A Large Pseudoautosomal Region on the Sex Chromosomes of the Frog *Silurana tropicalis*

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Abstract

Sex chromosome divergence has been documented across phylogenetically diverse species, with amphibians typically having cytologically nondiverged (“homomorphic”) sex chromosomes. With an aim of further characterizing sex chromosome divergence of an amphibian, we used "RAD-tags" and Sanger sequencing to examine sex specificity and heterozygosity in the Western clawed frog *Silurana tropicalis* (also known as *Xenopus tropicalis*). Our findings based on approximately 20 million genotype calls and approximately 200 polymerase chain reaction-amplified regions across multiple male and female genomes failed to identify a substantially sized genomic region with genotypic hallmarks of sex chromosome divergence, including in regions known to be tightly linked to the sex-determining region. We also found that expression and molecular evolution of genes linked to the sex-determining region did not differ substantially from genes in other parts of the genome. This suggests that the pseudoautosomal region, where recombination occurs, comprises a large portion of the sex chromosomes of *S. tropicalis*. These results may in part explain why African clawed frogs have such a high incidence of polyploidization, shed light on why amphibians have a high rate of sex chromosome turnover, and raise questions about why homomorphic sex chromosomes are so prevalent in amphibians.

Key words: sex chromosome, pseudoautosomal region, recombination, sex determination, African clawed frogs, *Xenopus tropicalis*.

Introduction

Sex can be advantageous because it decouples beneficial from deleterious mutations via recombination, which increases the variance in fitness effects of linked mutations, and thus the efficiency with which natural selection operates. In species with genetic sex determination, developmental differences between the sexes are initiated by genetic differences between the sex chromosomes. In some lineages, the genes responsible for triggering sex determination vary, and the sex chromosomes (which carry the sex-determining region) are routinely reassigned from one to another ancestral pair of autosomal chromosomes (Fridolfsson et al. 1998; Ross et al. 2009; Evans et al. 2012; Pease and Hahn 2012). Ironically, suppression of recombination within a sex-specific region is often favored by natural selection, lest a sex-specific, sex-determining allele loses its sex specificity.

The origin of sex chromosomes could be initiated by sexual antagonism (van Doorn and Kirkpatrick 2007), and in many species, this is associated with cessation of recombination between a portion of the sex chromosomes that makes possible unisexual inheritance of a key genomic region that triggers sex determination. Cessation of recombination between the sex chromosomes can be achieved by reducing or eliminating homology (Charlesworth 1991), for example, through point mutations, inversion, deletion, or insertion of DNA. Strikingly, the extent of the nonrecombining region may increase overtime,
although this is not necessarily the case (Charlesworth et al. 2005; Bergero and Charlesworth 2009). The expansion of nonrecombining regions may be influenced by the nature of evolution in nonrecombining genomic regions, which is influenced by Muller’s ratchet, background selection, Hill–Robertson effects, and genetic hitchhiking of deleterious alleles with beneficial mutations (Charlesworth B and Charlesworth D 2000). Suppressed recombination between sex chromosomes thus has important implications for genome evolution, speciation, and adaptation.

Sex chromosome “degeneration” can be associated with sex chromosome divergence resulting from suppressed recombination and involves the loss of coding regions, the accumulation of repetitive regions, and structural changes such as insertions, deletions, and inversions on the sex-specific chromosome (the Y or W). Thus, degenerate sex chromosomes have differences that extend beyond the fundamental difference in the presence or absence of a sex-determining allele. However, evolutionarily young sex chromosomes start out being similar to each other because they originated from an essentially identical pair of autosomal chromosomes. In the medaka fish, for example, the male-specific region of the Y chromosome contains a newly evolved sex-determining locus (dmrt1bY) and is only approximately 258,000-bp long (Kondo et al. 2006). Similarly, the sex chromosomes of the tiger pufferfish appear to be distinguished by only one nonhomologous substitution (Kamiya et al. 2012). In theory, natural selection may drive the expansion of the sex-specific region of suppressed recombination (Charlesworth et al. 2005), and old sex chromosomes may evolve distinct suites of genes with unique, nonhomologous, and/or extensively diverged functions. In humans, for example, almost all the ancestral genes persist on the X chromosome but have been lost on the Y chromosome (Skaletsky et al. 2003). The limit of divergence is achieved if the sex-specific sex chromosome is lost altogether, as occurred in the Ryukyu spiny rat (Kuroiwa et al. 2010). In contrast, however, the sex chromosomes of ratite birds and boid snakes are old, but each pair is morphologically nondiverged (homomorphic), suggesting that the size of the region of suppressed recombination on the sex-specific sex chromosome does not necessarily expand over time (Matsubara et al. 2006; Tsuda et al. 2007).

