Clustering of Sex-Biased Genes and Transposable Elements in the Genome of the Medaka Fish *Oryzias latipes*

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Accepted: 4 October 2021

Abstract

Although genes with similar expression patterns are sometimes found in the same genomic regions, almost nothing is known about the relative organization in genomes of genes and transposable elements (TEs), which might influence each other at the regulatory level. In this study, we used transcriptomic data from male and female gonads of the Japanese medaka *Oryzias latipes* to define sexually biased genes and TEs and analyze their relative genomic localization. We identified 20,588 genes expressed in the adult gonads of *O. latipes*. Around 39% of these genes are differentially expressed between male and female gonads. We further analyzed the expression of TEs using the program SQuIRE and showed that more TE copies are overexpressed in testis than in ovaries (36% vs. 10%, respectively). We then developed a method to detect genomic regions enriched in testis- or ovary-biased genes. This revealed that sex-biased genes and TEs are not randomly distributed in the genome and a part of them form clusters with the same expression bias. We also found a correlation of expression between TE copies and their closest genes, which increases with decreasing intervening distance. Such a genomic organization suggests either that TEs hijack the regulatory sequences of neighboring sexual genes, allowing their expression in germ line cells and consequently new insertions to be transmitted to the next generation, or that TEs are involved in the regulation of sexual genes, and might therefore through their mobility participate in the rewiring of sex regulatory networks.

Key words: Transposable elements, gonads, fish, genome organization, RNA-seq.

Significance

We analyzed RNA-seq data from medaka fish gonads, and estimated gene and transposable element (TE) expression. We report that sex-biased genes in *Oryzias latipes* gonads are arranged in clusters in the genome along with TEs. As germline expression of TEs is essential for their transmission to the next generation, we investigate the different mechanisms leading to this expression, either by being coexpressed with sexual genes, or by bringing their own regulatory sequences that will coregulate surrounding genes. Our findings are in favor of the second hypothesis. We also show that our method can discover particular TE families that are good candidates to control gene expression.

Introduction

With ~26,000 species (Nelson et al. 2016), teleost fish form the largest group of extant vertebrates. They present a high diversity of morphology, physiology, and behavior, this diversity also affecting their sexual development and function (Volff et al. 2007; Kobayashi et al. 2013). Many sexual modes exist in fish, from hermaphroditism, where one individual can have both sexes, either simultaneously or sequentially, to
gonochorism (individuals are either male or female). Sex determination, corresponding to the process by which the future sex is decided in gonochoristic species, is also diverse in teleosts. Several systems exist, ranging from environmental sex determination (where sex can be determined by water temperature for example) to genetic sex determination (GSD, where sex is controlled by a particular gene or a particular set of genes) with or without sex chromosomes (Bachtrog et al. 2014). In some fish species, both environmental and GSD occur and interact. In the medaka fish *O. latipes*, for instance, where sex is controlled by an XY chromosome system, high temperatures trigger female-to-male sex reversal (Adolfi et al. 2019). In mammals, the mechanism of GSD is highly conserved and ancient (~180–210 Ma; Waters et al. 2007), with almost all mammals using *Sry* as the male master sex-determining gene present on the Y chromosome. In contrast, in fish with GSD, different master sex determination genes can exist in different species, and in most species the master gene is still unknown (Kikuchi and Hamaguchi 2013). In *O. latipes* and in its related species *Oryzias curvinotus*, sex is controlled by the *dmrt1by* gene located on the Y chromosome. The *dmrt1by* gene appeared ~10 Ma in the common ancestor of *O. curvinotus* and *O. latipes* and was subsequently lost in *Oryzias luzonensis*, the sister species of *O. curvinotus* (Kikuchi and Hamaguchi 2013). In *O. luzonensis*, the master sex-determining gene is *Gsdf*, coding for the gonadal soma-derived factor. This gene is located on both X and Y chromosomes but in two different allelic forms, with the *Gsdf* allele triggering male differentiation. In *Oryzias dancena*, the master sex-determining gene is *Sox3Y* that evolved from *Sox3* as *Sry* did in mammals (Takehana et al. 2014; Herpin and Schartl 2015). The *Oryzias* group thus illustrates the high variability of master sex-determining genes that can control GSD in fish.

Sexual genes can be involved in sex determination (i.e., when sex is defined), but also in sexual differentiation (i.e., when the undifferentiated gonads become testes or ovaries) or sexual function. A way to detect such genes is to analyze gene expression between males and females, usually in the gonads, but not only (Grath and Parsch 2016), and to retrieve differentially expressed genes, called sex-biased genes, that is genes more expressed in males than in females or vice versa. An evolutionary conserved feature of sex-biased genes is their fast evolution, due to stronger positive selection, as observed in *Drosophila* (Assis et al. 2012), *Caenorhabditis elegans* (Cutter and Ward 2005), fish (Yang et al. 2016), and primates (Khaitovich et al. 2005), showing that this trend is conserved throughout evolution. Moreover, sex-biased genes often appear not randomly distributed on chromosomes. In mice and flies, female-biased genes are preferentially located on the X chromosome (Meisel et al. 2012). In *Drosophila* and mouse, testis-biased genes tend to co-localize in the genome and form clusters (Boutanaev et al. 2002; Li et al. 2005; Dorus et al. 2006).

Teleost fish genomes also harbor a large diversity of transposable element (TE) families compared with other vertebrates, particularly birds and mammals (Chalopin et al. 2015). TEs are sequences able to insert in the genome. They are often repeated and found in the genome of all eukaryotes analyzed to date. The high TE diversity observed in fish constitutes an important source of potential regulatory motifs for host genes. Indeed, if the majority of TE insertions are neutral or deleterious for the host, some can also be selected for adaptive functions. Several examples have been described of TEs with major roles in the rewiring of gene regulatory networks (Feschotte 2008; Lynch et al. 2011; Rebollo et al. 2012; Sorrells and Johnson 2015; Chuong et al. 2017), some of which are related to sex (Herpin et al. 2010; Ellison and Bachtrog 2013; Elison and Bachtrog 2015; Dechaud et al. 2019). Interestingly, the expression of *dmrt1by*, the master sex-determining gene of *O. latipes*, is partly controlled by a regulatory sequence carried by a TE called *Izanagi* (Herpin et al. 2010). This TE-derived enhancer allows the tightly regulated expression of *dmrt1by* limited to a short period of time before hatching, when sex determination occurs.

TEs are not randomly distributed in the genome. Patterns of TE insertions result from insertion preferences, selection, and genetic drift (Bourque et al. 2018). Because only TEs that insert in germline cells can be fixed in the genome to be transmitted to the next generation, these patterns could be particularly influenced by the structure of the chromatin in these cells, and thus related to gene expression. Conversely, TEs could bring regulatory elements with them and modify the expression of neighboring genes, participating in the evolution of regulatory networks in germ cells but also in gonads in general. As a first step to disentangle these potential functional regulatory links between sexual genes and TEs, we investigate here the localization of sex-biased TEs with respect to sex-biased genes. As sexual development is particularly diverse in *Oryzias* and TEs are highly diverse in teleost fish, we decided to use *O. latipes* as a model of study. We generated RNA-seq data from male and female adult gonads of *O. latipes*, and identified genes and TEs with sex-biased expression. Although gene expression in the genome is equivalently biased between males and females, TEs globally present a clear male-biased expression in gonads. We show that the closer the genes and TEs are, the more similar is their expression bias. Additionally, TEs located in sex-biased gene clusters tend to follow the cluster expression bias. Finally, we find that some male-biased TE families are enriched in male-biased gene clusters. These families constitute good candidates for TEs potentially involved in sexual gene regulation and its variability. Altogether, our study constitutes a first step toward a better understanding of the mutual regulatory influence between genes and TEs in the gonads.
Results

Identification of Gonadal Sex-Biased Genes

We first identified sex-biased genes by sequencing the transcriptome of three testis and three ovary replicates of *O. latipes*. These gonadal tissues are composed of both germ-line and somatic cells, as the two populations cannot be simply separated by dissection. However, as we are interested here in sexual function in general, and not only in germ cells, this is not limiting for our study. In addition, this did not prevent identification of sex-biased germ cell genes, for instance genes expressed in spermatogenesis (see below). Because gonads are specialized tissues that were not used to construct the reference genome annotation, this annotation (25,167 protein coding and noncoding genes) could lack some transcripts expressed in our data. To take this into account, we decided to apply the “new tuxedo” approach (Pertea et al. 2016), which is based on the reconstruction of a transcript annotation from the coordinates of the read alignments on the genome and the comparison of this new annotation with the reference from refseq (NCBI reference ASM223467v1; see Materials and Methods). We detected 45,444 expressed transcripts corresponding to 27,096 genes according to the pipeline. These gene models contain protein coding genes, noncoding genes, miss-assembled transcripts due to bioinformatic predictions, TEs, or any other type of expressed polyA RNA. We filtered these gene models to generate a set of coding genes and a set of noncoding genes. Among the 17,254 coding genes detected, 16,586 were already present in the refseq annotation (96.1%), and among the 3,334 noncoding genes detected, 1,845 were already present in the refseq annotation (55.3%).

