genes in *C. gigas*, which has been recovered also in our analysis for the entire Ostreida clade (*Fox-OG36/NA*; **Fig. S3**). Similarly, a gene orthology group putatively

specific to Pteriomorphia has been identified among Sox genes (*Sox-OG1/NA*). Of course, these genes deserve as much attention as their widely-distributed paralogs, as they may constitute true group-specific expansions and may play fundamental roles in some biological processes. However, they have not been discussed here or included in **Fig. 4.2** for clarity purposes, but they are freely available in supplementary materials.

Overall, our analysis clearly shows the importance of adopting a wide-angle approach when characterising the members of a gene family, especially for large ones such as the Fox genes (**Schomburg et al., 2022**). As a matter of fact, the presence of duplication events and orphan genes needs to be addressed with a broad taxonomic dataset, in order to account for possible mis-annotations, gene phylogenetic mis-placements, and sequence heterogeneity. Additionally, many reference species need to be included for the gene identification process, in order to consider distantly-related genes and obtain a solid annotation. Our gene annotation pipeline also resulted to be very solid, even with non-model organisms and sub-optimal genomic and transcriptomic resources, as they are those of bivalves. As a matter of fact, by running the same pipeline on two additional datasets composed of mammal and fruit fly genomes, we were able to obtain high-quality orthology groups in accordance with previous knowledge on the clades (**Fig. S5 and S10; Tab. S8 and S9**), with little or no manual curation needed. Furthermore, this represents also the first broad analysis of DSFGs in both mammals and fruit flies, as so far attention has been mainly dedicated to single well-studied organisms or little clades (e.g., **Jackson et al., 2010**).

4.4.2 High amino acid sequence divergence identifies putative sex-determining genes

Sex-biased genes tend to evolve more rapidly than unbiased genes at the level of their protein sequences. Accelerated rates have been observed in both male-biased genes (reviewed in **Parsch and Ellegren, 2013; Grath and Parsch, 2016**) and female-biased genes (e.g., **Papa et al., 2017; Ghiselli et al., 2018**), but also in SRGs and primary SDGs (**O’Neil and Belote, 1992; Whitfield et al., 1993; de Bono and Hodgkin, 1996**). For example, it has been shown that *Dm-W*, *Dmy*, and *Sry* (which are SDGs in the African clawed frog *X. laevis*, in the medaka fish *O. latipes*, and in eutherians, respectively) all have higher substitution rates than their paralogues (*Dmrt-1* for *Dm-W* and *Dmy*, *Sox-3* for *Sry*), particularly when considering their DNA-binding domains

(Mawaribuchi et al., 2012). Similarly, both a burst of positive selection and a relaxation of purifying selection has been detected in *Drosophila Sxl* in correspondence with its recruitment at the top of the sex-determining cascade. The same signs of relaxed purifying selection have been found in the downstream targets of *Sxl*, that is, *tra* and *dsx*, despite no evidence of positive selection has been detected (Mullon et al., 2012).

Considering these shared features of SRGs and SDGs, we decided to look for signs of accelerated sequence evolution in DSFGs of bivalves, in order to evaluate if any of them could be associated with SD by employing the tools of molecular evolution. However, we wanted to analyse patterns of sequence evolution not only among putative SRGs and their close paralogs (as done for Dmrt genes in Chapter 3), but also considering the genomic context in which these genes evolve. In fact, our aim was to check whether higher rates of sequence evolution of SRGs hold true also when compared to other genes not involved in SD and not belonging to the same gene family. To do so, we obtained the AASD median values of >11 k SCOs from bivalve genomes (Fig. 4.3A), in order to build a statistical distribution to be used as a reference: if SRGs/SDGs (in this case, DSFGs) truly evolve faster than other genes, we may expect them to fall within the 5 % (or even 1 %) upper quantile of the distribution (Fig. 4.3B), i.e., within highly divergent genes (Group 1 and Group 2 genes of the distribution; see Section 4.2). We chose to use the AASD as a metric of sequence evolution (instead of the tip-to-tip distances of phylogenetic trees, which account for more comprehensive evolutionary models) in order to save computational time. As a matter of fact, the AASD median values proved to be a good approximation of the tip-to-tip median distances in 200 randomly-selected genes (Fig. 4.3C; $R = 0.84, p < 2.2 \times 10^{-6}$).

Among DSFGs, three fell within the 5 % upper quantile, namely *Dmrt-1L*, *Sox-H*, and *Sox-F*. Interestingly, *Dmrt-1L* and *Sox-H* have been already proposed to be involved in the male SD pathway of *C. gigas* (inset in Fig. 4.3B; Zhang et al., 2014), on the basis of DGE analyses. Specifically, *Sox-H* (*CgSoxH*) would play a major role in *C. gigas* SD, by interacting with *Dmrt-1L* (*CgDsx*) and determining the onset of the male phenotype development; at the same time, both *Sox-H* and *Dmrt-1L* would inhibit *Fox-L2* (*CgFoxL2*), which instead is necessary to start the female phenotype development. *Dmrt-1L* and *Sox-H* have been appointed several other times to be involved in male-gonad development and differentiation, through DGE (e.g., Teaniniuraitemoana et al., 2014; Capt et al., 2018; Afonso et al., 2019), mRNA-ISH (e.g., Naimi et al., 2009; Li, Zhang, et al., 2018; Liang et al., 2019; Yue et al., 2021) and RNA

interference (mRNA-ISH; **Liang et al., 2019; Sun et al., 2022**). Therefore, the high AASD of *Dmrt-1L* and *Sox-H* is coherent with previous works, strengthening their role as putative SRGs.

The relationship between high gene AASD and the involvement in SD is particularly enforced when looking at the patterns of AASD in the test datasets, which corroborates the solidity of our analysis: (i) from one side, in the mammal dataset—which represents a strictly genetic SD system, thus with a master and rapidly-evolving SDG, one of the genes from the 5 % upper quantile of the distribution is *Sry* (**Fig. 4.4A** and **4.4B**), the male sex-determining gene in eutherians (**inset** in **Fig. 4.4B**); (ii) from the other side, in the fruit fly dataset—which represents a chromosomal SD system, thus without any expected difference in the rates of sequence evolution among SRGs, none of the DSFG exhibit significantly high AASD (**Fig. 4.4C** and **4.4D**), including the downstream effector *dsx* (**inset** in **Fig. 4.4D**). Also *Sxl* and *tra*, both involved in the SD pathway of *Drosophila* (**inset** in **Fig. 4.4D**) do not belong to the group of highly-divergent genes, as they have a mean amino acid divergence of about 0.09 and 0.9, respectively (Group 3; **Fig. 4.4D**). Therefore, it can be argued that both *Dmrt-1L* and *Sox-H* may not only be SRGs, but may participate in bivalve SD as primary SDGs, which is reflected in their high AASD, as it is observed for *Sry* in mammals. As a matter of fact, if they were involved in SD just as intermediate actors of the signalling cascade, then we should have not observed a high AASD, as *Drosophila Sxl*, *tra*, and *dsx* seem to suggest. Overall, these patterns of molecular evolution concerning SRGs and SDG are also supported by the way SD regulatory networks evolve. As a matter of fact, it has been proposed that the sex-determining cascades tend to arise and be established with a bottom-up mechanism (**Wilkins, 1995; Mullon et al., 2012; Beukeboom and Perrin, 2014; Capel, 2017**). This means that the regulative relationships among genes at the bottom of the cascade are settled up prior to the regulative relationships among genes at the top and, consequently, upstream regulators are progressively recruited to fine-tune diverse SD signals. These evolutionary patterns eventually produce gene-regulatory networks in which the divergence of the upstream triggers is higher than that of downstream effectors, in terms of both identity and sequence composition (**Beukeboom and Perrin, 2014**). This mechanism has been proposed for *Drosophila* species (**Mullon et al., 2012**), *C. elegans* (**Stothard and Pilgrim, 2003**), and vertebrates, despite in the latter case has been questioned several times (reviewed in **Capel, 2017**).

At this point, two main objections can be moved against our approach: (1) the distribution

of AASD is not appropriate for this kind of inference, as it does not represent the true gene evolutionary (or substitution) rates (which instead are those usually employed when dealing with SRGs and SDGs); (2) the three datasets are not comparable one to each other, as they take into consideration very different animal groups, with different taxonomic rankings and different divergence times (thus, the patterns of AASD are the products of other confounding factors not directly related to SD). Concerning the first objection, we are aware that the AASD does not represent the evolutionary rate itself, but rather its product. However, the two features are tightly linked, as on the long term highly-divergent proteins tend to be produced by genes with high evolutionary (or substitution) rates (**Echave et al., 2016**). By performing a GO-term enrichment, it emerged that highly-divergent genes of the mammal dataset are mainly involved in the immune response and male spermatogenesis (**Tab. 4.1** and **S10**), which are two processes notoriously connected with rapid sequence evolution (i.e., higher evolutionary rates; **Swanson and Vacquier, 2002**; **Murat et al., 2023**; **Vinkler et al., 2023**). Similarly, highly-divergent genes from the fruit fly dataset show an enrichment for GO-terms associated with meiotic-related functions (such as the formation of the synaptonemal complex by the products of *c(2)M*, *c(3)G*, *corona*, and *corolla* genes; **Tab. 4.1** and **S10**), which again are known to be rapidly evolving (**Hemmer and Blumenstiel, 2016**). In other words, the test datasets—which include well-studied and characterised model systems, allow us to directly link the high AASD (as computed in this work) with high rates of sequence evolution (as found in previous works), as they represent well-studied and characterised model systems. This consideration can thus be extended also to the bivalve dataset: highly-divergent genes in terms of AASD, which include some DSFGs and show an enrichment for GO-terms associated to macromolecule metabolism and morphological development (**Tab. 4.1** and **S10**), are also genes with accelerated substitution rates.

Concerning the second objection, we chose two test datasets with different characteristics as we wanted to check the extent of our hypothesis, that is, molecular evolution can be used to look for putative primary SDGs in taxonomic-wide analyses. The difference in divergence times and taxonomy ranks for bivalves and therians (Late Cambrian, about 498 million years ago [Mya], **Song et al., 2023**; and Early Mesozoic, 166–123 Mya, **Álvarez-Carretero et al., 2022**, respectively) seems to not influence the sequence diversity of SRGs, as both *Dmrt-1L/Sox-H* for bivalves and *Sry* for mammals exhibit high AASD with respect to their own distributions, regardless of their age. *Dmrt-1L* and *Sox-H* (which are mollusc- and Bilateria-specific, re-

spectively) are undoubtedly older than *Sry* (which, instead, emerged in the Theria common ancestor; **Foster et al., 1992**), but each of them can be considered a highly-divergent gene in bivalves and mammals, respectively (i.e., genes that are included in the 5 % upper quantile of bivalve and mammal AASD distributions). Conversely, the difference in divergence times and taxonomic ranks for *Drosophila* (Paleocene/Eocene boundary, about 56 Mya; **Russo et al., 2013**) may seem to be influencing the results for the dataset, resulting in a false negative. In other words, it can be argued that: (i) the genes included in the SD cascade of *Drosophila* (such as *Sxl*, *tra*, and *dsx*; **inset** in **Fig. 4.4D**) have a high AASD, which however has not been detected by our methodological approach (for example, this may be traced back to the young diversification age of *Drosophila* species if compared to bivalves); (ii) the species included in the analysis are all congeneric, thus the sequence differentiation of SRGs may exist not at the amino acid level but at the nucleotide one. To better disentangle this issue and further discuss the fruit fly dataset, we repeated the analysis of the AASD only on species of the *Crassostrea* genus (*C. gigas*, *Crassostrea angulata*, *Crassostrea ariakensis*, and *Crassostrea virginica*), which are much younger (Middle Cretaceous, less than 100 Mya; **Qi et al., 2023**), thus comparable to *Drosophila*. Results showed that, even when analysing a smaller bivalve dataset, encompassing only 4 species of recent origin, the high AASD of *Dmrt-1L* persists, that is, *Dmrt-1L* is still grouped together with highly-divergent genes (**Fig. S15**). The same has not been recovered for *Sox-H*, which fell in genes from Group 3 (the group corresponding to the 95 % interval of the AASD distribution) but still have the second highest AASD median value among DSFGs (**Fig. S15**).

Of course we should not expect that highly-divergent genes are only those involved in SD, but may participate also in other processes (as discussed earlier and shown by GO-term enrichments; **Tab. 4.1** and **S10**). Besides the genes of interest for SD (*Dmrt-1L/Sox-H* for bivalves, and *Sry* for mammals), also other components of the DSFG families have been retrieved with a high AASD, despite they have never been linked directly to SD so far: *Sox-F* in bivalves (**Fig. 4.3B**) and *Fox-D4* in mammals (**Fig. 4.4B**). This implies that our approach can't be used to unambiguously identify SDGs alone, as high AASD is exhibited also by many other genes. Instead, the analysis is meant to be used to detect highly-divergent genes and, subsequently, by comparison with literature and a more thorough and focused functional investigation, putative SDGs among them. In this sense, the mammal dataset exemplify the importance of putting the results of our pipeline (as those of any other comparative genomics analysis) into the correct

evolutionary and genomic context: among DSFGs of mammals, two genes exhibit high AASD, one of which is directly related to SD (*Sry*), while the other has a function connected with neural development (*Fox-D4*; **Klein et al., 2013**). Thus, the high AASD may arise either because of the involvement in the upper SD pathway or because of other life-history traits connected with the gene, respectively. Regarding bivalves, *Dmrt-1L* and *Sox-H* show a sharp connection with SD as a putative primary SDG, either when considering their molecular evolutionary features or when looking at their gene expression and possible function in gonad development (**Naimi et al., 2009; Teaniniuraitemoana et al., 2014; Zhang et al., 2014; Capt et al., 2018; Li, Zhang, et al., 2018; Afonso et al., 2019; Liang et al., 2019; Yue et al., 2021**). It is difficult to further speculate on the actual involvement in SD of *Dmrt-1L* and *Sox-H* without any additional information on their biology. Nonetheless, molecular evolution proves to be a valuable tool to investigate genes putatively involved in SD, and to identify major targets onto which dedicate future research effort.

4.5 Conclusions

Genes functioning in reproductive processes, and particularly SD, are often among the most variable in animal genomes, in terms of both sequence composition and regulatory interactions (**Swanson and Vacquier, 2002; Bachtrog et al., 2014**). Such high evolutionary rates may be traced back both to adaptive evolution (either as natural or sexual selection) or to non-adaptive processes (**Vicoso and Charlesworth, 2006; Meisel and Connallon, 2013; Parsch and Ellegren, 2013; Grath and Parsch, 2016**), and often results in striking differences in reproductive and sexual systems even among closely-related species. In the present work we took advantage of this characteristic to identify SDGs in bivalves among the DSFG families. By comprehensively analysing the phylogenetic history and AASD in a broad taxonomic dataset, we appointed *Dmrt-1L* and *Sox-H* as putative SDGs, thus confirming results in previous works that found them to be transcribed in a male-biased manner and/or strongly involved in male-gonad formation (**Naimi et al., 2009; Teaniniuraitemoana et al., 2014; Zhang et al., 2014; Capt et al., 2018; Li, Zhang, et al., 2018; Afonso et al., 2019; Liang et al., 2019; Yue et al., 2021**). Future studies would now need to further investigate their evolutionary history. For example, considering that SRGs tend to accumulate in the genomic neighbourhood where primary SDGs are located (**Capel, 2017**), analysing the genomic location of DSFGs in bivalve genomes may

provide enlightening results. Similarly, revealing the genetic interactions of *Dmrt-1L* and *Sox-H*, through functional and genome editing assays, would undoubtedly benefit our understanding of their role in the sexual processes of bivalves.

Chapter 5

Localisation of three sex-related genes and the germline marker Vasa/Vasa in the early developmental stages of *Mytilus* *galloprovincialis*

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5.1 Introduction

Despite the huge socio-economic and scientific importance of bivalves, the knowledge concerning the genetic and molecular bases of their sex determination (SD) system is scarce and overlooked (**Breton et al., 2018; Nicolini, Ghiselli, et al., 2023**). Several components of the Dmrt, Sox, and Fox gene (DSFG) families have been appointed as directly involved in SD by many works, mainly thanks to differential gene expression (DGE) analyses (e.g., **Milani et al., 2013; Zhang et al., 2014; Capt et al., 2018; Shi et al., 2018**), mRNA/protein visualisation (**Li, Liu, et al., 2018; Liang et al., 2019; Wang et al., 2020; Sun et al., 2022**), RNA interference (RNAi; **Liang et al., 2019; Wang et al., 2020; Sun et al., 2022**) and quantitative real-time polymerase chain reaction (qRT-PCR; **Li, Liu, et al., 2018; Liang et al., 2019; Wang et al., 2020; Sun et al., 2022**). For example, **Li et al. (2018)** found that *Fox-L2* and *Dmrt 1-like* (*Dmrt-1L*) are predominantly transcribed in ovaries and testes, respectively, of the Yesso scallop *Patinopacten yessoensis*, and that they contribute to establish the sexual identity of immature follicles at the molecular level prior to the morphological level. **Liang et al. (2019)** showed that *Sox-2* is involved in the differentiation of male gonads and spermatogenesis of the scallop *Chlamys farreri*, and that the knocked-out phenotype results in severe loss of both germ-cell mass and spermatogonia. **Wang et al. (2020)** speculated that *Fox-L2* is involved in the sex differentiation of female gonads in the freshwater mussel *Hyriopsis cumingii*. Overall, considerable effort has been made to characterise the transcription patterns of DSFGs of interest during the adult stage of bivalves, covering various reproductive phases, while little attention has been given to the embryo and larval stages. Nonetheless, early animal development may represent a crucial moment to the establishment of the sexual identity, as the transcription of sex-determination related genes (SRGs) and SD itself begins much earlier than the onset of gonad development and differentiation (even as early as the zygote formation; **Richardson et al., 2023**). In mammals, for example, the transcription of SRGs can be detected during the embryo preimplantation stage (before 4.5 days post fertilization [dpf]; reviewed in **Richardson et al., 2023**), while *Sex-determining region of chromosome Y* (*Sry*) realises its function as the male sex-determining gene (SDG) at 10.5 days post coitum (**Beukeboom and Perrin, 2014**). In *Drosophila melanogaster*, the early female splicing variant of *Sex-lethal* (*Sxl*)—which is the top regulator of the SD cascade and is activated by a mechanism of chromosome counting (**inset** in **Fig. 4.4D**), is transcribed during the syncytial stages of the embryo (i.e., before 2 hours post fertilization [hpf]; **Salz and Erickson, 2010**), when it establishes the sexual identity of the embryo through a cell-autonomous

mechanism. Therefore, the study of bivalve SD necessarily requires to consider also the early stages of embryonic and larval development, in order to obtain a comprehensive scenario of the process. Among bivalves, several species may constitute a model system particularly suitable to study the SD process during embryogenesis, because of the presence of the doubly uniparental inheritance (DUI) of mitochondria. This process—which involves the uniparental transmission of the maternal and paternal mitochondrial genomes through eggs and sperm, respectively, allows for an *a-priori* detection of the sexual identity of developing embryos, as early as the first cleavage division of the zygote: in female embryos, the sperm-inherited mitochondria assume a dispersed pattern between blastomeres; conversely, in males the sperm-inherited mitochondria stay assembled together, remain within one blastomere, and are eventually included in primordial germ cells (PGCs; **Zouros, 2013; Ghiselli et al., 2019**).

Here we sought to expand the knowledge on the process of bivalve SD, by employing the Mediterranean mussel *Mytilus galloprovincialis* as a study system, which is a species exhibiting DUI. Particularly, we aimed to investigate the transcription patterns of three DSFG (namely, *Dmrt-1L*, *Sox-H*, and *Fox-L2*; hereafter referred to as ‘SRG’) during embryonic and early larval stages. To this purpose, (i) we first performed a time-series DGE analysis by using the RNA-sequencing data published by **Miglioli et al. (2024)**; afterwards, (ii) we investigated the temporal and spatial transcription patterns of the DSFGs of interest through mRNA *in-situ* hybridization chain reaction (HCR). To obtain a more comprehensive developmental context for the transcription patterns of DSFGs, (iii) we also traced for the first time in *M. galloprovincialis* the process of the germline specification through mRNA *in-situ* HCR and immunolocalization of *Vasa*/Vasa, which is a traditionally-recognised marker of PGCs and germ cells (GCs) across Metazoa (**Extavour and Akam, 2003**). The specification and differentiation of GCs (which are part of the gonadal tissue in adults) is in fact a critical process in sexually reproducing multicellular organisms, as it provides the groundwork for the subsequent differentiation of sexually dimorphic gametes. Therefore, understanding the developmental pathway leading to the establishment of PGCs and GCs is essential to fully characterise the sex-determining process and how the sexual fate of PGCs/GCs is directed.

5.2 Materials and methods

5.2.1 Time-series gene expression

Miglioli et al. (2024) recently produced one of the very first detailed developmental transcriptomes of the Mediterranean mussel *M. galloprovincialis*, spanning from the unfertilized oocyte to the larval stage at 72 hpf, with time points sampled every 4 hpf. A total of thirty different mRNA libraries was sequenced, consisting of fifteen developmental time points per two biological replicates each (**Tab. S11**). These data are extremely useful to thoroughly investigate the transcription patterns of genes throughout the first three days of the *M. galloprovincialis* development, to quantify the transcription level of target genes to be investigated with mRNA HCR experiments and to have an overview of the possible outcome from such analysis.

Raw reads were downloaded from the Sequence Read Archive (SRA) in NCBI (BioProject: PRJNA996031) and trimmed using Trimmomatic v0.39 (**Bolger et al., 2014**; LEADING:5 TRAILING:5 SLIDINGWINDOW:4:15 MINLEN:65). Read quality was checked using FastQC v0.12.1 (**Andrews et al., 2010**). Trimmed reads were mapped against the *M. galloprovincialis* annotated genome (GCA_900618805.1; **Gerdol et al., 2020**) using STAR v2.7.10b (**Dobin et al., 2013**) in ‘alignReads’ mode with default parameters. The resulting gene count matrix was extracted with StringTie v2.2.1 (**Pertea et al., 2015, 2016**) in expression estimation mode followed by the python script ‘prepDE.py’ (-l 99).

The resulting matrix was processed in R. Raw gene counts were normalised using the median of ratios method as implemented by the ‘DESeq2’ package (**Love et al., 2014**), and then transformed through the DESeq2 variance stabilising transformation (vst). Transformed gene counts were used to run a principal component analyses (PCA) and visualise sample clustering, and to plot expression values of *Vasa*, *Dmrt-1L*, *Sox-H*, and *Fox-L2* (hereafter collectively referred to as ‘target genes’). Normalised gene counts were instead used to run a time-series DGE analysis in ‘maSigPro’ (**Conesa et al., 2006**).

The entire pipeline was automated through custom python and bash scripts, which are available in a private repository on GitHub.

5.2.2 Sample collection, MitoTracker staining and fixation

Adult mussels were hand collected from various locations surrounding the AltaSea institute at the port of Los Angeles (CA, USA). Sampling took place during the spawning season of the species in California, i.e., from Oct, 2023 to Jan, 2024.

Selected mussels were thoroughly cleaned from epibionts and placed in ice for approximately 30–60 minutes, then transferred in filtered artificial sea water (FASW) at 16 °C and acclimatized for 30 minutes. All the individuals were then placed in a common tank and spawning was induced by cyclical thermal shock, that is, by exposing mussels alternatively to FASW at 24–26 °C and 14–16 °C for a time of 30–40 min each. As soon as mussels started spawning, individuals were promptly removed from the common tank, carefully washed, air dried to remove contaminant gametes from the shell, and then allowed to continue spawning in isolated containers of about 250 mL with 16 °C FASW.

Both single and multiple crosses were performed: two males (M1, M2) and two females (F1, F2) were employed for single crosses; six males and six females were employed for multiple crosses, and gametes from the same sex were mixed. One hour after the spawning started, oocytes were filtered through a 75 over a 30 µm mesh, and aged in 1 L of FASW for 40–60 min, to allow them to assume a proper circular shape. Oocyte abundance was estimated under a stereomicroscope by eye counting the number of gametes in five aliquots of 1 mL, and then calculating the mean value. Sperm mitochondria were labelled with MitoTracker Red CMXRos (Thermo Fisher Scientific) at a working concentration of 500 nM for 30 min. MitoTracker is a fluorescent, vital and fixation-resistant mitochondrial dye and was used to detect the sex of developing embryos (as early as the two-blastomere stage) according to the distribution pattern of sperm mitochondria (**Cao et al., 2004; Obata and Komaru, 2005**). From this step onward, samples were always kept in the dark.

Fertilisation was performed by mixing oocytes and sperm at a ratio of 1:10. Fertilisation success was checked after 20–30 min by the formation of polar bodies. The suspension was then carefully washed on a 30 µm mesh to remove excess sperm, and brought to a concentration of 250 zygotes/mL. The resulting suspension was transferred into cell-culture flasks of 40 mL and embryos/larvae were reared at 16 ± 1 °C in the dark. Water was changed every 24 h. After 48 hpf, larvae were fed with the unicellular microalgae *Isochrysis galbana*, at a final concentration of about 10⁵ cells/mL following **Helm et al. (2004)**.

Embryos/larvae were sampled at 1, 2, 3 and 4 hpf, and then every 12 h until 72 hpf. Proper development and vitality were checked under a stereomicroscope at every sampling time. After concentration with a mesh of proper size, embryos/larvae were fixed in 3.2% paraformaldehyde (PFA) in 1× phosphate-buffered saline (PBS; 128 mM NaCl, 2 mM KCl, 8 mM Na₂HPO₄ · 2 H₂O, 2 mM KH₂PO₄) at 4 °C overnight under constant and gentle shaking. Fixed samples were washed 3×20 min in 1× PBS with 0.1% Tween 20 (PBS-Tw) and then dehydrated 3×30 min in absolute methanol at room temperature (RT). Dehydrated samples were stored at –20 °C until usage.

5.2.3 mRNA *in-situ* hybridization chain reaction (HCR)

HCR probe design

Vasa, *Dmrt-1L*, *Sox-H*, and *Fox-L2* spliced-transcript nucleotide sequences of *M. galloprovincialis* were obtained from the previous analyses with OrthoFinder v2.5.5 (**Emms and Kelly, 2019**) on annotated bivalve genomes and transcriptomes (see **Chapter 4**). Accession numbers of spliced transcripts are 10B017427, 10B093608, 10B014180, and 10B094018, respectively. The ‘insitu_probe_generator’ script from the Ozpolat Lab (**Kuehn et al., 2022**) was used to generate pairs of probes specifically designed for third-generation HCR (**Choi et al., 2018**). The built-in BLASTN search against the annotated *M. galloprovincialis* transcriptome was employed to check for putative off-target bindings of probe pairs. B1-488, B2-647, B3-546, and B4-700 pairs of HCR amplifiers and fluorophores were chosen, as reported in **Tab. 5.1**. Resulting probes were synthesised by Integrated DNA Technologies (IDT™) in separate oligo pools.

mRNA *in-situ* HCR and microscope imaging

mRNA *in-situ* HCR in *M. galloprovincialis* embryos was performed following **Miglioli et al. (2024)**. All the steps were carried out in the dark to prevent MitoTracker from fading. Probe hybridization buffer, probe wash buffer and amplification buffer were manufactured by Molecular Instruments, Inc.

Dehydrated samples stored in methanol were washed 4×5 min and 1×10 min in PBS-Tw. Samples were then permeabilized for 30 min in a detergent solution (1% sodium dodecyl sulfate [SDS], 0.5% Tween 20, 50 mM Tris–HCl, 1 mM ethylenediaminetetraacetic acid (EDTA), 150 mM NaCl), and washed again 2×5 min in PBS-Tw. Samples were prepared for the HCR

Table 5.1 – Characteristics of fluorescent dyes used for each labelled target. HCR amplifiers and the number of probe sets (as in Tab. S12) are reported when applicable. Dyes for both *Vasa* and Vasa are reported.

Target	Dye	HCR amplifier	HCR probe pairs	Excitation (nm)	Emission (nm)
dsDNA (nuclei)	DAPI	–	–	360	460
Sperm mitochondria	MitoTracker Red CMXRos	–	–	575	600
<i>Vasa</i> /Vasa	ALEXA-488/-488	B1/–	33/–	499	520
<i>Dmrt-1L</i>	ALEXA-647	B2	18	653	670
<i>Sox-H</i>	ALEXA-546	B3	22	557	575
<i>Fox-L2</i>	ALEXA-700	B4	28	685	700

detection stage by incubation in the probe hybridization buffer for 30 min at 37 °C. Detection stage was then performed with 4 nM of each probe set in hybridization solution overnight (>12 h) at 37 °C.

Excess probes were removed by washing 4×20 min with probe wash buffer at 37 °C and 3×5 min with 5× saline-sodium citrate (150 mM NaCl, 17 mM Na₃C₆H₅O₇) with 0.1% Tween 20 (SSC-Tw) at RT. Samples were incubated for 30 min in the amplification buffer at RT. Hairpins were heated at 95 °C for 90 s and then snap-cooled at RT for 30 min. The amplification step of HCR was performed with 6 pmol of each hairpin in the amplification buffer overnight (>12 h) at RT.

Excess hairpins were removed by washing 2×5 min, 2×10 min, and 1×5 min with SSC-Tw. If not immediately mounted on slides, samples were stored in SSC-Tw at 4 °C. Otherwise, samples were immersed first in 50% glycerol and then in 75% glycerol, each for 30–60 min, and then mounted with VECTASHIELD®PLUS Antifade Mounting Medium with DAPI (H-2000). Slides were imaged on a Stellaris 5 Confocal Package system with the software Las X (Leica Microsystems). Each dye was imaged sequentially in a separate channel, to enhance the yield and avoid crosstalks. **Tab. 5.1** summarises the excitation and emission peaks for each dye. Images were then manipulated and post-produced using Fiji v2.14.0.

5.2.4 Immunolocalization of Vasa

M. galloprovincialis Vasa sequence was manually inspected through multiple sequence alignment with Vasa from other bivalves (data from **Chapter 4**) and several reference species (*Danio rerio* [Ddx4: NP_571132.1]; *Homo sapiens* [Ddx4: NP_077726.1]; *Mus musculus* [Ddx4: NP_001139357.1]; *D. melanogaster* [Vasa: NP_001260458.1]; *Caenorhabditis elegans* [GLH-1: NP_001262379.1, GLH-2: NP_491876.1, GLH-3: NP_491681.1, GLH-4: NP_491207.3]), to support commercial antibody specificity in *M. galloprovincialis*. The Vasa sequence from *D. rerio* was included as the polyclonal antibody was generated using the zebrafish protein variant (manufacturer indications; ab209710 by Abcam Limited). A maximum likelihood (ML) phylogenetic tree of Vasa genes and its paralog Ddx3 (reference genes: *D. rerio* [Ddx3Xa: NP_001119895.1, Ddx3Xb: NP_571016.2]; *H. sapiens* [DDX3X: NP_001180346.1]; *M. musculus* [Pl10/Ddx3Xl: NP_149068.1]; *D. melanogaster* [Belle: NP_001262379.1]; *C. elegans* [LAF-1: NP_001254859.1, VBH-1: NP_001021793.1]) was built using IQTREE. The hidden Markov model (HMM) profile of the Asp-Glu-Ala-Asp/Asp-Glu-Ala-His box (DEAD/DEAH-box) signature domain for the amino acid guided alignment step, was built after the corresponding Pfam full database (PF00270). Methods are the same as in **Chapter 4**.

Vasa immunolocalization in *M. galloprovincialis* embryos was performed following **Milani et al. (2011)** with modifications. All the steps were carried out in the dark to prevent MitoTracker fluorescence from fading. Dehydrated samples stored in methanol were rinsed 3×10 min and 1×2 h in 1× Tris-buffered saline (TBS; 10 mM Tris–HCl, 155 mM NaCl), following an additional wash for 10 min with PBS. Samples were then digested for 6 min and 30 s with 0.01 % pronase E (Merck) in PBS, and washed again 2×5 min in PBS. Permeabilization was performed in 1× TBS with Triton X-100 (TBS-Tx) 0.1 % for 5 min at RT and in TBS-Tx 1 % overnight at 4 °C.

After an additional rinse for 5 min in TBS-Tx 0.1 %, non-specific binding sites were blocked with a TBS-Tx 0.1 % solution containing 3 % bovine serum albumin (BSA). Samples were then incubated at 4 °C for 32–48 h with primary anti-VASA/VAS antibody (polyclonal anti-VASA developed in rabbit; ab209710 by Abcam Limited), diluted 1:100.

Excess primary antibody was rinsed from samples with 4×30 min in TBS-Tx 0.1 %, followed by an incubation of 1 h in TBS-Tx 0.1 % containing 3 % BSA. Samples were then incubated at 4 °C for 24–32 h with secondary antibody HRP anti-rabbit in goat (Santa Cruz Biotechnology Inc.), diluted 1:400.

Excess secondary antibody was rinsed with 4×30 min in TBS-Tx 0.1% and 1×1 h in 1%. Samples were immersed first in 50% glycerol and then in 75% glycerol, each for 30–60 min, and then mounted with VECTASHIELD®PLUS Antifade Mounting Medium with DAPI (H-2000). Slides were imaged on a Nikon A1R+ HD25 confocal microscope. Each dye was imaged sequentially in a separate channel, to enhance the yield and avoid any crosstalks. **Tab. 5.1** summarises the excitation and emission peaks for each dye. Images were then manipulated and post-produced using Fiji v2.14.0.

5.3 Results

5.3.1 Differential gene expression analysis of *Vasa* and SRGs in embryo time-series

Over 24 M reads were mapped for each RNA-sequencing library (86.58% of the total input reads), with an average of 26.8 M (**Tab. S11**). Of these, an average of 22 M reads were uniquely mapped (71.86% of the total input reads), while an average of 4.5 M were multi-mapped (14.72% of the total input reads). The average of unmapped reads was 4.1 M (13.42% of the total input reads; **Tab. S11**). The PCA on normalised read counts returned well-clustered experimental groups for time points between 8 and 36 hpf, while for stages before 8 hpf and after 36 hpf, experimental groups are more homogeneous among each other (**Fig. 5.1A**). This situation may reflect major developmental dynamics during embryogenesis and larval development. As a matter of fact, before 8 hpf, the embryo undergoes segmentation and no big morphogenetic movements are usually detected. Between 8 and 13 hpf, instead, gastrulation begins, the embryo experiences strong morphogenetic rearrangements (such as the formation of embryonic layers) and the trochophore larva develops, all processes which are expected to be detected also at the molecular level. After 36 hpf, instead, the larva does not show any dramatic morphogenetic event, as the D-veliger is almost formed and the gross advanced larval morphology is established. The hierarchical clustering of differentially expressed genes computed by **Miglioli et al. (2024)** is concordant with this view.

Transcription levels of *Vasa*, SRGs, *Fox-B2*, and *Wnt-8a* were plotted individually (**Fig. 5.1B**) to obtain a proxy of the expected outcome of HCR. *Fox-B2* and *Wnt-8a* were employed as control genes to get support for handling of data and of the pipeline, as they were also analysed

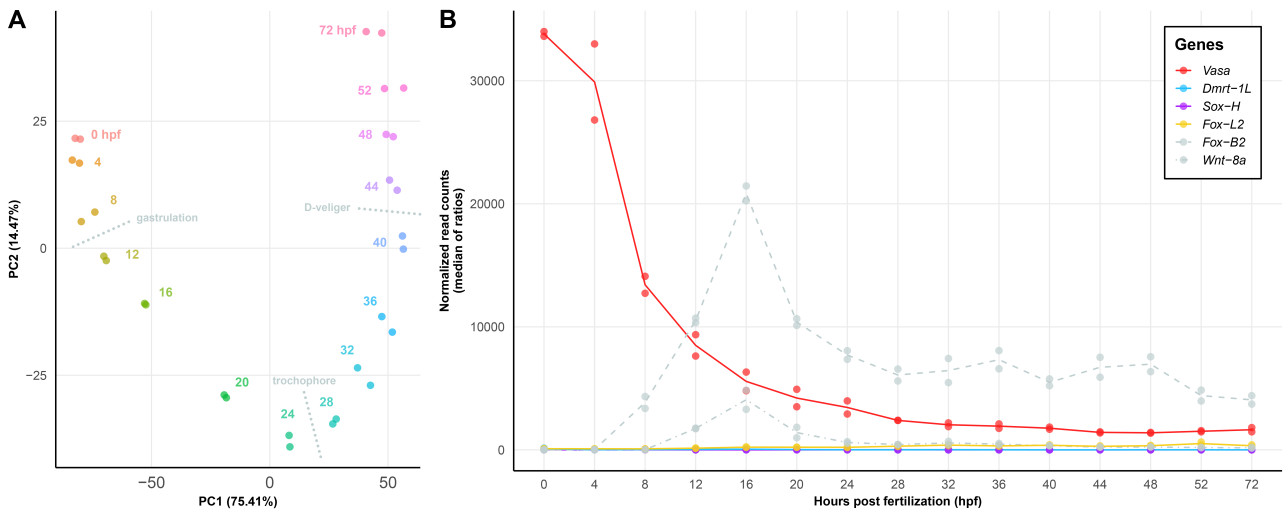


Figure 5.1 – PCA of DESeq2-normalised read counts (A) and transcription levels of target and reference genes (B). (A) Principal components (PCs) 1 and 2 are plotted in the x and y axes, respectively; the proportion of variance explained by each PC is shown in parentheses. Sampled time-points are shown in different colours and are indicated by the hours hours post fertilization (hpf). Major developmental transitions are marked with dotted lines. PCA has been performed on vst-transformed, normalised read counts (DESeq2 median of ratios). (B) Transcription levels of target (*Vasa*, *Dmrt-1L*, *Sox-H*, and *Fox-L2*) and reference genes (*Fox-B2* and *Wnt-8a*) as expressed by normalised read counts (DESeq2 median of ratios).

by Miglioli et al. (2024). The transcription of both genes starts at 4 and 8 hpf, respectively, reaches a peak at 16 hpf, and then constantly decreases (Fig. 5.1B). *Vasa* transcripts are highly abundant in unfertilized oocytes and in embryos 4 hpf, then constantly decrease throughout time; conversely, *Fox-L2* transcripts increase from 12 hpf onward (Fig. 5.1B). Both *Dmrt-1L* and *Sox-H*, instead, show low or null levels of transcriptions throughout the entire time series (Fig. 5.1B).

The maSigPro DGE analysis of the *M. galloprovincialis* developmental time series found 13,067 differentially expressed genes (about 17% of the analysed genes) and clustered them into 9 different groups, according to their specific transcriptional profiles (Fig. 5.2). Among the genes of interest, only *Vasa* and *Fox-L2* showed a significantly different transcriptional profile throughout the time series, and were included in clusters 3 and 1, respectively. As already discussed, *Vasa* and *Fox-L2* transcription levels show an opposite tendency, with the former decreasing and the latter increasing throughout time. Both *Dmrt-1L* and *Sox-H* were not found to be differentially transcribed by maSigPro and, thus, were not included in any cluster. The same holds true for *Wnt-8a* and *Fox-B2*.

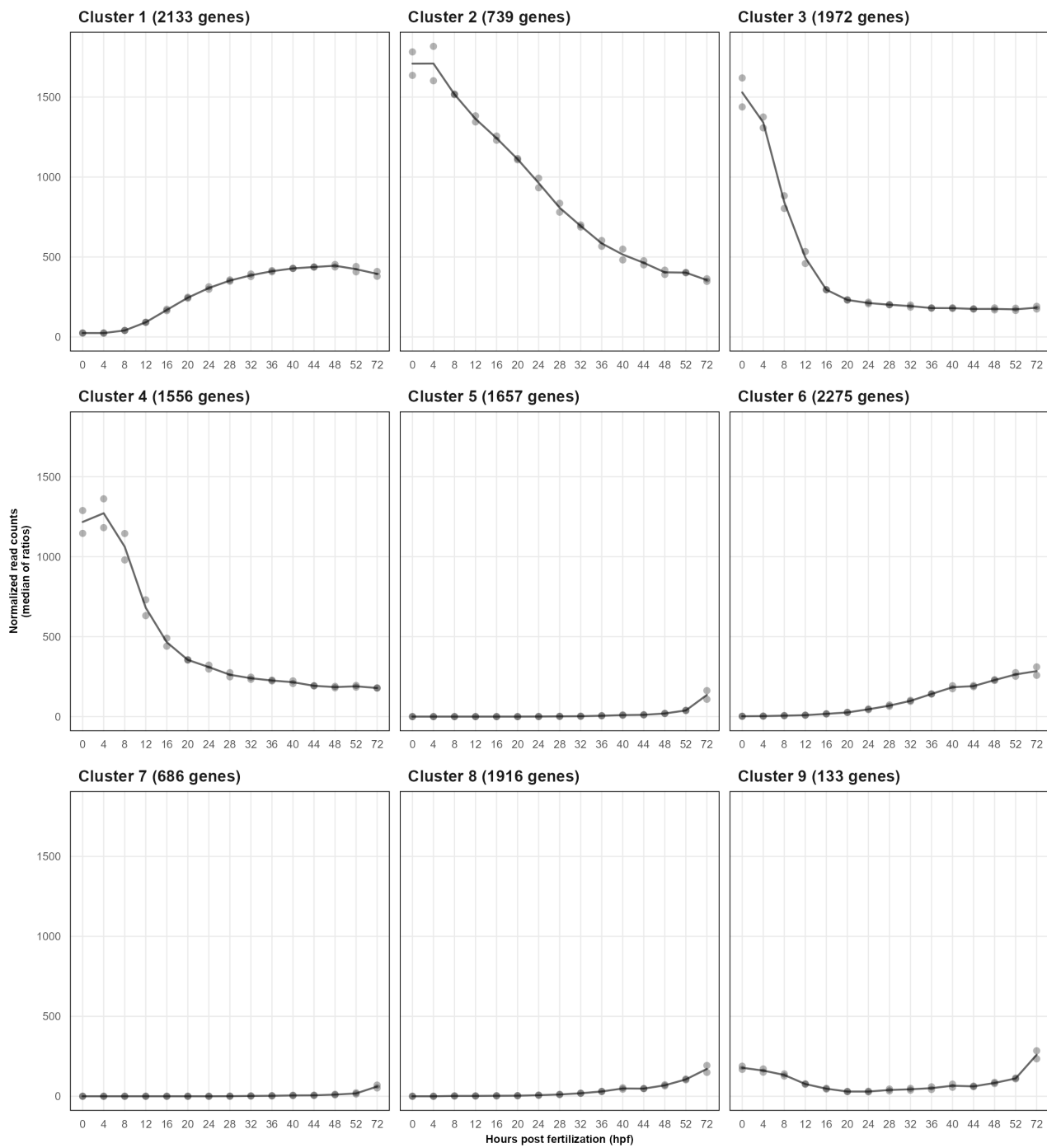


Figure 5.2 – Transcription patterns of differentially-expressed genes as inferred by maSigPro. Genes are divided into 9 different clusters according to their transcription patterns throughout 15 sampled time points. Median values of the two biological replicates are shown for each time point and represented by points. Mean values are shown for each time point and represented by solid lines. *Vasa* is included in Cluster 3, while *Fox-L2* in Cluster 1.

Table 5.2 – Number of imaged samples, divided by developmental stage, experiment, and sex.

Stage	Experiment	Females	Males	Undetermined	Total
Oocytes	HCR	–	–	–	11
2-cell embryos	HCR	8	9	1	18
4-cell embryos	HCR	9	3	0	12
8-cell embryos	HCR	11	3	0	14
12-hpf embryos	HCR	7	6	0	13
Total embryos	HCR	35	21	1	57
24-hpf larvae	HCR	0	1	11	12
48-hpf larvae	HCR	0	0	11	11
72-hpf larvae	HCR	1	1	8	10
Total larvae	HCR	1	2	30	33
Oocytes	Negative control	–	–	–	5
2-cell embryos	Negative control	7	2	0	9
4-cell embryos	Negative control	7	2	0	9
8-cell embryos	Negative control	5	1	0	6
12-hpf embryos	Negative control	0	0	3	3
Total embryos	Negative control	19	5	3	27
Total larvae	Negative control	0	0	4	4
Total imaged samples	All	55	28	38	137

5.3.2 mRNA *in-situ* HCR of Vasa and SRGs

Overall, a total of 80 adult *M. galloprovincialis* individuals were sampled and staged for thermal-shock induced spawning. Of these, 8 males and 8 females were eventually selected as parents for single (2 of each sex) and multiple (6 of each sex) crosses, on the basis of their gamete quality (i.e., presence sperm motility, and oocyte transparency and rounded shape). MitoTracker labelling was successfully retained in developing embryos of *M. galloprovincialis* until 12 hpf. After that stage, the stained sperm mitochondria were difficult to detect, and so was the dispersal pattern to establish the sexual identity.

After embryo rearing, fixation, and mRNA *in-situ* HCR of target genes, a total of 16 oocytes, 81 embryos and 33 mussel larvae were imaged (**Tab. 5.2**). Of these, on the basis of sperm mitochondria dispersal patterns, 55 were females (dispersed pattern), 28 were males (aggregated pattern) and 38 were of indeterminable sex (ambiguous pattern or unlabelled sperm mitochondria). For each stage, negative controls were also imaged (final count of 36), by staining just sperm mitochondria with MitoTracker and nuclei with DAPI, and going through the HCR protocol without adding probes in the hybridization step. A total of 137 samples were imaged.

The ‘insitu_probe_generator’ script (**Kuehn et al., 2022**) generated: (i) 33 probe pairs

conjugated with hairpin B1 and ALEXA-488 for *Vasa*; (ii) 32 probe pairs conjugated with hairpin B2 and ALEXA-647 for *Dmrt-1L*; (iii) 27 probe pairs conjugated with hairpin B3 and ALEXA-546 for *Sox-H*; and (iv) 28 probe pairs conjugated with hairpin B4 and ALEXA-700 for *Fox-L2* (**Tab. 5.1** and **S12**). HCR labelling of genes of interest proved to be concordant with results obtained from RNA-seq analysis (see **Section 5.3.1**; **Fig. 5.1B**). Concerning *Vasa*, it has been detected throughout every sampled stage (**Fig. 5.3A**): transcripts were identified homogeneously in the cytoplasm of unfertilized oocytes, 2-, 4-, and 8-cell embryos; in gastrulae, *Vasa* is located mainly in the ingressed cells; in trochophores, it forms a cup-like structure in the region opposite to the shell-field; in D-larvae, it is mainly retained in two central areas adjacent to the valves (right and left sides of the larvae) in a sort of a comma-shaped region. Concerning *Dmrt-1L* (**Fig. 5.3B**), final images were quite noisy and showed putative non-specific staining at the level of embryo external surface and larvae shell, which may have interfered with the true signal of HCR for this gene; in any case, no clear labelling distribution pattern was found in embryos of both sexes. *Sox-H* mRNAs (**Fig. 5.3C**) were not detected during the imaged developmental stages. Conversely, *Fox-L2* transcripts have been detected starting from the 8-cell stage—where they are homogeneously present, to the D-veliger larvae—where they appear to be mostly co-localized with *Vasa* (**Fig. 5.3D** and **E**). Imaging of control samples (i.e., without mRNA *in-situ* staining) can be found in **Fig. S16**.

5.3.3 Immunolocalization of Vasa

To determine whether the commercial polyclonal antibody (ab209710 by Abcam Limited) could successfully bind *M. galloprovincialis* Vasa, we conducted a phylogenetic analysis (**Fig. 5.4A**) and a multiple sequence alignment inspection (**Fig. 5.4B**) of Vasa/Ddx4 proteins, along with its paralog Ddx3, starting from the bivalve curated genome and transcriptome dataset analysed in **Chapter 4**. We retrieved three different Vasa sequences in the *M. galloprovincialis* genome (**Tab. S13**). Two of them (VDI03911.1 and VDI03912.1) are splicing variants of the same mRNA (acc. no. 10B017427) investigated through DGE and mRNA *in-situ* HCR in previous sections. Both variants are constituted by 17 exons and differ from each other for only eight leading amino acids at the protein N-terminus (**Fig. 5.4C**). Their DEAD/DEAH-box and C-terminal domains show high levels of sequence conservation with respect to *D. rerio* Vasa (**Fig. 5.4C**). Concerning the other Vasa *M. galloprovincialis* sequence (VDI58335.1), it originates from a separate genomic locus and appears very short (105 amino acid positions) if compared to other

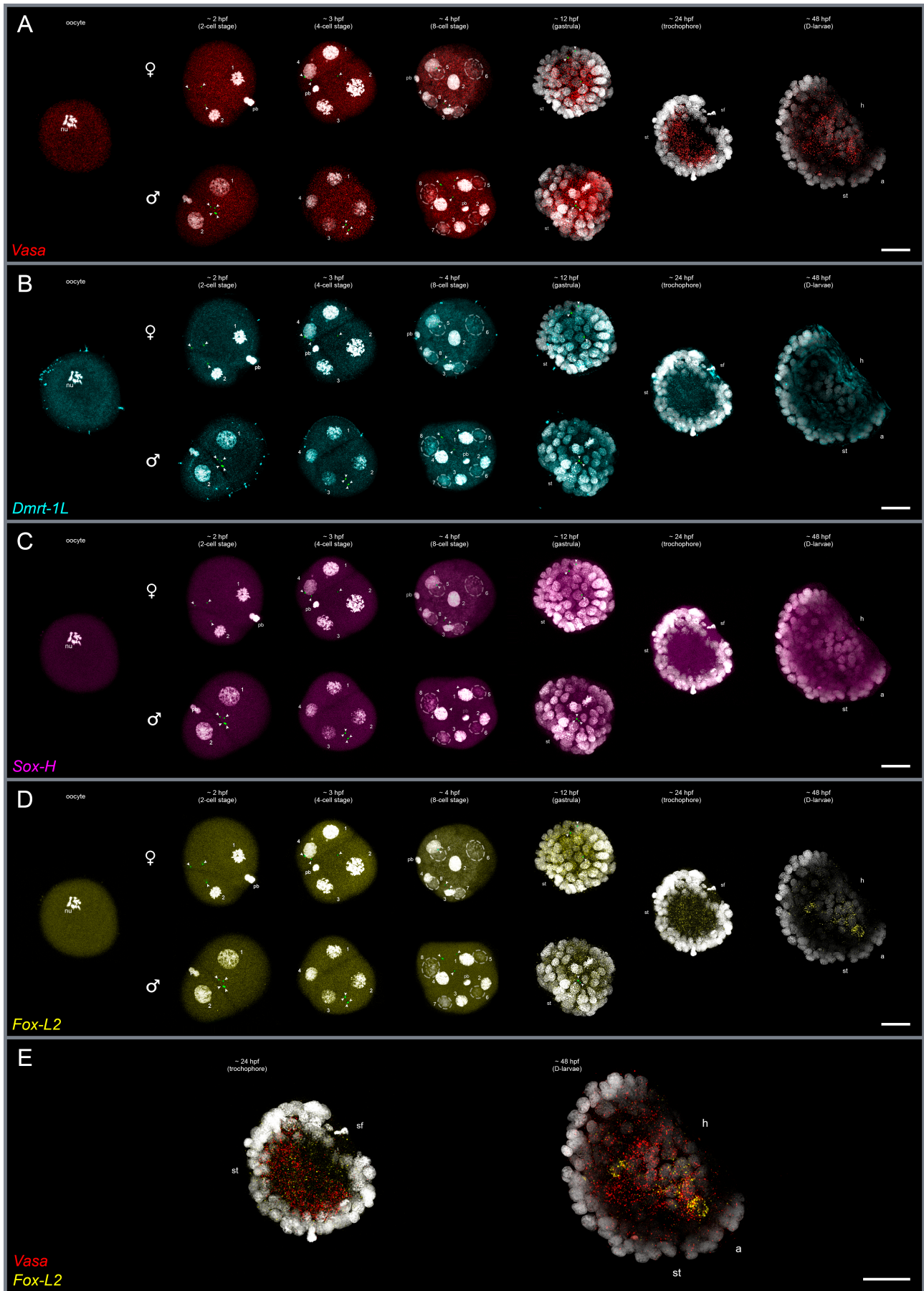


Figure 5.3 – Caption on next page.

Figure 5.3 – mRNA *in-situ* HCR of *Vasa* (A), *Dmrt-1L* (B), *Sox-H* (C), *Fox-L2* (D), and *Vasa+Fox-L2* (merged; E) in several developmental stages of *M. galloprovincialis*. Nuclei are shown in white; in the 2-, 4-, and 8-cell stages, nuclei are also marked with numbers; in the 8-cell stage, nuclei of blastomeres in the background are highlighted with dashed circles. Sperm mitochondria, when stained (shown in green), are marked with arrowheads. For 2-cell, 4-cell, 8-cell, and gastrula stages, embryos of both sexes are reported (top rows: females; bottom rows: males). a: anus; h: hinge; nu: oocyte nucleus; pb: polar body; sf: shell field; st: stomodeum. Scale bar: 20 µm. (Figure on previous page.)

Vasa from bivalves (>800; data not shown). Additionally, it does not exhibit any complete DEAD/DEAH-box domain (as per CDD domain annotation). This considered, the additional Vasa gene (and its relative protein) may be an artefact due to genome mis-assembly and/or mis-annotation, or a non-functional gene. Thus, we argue that it should not affect the correct immunolocalization of the *M. galloprovincialis* Vasa proteins, and will not be further discussed.

Unfortunately, the amount of available samples and antibodies for the experiment was limited. Therefore, we managed to acquire just two oocytes, two 2-cell embryos, and one gastrula. Nonetheless, from the obtained images, we found that Vasa proteins are apparently missing from the oocytes, but can be detected at increasing levels on embryos after 4 hpf (**Fig. 5.5**) and during gastrulation, with a localization matching that of *Vasa* mRNAs (**Fig. 5.4A**). Imaging of control samples (i.e., without primary antibody reaction) can be found in **Fig. S17**.

5.4 Discussion

5.4.1 Sperm mitochondria are not detected after 12 hpf because of Mito-Tracker misincorporation or fading

Because of the presence of the unique DUI of mitochondria, *M. galloprovincialis* offers a compelling system to investigate SD during the early stages of embryogenesis. As a matter of fact, the sexual fate of an embryo appears to be established as soon as the first cleavage division, according to the dispersal pattern of sperm mitochondria (**Saavedra et al., 1997; Cao et al., 2004**): (i) if the embryo is going to develop into a female, sperm mitochondria can be found scattered across different blastomeres; (ii) if the embryo is going to develop into a male, sperm mitochondria are found aggregated all in the same blastomere (usually the macromere), being subsequently transferred to PGCs as part of the germ plasm. Therefore, in order to be able to

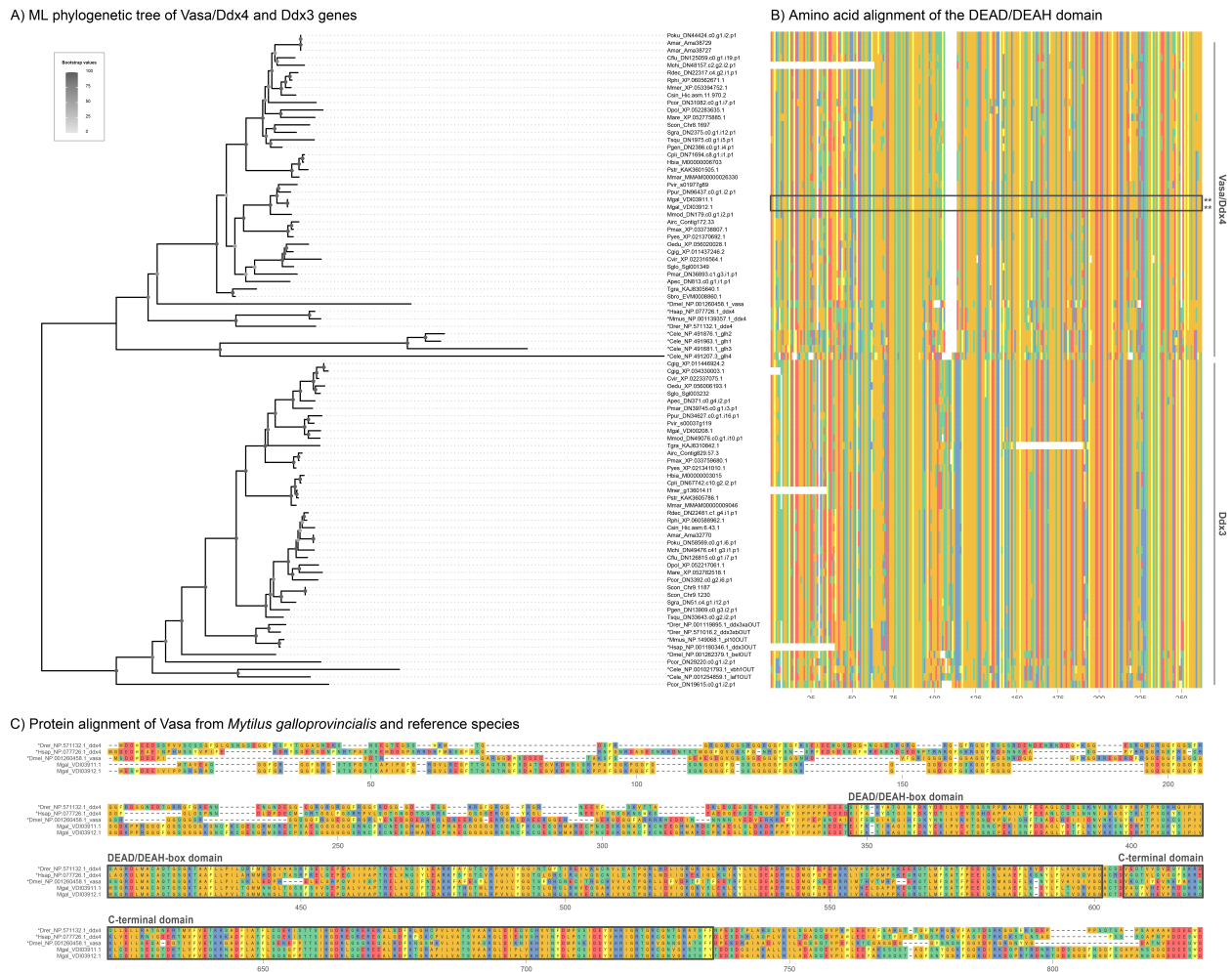


Figure 5.4 – ML phylogenetic tree of Vasa/Ddx3 and Ddx4 proteins from bivalves and reference species (A), along with the amino acid alignment of the relative DEAD/DEAH-box (B) and of Vasa proteins from *M. galloprovincialis* and reference species (C). (A) The tree has been rooted considering the Ddx4 clade as the outgroup. Reference genes from *D. rerio*, *H. sapiens*, *M. musculus*, *D. melanogaster*, and *C. elegans* are marked with an asterisk (*) at the beginning of the tip. Bootstrap values are shown for each node. (B) The alignment of the DEAD/DEAH-box is shown for each tip. The signature DEAD (Asp-Glu-Ala-Asp) motif can be found at positions 198–201. Vasa sequences from *M. galloprovincialis* are highlighted with a solid rectangle and two asterisks (**) on the right. (C) The alignment of complete Vasa sequences from *M. galloprovincialis*, *D. rerio*, *H. sapiens*, and *D. melanogaster* is shown. *C. elegans* has not been included for clarity purpose, because the species has multiple Vasa orthologs. The signature DEAD/DEAH-box and C-terminal-associated domains are highlighted with solid rectangles. Note that position coordinates are not the same between (B) and (C). Colours of amino acid residues in both (B) and (C) correspond to the ‘Chemistry_AA’ scheme from the R package ‘ggmsa’, which highlights the amino acid side-chain chemistry (Zhou et al., 2022). Bivalve species IDs as in Tab. S1. Cele: *Caenorhabditis elegans*; Drer: *Danio rerio*; Dmel: *Drosophila melanogaster*; Hsap: *Homo sapiens*; Mmus: *Mus musculus*. Full descriptions of gene names, accession numbers, and species can be found in Tab. S13.

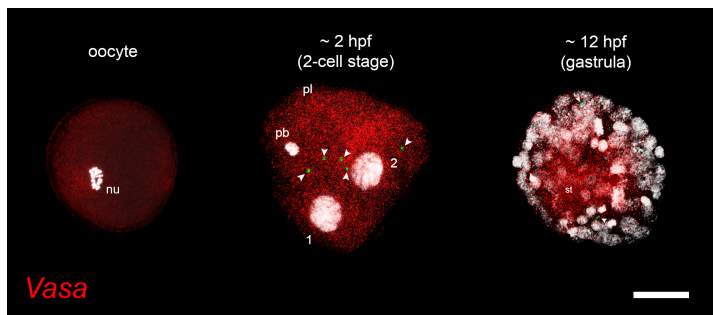


Figure 5.5 – Immunolocalization of Vasa in *M. galloprovincialis* oocyte and embryos. Nuclei are shown in white. Sperm mitochondria (in green) are marked with arrowheads. nu: oocyte nucleus; pl: polar lobe; pb: polar body; st: stomodeum. Scale bar: 20 μ m.

establish the sexual identity of embryos and link it to any differential transcriptions of DSFGs, we labelled sperm mitochondria with MitoTracker (prior to oocyte fertilisation), and check for their dispersal patterns throughout the various sampled stages. Note that the processes of SD and DUI are not necessarily causally linked (**Kenchington et al., 2009**). Despite being successfully retained in developing embryos up until 12 hpf, the MitoTracker fluorescence was difficult to detect in later stages, and so was the mitochondrial dispersal pattern. This phenomenon may have been caused by: (i) misincorporation of MitoTracker during pre-fertilization sperm incubation; (ii) MitoTracker fading after intense manipulation of samples; (iii) MitoTracker dye being incompatible with proper embryo development, i.e., labelled embryos not surviving after 12 hpf. Based on previous studies showing that MitoTracker labelling of sperm mitochondria (including the relative dispersed and aggregated patterns) can be observed up until the late D-larva stage in *Mytilus edulis* (72 hpf; **Cao et al., 2004**), we argue that (i) and (ii) are the most likely explanation for the dye not being detected in samples after 12 hpf. Therefore, MitoTracker has not interfered with the correct development of embryos. However, it must be considered that we employed a rosamine-based MitoTracker dye (MitoTracker Red; which better resists aldehydic fixation), while **Cao et al. (2004)** used a carbocyanine-based Mito-Tracker dye (MitoTracker Green). This may have determined different effects on cell vitality, thus making results not comparable to each other. As a matter of fact, despite MitoTracker dyes are life-compatible as per manufacturer's indications, **Minamikawa et al. (1999)** showed that rosamine-based MitoTracker dyes have photosensitising effects on cells. This means that cells labelled with MitoTracker Red may be committed to apoptosis if exposed to intense light, which induce the loss of the mitochondrial membrane potential and consequent mitochondria swelling. However, based on our experimental conditions, we argue that MitoTracker Red photosensitisation has had a minimal effect, if any, on embryo development. As a matter of fact, after sperm MitoTracker staining, samples were kept in the dark throughout the entire sampling period (MitoTracker Red cytotoxicity is not evident in the dark; **Minamikawa et al.,**

1999), with limited exposure to environmental light just in correspondence with water changes; furthermore, the photosensitisation has been shown to significantly increase with light doses exceeding 0.4 J/cm^2 (cell-colony mortality rate of 60–80 % compared to control colonies; **Minnamikawa et al., 1999**), which is higher than typical environmental light exposure (the solar constant is measured at around 1.362 kW/m^2 , equivalent to $0.1367\text{ J/cm}^2\text{ s}$); (iii) only a little proportion of mitochondria (the 5 sperm-derived mitochondria) have been labelled with MitoTracker, while the oocyte-derived ones remained unlabelled. Altogether, we think that MitoTracker Red staining did not determine a cytotoxic effect on *M. galloprovincialis* embryos and the consequent survival of only unlabelled embryos. Thus, we conclude that MitoTracker was not properly detected on samples older than 12 hpf because of misincorporation since the beginning or the dye fading. However, we acknowledge that a formal survival and vitality test should be performed on *M. galloprovincialis* embryos marked with MitoTracker Red, in order to exclude any possible cytotoxic effect.

5.4.2 Exploring the processes of SD in the *M. galloprovincialis* early development

To date, the molecular basis of bivalve SD has been investigated mainly in adult tissues (e.g., **Li, Zhang, et al., 2018; Liang et al., 2019; Wang et al., 2020; Sun et al., 2022; Wang et al., 2022**). As a matter of fact, considering that in many bivalve species gonads form anew at the beginning of every reproductive season from several populations of PGCs (**Filanti et al., 2021**), it can be speculated that the sexual identity may be established in correspondence with each new gonad formation. This observation would also explain the process by which many bivalve species are capable of sex changes and sex reversal from one reproductive season to the other (**Breton et al., 2018**). Nonetheless, animal SD is a key developmental process often triggered soon after fertilisation and occurring throughout the early development, as can be observed for example in mammals and fruit flies (**Salz and Erickson, 2010; Beukeboom and Perrin, 2014; Richardson et al., 2023**). Consequently, a full understanding of SD in bivalves needs to account also for the events taking place during embryo and larval life stages. To our best knowledge, the only investigation of bivalve SRGs during non-adult stages comes from the Pacific oyster *Crassostrea gigas* (**Naimi et al., 2009**), where the transcription levels of *Vasa*, *Dmrt-1L*, and *Fox-L2* have been investigated through qRT-PCR. In this work, however, only stages between 7 dpf larvae and 4-month-old spats have been tested, and a direct association

of the *Dmrt-1L/Fox-L2* transcription levels with SD could not be established. As a matter of fact, sexes cannot be differentiated in oysters before the onset of gametogenesis, and thus the sex of developing embryos/larvae/spats cannot be properly established (Naimi et al., 2009).

In this work, we aimed to expand the knowledge of bivalve SD by investigating for the first time the transcription patterns of three bivalve SDG candidates—belonging to the DSFG families, during the embryogenesis and early larval development of the Mediterranean mussel *M. galloprovincialis*. This species allows to infer sex of developing embryos by tracing the sperm mitochondria distribution patterns (see Section 5.1; Section 5.4.1). To this purpose, we employed an explorative investigation through a DGE analysis, and mRNA *in-situ* HCR. Our experimental setting, which included the sperm mitochondria labelling, allowed us to *a priori* establish the sex of developing embryos and larvae, and thus to link any differential transcription pattern of DSFGs to the sexual identity.

The DGE analysis showed that the inferred transcription levels of control genes, *Fox-B2* and *Wnt-8a* (Fig. 5.1B), are coherent with the ones reported by Miglioli et al. (2024), indicating that the results obtained from other genes can be considered reliable. The low or null transcription levels of both *Dmrt-1L* and *Sox-H* (Fig. 5.1B) may derive from the absence of transcription itself. However, it must be taken into account that *M. galloprovincialis* shows a mather-dependent sex ratio (Saavedra et al., 1997), that is, the percentage of females and males in the progeny is tightly linked to the mother’s nuclear genome, while being independent from the father’s. Thus, considering that Miglioli et al. (2024) do not specify the sex-ratio of the sequenced embryo pool, the possibility that the low expression levels of *Dmrt-1L* and *Sox-H* may be caused by some sex-biased related effect cannot be ruled out. Nonetheless, mRNA *in-situ* HCR supports also for our samples the scenario depicted by the DGE analysis, that is, the two genes are likely not transcribed, as no unambiguous signal was detected (Fig. 5.3B–C). Concerning *Dmrt-1L*, we think that an additional and more thorough investigation is needed, as the confocal imaging step seemed to have been affected by autofluorescent signals coming from the embryo surface and the larval shell, and/or by a-specific binding of probes (Fig. 5.3B). Thus, to obtain more reliable results, a new mRNA *in-situ* HCR experiment on *Dmrt-1L* should be designed, possibly using a set of amplifiers and fluorophores which is different from the ones employed here (B2-647; Tab. 5.1).

The transcription levels of *Fox-L2* are opposed to those of *Vasa* (see Section 5.3; Sec-

tion 5.4.3; Fig. 5.1A): the gene is not transcribed up until about 12 hpf, i.e., corresponding to gastrulation; from this stage onward, the gene transcription is detected homogeneously all over the embryo, and then becomes restricted to two regions located at both sides of the D-veliger larvae (**Fig. 5.3D**). In particular, in this stage of development, *Fox-L2* appears to be co-localised with Vasa (**Fig. 5.3E**), suggesting a role in PGC specification and/or differentiation. No sex-biased transcription has been detected for *Fox-L2*, though it must be considered that after 12 hpf we were not able to confidently establish the sexual identity of embryos/larvae through the localization of sperm-derived mitochondria (see **Section 5.4.1**). Thus, although it is tempting to speculate a possible role of *Fox-L2* in the specification of female gonads, as proposed by **Zhang et al. (2014)** in *C. gigas*, no definitive conclusions can be drawn at this time.

All considered, this work suggests two scenarios: (1) *Dmrt-1L*, *Sox-H*, and/or *Fox-L2* are truly SDGs, or in any case are top regulators in the bivalve SD process (as proposed by previous authors [**Zhang et al., 2014; Li, Zhang, et al., 2018**] and by the comparative genomics analysis of **Chapter 4**), though in *M. galloprovincialis* their activation (hence, SD) does not occur in early development, but in later stages; (2) *Dmrt-1L*, *Sox-H*, and *Fox-L2* are not SDGs, but are involved in gonad differentiation and maintenance in adult individuals (as found in previous works). That said, the two possibilities should not be viewed as mutually exclusive. As a matter of fact, *Dmrt-1L*, *Sox-H*, and/or *Fox-L2* may be required during *M. galloprovincialis* development for early SD, which however occur at advanced larval/spat stages, as observed in *C. gigas* (SD occurs at 40–60 dpf; **Naimi et al., 2009; Santerre et al., 2013**), but they are also required in adults to allow PGCs to initiate the sex-specific gonad development and differentiation at every reproductive season. As a matter of fact, a similar expression pattern has been seen for *M. galloprovincialis* *Vasa*, whose transcription is detectable in PGCs at a low level during the non-reproductive season, and at a high level both in immature mussels and in the reproductive season (**Obata et al., 2010**). Thus, it can be speculated that the genes triggering the SD cascade may be following similar paths.