Frog Sex Chromosomes

Most species of amphibians have homomorphic sex chromosomes (reviewed in Schmid et al. 2010). One possible explanation for the high incidence of homomorphic sex chromosomes in amphibians is that their sex chromosomes tend to be young because there has been frequent switching of the sex-determining locus during evolution. Consistent with this explanation is the inference that sex chromosomes have changed (“turned over”) approximately 32 times or more during evolution based on variation in male (XY) versus female (ZW) heterogamy (Evans et al. 2012; Schmid et al. 2010). Another explanation for widespread homomorphic sex chromosomes is that there is periodic recombination between the sex chromosomes over most of their length without changes in the sex-determining locus. In one group of hylid frogs, for instance, genomic regions that are tightly linked to the sex-determining locus are not substantially diverged between males and females, indicating that the sex chromosomes of these frogs recombine, at least occasionally, over most of their length (Stöck et al. 2011). This suggests that sex chromosomes of these frogs have large “pseudoautosomal” regions where inheritance of genetic information resembles autosomal genes and where recombination prevents divergence between the sex chromosomes. Thus, frequent turnover and recombination clearly both play a role in homomorph of amphibian sex chromosomes, but which phenomenon plays the dominant role remains an open question.

The only known amphibian sex-determining gene is called DM-W and was discovered in the African clawed frog *Xenopus laevis* (Yoshimoto et al. 2008). DM-W is female specific and originated after divergence from the sister genus *Silurana* but before diversification of most or all extant species of *Xenopus* (Bewick et al. 2011). Because *Silurana tropicalis* (also known as *X. tropicalis*) lacks DM-W, sex determination in this species must be triggered by another as yet unidentified genetic locus. A high-quality draft genome sequence is available for *S. tropicalis* that was generated from a female (Hellsten et al. 2010), but the sex-specific region of this genome has not been characterized. Using amplified fragment length polymorphisms (AFLPs), Olmstead et al. (2010) identified 22 AFLPs linked to the sex-determining locus in the “golden” strain of *S. tropicalis* (table 1) and proposed that females are the heterogametic sex in this strain. Four of these 22 AFLPs placed to the distal tip of chromosome/linkage group 7 in a linkage map developed by Wells et al. (2011), also represented by scaffold 7 in version 7.1 of the *S. tropicalis* genome sequence. However, the linkage map contains a large (15 cM) gap between the most distal two markers where the sex-determining region is likely to reside (Wells et al. 2011). Many of the other sex-linked AFLPs identified by Olmstead et al. (2010) map to other major or small chromosome/linkage groups (table 1); this is presumably because some scaffolds are chimera (e.g., a portion of scaffold 2 in version 7.1 is probably actually derived from *S. tropicalis* chromosome 7) and because linkage relationships between some small contigs (“orphan scaffolds”) and the larger scaffolds have not yet been established. Furthermore, additional experiments with other strains suggest that sex determination may occur through the action of multiple alleles at one locus or multiple tightly linked genes (Olmstead A, personal communication).