On the whole, 40% of coding genes were found to be sex-biased, including 3,600 (20.9%) genes overexpressed in testis compared with ovary, and 3,293 (19.1%) genes overexpressed in ovary compared with testis. The remaining 10,361 genes were not differentially expressed (Supplementary fig. 1, Supplementary Material online). The coding transcriptome is thus equivalently biased between testis and ovary. For what follows, we define genes differentially expressed between male and female gonads as “sex-biased genes.” We assume that they could be involved in sexual differentiation, maintenance, or function.

With respect to the expression of particular genes, we compared the patterns we obtained with previous studies drawn in medaka only. Indeed, recent observations suggest that sexual gene expression might vary greatly between species, which would reflect the rapid evolution of this pathway (Herpin and Scharl 2015). Some studies analyzed the expression of few genes in the gonads of *O. latipes* by RT-qPCR (Nakamoto et al. 2006; Herpin et al. 2013; Horie et al. 2016; Kobayashi et al. 2017). The main genes that have been studied in detail are: *dmrt1*, the ancestral paralog of *dmrt1by*; *gsdf*, the gonadal soma-derived factor; and *foxl2*, a transcription factor involved in ovarian development. Both *dmrt1* and *gsdf* are involved in testis development. *dmrt1*, along with *gsdf*, was always found to be overexpressed in testis compared with ovary, which corresponds to our data (Supplementary fig. 1, Supplementary Material online). *foxl2* was found to be highly expressed in ovary in previous studies, and is coherently detected as ovary-biased using our RNAseq data (Supplementary fig. 1, Supplementary Material online). Additionally, reverse transcriptase–polymerase chain reaction (RT-PCR) experiments detected an expression of *amh* and *sox9b* in both gonads, and of aromatase in ovary only (Kurokawa et al. 2007). We observe similar patterns of expression for these genes using RNAseq data.

In the case of noncoding genes, 33.2% were found to be sex-biased, including 695 (20.8%) genes overexpressed in testis, and 412 (12.4%) genes overexpressed in ovary compared with testis. The noncoding transcriptome is thus more male-biased than the coding transcriptome, with a lower contribution of ovary-biased genes.

Identification of TE Copies and Families with Sex-Biased Gonadal Expression

To analyze TE expression relative to gene expression, we then characterized gonadal expression of TEs at the single copy level. Our annotation of TEs (supplementary data 1 and 2, Supplementary Material online) covers 34% of the medaka genome, which corresponds to a previously obtained coverage (Chalopin et al. 2015). SQuiRE (Yang et al. 2019) allows retrieval of the expression of each TE locus from RNAseq data. We identified 37,108 loci as expressed TE copies (corresponding to 3.7% of all TE loci). Among them, 13,325 (35.9%) were found to be testis-biased (adjusted \(P < 0.05, \log_2 \text{fold-change} (\log_2 \text{FC}) < -1\)), whereas 3,842 (10.4%) were ovary-biased (adjusted \(P < 0.05, \log_2 \text{FC} > 1\)). Therefore, whereas the same proportion of coding genes was found to be testis- or ovary-biased, TE expression appears clearly biased toward testis.

We further searched for TE families enriched in copies with testis- or ovary-biased expression. We found 22 families with global testis-biased expression (\(\chi^2\) comparing the proportion of testis-biased copies in a given family to the proportion of testis-biased copies genome-wide: \(P < 0.05/n\_families\) and >50% of testis-biased copies), and 19 families with global ovary-biased expression (\(\chi^2\) comparing the proportion of ovary-biased copies in a given family to the proportion of ovary-biased copies genome-wide: \(P < 0.05/n\_families\) and >50% of ovary-biased copies, supplementary data 3 and fig. 2, Supplementary Material online) among a total of 1,164 families presenting at least one expressed copy. Interestingly, the majority of the sex-biased families correspond to class I long terminal repeats (LTR) elements: ovary-biased families comprise 13/14 LTRs and 1/14 long interspersed nuclear elements (LINEs), and testis-biased families...
10/16 LTRs, 4/16 DNA transposons, 1/16 LINE, and 1/16 Rolling Circles. Only LTR elements are overrepresented among ovary-biased families compared with what would be expected from their genome-wide proportion ($\chi^2, P < 10^{-6}$). We generated a phylogeny of the LTR RT from family consensus sequences to confirm our annotation of the families, and to test if the expression pattern of the sequences is linked to their evolutionary relationships (supplementary fig. 3, Supplementary Material online). As most of the biased families are Gypsy elements, we more specifically generated a phylogeny of expressed Gypsy TE copies, irrespective of sex bias (fig. 1). We only used expressed TE copies for which we were able to detect an RT sequence. Plotting of expression patterns on the molecular phylogeny of TE copies showed that related TE copies often have similar sex-biased expression. To test if this could be explained by shared insertional preferences that would target similar TEs to similar expression environments, we analyzed sequences surrounding insertions of four subtrees (two mainly male-biased, one mainly female-biased, and one mainly non-biased; supplementary fig. 4, Supplementary Material online). The Guanine–Cytosine content was not significantly different between subtrees, and no insertion sequence specificity or preference could be detected (supplementary figs. 4 and 5, Supplementary Material online). These observations suggest that the expression bias may be explained by the sequence of the TE itself, that is, that TEs would harbor regulatory sequences shared between phylogenetically related copies that co-regulate their expression. Analyzing Gypsy copies of a male-biased subtree of the previous phylogeny, we observed that at least 10 of them (42%) present LTRs and complete open reading frames (ORFs), suggesting that they are only mildly corrupted and might correspond to recent and autonomously transcribed insertions (supplementary fig. 5A, Supplementary Material online). However, the phylogeny also presents some examples of closely related copies with different expression patterns. This

**Fig. 1.**—Phylogeny of expressed Gypsy TE copies of medaka using the amino acid sequence of the RT. Tip colors correspond to the expression bias (red: ovary; blue: testis; black: locus expressed but not biased). The location of biased Gypsy families is indicated. Note that copies with the same expression bias locally group together.
might be explained either by the loss of regulatory sequences by some copies, and/or by the influence of regulatory sequences from neighboring genes. We further drew a phylogeny including nonexpressed copies, but no particular clustering of expressed sequences compared with nonexpressed ones could be observed (supplementary fig. S5B, Supplementary Material online).

Sex-Biased Genes and TEs Form Clusters in the Genome

To characterize the genomic distribution of sex-biased genes and TEs, we first asked if they physically clustered in the medaka genome. To unravel possible clusters of coexpressed genes, we applied a method that previously demonstrated the existence of large clusters of coexpressed genes in the Drosophila genome (Boutanaev et al. 2002). Briefly, stretches of adjacent genes or TEs with the same expression bias are counted across the genome. A stretch stops as soon as a gene or TE is found with a different expression bias: features within a stretch all present the same bias. For example, in the *O. latipes* genome, the biggest stretch of consecutive male-biased genes is 10 genes long (and spans 36 kb), and a maximum of 8 consecutive female-biased genes is found (spanning 133 kb). For TEs, a maximum of 32 and 15 male- and female-biased copies are found, respectively. The observed number of stretches was compared with an expected number computed from random distributions of genes in the genome (fig. 2, brown bars). We observed that testis- and ovary-biased genes are both not randomly distributed in the genome because they do not follow the expected random distribution in terms of stretches of adjacent genes with similar expression bias. We found more clusters of at least three genes than expected if genes were randomly distributed (fig. 2). Moreover, for testis-biased genes, we observed stretches of