5.4.3 Primordial germ cells are specified by both preformation and epigenesis in *M. galloprovincialis*

The process of gonad specification (including PGCs) in bivalves have been studied in several species, both in adults (e.g., **Fabioux, Pouvreau, et al., 2004; Obata et al., 2010; Filanti et al., 2021**) and during early development (**Woods, 1932; Fabioux, Huvet, et al., 2004; Kakoi et al., 2008**). In the pea clam *Sphaerium striatum* (**Woods, 1932**), the specification of the germline is traced back to the unfertilized oocyte, where the germline determinants (in the form of electron-dense granules, mainly containing mitochondria) are included in an asymmetric region of the cytoplasm (the germ plasm). The zygote segmentation then segregates the germ plasm in single blastomeres, until in the gastrula it is found only in two quiescent PGCs, which derives from the 4d blastomere (nomenclature as per **Lyons et al., 2012**). In the Pacific oyster *C. gigas* (**Fabioux, Huvet, et al., 2004**), a similar process of PGC formation has been described. The germline marker *Vasa* is found to be maternally transmitted to the embryo and deposited in a region (the germ plasm) at the vegetal pole of the oocyte. With the onset of segmentation, *Vasa* progressively segregates in single blastomeres, until it is found only in two separated cell clumps at both sides of the pericardic region in the D-larva (**Fig. 5.6A**), where they will form PGCs. The two cell clumps derive again from the 4d blastomere and, in adult oysters, they will periodically proliferate and migrate to the adjacent connective tissue to build gonad acini during the reproductive season (**Fabioux, Pouvreau, et al., 2004; Milani et al., 2017**). A different mechanism has instead been proposed for the Japanese spiny oyster *Saccostrea kegaki* (**Kakoi et al., 2008**). Here, *Vasa* is said to be transcribed all over the embryo until the 8-cell stage (data are not available on the original publication), becoming progressively more restricted to certain blastomeres only after the 50-cell stage. With gastrulation, *Vasa* is detected only at the posterior mesoderm, which derives from the 4d blastomere. In the present work, in the attempt to investigate the transcription patterns of several SDGs, we also characterised the emergence of PGCs in the early development of *M. galloprovincialis* by mean of *Vasa/Vasa* localization, thus providing an additional description of PGC development in bivalves.

According to the DGE analysis, *Vasa* shows a transcription pattern typical of maternal factors (**Xu et al., 2018**), which are stored as transcripts in the oocytes during oogenesis and then constantly decrease throughout embryo segmentation, gastrulation and early larval development, down to undetectable levels (**Fig. 5.1B**). mRNA *in-situ* HCR confirmed these results,

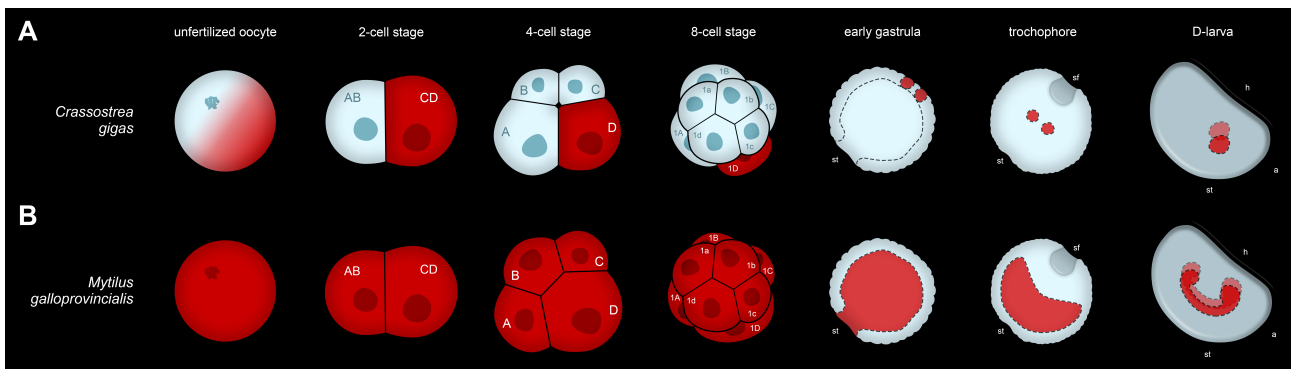


Figure 5.6 – Comparison of Vasa localization (in red) during the Pacific oyster *C. gigas* (A) and the Mediterranean mussel *M. galloprovincialis* (B) early development. Drawings not in scale. Data of *C. gigas* from **Fabioux et al. (2004)**. Blastomere nomenclature as per **Lyons et al. (2012)**. a: anus; h: hinge; st: stomodeum; sf: shell field.

showing that *Vasa* mRNA is located all over the cytoplasm of the oocyte and in all the blastomeres up until the gastrulation stage, when cells positive to *Vasa* move to constitute internal cell layers; following additional morphogenetic movements, *Vasa*-positive cells are eventually present only in two limited regions at both the lateral sides of the D-veliger (**Fig. 5.3A**). On the contrary, immunolocalization showed a different temporal distribution pattern compared to its mRNA, revealing that *Vasa* does not occur in the oocyte, but that its translation begins at low level only with segmentation of the zygote and then increases with gastrulation (**Fig. 5.5**). This different localisation is likely the result of a delay in *Vasa* mRNA translation, which is activated only during the embryo segmentation and grows with the increase of cell number. Nonetheless, this different localisation pattern of mRNA and protein may also have been determined by the differential transcription/translation of the two *Vasa* splicing variants annotated in the *M. galloprovincialis* genome, or by a non-specific binding of either the HCR DNA probes or the primary polyclonal antibody. However, considering that the two *Vasa*/Vasa variants are mostly identical, except for eight leading amino acids at the protein N-terminus (**Fig. 5.4C**), in our experiments we should have been able to target both. As a matter of fact, identifying any differential expression between the two variants would be almost impossible, either through mRNA *in-situ* HCR or immunolocalization. Accordingly, the DGE analyses failed in retrieving any dissimilarity in the transcription levels of the two splicing variants (data not shown), as the experiment was based on short-read sequencing (**Miglioli et al., 2024**). Regarding the HCR probes, they have been specifically designed on the complete *Vasa* mRNA spliced sequence, which has already been proven to specifically label PGCs in a previous analysis on *M. gallo-*

provincialis sub-adults and adults individuals (**Obata et al., 2010**). Regarding the commercial antibody, given the high sequence similarity between the Vasa proteins from *M. galloprovincialis* and *D. rerio* (the protein used to produce the antibody; see **Section 5.2**), at least in their core regions (i.e., the DEAD/DEAH-box and the C-terminal domains; **Fig. 5.4C**), we are confident that the immunolocalization procedure correctly labelled Vasa proteins; furthermore, the same set of polyclonal antibodies has been shown to successfully target PGCs/GCs in another bivalve species, *Ruditapes philippinarum*, and its specificity has been supported by Western blotting (**Filanti et al., 2021**). We thus consider our results to be strongly reliable in the correct localization of *Vasa*/Vasa.

Altogether, the present study shows a process of germline specification in the Mediterranean mussel which resembles that of *S. kegaki*, but differs from the one described in *C. gigas* (**Fig. 5.6A**) and *S. striatum*. As a matter of fact, contrary to these latter two species, in *M. galloprovincialis* *Vasa* transcripts do not form any evident gradient in the oocyte and early stages of embryogenesis (until 8-cell stage), while becoming progressively more restricted to specific cell populations only later in development (**Fig. 5.3**; ??). Particularly, with the onset of gastrulation, *Vasa*-positive cells are internalised in the developing gastrula, and *Vasa* is thus retained only by the inner cell layers. Once the embryo metamorphoses into a trochophore larva, *Vasa* transcripts arrange in a cup-like structure in the region opposite to the shell field (the ventral side), while in the early D-veliger *Vasa* is present only in two lateral regions next to the valves. Here, PGCs are going to form, eventually constituting two symmetrical linear clumps at the base of the dorsal mantle (**Obata et al., 2010**), which is going to represent the primary source of stem cells for gonad acini formation at every reproductive season (**Obata et al., 2010**). This mechanism is also reflected in the localization of Vasa proteins, which do not show any clear gradient in the oocyte (from which is absent) and at least up until gastrulation. Altogether, these findings suggest that in the Mediterranean mussel, *Vasa*/Vasa may segregate in the PGCs not only because of their inheritance as maternal factors (through preformation), but also in response to some external (and unknown) zygotic signal (through epigenesis). Therefore, *Vasa* alone do not allow to identify the presumptive primordial germ cells (pPGCs) or the PGCs during the earliest stages of development, as instead it has been shown in adult individuals (i.e., upon PGC formation; **Obata et al., 2010**). Given that *Vasa*/Vasa mark a population of cells instead of few blastomeres (those constituting the pPGCs), *Vasa* may consequently play a role also in the broader field of stem cell specification during embryogenesis, as shown in the

marine polychaete *Platynereis dumerilii* (**Rebscher et al., 2007**) and the sea urchin *Strongylocentrotus purpuratus* (**Voronina et al., 2008**). In these two species, the germline is specified via an intermediate process relying on both preformation and epigenesis, which can be considered a ‘two-step process’ (**Rebscher et al., 2007; Kumano, 2015**). In this model, a lineage of pluripotent stem cells (PSCs) first segregates during early embryogenesis, and then produces PGCs and other mesodermal somatic structures by unequal cell division. The germline markers, including *Vasa*, are thus localised in both the PSCs and in the descendant PGCs. A similar pattern of *Vasa* localisation (i.e., ubiquitously present in oocytes and during early cleavage of the embryo, then progressively restricted to specific blastomeres) has also been shown in the snail *Ilyanassa obsoleta* (**Swartz et al., 2008**) and the abalone *Haliotis asinina* (**Kranz et al., 2010**), despite not being directly linked to PSC specification.

5.5 Conclusion

In the present work, we hypothesise that the PGC specification in *M. galloprovincialis* follows a two-step process (i.e., a combination of both preformation and epigenesis), which involves the PGCs to be formed only after embryogenesis (**Kumano, 2015**). A similar process may also be hypothesised for *S. kegaki* (**Kakoi et al., 2008**). This mechanism may explain why *Dmrt-1L* and *Sox-H*, if confirmed as SDGs, are not transcribed during the investigated developmental stages. SD would in fact happen only upon PGC commitment, thus during advanced larval development. The present work represents the first attempt to characterise the spatial localisation of three DSFGs in the Mediterranean mussel embryonic and larval development, along with *Vasa/Vasa*, and proves the importance of considering also the developmental stages when investigating new species in a comparative framework. Adopting such an evolutionary developmental perspective may in fact reveal new processes and patterns in animal biology, even when considering closely-related species. As a matter of fact, on the basis of available studies, it has been previously proposed that PGC specification is generally based on epigenesis in gastropods and preformation in bivalves (**Obata and Komaru, 2012**), even though the underlying mechanisms may be species-specific (**Obata and Komaru, 2012**). However, the model represented by the Mediterranean mussel showed that the process of PGC specification may be more diverse in bivalves than expected: preformation happens in *C. gigas* and *S. striatum*, and the two-step process happens instead in *S. kegaki*, and, as we propose, in *M. galloprovincialis*. Therefore,

results provided by the present work support the idea that the traditional preformation and epigenesis should not be accounted as mutually-exclusive phenomena nor as the only mechanisms of PGC formation (**Extavour, 2007; Kumano, 2015**). Clearly, given the unavailability of any PGC marker in bivalve embryos and larvae (and in mollusc in general), at the moment it is not possible to unambiguously establish the emergence and commitment of PGCs during embryogenesis (**Rebscher, 2014**), especially if based only on few germline genes. As a matter of fact, PGCs may share certain genetic markers (e.g., *Vasa*, *Nanos*, *Piwi*, and *Pl-10*) also with some stem cell lineages (**Extavour and Akam, 2003; Extavour, 2007; Rebscher et al., 2007; Voronina et al., 2008; Rebscher, 2014; Piccinini and Milani, 2023**). Thus, more comprehensive investigations are needed to fully and unambiguously characterise the emergence of the germline in bivalve embryos, for example through the examination of the histological and cytological morphology and of genetic regulations (**Extavour and Akam, 2003**). A similar scenario holds true also for SD and SDGs. In fact, we should not expect that the sex-determining process, together with its underlying gene regulatory networks and the timing of its expression, is the same across the entire bivalve diversity. SD is indeed one of the most variable developmental processes, despite its importance in the morphological development of an organism (**Capel, 2017**). However, it can be expected that the main actors, being them genetics or environmental or of multiple origin, are conserved, at least in having a role along the whole SD process (**Capel, 2017**). Future studies would thus need to further address the functions of the main DSFG candidates, as well as the modes of SD and germline development, through cutting-edge techniques (such as single-cell RNA-sequencing) and possibly also encompassing various life stages. In this sense, it is tempting to consolidate the role of the Mediterranean mussel as a model system for SD and germline studies by taking advantage not only of the DUI of mitochondria as a proxy for the sexual identity, but also of the ability of the species to produce a sex-biased offspring with the sole maternal influence (**Saavedra et al., 1997**). This would allow a more thorough and straightforward investigation of the determinants influencing the sex and germline specification of developing mussels, by means of targeted RNA-sequencing and transcript/protein localisation.

Chapter 6

Conclusions

The main objective of this PhD thesis was to investigate bivalve sex determination (SD) through the lens of evolutionary and integrative biology. Bivalves is a group of animals characterised by highly heterogeneous sexual and reproductive modes, with strictly gonochoristic species, obligate and facultative hermaphrodites (either protandrous, protogynous and bidirectional), as well as androgenetic systems. Both genetic and environmental factors seem to influence the sexual identity, at various degrees according to the species, and heteromorphic sex chromosomes (HeSCs) seem to have not been selected throughout the bivalve evolutionary history. Therefore, a rigorous comparative approach is essential to unravel the extreme complexity that regulates bivalve SD. Particularly, by combining bioinformatics with *wet-lab* techniques, including genomics, phylogenetics, molecular evolution analyses, differential gene expression (DGE), mRNA *in-situ* hybridization chain reaction (HCR), and immunolocalization, this work lays the foundation to understanding how sex-determination related genes (SRGs), with a special focus on the Dmrt, Sox, and Fox gene (DSFG) families, have evolved and may function in sex-determining processes across the bivalve taxonomic diversity.

In **Chapter 3**, the emerging role of bivalves as model organisms for SD studies has been emphasised through a critical examination of the current knowledge. The complexity of bivalve reproduction and sexual systems underscored the need to view SD not as a binary and stationary process, but rather as a highly dynamic continuum influenced by multiple genetic and environmental factors. Adopting this broader perspective will allow for a more effective investigation of the biology of SD.

In **Chapter 4**, the molecular evolution of SRGs across a range of bivalve species was analysed. The findings revealed patterns of accelerated amino acid sequence divergence (AASD)

in key SRGs, namely *Dmrt 1-like* (*Dmrt-1L*) and *Sox-H*, supporting the hypothesis that these genes are deeply involved in SD mechanisms, possibly even as primary sex-determining genes (SDGs). Thanks to a comparative study which encompassed the analysis of additional control datasets—mammals and *Drosophila*, the validity of the results has been confirmed and discussed in the light of a broader framework. This comparative approach allowed for the identification of evolutionary convergences and divergences, advancing our understanding of the patterns of molecular evolution in animal SRGs.

Chapter 5 focused on gene expression studies in the Mediterranean mussel *Mytilus gallo-provincialis*, offering insights into the SD process in early development. Particularly, *Dmrt-1L* and *Sox-H* appear to not be expressed during these stages, while *Fox-L2* transcription starts only with the onset of gastrulation. This suggests that either these genes are not top regulators of SD, or that SD occurs only later in development, thus their expression is not found during the analysed stages. The latter interpretation would be in line with the pattern of primordial germ cell (PGC) specification in *M. galloprovincialis*, which begins only in correspondence with the onset of gastrulation, thus not following a strict preformation model as in other studied bivalves.

Overall, this thesis further demonstrates that bivalves, with their vast reproductive and sexual diversity, serve as ideal models for investigating the complexity of SD. By integrating genomic analyses with developmental biology, this work provides a new framework for understanding how SRG evolve and function in diverse species. Future studies could build on these insights by exploring the functional roles of SRGs through other advanced techniques (such as CRISPR-Cas9), thus expanding our understanding of the genetic underpinnings of SD and differentiation. This integrative approach has the potential to unlock new knowledge not only in bivalves but across a wide array of species, deepening our understanding of the evolutionary forces shaping reproductive biology in animals.

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Appendix

The appendix includes the titles and abstracts of the papers published during my PhD that are not part of this thesis.

Taxonomic revision of the Australian stick insect genus *Candovia* (Phasmida: Necrosciinae): insight from molecular systematics and species-delimitation approaches.

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Abstract. The Phasmida genus *Candovia* comprises nine traditionally recognized species, all endemic to Australia. In this study, *Candovia* diversity is explored through molecular species-delimitation analyses using the *COI_{Fol}* gene fragment and phylogenetic inferences leveraging seven additional mitochondrial and nuclear loci. Molecular results were integrated with morphological observations, leading us to confirm the already described species and to the delimitation of several new taxa and of the new genus *Paracandovia*. New *Candovia* species from various parts of Queensland and New South Wales are described and illustrated (*C. alata* sp. nov., *C. byfieldensis* sp. nov., *C. dagleishae* sp. nov., *C. eungellensis* sp. nov., *C. karasi* sp. nov., *C. koensi* sp. nov. and *C. wollumbinensis* sp. nov.). New combinations are proposed and species removed from synonymy with the erection of the new genus *Paracandovia* (*P. cercata* stat. rev., comb. nov., *P. longipes* stat. rev., comb. nov., *P. pallida* comb. nov., *P. peridromes* comb. nov., *P. tenera* stat. rev., comb. nov.). Phylogenetic analyses suggest that the egg capitulum may have independently evolved multiple times throughout the evolutionary history of these insects. Furthermore, two newly described species represent the first taxa with fully developed wings in this previously considered apterous clade.

Comparative genomics of *Hox* and *ParaHox* genes among major lineages of Branchiopoda with emphasis on tadpole shrimps.

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Abstract. *Hox* and *ParaHox* genes (HPHGs) are key developmental genes that pattern regional identity along the anterior-posterior body axis of most animals. Here, we identified HPHGs in tadpole shrimps (Pancrustacea, Branchiopoda, Notostraca), an iconic example of the so-called “living fossils” and performed a comparative genomics analysis of HPHGs and the *Hox* cluster among major branchiopod lineages. Notostraca possess the entire *Hox* complement, and the *Hox* cluster seems to be split into two different subclusters, although we were not able to support this finding with chromosome-level assemblies. However, the genomic structure of *Hox* genes in Notostraca appears more derived than that of *Daphnia* spp., which instead retains the plesiomorphic condition of a single compact cluster. Spinicaudata and *Artemia franciscana* show instead a *Hox* cluster subdivided across two or more genomic scaffolds with some orthologs either duplicated or missing. Yet, branchiopod HPHGs are similar among the various clades in terms of both intron length and number, as well as in their pattern of molecular evolution. Sequence substitution rates are in fact generally similar for most of the branchiopod *Hox* genes and the few differences we found cannot be traced back to natural selection, as they are not associated with any signals of diversifying selection or substantial switches in selective modes. Altogether, these findings do not support a significant stasis in the Notostraca *Hox* cluster and further confirm how morphological evolution is not tightly associated with genome dynamics.

Multiple and diversified transposon lineages contribute to early and recent bivalve genome evolution.

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Abstract. **Background.** Transposable elements (TEs) can represent one of the major sources of genomic variation across eukaryotes, providing novel raw materials for species diversification and innovation. While considerable effort has been made to study their evolutionary dynamics across multiple animal clades, molluscs represent a substantially understudied phylum. Here, we take advantage of the recent increase in mollusc genomic resources and adopt an automated TE annotation pipeline combined with a phylogenetic tree-based classification, as well as extensive manual curation efforts, to characterize TE repertoires across 27 bivalve genomes with a particular emphasis on DDE/D class II elements, long interspersed nuclear elements (LINEs), and their evolutionary dynamics. **Results.** We found class I elements as highly dominant in bivalve genomes, with LINE elements, despite less represented in terms of copy number per genome, being the most common retroposon group covering up to 10 % of their genome. We mined 86,488 reverse transcriptases (RVT) containing LINE coming from 12 clades distributed across all known superfamilies and 14,275 class II DDE/D-containing transposons coming from 16 distinct superfamilies. We uncovered a previously underestimated rich and diverse bivalve ancestral transposon complement that could be traced back to their most recent common ancestor that lived about 500 million years ago [Mya]. Moreover, we identified multiple instances of lineage-specific emergence and loss of different LINEs and DDE/D lineages with the interesting cases of CR1-Zenon, Proto2, RTE-X, and Academ elements that underwent a bivalve-specific amplification likely associated with their diversification. Finally, we found that this LINE diversity is maintained in extant species by an equally diverse set of long-living and potentially active elements, as suggested by their evolutionary history and transcription profiles in both male and female gonads. **Conclusions.** We found that bivalves host an exceptional

diversity of transposons compared to other molluscs. Their LINE complement could mainly follow a “stealth drivers” model of evolution where multiple and diversified families are able to survive and co-exist for a long period of time in the host genome, potentially shaping both recent and early phases of bivalve genome evolution and diversification. Overall, we provide not only the first comparative study of TE evolutionary dynamics in a large but understudied phylum such as Mollusca, but also a reference library for ORF-containing class II DDE/D and LINE elements, which represents an important genomic resource for their identification and characterization in novel genomes.

Towards a time-tree solution for Branchiopoda diversification: a jackknife assessment of fossil age priors.

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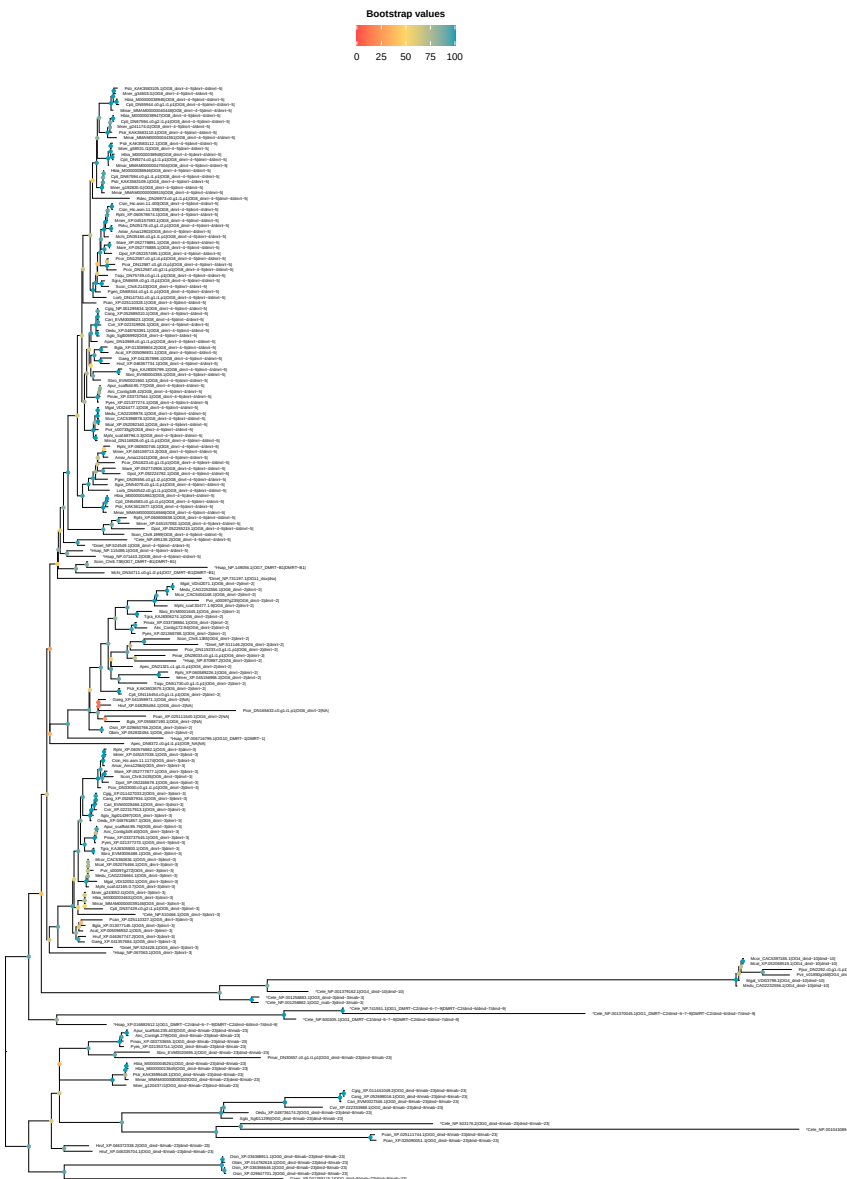
Submitted for peer-review.

Abstract. An understanding of Branchiopoda's evolutionary history is crucial for a comprehensive knowledge of the Pancrustacea tree of life, given their close evolutionary relationship with Hexapoda. Despite significant advances in molecular and morphological phylogenetics that have resolved much of the branchiopod backbone topology, a reliable temporal framework remains elusive. Key challenges include a sparse fossil record, long-term morphological stasis, and past topological inconsistencies. Leveraging a Bayesian Inference approach and the most extensive phylogenomic dataset for branchiopod to date, encompassing 46 species and over 130 genes, we inferred a time-calibrated phylogenetic tree. Furthermore, to strengthen the confidence in our divergence times estimation, we assessed the impact of age priors, topological uncertainties, and gene trees which are discordant from the species trees. Our results are largely consistent with the fossil record and with previous studies, indicating that Branchiopoda originated between 400 and 500 Mya, and the orders of large branchiopods diversified during the Mesozoic. Concerning Cladocera, results remain problematic, with a sharper uncertainty in the diversification time with respect to the fossil record. Though, the jackknife resampling of fossils and the other sensitivity analyses proved our calibration method to be robust, suggesting that the difficulties in obtaining a paleontological-consistent time tree may be hindered by the variability in branchiopod substitution rates and topological instability within certain clades.

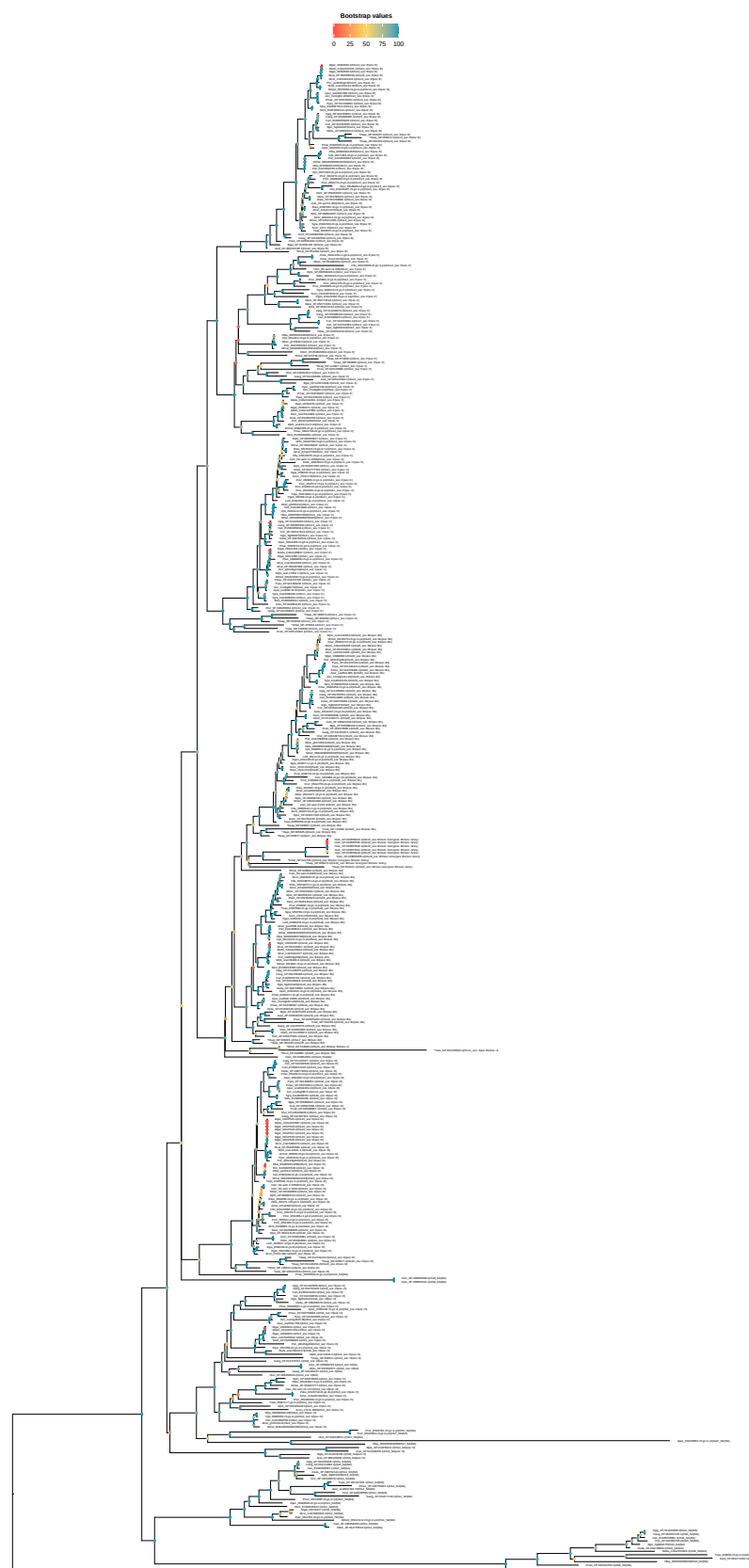
Supplementary figures

High-quality supplementary figures are available at the following GitHub repository:

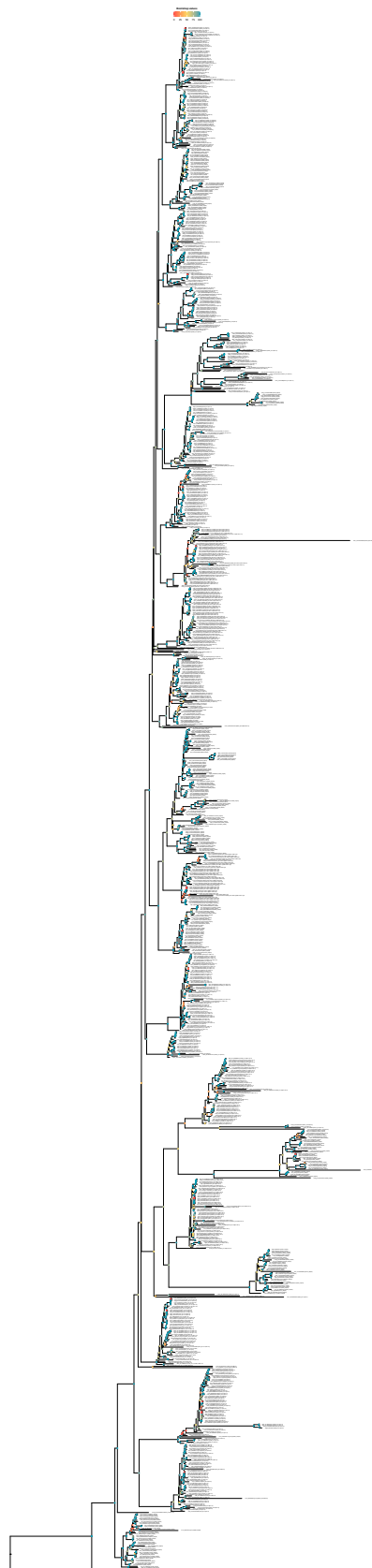
https://github.com/filonico/phd_thesis_tex



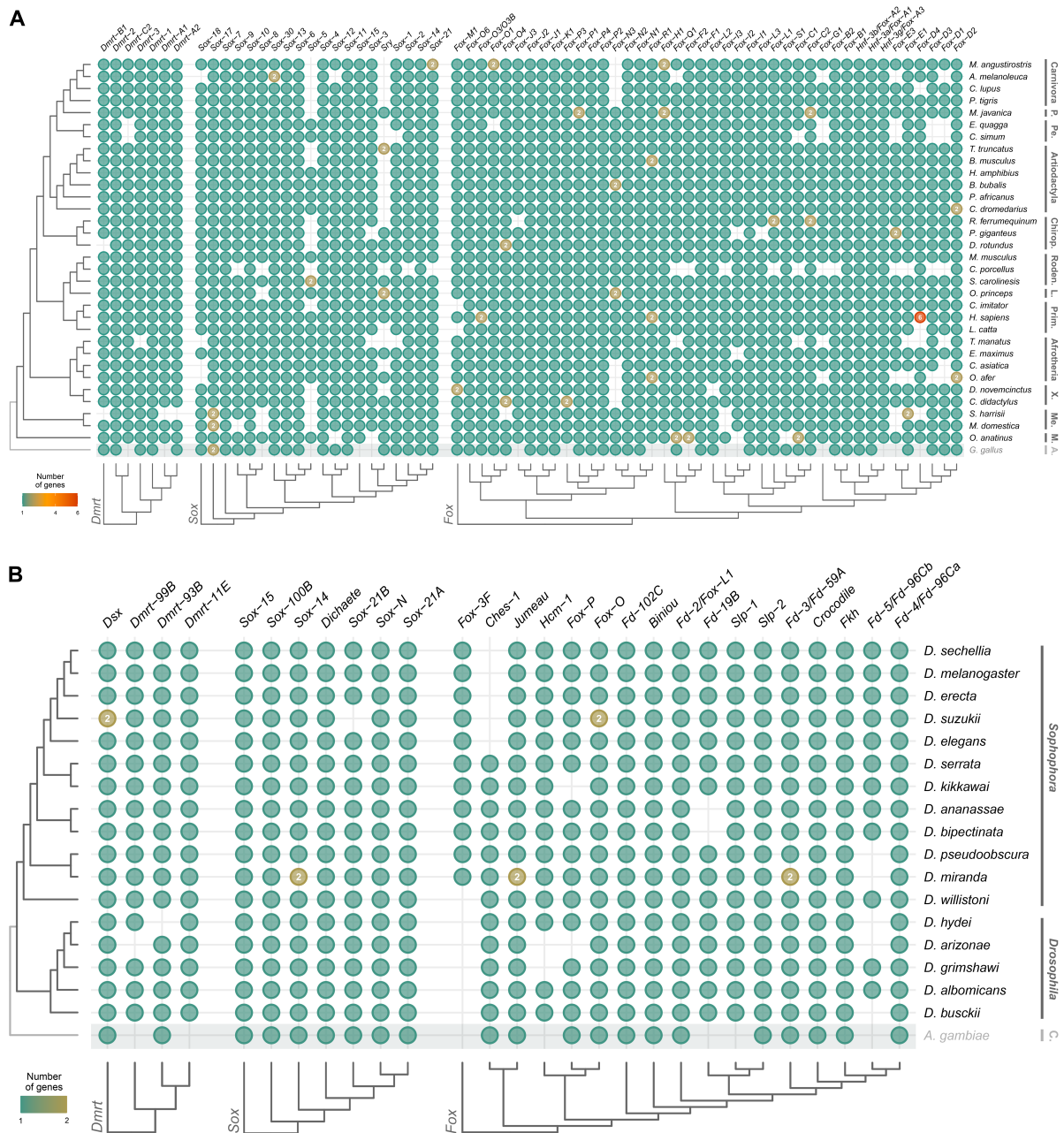
Supplementary Figure S1 – maximum likelihood (ML) phylogenetic tree of the Dmrt gene family in molluscs, including the possvm orthology inference. Reference genes from *Homo sapiens*, *Caenorhabditis elegans*, and *Drosophila melanogaster* are marked with an asterisk at the beginning of the tip names. Species ID can be found in **Tab. S1**. The tree has been midpoint rooted. Bootstrap values are shown for each node.



Supplementary Figure S2 – ML phylogenetic tree of the Sox gene family in molluscs, including the possvm orthology inference. Reference genes from *H. sapiens*, *C. elegans*, and *D. melanogaster* are marked with an asterisk at the beginning of the tip names. Species ID can be found in **Tab. S1**. Bootstrap values are shown for each node.

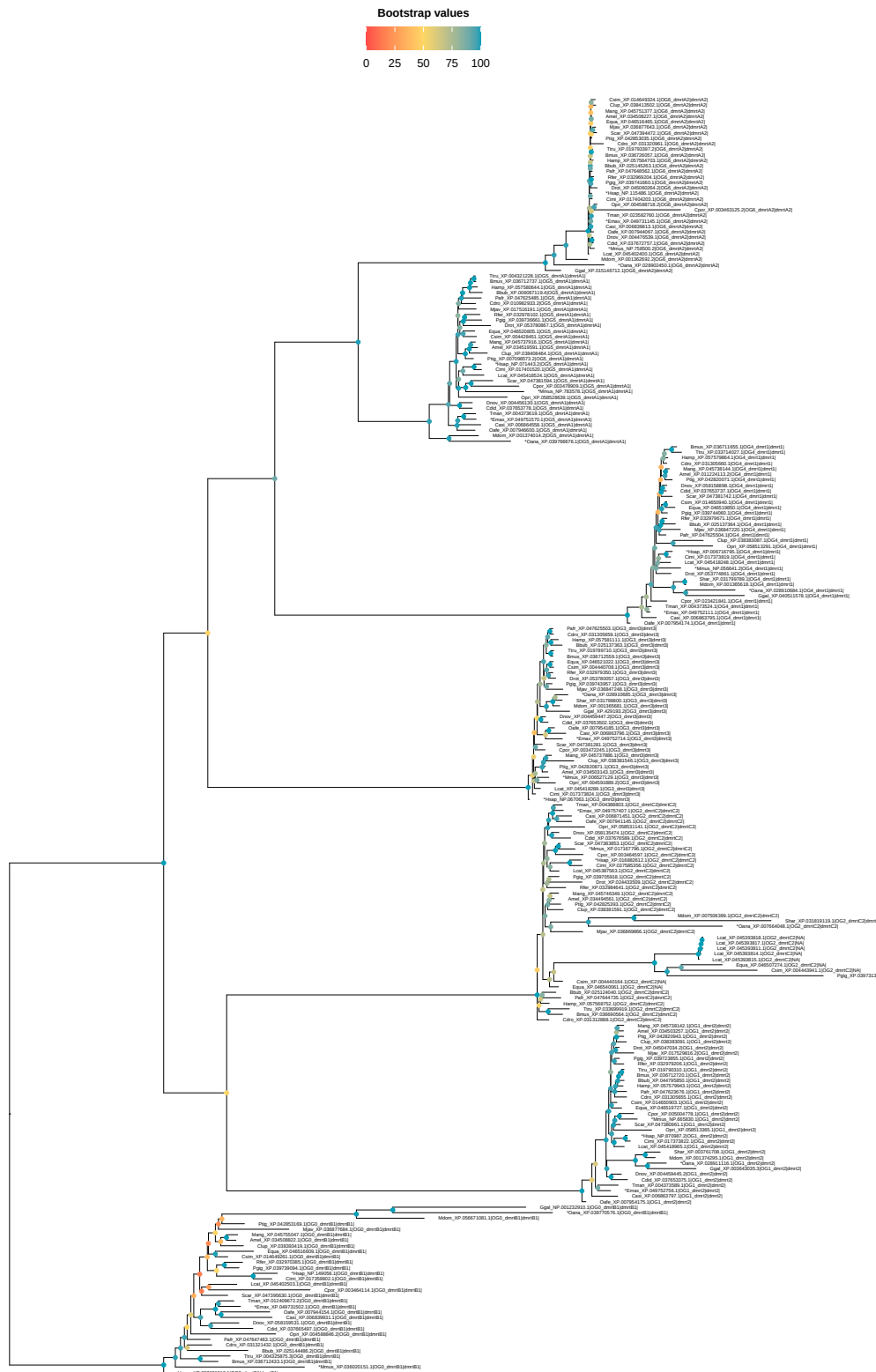


Supplementary Figure S3 – ML phylogenetic tree of the Fox gene family in molluscs, including the possvm orthology inference. Reference genes from *H. sapiens*, *C. elegans*, and *D. melanogaster* are marked with an asterisk at the beginning of the tip names. Species ID can be found in **Tab. S1**. Bootstrap values are shown for each node.

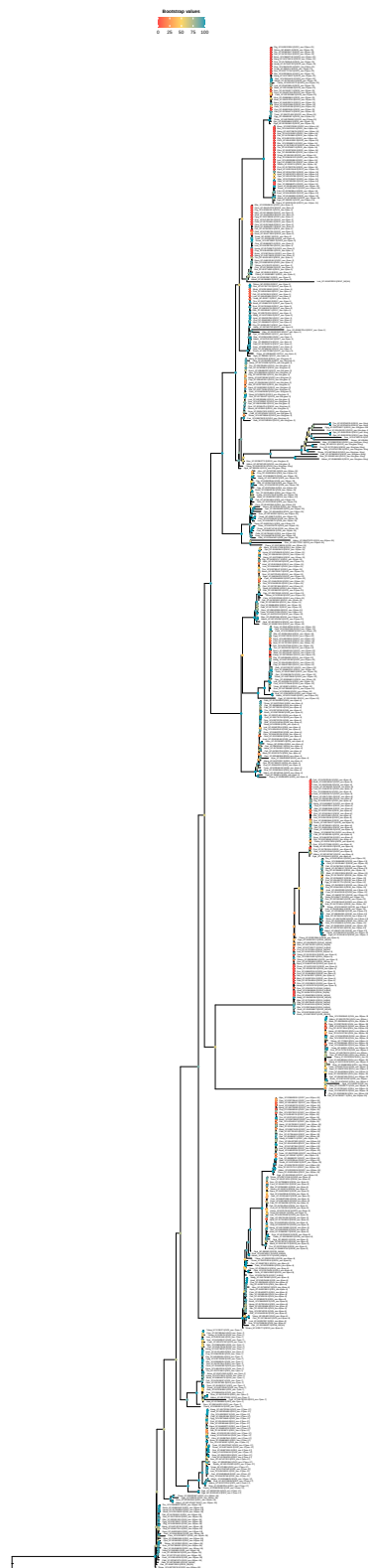


Supplementary Figure S4 – The DSFG complement in Mammalia (A) and *Drosophila* spp. (B).

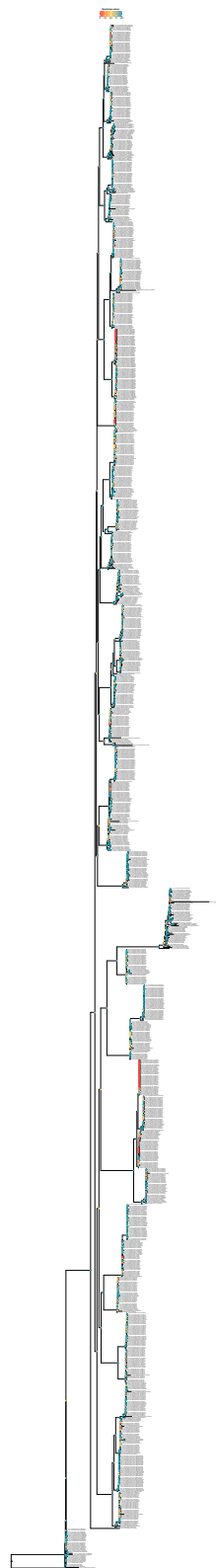
Presence/absence of genes in various species are indicated by filled circles. Numbers inside each circle specify genes with 2 or more copies. The shaded area highlights outgroup species, *Gallus gallus* (Aves) for mammals and *Anopheles gambiae* (Culicidae) for fruit flies. The phylogenetic tree of analysed species, as inferred from literature, is shown on the left, while major taxonomic groups are reported on the right. All species are represented by genomic data. Dmrt, Sox, and Fox gene (DSFG) trees are shown on the bottom (full trees can be found in Fig. S5 and S7). Full species names for both mammals and fruit flies, along with all assembly and taxonomic information, can be found in Tab. S4 and S5, respectively. A.: Aves; Chirop.: Chiroptera; L.: Lagomorpha; M.: Monotremata; Me.: Metatheria; P.: Pholidota; Pe.: Perissodactyla; Prim.: Primates; Roden.: Rodentia; X.: Xenarthra; C.: Culicidae.



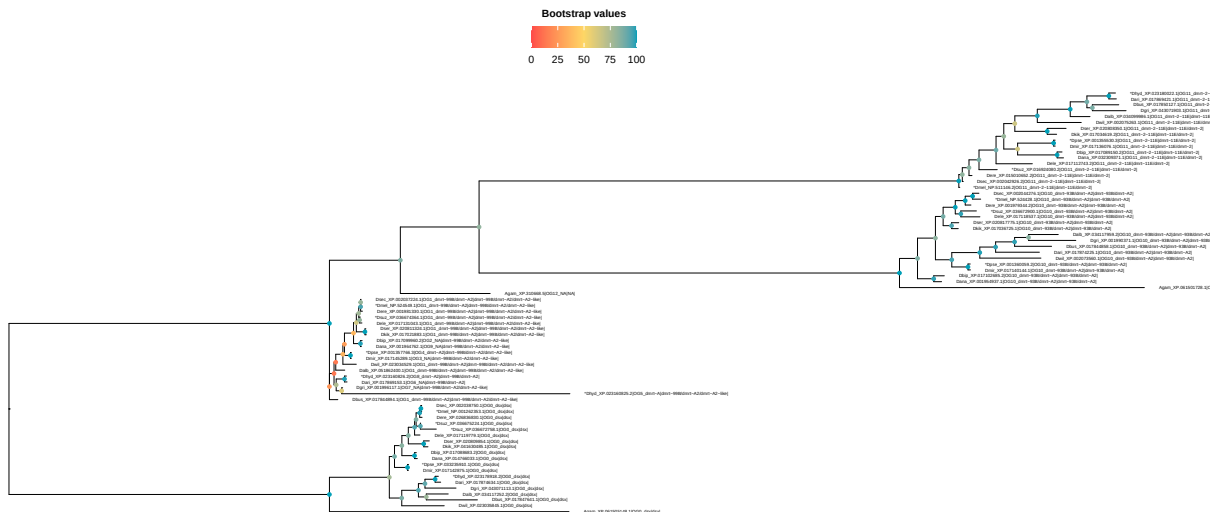
Supplementary Figure S5 – ML phylogenetic tree of the *dsx* and *mab-3* related transcription factor (Dmrt) gene family in mammals, including the Possvm orthology inference. Reference genes from *H. sapiens*, *Mus musculus*, *Elephas maximus indicus*, and *Ornithorhynchus anatinus* are marked with an asterisk at the beginning of the tip names. Species ID can be found in **Tab. S4**. The tree has been midpoint rooted. Bootstrap values are shown for each node.



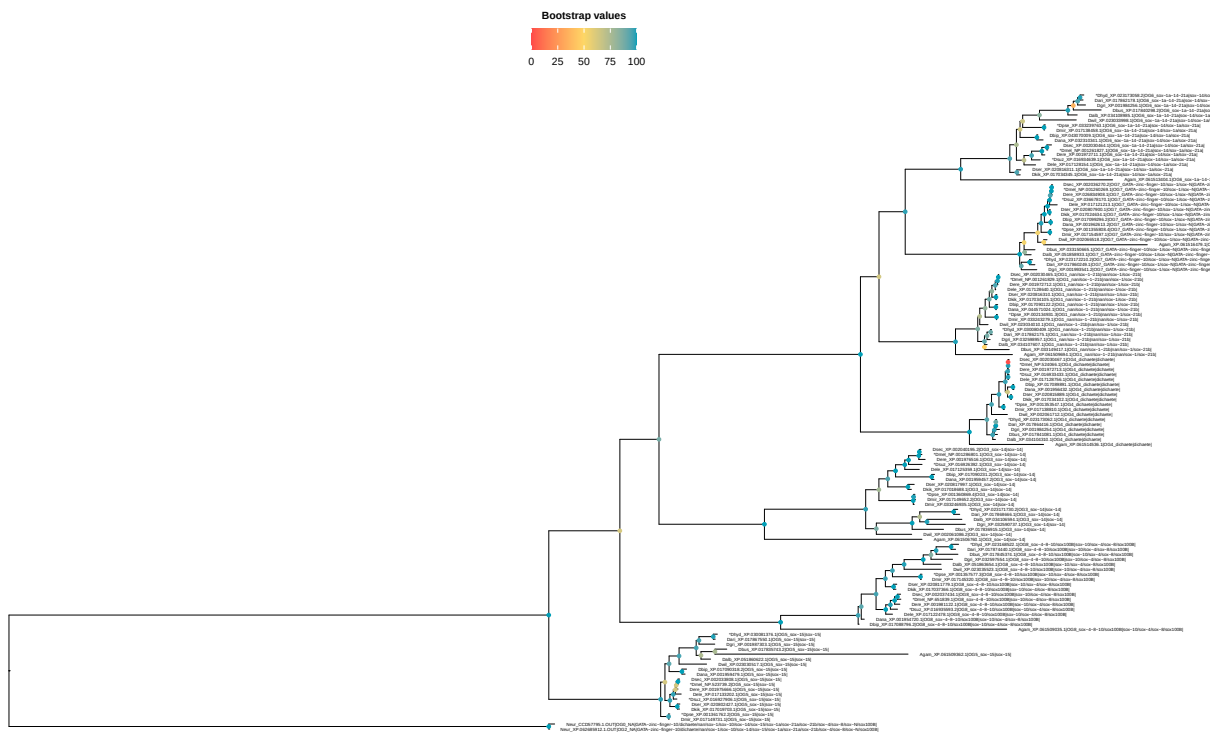
Supplementary Figure S6 – ML phylogenetic tree of the *Sry*-related HMG-box (Sox) gene family in mammals, including the Possvm orthology inference. Reference genes from *H. sapiens*, *M. musculus*, *E. maximus indicus*, and *O. anatinus* are marked with an asterisk at the beginning of the tip names. Species ID can be found in **Tab. S4**. Bootstrap values are shown for each node.



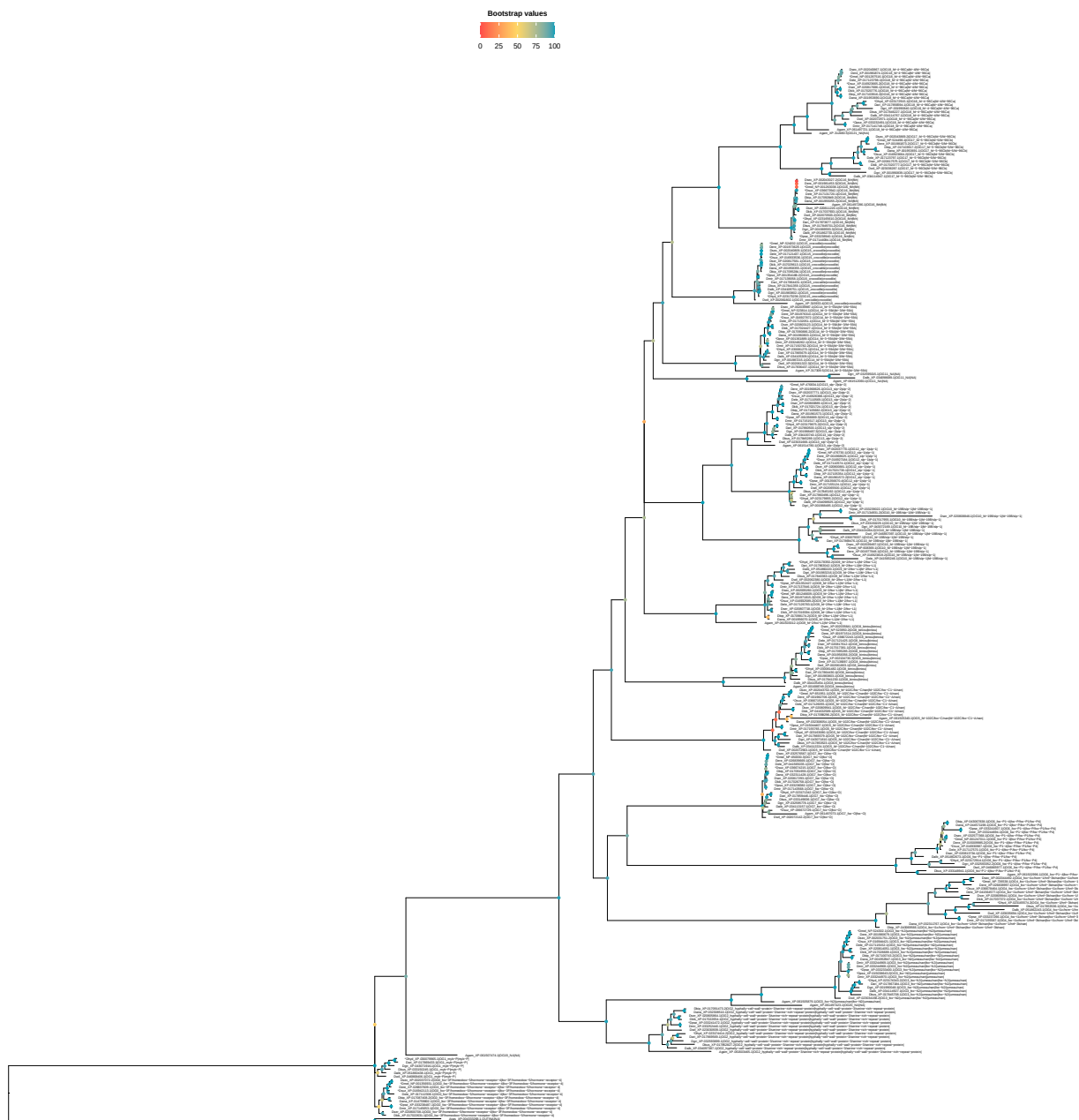
Supplementary Figure S7 – ML phylogenetic tree of the forkhead box (Fox) gene family in mammals, including the Possvm orthology inference. Reference genes from *H. sapiens*, *M. musculus*, *E. maximus indicus*, and *O. anatinus* are marked with an asterisk at the beginning of the tip names. Species ID can be found in **Tab. S4**. Bootstrap values are shown for each node.



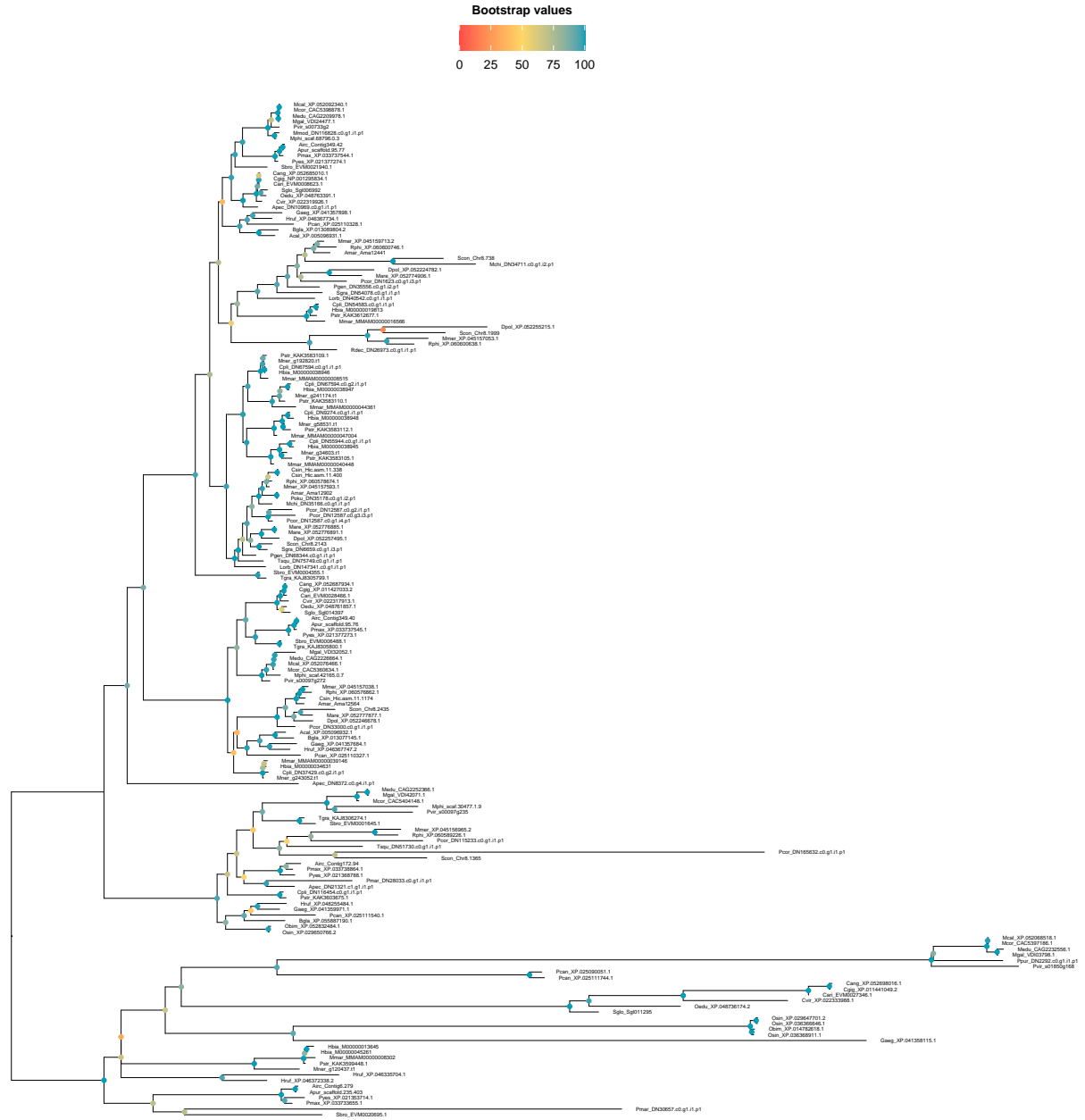
Supplementary Figure S8 – ML phylogenetic tree of the Dmrt gene family in fruit flies, including the Possvm orthology inference. Reference genes from *D. melanogaster*, *Drosophila hydei*, *Drosophila pseudoobscura*, and *Drosophila suzukii* are marked with an asterisk at the beginning of the tip names. Species ID can be found in **Tab. S5**. The tree has been midpoint rooted. Bootstrap values are shown for each node.



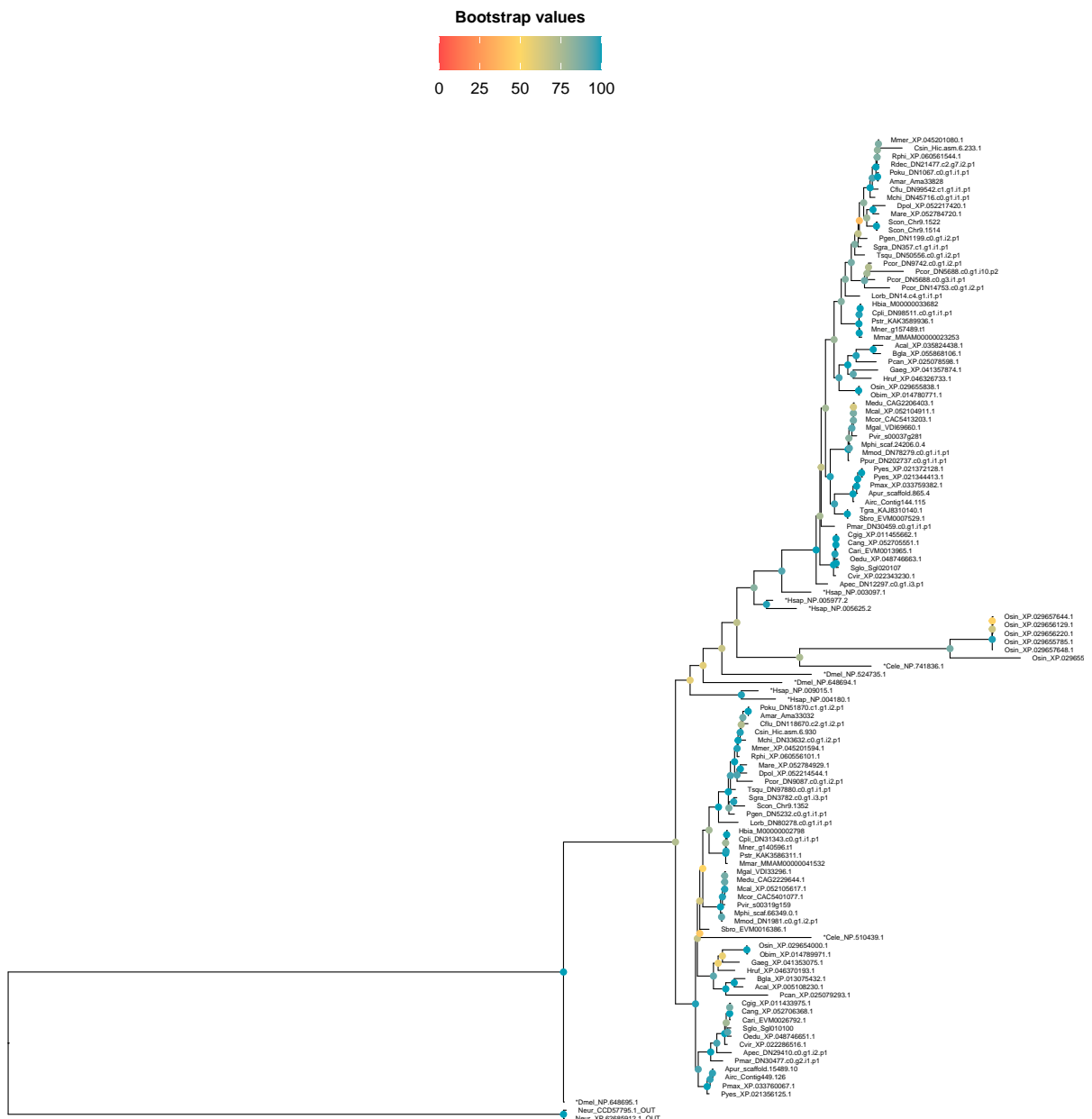
Supplementary Figure S9 – ML phylogenetic tree of the Sox gene family in fruit flies, including the Possvm orthology inference. Reference genes from *D. melanogaster*, *D. hydei*, *D. pseudoobscura*, and *D. suzukii* are marked with an asterisk at the beginning of the tip names. Species ID can be found in **Tab. S5**. Bootstrap values are shown for each node.



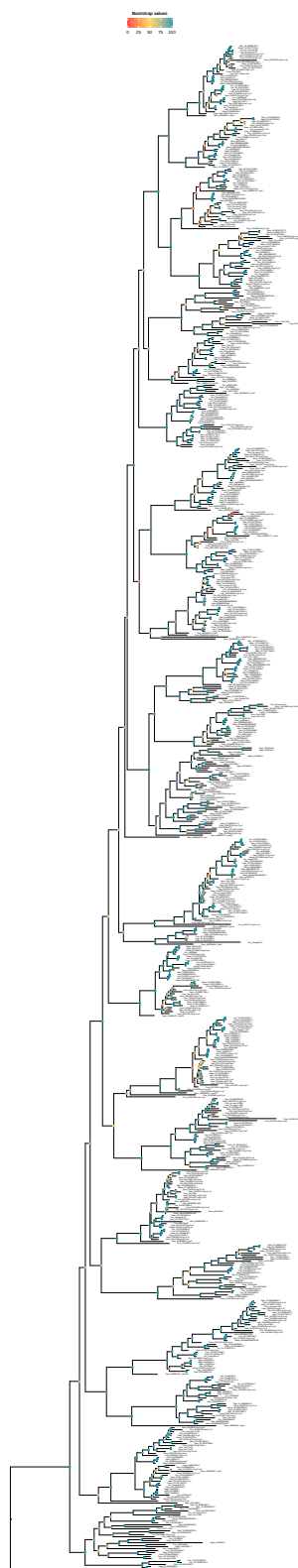
Supplementary Figure S10 – ML phylogenetic tree of the Fox gene family in fruit flies, including the Possvm orthology inference. Reference genes from *D. melanogaster*, *D. hydei*, *D. pseudoobscura*, and *D. suzukii* are marked with an asterisk at the beginning of the tip names. Species ID can be found in **Tab. S5**. Bootstrap values are shown for each node.



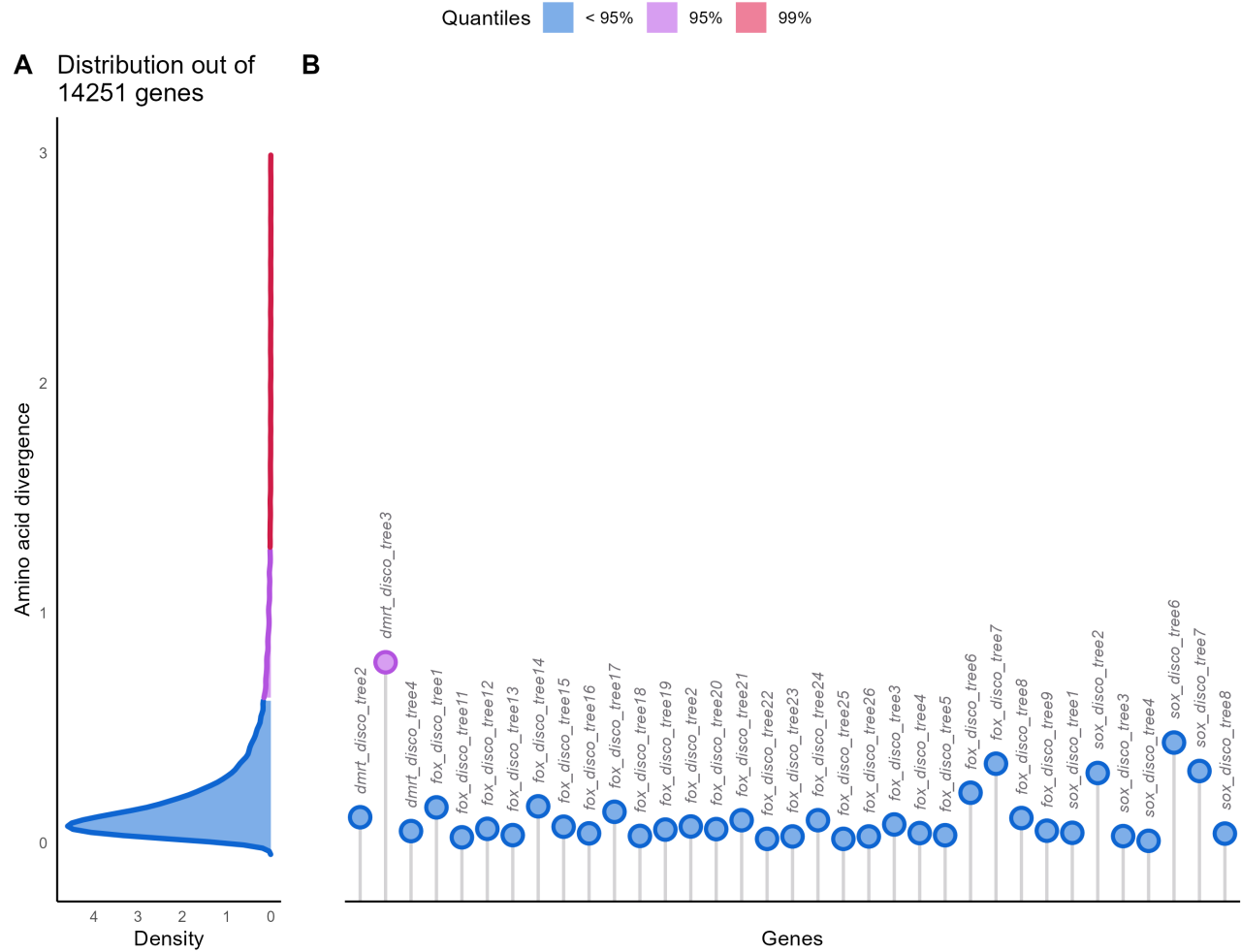
Supplementary Figure S11 – ML phylogenetic tree of the Dmrt gene family in mollusc species.
 Species ID can be found in **Tab. S1**. The tree has been midpoint rooted. Bootstrap values are shown for each node.



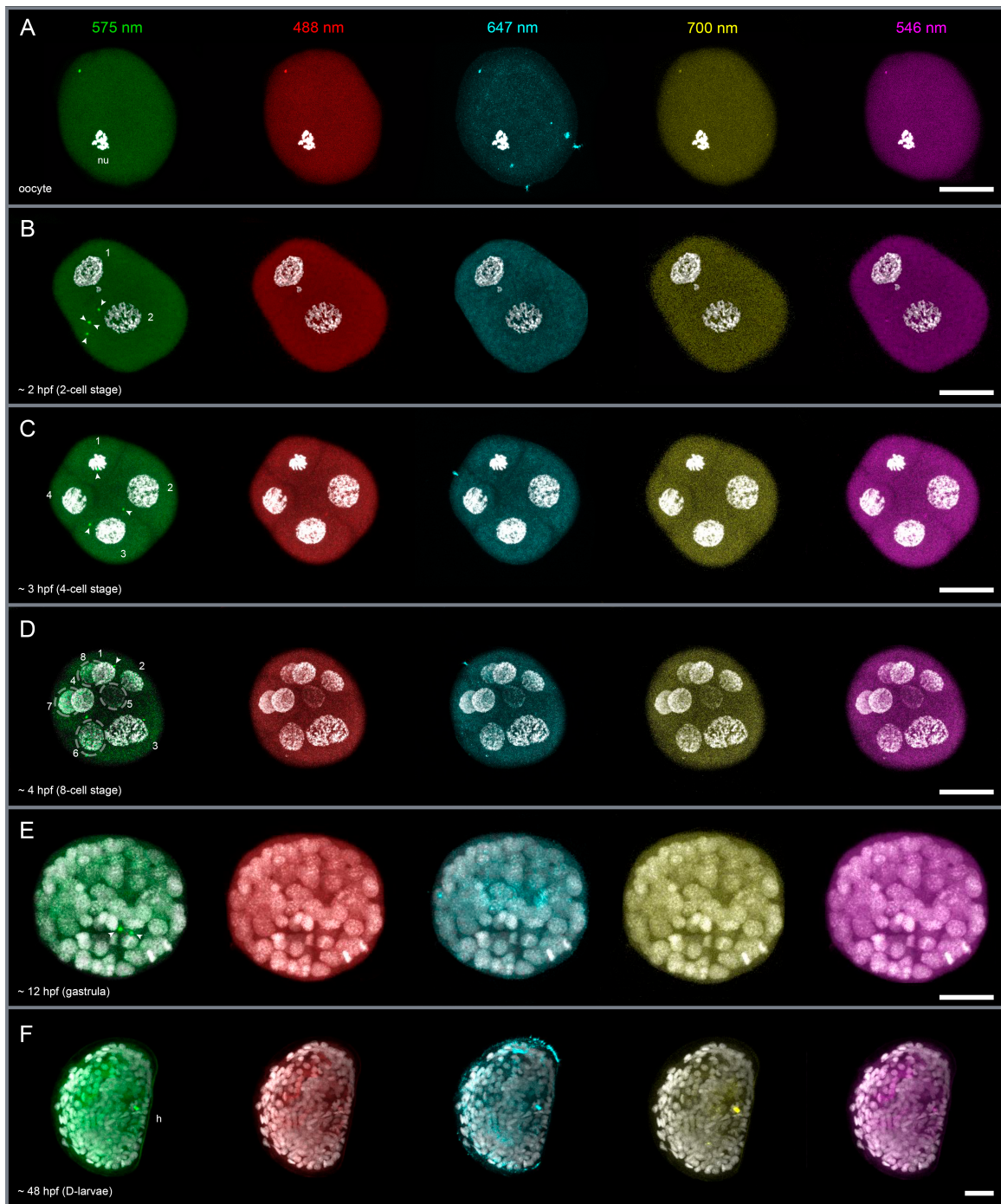
Supplementary Figure S12 – ML phylogenetic tree of *Sox-B1* and *Sox-B2* genes in mollusc and reference species. Reference genes from *H. sapiens*, *C. elegans*, and *D. melanogaster* are marked with an asterisk at the beginning of the tip names. Species ID can be found in **Tab. S1**. Bootstrap values are shown for each node.