The goal of this study is to further characterize the sex chromosomes of *S. tropicalis* in terms of the size and level
of divergence of the sex-specific region, and to compare molecular evolution and expression of sex-linked and nonsex-linked genes. To this end, we used restriction-site associated DNA (RAD) tags (Baird et al. 2008), a reduced representation next-generation sequencing approach, to genotype millions of homologous nucleotide positions in male and female individuals including positions that are monomorphic in both sexes, polymorphic in one or both sexes, and positions in which a genotype inference (i.e., homozygous or heterozygous) was only possible in one sex due either to sex specificity of the genotyped position or differences in coverage of that position between the sexes. The RAD tag approach produces sequences of thousands of small regions that are adjacent to a rarely cutting restriction enzyme site. Because the sequenced portions of the genome are associated with restriction enzyme sites, many homologous sequences are obtained from multiple individuals. Missing data among individuals can arise in unusual cases where mutation generates polymorphism in the presence or absence of the restriction enzyme sites or because of variation among individuals in the depth of sequencing coverage for a particular region. Our analysis incorporated information on sex-linked regions from Olmstead et al. (2010), information from a laser-dissected chromosome arm 7p from a male individual (Seifertova et al., submitted), which is linked to the sex-linked region identified by Olmstead et al. (2010), and the most recent genome assembly (version 7.1, reference accession PRJNA12348). This study thus provides, for the first time, a comprehensive perspective on the extent of sex chromosome divergence in this species by evaluating the distribution of homozygous and heterozygous genotypes, molecular evolution, and gene expression of sex chromosomes in the context of the rest of the genome.

Materials and Methods

Four female and four male S. tropicalis individuals were obtained from Xenopus Express (Brooksville, FL). Sex was confirmed by dissection and species assignment achieved by comparing between 809 and 812 bp of mitochondrial DNA sequence from a portion of the 16S gene from each sample to homologous sequence data from all other known species of African clawed frog (Evans et al. 2011). We performed a phylogenetic analysis on these 8 sequences, 27 sequences from a larger scaffold that is not interrupted by unknown sequence or the entire scaffold was assumed to be sex linked (ALL).
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UWBM5969), and sequences from 6 individuals from the “golden” strain used by Olmstead et al. (2010) that were provided by Richard Harland. We used an X. laevis sequence from South Africa as an outgroup in this analysis, and the total alignment length was 817 bp. Model selection for phylogenetic analysis was accomplished using MrModeltest2 (Nylander 2004). Phylogenetic analysis was performed with MrBayes version 3.1.2 (Huelsenbeck and Ronquist 2001) using the best-fit model based on the Akaike Information Criterion, with two independent Markov chain Monte Carlo (MCMC) runs, each for 2,000,000 generations. Convergence of the MCMC runs on the posterior distribution was assessed by inspecting parameter trends and effective sample sizes using Tracer version 1.5 (Rambaut and Drummond 2007). Based on these analyses, a burn-in of 500,000 generations was discarded before constructing a consensus tree with MrBayes.

Genomic DNA was extracted from liver using QIAGEN DNeasy kit, purified using QIAGEN’s spin purification protocol, and RAD tag library preparation performed by Floragenex, Inc (Eugene, OR). For each individual, two libraries were generated—one used the restriction enzyme SbfI and another used NotI. The RAD tag libraries were multiplexed on three Illumina flow cells using individual barcodes, and Illumina sequencing was performed at the University of Oregon. These data have been deposited in GenBank (accession number SRP022004).

Illumina sequence reads were sorted by barcode with RADtools v1.2.4 using the “fuzzy_MID” option, which assigns reads with barcode errors to the nearest barcode (Baxter et al. 2011). Data from each individual were independently aligned to the S. tropicalis genome using bwa-0.6.2 (Li and Durbin 2009) and samtools.0.1.18 (Li et al. 2009). The “MarkDuplicates” function in picard (http://picard.sourceforge.net) was used to mark putative PCR-amplified duplicates, which were then excluded from the genotyping analysis with an aim of minimizing genotyping error. The Genome Analysis Toolkit (GATK) version 2.2-15 was then used to realign indels using the “RealignerTargetCreator” and “IndelRealigner” functions (McKenna et al. 2010; DePristo et al. 2011). The “FixMateInformation” function of picard was then used to adjust mate pair alignments.