![Fig. 2.—Stretches of consecutive sex-biased genes and TEs along the genome of *O. latipes*. First row: stretches of sex-biased genes. Second row: stretches of sex-biased TEs, as identified by SQuIRE. The first column represents testis-biased sequences, and the second ovary-biased sequences. The observed number of stretches of a given length (in gene number) in the genome are shown in red and blue for females and males, respectively. One thousand random genomes were generated using the same proportion of male- and female-biased genes or TEs to estimate the stretch sizes if genes or TEs were randomly distributed. The median number of expected stretches is shown in brown. Error bars represent the fluctuation interval containing 95% of the values generated by the bootstraps.](image-url)
nine or ten genes, while such arrangement is never predicted among 1,000 random genomes. Results are given for coding and noncoding genes grouped together in the top panel of figure 2 (for coding and noncoding genes treated independently, see supplementary fig. 6, Supplementary Material online). We performed the same analysis with TE copies (fig. 2, bottom panel). We observed a nonrandom distribution for both testis and ovary sex-biased TEs as already found for genes. However, TE clusters contained markedly more elements than gene clusters: many comprised more than 8 TE copies, and up to 32 for male-biased TEs, while a maximum of 11 copies were predicted in a consecutive location in the random genomes. No particular enrichment of specific families could be observed in these clusters. As there are fewer ovary-biased TEs, the highest stretch sizes are lower than for male-biased TEs (15 and 32 consecutive biased TEs, respectively), but still higher than the maximum of 5 consecutive biased TEs expected at random under the null hypothesis. Globally, genes and TEs are thus not randomly distributed in the genome and tend to group into clusters with the same sex-biased expression. This is also true for coding and noncoding genes if analyzed separately (supplementary fig. 6, Supplementary Material online).

As mentioned before, this first method of identifying clusters of coexpressed genes is quite inflexible, a single gene with a different expression profile disrupting stretches and potentially hindering the identification of interesting regions regrouping genes with the same expression bias. This is why we applied a second and complementary search to identify regions showing a high density of sex-biased genes or TEs. We used the Gene clusters method previously developed by our team (Toubiana et al. 2020), which calculates the local mean log₂FC of the transcripts in a sliding window, and detects significantly biased regions (\( P_{adj} < 0.05 \)) using a bootstrap approach. We applied this to all gene transcripts, including coding and noncoding ones. We were able to identify 32 male-biased regions spread over 17 (out of 24) chromosomes (fig. 3A, supplementary table 1 and fig. 7, Supplementary Material online), and covering 3.94% of the genome. The method also uncovered 18 female-biased regions, spread over 13 chromosomes and

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**Fig. 3.**—Sex-biased gene expression clusters in *O. latipes* genome. (A) Genomic location of gene clusters built from coding and noncoding genes. (B) Percentage of testis- or ovary-biased genes in each type of clusters. (C) Chromosome 13 profile obtained with Gene clusters representing the expression bias of all genes across sliding windows along the genome. The mean log₂FC is represented for each window along the chromosome. The significantly male- and female-biased regions are shown in blue and red, respectively. The size of one window is shown, at scale, on the right of the plot.
covering 2.48% of the genome (supplementary table 1, Supplementary Material online, fig. 3A). Hence, about 6% of the genome of the medaka consists of sex-biased regions with respect to gene expression. In order to investigate their functions, we manually inspected all genes present in the sex-biased clusters in the medaka genome. We recovered genes known to be involved in male sexual functions in male-biased clusters, including dmrt1a (the autosomal paralog of the master sex-determining gene dmrt1bY), mom3, frizzled-4, ucq2, and lrguk, and genes with known female sexual functions in female-biased clusters, such as zonadhesin, bucky ball, bokb, and hsd17b1 (supplementary table 2, Supplementary Material online). We also tested the association of genes in sex-biased clusters with Gene Ontology (GO) terms. At the genome-wide scale, male-biased genes were significantly associated with "spermatogenesis," "cilium assembly and function," and "protein polyglutamylation" (Bobiniec et al. 1999, supplementary fig. 8, Supplementary Material online), and female-biased genes with "acrosome reaction," "oogenesis," and "binding of sperm to zona pellucida" (supplementary fig. 9, Supplementary Material online), indicating a global link in our analysis data between sex-biased expression and gonadal/germ cell function. In contrast, only the term "spermatogenesis" was significantly associated with genes in male-biased clusters (supplementary fig. 10, Supplementary Material online), and no significant GO term linked to sexual function and reproduction was found for genes in female-biased clusters (supplementary fig. 11, Supplementary Material online). In addition to the low power in the GO enrichment analysis due to a limited number of genes, this might also indicate that genes in sex-biased clusters, compared with sex-biased genes in general, are enriched in genes with so far uncharacterized sexual functions (possibly lineage-specific and evolving more rapidly), with functions that are less sex-specific than germ cell functions, or with functions with more recently recruited to the gonads.

We called “gene clusters” the regions presenting a significantly higher mean differential expression of genes. Eight hundred ten sex-biased genes (3.9% of all genes, and 10.1% of all sex-biased genes) are located in a cluster with the same expression bias. The size and composition of these regions calculated using coding genes only, noncoding genes only, or both are described in supplementary table 1, Supplementary Material online. Among the 828 genes located in a testis-biased region, 353 (42.6%) are testis-biased, whereas 78 (9.4%) are ovary-biased (fig. 3B). Ovary-biased regions contain 457 genes with 165 (36.1%) female-biased genes and 47 (10.3%) male-biased genes (fig. 3B). As an example, we observe on chromosome 13 of O. latipes two main ovary-biased regions of ∼1 Mb with a mean log2FC of −5 (fig. 3C). For the detailed gene cluster profiles for each chromosome, see supplementary data 4–7, Supplementary Material online.

In contrast to the situation observed in Drosophila (Boutanaev et al. 2002; Ellison and Bachtrog 2013), we did not observe any particular trend on chromosome 1, which is the X chromosome. The Y chromosome of O. latipes differs from the X by a 250-kb insertion that includes only one gene, the master sex-determining gene dmrt1bY. The rest of the X and Y chromosomes are very similar. The Y-specific region is not represented in the reference genome we used (thus containing an X but not a Y chromosome; https://www.ncbi.nlm.nih.gov/assembly/GCF_002234675.1, last accessed September 2021). If we look at the cluster profile obtained from TE expression, we observe on the X chromosome a testis-biased TE cluster surrounding the insertion breakpoint of the Y-specific region (supplementary data 7, chromosome 1, and fig. 12, Supplementary Material online). This cluster was thus present next to the region of insertion, and the insertion seems to have occurred in a region that was already male-biased.

**Neighboring Genes and TEs Share Correlated Expression Bias between Male and Female Gonads**

As we could determine the expression of TEs at the copy level, we next asked if neighboring genes and TEs shared similar expression patterns that would reflect a coregulation (originating from an enhancer of the gene or from the TE, or from the epigenetic state of the whole region). To do so, we assessed whether the sex bias in expression of adjacent genes (coding and noncoding: fig. 4, coding or noncoding: supplementary fig. 13, Supplementary Material online) and TEs is correlated. Only genes and TEs expressed in gonads were used to test the hypothesis. We calculated the Pearson correlation coefficient of log2FC of gene–TE pairs with different distance categories (border to border gene–TE distance 10 p–1 kb, 1–5 kb, 5–50 kb, and 50–500 kb). A given TE copy can be associated to all genes within the distance selected, and conversely a given gene can be associated to all TEs in the distance selected. We calculated the correlation using the copies grouped by TE family for families containing at least five TE copies, and making at least ten gene–TE pairs (i.e., ten TEs with one gene, or five TEs with two genes). The correlation coefficient of each family is represented depending on the distance considered to create the gene–TE pairs (fig. 4). We observed that the closer TEs and genes are located, the higher the correlation coefficient is (one-way analysis of variance, P<1e–10). The same result was obtained using only the coding or the noncoding genes (supplementary fig. 13, Supplementary Material online).