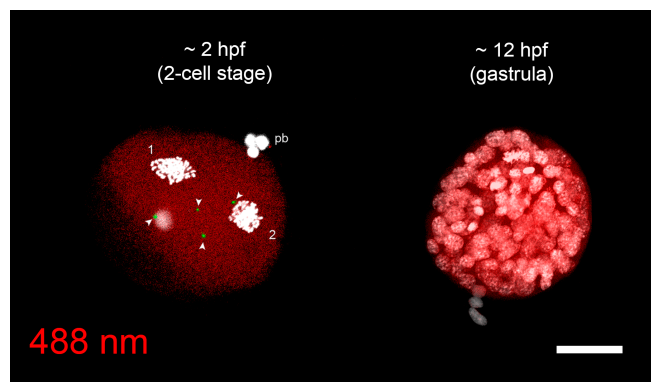
Supplementary Figure S14 – ML phylogenetic tree of the Fox gene family in bivalves and the sea urchin *Strongylocentrotus purpuratus* (Spur). Reference genes from *S. purpuratus* are marked with an asterisk at the beginning of the tip names. Species ID can be found in **Tab. S1**. *S. purpuratus* genes are those given by **Tu et al., 2006**. Bootstrap values are shown for each node.



Supplementary Figure S15 – Distribution of amino acid sequence divergence (AASD) of single-copy orthogroups in *Crassostrea gigas*, *Crassostrea angulata*, *Crassostrea ariakensis*, and *Crassostrea virginica* (A), including DSFG (B). The distribution of AASD in *Crassostrea* has been computed on the median values of pairwise distances of over 14 k single-copy orthogroups (SCOs). Circle heights of DSFGs show the median value of their AASD. *Dmrt-1L* genes are indicated as ‘dmrt₋disco₋tree3’.



Supplementary Figure S16 – MitoTracker staining and negative controls of mRNA *in-situ* hybridization chain reaction (HCR) in *Mytilus galloprovincialis* (A) oocyte, (B) 2-cell male embryo, (C) 4-cell female embryo, (D) 8-cell female embryo, (E) 12 hours post fertilization [hpf] embryo (gastrula), and (F) 48 hpf larvae (D-veliger). Nuclei are shown in white; in the 2-, 4-, and 8-cell stages, nuclei are also marked with numbers; in the 8-cell stage, nuclei of blastomeres in the background are highlighted with dashed circles. Sperm mitochondria, when stained (shown in green), are marked with arrowheads. Acquisition channels are indicated on top, and colours are the same as in Fig. 5.3. h: hinge; nu: oocyte nucleus. Scale bar: 20 µm.



Supplementary Figure S17 – Mito-Tracker staining and negative controls of Vasa immunolocalization in *M. galloprovincialis* embryos. Nuclei are shown in white. Sperm mitochondria (in green) are marked with arrowheads. pb: polar body. Scale bar: 20 μ m.

Supplementary tables

All the supplementary tables are available in a parsable version at the following GitHub repository:

https://github.com/filonico/phd_thesis_tex

Supplementary Table S1 – Genomic and transcriptomic data of bivalves and other molluscs. For each species, the relative ID, taxonomic information, BUSCO statistics, NCBI accession number, and source publication are reported. Biv: Bivalvia; Ca: Cenogastropoda; Cep: Cephalopoda; Co: Coleoidea; Gas: Gastro poda; Gen: Genome; He: Heterobranchia; Im: Imparidentia; Im: Heterobranchia; Ne: Neomphaliones; Pa: Palaeoheterodonta; Pt: Pteriomorpha; Tra: Transcriptome; Ve: Veti gastropoda

Species	ID	Class	Group	Order	Type	Reduced dataset	BUSCO statistics (metazoa_odb10)	NCBI acc. no.	Reference	Annotation source
<i>Magallana (Crassostrea) angulata</i>	Cang	Biv	Pt	Ostreida	Gen	No	C:99.1%[S:97.1%,D:2.0%] F:0.3%[M:0.6%]	GCF_025612915.1	Teng et al., 2023	<u>NCBI</u>
<i>Magallana (Crassostrea) gigas</i>	Cgig	Biv	Pt	Ostreida	Gen	Yes	C:98.2%[S:93.1%,D:5.1%] F:0.4%[M:1.4%]	GCF_902806645.1	Penaloza et al., 2021	<u>NCBI</u>
<i>Magallana (Crassostrea) ariakensis</i>	Cari	Biv	Pt	Ostreida	Gen	No	C:94.8%[S:91.2%,D:3.6%] F:0.7%[M:4.5%]	GCA_020567875.1	Li et al., 2021	<u>FigShare</u>
<i>Crassostrea virginica</i>	Cvir	Biv	Pt	Ostreida	Gen	Yes	C:98.2%[S:73.1%,D:25.1%] F:0.3%[M:1.5%]	GCF_002022765.2	Gómez-Chiarri et al., 2015	<u>NCBI</u>
<i>Ostrea edulis</i>	Oedu	Biv	Pt	Ostreida	Gen	Yes	C:98.7%[S:97.8%,D:0.9%] F:0.5%[M:0.8%]	GCF_947568905.1	Darwin Tree of Life	<u>NCBI</u>
<i>Saccostrea glomerata</i>	Sglo	Biv	Pt	Ostreida	Gen	No	C:89.1%[S:85.5%,D:3.6%] F:4.9%[M:6.0%]	GCA_003671525.1	Powell et al., 2018	<u>dbSRG</u>
<i>Atrina pectinata</i>	Apct	Biv	Pt	Ostreida	Tra	Yes	C:95.6%[S:93.1%,D:2.5%] F:1.9%[M:2.5%]	DRR348924,-25,-26	Shimizu et al., 2022	–
<i>Pinctada margaritifera</i>	Pmar	Biv	Pt	Ostreida	Tra	Yes	C:94.3%[S:93.9%,D:0.4%] F:1.7%[M:4.0%]	SRR1039667 SRR1041217	Teaniniuraitemoana et al., 2014	–
<i>Mytilus galloprovincialis</i>	Mgal	Biv	Pt	Mytilida	Gen	Yes	C:80.5%[S:50.4%,D:30.1%] F:8.6%[M:10.9%]	GCA_900618805.1	Gerdol et al., 2020	<u>NCBI</u>
<i>Mytilus edulis</i>	Medu	Biv	Pt	Mytilida	Gen	No	C:83.8%[S:70.9%,D:12.9%] F:5.1%[M:11.1%]	GCA_905397895.1	Corrochano-Fraile et al., 2022	<u>NCBI</u>
<i>Mytilus unguliculatus (conicus)</i>	Mcor	Biv	Pt	Mytilida	Gen	No	C:80.8%[S:78.8%,D:2.0%] F:4.3%[M:14.9%]	GCA_011752425.2	Yang et al., 2021	<u>NCBI</u>
<i>Mytilus californianus</i>	Mcal	Biv	Pt	Mytilida	Gen	No	C:96.2%[S:95.0%,D:1.2%] F:0.4%[M:3.4%]	GCF_021869535.1	Paggeot et al., 2022	<u>NCBI</u>
<i>Perna viridis</i>	Pvir	Biv	Pt	Mytilida	Gen	Yes	C:99.4%[S:99.0%,D:0.4%] F:0.2%[M:0.4%]	GCA_018227765.1	Inoue et al., 2021	<u>Google Drive</u>
<i>Modiolus modiolus</i>	Mmod	Biv	Pt	Mytilida	Tra	Yes	C:95.7%[S:92.3%,D:3.4%] F:2.1%[M:2.2%]	SRR5043294	Meng et al., 2018	–
<i>Modiolus philippinarum</i>	Mphi	Biv	Pt	Mytilida	Gen	No	C:64.9%[S:63.0%,D:1.9%] F:18.8%[M:16.3%]	GCA_002080025.1	Sun et al., 2017	<u>Dryad</u>
<i>Perumytilus purpuratus</i>	Ppur	Biv	Pt	Mytilida	Tra	Yes	C:84.2%[S:83.3%,D:0.9%] F:11.8%[M:4.0%]	SRR4343820	Briones et al., 2018	–
<i>Argopecten irradians concentricus</i>	Airc	Biv	Pt	Pectinida	Gen	Yes	C:94.9%[S:94.0%,D:0.9%] F:3.6%[M:1.5%]	GCA_004382765.1	Liu et al., 2020	<u>Dryad</u>
<i>Argopecten purpuratus</i>	Apur	Biv	Pt	Pectinida	Gen	No	C:89.2%[S:88.6%,D:0.6%] F:5.0%[M:5.8%]	–	Liu et al., 2020	<u>GigaDB</u>

Tab. S1 continued from previous page

Species	ID	Class	Group	Order	Type	Reduced dataset	BUSCO statistics ('metazoa_odb10')	NCBI acc. no.	Reference	Annotation source
<i>Pecten maximus</i>	Pmax	Biv	Pt	Pectinida	Gen	Yes	C:98.5%[S:94.5%,D:4.0%], F:0.4%,M:1.1%	GCF_902052985.1	Kenny et al., 2020	NCBI
<i>Mizuhoplecten (Patinoplecten) yessensis</i>	Pyes	Biv	Pt	Pectinida	Gen	Yes	C:98.3%[S:96.1%,D:2.2%], F:0.5%,M:1.2%	GCF_002113885.1	Wang, Zhang, et al., 2017	NCBI
<i>Anadara (Scapharca) broughtoni</i>	Sbro	Biv	Pt	Arcida	Gen	Yes	C:91.2%[S:85.6%,D:5.6%], F:2.6%,M:6.2%	—	Bai et al., 2019	GigaDB
<i>Tegillarca granosa</i>	Tgra	Biv	Pt	Arcida	Gen	Yes	C:70.6%[S:61.3%,D:9.3%], F:11.7%,M:17.7%	GCA_029721355.1	—	NCBI
<i>Ruditapes decussatus</i>	Rdec	Biv	Im	Venerida	Tra	Yes	C:84.8%[S:84.1%,D:0.7%], F:7.3%,M:7.9%	SRR527740,-41,-43,-44,-47,-51,-52,-57	Ghiselli et al., 2018	—
<i>Ruditapes philippinarum</i>	Rphi	Biv	Im	Venerida	Gen	Yes	C:97.8%[S:85.5%,D:12.3%], F:0.7%,M:1.5%	GCF_026671515.1	Xu, Marteossi, et al., 2022	NCBI
<i>Mercenaria mercenaria</i>	Mmer	Biv	Im	Venerida	Gen	No	C:96.0%[S:89.8%,D:6.2%], F:1.0%,M:3.0%	GCF_021730395.1	Farhat et al., 2022	NCBI
<i>Cyclina sinensis</i>	Csin	Biv	Im	Venerida	Gen	Yes	C:94.1%[S:83.9%,D:10.2%], F:1.8%,M:4.1%	GCA_012932295.1	Wei et al., 2020	Dryad
<i>Calyptogena (Archivesica) marisnica</i>	Amar	Biv	Im	Venerida	Gen	No	C:82.1%[S:80.1%,D:2.0%], F:6.0%,M:11.9%	GCA_014843695.1	Ip et al., 2021	FigShare
<i>Phreagena okutanii</i>	Poku	Biv	Im	Venerida	Tra	Yes	C:92.9%[S:85.8%,D:7.1%], F:3.0%,M:4.1%	SRR7156763,-64,-65,-66,-67,-68	Lan et al., 2019	—
<i>Corbicula fluminea</i>	Cflu	Biv	Im	Venerida	Tra	Yes	C:83.7%[S:79.9%,D:3.8%], F:10.3%,M:6.0%	SRR1559272,SRR512046	González et al., 2015 Zhu et al., 2019	—
<i>Mactra chinensis</i>	Mchi	Biv	Im	Venerida	Tra	Yes	C:81.5%[S:80.8%,D:0.7%], F:10.2%,M:8.3%	SRR1263980	—	—
<i>Mya arenaria</i>	Mare	Biv	Im	Myida	Gen	Yes	C:98.5%[S:80.4%,D:18.1%], F:0.4%,M:1.1%	GCF_026914265.1	Hart et al., 2023	NCBI
<i>Dreissena polymorpha</i>	Dpol	Biv	Im	Myida	Gen	Yes	C:97.2%[S:80.1%,D:17.1%], F:0.4%,M:2.4%	GCF_020536995.1	McCartney et al., 2022	NCBI
<i>Pisidium coreanum</i>	Pcor	Biv	Im	Sphaeriida	Tra	Yes	C:92.7%[S:90.0%,D:2.7%], F:2.5%,M:4.8%	SRR6474597	—	—
<i>Solen grandis</i>	Sgra	Biv	Im	Adapedonta	Tra	Yes	C:94.5%[S:81.6%,D:12.9%], F:3.6%,M:1.9%	SRR5484647, SRR5485368, SRR5499447	Nie et al., 2018	—
<i>Sinonovacula constricta</i>	Scon	Biv	Im	Adapedonta	Gen	Yes	C:90.8%[S:79.2%,D:11.6%], F:3.5%,M:5.7%	GCA_007844125.1	Ran et al., 2019	—
<i>Panopea generosa</i>	Pgen	Biv	Im	Adapedonta	Tra	Yes	C:84.1%[S:81.9%,D:2.2%], F:9.7%,M:6.2%	SRR12216869,-70	Putnam et al., 2022	Dryad
<i>Tridacna squamosa</i>	Tsqu	Biv	Im	Cardiida	Tra	Yes	C:89.7%[S:86.9%,D:2.8%], F:3.5%,M:6.8%	SRR10824662,-65	Li, Zhou, et al., 2020	—
<i>Loripes orbiculatus</i>	Lorb	Biv	Im	Lucinida	Tra	Yes	C:76.1%[S:74.9%,D:1.2%], F:14.3%,M:9.6%	SRR1002336,-38,-39,-47	Yuen et al., 2019	—
<i>Hyriopsis bivalvata (Unio delphinus)</i>	Hbia	Biv	Pa	Unionida	Gen	Yes	C:97.5%[S:94.9%,D:2.6%], F:2.0%,M:0.5%	GCA_029339505.1	Gomes-dos-Santos et al., 2023	FigShare

Tab. S1 continued from previous page

Species	ID	Class	Group	Order	Type	Reduced dataset	BUSCO statistics ('metazoa_odb10')	NCBI acc. no.	Reference	Annotation source
<i>Cristaria pilicata</i>	Cpli	Biv	Pa	Unionida	Tra	Yes	C:93.6%[S:92.8%,D:0.8%] F:2.1%,M:4.3%	SRR2175668 SRR3095781	Pathak et al., 2016 Wang, Liu, and Wu, 2017	—
<i>Megalonaia nervosa</i>	Mner	Biv	Pa	Unionida	Gen	Yes	C:65.0%[S:63.3%,D:1.7%] F:14.0%,M:21.0%	GCA_01617855.1	Rogers et al., 2021	Dryad
<i>Potamius streckersoni</i>	Pstr	Biv	Pa	Unionida	Gen	Yes	C:94.9%[S:93.4%,D:1.5%] F:1.2%,M:3.9%	GCA_01674295.1	Smith, 2021	NCBI
<i>Margaritifera margaritifera</i>	Mmar	Biv	Pa	Unionida	Gen	Yes	C:92.6%[S:92.1%,D:0.5%] F:3.0%,M:4.4%	GCA_015947965.1	Gomes-dos-Santos et al., 2021	FigShare
<i>Aplysia californica</i>	Acal	Gas	He	Aplysiida	Gen	No	C:97.8%[S:97.0%,D:0.8%] F:0.7%,M:1.5%	GCF_000002075.1	Knudsen et al., 2006	NCBI
<i>Biomphalaria glabrata</i>	Bglia	Gas	He	—	Gen	No	C:98.9%[S:98.2%,D:0.7%] F:0.1%,M:1.0%	GCF_947242115.1	—	NCBI
<i>Pomacea canaliculata</i>	Pcan	Gas	Ca	Architaenioglossa	Gen	No	C:98.2%[S:97.0%,D:1.2%] F:0.4%,M:1.4%	GCF_003073045.1	Liu et al., 2018	NCBI
<i>Gigantopelta aegis</i>	Gaeg	Gas	Ne	Neomphalida	Gen	No	C:98.4%[S:94.2%,D:4.2%] F:0.8%,M:0.8%	GCF_016097555.1	Lan et al., 2021	NCBI
<i>Haliotis rufescens</i>	Hruf	Gas	Ve	Lepetellida	Gen	No	C:99.0%[S:98.3%,D:0.7%] F:0.0%,M:1.0%	GCF_023055435.1	—	NCBI
<i>Octopus bimaculoides</i>	Obim	Cep	Co	Octopoda	Gen	No	C:94.9%[S:94.4%,D:0.5%] F:2.3%,M:2.8%	GCF_001194135.2	Albertin et al., 2015	NCBI
<i>Octopus sinensis</i>	Osin	Cep	Co	Octopoda	Gen	No	C:98.1%[S:96.9%,D:1.2%] F:0.9%,M:1.0%	GCF_006345805.1	Li, Bian, et al., 2020	NCBI

Supplementary Table S2 – Dmrt, Sox, and Fox gene (DSFG) family and domain identifiers (IDs) in PANTHER and CDD, respectively. After having retrieved putative DSFGs on the basis of hidden Markov model (HMM) profiles, IDs have been used to retain only reliable hits.

Gene family	PANTHER/CDD	ID	Description
Dmrt	CDD	gml CDD 214606	Doublesex DNA-binding motif
	CDD	gml CDD 425850	DM DNA binding domain
	PANTHER	PTHR12322	DOUBLESEX AND MAB-3 RELATED TRANSCRIPTION FACTOR DMRT PROTEIN CBR-MAB-23
	PANTHER	PTHR12322-SF116	DOUBLESEX- AND MAB-3-RELATED TRANSCRIPTION FACTOR 1
	PANTHER	PTHR12322-SF118	DOUBLESEX- AND MAB-3-RELATED TRANSCRIPTION FACTOR DMD-4
	PANTHER	PTHR12322-SF123	DOUBLESEX- AND MAB-3-RELATED TRANSCRIPTION FACTOR 2
	PANTHER	PTHR12322-SF53	DOUBLESEX- AND MAB-3-RELATED TRANSCRIPTION FACTOR A1
	PANTHER	PTHR16897-SF71	STRESS RESPONSE PROTEIN NST1
	PANTHER	PTHR46888-SF11	RIBONUCLEASE H
	CDD	gml CDD 432488	SOX transcription factor
	CDD	gml CDD 432558	Sox developmental protein N terminal
	CDD	gml CDD 438790	high mobility group (HMG)-box found in group B SRY-related high-mobility group (HMG) box (Sox) transcription factors
	CDD	gml CDD 438820	high mobility group (HMG)-box found in sex-determining region Y (SRY)-box (SOX) family transcription factors
Sox	CDD	gml CDD 438837	high mobility group (HMG)-box found in group A, group B and group G of SRY-related high-mobility group (HMG) box (Sox) transcription factors
	CDD	gml CDD 438838	high mobility group (HMG)-box found in group C SRY-related high-mobility group (HMG) box (Sox) transcription factors
	CDD	gml CDD 438839	high mobility group (HMG)-box found in group D SRY-related high-mobility group (HMG) box (Sox) transcription factors
	CDD	gml CDD 438840	high mobility group (HMG)-box found in group E SRY-related high-mobility group (HMG) box (Sox) transcription factors
	CDD	gml CDD 438841	high mobility group (HMG)-box found in group F SRY-related high-mobility group (HMG) box (Sox) transcription factors
	CDD	gml CDD 438842	high mobility group (HMG)-box found in sex-determining region Y (SRY)-box 30 (SOX30) and similar proteins
	CDD	gml CDD 438843	high mobility group (HMG)-box found in sex-determining region Y (SRY)-box 30 (SOX30) and similar proteins
	CDD	gml CDD 438844	high mobility group (HMG)-box found in sex-determining region Y (SRY)-box 15 (SOX15) and similar proteins
	CDD	gml CDD 438845	high mobility group (HMG)-box found in sex determining region Y (SRY)-box 4 (SOX4) and similar proteins
	CDD	gml CDD 438846	high mobility group (HMG)-box found in sex determining region Y (SRY)-box 11 (SOX11) and similar proteins
	CDD	gml CDD 438847	high mobility group (HMG)-box found in sex determining region Y (SRY)-box 12 (SOX12) and similar proteins
	CDD	gml CDD 438849	high mobility group (HMG)-box found in sex determining region Y (SRY)-box 7 (SOX7) and similar proteins
	CDD	gml CDD 438850	high mobility group (HMG)-box found in sex determining region Y (SRY)-box 17 (SOX17) and similar proteins
	PANTHER	PTHR10270-SF14	TRANSCRIPTION FACTOR SOX-14
	PANTHER	PTHR10270-SF161	SOX DOMAIN-CONTAINING PROTEIN DICHAETE-RELATED
	PANTHER	PTHR10270-SF199	SEX-DETERMINING REGION Y PROTEIN
	PANTHER	PTHR10270-SF231	TRANSCRIPTION FACTOR SOX-2
	PANTHER	PTHR10270-SF27	TRANSCRIPTION FACTOR SOX-4
	PANTHER	PTHR10270-SF313	TRANSCRIPTION FACTOR SOX-21
	PANTHER	PTHR10270-SF315	TRANSCRIPTION FACTOR SOX-1A-RELATED
	PANTHER	PTHR10270-SF317	TRANSCRIPTION FACTOR SOX-1B-RELATED
	PANTHER	PTHR10270-SF322	TRANSCRIPTION FACTOR SOX-3
	PANTHER	PTHR10270-SF324	TRANSCRIPTION FACTOR SOX-3
	PANTHER	PTHR10270-SF326	TRANSCRIPTION FACTOR SOX-3
	PANTHER	PTHR10270-SF328	TRANSCRIPTION FACTOR SOX-3
	PANTHER	PTHR10270-SF329	TRANSCRIPTION FACTOR SOX-3
	PANTHER	PTHR45789-SF2	F118025P1
	PANTHER	PTHR45789-SF72	TRANSCRIPTION FACTOR SOX-9
	PANTHER	PTHR45803-SF1	SOX100B
	PANTHER	PTHR45803-SF5	TRANSCRIPTION FACTOR SOX-30
	PANTHER	PTHR45803	
	PANTHER	PTHR47219-SF1	
	PANTHER	PTHR47219-SF29	
Fox	CDD	gml CDD 410788	Forkhead (FH) domain found in Forkhead box (FOX) family of transcription factors and similar proteins
	CDD	gml CDD 410789	Forkhead (FH) domain found in Forkhead box protein A (FOXA) subfamily
	CDD	gml CDD 410790	Forkhead (FH) domain found in Forkhead box protein B (FOXB) subfamily
	CDD	gml CDD 410791	Forkhead (FH) domain found in Forkhead box protein C (FOXC) subfamily
	CDD	gml CDD 410792	Forkhead (FH) domain found in Forkhead box protein D (FOXD) subfamily
	CDD	gml CDD 410793	Forkhead (FH) domain found in Forkhead box protein E (FOXE) subfamily
	CDD	gml CDD 410794	Forkhead (FH) domain found in Forkhead box protein F (FOXF) subfamily
	CDD	gml CDD 410795	Forkhead (FH) domain found in Forkhead box protein G (FOXG) subfamily
	CDD	gml CDD 410796	Forkhead (FH) domain found in Forkhead box protein H (FOXH) subfamily
	CDD	gml CDD 410797	Forkhead (FH) domain found in Forkhead box protein J (FOXJ1) and similar proteins
	CDD	gml CDD 410798	Forkhead (FH) domain found in Forkhead box proteins FOXJ2, FOXJ3 and similar proteins
	CDD	gml CDD 410799	Forkhead (FH) domain found in Forkhead box protein I (FOXI) subfamily
	CDD	gml CDD 410800	Forkhead (FH) domain found in Forkhead box protein K (FOXK) subfamily

Tab. S2 continued from previous page

Gene family	PANTHER/CDD	ID	Description
CDD	gml CDD 410801		Forkhead (FH) domain found in Forkhead box protein L1 (FOXL1) and similar proteins
CDD	gml CDD 410802		Forkhead (FH) domain found in Forkhead box protein L2 (FOXL2) and similar proteins
CDD	gml CDD 410803		Forkhead (FH) domain found in the Forkhead box protein M (FOXM1) subfamily
CDD	gml CDD 410804		Forkhead (FH) domain found in Forkhead box protein N1 (FOXNL1) and similar proteins
CDD	gml CDD 410805		Forkhead (FH) domain found in Forkhead box protein N2 (FOXNL2) and similar proteins
CDD	gml CDD 410806		Forkhead (FH) domain found in the Forkhead box protein O (FOXO) subfamily
CDD	gml CDD 410807		Forkhead (FH) domain found in the Forkhead box protein P (FOXP) subfamily
CDD	gml CDD 410808		Forkhead (FH) domain found in Forkhead box protein Q1 (FOXQ1) and similar proteins
CDD	gml CDD 410809		Forkhead (FH) domain found in Forkhead box protein Q2 (FOXQ2) and similar proteins
CDD	gml CDD 410810		Forkhead (FH) domain found in the Forkhead box protein R (FOXR) subfamily
CDD	gml CDD 410811		Forkhead (FH) domain found in Forkhead box protein S1 (FOXS1)
CDD	gml CDD 410812		Forkhead (FH) domain found in Forkhead box protein A1 (FOXA1) and similar proteins
CDD	gml CDD 410813		Forkhead (FH) domain found in Forkhead box protein A2 (FOXA2) and similar proteins
CDD	gml CDD 410814		Forkhead (FH) domain found in Forkhead box protein A3 (FOXA3) and similar proteins
CDD	gml CDD 410816		Forkhead (FH) domain found in Forkhead box protein B1 (FOXB1) and similar proteins
CDD	gml CDD 410817		Forkhead (FH) domain found in the Forkhead box protein B2 (FOXB2) and similar proteins
CDD	gml CDD 410818		Forkhead (FH) domain found in Forkhead box protein C1 (FOXC1) and similar proteins
CDD	gml CDD 410819		Forkhead (FH) domain found in Forkhead box protein C2 (FOXC2) and similar proteins
CDD	gml CDD 410820		Forkhead (FH) domain found in Forkhead box proteins FOXD1, FOXD2 and similar proteins
CDD	gml CDD 410821		Forkhead (FH) domain found in Forkhead box protein D3 (FOXD3) and similar proteins
CDD	gml CDD 410822		Forkhead (FH) domain found in Forkhead box protein D4 (FOXD4) and similar proteins
CDD	gml CDD 410823		Forkhead (FH) domain found in Forkhead box protein E1 (FOXE1) and similar proteins
CDD	gml CDD 410824		Forkhead (FH) domain found in Forkhead box protein F2 (FOXF2) and similar proteins
CDD	gml CDD 410825		Forkhead (FH) domain found in Forkhead box protein G2 (FOGX2) and similar proteins
CDD	gml CDD 410826		Forkhead (FH) domain found in Forkhead box protein I1 (FOXI1) and similar proteins
CDD	gml CDD 410827		Forkhead (FH) domain found in Forkhead box protein K1 (FOXK1) and similar proteins
CDD	gml CDD 410828		Forkhead (FH) domain found in Forkhead box protein K2 (FOXK2) and similar proteins
CDD	gml CDD 410829		Forkhead (FH) domain found in Forkhead box protein N1 (FOXNL1)
CDD	gml CDD 410830		Forkhead (FH) domain found in Forkhead box protein N2 (FOXNL2)
CDD	gml CDD 410831		Forkhead (FH) domain found in Forkhead box protein N3 (FOXNL3)
CDD	gml CDD 410832		Forkhead (FH) domain found in Forkhead box protein O1 (FOXO1)
CDD	gml CDD 410833		Forkhead (FH) domain found in Forkhead box protein O3 (FOXO3)
CDD	gml CDD 410834		Forkhead (FH) domain found in Forkhead box protein O4 (FOXO4) and similar proteins
CDD	gml CDD 410835		Forkhead (FH) domain found in Forkhead box protein O6 (FOXO6) and similar proteins
CDD	gml CDD 410836		Forkhead (FH) domain found in Forkhead box protein P1 (FOXP1)
CDD	gml CDD 410837		Forkhead (FH) domain found in Forkhead box protein P2 (FOXP2)
CDD	gml CDD 410838		Forkhead (FH) domain found in Forkhead box protein P3 (FOXP3) and similar proteins
CDD	gml CDD 410839		Forkhead (FH) domain found in Forkhead box protein P4 (FOXP4) and similar proteins
PANTHER	gml CDD 410841		PANTHER
PANTHER	PTHR1 1829		PANTHER
PANTHER	PTHR1 1829-SF142		PANTHER
PANTHER	PTHR1 1829-SF156		PANTHER
PANTHER	PTHR1 1829-SF206		PANTHER
PANTHER	PTHR1 1829-SF209		PANTHER
PANTHER	PTHR1 1829-SF335		PANTHER
PANTHER	PTHR1 1829-SF340		PANTHER
PANTHER	PTHR1 1829-SF342		PANTHER
PANTHER	PTHR1 1829-SF448		PANTHER
PANTHER	PTHR1 1829-SF361		PANTHER
PANTHER	PTHR1 1829-SF398		PANTHER
PANTHER	PTHR1 1829-SF399		PANTHER
PANTHER	PTHR1 1829-SF401		PANTHER
PANTHER	PTHR1 3962-SF17		PANTHER
PANTHER	PTHR1 3962-SF20		PANTHER
PANTHER	PTHR1 3962-SF22		PANTHER
PANTHER	PTHR1 3962-SF26		PANTHER
PANTHER	PTHR4 5767-SF2		PANTHER
PANTHER	PTHR4 5796-SF3		PANTHER
PANTHER	PTHR4 5796-SF4		PANTHER
PANTHER	PTHR4 5881-SF3		PANTHER
PANTHER	PTHR4 5881-SF4		PANTHER
PANTHER	PTHR4 6078		PANTHER
PANTHER	PTHR4 6262		PANTHER
PANTHER	PTHR4 6262-SF2		PANTHER
Fox			

Tab. S2 continued from previous page

Gene family	PANTHER/CD0	ID	Description
Fox	PANTHER	PTHR46617	FORKHEAD BOX PROTEIN G1
	PANTHER	PTHR46617_SF3	FORKHEAD BOX PROTEIN G1
	PANTHER	PTHR46721	FORKHEAD BOX PROTEIN N1
	PANTHER	PTHR46721_SF2	FORKHEAD BOX PROTEIN N1
	PANTHER	PTHR46805	FORKHEAD BOX PROTEIN J1
	PANTHER	PTHR46878	FORKHEAD BOX PROTEIN M1
	PANTHER	PTHR46878_SF1	FORKHEAD BOX PROTEIN M1
	PANTHER	PTHR47316	FORKHEAD BOX PROTEIN H1
	PANTHER	PTHR47316_SF1	FORKHEAD BOX PROTEIN H1

Supplementary Table S3 – List of DSFGs from reference species used to assess the identity of DSFGs in molluscs. NCBI accession numbers are reported in parenthesis. Each row represents an orthology group.

<i>Homo sapiens</i>	<i>Drosophila melanogaster</i>	<i>Caenorhabditis elegans</i>	Group
Dmrt gene family			
<i>DMRT1</i> (NP_068770.2)	–	–	1
<i>DMRT2</i> (NP_006548.1)	<i>dmyt11E</i> (NP_511146.2)	–	2
<i>DMRT3</i> (NP_067063.1)	<i>dmyt93B</i> (NP_524428.1)	<i>dmd-4</i> (NP_510466.1)	3
<i>DMRT4/A1</i> (NP_071443.2)	<i>dmyt99b</i> (NP_524549.1)	<i>dmd-5</i> (NP_495138.2)	A1/2
<i>DMRT5/A2</i> (NP_115486.1)	–	–	–
<i>DMRT6/B1</i> (NP_149056.1)	–	–	–
<i>DMRT7/C2</i> (NP_001035373.1)	–	–	–
<i>DMRT8/C1</i> (NP_149042.2)	–	–	–
–	<i>dsx</i> (NP_731197.1)	–	–
–	–	<i>mab-3</i> (NP_001256882.1)	–
–	–	<i>dmd-3</i> (NP_001256883.1)	–
–	–	<i>dmd-6</i> (NP_001370045.1)	–
–	–	<i>dmd-7</i> (NP_741551.1)	–
–	–	<i>dmd-8</i> (NP_503176.2)	–
–	–	<i>dmd-9</i> (NP_500305.1)	–
–	–	<i>dmd-11</i> (NP_001379162.1)	–
–	–	<i>mab-23</i> (NP_001041089.1)	–
Sox gene family			
<i>SRY</i> (NP_003131.1)	–	–	A
<i>SOX3</i> (NP_005625.2)	<i>dichaete</i> (NP_524066.1)	<i>sox3</i> (NP_510439.1)	
<i>SOX2</i> (NP_003097.1)	<i>soxN</i> (NP_524735.1)	<i>sox2</i> (NP_741836.1)	B1
<i>SOX1</i> (NP_005977.2)	–	–	–
<i>SOX14</i> (NP_004180.1)	<i>sox21a</i> (NP_648694.1)	–	B2
<i>SOX21</i> (NP_009015.1)	<i>sox21b</i> (NP_648695.1)	–	–
<i>SOX11</i> (NP_003099.1)	<i>sox14</i> (NP_476894.1)	<i>sem-2</i> (NP_740846.1)	C
<i>SOX12</i> (NP_008874.2)	–	–	–
<i>SOX4</i> (NP_003098.1)	–	–	–
<i>SOX13</i> (NP_005677.2)	<i>sox10f</i> (NP_726612.1)	<i>egl-13</i> (NP_001024918.1)	D
<i>SOX5</i> (NP_008871.3)	–	–	–
<i>SOX6</i> (NP_001139291.2)	–	–	–
<i>SOX9</i> (NP_000337.1)	<i>sox110b</i> (NP_651839.1)	–	E
<i>SOX8</i> (NP_055402.2)	–	–	–
<i>SOX10</i> (NP_008872.1)	–	–	–
<i>SOX18</i> (NP_060889.1)	<i>sox15</i> (NP_523739.2)	–	F
<i>SOX7</i> (NP_113627.1)	–	–	–
<i>SOX17</i> (NP_071899.1)	–	–	–
<i>SOX15</i> (NP_008873.1)	–	–	G
<i>SOX30</i> (NP_848511.1)	–	–	H
Fox gene family			
<i>FOXA1/HNF-3α</i> (NP_004487.2)	<i>forkhead/fkh</i> (NP_524542.1)	<i>pha-4/Ce-fkh1</i> (NP_001041114.1)	A
<i>FOXA2/HNF-3β</i> (NP_068556.2)	–	–	–
<i>FOXA3/HNF-3γ</i> (NP_004488.2)	–	–	–
<i>FOXB1</i> (NP_036314.2)	<i>fd96Ca/fd4</i> (NP_524495.1)	<i>lin-31</i> (NP_494704.1)	B
<i>FOXB2</i> (NP_001013757.1)	<i>fd96Cb/fd5</i> (NP_524496.1)	–	–
<i>FOXC1/MF1/FKHL7</i> (NP_001444.2)	<i>crocodile/fd1</i> (NP_524202.1)	–	C
<i>FOXC2/MFH1</i> (NP_005242.1)	–	–	–
<i>FOXD1/FREAC4</i> (NP_004463.1)	<i>fd59A/fd3</i> (NP_523814.1)	<i>unc-130</i> (NP_496411.1)	D
<i>FOXD2/FREAC9</i> (NP_004465.3)	–	–	–
<i>FOXD3</i> (NP_036315.1)	–	–	–
<i>FOXD4</i> (NP_997188.2)	–	–	–
<i>FOXE1/TITF2</i> (NP_004464.2)	–	–	E
<i>FOXE3</i> (NP_036318.1)	–	–	–
<i>FOXF1</i> (NP_001442.2)	<i>binious/FoxF</i> (NP_523950.2)	<i>let-381/F26B1.7</i> (NP_491826.1)	F
<i>FOXF2</i> (NP_001443.1)	–	–	–
<i>FOXG1/BF1/HBF2</i> (NP_005240.3)	<i>slp1</i> (NP_476730.1) <i>slp2</i> (NP_476834.1) <i>fd19B/cg9571</i> (NP_608369.1)	<i>fkh2/T14G12.4</i> (NP_508644.1)	G
<i>FOXH1/FAST1</i> (NP_003914.1)	–	–	H
<i>FOXI1/FREAC6/HFH3</i> (NP_036320.2)	–	–	I
<i>FOXJ1</i> (NP_001445.2)	–	–	J1
<i>FOXJ2</i> (XP_011519063.1)	–	–	J2
<i>FOXJ3</i> (XP_005270689.1)	–	–	J3
<i>FOXKL1/LF1</i> (NP_001032242.1)	<i>foxK/LD16137</i> (NP_001261701.1)	–	K
<i>FOXK2</i> (NP_004505.2)	–	–	–
<i>FOXL1</i> (NP_005241.1)	<i>foxL1/fd2</i> (NP_523912.1)	–	L1

Tab. S3 continued from previous page

<i>Homo sapiens</i>	<i>Drosophila melanogaster</i>	<i>Caenorhabditis elegans</i>	Group
Fox gene family			
<i>FOXL2</i> (NP_075555.1)	–	–	L2
<i>FOXM1</i> (NP_001400854.1)	–	–	M
<i>FOXN1/WHN</i> (NP_001356298.1)	<i>jumeau</i> (NP_524302.1)	–	N1/4
<i>FOXN4</i> (NP_998761.2)			
<i>FOXN2/HTLF</i> (NP_001362376.1)	<i>ches-1</i> (NP_511071.3)	–	N2/3
<i>FOXN3/CHES1</i> (NP_001078940.1)			
<i>FOXO1</i> (NP_002006.2)	–	<i>daf-16</i> (NP_001364785.1)	O
<i>FOXO3</i> (NP_963853.1)			
<i>FOXO3B</i> (NP_001355064.1)			
<i>FOXP1</i> (NP_001231739.1)			
<i>FOXP2</i> (NP_683696.2)			
<i>FOXP3</i> (NP_054728.2)	<i>foxP/cg16899</i> (NP_001247011.1)	<i>F26D12.1</i> (NP_001293813.1)	P
<i>FOXP4</i> (XP_011512591.1)			
<i>FOXQ/HFH11</i> (NP_150285.3)	–	–	Q1
–	<i>fd102C/cd11152</i> (NP_651951.1)	<i>fkh-10/C25A1.2</i> (NP_492676.2)	Q2
<i>FOXSI/FREAC10</i> (NP_004109.1)	–	–	S
–	–	<i>PES-1</i> (NP_001023406.1)	–
–	–	<i>B0286.5/FKH-6</i> (NP_494775.1)	–
–	–	<i>F40H3.4/FKH-8</i> (NP_001254107.1)	–
–	–	<i>C29F7.4/FKH-3</i> (NP_001294822.1)	–
–	–	<i>K03C7.2/FKH-9</i> (NP_001024760.1)	–

Supplementary Table S4 – Genomic data of mammals used to retrieve DSFGs and compute amino acid sequence divergence (AASD) of single-copy orthogroups (SCOs). For each species, the relative ID, taxonomic information, BUSCO statistics, NCBI accession number, and source publication are reported.

Species	ID	Class	Group	Order	Type	BUSCO statistics ('mammalia_odb10')	NCBI acc. no.	Reference
<i>Gallus gallus</i>	Ggal	Aves	Neognathae	Galliformes	Genome	C.99.0%[S:98.6%:D:0.4%][F:0.2%:M:0.8%	GCF_016699485.2	Vertebrate Genome Project
<i>Chryschloris asiatica</i>	Casi	Mammalia	Afrotheria	Afroscorida	Genome	C.98.0%[S:97.4%:D:0.6%][F:1.1%:M:0.9%	GCF_000296735.1	Murata et al., 2003
<i>Elephas maximus indicus</i>	Emax	Mammalia	Afrotheria	Proboscidea	Genome	C.98.0%[S:98.3%:D:0.6%][F:0.4%:M:0.7%	GCF_024166365.1	Vertebrate Genome Project
<i>Trichechus manatus latirostris</i>	Tman	Mammalia	Afrotheria	Sirenia	Genome	C.96.1%[S:95.7%:D:0.4%][F:1.8%:M:2.1%	GCF_000243295.1	Foote et al., 2015
<i>Orycteropus afer afer</i>	Oafe	Mammalia	Euarchontoglires	Tubulidentata	Genome	C.96.5%[S:96.0%:D:0.5%][F:1.9%:M:1.6%	GCF_000298275.1	–
<i>Ochetona princeps</i>	Opri	Mammalia	Euarchontoglires	Lagomorpha	Genome	C.98.3%[S:96.4%:D:1.9%][F:0.5%:M:1.2%	GCF_030433755.1	Vertebrate Genome Project
<i>Cebus imitator</i>	Cimi	Mammalia	Euarchontoglires	Primates	Genome	C.97.3%[S:95.1%:D:2.2%][F:1.7%:M:1.0%	GCF_001604975.1	Orkin et al., 2021
<i>Homo sapiens</i>	Hsap	Mammalia	Euarchontoglires	Primates	Genome	C.99.6%[S:97.3%:D:2.3%][F:0.2%:M:0.2%	GCF_000001405.40	Genome Reference Consortium
<i>Lemur catta</i>	Lcat	Mammalia	Euarchontoglires	Primates	Genome	C.98.3%[S:97.2%:D:1.1%][F:0.4%:M:1.3%	GCF_020740605.2	Vertebrate Genome Project
<i>Cavia porcellus</i>	Cpor	Mammalia	Euarchontoglires	Rodentia	Genome	C.96.4%[S:95.7%:D:0.7%][F:1.7%:M:1.9%	GCF_0001151735.1	The Genome Sequencing Platform
<i>Mus musculus</i>	Mmus	Mammalia	Euarchontoglires	Rodentia	Genome	C.99.4%[S:98.7%:D:0.7%][F:0.2%:M:0.4%	GCF_000001635.27	Genome Reference Consortium
<i>Sciurus carolinensis</i>	Scar	Mammalia	Euarchontoglires	Rodentia	Genome	C.99.1%[S:96.9%:D:2.2%][F:0.3%:M:0.6%	GCF_902686445.1	Mead et al., 2020
<i>Bubalus bubalis</i>	Bbub	Mammalia	Laurasatheria	Artiodactyla	Genome	C.98.7%[S:97.0%:D:1.7%][F:0.6%:M:0.7%	GCF_019923935.1	Deng et al., 2016
<i>Balaenoptera musculus</i>	Bmus	Mammalia	Laurasatheria	Artiodactyla	Genome	C.98.4%[S:95.7%:D:2.7%][F:0.6%:M:1.0%	GCF_00987245.2	Genome 10K
<i>Camelops dromedarius</i>	Cdro	Mammalia	Laurasatheria	Artiodactyla	Genome	C.99.7%[S:98.3%:D:0.4%][F:0.7%:M:0.6%	GCF_000083125.2	Elbers et al., 2019
<i>Hippopotamus amphibius kiboko</i>	Hamp	Mammalia	Laurasatheria	Artiodactyla	Genome	C.98.7%[S:95.2%:D:3.5%][F:0.5%:M:0.8%	GCF_030208045.1	Vertebrate Genome Project
<i>Phacochoerus africanus</i>	Paf	Mammalia	Laurasatheria	Artiodactyla	Genome	C.98.8%[S:98.3%:D:0.5%][F:0.6%:M:0.6%	GCF_016906955.1	–
<i>Tursiops truncatus</i>	Trtu	Mammalia	Laurasatheria	Artiodactyla	Genome	C.97.3%[S:95.2%:D:2.1%][F:1.1%:M:1.6%	GCF_011762595.1	Xiong et al., 2009
<i>Alliropoda melanoleuca</i>	Amel	Mammalia	Laurasatheria	Carnivora	Genome	C.97.3%[S:96.6%:D:0.7%][F:1.3%:M:1.4%	GCF_000207445.2	Fan et al., 2019
<i>Canis lupus familiaris</i>	Clup	Mammalia	Laurasatheria	Carnivora	Genome	C.98.5%[S:96.7%:D:1.8%][F:0.6%:M:0.9%	GCF_011100685.1	Wang et al., 2021
<i>Mirounga angustirostris</i>	Nang	Mammalia	Laurasatheria	Carnivora	Genome	C.96.7%[S:94.5%:D:2.2%][F:1.9%:M:1.4%	GCF_021288785.2	Moreno et al., 2024
<i>Panthera tigris</i>	Ptig	Mammalia	Laurasatheria	Carnivora	Genome	C.99.4%[S:98.9%:D:0.5%][F:0.3%:M:0.3%	GCF_018350195.1	Bredemeyer et al., 2023
<i>Desmodus rotundus</i>	Drot	Mammalia	Laurasatheria	Chiroptera	Genome	C.98.2%[S:97.2%:D:1.0%][F:0.5%:M:1.3%	GCF_0022682495.1	Bat 1K
<i>Pteropus giganteus</i>	Ppig	Mammalia	Laurasatheria	Chiroptera	Genome	C.97.2%[S:96.9%:D:0.3%][F:1.1%:M:1.7%	GCF_90272225.1	Fouret et al., 2020
<i>Rhinolophus ferrumequinum</i>	Rfer	Mammalia	Laurasatheria	Chiroptera	Genome	C.99.2%[S:97.9%:D:1.3%][F:0.3%:M:0.5%	GCF_004115265.2	Vertebrate Genome Project
<i>Ceratotherium simum simum</i>	Csim	Mammalia	Laurasatheria	Perissodactyla	Genome	C.98.8%[S:98.6%:D:0.2%][F:0.9%:M:0.3%	GCF_000283155.1	–
<i>Equus quagga</i>	Equa	Mammalia	Laurasatheria	Perissodactyla	Genome	C.98.5%[S:95.0%:D:3.5%][F:0.5%:M:1.0%	GCF_02161505.1	Vilstrup et al., 2013
<i>Manis javanica</i>	Mjav	Mammalia	Laurasatheria	Pholidota	Genome	C.95.7%[S:93.7%:D:2.0%][F:1.9%:M:2.4%	GCF_01457053.1	–
<i>Sarcophilus harrisi</i>	Shar	Mammalia	Metatheria	Dasyuromorphia	Genome	C.95.5%[S:94.5%:D:1.0%][F:0.9%:M:3.6%	GCF_90263505.1	Stammnitz et al., 2023
<i>Monodelphis domestica</i>	Mdom	Mammalia	Metatheria	Didelphimorphia	Genome	C.95.1%[S:92.3%:D:2.8%][F:0.9%:M:4.0%	GCF_027887165.1	Vertebrate Genome Project
<i>Ornithorhynchus anatinus</i>	Oana	Mammalia	Prototheria	Monotremata	Genome	C.92.3%[S:91.2%:D:1.1%][F:1.4%:M:6.3%	GCF_004115215.2	Zhou et al., 2021
<i>Dasyurus noveminctus</i>	Dnov	Mammalia	Xenarthra	Cingulata	Genome	C.96.9%[S:94.3%:D:2.6%][F:0.7%:M:2.4%	GCF_030445035.1	Vertebrate Genome Project
<i>Choloepus didactylus</i>	Cdid	Mammalia	Xenarthra	Pilosa	Genome	C.97.8%[S:91.9%:D:5.9%][F:0.7%:M:1.5%	GCF_015220235.1	Vertebrate Genome Project

Supplementary Table S5 – Genomic data of *Drosophila* used to retrieve DSFGs and compute AASD of SCOs. For each species, the relative ID, taxonomic information, BUSCO statistics, NCBI accession number, and source publication are reported.

Species	ID	Family	Subgenus	Type	BUSCO statistics ('diptera_odb10')	NCBI acc. no.	Reference
<i>Anopheles gambiae</i>	Agam	Culicidae	Cellia	Genome	C:99.4%[S:99.1%,D:0.3%],F:0.1%,M:0.5%	GCF_943734735.2	Habtewold et al., 2023
<i>Drosophila sechellia</i>	Dsec	Drosophilidae	Sophophora	Genome	C:99.9%[S:99.3%,D:0.6%],F:0.0%,M:0.1%	GCF_004382195.2	Chakraborty et al., 2021
<i>Drosophila melanogaster</i>	Dmel	Drosophilidae	Sophophora	Genome	C:100.0%[S:99.7%,D:0.3%],F:0.0%,M:0.0%	GCF_000001215.4	Hoskins et al., 2015
<i>Drosophila erecta</i>	Dere	Drosophilidae	Sophophora	Genome	C:99.9%[S:99.5%,D:0.4%],F:0.0%,M:0.1%	GCF_003286155.1	Dong et al., 2022
<i>Drosophila suzukii</i>	Dsuz	Drosophilidae	Sophophora	Genome	C:99.7%[S:96.5%,D:3.2%],F:0.1%,M:0.2%	GCF_013340165.1	Paris et al., 2020
<i>Drosophila elegans</i>	Dele	Drosophilidae	Sophophora	Genome	C:99.8%[S:99.5%,D:0.3%],F:0.1%,M:0.1%	GCF_018152505.1	Kim et al., 2021
<i>Drosophila serrata</i>	Dser	Drosophilidae	Sophophora	Genome	C:99.9%[S:97.5%,D:2.4%],F:0.0%,M:0.1%	GCF_002093755.2	Allen et al., 2017
<i>Drosophila kikkawai</i>	Dkik	Drosophilidae	Sophophora	Genome	C:100.0%[S:99.1%,D:0.9%],F:0.0%,M:0.0%	GCF_018152535.1	Kim et al., 2021
<i>Drosophila bipunctata</i>	Dbip	Drosophilidae	Sophophora	Genome	C:99.9%[S:99.2%,D:0.7%],F:0.0%,M:0.1%	GCF_018153845.1	Kim et al., 2021
<i>Drosophila ananassae</i>	Dana	Drosophilidae	Sophophora	Genome	C:99.6%[S:99.3%,D:0.3%],F:0.0%,M:0.4%	GCF_017639315.1	Tvedte et al., 2021
<i>Drosophila pseudoobscura</i>	Dpse	Drosophilidae	Sophophora	Genome	C:99.7%[S:98.8%,D:0.9%],F:0.1%,M:0.2%	GCF_009870125.1	Liao et al., 2021
<i>Drosophila miranda</i>	Dmir	Drosophilidae	Sophophora	Genome	C:99.8%[S:85.6%,D:14.2%],F:0.1%,M:0.1%	GCF_003369915.1	Mahajan et al., 2018
<i>Drosophila willistoni</i>	Dwil	Drosophilidae	Sophophora	Genome	C:99.6%[S:98.4%,D:1.2%],F:0.0%,M:0.4%	GCF_018902025.1	Ranz et al., 2023
<i>Drosophila arizonae</i>	Dari	Drosophilidae	Drosophila	Genome	C:95.7%[S:95.3%,D:0.4%],F:1.2%,M:3.1%	GCF_001654025.1	Sanchez-Flores et al., 2016
<i>Drosophila hydei</i>	Dhyd	Drosophilidae	Drosophila	Genome	C:99.7%[S:97.5%,D:2.2%],F:0.1%,M:0.2%	GCF_003285905.1	Dong et al., 2022
<i>Drosophila grimshawi</i>	Dgri	Drosophilidae	Drosophila	Genome	C:99.9%[S:99.2%,D:0.7%],F:0.0%,M:0.1%	GCF_018153295.1	Kim et al., 2021
<i>Drosophila albomicans</i>	Dalb	Drosophilidae	Drosophila	Genome	C:99.9%[S:99.1%,D:0.8%],F:0.0%,M:0.1%	GCF_009650485.2	Mai et al., 2020
<i>Drosophila busckii</i>	Dbus	Drosophilidae	Drosophila	Genome	C:98.1%[S:97.4%,D:0.7%],F:0.3%,M:1.6%	GCF_011750605.1	Renschler et al., 2019

Supplementary Table S6 – Complete set of DSFGs in bivalves. For each gene, the species ID (Sp. ID) as in Tab. S1, the accession number (Gene ID), the Possvm-based annotation, and the CDD domains (including their Psm-ID) are indicated.

Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Airc	Contig6_279	Dmrt	Dmrt-OGO/NA	Doublesex DNA-binding motif (214606)	-	Annotated as Dmrt-1L
	scaffold_235_403	Dmrt	Dmrt-OGO/NA	Doublesex DNA-binding motif (214606)	-	Annotated as Dmrt-1L
Apur	XP_052698016.1	Dmrt	Dmrt-OGO/NA	Doublesex DNA-binding motif (214606)	-	Annotated as Dmrt-1L
Cang	EVMO0027346.1	Dmrt	Dmrt-OGO/NA	Doublesex DNA-binding motif (214606)	-	Annotated as Dmrt-1L
Carl	XP_011441049.2	Dmrt	Dmrt-OGO/NA	Doublesex DNA-binding motif (214606)	-	Annotated as Dmrt-1L
Cgig	XP_022333988.1	Dmrt	Dmrt-OGO/NA	Doublesex DNA-binding motif (214606)	-	Annotated as Dmrt-1L
Cvir	XP_041358115.1	Dmrt	Dmrt-OGO/NA	Doublesex DNA-binding motif (214606)	-	Annotated as Dmrt-1L
Gaeq	M00000013645	Dmrt	Dmrt-OGO/NA	Doublesex DNA-binding motif (214606)	-	Annotated as Dmrt-1L
Hbia	M00000045261	Dmrt	Dmrt-OGO/NA	Doublesex DNA-binding motif (214606)	-	Annotated as Dmrt-1L
Hbifia	XP_046372338.2	Dmrt	Dmrt-OGO/NA	Doublesex DNA-binding motif (214606)	-	Annotated as Dmrt-1L
Hruf	XP_0463535704.1	Dmrt	Dmrt-OGO/NA	Doublesex DNA-binding motif (214606)	-	Annotated as Dmrt-1L
Hruf	XP_052068518.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Mcal	CAC3397186.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Mcor	CAG2232556.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Medu	VD103798.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606; partial)
Mgar	MMAN00000008302	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Mmar	g120437_t1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Mner	Obim	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Oedu	XP_048736174.2	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Osin	XP_046366891.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Osin	XP_0463666446.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Osin	XP_0396477012.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Pcan	XP_025090051.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Pcan	XP_025111744.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Pmar	DN306577_c0_g1.i.p1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Pmar	XP_037336565.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Pmar	PPur	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Pstr	KAK3994948.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606; partial)
Pvir	s01850g168	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Pyes	XP_021353714.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Sbro	EVM0020695.1	Dmrt	Dmrt-OG4/NA	Doublesex DNA-binding motif (214606)	-	Doublesex DNA-binding motif (214606)
Sgio	Sgi011295	Dmrt	Dmrt-2	Dmrt-2	-	-
Arc	Contig172_94	Dmrt	Dmrt-2	Dmrt-2	-	-
Apcc	DN1321_c0_g1.i.p1	Dmrt	Dmrt-2	Dmrt-2	-	-
Bgia	XP_055887190.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Cphi	DN116434_c0_g1.i.p1	Dmrt	Dmrt-2	Dmrt-2	-	-
Gaeq	XP_052832484.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Hruf	XP_048255484.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Mcor	CAC504148.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Medu	CAG2252366.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Mgar	VD142071.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Mmar	XP_045156965.2	Dmrt	Dmrt-2	Dmrt-2	-	-
Mphi	Scaf.304771.9	Dmrt	Dmrt-2	Dmrt-2	-	-
Obim	XP_052832484.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Osin	XP_029650766.2	Dmrt	Dmrt-2	Dmrt-2	-	-
Pcan	DN1165632_c0_g1.i.p1	Dmrt	Dmrt-2	Dmrt-2	-	-
Pcor	DN151233_c0_g1.i.p1	Dmrt	Dmrt-2	Dmrt-2	-	-
Pcor	DN280333_c0_g1.i.p1	Dmrt	Dmrt-2	Dmrt-2	-	-
Pmax	XP_033738864.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Pstr	KAK3603675.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Pvir	s0097g235	Dmrt	Dmrt-2	Dmrt-2	-	-
Pyes	XP_021368788.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Rphi	XP_060589226.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Sbro	EVM0001645.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Scon	Ch8_1365	Dmrt	Dmrt-2	Dmrt-2	-	-
Tgra	KAJ3306274.1	Dmrt	Dmrt-2	Dmrt-2	-	-
Tsqu	DN51730_c0_g1.i.p1	Dmrt	Dmrt-2	Dmrt-2	-	-
Acal	XP_030509632.1	Dmrt	Dmrt-3	Dmrt-3	-	-
Acal	Contig349_40	Dmrt	Dmrt-3	Dmrt-3	-	-
Amar	Ama12564	Dmrt	Dmrt-3	Dmrt-3	-	-
Apur	scaf.95_76	Dmrt	Dmrt-3	Dmrt-3	-	-
BglA	XP_013077145.1	Dmrt	Dmrt-3	Dmrt-3	-	-
Cang	XP_052687934.1	Dmrt	Dmrt-3	Dmrt-3	-	-

Tab. S6 continued from previous page

Sp ID	Gene ID	Group	Annotation	Main catalytic domain (P _{psm-ID})	Additional domains (P _{psm-ID})	Notes
Cari	EVM0028466.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Ceglg	XP_011427033.2	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Cphl	DN3729_c0_g1_i1_p1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	-	
Csin	Hic-asn..11.i1r4	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Cvir	XP_022317913.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Dpol	XP_052246678.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Gaege	XP_041357684.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Hbia	M0000034631	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Hruf	XP_046367747.2	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mare	XP_052277877.2	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mcal	XP_052076466.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mcor	CAC5360634.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Medu	CAG2226664.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mgal	VD32052.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mmar	MMAN00000039146	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mmer	XP_045157038.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Miner	g243_052_t1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mphi	seaf_42165_0.7	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Oedu	XP_048761857.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Pcan	PN_05110327.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Pcor	DN33000_c0_g1_i1_p1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Pmax	XP_033737545.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Pvir	s00097g272	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Rph1	XP_060576862.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Sbro	EVMM006488.1	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Scro	Ch8_2435	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Seflo	Sej014397	Dmrt	Dmrt-3	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Tgra	KAJ8305800.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Acal	XP_005096931.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Airc	Contig349_42	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Amar	Ama12441	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Amar	Ana12902	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Apur	DN10969_c0_g1_i1_p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Apur	scaffoff_95.777	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Bela	XP_013089804.2	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Cang	NP_001295334.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Cari	EVM0008823.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Ceglg	DN67594_c0_g1_i1_p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Cphi	DN159544_c0_g1_i1_p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Cphi	DN54583_c0_g1_i1_p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Cphi	DN9274_c0_g1_i1_p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Cphi	DN67594_c0_g2_i1_p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Cphi	Hic-asn..11.400	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Cphi	Csin..11.338	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Cvir	XP_022319926.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Dpol	XP_052255315.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Dpol	XP_052224782.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Dpol	XP_052257495.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Gaege	XP_041357789.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Hbia	M0000038945	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Hbia	M0000038946	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Hbia	M0000019813	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Hbia	M0000038947	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Hruf	DN147341_c0_g1_i1_p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Lorb	DN40542_c0_g1_i1_p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Lorb	XP_052776891.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mare	XP_052776490.6	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mare	XP_052776885.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mcal	XP_0520922340.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mchi	DN35166_c0_g1_i1_p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mcor	CAC5398878.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Medu	CAG2209978.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mgal	VD124477.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mmar	MMAM000000040448	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270600)	
Mmar	MMAM000000003815	Mmar	-	-	-	

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (P _{ssm-ID})	Additional domains (P _{ssm-ID})	Notes
Mmar	MMAN00000016566	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600, 270602)	
Mmar	MMAN00000047004	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600, 270602)	
Mmar	MMAN00000044361	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600, 270602)	
Mmer	XP_045157593_1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Mmer	XP_045157053_1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Mmer	DN16828_c0.g1.i1.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Mmed	g346_03.tl	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Mmer	g585_31.tl	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602)	
Mmer	g241_174.tl	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Mmer	g192_820.tl	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Mphi	scfa_68796_0.3	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Oedu	XP_048763391_1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270465)	
Pcan	PCan_025110328.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Pcor	DN12587_c0.g13.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602)	
Pcor	DN1623_c0.g1.i13.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600, 270602)	
Pcor	DN15587_c0.g1.i4.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602)	
Pgen	DN68344_c0.g1.i1.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Pgen	XP_033737544_1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270465)	
Pmax	DN35556_c0.g1.i2.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Poku	DN35178_c0.g1.i2.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Pstr	KAK3612677.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Pstr	KAK388310.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602, 270465)	
Rph1	XP_06000638.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Rph1	XP_060878674.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Sbro	EVM0004355.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602)	
Sbro	EVM0021940.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270602)	
Scon	Ch8-1999	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600, 270465)	
Scon	Ch8-2143	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Sglo	Sg106992	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600, 270465)	
Sgra	DN54078_c0.g1.i1.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Sgra	DN66569_c0.g1.i3.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600, 270465)	
Tgra	KAJ3305789.1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600)	
Tsqu	DN75749_c0.g1.i1.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600, 270465)	
Apec	DN8372_c0.g4.i1.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600, 270465)	
Mcchi	DN34711_c0.g1.i2.p1	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600, 270465)	
Scon	Ch8-738	Dmrt	Dmrt-4/5	Doublesex DNA-binding motif (214606)	CUE-like DMA domain (270553, 270601, 270600, 270465)	
Acal	XP_05397243.2	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Airc	Contig_157	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Amar	Ama08751	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Apcc	DN107972_c0.g1.i1.p1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Apur	scaffold_124.7	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Brg1	XP_013067134.2	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552, partial)	
Cang	XP_052701295.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Cari	EVM0004613.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Cflu	DN101169_c0.g1.i1.p1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Cgig	XP_011413445.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Cpli	DN47094_c0.g4.i1.p1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Csin	Hic_asn_10.437	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Cvir	XP_02233332.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Cvir	XP_022334050.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Dpol	XP_052272379.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552, partial)	
Gaeq	XP_041352454.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
M0000018167	Fox	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Hbia	XP_04631021.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Hraf	XP_052769228.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Mcal	XP_052106467.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552, partial)	
Mchi	DN35555_c0.g4.i1.p1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552, partial)	
Mcor	CAC5370406.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Medu	CAG2201348.1	Fox-A	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Pssm-ID)	Additional domains (Pssm-ID)	Notes
Mgal	VD11457.1	Fox	Fox-A	Forkhead domain A1 (410812)	-	
Mmar	MMAM00000008663	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552; partial)	
Mmer	XP_045173733.1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552; partial)	
Mmod	DN103780_c0.g1.i1.p1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872; partial); HNF3 C-terminal domain (430552)	
Mner	g192217.t1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Mphi	scaf4682.0	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal A1 (410812)	
Obim	XP_014788201.2	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872)	
Oedu	XP_048735259.1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552; partial)	
Pcan	XP_025090786.1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872; partial); HNF3 C-terminal domain (430552)	
Pcor	DN3402_c0.g1.i1.p1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872; partial)	
Pgen	DN17637_c0.g1.i1.p1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872; partial)	
Pmax	DN30866_c0.g1.i1.p1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Pstr	KAK3597847.1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552; partial)	
Pvir	XP_00068447	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Pyes	XP_021361791.1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Rphi	XP_060590754.1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Sbro	EVMM0003194.1	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Scor	Ch4_2670	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872); HNF3 C-terminal domain (430552)	
Sglo	Sglo00464	Fox	Fox-A	Forkhead domain A1 (410812)	Forkhead N-terminal (369872; partial); HNF3 C-terminal domain (430552)	
Sgra	DN78052_c0.g1.i1.p1	Fox	Fox-B	Forkhead domain A1 (410812)	Forkhead N-terminal (369872; partial)	
Acal	XP_005089018.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Airc	Contig636_38	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Ajrc	scaffold313.50	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Bglia	XP_013078204.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Cang	XP_032702033.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Cari	EVMM0003536.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Cflu	DN98613_c0.g1.i1.p1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Cgig	XP_011445364.2	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Csin	Hic_asn_16_1347	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Cvir	XP_022334612.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Dpol	XP_052233250.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Dpol	XP_052256324.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Dpol	XP_052281977.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Gaeq	XP_041361159.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Hbia	M00000029836	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Hrf	XP_0463585890.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Lorb	DN5589_c3.g1.i1.p1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Mare	XP_052791461.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Mcal	XP_052100219.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Mcor	CAC5382565.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Medu	CAG2229716.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Mgal	VD16970.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Mmar	MMAM00000015629	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Mmer	XP_042155305.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Mner	g250725.t1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Mphi	scaf10920.0.0	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Obim	XP_052832321.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Oedu	XP_048732871.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Osin	XP_029653697.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Pean	XP_025078261.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Pcor	DN23979_c0.g1.i1.p1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Pmax	DN23979_c1.g1.i1.p1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Pstr	KAK3607900.1	Fox	Fox-C	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Pvir	s001898g215	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Pyes	XP_021357620.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Rphi	XP_060564412.1	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Scon	Ch5_12	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Sglo	Sglo12012	Fox	Fox-B	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Tgra	KAJ3304921.1	Fox	Fox-C	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Acal	XP_005106277.1	Fox	Fox-C	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Airc	Config58.63	Fox	Fox-C	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Amar	Amar1094	Fox	Fox-C	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Apur	scaffold57.50	Fox	Fox-C	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Bglia	XP_055890240.1	Fox	Fox-C	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Cang	XP_052715579.1	Fox	Fox-C	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Cari	EVMM0002771.1	Fox	Fox-C	Forkhead domain B2 (410817)	Forkhead N-terminal (369872; partial)	
Cflu	DN95576_c0.g1.i1.p1	Fox	Fox-C	Forkhead domain C (410793)	Forkhead N-terminal (369872; partial)	

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Pssm-ID)	Additional domains (Pssm-ID)	Notes
Cggg	XP_011417585_2	Fox	Fox-C	Forkhead domain L1 (410801)		
Cpli	DN157725_c0_g1_i1.p1	Fox	Fox-C	Forkhead domain L1 (410801)		
Csin	Hic.asrn_17.1387	Fox	Fox-C	Forkhead domain L1 (410801)		
Csin	Hic.asrn_17.1443	Fox	Fox-C	Forkhead domain L1 (410801)		
Cvir	XP_022346235_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Dpol	XP_052247387_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Gaeg	XP_041377087_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Hbia	M0000031058	Fox	Fox-C	Forkhead domain L1 (410801)		
Hruf	XP_046372770_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Mare	XP_02819073_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Mcal	XP_052063314_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Mchi	DN13809_c0_g1_i1.p1	Fox	Fox-C	Forkhead domain L1 (410801)		
Mcor	CAC537404_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Medu	CAG220684_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Mgal	VD122482_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Mmar	MMAN0000035516	Fox	Fox-C	Forkhead domain L1 (410801)		
Mmer	XP_045194706_2	Fox	Fox-C	Forkhead domain L1 (410801)		
Mmer	g82158_t1	Fox	Fox-C	Forkhead domain L1 (410801)		
Mphi	scf.69501.0	Fox	Fox-C	Forkhead domain L1 (410801)		
Obim	XP_014786040_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Oedu	XP_048762038_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Osin	XP_029653806_2	Fox	Fox-C	Forkhead domain L1 (410801)		
Pcan	XP_025115697_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Pcor	DN14158_c0_g2_i1.p1	Fox	Fox-C	Forkhead domain L1 (410801)		
Pcor	DN14158_c0_g5_i1.p1	Fox	Fox-C	Forkhead domain L1 (410801)		
Pmax	XP_033755061_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Pstr	KAK390993_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Pvir	s020238_12	Fox	Fox-C	Forkhead domain L1 (410801)		
Pyes	XP_021346967_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Rphi	XP_06059704_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Sbro	EVM0022192_1	Fox	Fox-C	Forkhead domain L1 (410801)		
Scor	Chr1-1448	Fox	Fox-C	Forkhead domain L1 (410801)		
Sglo	Sg1009485	Fox	Fox-C	Forkhead domain L1 (410801)		
Tgra	KAJ3303551_1	Fox	Fox-C	Forkhead domain C2 (410819)		
Acal	XP_035824261_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Airc	Contig1003_15	Fox	Fox-D	Forkhead domain D4 (410822)		
Amar	Ama11686	Fox	Fox-D	Forkhead domain D3 (410821)		
Apc	DN87882_c0_g1_i1.p1	Fox	Fox-D	Forkhead domain D4 (410822)		
Apur	scf10d13962_11	Fox	Fox-D	Forkhead domain D4 (410822)		
Bglia	XP_013096936_2	Fox	Fox-D	Forkhead domain D4 (410822)		
Cang	XP_052688370_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Cari	EVMM0005770_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Ceig	XP_011446328_2	Fox	Fox-D	Forkhead domain D4 (410822)		
Cphl	DN233774_c0_g1_i1.p1	Fox	Fox-D	Forkhead domain D4 (410822)		
Csin	XP_046329290_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Csin	Hic.asrn_11.1425	Fox	Fox-D	Forkhead domain D4 (410822)		
Cvir	XP_022316146_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Dpol	XP_052256035_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Dpol	XP_052256891_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Gaeg	XP_041356731_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Hbia	M0000030583	Fox	Fox-D	Forkhead domain D4 (410822)		
Hruf	XP_042075808_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Lorb	CAC5382691_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Mare	XP_052777467_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Mcal	XP_052777725_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Mcal	XP_052095202_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Mgal	VD107735_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Mgal	VDH9306_1	Fox	Fox-D	Forkhead domain D3 (410821)		
Mcor	CAC5407497_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Medu	CAG220466_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Medu	CAG2248150_1	Fox	Fox-D	Forkhead domain D4 (410822)		
CAG2203862_1		Fox	Fox-D	Forkhead domain D4 (410822)		
Mgal	VDH9306_1	Fox	Fox-D	Forkhead domain D4 (410822)		
Mmar	MMAN0000002467	Fox	Fox-D	Forkhead domain D4 (410822)		
Mmer	XP_045157253_2	Fox	Fox-D	Forkhead domain D4 (410822)		
Mmer	g192986_t1	Fox	Fox-D	Forkhead domain D4 (410822)		
Mphi	scf.69491.12	Fox	Fox-D	Forkhead domain D4 (410822)		
Mphi	scf.42856_0.4	Fox	Fox-D	Forkhead domain D4 (410822)		
Obim	XP_052826256_1	Fox	Fox-D	Forkhead domain D4 (410822)		