Following “Best Practices” guidelines on the GATK website and forum (http://gatkforums.broadinstitute.org/) for analysis of genomes that lack known single-nucleotide polymorphisms (SNPs), the “UnifiedGenotyper,” “BaseRecalibrator,” and “PrintReads” functions of GATK were used to iteratively genotype, recalibrate base quality scores, and generate new input (bam) files, using the genotype files generated from “UnifiedGenotyper” as known polymorphic positions to be ignored for base recalibration in each iteration. Convergence was reached by the fifth iteration, in that variable positions recovered from this analysis were 99.8% identical to those from the fourth iteration. The “VariantFiltration” and “SelectVariants” functions of GATK were then used to identify and exclude genotyped positions that 1) were within 10 bp of an insertion/deletion, 2) had a Phred genotype quality score (Ewing and Green 1998) of less than 30, which means that we removed positions that had a probability of error of greater than 0.001, or 3) had more than one-tenth of the reads mapping equally well to another position and where there were at least four of these reads.

The S. tropicalis genome assembly 7.1 consists of 7,730 scaffolds aggregated from 55,234 contigs connected by “N”s within each scaffold. The total number of bases is 1,437,594,934, of which 5% (n = 71,599,926) are “N”s. This assembly includes 14 large “super scaffolds” that were assembled using meiotic map, synteny, and cytological data, corresponding to the 10 haploid chromosomes, with some chromosomes being represented by multiple scaffolds (3a and 3b; 5a and 5b; and 8a, 8b, and 8c). The rest of the scaffolds are “orphan scaffolds” whose chromosomal locations are not yet known. We divided the genomic regions into five mutually exclusive groups based on 1) the inferred level of recombination with the sex-determining region by Olmstead et al. (2010), 2) the linkage groups in the genome assembly 7.1 (table 1), and 3) the results of the Illumina sequencing of the dissected petite arm of chromosome 7 (Seifertova et al., submitted). The first of the five groups (“completely sex linked”) included contigs from assembly 7.1 that contain regions that had no recombination (0%) with the sex-determining region in Olmstead et al. (2010). This means that recombination between an AFLP polymorphism and the sex-determining region was not observed in any of 300 individuals assayed by Olmstead et al. (2010). The second group (“partially sex-linked”) included contigs from assembly 7.1 that contain regions that had a recombination rate of more than 0% and less than 3.0% in Olmstead et al. (2010). The third group (“chromosome 7p”) contained sections of scaffolds in assembly 7.1 that are located on chromosome 7p according to Seifertova et al. (submitted) and not in the “completely sex linked” or “partially linked” categories. The fourth group (“non-7p chromosomes”) contained the remaining sections on the chromosome-scale scaffolds in Assembly 7.1, including the portion of scaffold 7 that did not map to chromosome 7p. The fifth group (“other orphans”) contained orphan scaffolds in assembly 7.1 that 1) have not been linked to a chromosome, 2) have no evidence of sex linkage according to Olmstead et al. (2010), and 3) did not map to chromosome arm 7p according to Seifertova et al. (submitted). More specific information on the scaffold or scaffold portions in each of these groups is provided in table 1.

Genome-Wide Distribution of Genotypes in Female and Male S. tropicalis

For genomic regions in each of the six categories described earlier, we tabulated genotype patterns for three scenarios (fig. 1) in 500,000-bp windows across the S. tropicalis
Determining system. We call these patterns heterozygous in all males and no females (for an XY sex-determining system) or females and no males (for a ZW sex-determining system) or nation generates positions that are either heterozygous in all between the sex chromosomes due to suppressed recombination.

Some evolution in the following ways. First, divergence in males and females, for each window, we divided these counts by the total number of positions in each window for variation in coverage, we standardized the counts in each window by the sum of the number of positions in each window for which genotype data are available for 1) at least one female and at least one male, 2) at least one female but no males, and 3) at least one male but no females. Thus, by evaluating these three genotype scenarios in genomic windows across the S. tropicalis genome assembly, we attempted to identify genomic windows that either had significantly more heterozygous positions in one sex (Scenario 1), that had heterozygous positions only in one sex (Scenario 2), or that had homozygous positions in only one sex and no homologous genotypes in the other (Scenario 3). Higher values for each ratio are suggestive of genotype patterns characteristic of diverged sex chromosomes.