We complementarily asked if clusters of sex-biased genes could concentrate TEs with the same sex bias in gonadal expression. No clear common pattern of gene and TE distribution emerged from the inspection of gene clusters, indicating the diversity of their structures. In some clusters hotspots of TEs were observed (e.g., supplementary fig. 7A, Supplementary Material online), whereas in others, TEs showed no marked local enrichment (e.g., supplementary fig. 7B, Supplementary Material online).
On average, TE copies located in testis-biased regions are overexpressed in testis (mean log2FC = 2.96), and significantly more than TEs not located in sex-biased clusters (mean log2FC = 1.35, Student’s t-test P < 1e–16). TEs located in female-biased regions also present on average overexpression in testis (mean log2FC = 0.77), but at a lower level than TEs not located in sex-biased clusters (mean log2FC = 1.35, Student’s t-test P = 4.2e–6; supplementary fig. 14, Supplementary Material online). As the number of copies compared between both groups is large, a low P value is not really informative, and it is important to check for the size of the effect (mean log2FC 0.77 vs. 2.95 vs. 1.35). TEs located in ovary-biased regions are still more testis-biased than ovary-biased, which reflects the previous observation that the general trend of the genome is a testis-biased expression of TEs. Of the genome, 3.94% and 2.48% are considered as male and female sex-biased gene clusters, respectively; 5.84% of expressed TEs and 4.11% of all TEs (expressed or not) are located in male-biased gene clusters, whereas 1.81% of expressed TEs and 2.40% of all TEs are located in female-biased gene clusters. Overall, the results indicate that globally TEs do not preferentially integrate into sex-biased clusters (this was also observed considering only recent insertions, to take into account the possibility that clusters change their position during time; data not shown). A slight enrichment might be observed for expressed TEs in male-biased clusters, which might reflect preferential insertion (maybe due to open chromatin in testis) and/or positive selection of insertions in these regions, but this minor effect requires further investigation to assess its significance. To double-check for the presence of sex-biased TEs in sex-biased regions, we investigated if sex-biased TE copies were more likely to be inserted in regions with a similar sex-bias gene expression than somewhere else in the genome. For that, we tested the possible relationship between the localization of the copy and its expression (supplementary fig. 15, Supplementary Material online). Most of the 37,038 TE copies analyzed are located in unbiased regions and present unbiased expression (supplementary fig. 15, Supplementary Material online, 18,884 copies). We computed the expected copy numbers if there were no association between both localization and expression (supplementary fig. 15, Supplementary Material online, gray values), and tested the difference between observed and expected counts using a χ² test of independence (P < 1e–125). Again, as the number of copies is very high, it is important to consider the size of the effect (supplementary fig. 15, Supplementary Material online, ratio values) and not only the P value. We observed an enrichment of 1.72 (1,293/773) for testis-biased TE copies present in testis-biased regions, showing that there are more testis-biased TEs in testis-biased regions than expected at random. For ovary-biased copies within ovary-biased regions the enrichment is 1.58, which means that there are more ovary-biased TEs located in ovary-biased gene clusters than expected at random. On the contrary, testis-biased regions should contain approximately 217 ovary-biased copies, but contain only 114, making these regions depleted in ovary-biased copies (fold change ~0.53). Conversely, only 163 testis-biased copies are found in ovary-biased gene clusters (vs. 215 expected). Thus, these regions are depleted for testis-biased TEs (fold change ~0.76).

Finally, we tested if TE families harboring a large proportion of sex-biased copies are more likely to be located in sex-biased regions. Independently of the expression bias of their copies, and taking into account both expressed and nonexpressed copies, 19 TE families were enriched in testis-biased regions,
whereas 9 families appeared enriched in ovary-biased regions (fig. 5A). In figure 5B, we show the same data but coloring TE families if the expression of their copies is significantly biased in one sex. Overlapping the two graphic representations of the data (fig. 5C), we observe that four of the families enriched in male-biased regions are often enriched in male-biased copies. In contrast, families enriched in female-biased regions are not necessarily made up of copies overexpressed in ovaries. (C) Intersection between A and B. The five TE families with both significant testis-biased expression and testis cluster localization are shown in blue.

Discussion

Global Gene Expression Analysis Reveals a Similar Proportion of Testis- and Ovary-Biased Coding Genes in the Medaka

In this study we investigated gene and TE expression in the gonads of the Japanese medaka *O. latipes*. Compared with other tissues, gonads are the most sex-biased organs in terms of gene expression (Böhne et al. 2014; Tsakogiannis et al. 2018). These organs thus constitute a good model to study sex-dependent gene expression regulations. As the dissection of medaka gonads does not allow physical separation of the germline from the soma, our data comprise both types of cells and should be interpreted as a gonadal and not as a germline expression analysis. Several studies previously analyzed RNA-seq data in teleost fish, mainly from gonads (Böhne et al. 2014; Liu et al. 2015; Robledo et al. 2015; Bar et al. 2016; Zeng et al. 2016; Wang et al. 2017; Tao et al. 2018; Tsakogiannis et al. 2018) or from brain (Böhne et al. 2014; Liu et al. 2015; Saaristo et al. 2017; Beal et al. 2018; Tsakogiannis et al. 2018; Wu et al. 2019; Shen et al. 2020). Here we found that 40% of protein-coding genes present a sex-biased expression in medaka gonads, with 21% being testis- and 19% ovary-biased. In cichlid fish, by analyzing four species, 66% of the coding genes were found differentially expressed, with slightly more testis-biased genes (Böhne et al. 2014). In the Japanese fugu, only 3.7% of the coding genes were found differentially expressed between gonads, but still with more testis-biased genes (Wang et al. 2017).
proportion of gonad-biased genes thus greatly varies between fish species and probably also depends on the conditions and parameters of analysis. Many technical biases could explain these variations, because RNA-seq data analysis relies on several parameters including the type of pipeline used, the availability of a reference genome of good quality, and the thresholds applied to decide if a gene expression is sex-biased. The observed variations could nevertheless also result from true biological variability, notably because of differences in gonad maturity between species: medaka is a constitutive spawner, whereas fugu and cichlids have breeding periods, and the bluehead wrass is a consecutive hermaphrodite. A common trend that persists is the slight bias toward testis overexpressed genes. However, this effect is not that important in medaka, and we can consider that genes are overexpressed in similar proportion between sexes in this species.

A Higher Proportion of TEs and Noncoding RNA Genes Are Overexpressed in Testis Compared with Ovary

As TEs are known to be particularly involved in the regulation of vertebrate noncoding RNAs (Kapusta et al. 2013), we included noncoding RNA genes in our analyses, either merged with protein-coding genes, or as a distinct class of genes. It should be noted that, because our sets of transcripts were generated using poly-A purification and because a large part of noncoding RNAs is not polyadenylated (Cheng 2005; Kapranov et al. 2010; Sun et al. 2018), noncoding genes are likely to be underrepresented in our data set. Although RNA genes were globally found to be slightly less differentially expressed than coding genes (33.2% vs. 40%, respectively), they appeared more biased toward testis (63% of differentially expressed noncoding RNA genes are testis-biased, vs. 52% of differentially expressed protein-coding genes). Some of them might correspond to IncRNAs important for spermatogenesis, as reported in tetrapods (Necsulea et al. 2014). PolyA-enriched RNA-seq data are also relevant to study TE expression (Lerat et al. 2017; Lanciano and Cristofari 2020), as both class 1 and class 2 TEs express polyadenylated RNAs (Deininger and Batzer 2002). Almost half of expressed TE insertions were found differentially expressed (46.3%), with a major bias toward testis compared with ovary (78% vs. 22% of biased TEs, respectively). It is expected that TEs are particularly expressed in the germine, because their spreading and fixation in the genome rely on their activity in these cells (Dechaud et al. 2019). The question may arise if the higher proportion of testis-versus ovary-biased TEs might be a consequence of the different proportions of germine and somatic cells in the male and female gonads. These proportions are not available so far from literature, and we can only roughly estimate from our own unpublished observations in medaka a ratio of soma versus germine cells of ca. 1:100 in testis and 100:1 in ovary. Male and female germ cells, however, do not contain the same amounts of RNA. Again, there are no data available concerning this point, but we can very roughly estimate the ratio of soma versus germ cell RNA amount to be of 1:100 in both types of gonads. This estimate is reinforced by the similar proportions of male-versus female-biased gene transcripts determined in gonads, suggesting that expression levels are not much biased by their intrinsic structure. The higher proportion of testis-biased TEs could rather suggest that in medaka the male gonad is more permissive than the female gonad in terms of TE expression. This idea is somehow in opposition to the observations concerning the control of TEs in fish by piRNAs, a class of non-coding RNAs molecules that induce TE mRNA degradation. Indeed, both in zebrafish (Houwing et al. 2007) and medaka (Kneitz et al. 2016), piRNAs are more expressed in testes than in ovaries. Consequently, we might expect lower levels of TE expression in testis compared with ovary. However, the steady-state TE mRNA levels we observe result from both transcription and repression. The higher TE expression measured in testis could therefore result from much higher initial transcription, somehow counterbalanced by piRNA-mediated repression. More precise investigations are needed to conclude on this point. In addition, some retrotransposon families were shown to escape methylation in testis in mammals (Molaro et al. 2011). Similar experiments are needed in fish to evaluate if differential methylation of TEs might be linked to their higher level of expression in testis compared with ovary.