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Pssm-ID)	Additional domains (Pssm-ID)	Notes
Oedu	XP_048762457.1	Fox	Fox-D	Forkhead domain D4 (410822)		
Pcan	XP_025110523.1	Fox	Fox-D	Forkhead domain D4 (410822)		
Pcor	DN15187_c0_g1.i5.p1	Fox	Fox-D	Forkhead domain D4 (410822)		
Pmar	DN31265_c0_g1.i1.p1	Fox	Fox-D	Forkhead domain D4 (410822)		
Pmax	XP_0337373945.1	Fox	Fox-D	Forkhead domain D4 (410822)		
Pstr	KAK3592139.1	Fox	Fox-D	Forkhead domain D4 (410822)		
Pvir	s01422g51	Fox	Fox-D	Forkhead domain D4 (410822)		
Pves	s0097g340	Fox	Fox-D	Forkhead domain D4 (410822)		
Rphi	XP_021345225.1	Fox	Fox-D	Forkhead domain D4 (410822)		
Sbro	XP_060538528.1	Fox	Fox-D	Forkhead domain D4 (410822)		
Scon	EVM0002351.1	Fox	Fox-D	Forkhead domain D4 (410822)		
Serlo	Chr8_2069	Fox	Fox-D	Forkhead domain D4 (410822)		
Tgra	Sg1013024	Fox	Fox-D	Forkhead domain D4 (410822)		
Airc	KAJ8306624.1	Fox	Fox-E	Forkhead domain E (410793)		
Contig989.19	Contig989.19	Fox	Fox-E	Forkhead domain E (410793)		
Amar	Am12850	Fox	Fox-E	Forkhead domain E (410793)		
Apur	scaffold_253.23	Fox	Fox-E	Forkhead domain E (410793)		
Cang	XP_0252888878.1	Fox	Fox-E	Forkhead domain E (410793)		
Cari	EVM0003339.1	Fox	Fox-E	Forkhead domain E (410793)		
Chlu	DN108936_c2_g1.i1.p1	Fox	Fox-E	Forkhead domain E (410793)		
Cejg	XP_011444776.2	Fox	Fox-E	Forkhead domain E (410793)		
Cvrr	XP_022319236.1	Fox	Fox-E	Forkhead domain E (410793)		
Dpol	XP_02286360.1	Fox	Fox-E	Forkhead domain E (410793)		
Hbia	M0000038943	Fox	Fox-E	Forkhead domain E (410793)		
Hruf	XP_046353578.2	Fox	Fox-E	Forkhead domain E (410793)		
Mare	XP_032778423.1	Fox	Fox-E	Forkhead domain E (410793)		
Mcal	XP_052075782.1	Fox	Fox-E	Forkhead domain E (410793)		
Mcor	CAC384360.1	Fox	Fox-E	Forkhead domain E (410793)		
Medu	CAG2217852.1	Fox	Fox-E	Forkhead domain E (410793)		
Medu	CAG2194171.1	Fox	Fox-E	Forkhead domain E (410793)		
Megl	CAG2199036.1	Fox	Fox-E	Forkhead domain E (410793)		
Mnar	VDH90460.1	Fox	Fox-E	Forkhead domain E (410793)		
Mmer	MMAAM0000033594	Fox	Fox-E	Forkhead domain E (410793)		
Mmod	DN1_045157592.2	Fox	Fox-E	Forkhead domain E (410793)		
Mner	g241_620_t1	Fox	Fox-E	Forkhead domain E (410793)		
Mphi	scaf:31587_0.4	Fox	Fox-E	Forkhead domain E (410793)		
Oedu	XP_048762291.1	Fox	Fox-E	Forkhead domain E (410793)		
Pmar	DN27017_c0_g1.i1.p1	Fox	Fox-E	Forkhead domain E (410793)		
Pmax	XP_045157592.1	Fox	Fox-E	Forkhead domain E (410793)		
Pstr	KAK3583103.1	Fox	Fox-E	Forkhead domain E (410793)		
Pvir	s00145g54	Fox	Fox-E	Forkhead domain E (410793)		
Pves	XP_021378858.1	Fox	Fox-F	Forkhead domain F (410794)		
Rdec	DN2595_c4_g1.i1.p1	Fox	Fox-F	Forkhead domain F (410794)		
Rphi	XP_060578867.1	Fox	Fox-F	Forkhead domain F (410794)		
Sbro	EVM0010028.1	Fox	Fox-F	Forkhead domain F (410794)		
Selo	Sg1009305	Fox	Fox-F	Forkhead domain F (410794)		
Acal	XP_005105969.2	Fox	Fox-F	Forkhead domain F (410794)		
Airc	Contig113.18	Fox	Fox-F	Forkhead domain F (410794)		
Amar	Ama39500	Fox	Fox-F	Forkhead domain F (410794)		
Amar	Ama32615	Fox	Fox-F	Forkhead domain F (410794)		
Apc	DN75342_c0_g1.i1.p1	Fox	Fox-F	Forkhead domain F (410794)		
Apur	scaffold_860_37	Fox	Fox-F	Forkhead domain F (410794)		
Bglj	XP_055892380.1	Fox	Fox-F	Forkhead domain F (410794)		
Cang	XP_052712246.1	Fox	Fox-F	Forkhead domain F (410794)		
Cari	EVM0011190.1	Fox	Fox-F	Forkhead domain F (410794)		
Cgig	XP_022335664.1	Fox	Fox-F	Forkhead domain F (410794)		
Cpli	DPN628_c0_g1.i1.p1	Fox	Fox-F	Forkhead domain F (410794)		
Csin	Hic_asrn_17_158	Fox	Fox-F	Forkhead domain F (410794)		
Cvrr	XP_022332755.1	Fox	Fox-F	Forkhead domain F (410794)		
Dpol	XP_052232755.1	Fox	Fox-F	Forkhead domain F (410794)		
Gaeq	XP_041375666.1	Fox	Fox-F	Forkhead domain F (410794)		
Hbia	M0000007664	Fox	Fox-F	Forkhead domain F (410794)		
Hruf	XP_046372649.1	Fox	Fox-F	Forkhead domain F (410794)		
Mcal	XP_052815084.1	Fox	Fox-F	Forkhead domain F (410794)		
Mcor	CAC587332.1	Fox	Fox-F	Forkhead domain F (410794)		
Medu	CAG2252875.1	Fox	Fox-F	Forkhead domain F (410794)		
Megl	VD121852.1	Fox	Fox-F	Forkhead domain F (410794)		

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Pssm-ID)	Additional domains (Pssm-ID)	Notes
Mmar	MMA/M0000030848	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Mmer	XP_045194642_1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Mmod	DN104261_c0.g1.i1.p1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Mmer	g106_129.t1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Mphi	scaf_40546_0.2	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Obim	XP_014777539_1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Oedu	XP_0487322202_1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Pcan	XP_025116010_1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F1 (410823)
Pcor	DN180603_c0.g1.i1.p1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Rphi	XP_060001663_1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Sbro	EVM0015186_1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Pmar	XP_03755005_1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Pmax	KAK3601654_1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Pstr	sl33835s10	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Pvir	XP_021558008_1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Pyes	DN129940_c0.g1.i1.p1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Tgra	KAJ8302829_1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Tsqu	DN137576_c0.g1.i1.p1	Fox	Fox-F	Fox	Fox-F	Forkhead domain F (410794)
Acal	XP_005099252_2	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Acar	Contig625_38	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Amar	Amal0381	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Apec	DN108336_c0.g1.i1.p1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Apuc	scaffold_36470_28	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Bglia	XP_055879295_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Cang	XP_052699015_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Cari	EVM0001891_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Cflu	DN104980_c0.g1.i2.p1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Cgig	XP_011427689_2	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Cipl	DN158419_c0.g1.i1.p1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Csin	Hic.asn_10.1034	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Cvir	XP_022334541_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Dpol	XP_052270224_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Dpol	XP_052270147_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Gaeq	XP_041354930_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Gaeq	XP_041354700_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Hbia	M00000035850	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Hruf	XP_046371537_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Mcal	XP_08259351_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Mcor	XP_062104484_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Medu	CAC5405696_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Mgal	VD124297_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Mmar	MMA/M0000030730	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Mmer	XP_045162348_2	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Mmod	DN60588_c0.g1.i1.p1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Mper	g13325_t1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Mphi	scaf_15017_0.3	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Obim	XP_02824454_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Oedu	XP_048737541_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Pcan	XP_025105677_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Pcan	XP_025106039_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Pmax	XP_025105724_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Poku	DN81635_c0.g2.i1.p1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Pcor	DN81635_c0.g1.i1.p1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Pvir	s03333s35	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Pyes	XP_0215637290_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Rphi	XP_060589805_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Sbro	EVM0011335_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)
Sbro	EVM0012606_1	Fox	Fox-G	Fox	Fox-G	Forkhead domain G (410795)

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Scon	Ch3_2805	Fox	Fox-G	Forkhead domain G (410795)		
Sgio	Sg1014601	Fox	Fox-G	Forkhead domain G (410795)		
Sgra	DN49488_c0_g1.i1.p1	Fox	Fox-G	Forkhead domain G (410795)		
Airc	Contig18244.2	Fox	Fox-H	Forkhead domain H (410796)		
Airc	Contig178.106	Fox	Fox-H	Forkhead domain H (410796)		
Amar	Ama16564	Fox	Fox-H	Forkhead domain H (410796)		
Amar	Ama05868	Fox	Fox-H	Forkhead domain H (410796)		
Cang	XP_052684756.1	Fox	Fox-H	Forkhead domain H (410796)		
Cari	EVM0021377.1	Fox	Fox-H	Forkhead domain H (410796)		
Cflu	DN138466_c0_g1.i1.p1	Fox	Fox-H	Forkhead domain H (410796)		
Cgig	XP_034313225.1	Fox	Fox-H	Forkhead domain H (410796)		
Hic.asn.14.811	Hic.asn.14.811	Fox	Fox-H	Forkhead domain H (410796)		
Cvir	XP_022314590.1	Fox	Fox-H	Forkhead domain H (410796)		
Hbia	M0000018729	Fox	Fox-H	Forkhead domain H (410796)		
Hruf	XP_048254913.1	Fox	Fox-H	Forkhead domain H (410796)		
Hruf	XP_048255113.1	Fox	Fox-H	Forkhead domain H (410796)		
Mcor	CAC409624.1	Fox	Fox-H	Forkhead domain H (410796)		
Mcor	CAC3397897.1	Fox	Fox-H	Forkhead domain H (410796)		
Mcor	CAC5397906.1	Fox	Fox-H	Forkhead domain H (410796)		
Mcor	CAC5403969.1	Fox	Fox-H	Forkhead domain H (410796)		
Medu	CAG2228903.1	Fox	Fox-H	Forkhead domain H (410796)		
Medu	CAG2188004.1	Fox	Fox-H	Forkhead domain H (410796)		
Medu	CAG2252853.1	Fox	Fox-H	Forkhead domain H (410796)		
Medu	CAG2220596.1	Fox	Fox-H	Forkhead domain H (410796)		
Mgal	VD162725.1	Fox	Fox-H	Forkhead domain H (410796)		
Mgal	VDH3947.1	Fox	Fox-H	Forkhead domain H (410796)		
Mgal	VD120844.1	Fox	Fox-H	Forkhead domain H (410796)		
Mmar	MMAN00000022684.	Fox	Fox-H	Forkhead domain H (410796)		
Mmer	XP_053378216.1	Fox	Fox-H	Forkhead domain H (410796)		
Mmer	XP_045194303.2	Fox	Fox-H	Forkhead domain H (410796)		
Mmer	XP_045198985.2	Fox	Fox-H	Forkhead domain H (410796)		
Mner	g213542.t1	Fox	Fox-H	Forkhead domain H (410796)		
Mphi	scf17325.0.4	Fox	Fox-H	Forkhead domain H (410796)		
Mphi	scf128666.1.1	Fox	Fox-H	Forkhead domain H (410796)		
Oedu	XP_048759429.2	Fox	Fox-H	Forkhead domain H (410796)		
Pcan	DPN16937_c0_g1.i1.p1	Fox	Fox-H	Forkhead domain H (410796)		
Pcor	XP_033755807.1	Fox	Fox-H	Forkhead domain H (410796)		
Pmax	KAK36038359.1	Fox	Fox-H	Forkhead domain H (410796)		
Pstr	s75596g33	Fox	Fox-H	Forkhead domain H (410796)		
Pvir	s00234g131	Fox	Fox-H	Forkhead domain H (410796)		
Rphi	XP_025075954.1	Fox	Fox-H	Forkhead domain H (410796)		
Rphi	XP_060567331.1	Fox	Fox-H	Forkhead domain H (410796)		
Rphi	XP_060604067.1	Fox	Fox-H	Forkhead domain H (410796)		
Sbro	EVMM0016618.1	Fox	Fox-H	Forkhead domain H (410796)		
Sbro	EVM0013817.1	Fox	Fox-H	Forkhead domain H (410796)		
Scon	Chr11.1359	Fox	Fox-H	Forkhead domain H (410796)		
Scon	Chr2.2082	Fox	Fox-H	Forkhead domain H (410796)		
Sgio	Sg1013003	Fox	Fox-H	Forkhead domain H (410796)		
Acal	XP_005108651.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Airc	Contig775.5	Fox	Fox-J1	Forkhead domain J1 (410797)		
Amar	Ama02822	Fox	Fox-J1	Forkhead domain J1 (410797)		
Apc	DN20109_c0_g1.i6.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Apur	scffold797.10	Fox	Fox-J1	Forkhead domain J1 (410797)		
Brla	XP_013064514.2	Fox	Fox-J1	Forkhead domain J1 (410797)		
Cang	XP_052689275.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Cari	EVM00033558.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Cflu	DN127407_c0_g2.i1.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Cgig	XP_011445341	Fox	Fox-J1	Forkhead domain J1 (410797)		
Cpli	DN65792_c0_g1.i1.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Csin	Hic.asn.0.1540	Fox	Fox-J1	Forkhead domain J1 (410797)		
Cvir	XP_023319181.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Dpol	XP_0223192968.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Geg	XP_041362703.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Hbia	M0000003225	Fox	Fox-J1	Forkhead domain J1 (410797)		
Hruf	XP_046330175.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Lorb	DN146717_c0_g1.i1.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mare	XP_052764656.1	Fox	Fox-J1	Forkhead domain J1 (410797)		

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Mare	XP_052816354.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mare	XP_052764667.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mare	XP_052775202.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mare	XP_052775217.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mare	XP_052775230.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mcal	XP_052066038.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mcchi	DN41582_c0_g1.i5.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mcor	CAC40574.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Medu	CAG2242807.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Meigal	VD112691.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mmar	MMAAM0000019873	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mmer	XP_045212365.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mmod	DN23659_c0_g1.i1.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mmer	gl198765_t1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Mphi	scfa:33310:2:9	Fox	Fox-J1	Forkhead domain J1 (410797)		
Obim	XP_052824622.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Oedu	XP_048763213.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Osin	XP_029638410.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Pcan	XP_025095423.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Pcor	DN4891_c0_g1.i2.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Pcor	DN480_c0_g2.i1.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Pcor	DN18891_c0_g2.i2.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Pgen	DN2399_c8_g1.i1.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Pmar	DN32837_c1_g1.i1.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Pmax	XP_033752519.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Poku	DN19777_c2_g1.i1.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Ppur	DN2521_c0_g1.i4.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Pstr	KAK3579229.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Pvir	s01693g10	Fox	Fox-J1	Forkhead domain J1 (410797)		
Pyes	XP_021351058.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Rdec	DN22834_c0_g1.i1.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Rohi	XP_0605887750.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Sbro	EVM0018668.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Scen	Chr1_3201	Fox	Fox-J1	Forkhead domain J1 (410797)		
Scen	Chr1_3198	Fox	Fox-J1	Forkhead domain J1 (410797)		
Sgio	Sgi000050	Fox	Fox-J1	Forkhead domain J1 (410797)		
Sgra	DN11939_c0_g1.i11.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Tgra	KA.J8318321.1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Tsqu	DN6625_c2_g1.i2.p1	Fox	Fox-J1	Forkhead domain J1 (410797)		
Acal	XP_005111247.3	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Arc	Contig53_201	Fox	Fox-J2/3	Forkhead domain J3 (410826)		
Amar	Ama34942	Fox	Fox-J2/3	Forkhead domain J3 (410826)		
Apur	scffold_11801_24	Fox	Fox-J2/3	Forkhead domain J3 (410826)		
Bjala	XP_013070049.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Cang	DN6625_c2_g1.i2.p1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Cari	EVMM0008910.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Cflu	DN58808_c0_g1.i1.p1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Cgig	XP_0114222959.2	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Cpli	DN78731_c5_g1.i2.p1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Csin	Hic_asn_4_381	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Cvir	XP_022341777.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Dpol	XP_052776138.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Gaeg	XP_041378546.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Hbia	M0000010754	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Hruf	XP_048247606.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Mare	XP_052759624.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Mcgal	XP_052759802.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Mcchi	DN34970_c0_g1.i2.p1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Mcor	CAC5378041.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Medu	CAG2221519.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Meigal	VD157447.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Mmer	MMAAM0000017129	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Mmer	XP_053378821.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Mmod	DN36910_c0_g1.i1.p1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Mphi	scfa:405760.4	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Obim	XP_052832979.1	Fox	Fox-J2/3	Forkhead domain FOXJ2, FOXJ3 (410798)		

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Oedu	XP_048739234.1	Fox	Fox-J2.3	Forkhead domain J3 (410826)		
Osin	XP_029651657.1	Fox	Fox-J2.3	Forkhead domain J3 (410826)		
Pcan	XP_025081218.1	Fox	Fox-J2.3	Forkhead domain J3 (410826)		
Pcor	DN2942_c0_g4.i15.p1	Fox	Fox-J2.3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Pcor	DN2942_c0_g4.i3.p1	Fox	Fox-J2.3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Pgen	DN5381_c0_g17.p1	Fox	Fox-J2.3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Pmar	DN41364_c0_g1.i3.p1	Fox	Fox-J2.3	Forkhead domain J3 (410826)		
Pmax	XP_033763328.1	Fox	Fox-J2.3	Forkhead domain J3 (410826)		
Poku	DN14771_c0_g2.i3.p1	Fox	Fox-J2.3	Forkhead domain J3 (410826)		
Ppur	DN2181_c0_g15.p1	Fox	Fox-J2.3	Forkhead domain J3 (410826)		
Pstr	KAK3883417.1	Fox	Fox-J2.3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Pvir	s0019g6	Fox	Fox-J2.3	Forkhead domain J3 (410826)		
Pyes	XP_021374633.1	Fox	Fox-J2.3	Forkhead domain J3 (410826)		
Rphi	XP_000577634.1	Fox	Fox-J2.3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Sbro	EVM0013081.1	Fox	Fox-J2.3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Chr12_800	SG1003279	Fox	Fox-J2.3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Scon	SG1003279	Fox	Fox-J2.3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Sglo	DN1042_c0_g1.i2.p1	Fox	Fox-J2.3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Sgra	KAJ3320220.1	Fox	Fox-J2.3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Tgra	DN10376_c0_g1.i2.p1	Fox	Fox-J2.3	Forkhead domain FOXJ2, FOXJ3 (410798)		
Tsqu	XP_005092494.1	Fox	Fox-K	Forkhead domain K (410800)		
Acal	scafid:85.43	Fox	Fox-K	Forkhead domain K (410800)		
Apur	XP_013090285.1	Fox	Fox-K	Forkhead domain K (410800)		
Bgia	XP_052688140.1	Fox	Fox-K	Forkhead domain K (410800)		
Cang	EVMM00000012630	Fox	Fox-K	Forkhead domain K (410800)		
Cari	XP_011416099.1	Fox	Fox-K	Forkhead domain K (410800)		
Cgig	DN61350_c0_g1.i1.p1	Fox	Fox-K	Forkhead domain K (410800)		
Cpli	XP_022316096.1	Fox	Fox-K	Forkhead domain K (410800)		
Cvir	XP_041362451.1	Fox	Fox-K	Forkhead domain K (410800)		
Gaeg	M00000000333	Fox	Fox-K	Forkhead domain K (410800)		
Hbrf	XP_048248693.1	Fox	Fox-K	Forkhead domain K (410800)		
Mmar	MMAM00000012630	Fox	Fox-K	Forkhead domain K (410800)		
Mphi	XP_145801.11	Fox	Fox-K	Forkhead domain K (410800)		
Obim	XP_014772374.1	Fox	Fox-L	Forkhead domain L (410800)		
Oedu	XP_048761057.2	Fox	Fox-L	Forkhead domain L (410800)		
Osin	XP_029646877.1	Fox	Fox-L	Forkhead domain L (410800)		
Rdec	DN21696_c3_g.i1.p1	Fox	Fox-L	Forkhead domain L (410800)		
Sgio	SG100589	Fox	Fox-L	Forkhead domain L (410800)		
Acal	XP_012940028.1	Fox	Fox-L	Forkhead domain L (410800)		
Arc	Contig56.64	Fox	Fox-L	Forkhead domain L (410800)		
Amar	AMA17914	Fox	Fox-L	Forkhead domain L (410800)		
Apec	DN74037_c0_g1.i1.p1	Fox	Fox-L	Forkhead domain L (410800)		
scaffold122.2	Fox	Fox-L	Fox-L	Forkhead domain L (410800)		
Apur	Bgia	Fox	Fox-L	Forkhead domain L (410800)		
Bgia	XP_055890278.1	Fox	Fox-L	Forkhead domain L (410800)		
Cang	XP_022718868.1	Fox	Fox-L	Forkhead domain L (410800)		
Cari	EVM001909.1	Fox	Fox-L	Forkhead domain L (410800)		
Cgig	XP_011417586.2	Fox	Fox-L	Forkhead domain L (410800)		
Cpli	DN15469_c0_g.i1.p1	Fox	Fox-L	Forkhead domain L (410800)		
Cvir	Hic.asn.17.1225	Fox	Fox-L	Forkhead domain L (410800)		
Lorb	XP_022346240.1	Fox	Fox-L	Forkhead domain L (410800)		
Mare	XP_052252043.1	Fox	Fox-L	Forkhead domain L (410800)		
Mcal	XP_052063315.1	Fox	Fox-L	Forkhead domain L (410800)		
Gaeg	CAC5374005.1	Fox	Fox-L	Forkhead domain L (410800)		
Hbrf	XP_046344397.2	Fox	Fox-L	Forkhead domain L (410800)		
Medu	CAG2206845.1	Fox	Fox-L	Forkhead domain L (410800)		
Mgal	VD122484.1	Fox	Fox-L	Forkhead domain L (410800)		
Mgal	VDH97507.1	Fox	Fox-L	Forkhead domain L (410800)		
Mmar	MNMAN0000002876	Fox	Fox-L	Forkhead domain L (410800)		
Mmer	XP_053402988.1	Fox	Fox-L	Forkhead domain L (410800)		
Mmod	DN51324_c0_g1.i1.p1	Fox	Fox-L	Forkhead domain L (410800)		
Mner	g286924.t1	Fox	Fox-L	Forkhead domain L (410800)		
Mphi	scf69950.0	Fox	Fox-L	Forkhead domain L (410800)		
Obim	XP_014785001.1	Fox	Fox-L	Forkhead domain L (410800)		

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Oedu	XP_048762056.2	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Pcan	XP_025076243.1	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Pcor	DN23326_0.g1.i1.p1	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Pcor	DN181497.co.g1.i1.p1	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Pmar	DN30135.co.g1.i1.p1	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Pmax	XP_033755354.1	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Ppar	DN78331.co.g1.i1.p1	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Pstr	KAK3390991.1	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Pvir	s02023g11	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Pyes	XP_021346465.1	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Rphi	XP_000080839.1	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Sbro	EVM0016190.1	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Scon	Chr19.1.868	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Sglo	Sg1009486	Fox	Fox-L1	Fox	Forkhead domain L1 (410801)	
Acal	XP_005101910.2	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Airc	Contig551.34	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Amar	Ama34673	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Apur	scaffold84.159	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Bgia	XP_055865110.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Cang	EVMD0021728.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Cari	DN127322_6.g2.i2.p1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Cflu	NP_001295827.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Cggg	DN75086_c5.g1.i2.p1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Cpli	Hic_asn_4.274	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Csin	XP_022345405.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Cvir	XP_022345173.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Dpol	XP_052212727.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Gaeq	XP_041378252.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Hbia	M0000035173	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Hrfu	XP_048250285.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Lorb	DN129129_co.g1.i1.p1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Mare	XP_052760962.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Mcal	XP_052082415.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Mcor	CAC401149.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Medu	CAG2239672.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Mgal	VD149865.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Mgal	VD149864.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Mmar	MMAN00000016212	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Mmer	XP_045161614.2	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Mmod	DN2410.co.g1.i1.p1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Mmer	g832_35.tl	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Mphi	scf50301.0.3	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Obim	XP_014785648.2	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Oedu	XP_048729555.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Pcan	XP_025083514.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Pcor	DN35937.co.g1.i1.p1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Pgen	DN134171_co.g1.i1.p1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Pmar	EVMD0017513.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Pmax	XP_033724493.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Pstr	KAK3602726.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Pvir	s00246g193	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Pyes	XP_021353421.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Rdec	DN21003.co.g1.i2.p1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Rphi	XP_006086301.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Sbro	EVM0014371.1	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Scon	Chr12.1684	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Sglo	Sg1005363	Fox	Fox-L2	Fox	Forkhead domain L2 (410802)	
Tsqu	DN37_c29_g1.i1.p1	Fox	Fox-N1/4	Fox	Forkhead domain N1 (410804)	
Acal	Contig281.47	Fox	Fox-N1/4	Fox	Forkhead domain N1 (410804)	
Amar	Ama23426	Fox	Fox-N1/4	Fox	Forkhead domain N1 (410804)	
Apuc	DN27027.co.g1.i2.p1	Fox	Fox-N1/4	Fox	Forkhead domain N1 (410804)	
Apuc	scaffold17.163	Fox	Fox-N1/4	Fox	Forkhead domain N1 (410804)	
Bria	XP_055899100.1	Fox	Fox-N1/4	Fox	Forkhead domain N1 (410804)	
Cang	XP_052711314.1	Fox	Fox-N1/4	Fox	Forkhead domain N1 (410804)	
Cari	EVM002431.1	Fox	Fox-N1/4	Fox	Forkhead domain N1 (410804)	
Cggg	XP_034503195.1	Fox	Fox-N1/4	Fox	Forkhead domain N1 (410804)	

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Pssm-ID)	Additional domains (Pssm-ID)	Notes
Cphi	DN7831.cl.g.i1.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Csin	Hic.asn.2.101	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Cvir	XP_022992787.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Dpol	XP_052270473.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Gaeg	XP_041365083.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Hbia	M0000027642	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Hruf	XP_048241610.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Mare	XP_052801997.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Mcal	XP_052066630.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Mchi	DN2972.cl.g.i1.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Mcor	CAC5383890.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Medu	CAG2257106.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Mgal	VDH93464.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Mgal	VDH93462.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Mgal	VDH93463.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Mmar	MMAN0000018109	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Mmer	DN2977.cl.g2.i1.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Mmod	scf.69935.0.10	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Mphi	XP_02925413.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Obim	XP_055996035.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Oedu	XP_029638459.2	Fox	Fox-N1/4	Forkhead domain N1 (410831)		
Osin	XP_025087495.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Pcan	DN55558.cl.g2.i1.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Pcor	DN55558.cl.g1.i1.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Pcor	DN1454626.co.i1.i1.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Pgen	DN30748.cl.g1.i1.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Pmar	XP_033751425.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Pmax	DN45531.0.cl.i1.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Poku	DN199653.cl.g1.i1.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Ppur	KAK3387366.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Pstr	S24333g45	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Pvir	XP_021371548.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Pyes	DN20122.cl.g1.i1.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Rdec	XP_060066682.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Rphi	EVM0009578.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Sbro	Chr14.2061	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Scon	Sg104456	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Sglo	DN11935.cl.g2.i3.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Sgra	KAJ8298705.1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Tgra	DN22139.cl.g1.i1.p1	Fox	Fox-N1/4	Forkhead domain N1 (410804)		
Tsqu	XP_050599217.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Acal	Contig117.153	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Airc	Amar9979	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Amar	DN11918.cl.g1.i10.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Apuc	scf.489.22	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Apur	XP_013084252.2	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Bgia	XP_052698143.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Cang	EV/M001646.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Carl	DN125734.cl.g1.i18.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Cflu	XP_044242425.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Cgig	DN79231.cl.g1.i7.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Cphi	XP_046351344.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Csin	Hic.asn.10.136	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Cvir	XP_022331167.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Dpol	XP_052270965.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Gaeg	XP_041353111.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Hbia	M0000030949	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Hruf	XP_046351344.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Mare	XP_052767715.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Mcal	XP_052107190.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Mchi	DN39446.cl.g1.i1.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Mcor	CAC5378437.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Medu	CAG2235611.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Mgal	VDI80286.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Mgal	VDI80287.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Mmar	MMAM00000016688	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Mmer	XP_053376884.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Mmod	DN2418.cl.g1.i131.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Mnre	g186153.t2	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Mphi	scaf_37509_0.5	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Obim	XP_0528222674.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Oedu	XP_0487356022.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Osin	XP_0296333483.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Pcan	XP_0251060888.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Pcor	DN195240_c0.g1.i1.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Pcor	DN5191_c0.g2.i7.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Pcor	DN18451_c0.g2.i3.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Pgen	DN14328_c0.g1.i3.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Pmar	DN42157_c1.g1.i3.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Pmax	XP_033734749.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Poku	DN17429_c4.g1.i2.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Ppur	DN5075_c0.g1.i1.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Pstr	KAK3995953.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Pvir	s00410g05	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Pyes	XP_021366964.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Rdec	DN2296.c2.g1.i1.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Rphi	XP_060552999.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Sgio	Sg1013452	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Sgra	DN13133_c0.g2.i8.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Tgra	KAJ93208641.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Tgra	KAJ93167271.1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Tsqu	DN75347_c0.g1.i2.p1	Fox	Fox-N2/3	Forkhead domain N3 (410833)		
Acal	XP_005112460.1	Fox	Fox-O	Forkhead domain O (410806)		
Airc	Config16.24	Fox	Fox-O	Forkhead domain O (410806)		
Amar	Amat07814	Fox	Fox-O	Forkhead domain O (410806)		
Apec	DN23636_c0.g1.i1.p1	Fox	Fox-O	Forkhead domain O (410806)		
Apur	scaffold_712.6	Fox	Fox-O	Forkhead domain O (410806)		
Bgia	XP_013095255.2	Fox	Fox-O	Forkhead domain O (410806)		
Cang	EVMD0000968.1	Fox	Fox-O	Forkhead domain O (410806)		
Cari	DN112955_c5.g3.15.p1	Fox	Fox-O	Forkhead domain O (410806)		
Cflu	XP_011414359.1	Fox	Fox-O	Forkhead domain O (410806)		
Cgg	DN72415_c5.g1.i1.p1	Fox	Fox-O	Forkhead domain O (410806)		
Cpli	XP_022287692.1	Fox	Fox-O	Forkhead domain O (410806)		
Cvir	XP_022287423.1	Fox	Fox-O	Forkhead domain O (410806)		
Dpol	XP_022243903.1	Fox	Fox-O	Forkhead domain O (410806)		
Geg	XP_041351714.1	Fox	Fox-O	Forkhead domain O (410806)		
Hbia	M0000024236	Fox	Fox-O	Forkhead domain O (410806)		
Hruf	XP_046374293.1	Fox	Fox-O	Forkhead domain O (410806)		
Lorb	DN142512_c0.g1.i1.p1	Fox	Fox-O	Forkhead domain O (410806)		
Mare	XP_052808484.1	Fox	Fox-O	Forkhead domain O (410806)		
Mcal	XP_052087664.1	Fox	Fox-O	Forkhead domain O (410806)		
Mchi	DN32908_c0.g1.i3.p1	Fox	Fox-O	Forkhead domain O (410806)		
Mcor	CAC592548.1	Fox	Fox-O	Forkhead domain O (410806)		
Mmar	MMAM00000004017	Fox	Fox-O	Forkhead domain O (410806)		
Mmer	XP_045195791.2	Fox	Fox-O	Forkhead domain O (410806)		
Mmod	DN145_c2.g1.i2.p1	Fox	Fox-O	Forkhead domain O (410806)		
Mner	g187283.t1	Fox	Fox-O	Forkhead domain O (410806)		
Obim	XP_025232486.1	Fox	Fox-O	Forkhead domain O (410806)		
Oedu	XP_048766900.1	Fox	Fox-O	Forkhead domain O (410806)		
Osin	XP_029657040.1	Fox	Fox-O	Forkhead domain O (410806)		
Pcan	DN22971_c0.g1.i3.p1	Fox	Fox-O	Forkhead domain O (410806)		
Pcor	DN44399_c0.g5.i1.p1	Fox	Fox-O	Forkhead domain O (410806)		
Pcor	DN7549_c0.g3.i2.p1	Fox	Fox-O	Forkhead domain O (410806)		
Pgen	DN24871_c0.g1.i3.p1	Fox	Fox-O	Forkhead domain O (410806)		
Pgen	DN24871_c0.g1.i3.p2	Fox	Fox-O	Forkhead domain O (410806)		
Pmar	DN44399_c0.g5.i1.p1	Fox	Fox-O	Forkhead domain O (410806)		
Pmax	XP_033740844.1	Fox	Fox-O	Forkhead domain O (410806)		
Poku	DN10962_c1.g2.i2.p1	Fox	Fox-O	Forkhead domain O (410806)		
Ppur	DN72510_c0.g1.i1.p1	Fox	Fox-O	Forkhead domain O (410806)		
Pstr	KAK3576955.1	Fox	Fox-O	Forkhead domain O (410806)		
Pvir	s000798.02	Fox	Fox-O	Forkhead domain O (410806)		
Pyes	XP_021377366.1	Fox	Fox-O	Forkhead domain O (410806)		
Rdec	DN30721_c0.g1.i1.p1	Fox	Fox-O	Forkhead domain O (410806)		
Rphi	XP_060589384.1	Fox	Fox-O	Forkhead domain O (410806)		
Scon	Chr2.491	Fox	Fox-O	Forkhead domain O (410806)		

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Sgra	Sgra	Fox	Fox-O	Forkhead domain O (410834)		
	DN5576_c0_g1.i1.p1	Fox	Fox-O	Forkhead domain O (410806)		
Tsqu	DN138882_c0_g1.i1.p1	Fox	Fox-O	Forkhead domain O (410806)		
Airc	Contig3461.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Airc	Contig330.72	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Apur	scaffold_376_108	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Cang	XP_052673828.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Cari	EV/M0015778.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Cgig	XP_011412452.2	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Cvir	XP_0123300144.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Dpol	XP_052234997.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Dpol	XP_052237166.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Gaeq	XP_041362068.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Hbia	M00000010651	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Hruf	XP_048236781.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Hruf	XP_046328651.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Mcal	XP_052089402.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Mcor	CAC5388114.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Medu	CAG2250347.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Mgal	VD105563.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Mgal	VD105564.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Mmar	MMAAM00000027087	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Mmer	DN9753_c0_g1.i1.p1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Mmod	scaf_56119_0.21	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Mphi	XP_048731527.2	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Oedu	DN32892_c0_g1.i1.p1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Pmar	XP_045182963.2	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Pmar	XP_033727511.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Petr	KAK3609024.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Pvir	s01298g51	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Pyes	XP_021354438.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Rlef	DN23702_c2_g1.i2.p1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Rphi	XP_060966633.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Sbro	EV/M0019544.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Scor	Chr1_409	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Sgio	Sgi00484	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Tgra	KAJ322379.1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Tsqu	DN207442_c0_g1.i1.p1	Fox	Fox-OG13/NA	Forkhead domain P (410807)		
Acal	XP_0005106916.3	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Airc	Contig85.21	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Amar	Ama19770	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Apur	scaffold_360_14	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Bglj	XP_013071662.2	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Cang	XP_0520776257.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Cari	EV/M001823.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Cgig	XP_011439389.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Cpli	DN34749_c0_g1.i1.p1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Csin	Hic_asn_12_159	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Cvir	XP_022296913.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Dpol	XP_052253230.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Dpol	XP_052253240.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Gaeq	XP_041347345.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Hbia	MWAV00000015843	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Hruf	XP_046382017.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Mare	XP_0522778846.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Mcal	XP_052062481.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Mcor	CAC54195.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Medu	CAG2199155.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Mgal	VD105805.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Mmar	MMAAM00000012410	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Mmer	XP_045166371.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Mphi	scaf_567833_0.2	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Obim	XP_014772941.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Oedu	XP_048742700.2	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Pcan	XP_025096321.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Pcor	DN7667_c0_g1.i1.p1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Pcor	DN7667_c0_g3.i1.p1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	
Pmax	XP_033744896.1	Fox	Fox-OG15/NA	Forkhead domain Q (410809)	Annotated as Fox-Q2b	

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Pstr	KAK3389497.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Pyes	XP_021371037.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Rphi	XP_060586724.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Sbro	EVM00050606.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Sbro	EVM0006433.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Scon	Ch7.1624	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Sgo	Sgo24307	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Airc	Contig.1425.7	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
scaffold	604.173	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Apur	XP_052676026.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Cang	EVM0023364.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Cari	DN107758.c5.g1.i2.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Cflu	XP_019927657.2	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Cgig	Cp1.	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Cp1.	DN70609.c0.g1.i2.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Cvnr	XP_022321288.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Cvir	XP_02295893.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Dpol	XP_052222923.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Gaeq	XP_041365712.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Hbia	M0000030826	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Hruf	DN243726.c0.g1.i1.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Lorb	XP_046364140.2	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mare	XP_052802309.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mcor	CA5370465.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Medu	CAG2202185.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Medu	CAG2246856.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mgal	VD12465.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mmar	MMAN00000032793	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mmer	XP_045177123.2	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mphi	scaf.70200.0.4	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Obim	XP_014771053.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Oedu	XP_048728661.2	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Osin	DM42205.c4.g1.i2.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Pmar	XP_033752233.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Pmax	DN4666.c0.g1.i2.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Ppur	DN4666.c0.g1.i2.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Pstr	KAK3581527.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Pvir	s00194g51	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Pyes	XP_0215353413.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Rdec	DN22502.c0.g4.i1.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Rphi	XP_006599562.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Sbro	XP_000601370.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Scon	EVM0002125.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Ch14.1628	DN2384.c1.g1.i1.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Tsqu	DN113069.c3.g1.i4.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Coli	DN76173.c0.g1.i2.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Airc	Hic.asm.2.1802	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Apec	Csin	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Apur	scaffold.333.74	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Cang	XP_052712245.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Cari	EVMM0012682.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Cflu	XP_041366058.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Gaeq	DN76173.c0.g1.i2.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Hbia	M0000014061	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Hruf	XP_046335487.2	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Lorb	XP_052286391.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mare	XP_052228333.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mcal	XP_052234337.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Dpol	XP_052064572.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mcor	CAC537062.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Medu	CAG2224977.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mmar	MMAN00000037791	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mmer	XP_053384136.1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Cvnr	DN27089.c0.g1.i1.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Mcab	g144.243.t1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Oedu	XP_048731286.2	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	
Pcor	DN5679.c0.g2.i1.p1	Fox	Fox-OG15/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2b	

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Pssm-ID)	Additional domains (Pssm-ID)	Notes
Pcor	DN15679_c0g1.i15.p1	Fox	Forkhead domain M (410803)	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Pcor	DN113056_c0g1.i1.p1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Pcor	DN113056_c0g2.i2.p1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Pgen	DN128413_c1.g1.i2.p1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Pmar	DN44947_c1.g1.i3.p1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Pmax	XP_033751305_1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Poku	DN37223_c1.g1.i8.p1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Ppur	DN3451_c0g1.i1.p1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Pstr	KAK3597624_1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Pvir	S00219g11	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Pyes	XP_021377259_1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Rdec	DN22152_c4.g1.i1.p1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Rphi	XP_060569990_1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Sbro	EVM0023670_1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Sgio	Sg1005561	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Sgra	DN54780_c0g1.i1.p1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Tgra	KAJ3299135_1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Tsau	DN6434_0.g1.i7.p1	Fox	Fox-OG2/NA	Fox-OG2/NA	Forkhead domain M (410803)	Annotated as Fox-M
Airc	Contig465_41	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Amar	Ama25953	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Apur	scaffid_367_41	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Cang	XP_02700156_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Cari	EVM0004465_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Cejig	XP_01143457_2	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Csin	Hic-asn_16_939	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Cvir	XP_02233426_3_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Dpol	XP_052278875_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Dpol	XP_052278876_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Dpol	XP_052278804_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Hrf	XP_046341176_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mare	XP_052791887_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mare	XP_052791890_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mare	XP_052791888_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mare	XP_052791891_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcal	XP_052099876_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcal	XP_052099855_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcor	CAC5419385_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcor	CAC5380823_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcor	CAC5370920_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcor	CAC5419389_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcor	CAC5419386_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcor	CAC5419381_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcor	CAC5419382_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcor	CAC5419383_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcor	CAC5419387_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcor	CAC5419380_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mcor	CAC5419388_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Medu	CAG22194460_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Medu	CAG2194706_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Medu	CAG2198066_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Medu	CAG2198058_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Medu	CAG2198055_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Medu	CAG2214461_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Medu	CAG2198060_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Medu	CAG2198061_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Medu	CAG2198064_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Medu	CAG2198062_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mgal	VDI32850_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mgal	VDI39859_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mgal	VDI15906_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mgal	VDI02348_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c
Mgal	VDI15903_1	Fox	Fox-OG28/NA	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Pssm-ID)	Additional domains (Pssm-ID)	Notes
Mgal	VD102347.1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Mgal	VD115905.1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Mgal	VD102349.1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Mgal	VD115904.1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Mimer	XP_053405097.1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Mmod	DN6982_C0_&1.13_p1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Mphi	scfa.15444.01	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Oedu	XP_056021213.1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Sbro	DN39963_C0_g1.i1.p1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Pmax	XP_053751006.1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Pmax	XP_033749723.1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Pvir	s03437g48	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Pvir	XP_021360588.1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Rphi	XP_060585777.1	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
EVM0013029.1	EV	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Ch5.396.1	C	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Scon	Sc013625	Fox	Fox-OG28/NA	Forkhead domain Q2 (410809)	Annotated as Fox-Q2c	
Sj01	XP_005102249.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Acal	Contig636.41	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Airc	Amar2012	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Amar	scaffid-313_52	Fox	Fox-OG39/NA	Forkhead domain E (410793)	Annotated as Fox-AB	
Apur	XP_055874245.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Bgla	XP_052699279.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Cang	EVMD001854.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Cari	XP_011644298.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Ccg5	XP_022334408.1	Fox	Fox-OG39/NA	Forkhead domain E (410793)	Annotated as Fox-AB	
Cvfr	XP_022334077.1	Fox	Fox-OG39/NA	Forkhead domain E (410793)	Annotated as Fox-AB	
Dpol	XP_0522785869.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Gaeq	XP_041375684.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Hbia	M00000015535	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Hruf	XP_046350707.2	Fox	Fox-OG39/NA	Forkhead domain E (410793)	Annotated as Fox-AB	
Hruf	XP_046350707.0.1	Fox	Fox-OG39/NA	Forkhead domain E (410793)	Annotated as Fox-AB	
Hruf	XP_046350686.2	Fox	Fox-OG39/NA	Forkhead domain E (410793)	Annotated as Fox-AB	
Hruf	XP_046350709.2	Fox	Fox-OG39/NA	Forkhead domain E (410793)	Annotated as Fox-AB	
Hruf	XP_046350688.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Hruf	XP_046350660.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Hruf	XP_046350087.2	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Hruf	XP_046350714.2	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Hruf	XP_046350712.2	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Hruf	XP_046350708.2	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Mare	XP_032795236.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Mcal	XP_022102496.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Mcor	CAC014394.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Medu	CAG2193762.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Mgal	VD13594.2.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Mgal	MMAN000000238.30	Fox	Fox-OG39/NA	Forkhead domain E (410793)	Annotated as Fox-AB	
Mimer	XP_045215157.2	Fox	Fox-OG39/NA	Forkhead domain E (410793)	Annotated as Fox-AB	
Mphi	scfa.711.0.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Oedu	XP_048737442.2	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Pcan	XP_025078030.1	Fox	Fox-OG39/NA	Forkhead domain E (410793)	Annotated as Fox-AB	
Pmax	XP_023750990.1	Fox	Fox-OG39/NA	Forkhead domain E (410793)	Annotated as Fox-AB	
Pstr	KAK3001439.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Tgra	KAK3601419.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Sbro	s001398g393	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Pvir	XP_021357612.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Pyes	XP_060589081.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Rphi	EVMD0003782.1	Fox	Fox-OG39/NA	Forkhead domain L1 (410801)	Annotated as Fox-AB	
Scon	Chr5.18	Fox	Fox-OG39/NA	Forkhead domain P2 (410839)	-	
Sj01	Sg100401	Fox	Fox-OG39/NA	Forkhead domain P2 (410839)	-	
Tgra	KAJ33049.16.1	Fox	Fox-OG39/NA	Forkhead domain P2 (410839)	-	
Airc	Contig12.132	Fox	Fox-P	Fox-P	-	
Apur	scaffid-507.3	Fox	Fox-P	Fox-P	-	
Bgla	XP_055882746.1	Fox	Fox-P	Fox-P	-	
Cang	EVMD001529.1	Fox	Fox-P	Fox-P	-	
Cari	KAJ33049.16.1	Fox	Fox-P	Fox-P	-	
Cflu	DN101403_c0_g2_i3_p1	Fox	Fox-P	Fox-P	-	
Cgig	XP_01141930.2	Fox	Fox-P	Fox-P	-	
Cipl	DN80534_c6_g1_i4_p1	Fox	Fox-P	Fox-P	-	
Csin	Hic.asm.12.158	Fox	Fox-P	Fox-P	-	

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Pssm-ID)	Additional domains (Pssm-ID)	Notes
Cvir	XP_02295365.1	Fox	Fox-P	Forkhead domain P (410807)	FOXP coiled-coil domain (465036)	
Dpol	XP_02252968.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Gaeq	XP_041347582.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Hruf	XP_048239143.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Lorb	DN59652.cl.g1.i4.p1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mare	XP_052780914.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mcal	XP_052101037.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mcal	XP_052083749.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mcal	XP_052100969.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mcal	XP_052061774.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mchi	DN46200.co.g1.i1.p1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mcor	CAC5419517.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Medu	CAG2199156.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mgal	VD14555.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mgal	VD150808.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mgal	VD150806.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mgal	VD150807.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mmar	MMAAM0000012411.	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mmer	XP_053376718.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mmod	DN2151.co.g1.i2.p1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mner	g93547.tl	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Mphi	scfa:39347.0.7	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Obim	XP_05282821.1	Fox	Fox-P	Forkhead domain P (410807)	FOXP coiled-coil domain (465036)	
Oedu	XP_055998748.1	Fox	Fox-P	Forkhead domain P (410807)	FOXP coiled-coil domain (465036)	
Osin	XP_036357858.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Pcan	XP_025106713.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Pcor	DN1820.co.g1.i26.p1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Pgen	DN3661.co.g1.i6.p1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Pmar	DN4268.co.g1.i1.p1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Pmax	XP_033745571.1	Fox	Fox-P	Forkhead domain P (410807)	FOXP coiled-coil domain (465036)	
Poku	DN88526.c2.g1.i1.p1	Fox	Fox-P	Forkhead domain P (410807)	FOXP coiled-coil domain (465036)	
Pstr	KAK3589495.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Pvir	s01329g.124	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Pves	XP_021363304.1	Fox	Fox-P	Forkhead domain P (410807)	FOXP coiled-coil domain (465036; partial)	
Rdec	DN8828.co.g1.i1.p1	Fox	Fox-P	Forkhead domain P (410807)	FOXP coiled-coil domain (465036)	
Rphi	XP_040586741.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Sbro	EVMM004295.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Scon	Chr7:2129	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036; partial)	
Sglo	Sg011345	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Tgra	KAJ830234.1	Fox	Fox-P	Forkhead domain P2 (410839)	FOXP coiled-coil domain (465036)	
Acal	XP_005099459.2	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Airc	Contig1420.28	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Amar	Ama2905	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Apur	Hic.asn.16.4	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Bglia	scaffid8332.35	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Cang	XP_0558665367.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Cari	DN17101.co.g1.i1.p1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Cgig	EVM0002665.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Cphi	XP_011425762.2	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Csin	DN105612.co.g1.i1.p1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Cvir	XP_022333968.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Dpol	XP_0522808961.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Gaeq	XP_041363029.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Hbia	XP_046373579.2	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Hruf	XP_052101305.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Mcal	DN2286.co.g1.i1.p1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Mchi	CAC5383792.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Mcor	CAG2191193.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Medu	VD174621.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Mgal	MMAM00000000686	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Mmar	XP_045215524.2	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Mner	g20053.tl	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Mphi	scfa:22910.1.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Obim	XP_014767584.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Pcan	XP_025078472.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Pcor	DN19235_c0.g1.i1.p1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Pmar	DN49466_c0.g1.i1.p1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Pmax	XP_033751003.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Pstr	KAK3959133.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Pvir	s00115g23	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Pyes	XP_021243668.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Rphi	XP_06051531.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Sbro	EVM0023378.1	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Scn	Ch5_1974	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Scn	Ch5_2105	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Sgio	Sg009183	Fox	Fox-Q2	Forkhead domain Q2 (410809)	Annotated as Fox-Q2a	
Acal	XP_005109004.3	Fox	-	Forkhead domain (410788)	-	
Airc	Contig879_9	Fox	-	Forkhead domain Q2 (410809)	-	
Amar	Ama25952	Fox	-	Forkhead domain Q2 (410809)	-	
Amar	Am227375	Fox	-	Forkhead domain Q2 (410809)	-	
Cang	XP_052876682.1	Fox	-	Forkhead domain H (410796)	-	
Cang	XP_0526800388.1	Fox	-	Forkhead domain H (410796)	-	
Cang	XP_052677368.1	Fox	-	Forkhead domain H (410796)	-	
Cari	EVM0001935.1	Fox	-	Forkhead domain H (410796)	-	
Cari	EVM0027332.1	Fox	-	Forkhead domain H (410796)	-	
Cgig	XP_034206826.1	Fox	-	Forkhead domain H (410796)	-	
Cgig	XP_011447567.2	Fox	-	Forkhead domain H (410796)	-	
Cpli	DN157619_c0.g1.i1.p1	Fox	-	Forkhead domain H (410796)	-	
Cvir	XP_022300767.1	Fox	-	Forkhead domain (410788)	-	
Dpol	XP_022300750.1	Fox	-	Forkhead domain (410788)	-	
Dpol	XP_052227296.1	Fox	-	Forkhead domain Q2 (410809)	-	
Gaeq	XP_041366967.1	Fox	-	Forkhead domain Q2 (410809)	-	
Gaeq	XP_041378820.1	Fox	-	Forkhead domain Q2 (410809)	-	
Gaeq	XP_041347225.1	Fox	-	Forkhead domain Q2 (410809)	-	
Gaeq	XP_041375925.1	Fox	-	Forkhead domain Q2 (410809)	-	
Gaeq	XP_041375913.1	Fox	-	Forkhead domain Q2 (410809)	-	
Gaeq	XP_041379015.1	Fox	-	Forkhead domain N1 (410804)	-	
Hbia	M0000018946	Fox	-	Forkhead domain Q2 (410809)	-	
Mare	XP_052791886.1	Fox	-	Forkhead domain Q2 (410809)	-	
Mare	XP_052771066.1	Fox	-	Forkhead domain Q2 (410809)	-	
Mcal	XP_052098820.1	Fox	-	Forkhead domain Q2 (410809)	-	
Mcor	CAC5419379.1	Fox	-	Forkhead domain Q2 (410809)	-	
Medu	CAG2194707.1	Fox	-	Forkhead domain Q2 (410809)	-	
Medu	CAG2208945.1	Fox	-	Forkhead domain L1 (410801)	-	
Mgal	VD115902.1	Fox	-	Forkhead domain Q2 (410809)	-	
Mgal	VD152978.1	Fox	-	Forkhead domain L1 (410801)	-	
Mmar	MMAM00000049704.	Fox	-	Forkhead domain Q2 (410809)	-	
Mmer	XP_045216636.2	Fox	-	Forkhead domain Q2 (410809)	-	
Mmer	XP_045189131.2	Fox	-	Forkhead domain Q2 (410809)	-	
Oedu	gl59704_t1	Fox	-	Forkhead domain Q2 (410809)	-	
Osin	XP_036359188.1	Fox	-	Forkhead domain Q2 (410809)	-	
Mphi	scfa:46189_0.0	Fox	-	Forkhead domain Q2 (410809)	-	
Mphi	scfa:15444_0.2	Fox	-	Forkhead domain Q2 (410809)	-	
Mphi	scfa:27787_1.10	Fox	-	Forkhead domain L1 (410801)	-	
Obim	XP_042777604.1	Fox	-	Forkhead domain M (410803)	-	
Pcor	XP_048739629.2	Fox	-	Forkhead domain H (410796)	-	
Pcor	DN115905_c0.g1.i1.p1	Fox	-	Forkhead domain P (410807)	-	
Pmax	XP_033750561.1	Fox	-	Forkhead domain Q2 (410809)	-	
Pstr	KAK3385306.1	Fox	-	Forkhead domain Q2 (410809)	-	
Pvir	s00585g48	Fox	-	Forkhead domain Q2 (410801)	-	
Pyes	XP_021248419.1	Fox	-	Forkhead domain Q2 (410809)	-	
Rdec	DN235206_c2.g1.i1.p1	Fox	-	Forkhead domain Q2 (410809)	-	
Rphi	XP_060585776.1	Fox	-	Forkhead domain Q2 (410809)	-	
Scn	Ch5_397	Fox	-	Forkhead domain Q2 (410809)	-	
Sgio	DN23960_c0.g1.i1.p1	Fox	-	Forkhead domain Q2 (410809)	-	
Tsqu	XP_035824685.1	Sox	-	High mobility group box (43838)	-	
Acal				Helix loop helix domain (19764)		

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Acal	XP_012946205.1	Sox	-	High mobility group box (438820)	-	
Acal	XP_005105939.1	Sox	-	High mobility group box (438820)	-	
Apec	DN48806_c0_g1.i1.p1	Sox	-	High mobility group box (438820)	-	
Apec	DN108003_c0_g1.i1.p1	Sox	-	High mobility group box (438820)	-	
scaffold_391_70	-	Sox	-	High mobility group box (438820)	-	
Apur	-	Sox	-	High mobility group box (438820)	-	
BglA	XP_013078241.2	Sox	-	High mobility group box (438820)	-	
BglA	XP_013078156.1	Sox	-	High mobility group box (438820)	-	
BglA	XP_052697278.1	Sox	-	High mobility group box (438820)	-	Helix loop helix domain (197674)
Cang	XP_027136921.1	Sox	-	High mobility group box (438820)	-	Helix loop helix domain (197674, partial)
Car1	EVM0018891.1	Sox	-	High mobility group box (438820)	-	
Cari	EVM005567.1	Sox	-	High mobility group box (438820)	-	
Cgig	XP_011425869.2	Sox	-	High mobility group box (438820)	-	
Cgig	XP_03435619.1	Sox	-	High mobility group box (438820)	-	
Cvir	XP_022330758.1	Sox	-	High mobility group box (438820)	-	
Cvir	XP_02233079.1	Sox	-	High mobility group box (438820)	-	
Dpol	XP_05227104.1	Sox	-	High mobility group box (438820)	-	
Gaeq	XP_041377139.1	Sox	-	High mobility group box (438820)	-	
Hbia	M00000038049	Sox	-	High mobility group box (438820)	-	
Hbia	M0000000498	Sox	-	High mobility group box (438820)	-	
Hruf	XP_04632995.1	Sox	-	High mobility group box (438820)	-	
Hruf	XP_048239511.1	Sox	-	High mobility group box (438820)	-	
Mcor	CAC3848822.1	Sox	-	High mobility group box (438820)	-	
Medu	CAG2253429.1	Sox	-	High mobility group box (438820)	-	
Mgal	VD178477.1	Sox	-	High mobility group box (438820)	-	
Mmod	DN13112_c0_g1.i1.p1	Sox	-	High mobility group box (438820)	-	
Obim	XP_04776519.1	Sox	-	High mobility group box (438820)	-	
Oedu	XP_048738250.2	Sox	-	High mobility group box (438820)	-	
Oedu	XP_048752144.2	Sox	-	High mobility group box (438820)	-	
Osin	XP_036358200.1	Sox	-	High mobility group box (438820)	-	
Osin	XP_029654541.1	Sox	-	High mobility group box (438820)	-	
Osin	XP_029656368.1	Sox	-	High mobility group box (438820)	-	
Osin	XP_029657644.1	Sox	-	High mobility group box (438820)	-	
Osin	XP_029656220.1	Sox	-	High mobility group box A, B and G (438837)	-	
Osin	XP_029657648.1	Sox	-	High mobility group box A, B and G (438837)	-	
Osin	XP_029655056.1	Sox	-	High mobility group box A, B and G (438837)	-	
Osin	XP_029655785.1	Sox	-	High mobility group box A, B and G (438837)	-	
Osin	XP_029656129.1	Sox	-	High mobility group box A, B and G (438837)	-	
Osin	XP_029654991.1	Sox	-	High mobility group box A, B and G (438837)	-	
Pcan	XP_025104729.1	Sox	-	High mobility group box (438820)	-	
Pcor	DN32281_c0_g1.i1.p1	Sox	-	High mobility group box (438820)	-	
Pcor	DN21964_c0_g1.i2.p1	Sox	-	High mobility group box (438820)	-	
Pmar	DN33290_c0_g1.i2.p1	Sox	-	High mobility group box (438820)	-	
Pmar	DN35008_c0_g1.i4.p1	Sox	-	High mobility group box (438820)	-	
Pmax	XP_032755621.1	Sox	-	High mobility group box (438820)	-	
Ppur	DN4784_c0_g1.i4.p1	Sox	-	High mobility group box (438820)	-	
Pyes	XP_021347224.1	Sox	-	High mobility group box (438820)	-	
Sbro	Sg1009175	Sox	-	High mobility group box (438820)	-	
Sglo	Sg1012029	Sox	-	High mobility group box (438820)	-	
Tsqu	DN639_c0_g1.i1.p1	Sox	-	High mobility group box (438820)	-	
Acal	XP_005108230.1	Sox	-	High mobility group box B (438790)	-	
Acal	XP_03524438.1	Sox	-	High mobility group box B (438790)	-	
Contig49_126	-	Sox	-	High mobility group box B (438790)	-	
Airc	Contig14_115	Sox	-	High mobility group box B (438790)	-	
Amar	Ama33032	Sox	-	High mobility group box B (438790)	-	
Amar	Ama33828	Sox	-	High mobility group box B (438790)	-	
Amar	DN32410_c0_g1.i2.p1	Sox	-	High mobility group box B (438790)	-	
Apec	DN12297_c0_g1.i3.p1	Sox	-	High mobility group box B (438790)	-	
Apec	scaffold_15389_10	Sox	-	High mobility group box B (438790)	-	
Apur	scaffold_865_4	Sox	-	High mobility group box B (438790)	-	
BglA	XP_013075432.1	Sox	-	High mobility group box B (438790)	-	
Cang	XP_055668106.1	Sox	-	High mobility group box B (438790)	-	
Cang	XP_052705681.1	Sox	-	High mobility group box B (438790)	-	
Cari	EVMO026792.1	Sox	-	High mobility group box B (438790)	-	
Cari	EVM0013965.1	Sox	-	High mobility group box B (438790)	-	
Cflu	DN11867_c0_g1.i2.p1	Sox	-	High mobility group box B (438790)	-	
Cflu	DN98542_c1_g1.i1.p1	Sox	-	High mobility group box B (438790)	-	
Cgig	XP_011433975.1	Sox	-	High mobility group box B (438790)	-	

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Cggg	XP_011455662.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Cpli	DN31343.c0.g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Cpli	DN95511.c0.g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Csin	Hic.asm.6.930	Sox	Sox-B1/2	High mobility group box B (438790)		
Csin	Hic.asm.6.233.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Cvir	XP_02286516.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Cvir	XP_022243230.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Dpol	XP_052214544.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Dpol	XP_052217420.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Gaeq	XP_041353075.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Gaeq	XP_041357874.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Hbia	M00000002798	Sox	Sox-B1/2	High mobility group box B (438790)		
Hbia	M00000033682	Sox	Sox-B1/2	High mobility group box B (438790)		
Hruf	XP_046370193.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Hruf	XP_046326733.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Lorb	DN80278.c0.g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Lorb	DN14_c4.g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mare	XP_052784929.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mare	XP_052784720.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mcal	XP_052105617.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mchi	XP_052104911.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mchi	DN36332.c0.g1.i2.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mcor	DN45716.c0.g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mcor	CAC5413203.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Medu	CAG229644.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Medu	CAG2206403.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mgal	VD33296.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mgal	VD16960.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mmar	MMAM00000041532	Sox	Sox-B1/2	High mobility group box B (438790)		
Mmar	MMAM00000023253	Sox	Sox-B1/2	High mobility group box B (438790)		
Mmer	XP_045201594.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mmer	XP_0405201080.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mmod	DN1981_c0.g1.i2.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mmod	DN73279.c0.g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mner	g140596.t1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mner	g157489.t1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mphi	scfa.663490.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Mphi	scfa.24206.0.4	Sox	Sox-B1/2	High mobility group box B (438790)		
Obim	XP_04789971.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Obim	XP_04780771.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Oedu	XP_048746651.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Oedu	XP_048746663.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Osin	XP_039654000.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Osin	XP_029655638.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pcan	XP_025079293.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pcan	XP_025078598.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pcor	DN9587.c0.g1.i2.p1	Sox	Sox-B1/2	High mobility group box B (438837)		
Pcor	DN14753.c0.g1.i2.p1	Sox	Sox-B1/2	High mobility group box A, B and C (438837)		
Pcor	DN5688_c0.g1.i10.p2	Sox	Sox-B1/2	High mobility group box B (438790)		
Pcor	DN5688_c0.g3.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pcor	DN9742_c0.g1.i2.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pgen	DN5232_c0.g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pgen	DN1199_c0.g1.i12.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pmar	DN30477.c0.g2.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pmar	DN30459_c0.g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pmax	XP_033760067.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pmax	XP_033759382.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Poku	DN51870.c1.g1.i2.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Poku	DN1067_c0.g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Ppur	DN202737.c0.g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pstr	KAK3586311.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pstr	KAK358936.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pvir	s00319g159	Sox	Sox-B1/2	High mobility group box B (438790)		
Pvir	s00319g281	Sox	Sox-B1/2	High mobility group box B (438790)		
Pyes	XP_021556125.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pyes	XP_021344413.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Pyes	XP_021372128.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Rdec	DN21477.c2.g7.i2.p1	Sox	Sox-B1/2	High mobility group box B (438790)		

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Rphi	XP_06056101.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Rphi	XP_060561544.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Sbro	EVM0016386.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Sbro	EVM0007529.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Scon	Ch9-1352	Sox	Sox-B1/2	High mobility group box B (438790)		
Scon	Ch9-1522	Sox	Sox-B1/2	High mobility group box B (438790)		
Scon	Ch9-1514	Sox	Sox-B1/2	High mobility group box B (438790)		
Sgio	Sg101000	Sox	Sox-B1/2	High mobility group box B (438790)		
Sgio	Sg1020107	Sox	Sox-B1/2	High mobility group box B (438790)		
Sgra	DN3782_c0_g1.i3.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Sgra	DN357_c1_g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Tgra	KAJ8310140.1	Sox	Sox-B1/2	High mobility group box B (438790)		
Tsqu	DN97880_c0_g1.i1.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Tsqu	DN50556_c0_g1.i2.p1	Sox	Sox-B1/2	High mobility group box B (438790)		
Airc	Contig80.70	Sox	Sox-C	High mobility group box C (438838)		
Amar	Ama12726	Sox	Sox-C	High mobility group box C (438838)		
Apec	DN12286_c0_g3.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Apar	scaffold_16_61	Sox	Sox-C	High mobility group box C (438838)		
Cang	XP_052689209.1	Sox	Sox-C	High mobility group box C (438838)		
Cari	EVMO025846.1	Sox	Sox-C	High mobility group box C (438838)		
Cflu	DN1262276_c0_g1.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Cgig	XP_011445203.1	Sox	Sox-C	High mobility group box C (438838)		
Cipli	DN19112_c0_g1.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Csin	Hic_asn_11_1009	Sox	Sox-C	High mobility group box C (438838)		
Cvir	XP_022317619.1	Sox	Sox-C	High mobility group box C (438838)		
Dpol	XP_052257395.1	Sox	Sox-C	High mobility group box C (438838)		
Gaeg	XP_041358324.1	Sox	Sox-C	High mobility group box C (438838)		
Hbia	XP_M000037669	Sox	Sox-C	High mobility group box C (438838)		
Hruf	XP_046365064.1	Sox	Sox-C	High mobility group box C (438838)		
Lorb	DN14941_c0_g1.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Mare	XP_052777703.1	Sox	Sox-C	High mobility group box C (438838)		
Mcal	XP_052087802.1	Sox	Sox-C	High mobility group box C (438838)		
Mchi	DN4498_c0_g4.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Mcor	CAC5424030.1	Sox	Sox-C	High mobility group box C (438838)		
Medu	CAG2189937.1	Sox	Sox-C	High mobility group box C (438838)		
Mgal	VD141453.1	Sox	Sox-C	High mobility group box C (438838)		
Mgal	VD114462.1	Sox	Sox-C	High mobility group box C (438838)		
Mmar	MMAW0000036315	Sox	Sox-C	High mobility group box C (438838)		
Mmer	XP_045158937.1	Sox	Sox-C	High mobility group box C (438838)		
Mmod	DN104308_c0_g1.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Mmer	g264_04.i1	Sox	Sox-C	High mobility group box C (438838)		
Mphi	scfa_17954.1.5	Sox	Sox-C	High mobility group box C (438838)		
Oedu	XP_048762549.1	Sox	Sox-C	High mobility group box C (438838)		
Osin	XP_039654195.1	Sox	Sox-C	High mobility group box C (438838)		
Pean	XP_025110204.1	Sox	Sox-C	High mobility group box C (438838)		
Pcor	DN2429_c2_g1.i2.p1	Sox	Sox-C	High mobility group box C (438838)		
Pcor	DN929_c2_f1.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Pcor	DN2572_c0_g1.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Pcor	DN353_c2_g3.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Pgen	DN788_c0_g1.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Pmar	DN29124_c0_g2.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Pmax	XP_033737425.1	Sox	Sox-C	High mobility group box C (438838)		
Poku	DN71015_c0_g2.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Ppur	DN88859_c0_g1.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Pstr	KAK3610995.1	Sox	Sox-C	High mobility group box C (438838)		
Pvir	s00145g243	Sox	Sox-C	High mobility group box C (438838)		
Pys	XP_021356242.1	Sox	Sox-C	High mobility group box C (438838)		
Rdec	DN52924_c0_g1.i1.p1	Sox	Sox-C	High mobility group box C (438838)		
Rphi	XP_060555827.1	Sox	Sox-C	High mobility group box C (438838)		
Sbro	EVM0006311.1	Sox	Sox-C	High mobility group box C (438838)		
Scon	Ch8-1790	Sox	Sox-C	High mobility group box C (438838)		
Sgio	Sg1000072	Sox	Sox-C	High mobility group box C (438838)		
Sgra	KAJ8306264.1	Sox	Sox-C	High mobility group box C (438838)		
Tgra	KAJ8306266.1	Sox	Sox-C	High mobility group box C (438838)		
Tsqu	DN11669_c1_g1.i2.p1	Sox	Sox-D	High mobility group box C (438839)		
Acal	XP_03524396.1	Sox	Sox-D	High mobility group box C (438839)		
Airc	Contig290.5.1	Sox	Sox-D	High mobility group box C (438839)		
Amar	Ama23921	Sox	Sox-D	High mobility group box C (438839)		