Another genotypic scenario for sex chromosomes is that a genomic region may be present only on the Z chromosome (with female heterogamy) or only on the X chromosome (with male heterogamy) (Scenarios 2a and 2b; fig. 1). No counterpart exists on the W chromosome (or Y chromosome) due to deletion, insertion, or divergence. To detect such a genomic region, we searched for regions with heterozygous positions present in one sex but not the other. For such positions, we required a genotype call in at least one individual of each sex but heterozygous calls to be present in only one sex. To account for variation in coverage in males and females, for each window we divided these counts by the total number of positions in each window for which genotype calls were made in at least one female and at least one male.

A third genotypic scenario for sex chromosomes is that a genomic region may be present only on the W chromosome or only on the Y chromosome (Scenarios 3a and 3b, fig 1). Thus, we searched for positions that had genotype calls only in females (or only in males) and that are all homozygous. To account for variation in coverage, we standardized the counts in each window by the sum of the number of positions in each window for which genotype data are available for 1) at least one female and at least one male, 2) at least one female but no males, and 3) at least one male but no females. Thus, by evaluating these three genotype scenarios in genomic windows across the S. tropicalis genome assembly, we attempted to identify genomic windows that either had significantly more heterozygous positions in one sex (Scenario 1), that had heterozygous positions only in one sex (Scenario 2), or that had homozygous positions in only one sex and no homologous genotypes in the other (Scenario 3). Higher values for each ratio are suggestive of genotype patterns characteristic of diverged sex chromosomes.

**Expression and Molecular Evolution**

As described in Chain et al. (2011), we estimated gene expression levels based on sequences across 26 expressed sequence tag (EST) libraries from the following tissues or developmental stages: egg, gastrula, neurula, embryo, tailbud, tadpole, metamorphosis, adipose tissue, bone, brain, head, heart, intestine, kidney, limb, liver, lung, ovary, oviduct, skeletal muscle, skin, spleen, stomach, tail, testis, and thymus. We summarized patterns of gene expression across EST libraries using the nonindependent “total,” “intensity,” and “evenness” statistics described in Chain et al. (2011). The “total” expression of a gene (I) is the proportion of times that a gene was sequenced in each EST library (I) summed across all libraries (I = ΣI). The “intensity” of expression (I) is the mean expression level from the perspective of a gene and is calculated following this equation: I = ΣI/ΣI. “Evenness” of expression (E) can be thought of as the “effective number” of tissues in which a gene is expressed and is calculated following this equation: E = T/I.
For a subset of the sex-linked and nonsex-linked genes, we also calculated the rate ratio of nonsynonymous to synonymous substitutions per site (dN/dS) along the S. tropicalis lineage using PAML version 4.5 (Yang 1997). This ratio was calculated using a maximum likelihood model that individually estimates dN/dS for each branch in a phylogeny, following Chain et al. (2011). Our phylogeny was estimated from sequences from S. tropicalis, X. laevis, and using sequences from another pipid frog (Pipa canvalhio or Hymenochirus curtipes) as an outgroup. To avoid undefined values, we added 0.02 to all dS values before calculating dN/dS, following Chain et al. (2011). We made this adjustment a priori by looking only at dS values, to make better use of the data. Because extreme values for dN and dS were occasionally estimated, we excluded from the analysis genes with an estimated dN or dS value above 2, and any genes whose available data comprised less than 100 synonymous positions.

We used a one-sided permutations to test whether the expression and molecular evolutionary statistics differed between genes that either (a) were or (b) were not on the same chromosome as the sex-determining locus. The permutations randomly divided the set of (a+b) values into two groups of size a and b and then calculated the difference between the averages of each group. We repeated this 1,000 times to generate a distribution for the null hypothesis that the values were drawn from the same underlying distribution, and then compared this with the observed differences, which is the test statistic of each test. A significant difference was inferred if the observed difference was greater than 95% of the differences from the permutations. Because these tests are one sided, the operands of the test statistic (i.e., the minuend and subtrahend of each difference) were defined according to specific expectations for sex chromosome degeneration discussed later.