Sex-Biased Genes and TEs Form Clusters in the Medaka Genome

We focused particularly on the relative localization of genes and TEs with respect to their sex-linked expression bias. Many studies already reported that sexual genes (Boutanaev et al. 2002; Li et al. 2005; Dorus et al. 2006) and genes in general (Lercher et al. 2002; Roy et al. 2002; Singer et al. 2005) are not randomly distributed on chromosomes. Regulatory constraints such as the presence of specific enhancers or local chromatin states that affect neighboring genes can, for instance, result in the conservation of syntenies, even between distant species (Fernier and Holland 2001; Kikuta et al. 2007; Pascual-Anaya et al. 2013). These functional constraints are now known to translate into topologically associated domains (Jordan Rowley and Corces 2018; de Wit 2019), which are regions preferentially interacting in the 3D organization of the genome. A way to define clusters is by counting successive genes presenting the expression pattern (or any other feature) of interest (Boutanaev et al. 2002; Lercher et al. 2002; Roy et al. 2002; Li et al. 2005; Singer et al. 2005; Dorus et al. 2006). Such a procedure is quite stringent, but can reveal candidate regions where large stretches of coregulated genes exist. Applying this approach, we showed that the O. latipes genome is enriched in stretches of more than three sex-biased genes, this observation being valid for both coding and non-coding genes. When considering TEs, the enrichment is even...
stronger, with stretches of more than 20 testis-biased ele-
ments. This suggests that some regions contain a higher den-
sity of sequences with sex-biased expression.

In a second and complementary attempt, we developed a
new method that evaluates the global expression bias of
genes along the chromosomes and identifies regions with a
mean expression bias significantly diverging from random
expectations. This method can be applied to any set of differ-
entially expressed genes, and not only to sex-biased genes,
the only requirements being the localization of the genes and
the associated log₂ FC of their transcripts. Applying this
sliding-window approach, we detected 32 and 18 clusters
enriched in genes with male- or female-biased expression,
respectively. These clusters contain about 10% of all sex-
bias genes. We can hypothesize that such organization is
linked to a common regulation of the clustered genes. The
notion of “synexpression groups” has been proposed in
eukaryotes, which corresponds to sets of coregulated genes
(Niehrs and Pollet 1999; Ramialison et al. 2012; Herpin et al.
2019). In the medaka, a pilot study including 560 genes
expressed during embryonic development showed that core-
gulated genes tend to share particular DNA motifs in their cis-
regulatory regions (Ramialison et al. 2012). These motifs allow
the genes to be tightly controlled in space and time during
development. About a third of the developmental syn-
expression groups presented pairs of genes distant from less
than ten genes, which is rarely observed by chance
(Ramialison et al. 2012). This revealed a slight tendency of
coregulated genes to group on the chromosomes. With the
finding of 50 sex-biased gene clusters encompassing 25 bi-
ased genes in the mean, our genome-wide analysis made a
big step in this observation. It also demonstrates that this
trend not only concerns developmental genes but also genes
functioning in mature organs. It would be interesting to com-
pare the domains we identified with data of Hi-C, for now
unavailable for medaka gonads (Nakamura et al. 2021).

A Cluster of Sex-Biased TEs Could Have Favored the Birth
of Sexual Chromosomes

In mammals, X and Y chromosomes stopped recombin-
ing 210 Ma (Waters et al. 2007), leading to an accumula-
tion of TEs and a loss of genes on the Y chromosome. In O. latipes,
the sex chromosomes are relatively young. X and Y chromo-
some are still recombining on their almost complete length
and their single main structural difference is a short Y
chromosome-specific region of 250 kb containing the master
sex-determining gene dmrt1by and its transcriptional regula-
tory region, a copy of the Izanagi transposon (Kondo 2006).
One theory predicts that an important early step in the evolu-
tion of the Y chromosome is the linkage of sexually antag-
onistic genes that are beneficial to males but not to females in
the Y-specific region (Charlesworth et al. 2005). This leads to
a loss of recombination between the X and Y chromosomes in
this region, and in consequence to the accumulation of TEs
that can no longer be purged by crossing-over with the ho-
logous region lacking the TEs (Charlesworth et al. 2005).
We found on the X chromosome a testis-biased cluster of TEs
surrounding the insertion breakpoint of the Y-specific region,
suggesting that the future Y chromosome of O. latipes
already accumulated male-biased TEs in this region before the
new master sex-determining gene was inserted. It is thus
tempting to speculate that this region enriched in sex-
bias TEs could have eased the recruitment and evolution
of the new male master-sex determining gene dmrt1by of
O. latipes by providing a favorable male transcriptional
environment.

Disentangling the Possible Functional Links between TEs
and Sexual Genes

It has been already established that TEs are generally not ran-
domly distributed in the genome. Some retrotransposons,
for instance, are able to target regions (either precise nucleotide
sequences or larger particular chromatin environments)
where they can insert without generating deleterious muta-
tions, thus limiting their counterselection (Sultana et al. 2017).
Some Ty retrotransposons target the upstream region of pol-
III transcribed genes in Saccharomyces cerevisiae, which allows
both their expression and their location in a “safe” environ-
ment, with no risk of disrupting essential genes (Guo et al.
2015; Spaller et al. 2016; Cheung et al. 2018). Spreading and
fixation of TEs is intrinsically linked with their activity in
gonads, and more precisely in germ cells (Dechaud et al.
2019). They could be positively selected if they insert in
a region allowing their expression in this tissue. Postintegra-
tion selection is also an important force modulat-
ing the location of TEs. Strongly deleterious insertions, such as
insertions disrupting essential genes, are rapidly removed
from the genome because individuals carrying them are
strongly disadvantaged (Medstrand et al. 2002). In contrast,
insertions with a positive impact on host fitness will be
retained by selection. TEs are now known to be able to modu-
late gene expression and to rewire entire regulatory net-
works (Feschotte 2008; Chuong 2013). It has already been
proposed that TEs harboring TFBS allowing their germline ex-
pression could serve as a “taxi” for regulatory elements to
also control the expression of surrounding genes (Rebollo
et al. 2012; Sundaram et al. 2014; Dechaud et al. 2019).
They thus constitute good candidates to be involved in the
fast evolution of sexual pathways in medaka. If such a positive
selection concerns several insertions of the same TE family,
these insertions can concomitantly appear enriched in differ-
ent regions where they bring advantages. Finally, most inser-
tions have limited impact on host fitness; they generally
undergo genetic drift and are eliminated through random
mutations. The combination of all these mechanisms can ul-
timately lead to an enrichment of TEs in particular genomic
regions, either due to insertional preferences, to their low impact in these regions, or on the contrary to an acquired positive functional role, for instance in host gene regulation. To get more insights into these different evolutionary processes, we asked if the location of sex-biased TEs in *O. latipes* could be related to that of sex-biased genes.

We showed here that the sex-biased expression of neighboring genes and TEs is correlated: the closer TE copies are to genes, the higher is the correlation of their expression. Interestingly, such a coexpression has been also observed for TEs located near antiviral response genes in human and mice cells, suggesting that this phenomenon might not be restricted to sexual genes (Macchietto et al. 2020). This observation allows hypothesizing a coregulation of both types of sequences by shared cis-regulatory elements, provided either by enhancers of the sexual genes and/or by the sex-biased TEs themselves. Both hypotheses are not mutually exclusive and further analyses will be necessary to understand the origin of this correlation. In the first hypothesis, TEs inserting close to sexual genes could co-opt regulatory sequences favoring their expression and by this way their transposition in gonads, particularly in germ cells for transmission to the next generation. Our observation that expressed TEs are slightly enriched in male-biased clusters could indeed reflect a preference of insertion of TEs in these regions. The effect, however, was minor, and not observed for female-biased clusters.

In our data, TE families enriched in copies mainly expressed in one sex are LTRs, for both testis and ovary. Endogenous retroviruses (ERV)s were previously shown to frequently give birth to enhancers in fast-evolving tissues (Simonti et al. 2017), and more particularly their LTRs that contain many TFBS (Teng et al. 2014; Thompson et al. 2016). Furthermore, LTR elements are known to escape repression in tissues such as testis or placenta, which are hypomethylated and thus allow a higher global transcriptional activity (Molaro et al. 2011; Chuong 2013). However, further experiments would be needed to demonstrate such a recruitment of LTR elements for regulatory purpose. Even if we could not identify any sequence insertion preference between related Gypsy elements with the same sex-biased expression, from our data we cannot completely eliminate a purely neutral model where TEs preferentially insert in regions of open chromatin and subsequently follow the expression of neighboring genes. About three times more TEs were found overexpressed in testes than in ovaries. This observation agrees with the transcription of various genomic elements known to occur in testis (Soumillon et al. 2013). If the majority of these transcripts are probably nonfunctional, this high level of expression could also favor the birth of new genes or regulatory elements in this organ, in particular from TEs (Simonti et al. 2017).