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Apec	DN1990_c0_g1.i10.p1	Sox	Sox-D	High mobility group box (438339)		
Apur	scaffold_393.i10	Sox	Sox-D	High mobility group box (438339)		
Bgia	XP_055899647.1	Sox	Sox-D	High mobility group box (438339)		
Cari	EVM0012405.1	Sox	Sox-D	High mobility group box (438339)		
Cflu	DN124582_c0.g1.i15.p1	Sox	Sox-D	High mobility group box (438339)		
Cgig	XP_011425377.1	Sox	Sox-D	High mobility group box (438339)		
Cph	DN64448_c0.g1.i1.p1	Sox	Sox-D	High mobility group box (438339)		
Hic.asn.2.1656	DN5537_c0.g2.i13.p1	Sox	Sox-D	High mobility group box (438339)		
Csin	Hic.asn.2.1600.2	Sox	Sox-D	High mobility group box (438339)		
Cvir	XP_022302926.1	Sox	Sox-D	High mobility group box (438339)		
Dpol	XP_052213125.1	Sox	Sox-D	High mobility group box (438339)		
Gaeq	XP_041367101.1	Sox	Sox-D	High mobility group box (438339)		
Hbia	M0000014008	Sox	Sox-D	High mobility group box (438339)		
Hruf	XP_046329046.1	Sox	Sox-D	High mobility group box (438339)		
Lorb	DN5537_c0.g2.i13.p1	Sox	Sox-D	High mobility group box (438339)		
Mare	XP_052800695.1	Sox	Sox-D	High mobility group box (438339)		
Mcal	XP_052065962.1	Sox	Sox-D	High mobility group box (438339)		
Mchi	DN3691_c1.g1.i4.p1	Sox	Sox-D	High mobility group box (438339)		
Mcor	CAC5666270.1	Sox	Sox-D	High mobility group box (438339)		
Medu	CAG2197887.1	Sox	Sox-D	High mobility group box (438339)		
Mgal	VD47525.1	Sox	Sox-D	High mobility group box (438339)		
Mgal	VD147529.1	Sox	Sox-D	High mobility group box (438339)		
Mgal	VD147528.1	Sox	Sox-D	High mobility group box (438339)		
Mgal	VD47527.1	Sox	Sox-D	High mobility group box (438339)		
Mgal	VD47526.1	Sox	Sox-D	High mobility group box (438339)		
Mgal	VD147530.1	Sox	Sox-D	High mobility group box (438339)		
Mmar	MMAW000000043.19	Sox	Sox-D	High mobility group box (438339)		
Mmer	XP_053384959.1	Sox	Sox-D	High mobility group box (438339)		
Mmod	DN558_c0.g1.i9.p1	Sox	Sox-D	High mobility group box (438339)		
Mner	g103.147..t2	Sox	Sox-D	High mobility group box (438339)		
Mphi	scaf_42181.1..3	Sox	Sox-D	High mobility group box (438339)		
Obim	XP_042828391.1	Sox	Sox-D	High mobility group box (438339)		
Oedu	XP_048779633.1	Sox	Sox-D	High mobility group box (438339)		
Osin	XP_029644081.1	Sox	Sox-D	High mobility group box (438339)		
Pcan	XP_025088657.1	Sox	Sox-D	High mobility group box (438339)		
Pcor	DN353_c2.g2.i1.p1	Sox	Sox-D	High mobility group box (438339)		
Pcor	DN1386_c1.g1.i1.p1	Sox	Sox-D	High mobility group box (438339)		
Pcor	DN13571_c0.g1.i1.p1	Sox	Sox-D	High mobility group box (438339)		
Pcor	DN1386_c1.g2.i12.p1	Sox	Sox-D	High mobility group box (438339)		
Pgen	DN24654_c0.g1.i2.p1	Sox	Sox-D	High mobility group box (438339)		
Pmar	DN40112_c0.g1.i4.p1	Sox	Sox-D	High mobility group box (438339)		
Pmax	XP_03751614.1	Sox	Sox-D	High mobility group box (438339)		
Poku	DN33319_c18.g4.i1.p1	Sox	Sox-D	High mobility group box (438339)		
Ppur	DN33319_c0.g1.i1.p1	Sox	Sox-D	High mobility group box (438339)		
Pstr	KAK3605548.1	Sox	Sox-D	High mobility group box (438339)		
Pvir	s00219g102	Sox	Sox-D	High mobility group box (438339)		
Pves	XP_021368061.1	Sox	Sox-D	High mobility group box (438339)		
Rdec	DN8993_c0.g1.i1.p1	Sox	Sox-D	High mobility group box (438339)		
Rphi	XP_0606604110.1	Sox	Sox-D	High mobility group box (438339)		
Sbro	EVM0000795.1	Sox	Sox-D	High mobility group box (438339)		
Scor	Ch14..562.1	Sox	Sox-D	High mobility group box (438339)		
Sgra	DN16138_c0.g1.i16.p1	Sox	Sox-D	High mobility group box (438339)		
Tgra	KAJ3298781.1	Sox	Sox-D	High mobility group box (438339)		
Tsqu	DN55031_c0.g1.i1.p1	Sox	Sox-E	High mobility group box E (438340)		
Acal	XP_0005102100.1	Sox	Sox-E	High mobility group box E (438340)		
Airc	Contig52..209	Sox	Sox-E	High mobility group box E (438340)		
Amar	Ama01107	Sox	Sox-E	High mobility group box E (438340)		
Apec	DN4330_c0.g1.i1.p1	Sox	Sox-E	High mobility group box E (438340)		
Apur	scaffold_488.7	Sox	Sox-E	High mobility group box E (438340)		
Bgia	XP_013091187.2	Sox	Sox-E	High mobility group box E (438340)		
Cang	XP_0528989355.1	Sox	Sox-E	High mobility group box E (438340)		
Cari	EVM00058246.1	Sox	Sox-E	High mobility group box E (438340)		
Cflu	DN116407_c5.g2..1..1.p1	Sox	Sox-E	High mobility group box E (438340)		
Cgig	NP_001295801.1	Sox	Sox-E	High mobility group box E (438340)		
Cphi	DN71393_c0.g2.i1.p1	Sox	Sox-E	High mobility group box E (438340)		
Cphi	DN71393_c0.g12.p1	Sox	Sox-E	High mobility group box E (438340)		
Csin	Hic.asn.0..353	Sox	Sox-E	High mobility group box E (438340)		
Cvir	XP_022312895.1	Sox	Sox-E	High mobility group box E (438340)		
Dpol	XP_052264387.1	Sox	Sox-E	High mobility group box E (438340)		

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Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Gaegeg	XP_04136238.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Hbia	M0000012324	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Hbia	M0000012325	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Hruf	XP_046559366.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Mare	XP_052786344.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Mare	XP_052783666.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Mcal	XP_052069336.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Mchi	DN42011_c0_g1.i2.p1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Mmod	CAC5402442.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Mcor	CAG2231021.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Medu	VD182092.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Mgal	VD182090.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Mmar	MMAM0000042410	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Mmer	XP_045213795.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Mmifod	DN78330_c0_g1.i1.p1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Mphi	scf:25414.0.6	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Oedu	XP_056019113.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Pcan	XP_025091262.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Pcor	DN4274_c0_g1.i3.p1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Pcor	DN96098_c0_g1.i1.p1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Pmar	DN30335_c0_g1.i1.p1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Pmax	XP_033139301.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Poku	DN87807_c0_g1.i7.p1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Pour	DN46000_c0_g1.i1.p1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Pstr	KAK3610785.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Pvir	sl.136484674	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Pyes	XP_021248843.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Rphi	XP_060604697.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Sbro	EVMM002110.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Scro	Chr1.75	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Sglo	Sg024297	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Sgra	DN22463_c0_g1.i1.p1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Tgra	KAJ8317914.1	Sox	Sox-E	High mobility group box E (438840)	Sox developmental protein N terminal (463586; partial)	
Tsqu	DN8973_c2_g1.i2.p1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Acal	XP_005107482.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Arc	Contig80.101	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Amar	Ama11616	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Apur	scaffold_546_32	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Bglia	XP_013074628.2	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Cang	XP_052985434.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Cari	EVMM0006823.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Cflu	DN139006_c0_g1.i1.p1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Cggg	XP_011448074.2	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Cipli	DN4414_c0_g1.i1.p1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Csin	Hic_asm_11.549	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Cvir	XP_022319962.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Dpol	XP_052274104.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Gaegeg	XP_041359436.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Hbia	M0000015459	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Hruf	XP_046357912.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Mare	XP_052774544.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Mare	XP_052774361.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Mcal	XP_052061059.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Mcor	CAC5414609.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Medu	CAG2242031.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Mgal	CAG2187650.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Mgal	VD150271.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Mmar	MMAM000000258.10	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Mmer	DN8495_c0_g1.i1.p1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Mmod	g115494_t1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Mphi	scf:611140.13	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Obim	XP_052925684.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Oedu	XP_048764319.2	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	
Pcan	XP_025109598.1	Sox	Sox-F	High mobility group box F (438840)	Sox developmental protein N terminal (463586; partial)	

Tab. S6 continued from previous page

Sp ID	Gene ID	Group	Annotation	Main catalytic domain (Psm-ID)	Additional domains (Psm-ID)	Notes
Pcor	DN11375_c0_g1.i1.p1	Sox	Sox-F	High mobility group box F (438841)		
Pcor	DN29649_c0_g1.i3.p1	Sox	Sox-F	High mobility group box F (438841)		
Pcor	DN5688_c2_g1.i3.p1	Sox	Sox-F	High mobility group box F (438841)		
Pgen	DN144332_c0_g1.i1.p1	Sox	Sox-F	High mobility group box F (438841)		
Pmar	DN24748_c0_g1.i1.p1	Sox	Sox-F	High mobility group box F (438841)		
Pmax	XP_033738287_1	Sox	Sox-F	High mobility group box F (438841)		
Poku	DN14229_c1_g2.i1.p1	Sox	Sox-F	High mobility group box F (438841)		
Pstr	KAK3383243_1	Sox	Sox-F	High mobility group box F (438841)		
Pvir	s00137g284	Sox	Sox-F	High mobility group box F (438841)		
Rdec	XP_021378109_1	Sox	Sox-F	High mobility group box F (438841)		
Rphi	DN53443_c0_g1.i1.p1	Sox	Sox-F	High mobility group box F (438841)		
Sbro	XP_060559438_1	Sox	Sox-F	High mobility group box F (438841)		
Scon	EVM0000861_1	Sox	Sox-F	High mobility group box F (438841)		
Sglo	Ch8-897	Sox	Sox-F	High mobility group box F (438841)		
Sgra	Sglo05442	Sox	Sox-F	High mobility group box F (438841)		
Airc	Contig1525_38	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Amar	Ama26724	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Apec	DN93182_c0_g1.i1.p1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Apur	scaffid768_3	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Cang	XP_0522703370_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Cari	EVM0018164_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Cggg	XP_011415859_3	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Cpli	DN80002_c0_g1.i1.p1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Csin	Hic_asn_15_1471	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Cvir	XP_3022338738_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Dpol	XP_052226448_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Gaeq	XP_041370217_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Gaeq	XP_041369137_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Hbia	M00000001184	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Hrfu	XP_046358320_2	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Mcal	XP_052099860_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Mcor	CA5406014_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Medu	CAG2257203_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Mgal	VD130324_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Mgal	VD130323_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Mmar	MMAM0000015662	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Mmer	XP_053407277_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Mner	g125_224_t1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Mphi	scaf12010_0_4	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Mphi	scaf59202_0_9	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Obim	XP_028332677_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Oedu	XP_056006679_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Osin	XP_0363668794_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Pcor	DN1864456_c0_g1.i1.p1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Pmar	DN40950_c1_g1.i2.p1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Pmax	XP_032756818_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Poku	DN16718_c0_g1.i6.p1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Ppur	DN7268_c0_g1.i1.p1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Pstr	KAK3382760_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Pvir	s00451g108	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Pyes	XP_021340986_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Rdec	DN22482_c4_g1.i1.p1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Rphi	XP_060578490_1	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Scon	Ch15_1899	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Sglo	Sglo010047	Sox	Sox-H	High mobility group box Sox-30 (438842)		
Tsqu	DN874.c7_g1.i1.p1	Sox	Sox-H	High mobility group box Sox-30 (438842)		

Supplementary Table S7 – Proportions of missing data in both bivalve species and DSFGs. Bivalve species represented by transcriptomic data are highlighted with an asterisk (*).

Species		Genes	
Species	% missing data (out of 33 DSFGs)	Group	% missing data (out of 43 species)
<i>A. irradians concentricus</i>	0.000000	<i>Dmrt-1L</i>	48.837209
<i>A. marissinica</i>	21.212121	<i>Dmrt-3</i>	30.232558
<i>A. pectinata*</i>	48.484848	<i>Dmrt-2</i>	55.813953
<i>A. purpuratus</i>	6.060606	<i>Dmrt-4/5</i>	6.976744
<i>C. angulata</i>	6.060606	<i>Fox-A</i>	13.953488
<i>C. ariakensis</i>	3.030303	<i>Fox-B</i>	30.232558
<i>C. fluminea*</i>	42.424242	<i>Fox-C</i>	23.255814
<i>C. gigas</i>	6.060606	<i>Fox-D</i>	20.930233
<i>C. plicata*</i>	21.212121	<i>Fox-E</i>	30.232558
<i>C. sinensis</i>	21.212121	<i>Fox-F</i>	18.604651
<i>C. virginica</i>	3.030303	<i>Fox-G</i>	16.279070
<i>D. polymorpha</i>	9.090909	<i>Fox-H</i>	41.860465
<i>H. bialata</i>	9.090909	<i>Fox-J1</i>	0.000000
<i>L. orbiculatus*</i>	63.636364	<i>Fox-J2/3</i>	9.302326
<i>M. arenaria</i>	21.212121	<i>Fox-L1</i>	18.604651
<i>M. californianus</i>	9.090909	<i>Fox-L2</i>	13.953488
<i>M. chinensis*</i>	57.575758	<i>Fox-N1/4</i>	6.976744
<i>M. coruscus</i>	0.000000	<i>Fox-N2/3</i>	6.976744
<i>M. edulis</i>	3.030303	<i>Fox-O</i>	13.953488
<i>M. galloprovincialis</i>	6.060606	<i>Fox-P</i>	9.302326
<i>M. margaritifera</i>	6.060606	<i>Fox-Q2</i>	30.232558
<i>M. mercenaria</i>	3.030303	<i>Fox-OG13/NA</i>	32.558140
<i>M. modiolus*</i>	36.363636	<i>Fox-OG15/NA</i>	34.883721
<i>M. nervosa</i>	27.272727	<i>Fox-OG16/NA</i>	30.232558
<i>M. philippinarum</i>	9.090909	<i>Fox-OG2/NA</i>	16.279070
<i>O. edulis</i>	6.060606	<i>Fox-OG28/NA</i>	39.534884
<i>P. coreanum*</i>	18.181818	<i>Fox-OG39/NA</i>	37.209302
<i>P. generosa*</i>	54.545455	<i>Sox-B1/2</i>	0.000000
<i>P. margaritifera*</i>	21.212121	<i>Sox-C</i>	0.000000
<i>P. maximus</i>	0.000000	<i>Sox-D</i>	4.651163
<i>P. okutanii*</i>	54.545455	<i>Sox-E</i>	9.302326
<i>P. purpuratus*</i>	54.545455	<i>Sox-F</i>	13.953488
<i>P. streckersoni</i>	6.060606	<i>Sox-H</i>	20.930233
<i>P. viridis</i>	3.030303		
<i>P. yessoensis</i>	3.030303		
<i>R. decussatus*</i>	51.515152		
<i>R. philippinarum</i>	3.030303		
<i>S. broughtonii</i>	12.121212		
<i>S. constricta</i>	12.121212		
<i>S. glomerata</i>	9.090909		
<i>S. grandis*</i>	54.545455		
<i>T. granosa</i>	42.424242		
<i>T. squamosa*</i>	48.484848		

Supplementary Table S8 – Complete set of DSFGs in mammals. For each gene, the species ID (Sp. ID) as in **Tab. S4**, the accession number (Gene ID), and the Possvm-based annotation are indicated.

Sp. ID	Gene ID	Group	Annotation	Sp. ID	Gene ID	Group	Annotation
Hamp	XP_057601513.1	Dmrt	Dmrt-B1	Cimi	XP_017353003.1	Fox	Fox-C1
Hsap	NP_149056.1	Dmrt	Dmrt-B1	Oana	XP_028912285.1	Fox	Fox-C1
Oafe	XP_007944154.1	Dmrt	Dmrt-B1	Dnov	XP_058141148.1	Fox	Fox-C1
Amel	XP_034508822.1	Dmrt	Dmrt-B1	Bbub	XP_025120521.1	Fox	Fox-C1
Bbub	XP_025144486.2	Dmrt	Dmrt-B1	Drot	XP_053776986.1	Fox	Fox-C1
Casi	XP_006839931.1	Dmrt	Dmrt-B1	Rfer	XP_032971521.1	Fox	Fox-C1
Pgig	XP_039739094.1	Dmrt	Dmrt-B1	Mdom	XP_007488114.2	Fox	Fox-C1
Cimi	XP_017359902.1	Dmrt	Dmrt-B1	Casi	XP_006870626.1	Fox	Fox-C1
Mmus	XP_036020151.1	Dmrt	Dmrt-B1	Pafr	XP_047652737.1	Fox	Fox-C1
Bmus	XP_036712433.1	Dmrt	Dmrt-B1	Bmus	XP_036726614.1	Fox	Fox-C1
Oana	XP_039770576.1	Dmrt	Dmrt-B1	Hamp	XP_057555406.1	Fox	Fox-C1
Scar	XP_047395630.1	Dmrt	Dmrt-B1	Lcat	XP_045407522.1	Fox	Fox-C1
Mjav	XP_036877684.1	Dmrt	Dmrt-B1	Opri	XP_058524289.1	Fox	Fox-C1
Ptig	XP_042853169.1	Dmrt	Dmrt-B1	Cdid	XP_037699408.1	Fox	Fox-C1
Lcat	XP_045402503.1	Dmrt	Dmrt-B1	Hsap	NP_001444.2	Fox	Fox-C1
Emax	XP_049731502.1	Dmrt	Dmrt-B1	Mjav	XP_036867643.1	Fox	Fox-C1
Ttru	XP_004325875.3	Dmrt	Dmrt-B1	Scar	XP_047413053.1	Fox	Fox-C1
Mang	XP_0475755047.1	Dmrt	Dmrt-B1	Cdro	XP_031291159.1	Fox	Fox-C1
Pafr	XP_047647463.1	Dmrt	Dmrt-B1	Mmus	NP_032618.2	Fox	Fox-C1
Opri	XP_004588846.2	Dmrt	Dmrt-B1	Emax	XP_049738278.1	Fox	Fox-C1
Cpor	XP_003464114.1	Dmrt	Dmrt-B1	Oafe	XP_007933566.1	Fox	Fox-C1
Dnov	XP_058159531.1	Dmrt	Dmrt-B1	Mang	XP_045722432.1	Fox	Fox-C1
Cdid	XP_037665497.1	Dmrt	Dmrt-B1	Ttru	XP_019800382.2	Fox	Fox-C1
Equa	XP_046516609.1	Dmrt	Dmrt-B1	Oana	XP_028909462.1	Fox	Fox-C1
Mdom	XP_056671081.1	Dmrt	Dmrt-B1	Equa	XP_046496475.1	Fox	Fox-C1
Csim	XP_014649261.1	Dmrt	Dmrt-B1	Pgig	XP_039725399.1	Fox	Fox-C1
Cdro	XP_031321432.1	Dmrt	Dmrt-B1	Shar	XP_023352145.2	Fox	Fox-C1
Rfer	XP_032970385.1	Dmrt	Dmrt-B1	Ptig	XP_042841201.1	Fox	Fox-C1
Clup	XP_038393419.1	Dmrt	Dmrt-B1	Amel	XP_034516093.1	Fox	Fox-C1
Tman	XP_012409672.2	Dmrt	Dmrt-B1	Clup	XP_038439885.1	Fox	Fox-C1
Ggal	NP_001232910.1	Dmrt	Dmrt-B1	Scar	XP_047385678.1	Fox	Fox-C2
Oafe	XP_007954175.1	Dmrt	Dmrt-B1	Mjav	XP_036849635.1	Fox	Fox-C2
Hsap	NP_870987.2	Dmrt	Dmrt-B1	Cdro	XP_031292483.1	Fox	Fox-C2
Dnov	XP_004459445.2	Dmrt	Dmrt-B1	Ttru	XP_033700545.1	Fox	Fox-C2
Shar	XP_003761708.1	Dmrt	Dmrt-B1	Tman	XP_023588281.1	Fox	Fox-C2
Cpor	XP_005004778.1	Dmrt	Dmrt-B1	Csim	XP_004437101.1	Fox	Fox-C2
Ttru	XP_019790310.1	Dmrt	Dmrt-B1	Pafr	XP_047648754.1	Fox	Fox-C2
Ggal	XP_003643035.3	Dmrt	Dmrt-B1	Mjav	XP_036849634.1	Fox	Fox-C2
Mdom	XP_001374295.1	Dmrt	Dmrt-B1	Rfer	XP_032982997.1	Fox	Fox-C2
Oana	XP_028911116.1	Dmrt	Dmrt-B1	Pgig	XP_039744659.1	Fox	Fox-C2
Tman	XP_004373589.1	Dmrt	Dmrt-B1	Ggal	NP_001382975.1	Fox	Fox-C2
Equa	XP_046519727.1	Dmrt	Dmrt-B1	Clup	XP_038394038.1	Fox	Fox-C2
Bmus	XP_036712720.1	Dmrt	Dmrt-B1	Opri	XP_058530600.1	Fox	Fox-C2
Cdid	XP_037653375.1	Dmrt	Dmrt-B1	Amel	XP_034495216.1	Fox	Fox-C2
Opri	XP_058513365.1	Dmrt	Dmrt-B1	Bmus	XP_036688981.1	Fox	Fox-C2
Lcat	XP_045418965.1	Dmrt	Dmrt-B1	Lcat	XP_045389072.1	Fox	Fox-C2
Cimi	XP_017373822.1	Dmrt	Dmrt-B1	Cpor	XP_005008627.1	Fox	Fox-C2
Rfer	XP_032979206.1	Dmrt	Dmrt-B1	Mmus	NP_038547.2	Fox	Fox-C2
Pgig	XP_039723855.1	Dmrt	Dmrt-B1	Oafe	XP_007937406.1	Fox	Fox-C2
Ptig	XP_042820943.1	Dmrt	Dmrt-B1	Cdid	XP_037671788.1	Fox	Fox-C2
Hamp	XP_057579943.1	Dmrt	Dmrt-B1	Bbub	XP_006080917.3	Fox	Fox-C2
Pafra	XP_047623676.1	Dmrt	Dmrt-B1	Emax	XP_049720645.1	Fox	Fox-C2
Cdro	XP_031305655.1	Dmrt	Dmrt-B1	Rfer	XP_032982975.1	Fox	Fox-C2
Mmus	NP_665830.1	Dmrt	Dmrt-B1	Mmus	NP_038547.2	Fox	Fox-C2
Csim	XP_014650903.1	Dmrt	Dmrt-B1	Ptig	XP_042825379.1	Fox	Fox-C2
Casi	XP_006863797.1	Dmrt	Dmrt-B1	Mang	XP_045746247.1	Fox	Fox-C2
Emax	XP_049752756.1	Dmrt	Dmrt-B1	Cimi	XP_017399160.1	Fox	Fox-C2
Mjav	XP_017529816.2	Dmrt	Dmrt-B1	Drot	XP_053772216.1	Fox	Fox-C2
Amel	XP_034503257.1	Dmrt	Dmrt-B1	Hsap	NP_005242.1	Fox	Fox-C2
Clup	XP_038383091.1	Dmrt	Dmrt-B1	Hamp	XP_057568935.1	Fox	Fox-C2
Mang	XP_045738142.1	Dmrt	Dmrt-B1	Mdom	XP_001365891.1	Fox	Fox-C2
Bbub	XP_044795850.1	Dmrt	Dmrt-B1	Dnov	XP_004450287.3	Fox	Fox-C2
Drot	XP_045047034.2	Dmrt	Dmrt-B1	Casi	XP_006860181.1	Fox	Fox-C2
Scar	XP_047380961.1	Dmrt	Dmrt-B1	Oana	XP_039769457.1	Fox	Fox-C2
Cdro	XP_031312888.1	Dmrt	Dmrt-C2	Shar	XP_031806033.1	Fox	Fox-C2
Clup	XP_038381591.1	Dmrt	Dmrt-C2	Oana	XP_028921129.1	Fox	Hnf-3g/Fox-A3
Csim	XP_004440164.1	Dmrt	Dmrt-C2	Hsap	NP_004488.2	Fox	Hnf-3g/Fox-A3
Casi	XP_006871451.1	Dmrt	Dmrt-C2	Pafra	XP_047645608.1	Fox	Hnf-3g/Fox-A3
Mang	XP_045746349.1	Dmrt	Dmrt-C2	Ptig	XP_042825317.1	Fox	Hnf-3g/Fox-A3
Bmus	XP_036690564.1	Dmrt	Dmrt-C2	Mjav	XP_017507570.1	Fox	Hnf-3g/Fox-A3
Lcat	XP_045387563.1	Dmrt	Dmrt-C2	Mmus	NP_032286.1	Fox	Hnf-3g/Fox-A3
Hamp	XP_057568752.1	Dmrt	Dmrt-C2	Casi	XP_006871384.1	Fox	Hnf-3g/Fox-A3
Cdid	XP_037676589.1	Dmrt	Dmrt-C2	Shar	XP_031817519.1	Fox	Hnf-3g/Fox-A3
Pafra	XP_047644735.1	Dmrt	Dmrt-C2	Oafe	XP_007941248.1	Fox	Hnf-3g/Fox-A3
Mmus	XP_017167796.1	Dmrt	Dmrt-C2	Ttru	XP_033701564.1	Fox	Hnf-3g/Fox-A3
Equa	XP_046507274.1	Dmrt	-	Mang	XP_045746361.1	Fox	Hnf-3g/Fox-A3
Oafe	XP_007941145.1	Dmrt	Dmrt-C2	Clup	XP_038384456.1	Fox	Hnf-3g/Fox-A3
Tman	XP_004388803.1	Dmrt	Dmrt-C2	Opri	XP_004597990.1	Fox	Hnf-3g/Fox-A3
Drot	XP_024433509.1	Dmrt	Dmrt-C2	Hamp	XP_057566925.1	Fox	Hnf-3g/Fox-A3
Oana	XP_007664048.1	Dmrt	Dmrt-C2	Rfer	XP_032982943.1	Fox	Hnf-3g/Fox-A3
Ttru	XP_033699919.1	Dmrt	Dmrt-C2	Dnov	XP_004481594.1	Fox	Hnf-3g/Fox-A3
Amel	XP_034494561.1	Dmrt	Dmrt-C2	Drot	XP_049757192.1	Fox	Hnf-3g/Fox-A3
Opri	XP_058531141.1	Dmrt	Dmrt-C2	Pgig	XP_039722184.1	Fox	Hnf-3g/Fox-A3
Pgig	XP_039731326.1	Dmrt	-	Tman	XP_004381648.1	Fox	Hnf-3g/Fox-A3
Lcat	XP_045393814.1	Dmrt	-	Mdom	XP_001364242.1	Fox	Hnf-3g/Fox-A3
Lcat	XP_045393815.1	Dmrt	-	Bmus	XP_036689576.1	Fox	Hnf-3g/Fox-A3
Lcat	XP_045393818.1	Dmrt	-	Equa	XP_046539522.1	Fox	Hnf-3g/Fox-A3
Scar	XP_047383853.1	Dmrt	Dmrt-C2	Bbub	XP_025125137.1	Fox	Hnf-3g/Fox-A3
Pgig	XP_039705918.1	Dmrt	Dmrt-C2	Csim	XP_004440228.1	Fox	Hnf-3g/Fox-A3
Hsap	XP_016882612.1	Dmrt	Dmrt-C2	Cimi	XP_017358010.1	Fox	Hnf-3g/Fox-A3
Equa	XP_046540061.1	Dmrt	-	Scar	XP_047384662.1	Fox	Hnf-3g/Fox-A3
Mjav	XP_036869866.1	Dmrt	Dmrt-C2	Cdid	XP_037675543.1	Fox	Hnf-3g/Fox-A3
Rfer	XP_032984641.1	Dmrt	Dmrt-C2	Lcat	XP_045387339.1	Fox	Hnf-3g/Fox-A3
Shar	XP_031819119.1	Dmrt	Dmrt-C2	Cdro	XP_010985920.2	Fox	Hnf-3g/Fox-A3
Emax	XP_049757407.1	Dmrt	Dmrt-C2	Cpor	XP_003464639.1	Fox	Hnf-3g/Fox-A3
Lcat	XP_045393817.1	Dmrt	-	Amel	XP_002928546.3	Fox	Hnf-3g/Fox-A3
Ptig	XP_042825393.1	Dmrt	Dmrt-C2	Drot	XP_024425331.1	Fox	Hnf-3g/Fox-A3

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Sp. ID	Gene ID	Group	Annotation	Sp. ID	Gene ID	Group	Annotation
Oana	XP_039770060.1	Fox	Fox-G1	Rfer	XP_032981124.1	Fox	Fox-N2
Pgig	XP_039726380.1	Fox	Fox-G1	Emax	XP_049728576.1	Fox	Fox-R2
Dnov	XP_058149613.1	Fox	Fox-G1	Tman	XP_004391284.1	Fox	Fox-R2
Mdom	XP_001364896.1	Fox	Fox-G1	Shar	XP_031815965.1	Fox	Fox-R1
Mang	XP_045757915.1	Fox	Fox-G1	Drot	XP_024426718.1	Fox	Fox-R1
Mjav	XP_017501197.2	Fox	Fox-G1	Mdom	XP_001380644.1	Fox	Fox-R1
Emax	XP_049754324.1	Fox	Fox-G1	Oana	XP_028931756.1	Fox	Fox-R1
Ttru	XP_019779874.1	Fox	Fox-G1	Casi	XP_006834052.1	Fox	Fox-R1
Cdid	XP_037690644.1	Fox	Fox-G1	Equa	XP_046540933.1	Fox	Fox-R1
Hsap	NP_005240.3	Fox	Fox-G1	Mmus	NP_001028641.1	Fox	Fox-R1
Shar	XP_031808132.1	Fox	Fox-G1	Cpor	XP_003472699.1	Fox	Fox-R1
Ptig	XP_042845427.1	Fox	Fox-G1	Bbub	XP_044785576.1	Fox	Fox-R1
Scar	XP_047396708.1	Fox	Fox-G1	Pgig	XP_039722531.1	Fox	Fox-R1
Clup	XP_038400306.1	Fox	Fox-G1	Rfer	XP_032976774.1	Fox	Fox-R1
Hamp	XP_057580105.1	Fox	Fox-G1	Ptig	XP_042814206.1	Fox	Fox-R1
Casi	XP_006835434.1	Fox	Fox-G1	Bmus	XP_036721689.1	Fox	Fox-R1
Opri	XP_058521348.1	Fox	Fox-G1	Bmus	XP_036715648.1	Fox	Fox-R1
Bmus	XP_036700190.1	Fox	Fox-G1	Cdro	XP_031299399.1	Fox	Fox-R1
Bbub	XP_025126917.1	Fox	Fox-G1	Pafr	XP_047609433.1	Fox	Fox-R1
Mmus	NP_001153584.1	Fox	Fox-G1	Hsap	XP_016873064.1	Fox	Fox-R1
Ggal	NP_989659.2	Fox	Fox-O1	Cdid	XP_037695751.1	Fox	Fox-R1
Cimi	XP_017397972.1	Fox	Fox-O1	Csim	XP_004427297.1	Fox	Fox-R1
Rfer	XP_032960350.1	Fox	Fox-O1	Emax	XP_049713316.1	Fox	Fox-R1
Casi	XP_006873780.1	Fox	Fox-O1	Mjav	XP_017505694.2	Fox	Fox-R1
Pgig	XP_039695386.1	Fox	Fox-O1	Hsap	NP_859072.1	Fox	Fox-R1
Dnov	XP_058132895.1	Fox	Fox-O1	Lcat	XP_045412286.1	Fox	Fox-R1
Ttru	XP_019792245.1	Fox	Fox-O1	Dnov	XP_023443955.1	Fox	Fox-R1
Oana	XP_001512968.3	Fox	Fox-O1	Amel	XP_034521812.1	Fox	Fox-R1
Hsap	NP_002006.2	Fox	Fox-O1	Hamp	XP_057603312.1	Fox	Fox-R1
Mang	XP_045725787.1	Fox	Fox-O1	Tman	XP_004385753.1	Fox	Fox-R1
Bmus	XP_036687548.1	Fox	Fox-O1	Oafe	XP_007934779.1	Fox	Fox-R1
Amel	XP_034519677.1	Fox	Fox-O1	Clup	XP_038391436.1	Fox	Fox-R1
Scar	XP_047409558.1	Fox	Fox-O1	Scar	XP_047373976.1	Fox	Fox-R1
Mmus	NP_062713.2	Fox	Fox-O1	Opri	XP_058519782.1	Fox	Fox-R1
Opri	XP_012782130.2	Fox	Fox-O1	Ttru	XP_033717853.1	Fox	Fox-R1
Cdid	XP_037656524.1	Fox	Fox-O1	Oafe	XP_007947037.1	Fox	Fox-R1
Mjav	XP_036867814.1	Fox	Fox-O1	Mang	XP_045748841.1	Fox	Fox-R1
Cdro	XP_031321792.1	Fox	Fox-O1	Cimi	XP_017378961.1	Fox	Fox-R1
Lcat	XP_045423168.1	Fox	Fox-O1	Lcat	XP_045383006.1	Fox	Fox-K2
Clup	XP_038429080.1	Fox	Fox-O1	Casi	XP_006869677.1	Fox	Fox-K2
Cpor	XP_023416198.1	Fox	Fox-O1	Mmus	XP_011247520.1	Fox	Fox-K2
Drot	XP_024424320.2	Fox	Fox-O1	Drot	XP_0244209582.3	Fox	Fox-K2
Mdom	XP_001368312.2	Fox	Fox-O1	Cpor	XP_013002484.1	Fox	Fox-K2
Shar	XP_003764601.3	Fox	Fox-O1	Tman	XP_023584943.1	Fox	Fox-K2
Csim	XP_004443237.1	Fox	Fox-O1	Scar	XP_047400835.1	Fox	Fox-K2
Tman	XP_023597991.1	Fox	Fox-O1	Opri	XP_058532091.1	Fox	Fox-K2
Emax	XP_049709381.1	Fox	Fox-O1	Oafe	XP_007957978.1	Fox	Fox-K2
Pafr	XP_047611907.1	Fox	Fox-O1	Hsap	NP_004505.2	Fox	Fox-K2
Bbub	XP_006065837.2	Fox	Fox-O1	Emax	XP_049715130.1	Fox	Fox-K2
Ptig	XP_007097795.2	Fox	Fox-O1	Cimi	XP_017380207.1	Fox	Fox-K2
Hamp	XP_057563333.1	Fox	Fox-O1	Hamp	XP_057574401.1	Fox	-
Oafe	XP_007943433.1	Fox	Fox-O1	Hamp	XP_057571581.1	Fox	-
Mang	XP_045725786.1	Fox	Fox-O1	Lcat	XP_045397194.1	Fox	Fox-K1
Ggal	XP_015134143.3	Fox	Fox-O4	Bmus	XP_036681953.1	Fox	Fox-K1
Casi	XP_006868595.1	Fox	Fox-O4	Pgig	XP_039734750.1	Fox	Fox-K1
Mdom	XP_056665439.1	Fox	Fox-O4	Equa	XP_046521295.1	Fox	Fox-K1
Csim	XP_004439923.1	Fox	Fox-O4	Hsap	NP_001032242.1	Fox	Fox-K1
Cdid	XP_037677588.1	Fox	Fox-O4	Emax	XP_049758808.1	Fox	Fox-K1
Hsap	NP_005929.2	Fox	Fox-O4	Oana	XP_028905334.1	Fox	Fox-K1
Emax	XP_049727631.1	Fox	Fox-O4	Opri	XP_012786071.2	Fox	Fox-K1
Ptig	XP_007099158.2	Fox	Fox-O4	Cimi	XP_017389267.1	Fox	Fox-K1
Drot	XP_053773393.1	Fox	Fox-O4	Drot	XP_053782069.1	Fox	Fox-K1
Bmus	XP_036696517.1	Fox	Fox-O4	Cpor	XP_023420906.1	Fox	Fox-K1
Oafe	XP_007957073.1	Fox	Fox-O4	Dnov	XP_004454003.2	Fox	Fox-K1
Cimi	XP_017372253.1	Fox	Fox-O4	Scar	XP_047389516.1	Fox	Fox-K1
Mjav	XP_017525885.1	Fox	Fox-O4	Csim	XP_004440941.1	Fox	Fox-K1
Rfer	XP_032969557.1	Fox	Fox-O4	Mmus	NP_951031.2	Fox	Fox-K1
Pafr	XP_047621021.1	Fox	Fox-O4	Shar	XP_031796898.1	Fox	Fox-K1
Mmus	NP_061259.1	Fox	Fox-O4	Hamp	XP_057604533.1	Fox	Fox-K1
Mang	XP_045735285.1	Fox	Fox-O4	Mang	XP_045731846.1	Fox	Fox-K1
Pgig	XP_039697652.1	Fox	Fox-O4	Casi	XP_006859933.1	Fox	Fox-K1
Scar	XP_047393133.1	Fox	Fox-O4	Clup	XP_038395513.1	Fox	Fox-K1
Oana	XP_039768253.1	Fox	Fox-O4	Cdid	XP_037669880.1	Fox	Fox-K1
Dnov	XP_058146903.1	Fox	Fox-O4	Pafr	XP_047635407.1	Fox	Fox-K1
Cdid	XP_037678715.1	Fox	Fox-O4	Tman	XP_004380896.3	Fox	Fox-K1
Amel	XP_002929104.1	Fox	Fox-O4	Amel	XP_011219634.2	Fox	Fox-K1
Tman	XP_023590438.1	Fox	Fox-O4	Mjav	XP_036866572.1	Fox	Fox-K1
Clup	XP_038443581.1	Fox	Fox-O4	Ttru	XP_019802935.2	Fox	Fox-K1
Cpor	XP_013009181.1	Fox	Fox-O4	Cdro	XP_031327189.1	Fox	Fox-K1
Lcat	XP_045394627.1	Fox	Fox-O4	Mdom	XP_056662440.1	Fox	Fox-K1
Hamp	XP_057575075.1	Fox	Fox-O4	Ptig	XP_04282249.1	Fox	Fox-K1
Bbub	XP_006076433.1	Fox	Fox-O4	Rfer	XP_032951743.1	Fox	Fox-K1
Ttru	XP_033705152.1	Fox	Fox-O4	Oafe	XP_007941047.1	Fox	Fox-K1
Equa	XP_046529856.1	Fox	Fox-O4	Ggal	XP_015149844.1	Fox	Fox-K1
Opri	XP_004595290.1	Fox	Fox-O4	Bbub	XP_025131152.1	Fox	Fox-K1
Drot	XP_053773348.1	Fox	Fox-O4	Hsap	NP_001400854.1	Fox	Fox-M1
Cdro	XP_010977055.2	Fox	Fox-O4	Lcat	XP_045410542.1	Fox	-
Pgig	XP_039724773.1	Fox	Fox-J3	Ggal	XP_046760565.1	Fox	-
Mdom	XP_007492986.2	Fox	Fox-J3	Oana	XP_028912611.1	Fox	Fox-K2
Lcat	XP_045401678.1	Fox	Fox-J3	Clup	XP_038402133.1	Fox	-
Shar	XP_031818195.1	Fox	Fox-J3	Amel	XP_019662714.1	Fox	-
Equa	XP_046517148.1	Fox	Fox-J3	Mang	XP_045744166.1	Fox	-
Mjav	XP_036858181.1	Fox	Fox-J3	Equa	XP_046532308.1	Fox	-
Oafe	XP_007953745.1	Fox	Fox-J3	Ptig	XP_042846258.1	Fox	-
Mmus	NP_766287.1	Fox	Fox-J3	Oafe	XP_007942676.1	Fox	-
Drot	XP_024410422.2	Fox	Fox-J3	Casi	XP_006839645.1	Fox	-
Tman	XP_023582916.1	Fox	Fox-J3	Mmus	XP_011246143.1	Fox	Fox-R2
Dnov	XP_004477472.1	Fox	Fox-J3	Opri	XP_058514738.1	Fox	-
Hsap	NP_005270689.1	Fox	Fox-J3	Scar	XP_047391089.1	Fox	-
Csim	XP_004443185.1	Fox	Fox-J3	Ggal	XP_015137672.3	Fox	-
Scar	XP_047408133.1	Fox	Fox-J3	Ggal	XP_015141419.2	Fox	-
Cimi	XP_017397267.1	Fox	Fox-J3	Ggal	XP_015144452.3	Fox	-

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Sp. ID	Gene ID	Group	Annotation		Sp. ID	Gene ID	Group	Annotation
Hamp	XP_057574707.1	Fox	Fox-P3		Oana	XP_039766108.1	Sox	Sox-30
Lcat	XP_045393738.1	Fox	Fox-P3		Cimi	XP_017372578.1	Sox	Sox-30
Pafn	XP_047620716.1	Fox	Fox-P3		Oafe	XP_007948121.1	Sox	Sox-30
Amel	XP_011221377.1	Fox	Fox-P3		Cdro	XP_031305277.1	Sox	Sox-30
Dnov	XP_023446473.1	Fox	Fox-P3		Emax	XP_049730174.1	Sox	Sox-30
Ptig	XP_007096061.2	Fox	Fox-P3		Hamp	XP_057558120.1	Sox	Sox-30
Opri	XP_058514303.1	Fox	Fox-P3		Lcat	XP_045408644.1	Sox	Sox-30
Clup	NP_001161933.1	Fox	Fox-P3		Shar	XP_031809461.1	Sox	Sox-30
Mang	XP_045729506.1	Fox	Fox-P3		Bbub	XP_006058618.4	Sox	Sox-30
Cdid	XP_037677000.1	Fox	Fox-P3		Drot	XP_024432555.2	Sox	Sox-30
Hsap	NP_054728.2	Fox	Fox-P3		Clup	XP_038390830.1	Sox	Sox-30
Cdro	XP_031301084.1	Fox	Fox-P3		Cpor	XP_003473352.1	Sox	Sox-30
Tman	XP_004376890.1	Fox	Fox-P3		Ttru	XP_019780387.1	Sox	Sox-30
Oana	XP_028909446.1	Fox	Fox-F2		Opri	XP_004587550.2	Sox	Sox-30
Oana	XP_0289112286.1	Fox	Fox-F2		Mdom	XP_001379720.1	Sox	Sox-30
Opri	XP_012786079.2	Fox	Fox-L3		Equa	XP_046521338.1	Sox	Sox-30
Mdom	XP_007498348.2	Fox	Fox-L3		Csim	XP_004428646.1	Sox	Sox-30
Drot	XP_045040885.2	Fox	Fox-L3		Cdid	XP_037655441.1	Sox	Sox-30
Rfer	XP_032952385.1	Fox	Fox-L3		Ggal	XP_414564.1	Sox	Sox-30
Scar	XP_047390130.1	Fox	Fox-L3		Dnov	XP_058162315.1	Sox	Sox-30
Oana	XP_001511921.2	Fox	Fox-L3		Scar	XP_047412799.1	Sox	Sox-30
Ttru	XP_004328913.2	Fox	Fox-L3		Mmus	NP_775560.1	Sox	Sox-30
Dnov	XP_058141662.1	Fox	Fox-L3		Amel	XP_019650665.2	Sox	Sox-30
Mmus	NP_001182057.1	Fox	Fox-L3		Pgig	XP_039732492.1	Sox	Sox-30
Hsap	NP_001361767.1	Fox	Fox-L3		Mjav	XP_036881068.1	Sox	Sox-30
Emax	XP_049761077.1	Fox	Fox-L3		Mang	XP_045743539.1	Sox	Sox-30
Mang	XP_045728168.1	Fox	Fox-L3		Rfer	XP_032952708.1	Sox	Sox-30
Mjav	XP_036864912.1	Fox	Fox-L3		Tman	XP_004371314.1	Sox	Sox-30
Shar	XP_012398242.1	Fox	Fox-L3		Hsap	NP_848511.1	Sox	Sox-30
Cimi	XP_017363768.1	Fox	Fox-L3		Casi	XP_006863994.1	Sox	Sox-30
Hamp	XP_057605555.1	Fox	Fox-L3		Amel	XP_034508849.1	Sox	Sox-30
Pafn	XP_047635063.1	Fox	Fox-L3		Scar	XP_047389638.1	Sox	Sox-8
Tman	XP_004386032.1	Fox	Fox-L3		Bmusp	XP_036681892.1	Sox	Sox-8
Csim	XP_004440991.1	Fox	Fox-L3		Pafn	XP_047635134.1	Sox	Sox-8
Equa	XP_046523569.1	Fox	Fox-L3		Drot	XP_024409185.2	Sox	Sox-8
Lcat	XP_045398087.1	Fox	Fox-L3		Mjav	XP_017502547.2	Sox	Sox-8
Cpor	XP_03469985.1	Fox	Fox-L3		Clup	XP_038394815.1	Sox	Sox-8
Bmusp	XP_036680082.1	Fox	Fox-L3		Rfer	XP_032957935.1	Sox	Sox-8
Bbub	XP_044792209.1	Fox	Fox-L3		Equa	XP_046523802.1	Sox	Sox-8
Clup	XP_038394791.1	Fox	Fox-L3		Amel	XP_034525488.1	Sox	Sox-8
Casi	XP_006859950.1	Fox	Fox-L3		Cimi	XP_017366444.1	Sox	Sox-8
Cdid	XP_037669698.1	Fox	Fox-L3		Csim	XP_004438275.1	Sox	Sox-8
Ptig	XP_042828254.1	Fox	Fox-L3		Bbub	XP_025130606.3	Sox	Sox-8
Ggal	XP_425229.3	Fox	Fox-L3		Lcat	XP_045397007.1	Sox	Sox-8
Cdro	XP_031327057.1	Fox	Fox-L3		Hamp	XP_057551434.1	Sox	Sox-8
Amel	XP_034525410.1	Fox	Fox-L3		Pgig	XP_039740781.1	Sox	Sox-8
Oafe	XP_007941358.1	Fox	Fox-L3		Hsap	NP_055402.2	Sox	Sox-8
Cdro	XP_031326970.1	Fox	Fox-P1		Mang	XP_045730580.1	Sox	Sox-8
Oafe	XP_007944266.1	Fox	Fox-P1		Cdro	XP_010989436.2	Sox	Sox-8
Ttru	XP_033721167.1	Fox	Fox-P1		Ptig	XP_042827523.1	Sox	Sox-8
Amel	XP_011217264.1	Fox	Fox-P1		Ttru	XP_033696379.1	Sox	Sox-8
Mdom	XP_007500148.1	Fox	Fox-P1		Bmusp	XP_036722097.1	Sox	-
Opri	XP_058534843.1	Fox	Fox-P1		Ttru	XP_019791993.1	Sox	-
Shar	XP_031801298.1	Fox	Fox-P1		Oana	XP_028908350.1	Sox	Sox-4
Pafn	XP_047635064.1	Fox	Fox-P1		Opri	XP_058523457.1	Sox	Sox-4
Mjav	XP_036875087.1	Fox	Fox-P1		Ggal	NP_99815.2	Sox	Sox-4
Hsap	NP_001231739.1	Fox	Fox-P1		Casi	XP_006860848.1	Sox	Sox-4
Casi	XP_006874684.1	Fox	Fox-P1		Lcat	XP_045407834.1	Sox	Sox-4
Emax	XP_049718943.1	Fox	Fox-P1		Hamp	XP_057551434.1	Sox	Sox-4
Mjav	XP_036860335.1	Fox	Fox-P1		Mmus	NP_033264.2	Sox	Sox-4
Pgig	XP_039707610.1	Fox	Fox-P1		Drot	XP_053776704.1	Sox	Sox-4
Drot	XP_045048358.1	Fox	Fox-P1		Oafe	XP_007950280.1	Sox	Sox-4
Mmus	XP_030110934.1	Fox	Fox-P1		Dnov	XP_012385001.3	Sox	Sox-4
Tman	XP_023583661.1	Fox	Fox-P1		Bbub	XP_025121975.3	Sox	Sox-4
Cimi	XP_017398291.1	Fox	Fox-P1		Hamp	XP_057555091.1	Sox	Sox-4
Rfer	XP_032987798.1	Fox	Fox-P1		Mmus	NP_033264.2	Sox	Sox-4
Scar	XP_047390767.1	Fox	Fox-P1		Drot	XP_053776704.1	Sox	Sox-4
Ptig	XP_042835351.1	Fox	Fox-P1		Oafe	XP_007950280.1	Sox	Sox-4
Oana	XP_039766065.1	Fox	Fox-P1		Dnov	XP_012385001.3	Sox	Sox-4
Equa	XP_046523540.1	Fox	Fox-P1		Clup	XP_038440195.1	Sox	Sox-4
Clup	XP_038421798.1	Fox	Fox-P1		Equa	XP_046497050.1	Sox	Sox-4
Mang	XP_054360611.1	Fox	Fox-P1		Amel	XP_034516148.1	Sox	Sox-4
Bmusp	XP_036724195.1	Fox	Fox-P1		Cdro	XP_031290990.1	Sox	Sox-4
Cpor	XP_005005313.1	Fox	Fox-P1		Emax	XP_049732340.1	Sox	Sox-4
Cdid	XP_037656169.1	Fox	Fox-P1		Csim	XP_004432138.1	Sox	Sox-4
Hamp	XP_057561364.1	Fox	Fox-P1		Mjav	XP_036850086.1	Sox	Sox-4
Lcat	XP_045385591.1	Fox	Fox-P1		Pgig	XP_039722376.1	Sox	Sox-4
Bbub	XP_044789782.1	Fox	Fox-P1		Ptig	XP_042841543.1	Sox	Sox-4
Csim	XP_004419976.1	Fox	Fox-P1		Bmusp	XP_036725088.1	Sox	Sox-4
Dnov	XP_004482088.1	Fox	Fox-P1		Tman	XP_023594118.1	Sox	Sox-4
Ggal	XP_040502196.1	Fox	Fox-P1		Pafn	XP_047651758.1	Sox	Sox-4
Tman	XP_004385433.1	Fox	Fox-N1		Mang	XP_045722257.1	Sox	Sox-4
Pafn	XP_047614767.1	Fox	Fox-N1		Rfer	XP_032971451.1	Sox	Sox-4
Cdid	XP_037665986.1	Fox	Fox-N1		Cdid	XP_037700378.1	Sox	Sox-4
Casi	XP_006874270.1	Fox	Fox-N1		Cimi	XP_017399572.1	Sox	Sox-4
Emax	XP_049715494.1	Fox	Fox-N1		Shar	XP_031802646.1	Sox	Sox-4
Scar	XP_047401997.1	Fox	Fox-N1		Shar	XP_003775357.4	Sox	Sox-15
Oafe	XP_007935559.1	Fox	Fox-N1		Dnov	XP_004468798.1	Sox	Sox-15
Ggal	XP_415816.5	Fox	Fox-N1		Mdom	XP_056673573.1	Sox	Sox-15
Mmus	XP_006532328.1	Fox	Fox-N1		Oana	XP_039766818.1	Sox	Sox-15
Oana	XP_039770471.1	Fox	Fox-N1		Clup	XP_038392636.1	Sox	Sox-15
Mdom	XP_001375832.1	Fox	Fox-N1		Mjav	XP_017527261.1	Sox	Sox-15
Dnov	XP_004450005.2	Fox	Fox-N1		Cdid	XP_037664941.1	Sox	Sox-15
Shar	XP_031823539.1	Fox	Fox-N1		Pgig	XP_039703166.1	Sox	Sox-15
Cpor	XP_023420823.1	Fox	Fox-N1		Pafn	XP_047613485.1	Sox	Sox-15
Lcat	XP_045381339.1	Fox	Fox-N1		Bmusp	XP_036692236.1	Sox	Sox-15
Bbub	XP_025136790.1	Fox	Fox-N1		Ttru	XP_033704061.1	Sox	Sox-15
Opri	XP_004593965.2	Fox	Fox-N1		Emax	XP_049714749.1	Sox	Sox-15
Bmusp	XP_036693663.1	Fox	Fox-N1		Opri	XP_004594843.2	Sox	Sox-15
Drot	XP_045055001.2	Fox	Fox-N1		Cpor	XP_034646299.1	Sox	Sox-15
Amel	XP_034496649.1	Fox	Fox-N1		Rfer	XP_032947056.1	Sox	Sox-15
Pgig	XP_039742030.1	Fox	Fox-N1		Cimi	XP_037595583.1	Sox	Sox-15

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Sp. ID	Gene ID	Group	Annotation		Sp. ID	Gene ID	Group	Annotation
Cdro	XP_010996180.2	Fox	Fox-13		Emax	XP_049729443.1	Sox	Sry
Mjav	XP_017536768.2	Fox	Fox-13		Cdro	XP_031300930.1	Sox	Sox-3
Oafe	XP_007952295.1	Fox	Fox-13		Pgig	XP_039735027.1	Sox	Sry
Drot	XP_024408151.2	Fox	Fox-13		Shar	XP_031801149.1	Sox	Sry
Oana	XP_028931929.2	Fox	Fox-L1		Lcat	XP_045393696.1	Sox	Sox-3
Bbub	XP_006047558.1	Fox	Fox-L1		Cimi	XP_017374345.1	Sox	Sry
Mdom	XP_056674514.1	Fox	Fox-L1		Hamp	XP_057574656.1	Sox	Sox-3
Oafe	XP_007937407.1	Fox	Fox-L1		Mmus	NP_033263.2	Sox	Sox-3
Mjav	XP_036849629.1	Fox	Fox-L1		Amel	XP_034504946.1	Sox	Sox-3
Rfer	XP_032984676.1	Fox	Fox-L1		Cdid	XP_037678811.1	Sox	Sox-3
Hamp	XP_057569166.1	Fox	Fox-L1		Bmus	XP_036696323.1	Sox	Sox-3
Pgig	XP_039744660.1	Fox	Fox-L1		Ttru	XP_033706224.1	Sox	Sry
Dnov	XP_058135467.1	Fox	Fox-L1		Bbub	XP_025132502.3	Sox	Sox-3
Csim	XP_004437100.1	Fox	Fox-L1		Cdid	XP_037680520.1	Sox	Sry
Casi	XP_006860180.1	Fox	Fox-L1		Cpor	XP_003464754.2	Sox	Sox-3
Opri	XP_004584130.2	Fox	Fox-L1		Scar	XP_047393176.1	Sox	Sry
Clup	XP_038394040.1	Fox	Fox-L1		Opri	XP_058515261.1	Sox	Sry
Hsap	NP_005241.1	Fox	Fox-L1		Pafr	XP_047620863.1	Sox	Sox-3
Drot	XP_024411663.1	Fox	Fox-L1		Mmus	NP_035694.1	Sox	Sry
Amel	XP_002913428.1	Fox	Fox-L1		Drot	XP_053773489.1	Sox	Sox-3
Cdro	XP_031292484.1	Fox	Fox-L1		Mdom	XP_007507433.1	Sox	Sox-3
Cimi	XP_017399161.1	Fox	Fox-L1		Scar	XP_047392867.1	Sox	Sox-3
Cdid	XP_037671791.1	Fox	Fox-L1		Oafe	XP_007957280.1	Sox	Sry
Ttru	XP_004312143.1	Fox	Fox-L1		Cimi	XP_017355427.1	Sox	Sox-3
Rfer	XP_032984241.1	Fox	Fox-L1		Ttru	XP_033706225.1	Sox	Sry
Ptig	XP_042825075.1	Fox	Fox-L1		Mang	XP_045726266.1	Sox	Sox-3
Bmus	XP_036688240.1	Fox	Fox-L1		Casi	XP_006875033.1	Sox	Sox-3
Tman	XP_004377800.1	Fox	Fox-L1		Opri	XP_058515255.1	Sox	Sry
Mang	XP_045746246.1	Fox	Fox-L1		Mjav	XP_036857050.1	Sox	Sry
Shar	XP_031806032.1	Fox	Fox-L1		Dnov	XP_058142693.1	Sox	Sox-8
Mmus	NP_032050.2	Fox	Fox-L1		Casi	XP_006873911.1	Sox	Sox-8
Pafr	XP_047648182.1	Fox	Fox-L1		Cdid	XP_037670405.1	Sox	Sox-8
Equa	XP_046536521.1	Fox	Fox-L1		Emax	XP_049759218.1	Sox	Sox-8
Scar	XP_047385703.1	Fox	Fox-L1		Tman	XP_023584110.1	Sox	Sox-8
Ggal	XP_001231599.5	Fox	Fox-L1		Oafe	XP_007955685.1	Sox	Sox-8
Emax	XP_049720646.1	Fox	Fox-L1		Mjav	XP_017510601.1	Sox	-
Lcat	XP_045389091.1	Fox	Fox-L1		Clup	XP_038432983.1	Sox	-
Ggal	XP_004948470.2	Fox	Fox-P4		Ptig	XP_042847878.1	Sox	-
Casi	XP_006860505.1	Fox	Fox-P4		Mang	XP_054363126.1	Sox	-
Equa	XP_046496143.1	Fox	Fox-P4		Amel	XP_019650885.1	Sox	-
Cdid	XP_037698927.1	Fox	Fox-P4		Ggal	NP_989664.1	Sox	Sox-1
Mdom	XP_056674181.1	Fox	Fox-P4		Mjav	XP_036880873.1	Sox	Sox-1
Oafe	XP_007934458.1	Fox	Fox-P4		Amel	XP_034519706.1	Sox	Sox-1
Opri	XP_058519292.1	Fox	Fox-P4		Lcat	XP_045422689.1	Sox	Sox-1
Mmus	XP_017173175.1	Fox	Fox-P4		Oana	XP_028904097.1	Sox	Sox-1
Rfer	XP_032956861.1	Fox	Fox-P4		Opri	XP_058526612.1	Sox	Sox-1
Hsap	XP_011512591.1	Fox	Fox-P4		Cdid	XP_037656244.1	Sox	Sox-1
Ptig	XP_042842255.1	Fox	Fox-P4		Scar	XP_047410448.1	Sox	Sox-1
Ttru	XP_033720169.1	Fox	Fox-P4		Mdom	XP_007501341.1	Sox	Sox-1
Mang	XP_045722826.1	Fox	Fox-P4		Equa	XP_046519750.1	Sox	Sox-1
Mjav	XP_036861021.1	Fox	Fox-P4		Ttru	XP_019776483.2	Sox	Sox-1
Cimi	XP_017402725.1	Fox	Fox-P4		Clup	XP_038425993.1	Sox	Sox-1
Clup	XP_038409938.1	Fox	Fox-P4		Cdro	XP_031322193.1	Sox	Sox-1
Shar	XP_031820748.1	Fox	Fox-P4		Shar	XP_031814590.1	Sox	Sox-1
Lcat	XP_045397843.1	Fox	Fox-P4		Oafe	XP_007940578.1	Sox	Sox-1
Pgig	XP_039710787.1	Fox	Fox-P4		Rfer	XP_032959657.1	Sox	Sox-1
Drot	XP_053769706.1	Fox	Fox-P4		Mmus	NP_033259.2	Sox	Sox-1
Bmus	XP_036725368.1	Fox	Fox-P4		Dnov	XP_058132394.1	Sox	Sox-1
Amel	XP_002914563.3	Fox	Fox-P4		Bmus	XP_036687021.1	Sox	Sox-1
Csim	XP_004424223.1	Fox	Fox-P4		Pafr	XP_047612043.1	Sox	Sox-1
Oana	XP_028925009.1	Fox	Fox-P4		Drot	XP_045042640.2	Sox	Sox-1
Hamp	XP_057555834.1	Fox	Fox-P4		Ptig	XP_042814197.1	Sox	Sox-1
Pafr	XP_047653356.1	Fox	Fox-P4		Cimi	XP_017357713.1	Sox	Sox-1
Cpor	XP_003473920.1	Fox	Fox-P4		Mang	XP_045723818.1	Sox	Sox-1
Scar	XP_047414507.1	Fox	Fox-P4		Casi	XP_006851579.1	Sox	Sox-1
Emax	XP_049747581.1	Fox	Fox-P4		Cpor	XP_005007254.3	Sox	Sox-1
Tman	XP_004379494.1	Fox	Fox-P4		Hsap	NP_005977.2	Sox	Sox-1
Dnov	XP_058162686.1	Fox	Fox-P4		Hamp	XP_057563730.1	Sox	Sox-1
Bbub	XP_025125991.2	Fox	Fox-P4		Pgig	XP_039721974.1	Sox	Sox-1
Cdro	XP_031290589.1	Fox	Fox-P4		Bbub	XP_025118363.1	Sox	Sox-1
Bmus	XP_036718879.1	Fox	Fox-P2		Emax	XP_049723386.1	Sox	Sox-1
Ttru	XP_019780690.1	Fox	Fox-P2		Mdom	XP_001373727.1	Sox	-
Lcat	XP_045420440.1	Fox	Fox-P2		Shar	XP_003762017.1	Sox	-
Ptig	XP_042826228.1	Fox	Fox-P2		Hamp	XP_057557078.1	Sox	Sox-5
Opri	XP_058510875.1	Fox	Fox-P2		Equa	XP_046497249.1	Sox	Sox-5
Csim	XP_004418850.1	Fox	Fox-P2		Bbub	XP_045021534.1	Sox	Sox-5
Shar	XP_031794706.1	Fox	Fox-P2		Csim	XP_014646859.1	Sox	Sox-5
Equa	XP_046526335.1	Fox	Fox-P2		Cdro	XP_031299822.1	Sox	Sox-5
Cdro	XP_031311093.1	Fox	Fox-P2		Tman	XP_023593782.1	Sox	Sox-5
Amel	XP_034526782.1	Fox	Fox-P2		Emax	XP_049741103.1	Sox	Sox-5
Ptig	XP_042826228.1	Fox	Fox-P2		Oafe	XP_007944956.1	Sox	Sox-5
Opri	XP_058510875.1	Fox	Fox-P2		Pafr	XP_047643337.1	Sox	Sox-5
Cpor	XP_013013912.1	Fox	Fox-P2		Casi	XP_006867010.1	Sox	Sox-5
Casi	XP_006859359.1	Fox	Fox-P2		Clup	XP_038439395.1	Sox	Sox-2
Oana	XP_028928988.1	Fox	Fox-P2		Emax	XP_046515240.1	Sox	Sox-2
Pafr	XP_047618861.1	Fox	Fox-P2		Csim	XP_004424739.1	Sox	Sox-2
Oafe	XP_007942246.1	Fox	Fox-P2		Hamp	XP_057595873.1	Sox	Sox-2
Hsap	NP_683696.2	Fox	Fox-P2		Emax	XP_049730807.1	Sox	Sox-2
Mdom	XP_007504226.1	Fox	Fox-P2		Casi	XP_006869879.1	Sox	Sox-2
Ggal	XP_025007337.1	Fox	Fox-P2		Ptig	XP_042855191.1	Sox	Sox-2
Dnov	XP_058153204.1	Fox	Fox-P2		Ggal	NP_990519.3	Sox	Sox-2
Bbub	XP_044802585.1	Fox	Fox-P2		Clup	XP_038439395.1	Sox	Sox-2
Cimi	XP_017390531.1	Fox	Fox-P2		Emax	XP_046515240.1	Sox	Sox-2
Rfer	XP_032954948.1	Fox	Fox-P2		Hamp	XP_057595873.1	Sox	Sox-2
Hamp	XP_057586867.1	Fox	Fox-P2		Oafe	NP_003097.1	Sox	Sox-2
Mmus	XP_036021645.1	Fox	Fox-P2		Shar	XP_031813587.1	Sox	Sox-2
Mjav	XP_036883368.1	Fox	Fox-P2		Mang	XP_04571963.1	Sox	Sox-2
Emax	XP_049750594.1	Fox	Fox-P2		Scar	XP_047420710.1	Sox	Sox-2
Cdid	XP_037692269.1	Fox	Fox-P2		Tman	XP_023592536.1	Sox	Sox-2
Drot	XP_053782699.1	Fox	Fox-P2		Bbub	XP_006056297.2	Sox	Sox-2
Scar	XP_047417295.1	Fox	Fox-P2		Rfer	XP_032989610.1	Sox	Sox-2
Mang	XP_045720337.1	Fox	Fox-P2		Lcat	XP_045395708.1	Sox	Sox-2
Tman	XP_004382624.1	Fox	Fox-P2		Opri	XP_058518203.1	Sox	Sox-2
					Oana	XP_028928807.1	Sox	Sox-2

Tab. S8 continued from previous page

Sp. ID	Gene ID	Group	Annotation	Sp. ID	Gene ID	Group	Annotation
Clup	XP_038413332.1	Fox	Fox-P2	Bmus	XP_036705414.1	Sox	Sox-2
Cdid	XP_037688024.1	Fox	-	Mmus	NP_035573.3	Sox	Sox-2
Dnov	XP_058148549.1	Fox	-	Cdid	XP_037695755.1	Sox	Sox-2
Ggal	XP_414186.5	Fox	Fox-F1	Pgig	XP_039706608.1	Sox	Sox-2
Emax	XP_049720665.1	Fox	Fox-F1	Oafe	XP_007934094.1	Sox	Sox-2
Mdom	XP_001365832.4	Fox	Fox-F1	Pafr	XP_047643165.1	Sox	Sox-2
Hsap	NP_001442.2	Fox	Fox-F1	Dnov	XP_004477690.1	Sox	Sox-2
Clup	XP_038394042.1	Fox	Fox-F1	Cimi	XP_017393930.1	Sox	Sox-2
Mjav	XP_0175111561.2	Fox	Fox-F1	Drot	XP_053774095.1	Sox	Sox-2
Cimi	XP_017399155.2	Fox	Fox-F1	Amel	XP_034518206.1	Sox	Sox-2
Bbub	XP_006047564.2	Fox	Fox-F1	Cdro	XP_010974669.2	Sox	Sox-2
Dnov	XP_058135594.1	Fox	Fox-F1	Mjav	XP_017496634.2	Sox	Sox-2
Drot	XP_053772163.1	Fox	Fox-F1	Ttru	XP_004311832.3	Sox	Sox-2
Tman	XP_004377797.2	Fox	Fox-F1	Oana	XP_028931165.1	Sox	Sox-14
Bmus	XP_036688072.1	Fox	Fox-F1	Mdom	XP_007943983.1	Sox	Sox-14
Oana	XP_0289311561.1	Fox	Fox-F1	Scar	XP_047419052.1	Sox	Sox-14
Pafr	XP_047645925.1	Fox	Fox-F1	Cdid	XP_037683894.1	Sox	Sox-14
Scar	XP_047385701.1	Fox	Fox-F1	Bmus	XP_036707689.1	Sox	Sox-14
Lcat	XP_045389195.1	Fox	Fox-F1	Mjav	XP_017531607.1	Sox	Sox-14
Shar	XP_003758519.2	Fox	Fox-F1	Casi	XP_006846671.1	Sox	Sox-14
Amel	XP_002913450.2	Fox	Fox-F1	Pafr	XP_047610665.1	Sox	Sox-14
Opri	XP_058530279.1	Fox	Fox-F1	Hamp	XP_057595112.1	Sox	Sox-14
Pgig	XP_039744668.1	Fox	Fox-F1	Amel	XP_002923375.1	Sox	Sox-14
Casi	XP_006860182.1	Fox	Fox-F1	Csim	XP_004419391.1	Sox	Sox-14
Mmus	NP_034556.2	Fox	Fox-F1	Cpor	XP_003476788.1	Sox	Sox-14
Cdid	XP_037671853.1	Fox	Fox-F1	Hsap	NP_004180.1	Sox	Sox-14
Cpor	XP_003460772.2	Fox	Fox-F1	Clup	XP_038426113.1	Sox	Sox-14
Equa	XP_046538978.1	Fox	Fox-F1	Bbub	XP_025147052.1	Sox	Sox-14
Ptig	XP_007082882.2	Fox	Fox-F1	Rfer	XP_032988974.1	Sox	Sox-14
Csim	XP_014648135.1	Fox	Fox-F1	Equa	XP_046511648.1	Sox	Sox-14
Cdro	XP_031292478.1	Fox	Fox-F1	Drot	XP_024422013.1	Sox	Sox-14
Rfer	XP_032985403.1	Fox	Fox-F1	Lcat	XP_045395088.1	Sox	Sox-14
Mang	XP_045746573.2	Fox	Fox-F1	Opri	XP_004588396.1	Sox	Sox-14
Hamp	XP_057568154.1	Fox	Fox-F1	Pgig	XP_039692761.1	Sox	Sox-14
Oafe	XP_007937340.1	Fox	Fox-F1	Ptig	XP_042813213.1	Sox	Sox-14
Ttru	XP_033700961.1	Fox	Fox-F1	Ttru	XP_019789673.1	Sox	Sox-14
Mdom	XP_056669097.1	Fox	-	Cimi	XP_017383705.1	Sox	Sox-14
Shar	XP_031805673.1	Fox	-	Emax	XP_049726472.1	Sox	Sox-14
Ggal	XP_425714.5	Fox	Fox-S1	Mang	XP_045719678.1	Sox	Sox-14
Clup	XP_038427866.1	Fox	Fox-S1	Oafe	XP_007935393.1	Sox	Sox-14
Hsap	NP_004109.1	Fox	Fox-S1	Ggal	NP_990092.1	Sox	Sox-14
Pgig	XP_039719969.1	Fox	Fox-S1	Tman	XP_004381489.1	Sox	Sox-14
Bmus	XP_036682343.1	Fox	Fox-S1	Dnov	XP_004453833.1	Sox	Sox-14
Mdom	XP_001364156.2	Fox	Fox-S1	Shar	XP_012400053.1	Sox	Sox-14
Amel	XP_002918325.1	Fox	Fox-S1	Mmus	NP_035570.1	Sox	Sox-14
Shar	XP_031800982.1	Fox	Fox-S1	Cdro	XP_010975059.1	Sox	Sox-14
Bbub	XP_0251119474.1	Fox	Fox-S1	Ggal	NP_001383603.1	Sox	Sox-21
Casi	XP_006860769.1	Fox	Fox-S1	Mang	XP_045726097.1	Sox	Sox-21
Cdro	XP_010992198.2	Fox	Fox-S1	Rfer	XP_032959615.1	Sox	Sox-21
Mang	XP_045741514.1	Fox	Fox-S1	Bmus	XP_036687740.1	Sox	Sox-21
Pafr	XP_047627330.1	Fox	Fox-S1	Dnov	XP_058132486.1	Sox	Sox-21
Dnov	XP_004464085.1	Fox	Fox-S1	Equa	XP_046520974.1	Sox	Sox-21
Opri	XP_004585763.2	Fox	Fox-S1	Drot	XP_053772754.1	Sox	Sox-21
Cpor	XP_0034776682.1	Fox	Fox-S1	Opri	XP_058526666.1	Sox	Sox-21
Mjav	XP_017509202.2	Fox	Fox-S1	Cdro	XP_037656473.1	Sox	Sox-21
Csim	XP_004442549.2	Fox	Fox-S1	Csim	XP_014637259.1	Sox	Sox-21
Rfer	XP_032950864.1	Fox	Fox-S1	Shar	XP_031814415.1	Sox	Sox-21
Equa	XP_046536068.1	Fox	Fox-S1	Ttru	XP_033699347.1	Sox	Sox-21
Scar	XP_047396619.1	Fox	Fox-S1	Tman	XP_004371490.1	Sox	Sox-21
Tman	XP_023581200.1	Fox	Fox-S1	Mmus	NP_808421.1	Sox	Sox-21
Lcat	XP_045385439.1	Fox	Fox-S1	Oana	XP_001512271.3	Sox	Sox-21
Emax	XP_049725743.1	Fox	Fox-S1	Clup	XP_038425312.1	Sox	Sox-21
Mmus	NP_034356.1	Fox	Fox-S1	Ptig	XP_042813926.1	Sox	Sox-21
Ttru	XP_033696063.1	Fox	Fox-S1	Amel	XP_034519700.1	Sox	Sox-21
Ptig	XP_007073610.2	Fox	Fox-S1	Casi	XP_006832065.1	Sox	Sox-21
Hamp	XP_057559118.1	Fox	Fox-S1	Hamp	XP_057563720.1	Sox	Sox-21
Drot	XP_024415665.3	Fox	Fox-S1	Cdro	XP_031322066.1	Sox	Sox-21
Cimi	XP_017370299.1	Fox	Fox-S1	Mang	XP_045726110.1	Sox	Sox-21
Oafe	XP_007932722.1	Fox	Fox-S1	Pgig	XP_039693387.1	Sox	Sox-21
Cdid	XP_037667492.1	Fox	Fox-S1	Mjav	XP_036880860.1	Sox	Sox-21
Ggal	XP_015135808.1	Fox	Fox-B2	Scar	XP_047410577.1	Sox	Sox-21
Pafr	XP_047625507.1	Fox	Fox-B2	Pafr	XP_047612344.1	Sox	Sox-21
Shar	XP_012398564.2	Fox	Fox-B2	Oafe	XP_007942198.1	Sox	Sox-21
Csim	XP_004439686.1	Fox	Fox-B2	Cimi	XP_017399045.1	Sox	Sox-21
Scar	XP_047379799.1	Fox	Fox-B2	Mdom	XP_007501455.1	Sox	Sox-21
Cdro	XP_031305772.1	Fox	Fox-B2	Bbub	XP_025118466.2	Sox	Sox-21
Equa	XP_046519944.1	Fox	Fox-B2	Lcat	XP_045423694.1	Sox	Sox-21
Bbub	XP_006067725.3	Fox	Fox-B2	Hsap	NP_009015.1	Sox	Sox-21
Cimi	XP_017398753.1	Fox	Fox-B2	Emax	XP_049709053.1	Sox	Sox-21
Mmus	NP_032049.1	Fox	Fox-B2	Scar	XP_047405480.1	Sox	Sox-5
Pgig	XP_039724438.1	Fox	Fox-B2	Opri	XP_058511892.1	Sox	Sox-5
Lcat	XP_045418554.1	Fox	Fox-B2	Scar	XP_047405480.1	Sox	Sox-5
Casi	XP_006834992.1	Fox	Fox-B2	Mmus	NP_006506994.1	Sox	Sox-5
Mdom	XP_003341714.1	Fox	Fox-B2	Bmus	XP_036693736.1	Sox	Sox-9
Mjav	XP_036847034.1	Fox	Fox-B2	Hamp	XP_057573119.1	Sox	Sox-9
Oana	XP_001508095.2	Fox	Fox-B2	Ggal	NP_989612.1	Sox	Sox-9
Mang	XP_045737891.1	Fox	Fox-B2	Drot	XP_024429046.1	Sox	Sox-9
Drot	XP_053776114.1	Fox	Fox-B2	Tman	XP_004374235.1	Sox	Sox-9
Dnov	XP_023440163.2	Fox	Fox-B2	Cimi	XP_017391090.1	Sox	Sox-9
Opri	XP_058513381.1	Fox	Fox-B2	Clup	NP_001002978.1	Sox	Sox-9
Ttru	XP_019789709.1	Fox	Fox-B2	Dnov	XP_058140625.1	Sox	Sox-9
Cdid	XP_037654560.1	Fox	Fox-B2	Opri	XP_004593006.1	Sox	Sox-9
Tman	XP_023593539.1	Fox	Fox-B2	Casi	XP_006872359.1	Sox	Sox-9
Amel	XP_034502902.1	Fox	Fox-B2	Cdro	XP_031304465.1	Sox	Sox-9
Bmus	XP_036712759.1	Fox	Fox-B2	Mjav	XP_017519930.2	Sox	Sox-9
Clup	XP_038381543.1	Fox	Fox-B2	Csim	XP_004432762.1	Sox	Sox-9
Emax	XP_049751166.1	Fox	Fox-B2	Ptig	XP_042822833.1	Sox	Sox-9
Hsap	NP_001013757.1	Fox	Fox-B2	Equa	XP_046530969.1	Sox	Sox-9
Rfer	XP_032978089.1	Fox	Fox-B2	Emax	XP_049717106.1	Sox	Sox-9
Hamp	XP_057580851.1	Fox	Fox-B2	Oana	XP_001506094.2	Sox	Sox-9
Oafe	XP_007936768.1	Fox	Fox-B2	Rfer	XP_032945891.1	Sox	Sox-9
Ptig	XP_042819403.1	Fox	Fox-B2	Ttru	XP_033703168.1	Sox	Sox-9