Results

Mitochondrial DNA Variation within S. tropicalis, Including the “Golded” Strain

We analyzed phylogenetic relationships among approximately 810-bp region of mitochondrial DNA from the commercially obtained S. tropicalis individuals that we used for RAD tags and PCR screens, six individuals from the golden strain used by Olmstead et al. (2010), and several other wild-caught S. tropicalis individuals and individuals from other Silurana species. An identical mitochondrial DNA sequence was obtained from the six golden strain individuals, one of the samples we used for RAD tag sequencing (a female), 20 of the samples we used for PCR screens (9 females and 11 males), and one individual sampled from Nigeria. Mitochondrial sequences from five samples used in the RAD tag sequencing (2 females and 3 males) were identical to each other and differed from the golden strain sequence by one nucleotide substitution. Mitochondrial sequences from two other samples used in RAD tag sequencing (1 female and 1 male) and seven samples used in PCR screens (4 females and 3 males) were identical to each other and differed from the golden strain mitochondrial sequence by a different single-nucleotide substitution than the previously mentioned sequence present in five of the RAD tag samples. Mitochondrial sequences from another sample from Nigeria differed from the golden strain mitochondrial sequence by two nucleotide substitutions. Phylogenetic analysis of these and other sequences indicates that the commercially obtained S. tropicalis samples used in this study form a well-supported clade that includes two sequences from Nigeria and the six sequences from the golden strain of S. tropicalis (fig. 2). This clade is possibly common in individuals east of the Dahomey Gap, a savannah corridor that interrupts the West African rain forest (Salzmann and Hoelzmann 2005).

Reduced Representation Genome-Wide Genotyping from RAD Tags

We used a reduced representation genome sequencing approach called “RAD tags” to sequence many small but homologous portions of the S. tropicalis genome in four female and four male individuals. An average of 9,696,525 Illumina reads were mapped in each individual, with the average number of reads mapped per female or per male being 9,445,507 and 9,947,543 reads, respectively. After excluding positions in the reference sequence with no data, within an individual the average depth of coverage was 18.4 reads per position. Genotypes were called for a total of 19,624,824 positions, and 193,199 SNPs (0.98%) were detected. For each position at which at least one genotype was called, an average of 7.14 out of 8 individuals were genotyped.

If S. tropicalis has a large female-specific genomic region on the W chromosome, we expected a higher proportion of the Illumina reads from females to map to the genome assembly because this assembly was generated from a female individual. Contrary to this expectation, a slightly higher proportion of reads from males (average per male individual 89.2%, range: 87.8–90.8%) than from females (average per female individual 87.9%; range: 85.0–90.8%) mapped to this genome assembly, arguing against there being a large female-specific region in the S. tropicalis genome. Another indication of a large female-specific genomic region on the W chromosome would be a substantially higher number of positions genotyped in females than in males. Out of a total of 19,624,824 positions that were genotyped with high confidence in at least one individual, slightly more genotypes were recovered in females than in males: 912,738 (4.7%) positions were genotyped only in one or more females, and 462,792 (2.4%) positions genotyped only in one or more males. However, in the 10 largest scaffolds, the number of genotype calls in at least one female was consistently 1.1–3.3% higher than the number of genotype calls in at least one male, with
scaffold 7 having 2.5% more genotype calls in females than males. This suggests that the higher number of unique genotype calls in females is primarily a technical artifact related to differences in coverage among individuals in the RAD tag libraries. The RAD tag data did not provide high-quality genotypes from any positions on 5,721 scaffolds, which together comprise 36,133,437 bp (~2.1% of the genome).