Finally, we demonstrated in our work that sex-biased TE copies are enriched in gene clusters with the same sex-biased expression. We particularly identified an Unknown TE family strongly biased toward testis expression, and preferentially localized in regions considered as male-biased. Of note, expressed copies of this family are enriched in binding motifs for SOX8 and HOXD13, two factors involved in male sexual function. This TE family constitutes thus an interesting candidate to investigate for a potential role in the evolution of sex chromosomes and/or the regulation of sexual development of the medaka fish.

**Materials and Methods**

**Experimental Animals**

Laboratory-reared medaka (*Oryzias latipes*) of the Carbio strain were used. All fish were kept under standard photoperiod cycle of 14 h/10 h light/dark at 26 °C (±1 °C). Animals were kept and sampled in accordance with the applicable EU and national German legislation governing animal experimentation, in particular all experimental protocols were approved through an authorization (568/300–1,870/13) of the Veterinary Office of the District Government of Lower Franconia, Germany, in accordance with the German Animal Protection Law (TierSchG).

**Sampling and Sequencing**

The gonads of *O. latipes* were dissected. As testes are small, the testes of three males were pooled in one replicate. We generated three testis replicates (3 x 3 fish) and three ovary replicates (3 x 1 fish). Total RNA was isolated using RNeasy Mini kit (Qiagen) following the manufacturer’s instructions. RNA quality was assessed by measuring the RNA Integrity Number (RIN) using an Agilent 2100 Electrophoresis Bioanalyzer Instrument (Agilent Technologies 2100 Expert). RNA samples with RIN>8 were used for sequencing. RNA sequencing libraries were constructed following the standard TruSeq Illumina mRNA library preparation protocol (www.illumina.com, last accessed November 2019; Illumina Inc., BGI, Hong Kong), with a read length of 100 and sequencing depth for paired end of 62–72 million reads.

**Genome and TEs Annotation**

The genome of *Oryzias latipes* strain Hd-rR was sequenced and assembled with chromosome length scaffolds (https://www.ncbi.nlm.nih.gov/assembly/GCF_002234675.1, last accessed September 2021). This genome is also annotated (ftp://ftp.ncbi.nlm.nih.gov/genomes/efseq/vertebrate_other/Oryzias_latipes/latest_assembly_versions/GCF_002234675.1_ASM223467v1, last accessed September 2021).

TEs were annotated using the following protocol. First, a TE consensus database was generated using RepeatModeler (http://www.repeatmasker.org/RepeatModeler, last accessed May 2021; 1,596 consensi). To avoid false positives we
removed short consensi under 80 nt (1,400 resulting consensi), we self-blasted each consensus to find potential satellite sequences (1,398 resulting consensi), we removed non-TE genes by blasting the consensi against NCBI (13,82 resulting consensi), and we removed the redundant consensi by blasting the bank against itself (947 consensi). We crossed the bank with LTRharvest (Ellinghaus et al. 2008) output to reannotate some ERV TEs, and added Gypsy, ERV, and Copia elements (1,262 resulting consensi). We also added two Helitron sequences from HelitronScanner (Xiong et al. 2014) that were not found by other programs (1,264 consensi).

Some SINE sequences were reannotated using SINE_scan (Mao and Wang 2017). We finally ran MITE-hunter (Han and Wessler 2010), but after manual checking we did not find any good consensus to add. The bank was used to annotate the genome with RepeatMasker (http://repeatmasker.org/, last accessed May 2021). All TE copies annotated by the method used to generate figure 4 is the same as described in Boutanaev et al. (2002). The genes of TEs are represented by their expression bias, either testis, “T,” ovary, “O,” or nonbiased, “N.” They are represented as a sequence on each chromosome, like, as an example:

-NNNOO
NNNNTTTTNNNTONNNTTTONNNNNTTTTNNONNNNTN-

For each bias, the number of stretches is then counted. In this example, for testis-biased genes, we observe five stretches of one gene, one stretch of three genes, and one stretch of five genes. This will be represented by the blue bars in figure 4. Then the genome is shuffled 1,000 times. For each shuffling, the same counts are computed and we finally plot
the median value with the 95% fluctuation interval. This is not a confidence interval of the mean, but represents the distribution of 95% of the values obtained through the shuffles. The same approach was applied for ovary-biased genes. The script used to generate such a barplot is available from the gitlab repos (see below: stretch_of_genes.R).

Detection of Gene Clusters

The search for such stretches, however, is not fully suitable to detect genomic regions enriched in biased genes. If two consecutive genes are separated by a long gene desert, for example, they can still belong to a common stretch, in spite of their important intervening distance. Additionally, a single gene with a different expression can split a cluster, hindering its identification in spite of the global common expression bias of surrounding genes. It is thus interesting to study gene clustering on chromosomes in a more relaxed manner. We thus developed a new method to determine if genes are randomly distributed on chromosomes, or if they are grouped according to their expression, by designing a bootstrapping approach. The pipeline is available at https://gitlab.com/Corend/gene_clusters_pyth (last accessed October 2021), and was already used in a study on waterstriders (Toubiana et al. 2020).

Step 1: Design of the Expression Profile

First, a sliding window is designed on the genome. The size and step of the sliding window can be set by the user through the -step and -window parameters in the pipeline. The window and step sizes have to be chosen carefully according to the gene density of the studied genome, to ensure a sufficient statistical power. We tested different values and retained a step size of 35 kb and a window size of 700 kb. Hence, each window overlaps with the next 20 windows. Then, using the bed file of genes provided by the user (-b) and the expression of each gene (-e), the mean log₂FC of the genes is calculated in each window of each chromosome. The result can be used to display the fold change across each chromosome (fig. 3C).

Step 2: Bootstrap Analysis

Randomly distributed genes can form clusters just by chance. To test whether a cluster is observable by chance, we designed a bootstrap approach. The bootstrap number can be adjusted by the user (-boot). We used 10,000 bootstraps for our analysis. For each bootstrap, all the genes in the genome are randomly redistributed at each locus. Then the mean log₂FC is calculated again in each window. After the bootstraps, we have for each window the observed mean log₂FC, and 10,000 theoretical mean log₂FC. We calculate how many times the observed value is superior to the bootstrap and how many times it is inferior. If the observed value is always superior to the bootstrap values, then this region can be considered as “significantly testis-biased.” On the other hand, if the observed value is inferior to the bootstraps, it is considered as “ovary-biased.” The output of the pipeline corresponds to the mean log₂FC for each window and its associated bootstrap value.

Step 3: Statistical Analysis

This part is not included in the pipeline so that the user can choose its own bootstrap threshold. We converted the bootstrap value in a P value: [min(Bootstrap superior, Bootstrap inferior) × 2/10,000. We then converted the P values (1 P value per window) in q values using Benjamini–Hochberg false discovery rate correction from the R package qvalue (Storey et al. 2019). We use a q value threshold of 5%, meaning that among the windows considered as enriched in biased genes, 5% are false positives. Using this threshold, we colored the regions on the plot (fig. 3C).

Enrichment of TEs in Biased Regions

We investigated the localization of the TE copies of each family (fig. 5). We tested if each family was significantly enriched in the ovary- or testis-biased regions using Fisher’s exact test according to the method described in Karaku¨lah and Suner (2017). We tested the association between TE copy location and TE family expression using a Fisher exact test. As we tested all the TE families (1,164), we applied a Bonferroni correction to the test by taking 0.05/1164 as a P value threshold. The ternary plot was generated using the script Ternary_plot.R. The data used in the script are available in supplementary data 14, Supplementary Material online.

Supplementary Material

Supplementary data are available at Genome Biology and Evolution online.

Acknowledgments

This work was funded by a grant from the French National Research Agency (ANR-16-CE92-0019—EVOBOSTER) to JNV and by the Deutsche Forschungsgemeinschaft (Scha 408/13-1) to MS in the ANR/DFG cofunding program. We would like to thank the French Institute of Bioinformatics—IFB CNRS UMS 3601 (funded as part of Investissement d’avenir program managed by Agence Nationale pour la Recherche, contract ANR-11-INBS-0013)—for providing life science data and tools, storage and computing resources on the IFB national service infrastructure in bioinformatics.