Tab. S8 continued from previous page

Sp. ID	Gene ID	Group	Annotation	Sp. ID	Gene ID	Group	Annotation
Oana	XP_001518148.2	Fox	Fox-B1	Amel	XP_019660087.2	Sox	Sox-9
Clup	XP_038436513.1	Fox	Fox-B1	Mang	XP_045743969.1	Sox	Sox-9
Shar	XP_003755656.2	Fox	Fox-B1	Pafr	XP_047614296.1	Sox	Sox-9
Ttru	XP_033708504.1	Fox	Fox-B1	Scar	XP_047398689.1	Sox	Sox-9
Mjav	XP_017523883.1	Fox	Fox-B1	Bbub	XP_025135172.2	Sox	Sox-9
Casi	XP_006831648.1	Fox	Fox-B1	Shar	XP_003768593.1	Sox	Sox-9
Rfer	XP_032964940.1	Fox	Fox-B1	Hsap	NP_000337.1	Sox	Sox-9
Tman	XP_004374676.1	Fox	Fox-B1	Mmus	NP_035578.3	Sox	Sox-9
Hsap	NP_036314.2	Fox	Fox-B1	Oafe	XP_007957879.1	Sox	Sox-9
Amel	XP_034516791.1	Fox	Fox-B1	Cdid	XP_037664998.1	Sox	Sox-9
Pgig	XP_039720931.1	Fox	Fox-B1	Lcat	XP_045425568.1	Sox	Sox-9
Cdro	XP_031308829.1	Fox	Fox-B1	Pgig	XP_039715158.1	Sox	Sox-9
Opri	XP_004578087.1	Fox	Fox-B1	Ggal	XP_015139949.1	Sox	Sox-10
Mang	XP_045755904.1	Fox	Fox-B1	Mmus	NP_035567.1	Sox	Sox-10
Cimi	XP_017387405.1	Fox	Fox-B1	Csim	XP_004437902.1	Sox	Sox-10
Dnov	XP_012376939.1	Fox	Fox-B1	Hsap	NP_008872.1	Sox	Sox-10
Equa	XP_046508174.1	Fox	Fox-B1	Mjav	XP_036862891.1	Sox	Sox-10
Mdom	XP_001365592.1	Fox	Fox-B1	Oafe	XP_007939944.1	Sox	Sox-10
Scar	XP_047396474.1	Fox	Fox-B1	Opri	XP_004589571.1	Sox	Sox-10
Emax	XP_049708649.1	Fox	Fox-B1	Scar	XP_047407584.1	Sox	Sox-10
Bmus	XP_036700853.1	Fox	Fox-B1	Mang	XP_045728505.1	Sox	Sox-10
Oafe	XP_007956505.1	Fox	Fox-B1	Cdro	XP_031319106.1	Sox	Sox-10
Bbub	XP_006043154.1	Fox	Fox-B1	Shar	XP_031797111.1	Sox	Sox-10
Ggal	XP_004943811.1	Fox	Fox-B1	Cimi	XP_017364446.1	Sox	Sox-10
Cpor	XP_013000887.1	Fox	Fox-B1	Ttru	XP_033721970.1	Sox	Sox-10
Ptig	XP_042843621.1	Fox	Fox-B1	Dnov	XP_004466267.1	Sox	Sox-10
Mmus	NP_071773.2	Fox	Fox-B1	Ptig	XP_042849176.1	Sox	Sox-10
Csim	XP_004421699.1	Fox	Fox-B1	Tman	XP_004373905.1	Sox	Sox-10
Hamp	XP_057575814.1	Fox	Fox-B1	Cpor	XP_003470533.2	Sox	Sox-10
Lcat	XP_045411390.1	Fox	Fox-B1	Hamp	XP_057595912.1	Sox	Sox-10
Cdid	XP_037691321.1	Fox	Fox-B1	Oana	XP_028934366.1	Sox	Sox-10
Drot	XP_024424310.1	Fox	Fox-B1	Amel	XP_034500270.1	Sox	Sox-10
Pafr	XP_047625395.1	Fox	Fox-B1	Casi	XP_006865257.1	Sox	Sox-10
Oana	XP_028925280.1	Fox	Hnf-3b/Fox-A2	Bbub	XP_006071428.1	Sox	Sox-10
Mdom	XP_001382097.1	Fox	Hnf-3b/Fox-A2	Cdid	XP_037702433.1	Sox	Sox-10
Clup	XP_038427348.1	Fox	Hnf-3b/Fox-A2	Lcat	XP_045408856.1	Sox	Sox-10
Amel	XP_002924170.1	Fox	Hnf-3b/Fox-A2	Equa	XP_046502255.1	Sox	Sox-10
Cimi	XP_017396970.1	Fox	Hnf-3b/Fox-A2	Pgig	XP_039733644.1	Sox	Sox-10
Mjav	XP_036878344.1	Fox	Hnf-3b/Fox-A2	Clup	XP_038406550.1	Sox	Sox-10
Ttru	XP_019783169.2	Fox	Hnf-3b/Fox-A2	Rfer	XP_032973012.1	Sox	Sox-10
Mang	XP_045739756.1	Fox	Hnf-3b/Fox-A2	Emax	XP_049740365.1	Sox	Sox-10
Shar	XP_003758186.1	Fox	Hnf-3b/Fox-A2	Drot	XP_024434928.2	Sox	Sox-10
Emax	XP_049725634.1	Fox	Hnf-3b/Fox-A2	Pafr	XP_047642247.1	Sox	Sox-10
Scar	XP_047397766.1	Fox	Hnf-3b/Fox-A2	Mdom	XP_001381534.3	Sox	Sox-10
Pgig	XP_039730271.1	Fox	Hnf-3b/Fox-A2	Bmus	XP_036720803.1	Sox	Sox-10
Equa	XP_046535478.1	Fox	Hnf-3b/Fox-A2	Hsap	NP_011519134.2	Sox	Sox-5
Hsap	NP_068556.2	Fox	Hnf-3b/Fox-A2	Lcat	XP_045410158.1	Sox	Sox-5
Mmus	NP_001277994.1	Fox	Hnf-3b/Fox-A2	Oana	XP_028925314.1	Sox	Sox-13
Lcat	XP_045385044.1	Fox	Hnf-3b/Fox-A2	Pafr	XP_047607942.1	Sox	Sox-13
Drot	XP_045050691.1	Fox	Hnf-3b/Fox-A2	Shar	XP_003767655.1	Sox	Sox-13
Cdid	XP_037668003.1	Fox	Hnf-3b/Fox-A2	Ggal	XP_015154614.2	Sox	Sox-13
Tman	XP_004376477.1	Fox	Hnf-3b/Fox-A2	Mdom	XP_056671743.1	Sox	Sox-13
Casi	XP_006860717.1	Fox	Hnf-3b/Fox-A2	Ttru	XP_033708141.1	Sox	Sox-13
Oafe	XP_007948556.1	Fox	Hnf-3b/Fox-A2	Cpor	XP_013013443.1	Sox	Sox-13
Pafr	XP_047627543.1	Fox	Hnf-3b/Fox-A2	Bmus	XP_036684089.1	Sox	Sox-13
Hamp	XP_057558353.1	Fox	Hnf-3b/Fox-A2	Mang	XP_054363913.1	Sox	Sox-13
Dnov	XP_023439993.2	Fox	Hnf-3b/Fox-A2	Tman	XP_023586088.1	Sox	Sox-13
Bmus	XP_036681859.1	Fox	Hnf-3b/Fox-A2	Hamp	XP_057585117.1	Sox	Sox-13
Rfer	XP_032950275.1	Fox	Hnf-3b/Fox-A2	Cdro	XP_031294667.1	Sox	Sox-13
Cpor	XP_003476486.1	Fox	Hnf-3b/Fox-A2	Drot	XP_024430737.2	Sox	Sox-13
Opri	XP_004585707.2	Fox	Hnf-3b/Fox-A2	Dnov	XP_058130986.1	Sox	Sox-13
Bbub	XP_044783908.1	Fox	Hnf-3b/Fox-A2	Mjav	XP_017506787.2	Sox	Sox-13
Cdro	XP_031290193.1	Fox	Hnf-3b/Fox-A2	Amel	XP_034522993.1	Sox	Sox-13
Csim	XP_004442004.1	Fox	Hnf-3b/Fox-A2	Mmus	XP_011246247.1	Sox	Sox-13
Ptig	XP_042835681.1	Fox	Hnf-3b/Fox-A2	Scar	XP_047376454.1	Sox	Sox-13
Ggal	XP_046794381.1	Fox	Hnf-3b/Fox-A2	Ptig	XP_007086583.2	Sox	Sox-13
Oana	XP_028902569.1	Fox	Fox-E3	Clup	XP_038441905.1	Sox	Sox-13
Emax	XP_049731498.1	Fox	Fox-E3	Oafe	XP_007954219.1	Sox	Sox-13
Hsap	NP_036318.1	Fox	Fox-E3	Opri	XP_058525035.1	Sox	Sox-13
Mang	XP_045751381.1	Fox	Fox-E3	Emax	XP_049714298.1	Sox	Sox-13
Hamp	XP_057562286.1	Fox	Fox-E3	Equa	XP_046537707.1	Sox	Sox-13
Drot	XP_053777352.1	Fox	Fox-E3	Casi	XP_006834281.1	Sox	Sox-13
Ttru	XP_033709174.1	Fox	Fox-E3	Hsap	NP_005677.2	Sox	Sox-13
Pafr	XP_047647470.1	Fox	Fox-E3	Cimi	XP_037593260.1	Sox	Sox-13
Ptig	XP_042852491.1	Fox	Fox-E3	Bbub	XP_025133526.2	Sox	Sox-13
Clup	XP_038413565.1	Fox	Fox-E3	Cdid	XP_037680910.1	Sox	Sox-13
Rfer	XP_032971209.1	Fox	Fox-E3	Lcat	XP_045391966.1	Sox	Sox-13
Bmus	XP_036690899.1	Fox	Fox-E3	Csim	XP_004425199.1	Sox	Sox-13
Pgig	XP_039712722.1	Fox	Fox-E3	Rfer	XP_032948937.1	Sox	Sox-13
Bbub	XP_025144571.1	Fox	Fox-E3	Pgig	XP_039737775.1	Sox	Sox-13
Cdro	XP_031321430.1	Fox	Fox-E3	Ggal	XP_025006442.1	Sox	Sox-6
Tman	XP_023582802.1	Fox	Fox-E3	Emax	XP_049746477.1	Sox	Sox-6
Pgig	XP_039738800.1	Fox	Fox-E3	Ttru	XP_019804010.1	Sox	Sox-6
Mmus	NP_056573.1	Fox	Fox-E3	Hamp	XP_057604501.1	Sox	Sox-6
Mjav	XP_036877848.1	Fox	Fox-E3	Opri	XP_058519285.1	Sox	Sox-6
Casi	XP_006839917.1	Fox	Fox-E3	Rfer	XP_032975662.1	Sox	Sox-6
Dnov	XP_058160703.1	Fox	Fox-E3	Shar	XP_031797909.1	Sox	Sox-6
Lcat	XP_045401422.1	Fox	Fox-E3	Bbub	XP_04501884.1	Sox	Sox-6
Cimi	XP_017365578.1	Fox	Fox-E3	Dnov	XP_058161675.1	Sox	Sox-6
Cdid	XP_037672792.1	Fox	Fox-E3	Oana	XP_039767308.1	Sox	Sox-6
Opri	XP_004598546.1	Fox	Fox-E3	Hsap	NP_001139291.2	Sox	Sox-6
Scar	XP_047375686.1	Fox	Fox-E3	Bmus	XP_036717651.1	Sox	Sox-6
Oana	XP_001516678.2	Fox	Fox-D4	Mjav	XP_036882886.1	Sox	Sox-6
Pgig	XP_039744744.1	Fox	Fox-D4	Amel	XP_034501998.1	Sox	Sox-6
Hamp	XP_057576439.1	Fox	Fox-D4	Equa	XP_046493708.1	Sox	Sox-6
Pafr	XP_047625506.1	Fox	Fox-D4	Mdom	XP_007497087.2	Sox	Sox-6
Hsap	NP_001119806.1	Fox	Fox-D4	Cpor	XP_023419042.1	Sox	Sox-6
Emax	XP_049752498.1	Fox	Fox-D4	Pgig	XP_039707456.1	Sox	Sox-6
Cdid	XP_037706680.1	Fox	Fox-D4	Oafe	XP_007955001.1	Sox	Sox-6
Hsap	NP_954714.2	Fox	Fox-D4	Cdid	XP_037695265.1	Sox	Sox-6
Rfer	XP_032979406.1	Fox	Fox-D4	Casi	XP_006865846.1	Sox	Sox-6
Ttru	XP_033714036.1	Fox	Fox-D4	Cimi	XP_017381104.1	Sox	Sox-6

Tab. S8 continued from previous page

Sp. ID	Gene ID	Group	Annotation	Sp. ID	Gene ID	Group	Annotation
Ptig	XP_042820135.1	Fox	Fox-D4	Mmus	XP_036008726.1	Sox	Sox-6
Cdro	XP_010998361.2	Fox	Fox-D4	Ptig	XP_042815591.1	Sox	Sox-6
Mang	XP_045739597.1	Fox	Fox-D4	Pafr	XP_047632165.1	Sox	Sox-6
Scar	XP_047380997.1	Fox	Fox-D4	Tman	XP_023593283.1	Sox	Sox-6
Drot	XP_053785855.1	Fox	Fox-D4	Drot	XP_053780580.1	Sox	Sox-6
Hsap	NP_001078945.1	Fox	Fox-D4	Mang	XP_054368007.1	Sox	Sox-6
Amel	XP_034503132.1	Fox	Fox-D4	Csim	XP_014642191.1	Sox	Sox-6
Oafe	XP_007954771.1	Fox	Fox-D4	Scar	XP_047372506.1	Sox	Sox-6
Equa	XP_046519849.1	Fox	Fox-D4	Lcat	XP_045413549.1	Sox	Sox-6
Bbub	XP_025137372.3	Fox	Fox-D4	Cdro	XP_010975526.2	Sox	Sox-6
Dnov	XP_023444807.2	Fox	Fox-D4	Clup	XP_038425054.1	Sox	Sox-6
Bmus	XP_036712053.1	Fox	Fox-D4	Emax	XP_049725376.1	Sox	Sox-18
Hsap	NP_036316.1	Fox	Fox-D4	Casi	XP_006873390.1	Sox	-
Opri	XP_004591937.2	Fox	Fox-D4	Oafe	XP_007952883.1	Sox	-
Cimi	XP_017398902.1	Fox	Fox-D4	Mmus	NP_035577.1	Sox	Sox-8
Mdom	XP_001373972.1	Fox	Fox-D4	Opri	XP_004596857.2	Sox	-
Mmus	NP_032048.1	Fox	Fox-D4	Cpor	XP_003478479.1	Sox	-
Hsap	NP_997188.2	Fox	Fox-D4	Oana	XP_028913325.1	Sox	Sox-8
Hsap	NP_954586.4	Fox	Fox-D4	Ggal	NP_990062.2	Sox	-
Csim	XP_014650927.1	Fox	Fox-D4	Cpor	XP_023421073.1	Sox	-
Cpor	XP_013010584.1	Fox	Fox-D4	Cimi	XP_017378281.2	Sox	-
Mjav	XP_036847234.1	Fox	Fox-D4	Cdid	XP_037702077.1	Sox	-
Casi	XP_006863895.1	Fox	Fox-D4	Shar	XP_031794321.1	Sox	-
Mang	XP_045757341.1	Fox	-	Mdom	XP_016280870.1	Sox	-
Clup	XP_038401447.1	Fox	-	Ggal	XP_040553151.1	Sox	-
Amel	XP_019655962.1	Fox	-	Oana	XP_028913966.1	Sox	Sox-5
Clup	XP_038439326.1	Fox	-	Lcat	XP_045422920.1	Sox	-
Ggal	NP_001382914.1	Fox	Fox-C1				

Supplementary Table S9 – Complete set of DSFGs in *Drosophila*. For each gene, the species ID (Sp. ID) as in **Tab. S5**, the accession number (Gene ID), and the Possvm-based annotation are indicated.

Species ID	Gene ID	Group	Annotation	Species ID	Gene ID	Group	Annotation
Agam	XP_061505148.1	Dmrt	Dsx	Dbus	XP_017845192.1	Fox	Slp-1
Dere	XP_026836830.1	Dmrt	Dsx	Dgri	XP_001988485.1	Fox	Slp-1
Dkik	XP_041630485.1	Dmrt	Dsx	Dhyd	XP_023179855.2	Fox	Slp-1
Dmir	XP_017142875.1	Dmrt	Dsx	Dari	XP_017860496.1	Fox	Slp-1
Dgri	XP_043071131.1	Dmrt	Dsx	Dana	XP_001961572.2	Fox	Slp-1
Dsec	XP_002038750.1	Dmrt	Dsx	Dmir	XP_017155124.1	Fox	Slp-1
Dser	XP_020809854.1	Dmrt	Dsx	Dbp	XP_017105354.1	Fox	Slp-1
Dele	XP_017119779.1	Dmrt	Dsx	Dmel	NP_476730.1	Fox	Slp-1
Dalb	XP_034117252.2	Dmrt	Dsx	Dere	XP_001968625.1	Fox	Slp-1
Dwil	XP_023035845.1	Dmrt	Dsx	Dsec	XP_002037770.1	Fox	Slp-1
Dhyd	XP_023178918.2	Dmrt	Dsx	Dkik	XP_017021730.1	Fox	Slp-1
Dmel	NP_001262353.1	Dmrt	Dsx	Dpse	XP_001356670.4	Fox	Slp-1
Dpse	XP_033235910.1	Dmrt	Dsx	Dwil	XP_002065500.1	Fox	Slp-1
Dari	XP_017874634.1	Dmrt	Dsx	Dser	XP_020800881.1	Fox	Slp-1
Dbus	XP_017847641.1	Dmrt	Dsx	Dsuz	XP_016927184.1	Fox	Slp-1
Dbpip	XP_017088683.2	Dmrt	Dsx	Agam	XP_061514780.1	Fox	Slp-2
Dsuz	XP_036675224.1	Dmrt	Dsx	Dalb	XP_034100740.1	Fox	Slp-2
Dsuz	XP_036672758.1	Dmrt	Dsx	Dele	XP_017110569.1	Fox	Slp-2
Dana	XP_014766033.1	Dmrt	Dsx	Dere	XP_001968626.1	Fox	Slp-2
Dbus	XP_0178444894.1	Dmrt	Dmrt-99B	Dbus	XP_017845289.1	Fox	Slp-2
Dmel	NP_524549.1	Dmrt	Dmrt-99B	Dsec	XP_002037771.1	Fox	Slp-2
Dere	XP_001981330.1	Dmrt	Dmrt-99B	Dpse	XP_001356669.3	Fox	Slp-2
Dwil	XP_023034529.1	Dmrt	Dmrt-99B	Dmir	XP_017151517.1	Fox	Slp-2
Dkik	XP_017021833.1	Dmrt	Dmrt-99B	Dkik	XP_017021724.1	Fox	Slp-2
Dser	XP_020811324.1	Dmrt	Dmrt-99B	Dana	XP_001961573.1	Fox	Slp-2
Dalb	XP_051862400.1	Dmrt	Dmrt-99B	Dari	XP_017860500.1	Fox	Slp-2
Dsuz	XP_036674364.1	Dmrt	Dmrt-99B	Dser	XP_020800889.1	Fox	Slp-2
Dsec	XP_002037224.1	Dmrt	Dmrt-99B	Dbip	XP_017105682.2	Fox	Slp-2
Dele	XP_017131043.1	Dmrt	Dmrt-99B	Dhyd	XP_023179875.2	Fox	Slp-2
Dbib	XP_017099960.2	Dmrt	Dmrt-99B	Dsuz	XP_016926388.1	Fox	Slp-2
Dmir	XP_017145289.1	Dmrt	Dmrt-99B	Dwil	XP_023031666.1	Fox	Slp-2
Dpse	XP_001357766.3	Dmrt	Dmrt-99B	Dgri	XP_001998487.3	Fox	Slp-2
Dhyd	XP_023160825.2	Dmrt	Dmrt-99B	Dmel	NP_476834.1	Fox	Slp-2
Dari	XP_017869153.1	Dmrt	Dmrt-99B	Agam	NP_317309.5	Fox	Fd-3/Fd-59A
Dgri	XP_001996117.1	Dmrt	Dmrt-99B	Dbip	XP_017090886.2	Fox	Fd-3/Fd-59A
Dhyd	XP_023160826.2	Dmrt	Dmrt-99B	Dpse	XP_001361889.1	Fox	Fd-3/Fd-59A
Dana	XP_001964762.1	Dmrt	Dmrt-99B	Dmel	NP_523814.1	Fox	Fd-3/Fd-59A
Agam	XP_061501728.1	Dmrt	Dmrt-93B	Dsuz	XP_016927872.1	Fox	Fd-3/Fd-59A
Dalb	XP_034117959.2	Dmrt	Dmrt-93B	Dele	XP_017132051.1	Fox	Fd-3/Fd-59A
Dpse	XP_001360059.2	Dmrt	Dmrt-93B	Dser	XP_020803123.1	Fox	Fd-3/Fd-59A
Dkik	XP_017036725.1	Dmrt	Dmrt-93B	Dwil	XP_002061323.2	Fox	Fd-3/Fd-59A
Dbib	XP_017102685.2	Dmrt	Dmrt-93B	Dgri	XP_001987215.1	Fox	Fd-3/Fd-59A
Dbus	XP_017844858.1	Dmrt	Dmrt-93B	Dana	XP_001960803.1	Fox	Fd-3/Fd-59A
Dsuz	XP_036672900.1	Dmrt	Dmrt-93B	Dmir	XP_033248262.1	Fox	Fd-3/Fd-59A
Dser	XP_020817775.1	Dmrt	Dmrt-93B	Dalb	XP_034105309.1	Fox	Fd-3/Fd-59A
Dari	XP_017874225.1	Dmrt	Dmrt-93B	Dhyd	XP_030081270.1	Fox	Fd-3/Fd-59A
Dgri	XP_001990371.1	Dmrt	Dmrt-93B	Dmir	XP_017150782.2	Fox	Fd-3/Fd-59A
Dwil	XP_020273560.1	Dmrt	Dmrt-93B	Dkik	XP_017024427.1	Fox	Fd-3/Fd-59A
Dmel	NP_524428.1	Dmrt	Dmrt-93B	Dari	XP_017865679.1	Fox	Fd-3/Fd-59A
Dmir	XP_017140144.1	Dmrt	Dmrt-93B	Dsec	XP_002039987.1	Fox	Fd-3/Fd-59A
Dsec	XP_002044276.1	Dmrt	Dmrt-93B	Dere	XP_001976310.1	Fox	Fd-3/Fd-59A
Dele	XP_017118537.1	Dmrt	Dmrt-93B	Dbus	XP_017836437.1	Fox	Fd-3/Fd-59A
Dana	XP_001954937.1	Dmrt	Dmrt-93B	Agam	NP_315933.4	Fox	Crocodile
Dere	XP_001979344.2	Dmrt	Dmrt-93B	Dalb	XP_034109751.1	Fox	Crocodile
Dmel	NP_511146.2	Dmrt	Dmrt-11E	Dele	XP_017121437.1	Fox	Crocodile
Dsec	XP_002042926.2	Dmrt	Dmrt-11E	Dmel	NP_524202.1	Fox	Crocodile
Dwil	XP_020275263.1	Dmrt	Dmrt-11E	Dsec	XP_002040809.1	Fox	Crocodile
Dpse	XP_001355530.3	Dmrt	Dmrt-11E	Dsuz	XP_016933536.1	Fox	Crocodile
Dbib	XP_017089150.2	Dmrt	Dmrt-11E	Dari	XP_017864431.1	Fox	Crocodile
Dser	XP_020808350.1	Dmrt	Dmrt-11E	Dmir	XP_017139058.1	Fox	Crocodile
Dmir	XP_017136076.1	Dmrt	Dmrt-11E	Dhyd	XP_023173236.2	Fox	Crocodile
Dana	XP_032309371.1	Dmrt	Dmrt-11E	Dbib	XP_017095284.1	Fox	Crocodile
Dere	XP_015010652.2	Dmrt	Dmrt-11E	Dwil	XP_020261802.1	Fox	Crocodile
Dbus	XP_017850127.1	Dmrt	Dmrt-11E	Dser	XP_020817981.1	Fox	Crocodile
Dsuz	XP_016924080.2	Dmrt	Dmrt-11E	Dbus	XP_017841359.1	Fox	Crocodile
Dalb	XP_034099986.1	Dmrt	Dmrt-11E	Dpse	XP_001354188.2	Fox	Crocodile
Dgri	XP_043071903.1	Dmrt	Dmrt-11E	Dkik	XP_017029613.1	Fox	Crocodile
Dele	XP_017112743.2	Dmrt	Dmrt-11E	Dana	XP_001958351.1	Fox	Crocodile
Dhyd	XP_023180022.1	Dmrt	Dmrt-11E	Dgri	XP_001983802.1	Fox	Crocodile
Dkik	XP_017034619.2	Dmrt	Dmrt-11E	Dere	XP_001973629.1	Fox	Crocodile
Dari	XP_017869421.1	Dmrt	Dmrt-11E	Dmir	XP_017144084.1	Fox	Fkh
Agam	XP_310668.5	Dmrt	–	Dari	XP_017873677.1	Fox	Fkh
Dkik	XP_017022631.1	Fox	Fox-3F	Dpse	XP_033238940.1	Fox	Fkh
Dsec	XP_002037072.2	Fox	Fox-3F	Dwil	XP_002070503.2	Fox	Fkh
Dser	XP_02080708.1	Fox	Fox-3F	Dsuz	XP_036673542.1	Fox	Fkh
Dbib	XP_017087408.2	Fox	Fox-3F	Dalb	XP_051862733.1	Fox	Fkh
Dere	XP_026837609.1	Fox	Fox-3F	Dbib	XP_017092849.2	Fox	Fkh
Dana	XP_014759802.1	Fox	Fox-3F	Dbus	XP_017849701.2	Fox	Fkh
Dpse	XP_033238487.1	Fox	Fox-3F	Dsec	XP_002043027.2	Fox	Fkh
Dmir	XP_017145653.1	Fox	Fox-3F	Dgri	XP_001989953.3	Fox	Fkh
Dmel	NP_001356931.1	Fox	Fox-3F	Dana	XP_001955055.2	Fox	Fkh
Dele	XP_017112308.1	Fox	Fox-3F	Dkik	XP_017037933.1	Fox	Fkh
Dsuz	XP_016942113.1	Fox	Fox-3F	Dser	XP_020811220.1	Fox	Fkh
Dwil	XP_046868406.1	Fox	–	Dere	XP_001981453.3	Fox	Fkh
Dbus	XP_033150245.1	Fox	–	Agam	XP_061497286.1	Fox	Fkh
Dhyd	XP_030079965.1	Fox	–	Dmel	NP_001263038.1	Fox	Fkh
Dari	XP_017869433.1	Fox	–	Dele	XP_017131720.1	Fox	Fkh
Dgri	XP_043071916.1	Fox	–	Dhyd	XP_023165610.2	Fox	Fkh
Dalb	XP_051860438.1	Fox	–	Dalb	XP_034114947.1	Fox	Fd-5/Fd-96Cb
Agam	XP_061503465.1	Fox	Ches-1	Dele	XP_017123797.1	Fox	Fd-5/Fd-96Cb
Dbib	XP_017091473.2	Fox	Ches-1	Dgri	XP_001990839.1	Fox	Fd-5/Fd-96Cb
Dkik	XP_017022654.1	Fox	Ches-1	Dana	XP_001953691.1	Fox	Fd-5/Fd-96Cb
Dalb	XP_034097387.1	Fox	Ches-1	Dere	XP_001981873.2	Fox	Fd-5/Fd-96Cb
Dhyd	XP_023174414.2	Fox	Ches-1	Dkik	XP_017020777.1	Fox	Fd-5/Fd-96Cb
Dana	XP_032308510.1	Fox	Ches-1	Dwil	XP_023036287.1	Fox	Fd-5/Fd-96Cb
Dwil	XP_023030939.1	Fox	Ches-1	Dpse	XP_017103917.2	Fox	Fd-5/Fd-96Cb
Dser	XP_020805864.1	Fox	Ches-1	Dser	XP_020817575.1	Fox	Fd-5/Fd-96Cb

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Species ID	Gene ID	Group	Annotation	Species ID	Gene ID	Group	Annotation
Dgri	XP_032593899.1	Fox	Ches-1	Dsec	XP_002043869.2	Fox	Fd-5/Fd-96Cb
Dbus	XP_017852827.2	Fox	Ches-1	Dsuz	XP_016923664.2	Fox	Fd-5/Fd-96Cb
Dari	XP_017869569.1	Fox	Ches-1	Dmel	NP_524496.1	Fox	Fd-5/Fd-96Cb
Dpse	XP_03241472.1	Fox	Ches-1	Agam	XP_061497721.1	Fox	Fd-4/Fd-96Ca
Dmir	XP_03252440.1	Fox	Ches-1	Dana	XP_001953690.1	Fox	Fd-4/Fd-96Ca
Agam	XP_061505879.1	Fox	Jumeau	Dmel	NP_001287516.1	Fox	Fd-4/Fd-96Ca
Dalb	XP_034114927.1	Fox	Jumeau	Dsuz	XP_016923665.2	Fox	Fd-4/Fd-96Ca
Dgri	XP_001990048.1	Fox	Jumeau	Dpse	XP_033232491.1	Fox	Fd-4/Fd-96Ca
Dmir	XP_03244968.1	Fox	Jumeau	Dser	XP_020817686.1	Fox	Fd-4/Fd-96Ca
Dsec	XP_002031751.2	Fox	Jumeau	Dele	XP_017123798.1	Fox	Fd-4/Fd-96Ca
Dser	XP_020814051.1	Fox	Jumeau	Dkik	XP_017020776.1	Fox	Fd-4/Fd-96Ca
Dari	XP_017867184.1	Fox	Jumeau	Dwil	XP_002072971.1	Fox	Fd-4/Fd-96Ca
Dkik	XP_017026689.1	Fox	Jumeau	Dalb	XP_034114767.1	Fox	Fd-4/Fd-96Ca
Dana	XP_001952947.1	Fox	Jumeau	Dgri	XP_001990840.1	Fox	Fd-4/Fd-96Ca
Dmir	XP_03244970.1	Fox	Jumeau	Dbus	XP_017846227.1	Fox	Fd-4/Fd-96Ca
Dere	XP_001980679.1	Fox	Jumeau	Dbib	XP_017103916.2	Fox	Fd-4/Fd-96Ca
Dele	XP_017119152.1	Fox	Jumeau	Dhyd	XP_023172915.1	Fox	Fd-4/Fd-96Ca
Dsuz	XP_016944421.1	Fox	Jumeau	Dari	XP_017856594.1	Fox	Fd-4/Fd-96Ca
Dhyd	XP_023174343.2	Fox	Jumeau	Dsec	XP_002043867.1	Fox	Fd-4/Fd-96Ca
Dwil	XP_023034158.2	Fox	Jumeau	Dere	XP_001981874.1	Fox	Fd-4/Fd-96Ca
Dpse	XP_033233400.1	Fox	Jumeau	Dmir	XP_017141748.1	Fox	Fd-4/Fd-96Ca
Dpse	XP_015038643.2	Fox	Jumeau	Agam	XP_061507474.1	Fox	-
Dmir	XP_03244969.1	Fox	Jumeau	Agam	XP_061497423.1	Fox	-
Dmel	NP_524302.1	Fox	Jumeau	Agam	XP_312480.5	Fox	-
Dbib	XP_017100743.2	Fox	Jumeau	Dana	XP_044571024.1	Sox	Sox-21B
Dbus	XP_017845708.1	Fox	Jumeau	Dari	XP_017862175.1	Sox	Sox-21B
Dbib	XP_043069566.1	Fox	Hcm-1	Dhyd	XP_03080409.1	Sox	Sox-21B
Dsec	XP_002044492.1	Fox	Hcm-1	Dbus	XP_033149417.1	Sox	Sox-21B
Dere	XP_026838997.1	Fox	Hcm-1	Dere	XP_001972712.1	Sox	Sox-21B
Dhyd	XP_023165547.2	Fox	Hcm-1	Dmir	XP_033243279.1	Sox	Sox-21B
Dele	XP_041564577.1	Fox	Hcm-1	Dmel	NP_0012161829.1	Sox	Sox-21B
Dsuz	XP_036678464.1	Fox	Hcm-1	Dele	XP_017128640.1	Sox	Sox-21B
Dalb	XP_051862243.1	Fox	Hcm-1	Dpse	XP_002134931.3	Sox	Sox-21B
Dana	XP_032311767.1	Fox	Hcm-1	Dwil	XP_02304010.1	Sox	Sox-21B
Dpse	XP_033237266.1	Fox	Hcm-1	Dsec	XP_002030465.1	Sox	Sox-21B
Dmel	NP_726538.1	Fox	Hcm-1	Dalb	XP_034107607.1	Sox	Sox-21B
Dmir	XP_017155587.1	Fox	Hcm-1	Dbib	XP_017090122.2	Sox	Sox-21B
Dser	XP_020809944.1	Fox	Hcm-1	Dsec	XP_061509694.1	Sox	Sox-21B
Dbus	XP_017853536.1	Fox	Hcm-1	Dkik	XP_017034105.1	Sox	Sox-21B
Dwil	XP_023035694.1	Fox	Hcm-1	Dgri	XP_032598957.1	Sox	Sox-21B
Dkik	XP_017037372.1	Fox	Hcm-1	Dser	XP_020816310.1	Sox	Sox-21B
Dwil	XP_002072563.1	Fox	Fd-102C	Dgri	XP_032590737.1	Sox	Sox-14
Dhyd	XP_023163080.1	Fox	Fd-102C	Dhyd	XP_023171730.2	Sox	Sox-14
Dbib	XP_017098296.2	Fox	Fd-102C	Dbus	XP_017836915.1	Sox	Sox-14
Dmel	NP_651951.1	Fox	Fd-102C	Dpse	XP_001360869.4	Sox	Sox-14
Dsuz	XP_036671526.1	Fox	Fd-102C	Dkik	XP_017018688.1	Sox	Sox-14
Dele	XP_017126055.1	Fox	Fd-102C	Dbib	XP_017090231.2	Sox	Sox-14
Dbus	XP_017853523.1	Fox	Fd-102C	Dser	XP_020817997.1	Sox	Sox-14
Dsec	XP_002043702.1	Fox	Fd-102C	Dmir	XP_033246935.1	Sox	Sox-14
Agam	XP_061505343.1	Fox	Fd-102C	Dsec	XP_002040195.2	Sox	Sox-14
Dere	XP_001982706.1	Fox	Fd-102C	Dmel	NP_001286801.1	Sox	Sox-14
Dmir	XP_017155765.1	Fox	Fd-102C	Dmir	XP_017149652.2	Sox	Sox-14
Dana	XP_032308054.1	Fox	Fd-102C	Dsuz	XP_016926392.1	Sox	Sox-14
Dalb	XP_034112224.1	Fox	Fd-102C	Dari	XP_017868666.1	Sox	Sox-14
Dpse	XP_015044407.1	Fox	Fd-102C	Dalb	XP_034106594.1	Sox	Sox-14
Dkik	XP_041632589.1	Fox	Fd-102C	Dere	XP_001976516.1	Sox	Sox-14
Dgri	XP_043071810.1	Fox	Fd-102C	Dwil	XP_002061086.2	Sox	Sox-14
Dser	XP_020809941.1	Fox	Fd-102C	Dgri	XP_017125359.1	Sox	Sox-14
Dari	XP_017869379.1	Fox	Fd-102C	Dose	XP_001959457.2	Sox	Sox-14
Agam	XP_061502996.1	Fox	Fox-P	Dana	XP_001355471.1	Sox	Dichaete
Dhyd	XP_023172914.1	Fox	Fox-P	Dpse	XP_001355471.1	Sox	Dichaete
Dbus	XP_033148941.1	Fox	Fox-P	Del	XP_017128756.1	Sox	Dichaete
Dbib	XP_043067838.1	Fox	Fox-P	Dmir	XP_017138810.1	Sox	Dichaete
Dgri	XP_032593352.2	Fox	Fox-P	Dalb	XP_034104310.1	Sox	Dichaete
Dana	XP_044571198.1	Fox	Fox-P	Dsec	XP_002030467.1	Sox	Dichaete
Dalb	XP_051862673.1	Fox	Fox-P	Dgri	XP_001984254.1	Sox	Dichaete
Dmir	XP_033244694.1	Fox	Fox-P	Dana	XP_001956432.1	Sox	Dichaete
Dmel	NP_001247011.1	Fox	Fox-P	Dmel	NP_524066.1	Sox	Dichaete
Dere	XP_015009985.2	Fox	Fox-P	Dere	XP_001972713.1	Sox	Dichaete
Dele	XP_017117570.1	Fox	Fox-P	Dbib	XP_017089891.1	Sox	Dichaete
Dsuz	XP_016930987.1	Fox	Fox-P	Dgri	XP_061514536.1	Sox	Dichaete
Dpse	XP_033241607.1	Fox	Fox-P	Dbus	XP_017841081.1	Sox	Dichaete
Dwil	XP_046865977.1	Fox	Fox-P	Dkik	XP_017034102.1	Sox	Dichaete
Dsec	XP_032577368.1	Fox	Fox-P	Dwil	XP_002061712.1	Sox	Dichaete
Dser	XP_020813734.1	Fox	Fox-P	Dari	XP_017864416.1	Sox	Dichaete
Dwil	XP_002072142.2	Fox	Fox-O	Dhyd	XP_023173062.1	Sox	Dichaete
Agam	XP_061497073.1	Fox	Fox-O	Dser	XP_020815889.1	Sox	Dichaete
Dari	XP_017859446.1	Fox	Fox-O	Dsuz	XP_016933433.1	Sox	Dichaete
Dsuz	XP_036674215.1	Fox	Fox-O	Dere	XP_001975666.1	Sox	Sox-15
Dsuz	XP_036672729.1	Fox	Fox-O	Dele	XP_017133202.1	Sox	Sox-15
Dgri	XP_032595778.1	Fox	Fox-O	Dbus	XP_017835743.2	Sox	Sox-15
Dbib	XP_017091959.2	Fox	Fox-O	Dwil	XP_023030517.1	Sox	Sox-15
Dmir	XP_017143568.1	Fox	Fox-O	Dalb	XP_051860622.1	Sox	Sox-15
Dalb	XP_034113157.1	Fox	Fox-O	Dana	XP_001959479.1	Sox	Sox-15
Dsec	XP_032576567.1	Fox	Fox-O	Dbib	XP_017090318.2	Sox	Sox-15
Dhyd	XP_023171342.1	Fox	Fox-O	Dari	XP_017867550.1	Sox	Sox-15
Dbus	XP_033149836.1	Fox	Fox-O	Dmir	XP_017149731.1	Sox	Sox-15
Dere	XP_026839669.1	Fox	Fox-O	Dsec	XP_002033808.1	Sox	Sox-15
Dkik	XP_017026758.1	Fox	Fox-O	Dgri	XP_061509362.1	Sox	Sox-15
Dser	XP_020817293.1	Fox	Fox-O	Dmel	NP_523739.2	Sox	Sox-15
Dpse	XP_03236382.1	Fox	Fox-O	Dser	XP_020802427.1	Sox	Sox-15
Dele	XP_041565035.1	Fox	Fox-O	Dhyd	XP_030081376.1	Sox	Sox-15
Dana	XP_032311428.1	Fox	Fox-O	Dsuz	XP_016927906.1	Sox	Sox-15
Dmel	NP_650330.3	Fox	Fox-O	Dpse	XP_001361762.2	Sox	Sox-15
Agam	XP_001688749.2	Fox	Binioi	Dkik	XP_017019703.1	Sox	Sox-15
Dmel	NP_523950.2	Fox	Binioi	Dgri	XP_001987303.1	Sox	Sox-15
Dalb	XP_034105454.1	Fox	Binioi	Dsec	XP_002030464.1	Sox	Sox-21A
Dser	XP_020817612.1	Fox	Binioi	Dgri	XP_001984256.1	Sox	Sox-21A
Dele	XP_017121425.1	Fox	Binioi	Dsuz	XP_016934639.1	Sox	Sox-21A
Dkik	XP_017017381.1	Fox	Binioi	Dmir	XP_017138458.1	Sox	Sox-21A
Dana	XP_001958356.2	Fox	Binioi	Dkik	XP_017034345.1	Sox	Sox-21A
Dari	XP_017864430.1	Fox	Binioi	Dbus	XP_017840298.2	Sox	Sox-21A

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Species ID	Gene ID	Group	Annotation	Species ID	Gene ID	Group	Annotation
Dbus	XP_017841230.1	Fox	Biniou	Dari	XP_017862178.1	Sox	Sox-21A
Dwil	XP_002061803.1	Fox	Biniou	Dalb	XP_034108985.1	Sox	Sox-21A
Dgri	XP_001983803.1	Fox	Biniou	Dser	XP_020816311.1	Sox	Sox-21A
Dere	XP_001971514.2	Fox	Biniou	Agam	XP_061513404.1	Sox	Sox-21A
Dsuz	XP_036672243.1	Fox	Biniou	Dele	XP_017128154.1	Sox	Sox-21A
Dpse	XP_002134730.3	Fox	Biniou	Dana	XP_032310341.1	Sox	Sox-21A
Dhyd	XP_030081482.1	Fox	Biniou	Dhyd	XP_023173058.2	Sox	Sox-21A
Dbip	XP_017095285.2	Fox	Biniou	Dere	XP_001972711.1	Sox	Sox-21A
Dmir	XP_017139097.1	Fox	Biniou	Dbip	XP_043070009.1	Sox	Sox-21A
Dsec	XP_002035641.1	Fox	Biniou	Dwil	XP_023033998.1	Sox	Sox-21A
Agam	XP_061503212.1	Fox	Fd-2/Fox-L1	Dmel	NP_001261827.1	Sox	Sox-21A
Dmel	NP_001246609.1	Fox	Fd-2/Fox-L1	Dpse	XP_033239763.1	Sox	Sox-21A
Dhyd	XP_023178350.2	Fox	Fd-2/Fox-L1	Dwil	XP_002066518.2	Sox	Sox-N
Dbus	XP_017840363.1	Fox	Fd-2/Fox-L1	Dser	XP_020807900.1	Sox	Sox-N
Dkik	XP_017019394.1	Fox	Fd-2/Fox-L1	Dsuz	XP_036678170.1	Sox	Sox-N
Dser	XP_020807718.1	Fox	Fd-2/Fox-L1	Dbus	XP_033150665.1	Sox	Sox-N
Dari	XP_017863042.1	Fox	Fd-2/Fox-L1	Dsec	XP_002036270.2	Sox	Sox-N
Dsec	XP_002035260.1	Fox	Fd-2/Fox-L1	Dgri	XP_001993541.2	Sox	Sox-N
Dere	XP_001971815.2	Fox	Fd-2/Fox-L1	Dele	XP_017121213.1	Sox	Sox-N
Dalb	XP_051860220.1	Fox	Fd-2/Fox-L1	Dari	XP_017860249.1	Sox	Sox-N
Dana	XP_001956070.1	Fox	Fd-2/Fox-L1	Dalb	XP_051858933.1	Sox	Sox-N
Dsuz	XP_016932589.2	Fox	Fd-2/Fox-L1	Dkik	XP_017024634.1	Sox	Sox-N
Dele	XP_017126783.1	Fox	Fd-2/Fox-L1	Agam	XP_061516479.1	Sox	Sox-N
Dgri	XP_001983216.1	Fox	Fd-2/Fox-L1	Dmir	XP_017154597.1	Sox	Sox-N
Dpse	XP_001352427.1	Fox	Fd-2/Fox-L1	Dbip	XP_017099296.2	Sox	Sox-N
Dwil	XP_002062380.1	Fox	Fd-2/Fox-L1	Dere	XP_026834908.1	Sox	Sox-N
Dbip	XP_017098174.2	Fox	Fd-2/Fox-L1	Dmel	NP_001260269.1	Sox	Sox-N
Dmir	XP_017137846.1	Fox	Fd-2/Fox-L1	Dhyd	XP_023172210.2	Sox	Sox-N
Dere	XP_001977848.1	Fox	Fd-19B	Dpse	XP_001355808.4	Sox	Sox-N
Dhyd	XP_030079337.1	Fox	Fd-19B	Dana	XP_001962613.2	Sox	Sox-N
Dmel	NP_608369.1	Fox	Fd-19B	Dbus	XP_017845374.1	Sox	Sox-100B
Dsec	XP_002039497.1	Fox	Fd-19B	Dari	XP_017874440.1	Sox	Sox-100B
Dsuz	XP_016923819.2	Fox	Fd-19B	Agam	XP_061509035.1	Sox	Sox-100B
Dele	XP_041565246.1	Fox	Fd-19B	Dhyd	XP_023168522.1	Sox	Sox-100B
Dpse	XP_033239022.1	Fox	Fd-19B	Dkik	XP_017037336.1	Sox	Sox-100B
Dmir	XP_017134931.2	Fox	Fd-19B	Dana	XP_001954720.1	Sox	Sox-100B
Dari	XP_017869476.1	Fox	Fd-19B	Dele	XP_017122478.1	Sox	Sox-100B
Dgri	XP_043072169.1	Fox	Fd-19B	Dser	XP_020811779.1	Sox	Sox-100B
Dalb	XP_034101064.2	Fox	Fd-19B	Dbip	XP_017088796.2	Sox	Sox-100B
Dwil	XP_046867097.1	Fox	Fd-19B	Dmir	XP_017145320.1	Sox	Sox-100B
Dser	XP_020808848.1	Fox	Fd-19B	Dalb	XP_051863654.1	Sox	Sox-100B
Dkik	XP_017017955.1	Fox	Fd-19B	Dmel	NP_651839.1	Sox	Sox-100B
Dbus	XP_033150291.1	Fox	Fd-19B	Dsec	XP_002037434.1	Sox	Sox-100B
Agam	XP_061512060.1	Fox	-	Dere	XP_001981122.1	Sox	Sox-100B
Dalb	XP_034098689.1	Fox	-	Dsuz	XP_016935593.2	Sox	Sox-100B
Dgri	XP_032595015.1	Fox	-	Dwil	XP_023035523.1	Sox	Sox-100B
Dalb	XP_034098925.1	Fox	Slp-1	Dgri	XP_032597554.1	Sox	Sox-100B
Dele	XP_017110574.1	Fox	Slp-1	Dpse	XP_001357577.3	Sox	Sox-100B

Supplementary Table S10 – All the enriched GO terms for Group 1 and Group 2 genes of bivalves, mammals, and *Drosophila*.

Dataset	Group of genes	GO.id	Term	Annotated	Significant	Expected	Classic Fisher
Bivalvia	Group 1 + Group 2	GO:0060255	regulation of macromolecule metabolic process	737	59	31.91	0.0453
Bivalvia	Group 1 + Group 2	GO:0080090	regulation of primary metabolic process	673	53	29.14	0.0182
Bivalvia	Group 1 + Group 2	GO:0019219	regulation of nucleobase-containing compound metabolic process	541	41	23.42	0.0239
Bivalvia	Group 1 + Group 2	GO:0003551	DNA-templated transcription	571	39	24.72	0.0377
Bivalvia	Group 1 + Group 2	GO:0032774	RNA biosynthetic process	579	39	25.07	0.0449
Bivalvia	Group 1 + Group 2	GO:0051252	regulation of RNA metabolic process	517	37	22.38	0.0272
Bivalvia	Group 1 + Group 2	GO:0003535	regulation of DNA-templated transcription	490	35	21.22	0.0375
Bivalvia	Group 1 + Group 2	GO:2001141	regulation of RNA biosynthetic process	491	35	21.26	0.0384
Bivalvia	Group 1 + Group 2	GO:006950	response to stress	370	33	16.02	0.0195
Bivalvia	Group 1 + Group 2	GO:0032502	developmental process	261	27	11.30	0.0445
Bivalvia	Group 1 + Group 2	GO:0006468	protein phosphorylation	345	23	14.94	0.0248
Bivalvia	Group 1 + Group 2	GO:0031325	positive regulation of cellular metabolic process	125	17	5.41	0.0080
Bivalvia	Group 1 + Group 2	GO:0010604	positive regulation of macromolecule metabolic process	151	17	6.54	0.0405
Bivalvia	Group 1 + Group 2	GO:0051172	negative regulation of nitrogen compound metabolic process	117	16	5.07	0.0081
Bivalvia	Group 1 + Group 2	GO:0051173	positive regulation of nitrogen compound metabolic process	137	15	5.93	0.0245
Bivalvia	Group 1 + Group 2	GO:0066310	DNA recombination	66	14	2.86	0.0009
Bivalvia	Group 1 + Group 2	GO:0048513	animal organ development	83	12	3.59	0.0409
Bivalvia	Group 1 + Group 2	GO:0010629	negative regulation of gene expression	78	11	3.38	0.0005
Bivalvia	Group 1 + Group 2	GO:0023051	regulation of signal transduction	133	11	5.76	0.0287
Bivalvia	Group 1 + Group 2	GO:0045934	negative regulation of nucleobase-containing compound metabolic process	64	11	2.77	0.0364
Bivalvia	Group 1 + Group 2	GO:0009605	response to external stimulus	90	11	3.90	0.0454
Bivalvia	Group 1 + Group 2	GO:0044419	biological process involved in interspecies interaction between organisms	63	11	2.73	0.0476
Bivalvia	Group 1 + Group 2	GO:0006915	apoptotic process	95	10	4.11	0.0077
Bivalvia	Group 1 + Group 2	GO:0009966	regulation of signal transduction	120	10	5.20	0.0345
Bivalvia	Group 1 + Group 2	GO:0006417	regulation of translation	52	9	2.25	0.0003
Bivalvia	Group 1 + Group 2	GO:0045892	negative regulation of DNA-templated transcription	59	9	2.55	0.0297
Bivalvia	Group 1 + Group 2	GO:1902679	negative regulation of RNA biosynthetic process	59	9	2.55	0.0297
Bivalvia	Group 1 + Group 2	GO:0009607	response to biotic stimulus	55	9	2.38	0.0321
Bivalvia	Group 1 + Group 2	GO:0051253	negative regulation of RNA metabolic process	61	9	2.64	0.0372
Bivalvia	Group 1 + Group 2	GO:0006952	defense response	58	9	2.51	0.0416
Bivalvia	Group 1 + Group 2	GO:0006302	double-strand break repair	52	9	2.25	0.0486
Bivalvia	Group 1 + Group 2	GO:0080134	regulation of response to stress	43	8	1.86	0.0067
Bivalvia	Group 1 + Group 2	GO:0010564	regulation of cell cycle process	70	8	3.03	0.0102
Bivalvia	Group 1 + Group 2	GO:0042981	regulation of apoptotic process	72	8	3.12	0.0121
Bivalvia	Group 1 + Group 2	GO:0043067	regulation of programmed cell death	61	8	2.64	0.0400
Bivalvia	Group 1 + Group 2	GO:0048584	positive regulation of response to stimulus	58	9	2.51	0.0416
Bivalvia	Group 1 + Group 2	GO:0006310	cellular response to organic substance	52	7	2.25	0.0066
Bivalvia	Group 1 + Group 2	GO:0010628	import into cell	34	7	1.47	0.0266
Bivalvia	Group 1 + Group 2	GO:0045944	positive regulation of gene expression	38	6	1.65	0.0054
Bivalvia	Group 1 + Group 2	GO:1901987	positive regulation of transcription by RNA polymerase II	29	6	1.26	0.0237
Bivalvia	Group 1 + Group 2	GO:2000779	regulation of cell cycle phase transition	11	6	0.48	0.0243
Bivalvia	Group 1 + Group 2	GO:0051247	regulation of double-strand break repair	54	6	2.34	0.0282
Bivalvia	Group 1 + Group 2	GO:0051248	positive regulation of protein metabolic process	55	6	2.38	0.0305
Bivalvia	Group 1 + Group 2	GO:0098657	negative regulation of protein metabolic process	56	6	2.42	0.0330
Bivalvia	Group 1 + Group 2	GO:1902531	regulation of intracellular signal transduction	59	6	2.55	0.0412
Bivalvia	Group 1 + Group 2	GO:0044770	cell cycle phase transition	35	6	1.52	0.0467
Bivalvia	Group 1 + Group 2	GO:0000122	negative regulation of transcription by RNA polymerase II	31	5	1.34	0.0099
Bivalvia	Group 1 + Group 2	GO:0006402	mRNA catabolic process	35	5	1.52	0.0164
Bivalvia	Group 1 + Group 2	GO:1901990	regulation of mitotic cell cycle phase transition	18	5	0.78	0.0244
Bivalvia	Group 1 + Group 2	GO:1901698	response to nitrogen compound	5	4	0.22	0.0054
Bivalvia	Group 1 + Group 2	GO:0006401	RNA catabolic process	41	5	1.78	0.0307
Bivalvia	Group 1 + Group 2	GO:0030155	regulation of cell adhesion	44	5	0.91	0.0115
Bivalvia	Group 1 + Group 2	GO:0048568	embryonic organ development	11	4	0.48	0.0010
Bivalvia	Group 1 + Group 2	GO:0007517	muscle organ development	12	4	0.52	0.0113
Bivalvia	Group 1 + Group 2	GO:0051607	defense response to virus	13	4	0.56	0.0018
Bivalvia	Group 1 + Group 2	GO:0010569	regulation of double-strand break repair via homologous recombination	5	4	0.22	0.0054
Bivalvia	Group 1 + Group 2	GO:0042274	ribosomal small subunit biogenesis	21	4	0.91	0.0401
Bivalvia	Group 1 + Group 2	GO:0043066	negative regulation of apoptotic process	28	4	1.21	0.0310
Bivalvia	Group 1 + Group 2	GO:0043069	negative regulation of programmed cell death	29	4	1.26	0.0348
Bivalvia	Group 1 + Group 2	GO:0016477	cell migration	29	4	1.30	0.0388
Bivalvia	Group 1 + Group 2	GO:0032101	regulation of response to external stimulus	30	4	1.30	0.0388
Bivalvia	Group 1 + Group 2	GO:0050769	positive regulation of neurogenesis	5	3	0.22	0.0008
Bivalvia	Group 1 + Group 2	GO:0007368	determination of left/right symmetry	7	3	0.30	0.0025
Bivalvia	Group 1 + Group 2	GO:0001819	positive regulation of cytokine production	7	3	0.30	0.0025
Bivalvia	Group 1 + Group 2	GO:0070192	chromosome organization involved in meiotic cell cycle	7	3	0.35	0.035
Bivalvia	Group 1 + Group 2	GO:0045132	meiotic chromosome segregation	8	8		

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Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Bivalvia	Group 1 + Group 2	GO:0042326	negative regulation of phosphorylation	10	3	0.43	0.0077
Bivalvia	Group 1 + Group 2	GO:0082856	negative regulation of cell population proliferation	10	3	0.43	0.0077
Bivalvia	Group 1 + Group 2	GO:0022604	regulation of cell morphogenesis	10	3	0.43	0.0077
Bivalvia	Group 1 + Group 2	GO:0001894	tissue homeostasis	10	3	0.43	0.0077
Bivalvia	Group 1 + Group 2	GO:0033007	heart morphogenesis	10	3	0.43	0.0077
Bivalvia	Group 1 + Group 2	GO:0051093	negative regulation of developmental process	11	3	0.48	0.0102
Bivalvia	Group 1 + Group 2	GO:0001501	skelet system development	11	3	0.48	0.0102
Bivalvia	Group 1 + Group 2	GO:0042327	positive regulation of phosphorylation	12	3	0.52	0.0132
Bivalvia	Group 1 + Group 2	GO:0010562	positive regulation of phosphorus metabolic process	13	3	0.56	0.0166
Bivalvia	Group 1 + Group 2	GO:0045010	actin nucleation	13	3	0.56	0.0166
Bivalvia	Group 1 + Group 2	GO:0045937	positive regulation of phosphate metabolic process	13	3	0.56	0.0166
Bivalvia	Group 1 + Group 2	GO:0007127	meiosis I	14	3	0.61	0.0205
Bivalvia	Group 1 + Group 2	GO:0031400	negative regulation of protein modification process	14	3	0.61	0.0205
Bivalvia	Group 1 + Group 2	GO:0061982	meiosis I cell cycle process	14	3	0.61	0.0205
Bivalvia	Group 1 + Group 2	GO:0097190	apoptotic signaling pathway	16	3	0.69	0.0296
Bivalvia	Group 1 + Group 2	GO:0040008	regulation of growth	16	3	0.69	0.0296
Bivalvia	Group 1 + Group 2	GO:0051345	negative regulation of hydrolase activity	17	3	0.74	0.0348
Bivalvia	Group 1 + Group 2	GO:0010257	NADH dehydrogenase complex assembly	17	3	0.74	0.0348
Bivalvia	Group 1 + Group 2	GO:0032981	mitochondrial respiratory chain complex I assembly	17	3	0.74	0.0348
Bivalvia	Group 1 + Group 2	GO:0032880	regulation of protein localization	18	3	0.78	0.0405
Bivalvia	Group 1 + Group 2	GO:005976	polysaccharide metabolic process	18	3	0.78	0.0405
Bivalvia	Group 1 + Group 2	GO:0048729	tissue morphogenesis	19	3	0.82	0.0466
Bivalvia	Group 1 + Group 2	GO:0018022	peptidyl-lysine methylation	19	3	0.82	0.0466
Bivalvia	Group 1 + Group 2	GO:0000041	transition metal ion transport	19	3	0.82	0.0466
Bivalvia	Group 1 + Group 2	GO:0042488	Cdc42 protein signal transduction	2	2	0.09	0.0019
Bivalvia	Group 1 + Group 2	GO:0022600	digestive system process	2	2	0.09	0.0019
Bivalvia	Group 1 + Group 2	GO:007097	nuclear migration	2	2	0.09	0.0019
Bivalvia	Group 1 + Group 2	GO:0032232	negative regulation of actin filament bundle assembly	2	2	0.09	0.0019
Bivalvia	Group 1 + Group 2	GO:1905168	positive regulation of double-strand break repair via homologous recombination	2	2	0.09	0.0019
Bivalvia	Group 1 + Group 2	GO:0002064	epithelial cell development	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:0061383	tracheal morphogenesis	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:0010830	regulation of myoblast differentiation	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:0010833	telomere maintenance via telomerase lengthening	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:0000959	mitochondrial RNA metabolic process	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:0033617	mitochondrial cytochrome c oxidase assembly	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:2000179	positive regulation of neural precursor cell proliferation	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:0050777	negative regulation of immune response	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:0007095	mitotic G2 DNA damage checkpoint signaling	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:2000736	regulation of stem cell differentiation	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:0016233	telomere capping	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:0005910	negative regulation of DNA recombination	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:0051701	biological process involved in interaction with host	3	2	0.13	0.0054
Bivalvia	Group 1 + Group 2	GO:0050777	response to antibiotic	4	2	0.17	0.0106
Bivalvia	Group 1 + Group 2	GO:0046677	regulation of organ growth	4	2	0.17	0.0106
Bivalvia	Group 1 + Group 2	GO:0046620	regulation of stem cell differentiation	4	2	0.17	0.0106
Bivalvia	Group 1 + Group 2	GO:0030514	negative regulation of BMP signaling pathway	4	2	0.17	0.0106
Bivalvia	Group 1 + Group 2	GO:1901678	iron coordination entity transport	4	2	0.17	0.0106
Bivalvia	Group 1 + Group 2	GO:0007519	skeletal muscle tissue development	4	2	0.17	0.0106
Bivalvia	Group 1 + Group 2	GO:0032507	maintenance of protein location in cell	4	2	0.17	0.0106
Bivalvia	Group 1 + Group 2	GO:00097416	synapse assembly	4	2	0.17	0.0106
Bivalvia	Group 1 + Group 2	GO:0098781	ncRNA transcription	4	2	0.17	0.0106
Bivalvia	Group 1 + Group 2	GO:0035023	regulation of Rho protein signal transduction	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0061371	atrioventricular valve formation	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0003181	atrioventricular valve morphogenesis	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0090101	negative regulation of transmembrane receptor protein serine/threonine kinase signaling pathway	4	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0010001	glial cell differentiation	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0090288	negative regulation of cellular response to growth factor stimulus	4	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0030510	regulation of BMP signaling pathway	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0001947	determination of heart left/right asymmetry	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0035025	gliogenesis	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0042063	negative regulation of cell adhesion	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0042026	protein refolding	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0007129	homologous chromosome pairing at meiosis	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0031717	heart looping	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0030968	embryonic heart tube development	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:003188	endothelial reticulum unfolded protein response	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0003171	heart valve formation	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:003179	atrioventricular valve development	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0031711	heart valve morphogenesis	5	2	0.22	0.0171

Tab. S10 continued from previous page

Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Bivalvia	Group 1 + Group 2	GO:0003143	embryonic heart tube morphogenesis	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0009287	regulation of cellular response to growth factor stimulus	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0006538	skeletal muscle organ development	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0034620	cellular response to unfolded protein	5	2	0.22	0.0171
Bivalvia	Group 1 + Group 2	GO:0030509	BMP signaling pathway	6	2	0.26	0.0250
Bivalvia	Group 1 + Group 2	GO:0006360	transcription by RNA polymerase I	6	2	0.26	0.0250
Bivalvia	Group 1 + Group 2	GO:0045198	maintenance of protein location	6	2	0.26	0.0250
Bivalvia	Group 1 + Group 2	GO:0045143	homologous chromosome segregation	6	2	0.26	0.0250
Bivalvia	Group 1 + Group 2	GO:0001889	liver development	6	2	0.26	0.0250
Bivalvia	Group 1 + Group 2	GO:0071772	response to BMP	6	2	0.26	0.0250
Bivalvia	Group 1 + Group 2	GO:0001773	cellular response to BMP stimulus	6	2	0.26	0.0250
Bivalvia	Group 1 + Group 2	GO:0009092	regulation of transmembrane receptor protein serine/threonine kinase signaling pathway	6	2	0.26	0.0250
Bivalvia	Group 1 + Group 2	GO:0003170	heart valve development	6	2	0.26	0.0250
Bivalvia	Group 1 + Group 2	GO:0003012	muscle system process	6	2	0.26	0.0250
Bivalvia	Group 1 + Group 2	GO:0030490	maturatation of SSU-rRNA	7	2	0.30	0.0340
Bivalvia	Group 1 + Group 2	GO:0018105	peptidyl-serine phosphorylation	7	2	0.30	0.0340
Bivalvia	Group 1 + Group 2	GO:0032465	regulation of cytokinesis	7	2	0.30	0.0340
Bivalvia	Group 1 + Group 2	GO:0061448	connective tissue development	7	2	0.30	0.0340
Bivalvia	Group 1 + Group 2	GO:0018209	peptidyl-serine modification	7	2	0.30	0.0340
Bivalvia	Group 1 + Group 2	GO:0032102	negative regulation of response to external stimulus	7	2	0.30	0.0340
Bivalvia	Group 1 + Group 2	GO:0061008	hepatobiliary system development	7	2	0.30	0.0340
Bivalvia	Group 1 + Group 2	GO:0048638	regulation of developmental growth	7	2	0.30	0.0340
Bivalvia	Group 1 + Group 2	GO:0010212	response to ionizing radiation	7	2	0.30	0.0340
Bivalvia	Group 1 + Group 2	GO:0034329	cell junction assembly	7	2	0.30	0.0340
Bivalvia	Group 1 + Group 2	GO:1901652	response to peptide	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:0048732	gland development	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:0033157	regulation of intracellular protein transport	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:0051302	regulation of cell division	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:0001822	kidney development	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:0050808	synapse organization	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:0033967	cellular response to topologically incorrect protein	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:0031032	actomyosin structure organization	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:0001503	ossification	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:000271	polysaccharide biosynthetic process	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:008593	regulation of Notch signaling pathway	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:0045596	negative regulation of cell differentiation	8	2	0.35	0.0440
Bivalvia	Group 1 + Group 2	GO:0050691	regulation of defense response to virus by host	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1900220	semaphorin-plexin signaling pathway involved in bone trabecula morphogenesis	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0046621	negative regulation of organ growth	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048799	animal organ maturation	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0046402	O antigen metabolic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0052572	positive regulation of skeletal muscle fiber development	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0015743	positive regulation of extrinsic apoptotic signaling pathway via death domain receptors	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0043628	malate transport	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1904862	regulatory ncRNA 3'-end processing	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:000963	inhibitory synapse assembly	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048799	mitochondrial RNA processing	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0002230	positive regulation of defense response to virus by host	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048742	semaphorin-plexin signaling pathway involved in axon guidance	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0033688	regulation of osteoblast proliferation	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0033689	beta-lactam antibiotic catabolic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:001100	negative regulation of exit from mitosis	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:008625	extrinsic apoptotic signaling pathway via death domain receptors	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0071423	osteoblast proliferation	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0008653	lipopolysaccharide metabolic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0030653	beta-lactam antibiotic catabolic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0070212	protein poly-ADP-ribosylation	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0033687	maleate transmembrane transport	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1902287	protection from non-homologous end joining at telomere	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0031848	retrotransposon silencing	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0061668	mitochondrial ribosome assembly	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0035418	protein localization to synapse	1	1	0.04	0.0433

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Dataset	Group of genes	GO ID	Term	Annotated	Significant	Expected	Classic Fisher
Bivalvia	Group 1 + Group 2	GO:0061430	bone trabecula morphogenesis	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0006356	regulation of transcription by RNA polymerase I	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1901269	lipopolysaccharide metabolic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0035622	intrahepatic bile duct development	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0016444	somatic cell DNA recombination	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1901271	lipopolysaccharide biosynthetic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:005173	response to defences of other organism	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0003382	epithelial cell morphogenesis	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0070977	bone maturation	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0031571	mitotic G1 DNA damage checkpoint signaling	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0043247	telomere maintenance in response to DNA damage	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0010669	epithelial structure maintenance	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:009301	snRNA transcription	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0042149	cellular response to glucose starvation	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1902914	regulation of protein polyubiquitination	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1902915	negative regulation of protein polyubiquitination	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0032196	transposition	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0032197	retrotransposition	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0042149	retroh/ACA sno(s)RNA processing	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:007168	ribonucleoprotein granule cycle signaling pathway	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0017001	antibiotic catabolic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0030277	maintenance of gastrointestinal epithelium	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0004949	box H/ACA sno(s)RNA 3'-end processing	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1903513	endoplasmic reticulum to cytosol transport	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1900481	mRNA pseudouridine synthase	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0044819	mitotic G1/S transition checkpoint signaling	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0051155	positive regulation of striated muscle cell differentiation	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0007140	antibiotic resistance	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0050772	male meiotic nuclear division	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0009046	positive regulation of xenogenesis	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0099172	mitochondrial tRNA processing	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:00132065	presynapse organization	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0044819	maintenance of protein location in cell cortex	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0051155	mitotic G1/S transition checkpoint signaling	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0007140	positive regulation of striated muscle cell differentiation	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0050772	antibiotic resistance	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0009046	male meiotic nuclear division	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0099172	positive regulation of xenogenesis	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:00132065	mitochondrial tRNA processing	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:190152	presynapse organization	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:009722	maintenance of protein location in cell cortex	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0009046	mitotic G1/S transition checkpoint signaling	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0002033	positive regulation of muscle cell differentiation	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0052200	antigenic variation	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0016074	positive regulation of muscle tissue development	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0043931	ossification involved in bone maturation	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0035279	miRNA-mediated gene silencing by mRNA destabilization	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0061009	common bile duct development	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0099054	presynapse assembly	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048641	regulation of skeletal muscle tissue development	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048643	positive regulation of skeletal muscle tissue development	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1901861	regulation of muscle tissue development	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1901863	positive regulation of muscle tissue development	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048634	retrograde protein transport, ER to cytosol	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0051274	beta-glucan biosynthetic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0051278	snoRNA pseudouridine synthase	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0031120	sno(s)RNA 3'-end processing	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0033126	lipopolysaccharide biosynthetic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0033397	nuclear matrix anchoring at nuclear membrane	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0071966	fungi-type cell wall polysaccharide metabolic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0042783	evasion of host immune response	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0030970	retrograde protein transport, ER to cytosol	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048644	beta-glucan biosynthetic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048645	fungal-type cell wall polysaccharide biosynthetic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048147	lipopolysaccharide biosynthetic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0090292	nuclear matrix anchoring at nuclear membrane	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0071947	protein debiquitination involved in ubiquitin-dependent protein catabolic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0045943	(I->3)-beta-D-glucan metabolic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048144	(I->3)-beta-D-glucan biosynthetic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048145	lactam metabolic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0044650	adhesion of symbiont to host cell	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1902414	protein localization to cell junction	1	1	0.04	0.0433

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Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Bivalvia	Group 1 + Group 2	GO:0009272	fungi-type cell wall biogenesis	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0009243	O antigen biosynthetic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0043578	nuclear matrix organization	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0009445	lipid A biosynthetic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0044406	adhesion of symbiont to host	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0032978	protein insertion into membrane from inner side	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0032979	protein insertion into mitochondrial inner membrane from matrix	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0046493	lipid A metabolic process	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0072695	regulation of DNA recombination at telomere	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1900044	regulation of protein K63-linked ubiquitination	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0035871	protein K11-linked deubiquitination	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:1900045	negative regulation of protein K63-linked ubiquitination	1	1	0.04	0.0433
Bivalvia	Group 1 + Group 2	GO:0048239	negative regulation of DNA recombination at telomere	1	1	0.04	0.0433
Drosophila	Group 1 + Group 2	GO:0045132	meiotic chromosome segregation	64	11	2.63	0.0015
Drosophila	Group 1 + Group 2	GO:0006819	sister chromatid segregation	140	11	5.75	0.0293
Drosophila	Group 1 + Group 2	GO:0070192	chromosome organization involved in meiotic cell cycle	54	9	2.22	0.0085
Drosophila	Group 1 + Group 2	GO:0007131	reciprocal meiotic recombination	37	7	1.52	0.0007
Drosophila	Group 1 + Group 2	GO:0007143	female meiotic nuclear division	54	6	2.22	0.0227
Drosophila	Group 1 + Group 2	GO:0035967	cellular response to topologically incorrect protein	44	5	1.81	0.0333
Drosophila	Group 1 + Group 2	GO:0039966	response to topologically incorrect protein	47	5	1.93	0.0427
Drosophila	Group 1 + Group 2	GO:0007141	male meiosis	13	4	0.53	0.0015
Drosophila	Group 1 + Group 2	GO:0040543	positive regulation of pRNA transcription	3	3	0.12	0.0001
Drosophila	Group 1 + Group 2	GO:0010526	retrotransposon silencing	8	3	0.33	0.0033
Drosophila	Group 1 + Group 2	GO:0007130	synaptonemal complex assembly	10	3	0.41	0.0067
Drosophila	Group 1 + Group 2	GO:0030719	P granule organization	11	3	0.45	0.0089
Drosophila	Group 1 + Group 2	GO:0071218	cellular response to misfolded protein	12	3	0.49	0.0115
Drosophila	Group 1 + Group 2	GO:00051788	response to misfolded protein	12	3	0.49	0.0115
Drosophila	Group 1 + Group 2	GO:0007135	meiosis II	15	3	0.62	0.0217
Drosophila	Group 1 + Group 2	GO:0034508	centromere complex assembly	19	3	0.78	0.0409
Drosophila	Group 1 + Group 2	GO:0048136	male germ-line cyst formation	2	2	0.08	0.0017
Drosophila	Group 1 + Group 2	GO:0061964	negative regulation of entry into reproductive diapause	5	2	0.21	0.0155
Drosophila	Group 1 + Group 2	GO:0051382	kinetochore assembly	5	2	0.21	0.0155
Drosophila	Group 1 + Group 2	GO:0055116	entry into reproductive diapause	6	2	0.25	0.0226
Drosophila	Group 1 + Group 2	GO:0071712	ER-associated misfolded protein catabolic process	6	2	0.25	0.0226
Drosophila	Group 1 + Group 2	GO:0061963	regulation of entry into reproductive diapause	6	2	0.25	0.0226
Drosophila	Group 1 + Group 2	GO:0043984	histone H4-K16 acetylation	6	2	0.25	0.0226
Drosophila	Group 1 + Group 2	GO:0055115	entry into diapause	7	2	0.29	0.0308
Drosophila	Group 1 + Group 2	GO:1990834	response to odorant	7	2	0.29	0.0308
Drosophila	Group 1 + Group 2	GO:0051177	meiotic sister chromatid cohesion	8	2	0.33	0.0400
Drosophila	Group 1 + Group 2	GO:0042795	snRNA transcription by RNA polymerase II	8	2	0.33	0.0400
Drosophila	Group 1 + Group 2	GO:0022611	dormancy process	8	2	0.33	0.0400
Drosophila	Group 1 + Group 2	GO:0009301	snrRNA transcription	8	2	0.33	0.0400
Drosophila	Group 1 + Group 2	GO:0045144	meiotic sister chromatid segregation	8	2	0.33	0.0400
Drosophila	Group 1 + Group 2	GO:0001015	snRNA transcription by RNA polymerase II	1	1	0.04	0.0411
Drosophila	Group 1 + Group 2	GO:0010778	meiotic DNA repair synthesis involved in reciprocal meiotic recombination	1	1	0.04	0.0411
Drosophila	Group 1 + Group 2	GO:0072765	centromere localization	1	1	0.04	0.0411
Drosophila	Group 1 + Group 2	GO:0099302	sn(s)RNA transcription	1	1	0.04	0.0411
Drosophila	Group 1 + Group 2	GO:0051308	male meiosis chromosome separation	1	1	0.04	0.0411
Drosophila	Group 1 + Group 2	GO:0098663	centromere clustering	1	1	0.04	0.0411
Drosophila	Group 1 + Group 2	GO:0051415	microtubule nucleation by interphase microtubule organizing center	1	1	0.04	0.0411
Mammalia	Group 1 + Group 2	GO:0006955	immune response	1,297	145	48.02	0.0006
Mammalia	Group 1 + Group 2	GO:0098542	defense response to other organism	853	112	31.58	0.0207
Mammalia	Group 1 + Group 2	GO:0045087	innate immune response	647	82	23.95	0.0000
Mammalia	Group 1 + Group 2	GO:001817	regulation of cytokine production	630	51	23.33	0.0466
Mammalia	Group 1 + Group 2	GO:0042742	defense response to bacterium	233	45	8.63	0.0000
Mammalia	Group 1 + Group 2	GO:000954	inflammatory response	642	45	23.77	0.0174
Mammalia	Group 1 + Group 2	GO:191221	cytokine-mediated signaling pathway	382	44	14.14	0.0000
Mammalia	Group 1 + Group 2	GO:002250	adaptive immune response	342	41	12.66	0.0000
Mammalia	Group 1 + Group 2	GO:001819	positive regulation of cytokine production	402	41	14.88	0.0272
Mammalia	Group 1 + Group 2	GO:002697	regulation of immune effector process	308	37	11.40	0.0443
Mammalia	Group 1 + Group 2	GO:0042110	T cell activation	432	35	15.99	0.0256
Mammalia	Group 1 + Group 2	GO:0051607	defense response to virus	257	34	9.52	0.0000
Mammalia	Group 1 + Group 2	GO:0048232	male gamete generation	491	32	18.18	0.0226
Mammalia	Group 1 + Group 2	GO:007283	spontaneous	478	31	17.70	0.0280
Mammalia	Group 1 + Group 2	GO:0070661	leukocyte proliferation	273	29	10.11	0.0120
Mammalia	Group 1 + Group 2	GO:002449	lymphocyte mediated immunity	221	29	8.18	0.0483
Mammalia	Group 1 + Group 2	GO:0070663	regulation of leukocyte proliferation	212	25	7.85	0.0187

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Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Mammalia	Group 1 + Group 2	GO:0050727	regulation of inflammatory response	300	24	11.11	0.0024
Mammalia	Group 1 + Group 2	GO:0031349	positive regulation of defense response	240	24	8.89	0.0124
Mammalia	Group 1 + Group 2	GO:0020768	immune response-regulating cell surface receptor signaling pathway	177	22	6.55	0.0034
Mammalia	Group 1 + Group 2	GO:0050829	defense response to Gram-negative bacterium	66	17	2.44	0.0000
Mammalia	Group 1 + Group 2	GO:0071222	cellular response to lipopolysaccharide	164	17	6.07	0.0001
Mammalia	Group 1 + Group 2	GO:0010466	negative regulation of peptidase activity	163	16	6.04	0.0004
Mammalia	Group 1 + Group 2	GO:0002429	immune response-activating cell surface receptor signaling pathway	164	16	6.07	0.0024
Mammalia	Group 1 + Group 2	GO:1930555	regulation of tumor necrosis factor superfamily cytokine production	137	16	5.07	0.0124
Mammalia	Group 1 + Group 2	GO:0071706	tumor necrosis factor superfamily cytokine production	137	16	5.07	0.0124
Mammalia	Group 1 + Group 2	GO:0070665	positive regulation of leukocyte proliferation	132	16	4.89	0.0277
Mammalia	Group 1 + Group 2	GO:0045089	positive regulation of innate immune response	113	16	4.18	0.0322
Mammalia	Group 1 + Group 2	GO:0071356	cellular response to tumor necrosis factor	175	15	6.48	0.0022
Mammalia	Group 1 + Group 2	GO:0002695	negative regulation of leukocyte activation	148	15	5.48	0.0115
Mammalia	Group 1 + Group 2	GO:0002456	T cell mediated immunity	82	15	3.04	0.0161
Mammalia	Group 1 + Group 2	GO:0002705	positive regulation of leukocyte mediated immunity	113	15	4.18	0.0184
Mammalia	Group 1 + Group 2	GO:0032680	regulation of tumor necrosis factor production	133	15	4.92	0.0326
Mammalia	Group 1 + Group 2	GO:0032640	tumor necrosis factor production	133	15	4.92	0.0326
Mammalia	Group 1 + Group 2	GO:0050866	negative regulation of cell activation	165	15	6.11	0.0405
Mammalia	Group 1 + Group 2	GO:0031341	regulation of cell killing	71	14	2.63	0.0063
Mammalia	Group 1 + Group 2	GO:0001818	negative regulation of cytokine production	225	14	8.33	0.0407
Mammalia	Group 1 + Group 2	GO:0050830	defense response to Gram-positive bacterium	87	13	3.22	0.0000
Mammalia	Group 1 + Group 2	GO:0002286	T cell activation involved in immune response	94	13	3.48	0.0293
Mammalia	Group 1 + Group 2	GO:0050909	sensory perception of taste	52	13	1.93	0.0496
Mammalia	Group 1 + Group 2	GO:007259	cell surface receptor signaling pathway via JAK-STAT	134	12	4.96	0.0041
Mammalia	Group 1 + Group 2	GO:0019731	antigen receptor-mediated signaling pathway	130	12	4.81	0.0182
Mammalia	Group 1 + Group 2	GO:0042101	natural killer cell response	40	11	1.48	0.0000
Mammalia	Group 1 + Group 2	GO:0042102	defense response to T cell proliferation	66	11	2.44	0.0000
Mammalia	Group 1 + Group 2	GO:0050832	positive regulation of T cell proliferation	85	11	3.15	0.0003
Mammalia	Group 1 + Group 2	GO:0071346	defense response to fungus	48	11	1.78	0.0008
Mammalia	Group 1 + Group 2	GO:008584	cellular response to type II interferon	98	11	3.63	0.0010
Mammalia	Group 1 + Group 2	GO:0002820	male gonad development	120	11	4.44	0.0049
Mammalia	Group 1 + Group 2	GO:0043123	negative regulation of adaptive immune response	42	11	1.56	0.0077
Mammalia	Group 1 + Group 2	GO:0030101	positive regulation of canonical NF-κappaB signal transduction	162	11	6.00	0.0386
Mammalia	Group 1 + Group 2	GO:0042100	B cell proliferation	76	11	2.81	0.0449
Mammalia	Group 1 + Group 2	GO:0001580	detection of chemical stimulus involved in sensory perception of bitter taste	29	10	1.07	0.0000
Mammalia	Group 1 + Group 2	GO:0030593	neutrophil chemotaxis	80	10	2.96	0.0007
Mammalia	Group 1 + Group 2	GO:0009556	complement activation	47	10	1.74	0.0008
Mammalia	Group 1 + Group 2	GO:0032760	positive regulation of tumor necrosis factor production	84	10	3.11	0.0011
Mammalia	Group 1 + Group 2	GO:0071347	cellular response to interleukin-1	84	10	3.11	0.0011
Mammalia	Group 1 + Group 2	GO:0050729	positive regulation of inflammatory response	122	10	4.52	0.0152
Mammalia	Group 1 + Group 2	GO:0002823	negative regulation of adaptive immune response [...]	39	10	1.44	0.0306
Mammalia	Group 1 + Group 2	GO:0050688	regulation of defense response to virus	62	10	3.55	0.0091
Mammalia	Group 1 + Group 2	GO:0002718	regulation of cytokine production involved in immune response	59	9	4.59	0.0435
Mammalia	Group 1 + Group 2	GO:0002367	cytokine production involved in immune response	98	10	3.63	0.0435
Mammalia	Group 1 + Group 2	GO:0007339	binding of sperm to zona pellucida	35	9	1.30	0.0000
Mammalia	Group 1 + Group 2	GO:0061844	antimicrobial humoral immune response mediated by antimicrobial peptide	64	9	2.37	0.0006
Mammalia	Group 1 + Group 2	GO:0031640	T cell migration	64	9	2.37	0.0006
Mammalia	Group 1 + Group 2	GO:0050866	regulation of response to cytokine stimulus	63	9	2.33	0.0360
Mammalia	Group 1 + Group 2	GO:0001910	fusion movement involved in cell motility	96	9	3.55	0.0088
Mammalia	Group 1 + Group 2	GO:0002722	regulation of leukocyte mediated cytotoxicity	124	9	4.59	0.0408
Mammalia	Group 1 + Group 2	GO:0030317	flagellar production involved in immune response	114	9	4.74	0.0483
Mammalia	Group 1 + Group 2	GO:0097722	flagellated sperm motility	25	8	0.0256	0.0326
Mammalia	Group 1 + Group 2	GO:0038061	sperm motility	119	9	4.41	0.0027
Mammalia	Group 1 + Group 2	GO:0072678	non-canonical NF-κappaB signal transduction	39	8	2.55	0.0326
Mammalia	Group 1 + Group 2	GO:0060759	T cell migration	45	8	1.67	0.0002
Mammalia	Group 1 + Group 2	GO:0060294	regulation of response to cytokine stimulus	54	8	2.00	0.0008
Mammalia	Group 1 + Group 2	GO:007342	fusion of sperm to egg plasma membrane involved in single fertilization	60	8	2.22	0.0016
Mammalia	Group 1 + Group 2	GO:0045071	negative regulation of viral genome replication	65	8	2.41	0.0027
Mammalia	Group 1 + Group 2	GO:0002218	activation of innate immune response	39	8	1.44	0.0001
Mammalia	Group 1 + Group 2	GO:0032757	positive regulation of interleukin-8 production	45	8	1.67	0.0002
Mammalia	Group 1 + Group 2	GO:0030888	regulation of B cell proliferation	54	8	2.00	0.0008
Mammalia	Group 1 + Group 2	GO:0032722	positive regulation of chemokine production	60	8	2.22	0.0016
Mammalia	Group 1 + Group 2	GO:0010921	regulation of phosphatase activity	65	8	2.41	0.0027
Mammalia	Group 1 + Group 2	GO:0070998	chemokine-mediated signaling pathway	69	8	2.55	0.039
Mammalia	Group 1 + Group 2	GO:0029220	regulation of humoral immune response	36	8	1.33	0.0051
Mammalia	Group 1 + Group 2	GO:0043303	activation of innate immune response	43	8	1.59	0.0055
Mammalia	Group 1 + Group 2	GO:0002251	mast cell degranulation	22	8	0.81	0.0065
Mammalia	Group 1 + Group 2	GO:002886	organ or tissue specific immune response	45	8	1.67	0.0078
Mammalia	Group 1 + Group 2	GO:0043030	regulation of myeloid leukocyte mediated immunity	46	8	1.70	0.0092

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Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Mammalia	Group 1 + Group 2	GO:0002526	acute inflammatory response	83	8	3.07	0.0116
Mammalia	Group 1 + Group 2	GO:0032649	regulation of type II interferon production	88	8	3.26	0.0162
Mammalia	Group 1 + Group 2	GO:0032609	type II interferon production	88	8	3.26	0.0162
Mammalia	Group 1 + Group 2	GO:0050691	regulation of defense response to virus by host	37	8	1.37	0.0498
Mammalia	Group 1 + Group 2	GO:0042267	natural killer cell mediated cytotoxicity	44	7	1.63	0.0111
Mammalia	Group 1 + Group 2	GO:0048247	lymphocyte chemotaxis	49	7	1.81	0.0020
Mammalia	Group 1 + Group 2	GO:0042119	neutrophil activation	32	7	1.18	0.0079
Mammalia	Group 1 + Group 2	GO:0032945	negative regulation of mononuclear cell proliferation	63	7	2.33	0.0085
Mammalia	Group 1 + Group 2	GO:0002726	positive regulation of cytokine production involved in immune response	66	7	2.44	0.0108
Mammalia	Group 1 + Group 2	GO:0038456	response to interferon-beta	26	7	0.96	0.0125
Mammalia	Group 1 + Group 2	GO:0050853	B cell receptor signaling pathway	45	7	1.67	0.0168
Mammalia	Group 1 + Group 2	GO:0098586	cellular response to virus	78	7	2.89	0.0252
Mammalia	Group 1 + Group 2	GO:0046425	regulation of receptor signaling pathway via JAK-STAT	78	7	2.89	0.0252
Mammalia	Group 1 + Group 2	GO:0002385	mucosal immune response	19	7	0.70	0.0325
Mammalia	Group 1 + Group 2	GO:0032755	positive regulation of interleukin-6 production	83	7	3.07	0.0340
Mammalia	Group 1 + Group 2	GO:1901222	regulation of non-canonical NF-kappaB signal transduction	89	7	3.30	0.0468
Mammalia	Group 1 + Group 2	GO:0048245	eosinophil chemotaxis	17	6	0.63	0.0000
Mammalia	Group 1 + Group 2	GO:0048240	sperm capacitation	25	6	0.93	0.0002
Mammalia	Group 1 + Group 2	GO:0070269	pyroptotic inflammatory response	26	6	0.96	0.0003
Mammalia	Group 1 + Group 2	GO:0002230	positive regulation of defense response to virus by host	27	6	1.00	0.0004
Mammalia	Group 1 + Group 2	GO:0031295	T cell costimulation	27	6	1.00	0.0004
Mammalia	Group 1 + Group 2	GO:0003825	regulation of T-helper 1 type immune response	29	6	1.07	0.0006
Mammalia	Group 1 + Group 2	GO:002691	regulation of cellular extravasation	38	6	1.41	0.0025
Mammalia	Group 1 + Group 2	GO:0140374	antiviral innate immune response	41	6	1.52	0.0037
Mammalia	Group 1 + Group 2	GO:0002639	positive regulation of immunoglobulin production	44	6	1.63	0.0054
Mammalia	Group 1 + Group 2	GO:0032731	positive regulation of interleukin-1 beta production	46	6	1.70	0.0067
Mammalia	Group 1 + Group 2	GO:00190894	positive regulation of receptor signaling pathway via STAT	46	6	1.70	0.0067
Mammalia	Group 1 + Group 2	GO:0031113	heterotypic cell-cell adhesion	47	6	1.74	0.0074
Mammalia	Group 1 + Group 2	GO:002548	monocyte chemoattractant	50	6	1.85	0.0100
Mammalia	Group 1 + Group 2	GO:001961	positive regulation of cytokine-mediated signaling pathway	50	6	1.85	0.0100
Mammalia	Group 1 + Group 2	GO:0030449	regulation of complement activation	18	6	0.67	0.0108
Mammalia	Group 1 + Group 2	GO:0042531	positive regulation of tyrosine phosphorylation of STAT protein	51	6	1.89	0.0110
Mammalia	Group 1 + Group 2	GO:0032731	type I interferon-mediated signaling pathway	52	6	1.93	0.0121
Mammalia	Group 1 + Group 2	GO:001357	cellular response to type I interferon	52	6	1.93	0.0121
Mammalia	Group 1 + Group 2	GO:0001914	regulation of T cell mediated cytotoxicity	25	6	0.93	0.0126
Mammalia	Group 1 + Group 2	GO:0051293	establishment of spindle localization	54	6	2.00	0.0144
Mammalia	Group 1 + Group 2	GO:0051873	killing by host symbiont cells	23	6	0.85	0.0163
Mammalia	Group 1 + Group 2	GO:0060760	positive regulation of response to cytokine stimulus	57	6	2.11	0.0184
Mammalia	Group 1 + Group 2	GO:0150077	regulation of neuroinflammatory response	28	6	1.04	0.0195
Mammalia	Group 1 + Group 2	GO:0033430	response to type I interferon	58	6	2.15	0.0199
Mammalia	Group 1 + Group 2	GO:0051653	spindle localization	59	6	2.18	0.0215
Mammalia	Group 1 + Group 2	GO:001895	retina homeostasis	60	6	2.22	0.0232
Mammalia	Group 1 + Group 2	GO:0071260	cellular response to mechanical stimulus	60	6	2.22	0.0232
Mammalia	Group 1 + Group 2	GO:0050672	negative regulation of lymphocyte proliferation	62	6	2.30	0.0268
Mammalia	Group 1 + Group 2	GO:0032729	positive regulation of type II interferon production	62	6	2.30	0.0268
Mammalia	Group 1 + Group 2	GO:0042509	regulation of tyrosine phosphorylation of STAT protein	62	6	2.30	0.0268
Mammalia	Group 1 + Group 2	GO:0007260	tyrosine phosphorylation of STAT protein	66	6	2.44	0.0350
Mammalia	Group 1 + Group 2	GO:002227	innate immune response in mucosa	11	5	0.41	0.0000
Mammalia	Group 1 + Group 2	GO:0002830	positive regulation of type 2 immune response	14	5	0.52	0.0001
Mammalia	Group 1 + Group 2	GO:0038455	response to interferon-alpha	14	5	0.52	0.0001
Mammalia	Group 1 + Group 2	GO:0033005	positive regulation of mast cell activation	17	5	0.63	0.0003
Mammalia	Group 1 + Group 2	GO:001774	microglial cell activation	35	5	1.30	0.0089
Mammalia	Group 1 + Group 2	GO:0002701	negative regulation of production of molecular mediator of immune response	35	5	1.30	0.0089
Mammalia	Group 1 + Group 2	GO:0140632	canonical inflammasome complex assembly	35	5	1.30	0.0089
Mammalia	Group 1 + Group 2	GO:0006953	acute-phase response	38	5	1.41	0.0125
Mammalia	Group 1 + Group 2	GO:0032653	regulation of interleukin-10 production	45	5	1.67	0.0247
Mammalia	Group 1 + Group 2	GO:0032613	interleukin-10 production	45	5	1.70	0.0269
Mammalia	Group 1 + Group 2	GO:043666	regulation of phosphoprotein phosphatase activity	47	5	1.74	0.0292
Mammalia	Group 1 + Group 2	GO:0033006	negative regulation of tumor necrosis factor production	47	5	1.74	0.0292
Mammalia	Group 1 + Group 2	GO:1903556	negative regulation of tumor necrosis factor superfamily cytokine production	49	5	1.81	0.0342
Mammalia	Group 1 + Group 2	GO:0070228	regulation of lymphocyte apoptotic process	50	5	1.85	0.0369
Mammalia	Group 1 + Group 2	GO:0072683	T cell extravasation	13	4	0.48	0.0010
Mammalia	Group 1 + Group 2	GO:0002710	negative regulation of T cell mediated immunity	15	4	0.56	0.0018

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Dataset	Group of genes	GO ID	Term	Annotated	Significant	Expected	Classic Fisher	
Mammalia	Group 1 + Group 2	GO:0072540	T-helper 17 cell lineage commitment	17	4	0.63	0.0030	
Mammalia	Group 1 + Group 2	GO:0048624	positive regulation of T-helper cell differentiation	17	4	0.63	0.0030	
Mammalia	Group 1 + Group 2	GO:1903659	regulation of complement-dependent cytotoxicity	5	4	0.19	0.0040	
Mammalia	Group 1 + Group 2	GO:0008228	opsonization	19	4	0.70	0.0046	
Mammalia	Group 1 + Group 2	GO:0043302	positive regulation of leukocyte degranulation	20	4	0.74	0.0056	
Mammalia	Group 1 + Group 2	GO:001916	positive regulation of T cell mediated cytotoxicity	20	4	0.74	0.0056	
Mammalia	Group 1 + Group 2	GO:001562	response to protease	21	4	0.78	0.0067	
Mammalia	Group 1 + Group 2	GO:0021717	positive regulation of natural killer cell mediated immunity	21	4	0.78	0.0067	
Mammalia	Group 1 + Group 2	GO:1903901	negative regulation of viral life cycle	21	4	0.78	0.0067	
Mammalia	Group 1 + Group 2	GO:0038821	modulation of process of another organism	21	4	0.78	0.0067	
Mammalia	Group 1 + Group 2	GO:0043304	regulation of mast cell degranulation	23	4	0.85	0.0094	
Mammalia	Group 1 + Group 2	GO:0032740	positive regulation of interleukin-17 production	23	4	0.85	0.0094	
Mammalia	Group 1 + Group 2	GO:0043032	positive regulation of macrophage activation	23	4	0.85	0.0094	
Mammalia	Group 1 + Group 2	GO:0070498	interleukin-1-mediated signaling pathway	24	4	0.89	0.0110	
Mammalia	Group 1 + Group 2	GO:0010922	positive regulation of phosphatase activity	26	4	0.96	0.0145	
Mammalia	Group 1 + Group 2	GO:0019884	antigen processing and presentation of exogenous antigen	28	4	1.04	0.0188	
Mammalia	Group 1 + Group 2	GO:002446	neutrophil mediated immunity	29	4	1.07	0.0212	
Mammalia	Group 1 + Group 2	GO:0032743	positive regulation of interleukin-2 production	30	4	1.11	0.0238	
Mammalia	Group 1 + Group 2	GO:2000352	negative regulation of endothelial cell apoptotic process	30	4	1.11	0.0238	
Mammalia	Group 1 + Group 2	GO:1902225	response to NLRP3 inflammasome complex assembly	30	4	1.11	0.0265	
Mammalia	Group 1 + Group 2	GO:0043330	response to exogenous dsRNA	31	4	1.15	0.0265	
Mammalia	Group 1 + Group 2	GO:0032733	positive regulation of interleukin-10 production	32	4	1.18	0.0294	
Mammalia	Group 1 + Group 2	GO:0046006	positive regulation of protein dephosphorylation	32	4	1.18	0.0294	
Mammalia	Group 1 + Group 2	GO:0032814	regulation of activated T cell proliferation	32	4	1.18	0.0294	
Mammalia	Group 1 + Group 2	GO:0044546	regulation of natural killer cell activation	33	4	1.22	0.0325	
Mammalia	Group 1 + Group 2	GO:0050798	NLRP3 inflammasome complex assembly	33	4	1.22	0.0325	
Mammalia	Group 1 + Group 2	GO:0046636	negative regulation of alpha-beta T cell activation	34	4	1.26	0.0358	
Mammalia	Group 1 + Group 2	GO:0043331	response to dsRNA	35	4	1.30	0.0393	
Mammalia	Group 1 + Group 2	GO:0035307	positive regulation of protein dephosphorylation	36	4	1.33	0.0430	
Mammalia	Group 1 + Group 2	GO:0030890	positive regulation of B cell proliferation	36	4	1.33	0.0430	
Mammalia	Group 1 + Group 2	GO:0038095	Fc-receptor signaling pathway	37	4	1.37	0.0468	
Mammalia	Group 1 + Group 2	GO:0045959	negative regulation of complement activation, classical pathway	5	3	0.19	0.0005	
Mammalia	Group 1 + Group 2	GO:0038156	interleukin-3-mediated signaling pathway	5	3	0.19	0.0005	
Mammalia	Group 1 + Group 2	GO:0051838	cytokine by host of symbiont transduction	7	3	0.26	0.0016	
Mammalia	Group 1 + Group 2	GO:0006977	DNA damage response, signal transduction by p53 class mediator resulting in cell cycle arrest	8	3	0.30	0.0025	
Mammalia	Group 1 + Group 2	GO:0097527	GO:0006957	necrotic signaling pathway	8	3	0.30	0.0025
Mammalia	Group 1 + Group 2	GO:0008924	activation-induced cell death of T cells	8	3	0.37	0.0050	
Mammalia	Group 1 + Group 2	GO:1902563	regulation of neutrophil activation	10	3	0.37	0.0050	
Mammalia	Group 1 + Group 2	GO:0010526	retrotransposon silencing	10	3	0.37	0.0050	
Mammalia	Group 1 + Group 2	GO:0032754	negative regulation of neuroinflammatory response	10	3	0.37	0.0050	
Mammalia	Group 1 + Group 2	GO:2000551	positive regulation of interleukin-5 production	10	3	0.37	0.0050	
Mammalia	Group 1 + Group 2	GO:1901731	regulation of T-helper 2 cell cytokine production	10	3	0.37	0.0050	
Mammalia	Group 1 + Group 2	GO:0032736	natural killer cell activation involved in immune response	10	3	0.41	0.0067	
Mammalia	Group 1 + Group 2	GO:0007343	positive regulation of interleukin-13 production	11	3	0.41	0.0067	
Mammalia	Group 1 + Group 2	GO:0035723	complement activation, alternative pathway	14	3	0.52	0.0135	
Mammalia	Group 1 + Group 2	GO:010527	interleukin-15-mediated signaling pathway	11	3	0.52	0.0135	
Mammalia	Group 1 + Group 2	GO:0150079	negative regulation of neuroinflammatory response	14	3	0.52	0.0135	
Mammalia	Group 1 + Group 2	GO:0060046	regulation of acrosome reaction	14	3	0.52	0.0135	
Mammalia	Group 1 + Group 2	GO:002323	T-helper 2 cell differentiation	15	3	0.56	0.0165	
Mammalia	Group 1 + Group 2	GO:0030889	natural killer cell activation involved in immune response	15	3	0.56	0.0165	
Mammalia	Group 1 + Group 2	GO:0006957	negative regulation of B cell proliferation	16	3	0.59	0.0197	
Mammalia	Group 1 + Group 2	GO:0043306	complement activation, alternative pathway	16	3	0.59	0.0197	
Mammalia	Group 1 + Group 2	GO:1903027	positive regulation of opsonization	16	3	0.59	0.0197	
Mammalia	Group 1 + Group 2	GO:0020335	CD40 signaling pathway	17	3	0.63	0.0233	
Mammalia	Group 1 + Group 2	GO:0033008	positive regulation of mast cell activation involved in immune response	17	3	0.63	0.0233	
Mammalia	Group 1 + Group 2	GO:0045064	regulation of acrosome reaction	17	3	0.67	0.0272	
Mammalia	Group 1 + Group 2	GO:00023730	dendritic cell cytokine production	18	3	0.67	0.0272	
Mammalia	Group 1 + Group 2	GO:1901538	changes to DNA methylation involved in embryo development	18	3	0.70	0.0315	
Mammalia	Group 1 + Group 2	GO:0002888	positive regulation of myeloid leukocyte mediated immunity	19	3	0.70	0.0315	
Mammalia	Group 1 + Group 2	GO:002544	chronic inflammatory response	19	3	0.70	0.0315	
Mammalia	Group 1 + Group 2	GO:002693	positive regulation of cellular extravasation	17	3	0.63	0.0233	
Mammalia	Group 1 + Group 2	GO:002827	positive regulation of T-helper 1 type immune response	18	3	0.67	0.0272	
Mammalia	Group 1 + Group 2	GO:0045346	defense response to protozoan	19	3	0.70	0.0315	
Mammalia	Group 1 + Group 2	GO:0042832	MHC class II biosynthetic process	19	3	0.70	0.0315	

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Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Mammalia	Group 1 + Group 2	GO:0040019	positive regulation of embryonic development	19	3	0.70	0.0315
Mammalia	Group 1 + Group 2	GO:0060340	positive regulation of type I interferon-mediated signaling pathway	19	3	0.70	0.0315
Mammalia	Group 1 + Group 2	GO:0060333	type II interferon-mediated signaling pathway	19	3	0.70	0.0315
Mammalia	Group 1 + Group 2	GO:0032753	positive regulation of interleukin-4 production	20	3	0.74	0.0360
Mammalia	Group 1 + Group 2	GO:0045063	T-helper 1 cell differentiation	21	3	0.78	0.0409
Mammalia	Group 1 + Group 2	GO:0042104	positive regulation of activated T cell proliferation	22	3	0.81	0.0461
Mammalia	Group 1 + Group 2	GO:0002726	positive regulation of T cell cytokine production	22	3	0.81	0.0461
Mammalia	Group 1 + Group 2	GO:1903660	negative regulation of complement-dependent cytotoxicity	2	2	0.07	0.0014
Mammalia	Group 1 + Group 2	GO:0097528	execution phase of necrosis	2	2	0.07	0.0014
Mammalia	Group 1 + Group 2	GO:0030887	positive regulation of myeloid dendritic cell activation	2	2	0.07	0.0014
Mammalia	Group 1 + Group 2	GO:0009609	response to symbiotic bacterium	4	2	0.15	0.0078
Mammalia	Group 1 + Group 2	GO:0048006	antigen processing and presentation, endogenous lipid antigen via MHC class Ib	4	2	0.15	0.0078
Mammalia	Group 1 + Group 2	GO:0050859	negative regulation of B cell receptor signaling pathway	4	2	0.15	0.0078
Mammalia	Group 1 + Group 2	GO:0002765	immune response-inhibiting signal transduction	5	2	0.19	0.0127
Mammalia	Group 1 + Group 2	GO:0036015	response to interleukin-3	5	2	0.19	0.0127
Mammalia	Group 1 + Group 2	GO:0036016	cellular response to interleukin-3	5	2	0.19	0.0127
Mammalia	Group 1 + Group 2	GO:2000416	regulation of eosinophil migration	5	2	0.19	0.0127
Mammalia	Group 1 + Group 2	GO:0033634	positive regulation of cell-cell adhesion mediated by integrin	5	2	0.19	0.0127
Mammalia	Group 1 + Group 2	GO:0045630	positive regulation of T-helper 2 cell differentiation	5	2	0.19	0.0127
Mammalia	Group 1 + Group 2	GO:0043313	regulation of neutrophil degranulation	6	2	0.22	0.0186
Mammalia	Group 1 + Group 2	GO:0033141	positive regulation of peptidyl-serine phosphorylation of STAT protein	6	2	0.22	0.0186
Mammalia	Group 1 + Group 2	GO:0002826	negative regulation of T-helper 1 type immune response	6	2	0.22	0.0186
Mammalia	Group 1 + Group 2	GO:006300	protein-DNA covalent cross-linking repair	6	2	0.22	0.0186
Mammalia	Group 1 + Group 2	GO:0043341	MHC class I biosynthetic process	6	2	0.22	0.0186
Mammalia	Group 1 + Group 2	GO:0045343	regulation of MHC class I biosynthetic process	6	2	0.22	0.0186
Mammalia	Group 1 + Group 2	GO:0048007	antigen processing and presentation, exogenous lipid antigen via MHC class Ib	6	2	0.22	0.0186
Mammalia	Group 1 + Group 2	GO:0060332	positive regulation of response to type II interferon	6	2	0.22	0.0186
Mammalia	Group 1 + Group 2	GO:2000659	positive regulation of type II interferon-mediated signaling pathway	6	2	0.22	0.0186
Mammalia	Group 1 + Group 2	GO:0060545	regulation of interleukin-1-mediated signaling pathway	6	2	0.22	0.0186
Mammalia	Group 1 + Group 2	GO:0043307	positive regulation of necrotic cell processes	6	2	0.22	0.0186
Mammalia	Group 1 + Group 2	GO:002733	eosinophil activation	7	2	0.26	0.0254
Mammalia	Group 1 + Group 2	GO:0002735	regulation of myeloid dendritic cell cytokine production	7	2	0.26	0.0254
Mammalia	Group 1 + Group 2	GO:0003391	positive regulation of neutrophil extravasation	7	2	0.26	0.0254
Mammalia	Group 1 + Group 2	GO:0010940	positive regulation of interleukin-1-mediated signaling pathway	7	2	0.26	0.0254
Mammalia	Group 1 + Group 2	GO:00060467	negative regulation of necrotic cell death	7	2	0.26	0.0254
Mammalia	Group 1 + Group 2	GO:002733	negative regulation of fertilization	7	2	0.26	0.0254
Mammalia	Group 1 + Group 2	GO:0002735	prevention of polysomy	7	2	0.26	0.0254
Mammalia	Group 1 + Group 2	GO:002735	cellular response to type III interferon	7	2	0.26	0.0254
Mammalia	Group 1 + Group 2	GO:0002372	type III interferon-mediated signaling pathway	7	2	0.26	0.0254
Mammalia	Group 1 + Group 2	GO:0006210	myeloid dendritic cell cytokine production	7	2	0.26	0.0254
Mammalia	Group 1 + Group 2	GO:0001781	positive regulation of programmed necrotic cell death	7	2	0.26	0.0254
Mammalia	Group 1 + Group 2	GO:0043312	neutrophil apoptosis	8	2	0.30	0.0330
Mammalia	Group 1 + Group 2	GO:0042796	neutrophil degranulation	8	2	0.30	0.0330
Mammalia	Group 1 + Group 2	GO:0031358	snRNA transcription by RNA polymerase III	8	2	0.30	0.0330
Mammalia	Group 1 + Group 2	GO:0038196	response to type III interferon	8	2	0.30	0.0330
Mammalia	Group 1 + Group 2	GO:0002372	positive regulation of blood vessel endothelial cell proliferation involved in sprouting angiogenesis	8	2	0.30	0.0330
Mammalia	Group 1 + Group 2	GO:2000389	regulation of neutrophil extravasation	8	2	0.30	0.0330
Mammalia	Group 1 + Group 2	GO:0026774	negative regulation of acute inflammatory response	8	2	0.30	0.0330
Mammalia	Group 1 + Group 2	GO:0002638	negative regulation of immunoglobulin production	8	2	0.30	0.0330
Mammalia	Group 1 + Group 2	GO:0032650	interleukin-1 alpha production	8	2	0.30	0.0330
Mammalia	Group 1 + Group 2	GO:0032650	interleukin-1 alpha production	8	2	0.30	0.0330
Mammalia	Group 1 + Group 2	GO:0034297	interleukin-4-mediated signaling pathway	8	2	0.30	0.0330
Mammalia	Group 1 + Group 2	GO:0015158	microglial cell proliferation	9	2	0.33	0.0415
Mammalia	Group 1 + Group 2	GO:0071352	regulation of peptide-serine phosphorylation of STAT protein	9	2	0.33	0.0415
Mammalia	Group 1 + Group 2	GO:2000553	regulation of symbiosis	9	2	0.33	0.0415
Mammalia	Group 1 + Group 2	GO:0010918	positive regulation of mitochondrial membrane potential	9	2	0.33	0.0415
Mammalia	Group 1 + Group 2	GO:0033632	regulation of cell-cell adhesion mediated by integrin	9	2	0.33	0.0415
Mammalia	Group 1 + Group 2	GO:0045625	regulation of T-helper 1 cell differentiation	9	2	0.33	0.0415
Mammalia	Group 1 + Group 2	GO:1901625	regulation of T-helper 2 cell differentiation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0034349	cellular response to ergosterol	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:1920569	negative regulation of activation of Janus kinase activity	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0071660	positive regulation of IP-10 production	1	1	0.04	0.0370

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Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Mammalia	Group 1 + Group 2	GO:0044355	clearance of foreign intracellular DNA	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0071652	regulation of chemokine (C-C motif) ligand 1 production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0071654	positive regulation of chemokine (C-C motif) ligand 1 production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:002736	regulation of plasmacytoid dendritic cell cytokine production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:002737	negative regulation of plasmacytoid dendritic cell cytokine production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0071610	chemokine (C-C motif) ligand 1 production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0051673	disruption of plasma membrane integrity in another organism	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0150073	regulation of protein-gutamine gamma-glutamyltransferase activity	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0150074	positive regulation of protein-gutamine gamma-glutamyltransferase activity	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0032759	positive regulation of TRAIL production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:195151	negative regulation of voltage-gated sodium channel activity	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0160006	FC receptor-mediated immune complex endocytosis	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0032723	positive regulation of connective tissue growth factor production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:195154	negative regulation of membrane invagination	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0032745	positive regulation of interleukin-21 production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0160688	type I hypersensitivity	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:1904784	NLRP1 inflammasome complex assembly	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0035397	helper T cell enhancement of adaptive immune response	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:195223	epicardium morphogenesis	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0050902	leukocyte adhesive activation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0025158	lymphocyte chemotaxis across high endothelial venule	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0034156	negative regulation of toll-like receptor 7 signaling pathway	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:1919484	negative regulation of cell chemotaxis to fibroblast growth factor	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0120042	negative regulation of macrophage proliferation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0034125	negative regulation of MyD88-dependent toll-like receptor signaling pathway	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0060097	cytoskeletal rearrangement involved in phagocytosis, engulfment	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:2000422	regulation of eosinophil chemotaxis	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:00200423	positive regulation of eosinophil chemotaxis	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0090073	positive regulation of protein homodimerization activity	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0051977	lysophospholipid transport	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0032665	regulation of interleukin-21 production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0032679	regulation of TRAIL production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0032625	interleukin-21 production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0032639	TRAIL production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:001971	negative regulation of activation of membrane attack complex	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:003496	regulation of translational initiation by eIF2 alpha dephosphorylation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0036497	eIF2alpha dephosphorylation in response to endoplasmic reticulum stress	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:2000229	regulation of pancreatic stellate cell proliferation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0051041	positive regulation of calcium-independent cell-cell adhesion	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:2000231	positive regulation of pancreatic stellate cell proliferation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:1900450	negative regulation of glutamate receptor signaling pathway	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0018003	peptidyl-lysine N6-acetylation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:1903916	regulation of endoplasmic reticulum stress-induced eIF2 alpha dephosphorylation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:19103917	positive regulation of endoplasmic reticulum stress-induced eIF2 alpha dephosphorylation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0099046	clearance of foreign intracellular nucleic acids	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0150140	regulation of CD86 production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0150142	positive regulation of CD86 production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0150143	regulation of CD80 production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0150145	positive regulation of CD80 production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0036153	triacylglyceride acyl-chain remodeling	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0036155	triglyceride acyl-chain remodeling	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:1905675	negative regulation of adaptive immune memory response	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:00090320	regulation of chytromicron remnant clearance	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0090321	positive regulation of chytromicron remnant clearance	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0140121	Levy body formation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0140122	regulation of Levy body formation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:1902310	positive regulation of peptidyl-serine dephosphorylation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:2000545	negative regulation of endothelial cell chemotaxis to fibroblast growth factor	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0060101	negative regulation of phagocytosis, engulfment	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0070246	natural killer cell apoptotic process	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:1901251	positive regulation of lung goblet cell differentiation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:1901249	regulation of lung chondrocyte differentiation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:00310186	melatonin metabolic process	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0030187	melatonin biosynthetic process	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0072343	pancreatic stellate cell proliferation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0010185	regulation of cellular defense response	1	1	0.04	0.0370

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Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Mammalia	Group 1 + Group 2	GO:0010186	positive regulation of cellular defense response	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0002384	hepatic immune response	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:1903496	response to 11-deoxycorticosterone	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0002373	plasmacytoid dendritic cell cytokine production	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0098784	biofilm matrix organization	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0098786	biofilm matrix disassembly	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:002450	B cell antigen processing and presentation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0046603	negative regulation of mitotic centrosome separation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0002470	plasmacytoid dendritic cell antigen processing and presentation	1	1	0.04	0.0370
Mammalia	Group 1 + Group 2	GO:0072139	glomerular parietal epithelial cell differentiation	1	1	0.04	0.0370
Bivalvia	Group 1	GO:0080090	regulation of primary metabolic process	673	47	23.83	0.0231
Bivalvia	Group 1	GO:0019219	regulation of nucleobase-containing compound metabolic process	541	36	19.16	0.0270
Bivalvia	Group 1	GO:0051252	regulation of RNA metabolic process	517	32	18.31	0.0342
Bivalvia	Group 1	GO:0069550	response to stress	370	29	13.10	0.0270
Bivalvia	Group 1	GO:0035554	cellular response to stress	974	974	0.0419	0.0172
Bivalvia	Group 1	GO:0051172	negative regulation of nitrogen compound metabolic process	275	21	4.14	0.0075
Bivalvia	Group 1	GO:0031325	positive regulation of cellular metabolic process	117	15	4.43	0.0100
Bivalvia	Group 1	GO:0063110	DNA recombination	125	13	2.34	0.0100
Bivalvia	Group 1	GO:0051173	positive regulation of nitrogen compound metabolic process	66	13	2.34	0.0100
Bivalvia	Group 1	GO:0010629	negative regulation of gene expression	137	13	4.85	0.0336
Bivalvia	Group 1	GO:0048513	animal organ development	78	11	2.76	0.0172
Bivalvia	Group 1	GO:0045934	negative regulation of nucleobase-containing compound metabolic process	83	11	2.94	0.0172
Bivalvia	Group 1	GO:0006915	apoptotic process	64	11	2.27	0.0186
Bivalvia	Group 1	GO:0048892	negative regulation of DNA templated transcription	95	9	3.36	0.0062
Bivalvia	Group 1	GO:1902679	negative regulation of RNA biosynthetic process	59	9	2.09	0.0151
Bivalvia	Group 1	GO:0051253	negative regulation of RNA metabolic process	61	9	2.16	0.0192
Bivalvia	Group 1	GO:0064117	regulation of translation	52	8	1.84	0.0004
Bivalvia	Group 1	GO:0051726	regulation of cell cycle	75	8	2.66	0.0399
Bivalvia	Group 1	GO:0065009	regulation of molecular function	114	8	4.04	0.0485
Bivalvia	Group 1	GO:0045893	positive regulation of DNA templated transcription	67	7	2.37	0.0091
Bivalvia	Group 1	GO:0042981	regulation of apoptotic process	70	7	2.48	0.0115
Bivalvia	Group 1	GO:0043067	regulation of programmed cell death	72	7	2.55	0.0133
Bivalvia	Group 1	GO:000122	negative regulation of transcription by RNA polymerase II	31	5	1.10	0.0043
Bivalvia	Group 1	GO:0006402	mRNA catabolic process	35	5	1.24	0.0073
Bivalvia	Group 1	GO:0045944	positive regulation of transcription by RNA polymerase II	38	5	1.35	0.0103
Bivalvia	Group 1	GO:0071310	cellular response to organic substance	52	5	1.84	0.0359
Bivalvia	Group 1	GO:0006228	response to abiotic stimulus	53	5	1.88	0.0385
Bivalvia	Group 1	GO:0051248	negative regulation of protein metabolic process	55	5	1.95	0.0441
Bivalvia	Group 1	GO:0030155	regulation of cell adhesion	55	5	0.99	0.0004
Bivalvia	Group 1	GO:0045568	embryonic organ development	11	4	0.42	0.0006
Bivalvia	Group 1	GO:0051607	defense response to virus	12	4	0.46	0.0009
Bivalvia	Group 1	GO:0010569	regulation of double-strand break repair via homologous recombination	13	4	0.18	0.0036
Bivalvia	Group 1	GO:0000902	cell morphogenesis	5	4	1.10	0.0227
Bivalvia	Group 1	GO:000280	nuclear division	31	4	1.35	0.0441
Bivalvia	Group 1	GO:0050769	positive regulation of neurogenesis	38	4	0.95	0.0004
Bivalvia	Group 1	GO:0001819	positive regulation of cytokine production	5	3	0.18	0.0004
Bivalvia	Group 1	GO:0070192	chromosome organization involved in meiotic cell cycle	7	3	0.25	0.0014
Bivalvia	Group 1	GO:0073668	determination of left/right symmetry	7	3	0.25	0.0014
Bivalvia	Group 1	GO:0001894	tissue homeostasis	10	3	0.35	0.0044
Bivalvia	Group 1	GO:003007	heart morphogenesis	10	3	0.35	0.0044
Bivalvia	Group 1	GO:008285	negative regulation of cell population proliferation	10	3	0.35	0.0044
Bivalvia	Group 1	GO:0051093	negative regulation of developmental process	11	3	0.39	0.0059
Bivalvia	Group 1	GO:0015013	skeletal system development	11	3	0.46	0.0059
Bivalvia	Group 1	GO:0075117	muscle organ development	13	3	0.50	0.0119
Bivalvia	Group 1	GO:0061982	meiosis I cell cycle process	14	3	0.50	0.0119
Bivalvia	Group 1	GO:0007127	meiosis I	14	3	0.50	0.0119
Bivalvia	Group 1	GO:010257	NADH dehydrogenase complex assembly	17	3	0.60	0.0206
Bivalvia	Group 1	GO:0032981	mitochondrial respiratory chain complex I assembly	17	3	0.60	0.0206
Bivalvia	Group 1	GO:0022603	ribosomal small subunit biogenesis	17	3	0.74	0.0364
Bivalvia	Group 1	GO:0140013	regulation of anatomical structure morphogenesis	23	3	0.81	0.0460
Bivalvia	Group 1	GO:1905168	mitochondrial gene expression	23	3	0.81	0.0460
Bivalvia	Group 1	GO:0032488	positive regulation of double-strand break repair via homologous recombination	2	2	0.07	0.0013
Bivalvia	Group 1	GO:0022600	Cdc42 protein signal transduction	2	2	0.07	0.0013
Bivalvia	Group 1	GO:0035295	digestive system process	2	2	0.07	0.0013

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Dataset	Group of genes	GO ID	Term	Annotated	Significant	Expected	Classic Fisher
Bivalvia	Group 1	GO:0032232	negative regulation of actin filament bundle assembly	2	2	0.07	0.0013
Bivalvia	Group 1	GO:0033617	mitochondrial cytochrome c oxidase assembly	3	2	0.11	0.0037
Bivalvia	Group 1	GO:0045910	negative regulation of DNA recombination	3	2	0.11	0.0037
Bivalvia	Group 1	GO:2000179	regulation of neural precursor cell proliferation	3	2	0.11	0.0037
Bivalvia	Group 1	GO:0061383	trabecula morphogenesis	3	2	0.11	0.0037
Bivalvia	Group 1	GO:0051701	biological process involved in interaction with host	3	2	0.11	0.0037
Bivalvia	Group 1	GO:0010833	telomere maintenance via telomere lengthening	3	2	0.11	0.0037
Bivalvia	Group 1	GO:0007095	mitotic G2 DNA damage checkpoint signaling	3	2	0.11	0.0037
Bivalvia	Group 1	GO:0009959	mitochondrial RNA metabolic process	3	2	0.11	0.0037
Bivalvia	Group 1	GO:002064	epithelial cell development	3	2	0.11	0.0037
Bivalvia	Group 1	GO:0010001	glial cell differentiation	4	2	0.14	0.0071
Bivalvia	Group 1	GO:0046620	regulation of organ growth	4	2	0.14	0.0071
Bivalvia	Group 1	GO:0003190	atrioventricular valve formation	4	2	0.14	0.0071
Bivalvia	Group 1	GO:0098781	ncRNA transcription	4	2	0.14	0.0071
Bivalvia	Group 1	GO:0030514	negative regulation of BMP signaling pathway	4	2	0.14	0.0071
Bivalvia	Group 1	GO:0035023	regulation of Rho protein signal transduction	4	2	0.14	0.0071
Bivalvia	Group 1	GO:001947	synapse assembly	5	2	0.18	0.0116
Bivalvia	Group 1	GO:0001947	heart looping	5	2	0.18	0.0116
Bivalvia	Group 1	GO:0042026	atrioventricular valve formation	5	2	0.18	0.0116
Bivalvia	Group 1	GO:0003143	protein refolding	5	2	0.18	0.0116
Bivalvia	Group 1	GO:0061371	embryonic heart tube morphogenesis	5	2	0.18	0.0116
Bivalvia	Group 1	GO:0035050	determination of heart left/right asymmetry	5	2	0.18	0.0116
Bivalvia	Group 1	GO:0007162	embryonic heart tube development	5	2	0.18	0.0116
Bivalvia	Group 1	GO:0007129	negative regulation of cell adhesion	5	2	0.18	0.0116
Bivalvia	Group 1	GO:0036360	homologous chromosome pairing at meiosis	6	2	0.21	0.0170
Bivalvia	Group 1	GO:0001889	transcription by RNA polymerase I	6	2	0.21	0.0170
Bivalvia	Group 1	GO:0045143	liver development	6	2	0.21	0.0170
Bivalvia	Group 1	GO:0061008	homologous chromosome segregation	6	2	0.21	0.0170
Bivalvia	Group 1	GO:0010212	hepatobiliary system development	7	2	0.25	0.0233
Bivalvia	Group 1	GO:0030490	determination of cell adhesion	7	2	0.25	0.0233
Bivalvia	Group 1	GO:0061448	matured of SSU-RNA	7	2	0.25	0.0233
Bivalvia	Group 1	GO:0048732	connective tissue development	7	2	0.25	0.0233
Bivalvia	Group 1	GO:0001822	gland development	8	2	0.28	0.0304
Bivalvia	Group 1	GO:0045596	kidney development	8	2	0.28	0.0304
Bivalvia	Group 1	GO:0002711	negative regulation of cell differentiation	8	2	0.28	0.0304
Bivalvia	Group 1	GO:0002711	polysaccharide biosynthetic process	8	2	0.28	0.0304
Bivalvia	Group 1	GO:0005153	ossification	8	2	0.28	0.0304
Bivalvia	Group 1	GO:0045132	meiotic chromosome segregation	8	2	0.28	0.0304
Bivalvia	Group 1	GO:0016073	snrRNA metabolic process	8	2	0.28	0.0304
Bivalvia	Group 1	GO:0072001	renal system development	9	2	0.32	0.0304
Bivalvia	Group 1	GO:0060562	epithelial tube morphogenesis	9	2	0.32	0.0304
Bivalvia	Group 1	GO:0048562	embryonic organ morphogenesis	9	2	0.32	0.0304
Bivalvia	Group 1	GO:0042326	negative regulation of phosphorylation	10	2	0.35	0.0466
Bivalvia	Group 1	GO:0001934	positive regulation of protein phosphorylation	10	2	0.35	0.0466
Bivalvia	Group 1	GO:0022604	regulation of cell morphogenesis	10	2	0.35	0.0466
Bivalvia	Group 1	GO:0034134	Arp2/3 complex-mediated actin nucleation	10	2	0.35	0.0466
Bivalvia	Group 1	GO:0010669	epithelial structure maintenance	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0046621	negative regulation of organ growth	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0097222	mitochondrial mRNA polyadenylation	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0061668	mitochondrial ribosome assembly	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0048799	animal organ maturation	11	1	0.04	0.0354
Bivalvia	Group 1	GO:00444406	adhesion of symbiont to host	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0043247	telomere maintenance in response to DNA damage	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0061009	common bile duct development	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0006356	regulation of transcription by RNA polymerase I	11	1	0.04	0.0354
Bivalvia	Group 1	GO:1900044	regulation of protein K63-linked ubiquitination	11	1	0.04	0.0354
Bivalvia	Group 1	GO:1902285	semaphorin-plexin signalling pathway involved in neuron projection guidance	11	1	0.04	0.0354
Bivalvia	Group 1	GO:1900045	negative regulation of protein K63-linked ubiquitination	11	1	0.04	0.0354
Bivalvia	Group 1	GO:1902287	semaphorin-plexin signalling pathway involved in axon guidance	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0050772	positive regulation of axonogenesis	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0070977	bone maturation	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0031120	sno(s)RNA 3'-end processing	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0044650	sno(s)RNA pseudouridine synthetase	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0090646	adhesion of symbiont to host cell	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0016074	mitochondrial tRNA processing	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0032978	sno(s)RNA metabolic process	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0032979	protein insertion into membrane from inner side	11	1	0.04	0.0354
Bivalvia	Group 1	GO:1904862	mitochondrial inner membrane from matrix	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0072695	inhibitory synapse assembly	11	1	0.04	0.0354
Bivalvia	Group 1	GO:0072695	regulation of DNA recombination at telomere	11	1	0.04	0.0354

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Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Bivalvia	Group 1	GO:002230	positive regulation of defense response to virus by host	1	1	0.04	0.0354
Bivalvia	Group 1	GO:008653	lipopolysaccharide metabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0045943	positive regulation of transcription by RNA polymerase I	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0052572	response to host immune system process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0051274	beta-glucan biosynthetic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0051278	fungus-type cell wall polysaccharide biosynthetic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:1900220	semaphorinplexin signaling pathway involved in bone trabecula morphogenesis	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0086225	semaphorinplexin signaling pathway via death domain receptors	1	1	0.04	0.0354
Bivalvia	Group 1	GO:009301	snRNA transcription	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0015743	maleate transport	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0032065	maintenance of protein location in cell cortex	1	1	0.04	0.0354
Bivalvia	Group 1	GO:032196	retrotransposition	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0042149	cellular response to glucose starvation	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0052173	response to defenses of other organism	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0060704	(1->3)-beta-D-glucan metabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0007057	(1->3)-beta-D-glucan biosynthetic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0075136	presynapse assembly	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0099054	protein localization to synapse	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0035418	classification involved in bone maturation	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0043931	fungus-type cell wall polysaccharide metabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0071966	presynapse organization	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0099172	box H/ACA sno(s)RNA metabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0033979	regulation of protein polyubiquitination	1	1	0.04	0.0354
Bivalvia	Group 1	GO:1902914	negative regulation of protein polyubiquitination	1	1	0.04	0.0354
Bivalvia	Group 1	GO:1902915	protein deubiquitination involved in ubiquitin-dependent protein catabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0071947	regulation of defense response to virus by host	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0050691	lipid A metabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0046493	antibiotic catabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0017001	maleate transport	1	1	0.04	0.0354
Bivalvia	Group 1	GO:1901423	lipopolysaccharide metabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0070212	protein poly-ADP-ribosylation	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0070213	protein auto-ADP-ribosylation	1	1	0.04	0.0354
Bivalvia	Group 1	GO:1902414	protein localization to cell junction	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0011100	negative regulation of exit from mitosis	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0048144	fibroblast proliferation	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0048145	regulation of fibroblast proliferation	1	1	0.04	0.0354
Bivalvia	Group 1	GO:19048147	negative regulation of fibroblast proliferation	1	1	0.04	0.0354
Bivalvia	Group 1	GO:1901271	lipopolysaccharide biosynthetic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0036622	intraluminal bile duct development	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0030653	beta-lactam antibiotic metabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0030655	beta-lactam antibiotic catabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:00452200	response to host immune response	1	1	0.04	0.0354
Bivalvia	Group 1	GO:1900481	sno(s)RNA processing	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0091103	protection from non-homologous end joining at telomere	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0048634	lipopolysaccharide biosynthetic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0020333	regulation of muscle organ development	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0010526	retrotransposon silencing	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0046402	O antigen metabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0042783	evasion of host immune response	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0043144	sno(s)RNA processing	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0031848	protection from non-homologous end joining at telomere	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0061430	bone trabecula morphogenesis	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0020033	anticigenic variation	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0009963	mitochondrial RNA processing	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0007168	receptor giantry cyclase signalling pathway	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0043628	regulatory ncRNA 3'-end processing	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0035279	mRNA-mediated gene silencing by mRNA destabilization	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0016999	antibiotic metabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0072340	lactam catabolic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:1902041	regulation of extrinsic apoptotic signalling pathway via death domain receptors	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0009243	O antigen biosynthetic process	1	1	0.04	0.0354
Bivalvia	Group 1	GO:009245	lipid A biosynthetic process	1	1	0.04	0.0354

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Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Bivalvia	Group 1	GO:0033687	osteoblast proliferation	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0033688	regulation of osteoblast proliferation	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0033689	negative regulation of osteoblast proliferation	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0030277	maintenance of gastrointestinal epithelium	1	1	0.04	0.0354
Bivalvia	Group 1	GO:0048239	negative regulation of DNA recombination at telomere	1	1	0.04	0.0354
Drosophila	Group 1	GO:0051276	chromosome organization	331	18	11.07	0.0293
Drosophila	Group 1	GO:0054532	metabolic chromosome segregation	64	7	2.14	0.0053
Drosophila	Group 1	GO:0007131	reciprocal meiotic recombination	37	6	1.24	0.0113
Drosophila	Group 1	GO:0031146	SCF-dependent proteasomal ubiquitin-dependent protein catabolic process	57	5	1.91	0.0413
Drosophila	Group 1	GO:0007141	male meiosis I	13	4	0.43	0.0007
Drosophila	Group 1	GO:1902725	regulation of chromatin organization	35	4	1.17	0.0285
Drosophila	Group 1	GO:0042078	germline stem cell division	36	4	1.20	0.0312
Drosophila	Group 1	GO:0010526	retrotransposon silencing	8	3	0.27	0.0018
Drosophila	Group 1	GO:0071218	cellular response to misfolded protein	12	3	0.40	0.0065
Drosophila	Group 1	GO:0034508	centromere complex assembly	19	3	0.64	0.0242
Drosophila	Group 1	GO:0007080	mitotic metaphase chromosome alignment	23	3	0.77	0.0400
Drosophila	Group 1	GO:0007020	microtubule nucleation	23	3	0.77	0.0400
Drosophila	Group 1	GO:0031445	regulation of heterochromatin formation	25	3	0.84	0.0495
Drosophila	Group 1	GO:0120261	regulation of heterochromatin organization	25	3	0.84	0.0495
Drosophila	Group 1	GO:0048136	male germ-line cyst formation	2	2	0.07	0.0111
Drosophila	Group 1	GO:0061964	negative regulation of entry into reproductive diapause	5	2	0.17	0.0104
Drosophila	Group 1	GO:0051382	kinetochore assembly	5	2	0.17	0.0104
Drosophila	Group 1	GO:0055116	entry into reproductive diapause	6	2	0.20	0.0153
Drosophila	Group 1	GO:0071712	ER-associated misfolded protein catabolic process	6	2	0.20	0.0153
Drosophila	Group 1	GO:0061963	regulation of entry into reproductive diapause	6	2	0.20	0.0153
Drosophila	Group 1	GO:0043984	histone H4-K16 acetylation	6	2	0.20	0.0153
Drosophila	Group 1	GO:0055115	entry into diapause	7	2	0.23	0.0210
Drosophila	Group 1	GO:1990834	response to odorant	7	2	0.23	0.0210
Drosophila	Group 1	GO:0042795	snRNA transcription by RNA polymerase II	8	2	0.27	0.0273
Drosophila	Group 1	GO:00092611	dormancy process	8	2	0.27	0.0273
Drosophila	Group 1	GO:0071786	endothelial potassium ion homeostasis	8	2	0.30	0.0344
Drosophila	Group 1	GO:0030007	intracellular potassium ion homeostasis	9	2	0.30	0.0344
Drosophila	Group 1	GO:0051383	kinetochore organization	9	2	0.30	0.0344
Drosophila	Group 1	GO:0040020	regulation of meiotic nuclear division	9	2	0.30	0.0344
Drosophila	Group 1	GO:043967	histone H4 acetylation	9	2	0.30	0.0344
Drosophila	Group 1	GO:0036376	sodium ion export across plasma membrane	9	2	0.30	0.0344
Drosophila	Group 1	GO:0006883	intracellular sodium ion homeostasis	9	2	0.30	0.0344
Drosophila	Group 1	GO:0001015	snRNA transcription by RNA polymerase II	1	1	0.03	0.0335
Drosophila	Group 1	GO:0010778	meiotic DNA repair synthesis involved in reciprocal meiotic recombination	1	1	0.03	0.0335
Drosophila	Group 1	GO:0009302	sn(s)RNA transcription	1	1	0.03	0.0335
Drosophila	Group 1	GO:0051308	male meiosis chromosome separation	1	1	0.03	0.0335
Drosophila	Group 1	GO:0051415	microtubule nucleation by interphase microtubule organizing center	1	1	0.03	0.0335
Mammalia	Group 1	GO:0009695	immune response	1,297	120	38.75	0.0027
Mammalia	Group 1	GO:0045087	innate immune response	647	69	19.33	0.0000
Mammalia	Group 1	GO:0050778	positive regulation of immune response	419	47	12.52	0.0068
Mammalia	Group 1	GO:002250	adaptive immune response	342	39	10.22	0.0000
Mammalia	Group 1	GO:002694	regulation of leukocyte activation	456	39	13.62	0.0052
Mammalia	Group 1	GO:001819	positive regulation of cytokine production	402	37	12.01	0.0025
Mammalia	Group 1	GO:002274	cytokine-mediated signaling pathway	382	35	5.56	0.0294
Mammalia	Group 1	GO:0042742	leukocyte proliferation	273	24	11.41	0.0000
Mammalia	Group 1	GO:0042110	defense response to bacterium	233	32	6.96	0.0000
Mammalia	Group 1	GO:0050777	T cell activation	432	31	12.91	0.0133
Mammalia	Group 1	GO:002697	regulation of immune effector process	308	31	7.17	0.0057
Mammalia	Group 1	GO:0051607	adaptive immune response	456	39	10.22	0.0000
Mammalia	Group 1	GO:0002694	regulation of leukocyte activation	402	37	13.62	0.0000
Mammalia	Group 1	GO:001819	myeloid leukocyte activation	195	26	5.83	0.0294
Mammalia	Group 1	GO:000274	positive regulation of cytokine production	186	25	11.41	0.0000
Mammalia	Group 1	GO:002703	leukocyte-mediated signaling pathway	382	35	5.56	0.0294
Mammalia	Group 1	GO:0070661	leukocyte proliferation	273	24	8.16	0.0271
Mammalia	Group 1	GO:0019061	positive regulation of defense response	233	32	4.57	0.0449
Mammalia	Group 1	GO:0031349	mononuclear cell differentiation	432	31	7.17	0.0482
Mammalia	Group 1	GO:1903131	regulation of inflammatory response	308	31	7.68	0.0025
Mammalia	Group 1	GO:0050727	humoral immune response	300	21	8.96	0.0001
Mammalia	Group 1	GO:0009599	immune response-regulating cell surface receptor signaling pathway	177	21	5.29	0.0018
Mammalia	Group 1	GO:002766	cell killing	153	20	4.57	0.0449
Mammalia	Group 1	GO:002699	positive regulation of immune effector process	208	20	6.21	0.0477
Mammalia	Group 1	GO:0030217	T cell differentiation	255	18	7.62	0.0441
Mammalia	Group 1	GO:0045089	positive regulation of innate immune response	113	16	3.38	0.0139
Mammalia	Group 1	GO:0071222	cellular response to lipopolysaccharide	164	15	4.90	0.0045
Mammalia	Group 1	GO:1903555	regulation of tumor necrosis factor superfamily cytokine production	137	15	4.09	0.0045
Mammalia	Group 1	GO:0010466	negative regulation of peptidase activity	163	14	0.0004	0.0004