Author Contributions

C.D., S.M., J.-N.V., and M.N. designed the study. A.M.-B. performed wet-lab analyses; C.D. and S.M. analyzed the data; C.D. wrote the first draft of the paper; S.M., M.N., J.-

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N.V., A.M.-B., and M.S. contributed to the final version of the paper. All authors read and approved the final manuscript.

**Data Availability**

The data underlying this article will be shared on reasonable request to the corresponding author. Scripts used to generate the figures of the paper are available on the following gitlab repository: https://gitlab.com/Corend/te_gene_expression (last accessed October 2021).

**Literature Cited**

Adolfi MC, et al. 2019. Increase of cortisol levels after temperature stress activates dmr11a causing female-to-male sex reversal and reduced germ cell number in medaka. Mol Reprod Dev. 86(10):1–13.

Assis R, Zhou Q, Bachtrog D. 2012. Sex-biased transcriptome evolution in Drosophila. Genome Biol Evol. 4(11):1189–1200.

Bachtrog D, et al.; Tree of Sex Consortium. 2014. Sex determination: why so many ways of doing it? PLoS Biol. 12(7):e1001899.

Bar I, Cummins S, Elizur A. 2016. Transcriptome analysis reveals differentially expressed genes associated with germ cell and gonad development in the Southern bluefin tuna (Thunnus maccoyii). BMC Genomics. 17:217.

Beal AP, Martin FD, Hale MC. 2014. Using RNA-seq to determine patterns of sex-bias in gene expression in the brain of the sex-role reversed Gulf Pipefish (Syngnathus scovelli). Mar Genomics. 37:120–127.

Bobinny C, Marcaillou C, Debec A. 1999. Microtubule polymerlyamylase in Drosophila melanogaster brain and testis. Eur J Cell Biol. 78(9):671–674.

Böhme A, Sengstag T, Salzburger W. 2014. Comparative transcriptomics in east African cichlids reveals sex- and species-specific expression and new candidates for sex differentiation in fishes. Genome Biol Evol. 6(9):2567–2585.

Bourque G, et al. 2018. Ten things you should know about transposable elements. Genome Biol. 19:199.

Boutanaev AM, Kalmykova AI, Shevelyov YY, Nurminsky DI. 2002. Large DNA. 10:42.

Chuong EB. 2013. Retroviruses facilitate the rapid evolution of the mammalian placenta. BMC Genomics. 14:506.

Chuong EB. 2011. Retroviruses facilitate the rapid evolution of the mammalian placenta. J Mol Biol. 318(2):236–340.

Chuong EB. 2017. Regulatory activities of transposable elements: from conflicts to benefits. Nat Rev Genet. 18(2):71–86.

Collins M, et al. 2015. Transcriptional rewiring of the sex determining dmr1 gene duplicate by transposable elements. PLoS Genet. 11(2):e1005014.

Czernecki P, et al. 2016. A novel evolutionary conserved mechanism of RNA stability regulates synexpression of primordial germ cell-specific genes prior to the sex-determination stage in medaka. PLoS Biol. 14(7):e2001875.

Durbec P, et al. 2015. Plasticity of gene-regulatory networks controlling sex determination: of masters, slaves, usual suspects, newcomers, and usurpators. EMBO Rep. 16(10):1260–1274.

Ellingham D, Kurtz S, Willhoeft U. 2008. LTRharvest, an efficient and flexible software for de novo detection of LTR retrotransposons. BMC Bioinformatics. 9:18.

Ellison CE, Bachtrog D. 2013. Dosage compensation via transposable element mediated rewiring of a regulatory network. Science 342(660):846–850.

Ellison CE, Bachtrog D. 2015. Non-allelic gene conversion enables rapid evolutionary change at multiple regulatory sites encoded by transposable elements. eLife 4:e05899.

Ferrier DEK, Holland PWH. 2001. Ancient origin of the Hox gene cluster. Nat Rev Genet. 2(1):33–38.

Feschotte C. 2008. Transposable elements and the evolution of regulatory networks. Nat Rev Genet. 9(5):397–405.

Fraze AC, et al. 2015. Ballgown bridges the gap between transcriptome assembly and expression analysis. Nat Biotechnol. 33(3):243–246.

Grath S, Parsch J. 2016. Sex-biased gene expression. Annu Rev Genet. 50:29–44.

Guo Y, Singh PK, Levin HL. 2015. A long terminal repeat retrotransposon of Schizosaccharomyces japonicus integrates upstream of RNA pol III transcribed genes. Mob DNA. 6:19.

Han Y, Wessler SR. 2010. MITE-Hunter: a program for discovering miniature repeat transposable elements from genomic sequences. Nucleic Acids Res. 38(22):e199.

Herpin A, et al. 2010. Transcriptional rewiring of the sex determining dmr1 gene duplicate by transposable elements. PLoS Genet. 6(2):e1000844.

Herpin A, et al. 2013. Divergent expression regulation of gonad development genes in medaka shows incomplete conservation of the downstream regulatory network of vertebrate sex determination. Mol Biol Evol. 30(10):2328–2346.

Herpin A, et al. 2019. A novel evolutionary conserved mechanism of RNA stability regulates synexpression of primordial germ cell-specific genes prior to the sex-determination stage in medaka. PLoS Biol. 17(4):e3000185.

Herpin A, Schartl M. 2015. Plasticity of gene-regulatory networks controlling sex determination: of masters, slaves, usual suspects, newcomers, and usurpators. EMBO Rep. 16(10):1260–1274.

Horie Y, et al. 2016. Androgen induces gonadal soma-derived factor, Gsdf, in XX gonads correlated to sex-reversal but not Dmrt1 directly, in the teleost fish, northern medaka (Oryzias sakaizumii). Mol Cell Endocrinol. 436:141–149.

Houwing S, et al. 2007. A role for Poxi and piRNAs in germ cell maintenance and transposon silencing in Zebrafish. Cell 129(1):69–82.

Jordan Rowley M, Corces VG. 2018. Organizational principles of 3D genome architecture. Nat Rev Genet. 19(12):789–800.

Kapranov P, et al. 2010. The majority of total nuclear-encoded non-ribosomal RNA in a human cell is “dark matter” un-annotated RNA. BMC Biol. 8:149.

Kapusta A, et al. 2013. Transposable elements are major contributors to the origin, diversification, and regulation of vertebrate long noncoding RNAs. PLoS Genet. 9(4):e1003470.

Karaulikh G, Suner A. 2017. PlanTeNrichment: a tool for enrichment analysis of transposable elements in plants. Genomics 109(5–6):336–340.

Katoh K, Misawa K, Kuma K-I, Miyata T. 2002. MAFFT: a novel method for rapid multiple sequence alignment based on fast Fourier transform. Nucleic Acids Res. 30(14):3059–3066.

Katoh K, Standley DM. 2013. MAFFT multiple sequence alignment software version 7: improvements in performance and usability. Mol Biol Evol. 30(4):772–780.

Khaitovich P, et al. 2005. Parallel patterns of evolution in the genomes and transcriptomes of humans and chimpanzees. Science 309(5742):1850–1854.
Kikuchi K, Harnaguchi S. 2013. Novel sex-determining genes in fish and sex chromosome evolution: novel sex-determining genes in fish. Dev Dyn. 242(4):339–353.

Kikuchi K, et al. 2007. Genomic regulatory blocks encompass multiple neighboring genes and maintain conserved synteny in vertebrates. Genome Res. 17(5):545–555.

Kondo M, et al. 2006. Genomic organization of the sex-determining and adjacent regions of the sex chromosomes of medaka. Genome Res. 16(7):815–826.

Kurokawa H, et al. 2007. Germ cells are essential for sexual dimorphism in the medaka gonad. Proc Natl Acad Sci USA. 104(43):16958–16963.

Lanciano S, Cristofari G. 2020. Measuring and interpreting transposable element expression. Nat Rev Genet. 21(12):721–736.

Lercher MJ, Urrutia AO, Hurst LD. 2002. Clustering of housekeeping genes provides a unified model of gene order in the human genome. Nat Genet. 31(2):180–183.

Li Q, Lee BT, Zhang L. 2005. Genome-scale analysis of positional clustering of mouse testis-specific genes. BMC Genomics. 6:7.

Liu H, et al. 2015. Large-scale transcriptome sequencing reveals novel sex differentiation in adult medaka, Oryzias latipes. Aquat Toxicol. 191:209–218.

Kobayashi T, et al. 2017. Estrogen alters gonadal soma-derived factor (Gsdfl/Fox2) expression levels in the testes associated with testis-ova differentiation in adult medaka, Oryzias latipes. Aquat Toxicol. 191:209–218.

Kobayashi Y, Nagahama Y, Nakamura M. 2013. Diversity and plasticity of sex determination and differentiation in fishes. Sex Dev. 7(1–3):115–125.

Kondo M, et al. 2006. Genomic organization of the sex-determining and adjacent regions of the sex chromosomes of medaka. Genome Res. 16(7):815–826.

Lynch VJ, Leclerc RD, May G, Wagner GP. 2011. Transposon-mediated rewiring of gene regulatory network contributed to the evolution of pregnancy in mammals. Nat Genet. 43(11):1154–1159.

Macchiott MG, Langlois RA, Shen SS. 2020. Virus-induced transposable element expression up-regulation in human and mouse host cells. Life Sci Alliance. 3(2):e201900536.

Mao H, Wang H. 2017. SINE_scan: an efficient tool to discover short interspersed nuclear elements (SINEs) in large-scale genomic datasets. Bioinformatics 33(5):743–745.

Medstrand P, van de Lagemaat LN, Mager DL. 2002. Retroelement distributions in the human genome: variations associated with age and proximity to genes. Genome Res. 12(10):1483–1495.

Meisel RP, Malone JH, Clark AG. 2012. Disentangling the relationship between sex-biased gene expression and X-linkage. Genome Res. 22(7):1255–1265.

Molara O, et al. 2011. Sperm methylation profiles reveal features of epigenetic inheritance and evolution in primates. Cell 146(6):1029–1041.

Nakamoto M, Matsuda M, Wang D-S, Nagahama Y, Shibata N. 2006. Molecular cloning and analysis of gonadal expression of Fox2 in the medaka, Oryzias latipes. Biochem Biophys Res Commun. 344(1):353–361.

Nakamura R, et al. 2021. CTCF looping is established during gastrulation in medaka embryos. Genome Res. 31(6):968–980.

Neculca A, et al. 2014. The evolution of IncRNA repertoires and expression patterns in tetrapods. Nature 505(7485):635–640.

Nelson JS, Grande TC, Wilson MVH. 2016. Fishes of the World. 5th ed. Hoboken (NJ): Wiley.

Niehrs C, Pollet N. 1999. Synexpression groups in eukaryotes. Nature 402(6761):483–487.

Pascual-Anaya J, D’Aniello S, Kuratani S, Garcia-Fernández J. 2013. Evolution of Hox gene clusters in deuterostomes. BMC Dev Biol. 13:26.

Pertea M, et al. 2015. StringTie enables improved reconstruction of a transcriptome from RNA-seq reads. Nat Biotechnol. 33(3):290–295.

Pertea M, Kim D, Pertea GM, Leek JT, Salzberg SL. 2016. Transcript-level expression analysis of RNA-seq experiments with HISAT, StringTie and Ballgown. Nat. Protoc. 11(9):1650–1667.

Price MN, Dekel PS, Arkin AP. 2010. FastTree 2-approximately maximum-likelihood trees for large alignments. PLoS One 5(3):e9490.

Ramialison M, et al. 2012. Cis-regulatory properties of medaka synexpression groups. Development 139(5):917–928.

Rebollo R, Romanish MT, Mager DL. 2012. Transposable elements: an abundant and natural source of regulatory sequences for host genes. Annu Rev Genet. 46(1):21–42.

Robledo D, et al. 2015. Gene expression analysis at the onset of sex differentiation in turbot (Scophthalmus maximus). BMC Genomics. 16(1):973.

Roy PJ, Stuart JM, Lund J, Kim SK. 2002. Chromosomal clustering of muscle-expressed genes in Caenorhabditis elegans. Nature 416(6891):975–979.

Saaristo M, et al. 2017. Characterisation of the transcriptome of male and female wild-type guppy brains with RNA-Seq and consequences of exposure to the pharmaceutical pollutant, 17 alpha-ethinyl estradiol. Aquat Toxicol. 186:28–39.

Shen X, et al. 2020. Transcriptomic analyses reveal novel genes with sexually dimorphic expression in Takifugu rubripes brain during gonadal sex differentiation. Genes Genomics. 42(4):425–439.

Simonti CN, Pavlicev M, Capra JA. 2017. Transposable element exaptation into regulatory regions is rare, influenced by evolutionary age, and subject to pleiotropic constraints. Mol Biol Evol. 34(11):2856–2869.

Singer GAC, Lloyd AT, Huminiecki LB, Wolfe KH. 2005. Clusters of co-expressed genes in mammalian genomes are conserved by natural selection. Mol Biol Evol. 22(3):767–775.

Storrells TR, Johnson AD. 2015. Making sense of transcription networks. Cell 161(4):714–723.

Sounillon M, et al. 2013. Cellular source and mechanisms of high transcriptome complexity in the mammalian testis. Cell Rep. 3(6):2179–2190.

Spalter T, Kling E, Glückner G, Hillmann F, Winkler T. 2016. Convergent evolution of lTRNA gene targeting preferences in compact genomes. Mob DNA. 7(1):17.

Storey JD, Bass AJ, Dabney A, Robinson D. 2019. qvalue: Q-value estimation for false discovery rate control. Available from: http://github.com/jstorey/qvalue. Accessed 11 December 2021.

Sultana T, Zamborlini A, Cristofari G, Lesage P. 2017. Integration site selection by retroviruses and transposable elements in eukaryotes. Nat Rev Genet. 18(5):292–308.

Sun Q, Hao Q, Prasanth KV. 2018. Nuclear long noncoding RNAs: key regulators of gene expression. Trends Genet. 34(2):142–157.

Sundaram V, et al. 2014. Widespread contribution of transposable elements to the innovation of gene regulatory networks. Genome Res. 24(12):1963–1976.

Takehara Y, et al. 2014. Co-option of Sox3 as the male-determining factor on the Y chromosome in the fish Oryzias danzica. Nat Commun. 5:4157.

Tao W, et al. 2018. Transcriptome display during tilapia sex determination and differentiation as revealed by RNA-Seq analysis. BMC Genomics. 19:363.
Teng L, He B, Gao P, Gao L, Tan K. 2014. Discover context-specific combinatorial transcription factor interactions by integrating diverse ChIP-Seq data sets. Nucleic Acids Res. 42(4):e24.

Thompson PJ, Macfarlan TS, Lorincz MC. 2016. Long terminal repeats: from parasitic elements to building blocks of the transcriptional regulatory repertoire. Mol Cell. 62(5):766–776.

Toubiana W, Armisén D, Dechaud C, Arbore R, Khila A. 2020. Impact of male trait exaggeration on sex-biased gene expression and genome architecture in a water strider. BMC Biol. 19:89.

Tsakogiannis A, et al. 2018. The transcriptomic signature of different sexes in two protogynous hermaphrodites: insights into the molecular network underlying sex phenotype in fish. Sci Rep. 8(1):3564.

Volff J-N, Nanda I, Schmid M, Schartl M. 2007. Governing sex determination in fish: regulatory putsches and ephemeral dictators. Sex Dev. 1(2):85–99.

Wang Z, et al. 2017. Comparative RNA-Seq analysis of differentially expressed genes in the testis and ovary of Takifugu rubripes. Comp Biochem Physiol Part D Genomics Proteomics. 22:50–57.

Waters PD, Wallis MC, Graves JAM. 2007. Mammalian sex—origin and evolution of the Y chromosome and SRY. Semin Cell Dev Biol. 18(3):389–400.

Wu J-J, et al. 2019. Comparative transcriptome analysis reveals differentially expressed genes and signaling pathways between male and female red-tail catfish (Mystus wyckioides). Mar Biotechnol (NY). 21(4):463–474.

Xiong W, He L, Lai J, Dooner HK, Du C. 2014. HelitronScanner uncovers a large overlooked cache of Helitron transposons in many plant genomes. Proc Natl Acad Sci USA. 111(28):10263–10268.

Yang L, Zhang Z, He S. 2016. Both male-biased and female-biased genes evolve faster in fish genomes. Genome Biol Evol. 8(11):3433–3445.

Yang WR, Ardeljan D, Pacyna CN, Payer LM, Burns KH. 2019. SQuiRE reveals locus-specific regulation of interspersed repeat expression. Nucleic Acids Res. 47(5):e27.

Zeng Q, et al. 2016. Transcriptome display during testicular differentiation of channel catfish (Ictalurus punctatus) as revealed by RNA-seq analysis. Biol Reprod. 95(1):19.

Zhou Q, et al. 2006. Helitron transposons on the sex chromosomes of the platyfish Xiphophorus maculatus and their evolution in animal genomes. Zebrafish 3(1):39–52.

Associate editor: Judith Mank