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Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Mammalia	Group 1	GO:0002429	immune response-activating cell surface receptor signaling pathway	164	14	4.90	0.0052
Mammalia	Group 1	GO:0032680	regulation of tumor necrosis factor production	133	14	3.97	0.0142
Mammalia	Group 1	GO:0032640	tumor necrosis factor production	133	14	3.97	0.0142
Mammalia	Group 1	GO:0002705	positive regulation of leukocyte mediated immunity	133	14	3.38	0.0395
Mammalia	Group 1	GO:0050829	defense response to Gram-negative bacterium	66	13	1.97	0.0000
Mammalia	Group 1	GO:002444	myeloid leukocyte mediated immunity	90	13	2.69	0.0001
Mammalia	Group 1	GO:0051091	positive regulation of DNA-binding transcription factor activity	227	13	6.78	0.0191
Mammalia	Group 1	GO:0002456	T cell mediated immunity	82	13	2.45	0.0395
Mammalia	Group 1	GO:0031341	regulation of cell killing	71	12	2.12	0.0042
Mammalia	Group 1	GO:002695	negative regulation of leukocyte activation	148	12	4.42	0.0178
Mammalia	Group 1	GO:0002286	T cell activation involved in immune response	94	12	2.81	0.0194
Mammalia	Group 1	GO:0050866	negative regulation of cytokine production	165	12	4.93	0.0351
Mammalia	Group 1	GO:0001818	negative regulation of cytokine production	225	12	6.72	0.0381
Mammalia	Group 1	GO:0050830	defense response to Gram-positive bacterium	87	11	2.60	0.0001
Mammalia	Group 1	GO:0002275	myeloid cell activation involved in immune response	83	11	2.48	0.0010
Mammalia	Group 1	GO:0050909	sensory perception of taste	52	11	1.55	0.0289
Mammalia	Group 1	GO:0042102	positive regulation of T cell proliferation	85	10	2.54	0.0002
Mammalia	Group 1	GO:0043299	leukocyte degranulation	62	10	1.85	0.0005
Mammalia	Group 1	GO:0030101	natural killer cell activation	66	10	1.97	0.0009
Mammalia	Group 1	GO:0097696	cell surface receptor signaling pathway via ST _{AT}	139	10	4.15	0.0089
Mammalia	Group 1	GO:0016064	immunoglobulin mediated immune response	107	10	3.20	0.0321
Mammalia	Group 1	GO:0019724	B cell mediated immunity	108	10	3.23	0.0336
Mammalia	Group 1	GO:0032760	positive regulation of tumor necrosis factor production	84	9	2.51	0.0009
Mammalia	Group 1	GO:0006956	complement activation	47	9	1.40	0.0100
Mammalia	Group 1	GO:0050729	positive regulation of inflammatory response	122	9	3.64	0.0109
Mammalia	Group 1	GO:0051250	negative regulation of lymphocyte activation	124	9	3.70	0.0121
Mammalia	Group 1	GO:0050832	defense response to fungus	48	9	1.43	0.0126
Mammalia	Group 1	GO:0002823	negative regulation of adaptive immune response [-.-]	39	9	1.17	0.0174
Mammalia	Group 1	GO:0007342	fusion of sperm to egg plasma membrane involved in single fertilization	25	8	0.75	0.0000
Mammalia	Group 1	GO:0001580	detection of chemical stimulus involved in sensory perception of bitter taste	29	8	0.87	0.0000
Mammalia	Group 1	GO:0002218	activation of innate immune response	45	8	1.34	0.0000
Mammalia	Group 1	GO:0045649	regulation of type II interferon production	88	8	2.63	0.0048
Mammalia	Group 1	GO:0045576	masitin cell activation	54	8	1.61	0.0063
Mammalia	Group 1	GO:0071346	cellular response to type II interferon	98	8	2.93	0.0090
Mammalia	Group 1	GO:001959	regulation of cytokine-mediated signaling pathway	114	8	3.41	0.0209
Mammalia	Group 1	GO:0030317	flagellated sperm motility	119	8	3.55	0.0263
Mammalia	Group 1	GO:0097722	sperm motility	119	8	3.55	0.0263
Mammalia	Group 1	GO:0042100	B cell proliferation	76	9	2.27	0.0261
Mammalia	Group 1	GO:0009620	response to fungus	58	9	1.73	0.0323
Mammalia	Group 1	GO:0035036	sperm-egg recognition process	46	9	1.37	0.0400
Mammalia	Group 1	GO:0071887	leukocyte apoptotic process	98	9	2.93	0.0468
Mammalia	Group 1	GO:0007342	fusion of sperm to egg plasma membrane involved in single fertilization	25	8	0.75	0.0000
Mammalia	Group 1	GO:0001580	detection of chemical stimulus involved in sensory perception of bitter taste	29	8	0.87	0.0000
Mammalia	Group 1	GO:0002218	activation of innate immune response	45	8	1.34	0.0000
Mammalia	Group 1	GO:0032649	regulation of type II interferon production	88	8	2.63	0.0048
Mammalia	Group 1	GO:0045576	masitin cell activation	54	8	1.61	0.0063
Mammalia	Group 1	GO:0071346	cellular response to type II interferon	98	8	2.93	0.0090
Mammalia	Group 1	GO:001959	regulation of cytokine-mediated signaling pathway	114	8	3.41	0.0209
Mammalia	Group 1	GO:0030317	flagellated sperm motility	119	8	3.55	0.0263
Mammalia	Group 1	GO:0097722	sperm motility	119	8	3.55	0.0263
Mammalia	Group 1	GO:0035834	male gonad development	120	8	3.58	0.0275
Mammalia	Group 1	GO:0045546	development of primary male sexual characteristics	121	8	3.61	0.0287
Mammalia	Group 1	GO:0045759	regulation of response to cytokine stimulus	124	8	3.70	0.0325
Mammalia	Group 1	GO:0060294	cilium movement involved in cell motility	128	8	3.82	0.0382
Mammalia	Group 1	GO:0019739	binding of sperm to zona pellucida	35	7	1.05	0.0001
Mammalia	Group 1	GO:0019731	antibacterial humorall response	40	7	1.19	0.0002
Mammalia	Group 1	GO:0032757	positive regulation of interleukin-8 production	54	7	1.61	0.0114
Mammalia	Group 1	GO:0043303	positive regulation of chemoattractant production	44	7	1.28	0.0101
Mammalia	Group 1	GO:0032722	positive regulation of chemokine production	44	7	1.31	0.0114
Mammalia	Group 1	GO:002251	organ or tissue specific immune response	60	7	1.79	0.0020
Mammalia	Group 1	GO:0098586	cellular response to virus by host	22	7	0.66	0.0036
Mammalia	Group 1	GO:0030593	neutrophil chemotaxis	78	7	2.39	0.0099
Mammalia	Group 1	GO:0043303	neutrophil chemotaxis	43	7	1.28	0.0101
Mammalia	Group 1	GO:002279	neutrophil degranulation	29	6	0.87	0.0002
Mammalia	Group 1	GO:1904892	regulation of receptor signaling pathway via ST _{AT}	83	7	2.48	0.0120
Mammalia	Group 1	GO:0023886	regulation of myeloid leukocyte mediated immunity	45	7	1.34	0.0128
Mammalia	Group 1	GO:002448	masitin cell mediated immunity	46	7	1.37	0.0143
Mammalia	Group 1	GO:0050691	regulation of defense response to virus by host	37	7	1.11	0.0336
Mammalia	Group 1	GO:00335303	regulation of dephosphorylation	103	7	3.08	0.0346
Mammalia	Group 1	GO:0002825	regulation of T-helper 1 type immune response	29	6	0.87	0.0002
Mammalia	Group 1	GO:0042119	neutrophil activation	32	6	0.96	0.0003
Mammalia	Group 1	GO:0045071	negative regulation of viral genome replication	39	6	1.17	0.0010
Mammalia	Group 1	GO:002711	positive regulation of T cell mediated immunity	44	6	1.31	0.0019
Mammalia	Group 1	GO:0002639	positive regulation of immunoglobulin production	44	6	1.31	0.0019
Mammalia	Group 1	GO:0042267	natural killer cell mediated cytotoxicity	44	6	1.31	0.0019
Mammalia	Group 1	GO:0032731	positive regulation of interleukin-1 beta production	46	6	1.37	0.0023

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Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Mammalia	Group 1	GO:0060337	type I interferon-mediated signaling pathway	52	6	1.55	0.0044
Mammalia	Group 1	GO:0030888	regulation of B cell proliferation	54	6	1.61	0.0053
Mammalia	Group 1	GO:0051293	establishment of spindle localization	54	6	1.61	0.0053
Mammalia	Group 1	GO:0032729	positive regulation of type II interferon production	62	6	1.85	0.0103
Mammalia	Group 1	GO:0010921	regulation of phosphatase activity	65	6	1.94	0.0129
Mammalia	Group 1	GO:0027220	regulation of cytokine production involved in immune response	66	6	1.97	0.0138
Mammalia	Group 1	GO:0092395	mucosal immune response	19	6	0.57	0.0218
Mammalia	Group 1	GO:2000106	regulation of leukocyte apoptotic process	75	6	2.24	0.0245
Mammalia	Group 1	GO:0046425	regulation of receptor signaling pathway via JAK-STAT	78	6	2.33	0.0290
Mammalia	Group 1	GO:0032755	positive regulation of interleukin-6 production	83	6	2.48	0.0377
Mammalia	Group 1	GO:0061760	antifungal innate immune response	17	5	0.51	0.0001
Mammalia	Group 1	GO:0035458	response to interferon-beta	21	5	0.63	
Mammalia	Group 1	GO:0048240	sperm capacitation	25	5	0.75	0.0008
Mammalia	Group 1	GO:0001914	regulation of T cell mediated cytotoxicity	25	5	0.75	0.0008
Mammalia	Group 1	GO:0002330	positive regulation of defense response to virus by host	27	5	0.81	0.0011
Mammalia	Group 1	GO:0002830	positive regulation of type 2 immune response	14	5	0.42	0.0019
Mammalia	Group 1	GO:0046596	regulation of viral entry into host cell	31	5	0.93	0.0021
Mammalia	Group 1	GO:001774	microglial cell activation	35	5	1.05	0.0036
Mammalia	Group 1	GO:0140632	canonical inflammasome complex assembly	35	5	1.05	0.0036
Mammalia	Group 1	GO:0006953	acute-phase response	38	5	1.14	0.0052
Mammalia	Group 1	GO:0140374	antiviral innate immune response	41	5	1.22	0.0072
Mammalia	Group 1	GO:0001912	positive regulation of leukocyte mediated cytotoxicity	42	5	1.25	0.0080
Mammalia	Group 1	GO:0032720	positive regulation of tumor necrosis factor production	47	5	1.40	0.0127
Mammalia	Group 1	GO:1901774	heterotypic cell-cell adhesion	47	5	1.40	0.0127
Mammalia	Group 1	GO:1935556	negative regulation of tumor necrosis factor superfamily cytokine production	49	5	1.46	0.0151
Mammalia	Group 1	GO:0070228	regulation of lymphocyte apoptotic process	50	5	1.49	0.0164
Mammalia	Group 1	GO:0011961	positive regulation of cytokine-mediated signaling pathway	50	5	1.49	0.0164
Mammalia	Group 1	GO:0060760	positive regulation of response to cytokine stimulus	57	5	1.76	0.0273
Mammalia	Group 1	GO:0038034	signal transduction in absence of ligand	59	5	1.76	0.0311
Mammalia	Group 1	GO:0017912	extrinsic apoptotic signaling pathway in absence of ligand	59	5	1.76	0.0311
Mammalia	Group 1	GO:0071260	cellular response to mechanical stimulus	60	5	1.79	
Mammalia	Group 1	GO:0031640	cell killing of cells of another organism	64	5	1.91	0.0421
Mammalia	Group 1	GO:0061844	antimicrobial humoral immune response mediated by antimicrobial peptide	64	5	1.91	0.0421
Mammalia	Group 1	GO:0002227	innate immune response in mucosa	11	4	0.33	0.0002
Mammalia	Group 1	GO:0035455	response to interferon-alpha	14	4	0.42	0.0006
Mammalia	Group 1	GO:0042454	eosinophil chemoataxis	17	4	0.51	0.0014
Mammalia	Group 1	GO:0027177	T-helper 17 cell lineage commitment	21	4	0.63	0.0031
Mammalia	Group 1	GO:0032740	positive regulation of natural killer cell mediated immunity	23	4	0.69	0.0044
Mammalia	Group 1	GO:0043032	positive regulation of interleukin-17 production	23	4	0.69	0.0044
Mammalia	Group 1	GO:0070498	positive regulation of macrophage activation	24	4	0.72	0.0052
Mammalia	Group 1	GO:0070269	interleukin-1-mediated signaling pathway	26	4	0.78	0.0070
Mammalia	Group 1	GO:0031295	pyroptotic inflammatory response	27	4	0.81	0.0080
Mammalia	Group 1	GO:0031160	T cell costimulation	28	4	0.84	0.0091
Mammalia	Group 1	GO:0019884	antigen processing and presentation of exogenous antigen	29	4	0.87	0.0103
Mammalia	Group 1	GO:0002446	neutrophil mediated inflammatory response	29	4	0.90	0.0116
Mammalia	Group 1	GO:2000352	negative regulation of endothelial cell apoptosis	30	4	0.90	0.0116
Mammalia	Group 1	GO:0032743	positive regulation of interleukin-2 production	30	4	0.90	0.0116
Mammalia	Group 1	GO:1900225	regulation of NLRP3 inflammasome complex assembly	30	4	0.90	0.0116
Mammalia	Group 1	GO:0043330	response to exogenous dsRNA	31	4	0.93	0.0131
Mammalia	Group 1	GO:0046006	regulation of activated T cell proliferation	32	4	0.96	0.0146
Mammalia	Group 1	GO:0032814	regulation of natural killer cell activation	32	4	0.96	0.0146
Mammalia	Group 1	GO:0044546	NLRP3 inflammasome complex assembly	33	4	0.99	0.0162
Mammalia	Group 1	GO:0050798	activated T cell proliferation	34	4	1.02	0.0180
Mammalia	Group 1	GO:0006968	cellular defense response	34	4	1.02	0.0180
Mammalia	Group 1	GO:0002701	negative regulation of production of molecular mediator of immune response	35	4	1.05	0.0198
Mammalia	Group 1	GO:0043331	response to dsRNA	36	4	1.08	0.0218
Mammalia	Group 1	GO:002691	regulation of cellular extravasation	38	4	1.14	0.0260
Mammalia	Group 1	GO:2003551	regulation of endothelial cell apoptotic process	43	4	1.28	0.0388
Mammalia	Group 1	GO:0032653	regulation of interleukin-10 production	45	4	1.34	0.0447
Mammalia	Group 1	GO:0032613	positive regulation of T-helper cell differentiation	45	4	1.34	0.0447
Mammalia	Group 1	GO:0046624	regulation of phosphoprotein phosphatase activity	17	4	0.51	0.0479
Mammalia	Group 1	GO:0043666	positive regulation of receptor signaling pathway via STAT	46	4	1.37	0.0479
Mammalia	Group 1	GO:1904894	Fc epsilon receptor signaling pathway	5	3	0.15	0.0003
Mammalia	Group 1	GO:0045959	negative regulation of complement activation, classical pathway	5	3	0.15	0.0003
Mammalia	Group 1	GO:0051838	cytosis by host of symbiont cells	8	3	0.24	0.0013
Mammalia	Group 1	GO:0097527	neurotoxic signaling pathway	8	3	0.24	0.0013
Mammalia	Group 1	GO:0069242	activation-induced cell death of T cells	10	3	0.30	0.0027
Mammalia	Group 1	GO:1901731	positive regulation of platelet aggregation	10	3	0.30	0.0027

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Dataset	Group of genes	GO ID	Term	Annotated	Significant	Expected	Classic Fisher
Mammalia	Group 1	GO:2000551	regulation of T-helper 2 cell cytokine production	10	3	0.33	0.0027
Mammalia	Group 1	GO:007343	egg activation	11	3	0.33	0.0037
Mammalia	Group 1	GO:0035723	interleukin-15-mediated signaling pathway	11	3	0.33	0.0037
Mammalia	Group 1	GO:0150079	negative regulation of neuroinflammatory response	12	3	0.36	0.0048
Mammalia	Group 1	GO:0063233	natural killer cell activation involved in immune response	13	3	0.39	0.0061
Mammalia	Group 1	GO:0032197	retrotransposition	13	3	0.39	0.0061
Mammalia	Group 1	GO:0060046	regulation of acrosome reaction	13	3	0.39	0.0061
Mammalia	Group 1	GO:0072683	T cell extravasation	13	3	0.39	0.0061
Mammalia	Group 1	GO:0043306	positive regulation of mast cell degranulation	14	3	0.42	0.0075
Mammalia	Group 1	GO:006957	complement activation, alternative pathway	14	3	0.42	0.0075
Mammalia	Group 1	GO:0050855	regulation of B cell receptor signaling pathway	14	3	0.42	0.0075
Mammalia	Group 1	GO:0230355	CD40 signaling pathway	14	3	0.42	0.0075
Mammalia	Group 1	GO:1930277	regulation of opsonization	14	3	0.42	0.0075
Mammalia	Group 1	GO:0002710	negative regulation of T cell mediated immunity	15	3	0.45	0.0092
Mammalia	Group 1	GO:0043031	negative regulation of macrophage activation	15	3	0.45	0.0092
Mammalia	Group 1	GO:0046597	negative regulation of viral entry into host cell	16	3	0.48	0.0111
Mammalia	Group 1	GO:0052888	positive regulation of myeloid leukocyte mediated immunity	17	3	0.51	0.0132
Mammalia	Group 1	GO:0002822	positive regulation of T-helper 1 type immune response	18	3	0.54	0.0155
Mammalia	Group 1	GO:0045346	regulation of MHC class II biosynthetic process	18	3	0.54	0.0155
Mammalia	Group 1	GO:0042832	defense response to protozoan	19	3	0.57	0.0180
Mammalia	Group 1	GO:0040019	positive regulation of embryonic development	19	3	0.57	0.0180
Mammalia	Group 1	GO:0060340	positive regulation of type I interferon-mediated signaling pathway	19	3	0.57	0.0180
Mammalia	Group 1	GO:0006333	type II interferon-mediated signaling pathway	19	3	0.57	0.0180
Mammalia	Group 1	GO:0045342	MHC class II biosynthetic process	19	3	0.57	0.0180
Mammalia	Group 1	GO:0032753	positive regulation of interleukin-4 production	20	3	0.60	0.0207
Mammalia	Group 1	GO:0001916	positive regulation of T cell mediated cytotoxicity	20	3	0.60	0.0207
Mammalia	Group 1	GO:001562	response to protozoan	21	3	0.63	0.0236
Mammalia	Group 1	GO:1903901	negative regulation of viral life cycle	21	3	0.63	0.0236
Mammalia	Group 1	GO:0038821	modulation of process of another organism	21	3	0.63	0.0236
Mammalia	Group 1	GO:0027226	positive regulation of T cell cytokine production	22	3	0.66	0.0268
Mammalia	Group 1	GO:0042104	positive regulation of activated T cell proliferation	22	3	0.66	0.0268
Mammalia	Group 1	GO:0010743	regulation of macrophage derived foam cell differentiation	25	3	0.75	0.0374
Mammalia	Group 1	GO:0032673	regulation of interleukin-4 production	25	3	0.75	0.0374
Mammalia	Group 1	GO:0032633	interleukin-4 production	25	3	0.75	0.0374
Mammalia	Group 1	GO:0071354	cellular response to interleukin-6	26	3	0.78	0.0414
Mammalia	Group 1	GO:0010922	positive regulation of phosphatase activity	26	3	0.78	0.0414
Mammalia	Group 1	GO:1930219	regulation of cilium-dependent cell motility	28	3	0.84	0.0500
Mammalia	Group 1	GO:0070741	response to interleukin-6	28	3	0.84	0.0500
Mammalia	Group 1	GO:0060295	regulation of ciliary movement involved in cell motility	28	3	0.84	0.0500
Mammalia	Group 1	GO:0042269	regulation of natural killer cell mediated cytotoxicity	28	3	0.84	0.0500
Mammalia	Group 1	GO:0097528	execution phase of necrosis	2	2	0.06	0.0009
Mammalia	Group 1	GO:1936660	negative regulation of complement-dependent cytotoxicity	2	2	0.06	0.0009
Mammalia	Group 1	GO:0030887	positive regulation of myeloid dendritic cell activation	6	2	0.12	0.0051
Mammalia	Group 1	GO:0009609	response to symbiotic bacterium	4	2	0.12	0.0051
Mammalia	Group 1	GO:0048080	antigen processing and presentation, endogenous lipid antigen via MHC class Ib	4	2	0.12	0.0051
Mammalia	Group 1	GO:002765	immune response-inhibiting signal transduction	5	2	0.15	0.0084
Mammalia	Group 1	GO:0042630	positive regulation of T-helper 2 cell differentiation	5	2	0.15	0.0084
Mammalia	Group 1	GO:0043313	regulation of neutrophil degranulation	6	2	0.18	0.0123
Mammalia	Group 1	GO:0106300	protein-DNA covalent cross-linking repair	6	2	0.18	0.0123
Mammalia	Group 1	GO:0060545	positive regulation of necroptotic process	6	2	0.18	0.0123
Mammalia	Group 1	GO:0002826	negative regulation of T-helper 1 type immune response	6	2	0.18	0.0123
Mammalia	Group 1	GO:0033141	positive regulation of peptidyl-serine phosphorylation of STAT protein	6	2	0.18	0.0123
Mammalia	Group 1	GO:0001781	neutrophil apoptotic process	7	2	0.21	0.0169
Mammalia	Group 1	GO:002372	myeloid dendritic cell cytokine production	7	2	0.21	0.0169
Mammalia	Group 1	GO:0060335	regulation of myeloid dendritic cell cytokine production	7	2	0.21	0.0169
Mammalia	Group 1	GO:0002735	positive regulation of type II interferon-mediated signaling pathway	6	2	0.21	0.0169
Mammalia	Group 1	GO:2000659	antigen processing and presentation, exogenous lipid antigen via MHC class Ib	7	2	0.21	0.0169
Mammalia	Group 1	GO:0045341	type III interferon-mediated signalling pathway	7	2	0.21	0.0169
Mammalia	Group 1	GO:0045343	MHC class I biosynthetic process	6	2	0.21	0.0169
Mammalia	Group 1	GO:0060467	negative regulation of fertilization	7	2	0.21	0.0169
Mammalia	Group 1	GO:0060468	prevention of polyspermy	7	2	0.21	0.0169
Mammalia	Group 1	GO:0062100	positive regulation of programmed necrotic cell death	7	2	0.21	0.0169
Mammalia	Group 1	GO:0071358	cellular response to type III interferon	7	2	0.21	0.0169
Mammalia	Group 1	GO:0010940	positive regulation of necrotic cell death	7	2	0.21	0.0169
Mammalia	Group 1	GO:0042796	sRNA transcription by RNA polymerase III	8	8	0.24	0.0221

Tab. S10 continued from previous page

Dataset	Group of genes	GO ID	Term	Annotated	Significant	Expected	Classic Fisher
Mammalia	Group 1	GO:0043312	neutrophil degranulation	8	2	0.24	0.0221
Mammalia	Group 1	GO:1903589	positive regulation of blood vessel endothelial cell proliferation involved in sprouting angiogenesis	8	2	0.24	0.0221
Mammalia	Group 1	GO:0006977	DNA damage response, signal transduction by p53 class mediator resulting in cell cycle arrest	8	2	0.24	0.0221
Mammalia	Group 1	GO:0038110	interleukin-2-mediated signaling pathway	8	2	0.24	0.0221
Mammalia	Group 1	GO:0002638	negative regulation of immunoglobulin production	8	2	0.24	0.0221
Mammalia	Group 1	GO:0032650	regulation of interleukin-1 alpha production	8	2	0.24	0.0221
Mammalia	Group 1	GO:0035771	interleukin-4-mediated signaling pathway	8	2	0.24	0.0221
Mammalia	Group 1	GO:0032610	interleukin-1 alpha production	8	2	0.24	0.0221
Mammalia	Group 1	GO:0071352	cellular response to interleukin-2	8	2	0.24	0.0221
Mammalia	Group 1	GO:2000553	positive regulation of T-helper 2 cell cytokine production	8	2	0.24	0.0221
Mammalia	Group 1	GO:0034342	response to type III interferon	8	2	0.24	0.0221
Mammalia	Group 1	GO:0044406	adhesion of symbiont to host	8	2	0.27	0.0279
Mammalia	Group 1	GO:0048625	regulation of T-helper 1 cell differentiation	9	2	0.27	0.0279
Mammalia	Group 1	GO:0033139	regulation of peptidyl-serine phosphorylation of STAT protein	9	2	0.27	0.0279
Mammalia	Group 1	GO:0061518	microglial cell proliferation	9	2	0.27	0.0279
Mammalia	Group 1	GO:0070669	response to interleukin-2	9	2	0.27	0.0279
Mammalia	Group 1	GO:0010918	positive regulation of mitochondrial membrane potential	9	2	0.27	0.0279
Mammalia	Group 1	GO:1920563	regulation of neutrophil activation	10	2	0.30	0.0342
Mammalia	Group 1	GO:0002430	complement receptor mediated signaling pathway	10	2	0.30	0.0342
Mammalia	Group 1	GO:0032754	positive regulation of interleukin-5 production	10	2	0.30	0.0342
Mammalia	Group 1	GO:0010526	retrotransposon silencing	10	2	0.30	0.0342
Mammalia	Group 1	GO:0010528	regulation of transposition	10	2	0.30	0.0342
Mammalia	Group 1	GO:0010529	negative regulation of transposition	10	2	0.30	0.0342
Mammalia	Group 1	GO:1920563	regulation of neutrophil activation	10	2	0.30	0.0342
Mammalia	Group 1	GO:0002430	complement receptor mediated signaling pathway	11	2	0.33	0.0410
Mammalia	Group 1	GO:0032754	positive regulation of interleukin-5 production	11	2	0.33	0.0410
Mammalia	Group 1	GO:0097011	cellular response to granulocyte macrophage colony-stimulating factor stimulus	11	2	0.33	0.0410
Mammalia	Group 1	GO:0097012	response to granulocyte macrophage colony-stimulating factor	11	2	0.33	0.0410
Mammalia	Group 1	GO:0032306	regulation of prostaglandin secretion	12	2	0.36	0.0482
Mammalia	Group 1	GO:0032308	positive regulation of dendritic cell cytokine production	12	2	0.36	0.0482
Mammalia	Group 1	GO:0042501	serine phosphorylation of STAT protein	12	2	0.36	0.0482
Mammalia	Group 1	GO:0032736	positive regulation of interleukin-13 production	12	2	0.36	0.0482
Mammalia	Group 1	GO:1901857	positive regulation of cellular respiration	12	2	0.36	0.0482
Mammalia	Group 1	GO:1900226	macrophage proliferation	12	2	0.36	0.0482
Mammalia	Group 1	GO:0097170	negative regulation of NLRP3 inflammasome complex assembly	12	2	0.36	0.0482
Mammalia	Group 1	GO:0032306	positive regulation of nitric-oxide synthase biosynthetic process	12	2	0.36	0.0482
Mammalia	Group 1	GO:0032308	regulation of prostaglandin secretion	12	2	0.36	0.0482
Mammalia	Group 1	GO:0042501	positive regulation of dendritic cell cytokine production	12	2	0.36	0.0482
Mammalia	Group 1	GO:0060330	serine phosphorylation of STAT protein	12	2	0.36	0.0482
Mammalia	Group 1	GO:0060334	regulation of type II interferon-mediated signaling pathway	12	2	0.36	0.0482
Mammalia	Group 1	GO:0051770	peptidyl-lysine N6-acyltransferase	12	2	0.36	0.0482
Mammalia	Group 1	GO:0002306	hepatocyte immune response	12	2	0.36	0.0482
Mammalia	Group 1	GO:0032308	epicardium morphogenesis	12	2	0.36	0.0482
Mammalia	Group 1	GO:0042501	B cell antigen processing and presentation	12	2	0.36	0.0482
Mammalia	Group 1	GO:0060330	plasmacytoid dendritic cell antigen processing and presentation	12	2	0.36	0.0482
Mammalia	Group 1	GO:0060334	pancreatic stellate cell proliferation	12	2	0.36	0.0482
Mammalia	Group 1	GO:0018003	regulation of cellular defense response	12	2	0.36	0.0482
Mammalia	Group 1	GO:0002384	type I hypersensitivity	12	2	0.36	0.0482
Mammalia	Group 1	GO:1905223	helper T cell enhancement of adaptive immune response	12	2	0.36	0.0482
Mammalia	Group 1	GO:0024501	glomerular parietal epithelial cell differentiation	12	2	0.36	0.0482
Mammalia	Group 1	GO:0010186	Lewy body formation	12	2	0.36	0.0482
Mammalia	Group 1	GO:0010186	positive regulation of cellular defense response	12	2	0.36	0.0482
Mammalia	Group 1	GO:0098784	regulation of Lewy body formation	12	2	0.36	0.0482
Mammalia	Group 1	GO:0098786	biofilm matrix organization	12	2	0.36	0.0482
Mammalia	Group 1	GO:0024501	melanin matrix disassembly	12	2	0.36	0.0482
Mammalia	Group 1	GO:0016068	type I hypersensitivity	12	2	0.36	0.0482
Mammalia	Group 1	GO:0072343	melanin biosynthetic process	12	2	0.36	0.0482
Mammalia	Group 1	GO:0010185	positive regulation of IP-10 production	12	2	0.36	0.0482
Mammalia	Group 1	GO:0010186	cortoskeletal rearrangement involved in phagocytosis, engulfment	12	2	0.36	0.0482
Mammalia	Group 1	GO:0140122	regulation of Lewy body formation	12	2	0.36	0.0482
Mammalia	Group 1	GO:0140123	regulation of Lewy body formation	12	2	0.36	0.0482
Mammalia	Group 1	GO:0030186	melanin matrix disassembly	12	2	0.36	0.0482
Mammalia	Group 1	GO:0030187	type I hypersensitivity	12	2	0.36	0.0482
Mammalia	Group 1	GO:0072139	melanin biosynthetic process	12	2	0.36	0.0482
Mammalia	Group 1	GO:0140121	regulation of Lewy body formation	12	2	0.36	0.0482
Mammalia	Group 1	GO:0060097	cytoskeletal rearrangement involved in phagocytosis, engulfment	12	2	0.36	0.0482
Mammalia	Group 1	GO:0051041	regulation of calcium-i-dependent cell-cell adhesion	12	2	0.36	0.0482
Mammalia	Group 1	GO:2000422	regulation of eosinophil chemotaxis	12	2	0.36	0.0482
Mammalia	Group 1	GO:0032759	positive regulation of eosinophil chemotaxis	12	2	0.36	0.0482
Mammalia	Group 1	GO:0032723	positive regulation of connective tissue growth factor production	12	2	0.36	0.0482
Mammalia	Group 1	GO:1912051	cytoskeletal rearrangement involved in phagocytosis, engulfment	12	2	0.36	0.0482
Mammalia	Group 1	GO:0032745	positive regulation of calcium-i-independent cell-cell adhesion	12	2	0.36	0.0482
Mammalia	Group 1	GO:0051977	regulation of eosinophil chemotaxis	12	2	0.36	0.0482
Mammalia	Group 1	GO:1901247	lysophospholipid transport	12	2	0.36	0.0482
Mammalia	Group 1	GO:1901249	regulation of lung ciliated cell differentiation	12	2	0.36	0.0482
Mammalia	Group 1	GO:0070246	natural killer cell apoptosis process	12	2	0.36	0.0482

Tab. S10 continued from previous page

Dataset	Group of genes	GO.ID	Term	Annotated	Significant	Expected	Classic Fisher
Mammalia	Group 1	GO:0070247	regulation of natural killer cell apoptotic process	1	1	0.03	0.0299
Mammalia	Group 1	GO:1902310	positive regulation of peptidyl-serine dephosphorylation	1	1	0.03	0.0299
Mammalia	Group 1	GO:2000229	regulation of pancreatic stellate cell proliferation	1	1	0.03	0.0299
Mammalia	Group 1	GO:1904784	NLRP1 inflammasome complex assembly	1	1	0.03	0.0299
Mammalia	Group 1	GO:2000231	positive regulation of pancreatic stellate cell proliferation	1	1	0.03	0.0299
Mammalia	Group 1	GO:0150140	regulation of CD36 production	1	1	0.03	0.0299
Mammalia	Group 1	GO:0150142	positive regulation of CD86 production	1	1	0.03	0.0299
Mammalia	Group 1	GO:0150143	regulation of CD80 production	1	1	0.03	0.0299
Mammalia	Group 1	GO:0150145	positive regulation of CD80 production	1	1	0.03	0.0299
Mammalia	Group 1	GO:0032665	regulation of interleukin-21 production	1	1	0.03	0.0299
Mammalia	Group 1	GO:0032679	regulation of TRAIL production	1	1	0.03	0.0299
Mammalia	Group 1	GO:0032625	interleukin-21 production	1	1	0.03	0.0299
Mammalia	Group 1	GO:0032639	TRAIL production	1	1	0.03	0.0299
Mammalia	Group 1	GO:0090320	regulation of chylomicron remnant clearance	1	1	0.03	0.0299
Mammalia	Group 1	GO:0090321	positive regulation of chylomicron remnant clearance	1	1	0.03	0.0299
Mammalia	Group 1	GO:1900450	negative regulation of glutamate receptor signaling pathway	1	1	0.03	0.0299
Mammalia	Group 1	GO:0120042	negative regulation of macrophage proliferation	1	1	0.03	0.0299
Mammalia	Group 1	GO:1903916	regulation of endoplasmic reticulum stress-induced eIF2 alpha dephosphorylation	1	1	0.03	0.0299
Mammalia	Group 1	GO:1903917	positive regulation of endoplasmic reticulum stress-induced eIF2 alpha dephosphorylation	1	1	0.03	0.0299
Mammalia	Group 1	GO:0036496	regulation of translational initiation by eIF2 alpha dephosphorylation	1	1	0.03	0.0299
Mammalia	Group 1	GO:0036497	eIF2alpha dephosphorylation in response to endoplasmic reticulum stress	1	1	0.03	0.0299
Mammalia	Group 1	GO:1901625	cellular response to egestrol	1	1	0.03	0.0299
Mammalia	Group 1	GO:0160006	Fc receptor-mediated immune complex endocytosis	1	1	0.03	0.0299
Mammalia	Group 1	GO:0036153	triglyceride acyl-chain remodeling	1	1	0.03	0.0299
Mammalia	Group 1	GO:0036155	acylglycerol acyl-chain remodeling	1	1	0.03	0.0299
Mammalia	Group 1	GO:0046603	negative regulation of mitotic centrosome separation	1	1	0.03	0.0299
Mammalia	Group 1	GO:0050902	leukocyte adhesive activation	1	1	0.03	0.0299
Mammalia	Group 1	GO:0034156	negative regulation of toll-like receptor 7 signaling pathway	1	1	0.03	0.0299
Mammalia	Group 1	GO:1905151	negative regulation of voltage-gated sodium channel activity	1	1	0.03	0.0299

Supplementary Table S11 – Accession numbers and general statistics of RNA-sequencing libraries of *Mytilus galloprovincialis* developmental stages.

SRA acc. no.	Time point (hpf)	Developmental stage	Biological replicate	No. of trimmed reads	No. of uniquely mapped reads	% of uniquely mapping reads	No. of multi-mapped reads	% of multi-mapped reads	No. of unmapped reads	% of unmapped reads
SRR25387458	0	Unfertilized Egg	R2	31,614,834	22,677,680	71.73	4,525,340	14.31	4,411,814	13.96
SRR25387459	0	Unfertilized Egg	R1	31,229,282	22,930,909	73.43	3,631,800	11.63	4,666,573	14.94
SRR25387436	4	Zygote	R2	31,573,942	22,238,537	70.43	4,747,330	15.04	4,588,075	14.53
SRR25387447	4	Zygote	R1	31,215,045	22,657,854	72.59	3,980,706	12.75	4,576,485	14.66
SRR25387434	8	Embryo	R2	31,679,197	22,361,698	70.59	5,224,621	16.49	4,092,878	12.92
SRR25387435	8	Embryo	R1	31,137,192	22,231,464	71.40	4,558,377	14.63	4,347,351	13.96
SRR25387432	12	Gastrula 1	R2	30,684,472	21,477,819	70.00	5,351,506	17.44	3,855,147	12.57
SRR25387433	12	Gastrula 1	R1	31,006,745	22,158,048	71.46	4,808,473	15.51	4,040,224	13.03
SRR25387430	16	Gastrula 2	R2	31,129,558	21,172,217	68.01	5,943,113	19.10	4,014,228	12.89
SRR25387431	16	Gastrula 2	R1	31,108,790	22,366,338	71.90	4,797,011	15.42	3,945,441	12.68
SRR25387456	20	Trochophore 1	R2	31,313,006	21,923,726	70.01	5,234,475	16.71	4,154,805	13.27
SRR25387457	20	Trochophore 1	R1	31,153,366	22,818,864	73.25	4,258,642	13.67	4,075,860	13.08
SRR25387454	24	Trochophore 2	R2	31,237,678	21,993,262	70.41	5,095,540	16.32	4,148,876	13.29
SRR25387455	24	Trochophore 2	R1	30,568,998	22,102,616	72.30	4,376,490	14.31	4,089,892	13.38
SRR25387452	28	Trochophore 3	R2	31,251,736	22,218,911	71.10	4,737,140	15.16	4,295,685	13.74
SRR25387453	28	Trochophore 3	R1	30,151,120	21,916,806	72.69	4,155,633	13.78	4,075,661	13.53
SRR25387450	32	Advanced Trochophore 1	R2	31,450,472	22,501,615	71.55	4,808,549	15.29	4,140,308	13.16
SRR25387451	32	Advanced Trochophore 1	R1	30,344,616	22,168,913	73.06	4,088,100	13.47	4,087,603	13.47
SRR25387448	36	Advanced Trochophore 2	R2	31,281,625	22,138,801	70.77	5,099,606	16.30	4,043,238	12.92
SRR25387449	36	Advanced Trochophore 2	R1	30,446,461	22,282,130	73.18	4,164,126	13.68	4,000,205	13.13
SRR25387445	40	Advanced Trochophore 3	R2	31,286,823	22,253,466	71.13	4,808,187	15.37	4,225,170	13.50
SRR25387446	40	Advanced Trochophore 3	R1	30,427,358	22,021,178	72.37	4,139,754	13.61	4,266,426	14.02
SRR25387443	44	D-veliger 1	R2	30,469,689	21,902,055	71.88	4,591,610	15.06	3,976,024	13.05
SRR25387444	44	D-veliger 1	R1	30,486,877	22,346,398	73.30	4,187,438	13.74	3,953,041	12.96
SRR25387441	48	D-veliger 2	R2	30,209,384	21,152,507	70.02	5,122,804	16.95	3,934,073	13.03
SRR25387442	48	D-veliger 2	R1	30,492,559	22,356,764	73.32	4,132,649	13.55	4,003,146	13.12
SRR25387439	52	D-veliger 3	R2	30,581,805	22,287,023	72.88	4,273,235	13.98	4,021,547	13.15
SRR25387440	52	D-veliger 3	R1	30,540,062	22,508,657	73.70	3,938,157	12.90	4,093,248	13.40
SRR25387437	72	Late D-veliger	R2	31,144,538	22,754,899	73.06	4,155,579	13.35	4,234,060	13.60
SRR25387438	72	Late D-veliger	R1	31,504,212	23,356,674	74.14	3,830,231	12.16	4,317,307	13.70

Supplementary Table S12 – Set of HCR probes generated with the ‘insitu_probe_generator’ script from the Ozpolat Lab (Kuehn et al., 2022).

Pool name	Sequence
B1_Mgal_10B017427_vasa_33_Dla0	GAGGAGGGCAGAACCGaaAAATGCTGTAAAAACAAAACGTATA TTAAAATGTTGAAATGATTAAATAtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaAGAGTTCATCACGATTAAAGAAAGA ACTTAAAACATTACATCATAAGAtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaGCACATGATTAAAGGTAGCACTACT AAACAAAATAAGAACTGGATTGATTtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaGACGAACCTCTCCCAGACAGGCCG AAAAAATGCCCCCTTCCAATCataGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaTTCACTGCTGTTGTTCTTGTAAA CCAATAGTGAATTCAACTGAACAtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaGCAAGACATGGCTGGTTATAGTT ATTATGCCATTAGGTTAGTCGtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaACATCCACTGCTAAGTCGATG GTGACAACGTGTTGATTATATGtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaCATTGAAGAATTATCTGCTGAGAGT TGACACCCACTTTAGGTCGAAACAAAtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaGCATTCAATATGAATGGAGT ACAATTCCCTGCTTCCTCGCAGGTTAtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaAGAATGTGATTAATCCCAGGATTCC ATTAAATGGTTGAAATAACATGtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaTTGCTCTGGTTGGTCTTTCTGAT CTCCGGAACTCCCTGTGTTGCTGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaCTGAATCATCTGTGAGAACTGGT GTAACAAAGCCTTGGCAATTCCACCTaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaACATGTTAACAGAGGAATGTCTA ATAGATTGTGGAAGGTATAATTAtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaGCAAGGAAGTCAGCATTTCTTCT GGAAATTCAGACTGAGACAGTAAGtGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaATGCACATACTGTGCAACATCGGTA TTGACGTTCTGATCACGGGAACTtGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaTTGCACTGAACATAAGAGTTGCT CAACTCTGATTCTCTGGAAAtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaCTTCCCTTACCAATGACATCAATT TAAATATTCAGCTTCTAAACTGtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaCTACAGTCTTAACATTGCGCATG GTCCAAGAGAGGTTCCCATATAAtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaGAGAATGAACACTACCAGTCAGTCCAT ACTAAAGCCTGTGCTCCTGACTAtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaGACTTTTCACATTAAAGAAGG TGTAACAGTGTGGTCTATCATAGTtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaTGGTGTATGTTTAAATTTCA TTTTCATATTGTCAAAGTTTATGtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaGGGCCATATGCTCTCATAC AAAGACCTTCTGCTTAGGACAATCtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaCACTTCATTACATTGAAGCAGTT CATTTAGGACACTCTTGCCATGTGtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaCCTTGCCCTCACCACCTGATCCTC CCACTTCAACACATTGAAAGCAGTtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaCCTCCACCTTATTCCTCAAATC CCCCCTTGCTGCAAACTCACCATTaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaAGATATGCTCATGTCCTTACCCCT AGGCTTCTCCTCAAAAGCAGGGGTTaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaTCCTTGAGCACGGCCTACTTGGT TCTTCCAACACACGCCCCAAAtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaATTGTACTGTTAAGGTGTTGTT ATAACAAATTCTGTCAGGACTtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaACTTGTAATTTCATCAACACAGT AACTAACTAACATGCAAATTCTGtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaGTGAAAACAAAGAATGTTAAATT AAAAATATGTCACATGAACTTCAAtaGAAGAGTCTCCCTTACG GAGGAGGGCAGAACCGaaTTGAAACTTTAGAACATATTATA

Table S12 continued from previous page

Pool name	Sequence
B1_Mgal_10B017427_vasa_33_Dla0	ACAGTGTAAACATTTCTGCATGCCtaGAAGAGTCTCCCTTACG GAGGGAGGGCAGCAAACGGaaGTCAGTTATCTCTTTCAAAACAG CTTCTACTGCTGTATTGTGAATAATtaGAAGAGTCTCCCTTACG GAGGGAGGGCAGCAAACGGaaCAGACACTTCCCTGCCCTTCTC TGTGTAGTCCCCTCACCCCTTGTaaGAAGAGTCTCCCTTACG
B2_Mgal_10B093608_dmrt1l_32_Dla0	CCTCGTAATCCTCATCAaaaaaaaaggggggtttttttaaa AAAAAATAATGTCTCGATTTCAttaaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaAACAAAGCTCAATATGATATATC taaccaCTTCCAGATAACTTGAGaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaAAGTAACACAGctaaaaattaaaga CAGTTGAACCTTATTTCACCTTaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaattatacaaattataacatTGACTG ttataccTCCCTCATCTGTATCCAaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaAAAATCACCCCTGGTGGACAAA ACACATATTCAACCTTATTGGaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaATGAATTTGATAATCAAAGTACT CTATGATTATATCAAGTATCATTaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaatcatgtatattttttttttttttt TCAGACAAAGCATTGTATCtttggaaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaTAACCTTTGCAAAACATTCCCTT TGACCTGGACTTGCACCAAAATAaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaaacattttttttttttttttttttt ACGGGCACCAAAACATGTCAATTtcaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaATTTCAGACTCACATTCTTATGT attt CCTCGTAATCCTCATCAaaaaaaattttttttttttttttttttt TTAAGACACACTATCTGGCTTGGaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaACAGACAGcgaaactttttttttt TTCAATGATTCAAAAGTCTTAAaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaCATGCAGACAGACACTCTAACAA tggatgtAAAAATTACAGACTTCAaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaTCATTATTATACCTTCAAGGTGTGC TCAACCTCACCCCTCACAAATGCAAGaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaTTGAATTCTGGTtattttttttttt CTATAATTATTCACCTCTGAATTGaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaTTCTCTCTTATGACAATATCTT aatattt CCTCGTAATCCTCATCAaaactttttttttttttttttttttt CTCTGTATTAAACATTTCatcacaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaAAATTCTACAAATTtgccatgtat actt CCTCGTAATCCTCATCAaaATGAAATATGCACTAAatcaataa tttattt CCTCGTAATCCTCATCAaaTTGAATTCTGGTtattttttttttt CCTCGTAATCCTCATCAaaTTCTCAATATGTCCTTAAATCC tagttt CCTCGTAATCCTCATCAaaTCATTATTATGCACTAAACCGCC CCTCGTAATCCTCATCAaaATGCACTACAGAGATTAAACT GAAAGTCTAACAGATCAATTAAAaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaAGCCACGACTACAAAATTAAAGTT AAAAACGTTCTGCTCTTATCTGTaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaGTACAGTTGAATGTGGAAATATCATG CTTGTGGCTAGCTGGAGTCCATAaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaATTGTACGGCATACTTGGTGGCA TGGTCCCTGCATATATCCTGGGTTaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaTGTAAACAGTTGGTGGATAAAATCCA TGATGGTGGGTATGGTGTGGTAGaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaATTATTCTACACTCTGTGCAAC TGGCTACACAGAGGACTGACAATaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaCTCGCACTAGATTTTAAACACATG CTCTGTATGGACAATCTCTGTGaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaTCTGTCAAGATAATTGGATTGTTGT TTGACATCTACGGCAATAACAGCTaaATCATCCAGTAAACCGCC CCTCGTAATCCTCATCAaaacaattttttttttttttttttttt TTCTTCTCTATTGCTTgttctgtaaATCATCCAGTAAACCGCC

Table S12 continued from previous page

Pool name	Sequence
B2_Mgal_10B093608_dmrt1l_32_Dla0	CCTCGTAAATCCTCATCAaaCTGGCGTTAATACTAACAAACCTGA TTTTGAGCAGTAACCTATCTGTaaATCATCCAGTAAACCGCC CCTCGTAAATCCTCATCAaaATCTAACATGGACACCAGAGCAAG AAAAACATCCAAGCTGTGACCTTaaATCATCCAGTAAACCGCC CCTCGTAAATCCTCATCAaaTGCGGAATTGACGTTACTTC aaattacAACTTCTTCTCTCTaaATCATCCAGTAAACCGCC
B3_Mgal_10B014180_soxh_27_Dla0	GTCCCTGCCCTATATCTttCTTTCTTGAGACATCCACATT ATTAGTTATCGTCTCCCTTtCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttCTTGAGACATCCACATTAGTT TTATCGTCTCTCCCTTTCTTTtCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttCATCCACATTAGTTATCGTCTT TTCTCCCTTTCTCTGAtttCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttATTAGTTATCGTCTCTCCCTT CTTTCTCTCTGAGACATCCtCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttATATGCTCTACTAAAGACCTAGAGA TTTCTCTACAGAAGTAGGTAACATTtCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttTTCTCTGAGACATCCAC CAACATTAGTTATCGTCTCTCCtCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttTTCTCTGAGACATCCACATTAGT TAGTTATCGTCTCTCCCTTCCtCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttGAGACATCCACATTAGTTATCGT TGGCTCTCCCTTTCTCTGAtttCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttACATGCGTCCACGTGAGATTCTG CAATGGTATCATCTATGGACCCtCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttGTCTCTGGATGTTCTCTCAGC TGGCTGGGGAACGGGAACGGCAAttCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttGCAAGCTCTGAATCATCGGTAG GTCCCTGCCCTATATCTttGCAAGCTCTGAATCATCGGTAG GTCCCTGCCCTATATCTttCTGCTTCTTGAAG TGTCTGTTCTAAAGTGTGCTATGttCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttTCTTCTGCTTCTTGAAG TTTTCTGCTCTGCTGCTTCTTGAAG GTCCCTGCCCTATATCTttGGGTGCTTGTGATCTTCTGACAT GTCAGTATTGCTATAAGCAGACGTTtCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttGCATTACAGGCTTGCCTTCAATG TCATTGCTTTCTATCTGttCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttGACCACAGCGGCTCACCCCTCGGA TGCTGGAACAGCTTAAAGCTCTCAttCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttGTATTATTGTGACATTGAG GGTATTGAAAGGAAGTGGTTGGtCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttATCTGGAAAGCTTGAAGAAAG TGTAGTCAGGTATTCCGCTGTTGttCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttGAATGGCTGTCGTACATGGGATAT ATTCCatTCAGGAAATTCTGGCtCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttGGACAGACAATTGGTACCTTGAG ATTGGTGAACGATTGCTGTTGttCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttCATTACGGGTAATTGGTCTGTT GCATGTTACAGGTTGAATAGGCAATTCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttATCCCAGTACCTGGCATGAGAGTAG GAACCTGGACAGGTGTTCTACGtCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttGTGACATTCCCTGAGGTT tCCATCTGCTGACATAAGAAAATCTCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttGCAAGTATCGGCAATTGTT GCTTGAATGGTGTGCAACATTCAttCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttTTTAAGTGTGACATTCCCTGAGGTT CAGGTTGGGTGAATGACATACAACTCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttttttatgcaaAGTCCATTGCA TTCTGTTAAATTAGccgaatttCCACTCAACTTTAACCG GTCCCTGCCCTATATCTttTTATGAGTAAACATGTTGAAA CCCGGCGATAAAAGTTGCTGATTTtCCACTCAACTTTAACCG
B4_Mgal_10B094018_foxl2_28_Dla0	CCTCAACCTACCTCCAAcaagaaaaataacaataatataat TTGCATGGTAAGAAATTGcccttaatTCTCACCATATTGCTTC

Table S12 continued from previous page

Pool name	Sequence
B4_Mgal_10B094018_foxl2_28_Dla0	CCTCAACCTACCTCCAA CaaTAATTTCTTGTAGGCTCATAA ttaaaatatgaataaaattctCGATatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaCATT CAGGTATGATA AATATCTCA TTTTCTGGCACAGAATCGTGACACatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaTCCAGTAGGTATAATGCACCGCCTC ACTGTATGCCAATTATTACCTCTCatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaACTGGACCCATCGA ACTGTAGTCG TTGTTGCTGTAAGGAAAGGGAAA TattTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaTTCAATTGCTGATTGCTATTGCTTC AACTGGGAAATGTTGGTCAGGA TattTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaCCGGTGCGGAACACGGGACTGGTT TATAACTACAGGCCGCTAAACTGGGatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaTGGAGATGCAGAATTGGTGCAGA ACTGtataactgttatttgttgcattTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaTACGGTGGCGTGA AAAATACGGTT CACGGGAATATTGTAAGTACTGAGattTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaACTGTCCGTCACAA AATGGTT TTAAAGAAAATTGGTTATGAGAattTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaACGCATTCTCGTCGACGTGGTAG GAGAGATATCGATGCTGGTATGGTattTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaGGGTCTAAAGTCCAATAATTCCCTT CCTTCTCAAACATGTCCTCAAATGatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaTTACAAAACATTCACTCAGACTTAG GTTCTCCCCACCTTCTGGGACatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaCTTATTTCATAGTAGGGAAAT GTGACGAATACTATTGGCATCCTatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaACTGTTAATCTTTATCATGCAATT TTCATGATATATTGATAAATTCCACatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaAAGAATATGGTGGTTGACATCGGG TAATTGCCATAGCAATAAGAGCAACatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaTTGTTCTCTTCTGGTGTGTTT TTTTCACCGGTTTATTCACAGATTatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaCATTCTGGTGTACTATTCCCAT TCTTATCTCAAAGTAGTTTAttTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaTTAACATGTTAGGAATTGAGT TGAGCGATTCCACATTCCGCTAatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaCTGAAATTGGGATTCGAGAGGGT CACATTTCGAGTCATAAAGTTTatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaCCGTCAAACCTGTcatctaaatttg GTCTTAATTAAAATCCAAATCTGAatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaataatattagAATTACAAACACGC tattttcatgtttccatTTTgttaatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaCATCGATTAA Ttttttttttttttttt TCATAATGAAGTTTAtaTGTCTTTatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaTAACCTCTTCTTTGCTTCATT CTCAGGTTTAGTGA CTTCAAATCattTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaTAGTCACCAAATGTTTCTCATGG TTTCTTCTCTTTAATTCTGTCatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaCACTTCTCAGGATCTTAAACCTT GTCTTTCTTACTAAAGGTGACTTatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaTCAAACCTCATTGGCTGCATCTGAT TGCTGATACACCAGGACA ACTCTAatTCTCACCATATTGCTTC CCTCAACCTACCTCCAA CaaCGTTGTTGCTTTCTTGGTGG CGTTGTTGGTGA TTTGCTTGCATTatTCTCACCATATTGCTTC

Supplementary Table S13 – Set of Vasa/Ddx4 and Ddx3 sequences used in the phylogenetic analysis. For each sequence, the species, the gene accession number (Gene ID), the orthology group and the gene names for reference species are shown. Reference species are marked with an asterisk.

Species	Gene ID	Orthology group	Gene name
<i>Phreagena okutanii</i>	DN44424.c0.g1.i2.p1	Vasa/Ddx4	-
<i>Calyptogena (Archivesica) marissinica</i>	Ama38729	Vasa/Ddx4	-
<i>Calyptogena (Archivesica) marissinica</i>	Ama38727	Vasa/Ddx4	-
<i>Corbicula fluminea</i>	DN125059.c0.g1.i19.p1	Vasa/Ddx4	-
<i>Mactra chinensis</i>	DN48157.c2.g2.i2.p1	Vasa/Ddx4	-
<i>Ruditapes decussatus</i>	DN22317.c4.g2.i1.p1	Vasa/Ddx4	-
<i>Ruditapes philippinarum</i>	XP_060562671.1	Vasa/Ddx4	-
<i>Mercenaria mercenaria</i>	XP_053394752.1	Vasa/Ddx4	-
<i>Cyclina sinensis</i>	Hic.asm.11.970.2	Vasa/Ddx4	-
<i>Pisidium coreanum</i>	DN31082.c0.g1.i7.p1	Vasa/Ddx4	-
<i>Dreissena polymorpha</i>	XP_052283635.1	Vasa/Ddx4	-
<i>Mya arenaria</i>	XP_052775885.1	Vasa/Ddx4	-
<i>Sinonovacula constricta</i>	Chr8.1697	Vasa/Ddx4	-
<i>Solen grandis</i>	DN2375.c0.g1.i12.p1	Vasa/Ddx4	-
<i>Tridacna squamosa</i>	DN1975.c0.g1.i5.p1	Vasa/Ddx4	-
<i>Panopea generosa</i>	DN2386.c0.g1.i4.p1	Vasa/Ddx4	-
<i>Cristaria plicata</i>	DN71694.c8.g1.i1.p1	Vasa/Ddx4	-
<i>Hyriopsis bialata (Unio delphinus)</i>	M00000006703	Vasa/Ddx4	-
<i>Potamilus streckersoni</i>	KAK3601505.1	Vasa/Ddx4	-
<i>Margaritifera margaritifera</i>	MMAM00000026330	Vasa/Ddx4	-
<i>Perna viridis</i>	s01977g89	Vasa/Ddx4	-
<i>Perumytilus purpuratus</i>	DN96437.c0.g1.i2.p1	Vasa/Ddx4	-
<i>Mytilus galloprovincialis</i>	VDI03911.1	Vasa/Ddx4	-
<i>Mytilus galloprovincialis</i>	VDI03912.1	Vasa/Ddx4	-
<i>Modiolus modiolus</i>	DN179.c0.g1.i2.p1	Vasa/Ddx4	-
<i>Argopecten irradians concentricus</i>	Contig172.33	Vasa/Ddx4	-
<i>Pecten maximus</i>	XP_033738807.1	Vasa/Ddx4	-
<i>Patinopecten yessoensis</i>	XP_021370692.1	Vasa/Ddx4	-
<i>Ostrea edulis</i>	XP_056020028.1	Vasa/Ddx4	-
<i>Magallana (Crassostrea) gigas</i>	XP_011437246.2	Vasa/Ddx4	-
<i>Crassostrea virginica</i>	XP_022316564.1	Vasa/Ddx4	-
<i>Saccostrea glomerata</i>	Sgl001349	Vasa/Ddx4	-
<i>Pinctada margaritifera</i>	DN36893.c1.g3.i1.p1	Vasa/Ddx4	-
<i>Atrina pectinata</i>	DN813.c0.g1.i1.p1	Vasa/Ddx4	-
<i>Tegillarca granosa</i>	KAJ8305640.1	Vasa/Ddx4	-
<i>Anadara (Scapharca) broughtonii</i>	EVM0008860.1	Vasa/Ddx4	-
<i>Drosophila melanogaster*</i>	NP_001260458.1	Vasa/Ddx4	vasa
<i>Homo sapiens*</i>	NP_077726.1	Vasa/Ddx4	DDX4
<i>Mus musculus*</i>	NP_001139357.1	Vasa/Ddx4	Ddx4
<i>Danio rerio*</i>	NP_571132.1	Vasa/Ddx4	ddx4
<i>Caenorhabditis elegans*</i>	NP_491876.1	Vasa/Ddx4	glh-2
<i>Caenorhabditis elegans*</i>	NP_491963.1	Vasa/Ddx4	glh-1
<i>Caenorhabditis elegans*</i>	NP_491681.1	Vasa/Ddx4	glh-3

Table S13 continued from previous page

Species	Gene ID	Orthology group	Gene name
<i>Caenorhabditis elegans</i> *	NP_491207.3	Vasa/Ddx4	<i>glh-4</i>
<i>Magallana (Crassostrea) gigas</i>	XP_011446924.2	Ddx3	-
<i>Magallana (Crassostrea) gigas</i>	XP_034330003.1	Ddx3	-
<i>Crassostra virginica</i>	XP_022337075.1	Ddx3	-
<i>Ostrea edulis</i>	XP_056006193.1	Ddx3	-
<i>Saccostrea glomerata</i>	Sgl003232	Ddx3	-
<i>Atrina pectinata</i>	DN371.c0.g4.i2.p1	Ddx3	-
<i>Pinctada margaritifera</i>	DN39745.c0.g1.i3.p1	Ddx3	-
<i>Perumytilus purpuratus</i>	DN34627.c0.g1.i16.p1	Ddx3	-
<i>Perna viridis</i>	s00037g119	Ddx3	-
<i>Mytilus galloprovincialis</i>	VDI00208.1	Ddx3	-
<i>Modiolus modiolus</i>	DN49076.c0.g1.i10.p1	Ddx3	-
<i>Tegillarca granosa</i>	KAJ8310842.1	Ddx3	-
<i>Argopecten irradians concentricus</i>	Contig829.57.3	Ddx3	-
<i>Pecten maximus</i>	XP_033759680.1	Ddx3	-
<i>Patinopecten yessoensis</i>	XP_021341010.1	Ddx3	-
<i>Hyriopsis bialata (Unio delphinus)</i>	M00000003015	Ddx3	-
<i>Cristaria plicata</i>	DN67742.c10.g2.i2.p1	Ddx3	-
<i>Megalonaia nervosa</i>	g136014.t1	Ddx3	-
<i>Potamilus streckersoni</i>	KAK3605786.1	Ddx3	-
<i>Margaritifera margaritifera</i>	MMAM00000009046	Ddx3	-
<i>Ruditapes decussatus</i>	DN22481.c1.g4.i1.p1	Ddx3	-
<i>Ruditapes philippinarum</i>	XP_060588962.1	Ddx3	-
<i>Cyclina sinensis</i>	Hic.asm.6.43.1	Ddx3	-
<i>Calyptogena (Archivesica) marissinica</i>	Ama32770	Ddx3	-
<i>Phreagena okutanii</i>	DN58569.c0.g1.i6.p1	Ddx3	-
<i>Mactra chinensis</i>	DN49476.c41.g3.i1.p1	Ddx3	-
<i>Corbicula fluminea</i>	DN126815.c0.g1.i7.p1	Ddx3	-
<i>Dreissena polymorpha</i>	XP_052217061.1	Ddx3	-
<i>Mya arenaria</i>	XP_052782518.1	Ddx3	-
<i>Pisidium coreanum</i>	DN3392.c0.g2.i6.p1	Ddx3	-
<i>Sinonovacula constricta</i>	Chr9.1187	Ddx3	-
<i>Sinonovacula constricta</i>	Chr9.1230	Ddx3	-
<i>Solen grandis</i>	DN51.c4.g1.i12.p1	Ddx3	-
<i>Panopea generosa</i>	DN13909.c0.g3.i2.p1	Ddx3	-
<i>Tridacna squamosa</i>	DN33643.c0.g2.i2.p1	Ddx3	-
<i>Danio rerio</i> *	NP_001119895.1	Ddx3	<i>ddx3xa</i>
<i>Danio rerio</i> *	NP_571016.2	Ddx3	<i>ddx3xb</i>
<i>Mus musculus</i> *	NP_149068.1	Ddx3	<i>Pl10</i>
<i>Homo sapiens</i> *	NP_001180346.1	Ddx3	<i>DDX3X</i>
<i>Drosophila melanogaster</i> *	NP_001262379.1	Ddx3	<i>bel</i>
<i>Pisidium coreanum</i>	DN29220.c0.g1.i2.p1	Ddx3	-
<i>Caenorhabditis elegans</i> *	NP_001021793.1	Ddx3	<i>vbh-1</i>
<i>Caenorhabditis elegans</i> *	NP_001254859.1	Ddx3	<i>laf-1</i>
<i>Pisidium coreanum</i>	DN19615.c0.g1.i2.p1	Ddx3	-

Data availability

All the supplementary materials, as well as high-resolution figures and a parsable version of tables, are accessible online at the following GitHub repository:

https://github.com/filonico/phd_thesis_tex

Activity report

This is the report of the activities carried out during my 3-year PhD course (2021–2024).

Research activity

Here are the research activities not directly related to the main topic of the PhD thesis.

- Manual curation of long interspersed nuclear element (LINE) libraries of several bivalve species;
- comparative genomics analysis of Hox and ParaHox genes in branchiopod crustaceans;
- comparative genomics analysis of branchiopod crustaceans to investigate the molecular underpinnings of morphological stasis and genome size variations;
- molecular phylogenetics and Bayesian dating of branchiopod crustaceans;
- preparation of mRNA sequences of genes involved in body segmentation in *Triops cancriformis* (Pancrustacea, Branchiopoda), to be used to generate probes for mRNA *in-situ* HCR on larvae (in collaboration with the Patel Lab; Marine Biology Lab, Woods Hole, MA, USA);
- collection, fixation, and storing of juvenile stages of several stick insect (Insecta, Phasmida) species, to be used for mRNA *in-situ* HCR to investigate the temporal and spatial transcription of genes involved in wing morphogenesis (in collaboration with the Patel Lab; Marine Biology Lab, Woods Hole, MA, USA);
- preparation of a review on the evolutionary causes and consequences of trait loss reversals;
- preparation of mitotic chromosome plates in the red wood ant *Formica paralugubris* from cerebral ganglia of pre-pupae.

Visiting scholar

- Nuzhdin Lab (University of Southern California, Los Angeles, CA, UA; Aug 20th, 2023–Feb 20th, 2024), to accomplish the abroad period of my PhD;
- Juan Pasantes' lab (University of Vigo, Vigo, Spain; Jan 12th–22nd, 2023), for a specific training on chromosome mitotic plate preparation in bivalve species.

Teaching activity

- Practical class “CAFE: estimating gene family turnover across a phylogenetic tree” (Apr 23, 2024) for first-year students of the course “Molecular phylogenetics” pursuing a Master degree in “Bioinformatics” at the University of Bologna (Italy);
- Practical invertebrate zoology class (Sep, 2022–Jan, 2023) for first-year students pursuing a Bachelor degree in “Biological Sciences” at the University of Bologna (Italy).

Co-supervised thesis

- *Evaluation of different calibration methods on Branchiopoda (Crustacea) phylogeny.* Niccolò Righetti. Master degree in “Biodiversità ed evoluzione”, University of Bologna, Bologna (Italy). Supervisor: Andrea Luchetti. Co-supervisor: Filippo Nicolini. AA 2022–2023;
- *Filogenesi molecolare di alcune famiglie dell'ordine Phasmatodea con enfasi sulla famiglia Heteropterygidae (Bacilloidea).* Giacomo Orsini. Bachelor degree in “Scienze biologiche”, University of Bologna, Bologna (Italy). Supervisor: Andrea Luchetti. Co-supervisor: Simona Corneti, Filippo Nicolini. AA 2021–2022;
- *Filogenesi molecolare di specie appartenenti alle famiglie Heteropterygidae e Anisacanthidae (Phasmatodea, Bacilloidea).* Alessandro Siragusa Camacho. Bachelor degree in “Scienze biologiche”, University of Bologna, Bologna (Italy). Supervisor: Andrea Luchetti. Co-supervisor: Simona Corneti, Filippo Nicolini. AA 2021/2022;
- *Filogenesi molecolare di specie della famiglia Pseudophasmatidae.* Giovanni Amedeo Paselli. Bachelor degree in “Scienze biologiche”, University of Bologna, Bologna (Italy).

Supervisor: Barbara Mantovani. Co-supervisors: Simona Corneti, Filippo Nicolini. AA 2020–2021.

Courses and workshops

- *Establishing state-of-the-art mollusc genomics*. EMBO Workshop. Namur, Belgium. May 28th–31st, 2024;
- *Art (Science) Attack*. Physalia Courses. Online. May 20th–23rd, 2024;
- *Introduction to Python for biologists*. Physalia Courses. Online. Sep 2nd–5th–8th–2nd–8, 2023;
- *ITA *PHY phylogenetics workshop*. Trento, Italy. Jan 6th–9th, 2023;
- *Sex chromosome evolution*. Physalia Courses. Online. Jan 23rd–27th, 2023.

Awards and scholarships

- Travel grant to attend the “Evoluzione2024” congress in Naples (Italy). Stazione Zoologica Anton Dohrn. Sep 8th–11th, 2024;
- Travel grant to attend the EMBO workshop *Establishing state-of-the-art mollusc genomics* in Namur (Belgium). EMBO. May 28th–31st, 2024;
- Laura Bassi scholarship for editorial assistance to postgraduates and junior academics. Editing Press. Apr 13th, 2023.

Presentations at congresses

Oral presentations

- Nicolini F, Iannello M, Piccinini G, Ghiselli F, Luchetti A, Milani L. (2024). Advancing the study of bivalve sex determination in the light of comparative genomics. Establishing state-of-the-art mollusc genomics (EMBO workshop). Namur (Belgium). May 27-30, 2024;
- Nicolini F, Ghiselli F, Milani L, Luchetti A. (2023). Contrasting patterns of amino acid evolution and shared ancestry between putative sex-determining genes in bivalve molluscs.

EVOLMAR 2023. Online. Nov 14-17, 2023;

- Nicolini F, Ghiselli F, Milani L, Luchetti A. (2023). Sex-determination related genes in bivalves: novel acquisitions and high rates of sequence evolution. Evolution 2023 (Ernst Mayr Award symposium). Online. Jun 2-3, 2023.

Poster presentations

- Nicolini F, Iannello M, Piccinini G, ghiselli F, Nuzhdin S, Luchetti A, Milani L. (2024). How to detect sex-determining genes through molecular evolution: bivalves as a case study. Evoluzione 2024. Naples, Italy. Sep 8–11, 2024;
- Nicolini F, Ghiselli F, Milani L, Luchetti A. (2022). Clues of accelerated molecular evolution in gene families associated with sex determination in bivalves. SMBE 2023. Ferrara, Italy. Jul 24–27, 2023;
- Nicolini F, Ghiselli F, Milani L, Luchetti A. (2022). Clues of accelerated molecular evolution in gene families associated with sex determination in bivalves. SIBE/ISEB 2022. Ancona, Italy. Sep 4–7, 2022;
- Nicolini F, Martelossi J, Forni G, Mantovani B, Luchetti A. (2021) First insights and comparative genomics of Hox and ParaHox genes in tadpole shrimps. EuroEvoDevo 2022. Naples, Italy. May 31–Jun 3, 2022.

Invited talks

- *From comparative genomics to fluorescence imaging: a multi-disciplinary approach to study bivalve sex determination.* Auer Lab. University of Fribourg, Fribourg. Jul 26, 2024.

Outreach activity

- Editor and web writer for BioPills – the Italian community of life sciences (biopills.net/). Jul 2017–ongoing;
- Presenter at the European Researchers' Night 2024, University of Bologna, Bologna (Italy). Sep 27, 2024;

- Presenter at the BiGeA Day 2023, University of Bologna, Bologna (Italy). May 27, 2023;
- Opening Days, University of Bologna, Bologna (Italy). Nov 18, 2022.

Scientific publications

* equal contribution

- Righetti N*, Nicolini F*, Forni G, & Luchetti A. (2024). Towards a time-tree solution for Branchiopoda diversification: a jackknife assessment of fossil age priors. *Submitted for peer-review*
- Nicolini F, Ghiselli F, Luchetti A, & Milani L. (2023). Bivalves as emerging model systems to study the mechanisms and evolution of sex determination: a genomic point of view. *Genome Biology and Evolution*, 15(10), evad181. doi: [10.1093/gbe/evad181](https://doi.org/10.1093/gbe/evad181)
- Martelossi J, Nicolini F, Subacchi S, Pasquale D, Ghiselli F, & Luchetti A. (2023). Multiple and diversified transposon lineages contribute to early and recent bivalve genome evolution. *BMC Biology*, 21(1), 1–23. doi: [10.1186/s12915-023-01632-z](https://doi.org/10.1186/s12915-023-01632-z)
- Nicolini F, Martelossi J, Forni G, Savojardo C, Mantovani B, & Luchetti A. (2023). Comparative genomics of Hox and ParaHox genes among major lineages of Branchiopoda with emphasis on tadpole shrimps. *Frontiers in Ecology and Evolution*, 11, 23. doi: [10.3389/fevo.2023.1046960](https://doi.org/10.3389/fevo.2023.1046960)
- Forni G, Cussigh A, Brock PD, Jones BR, Nicolini F, Martelossi J, Luchetti A, & Mantovani B. (2023). Taxonomic revision of the Australian stick insect genus *Candovia* (Phasmida: Necrosciinae): insight from molecular systematics and species-delimitation approaches. *Zoological Journal of the Linnean Society*, 197(1), 189–210. doi: [10.1093/zoolinnean/zlac074](https://doi.org/10.1093/zoolinnean/zlac074)