Genome-Wide Genotype Patterns and Nucleotide Diversity Similar in Males and Females

We searched 500,000-bp windows for various genotypic patterns consistent with sex chromosome divergence expected under female or male heterogamy (fig. 1). In general, this effort failed to identify any regions with a pronounced genotypic signature of sex chromosome divergence expected by female heterogamy (table 2). One exception was a significant excess of windows with female-only homozygous genotypes (Scenario 3a) in orphan scaffolds, but we suspect this was an artifact related to the broader coverage in females. Most notably, portions of linkage groups 2 and 7 that were categorized as “partially sex-linked” and “completely sex-linked” to the sex-determining region based on Olmstead et al. (2010) did not exhibit a genotypic pattern consistent with degenerate sex chromosomes based on the RAD tag genotypes.

Considerable caution is needed in the interpretation of the average genotype frequencies in genomic windows for the “other orphans” category because in many cases the scaffold is smaller than the window size (500,000 bp), and the resulting truncated genomic windows are therefore expected to have an increased variance in the frequency of various genotypic patterns. Additionally, average genotype frequencies in these genomic windows could fail to detect small scaffolds that have genotypic patterns consistent with sex chromosome divergence. For example, “other orphans” had higher than expected values for Scenarios 2b and 3b, which are consistent with male heterogamy, and this is probably related to the small size of these scaffolds and consequent increase in the sex-specific genotypes in truncated windows for these scaffolds.

Additional insights are gained by examining nucleotide diversity in each sex within 500,000-bp windows. If a portion of the sex chromosomes is substantially diverged, we expected much higher average nucleotide diversity per site in one sex (females for female heterogamy) in genomic windows spanning this diverged region. However, average nucleotide diversity
Genotype Patterns Based on Sanger Sequencing

We amplified 65 portions (amplicons) of genomic regions identified by Olmstead et al. (2010) to be linked to the sex-determining region, including 18 and 46 amplicons from "completely sex-linked" and "partially sex-linked" regions, respectively (table 1, supplementary table S1 and fig. S3, Supplementary Material online). None had female-specific amplifications, allowing us to dismiss Scenario 3 (fig. 1) for all these regions. We sequenced 45 of these amplifications in multiple male and female individuals. SNPs or insertion/deletion polymorphisms were shared between males and females in at least one amplicon for essentially all scaffolds (no polymorphism was observed in amplicons from scaffold 144). This suggests that Scenario 1 is unlikely for these regions, with the caveat being that a heterozygous position could arise in both sexes in a region consistent with Scenario 1 through convergent evolution on the W and Z.

We also used PCR to examine an additional 173 regions that exhibited signs of sex linkage based on our analyses of the RAD tag data, including regions of chromosome 7p and elsewhere as detailed in supplementary figure S3 and table S1, Supplementary Material online. None had sex-specific amplifications, allowing us to dismiss Scenario 3 for all these regions. We sequenced 94 of these amplifications from male and female individuals. Thirty of these were not polymorphic in any of the individuals we sequenced. Forty-eight had polymorphisms in both sexes with none being shared across sexes. Ten had polymorphisms only in females and two had polymorphisms only in males.

Two amplifications were of particular interest. An amplification on scaffold 7 that spanned positions 10,128,301–10,129,920 was highly polymorphic in females but not males, although no polymorphism was fixed in females (out of seven females and three males sequenced; supplementary table S1, Supplementary Material online). This region failed to amplify in four females and three males. Another amplification, which targeted a region on scaffold 163 had 31 polymorphisms in three females but only one polymorphism in

diversity per site is essentially identical in males and females throughout these scaffolds, including chromosome arm 7p, which is linked to the sex-determining region (fig. 3). To explore the possibility that there could be variation within the RAD tag samples in sex chromosome divergence that corresponds with the three mitochondrial DNA haplotype groups detailed earlier, we explored nucleotide diversity in male and female individuals from each group. This analysis also did not identify a pronounced signature of sex chromosome divergence (supplementary figs. S1 and S2, Supplementary Material online).

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three males, but essentially all the female polymorphisms were present in only one individual.

**Gene Expression and Molecular Evolution**

Expression was detected in a total of 37,790 transcripts in at least one of the 26 EST libraries we surveyed (table 3). On the basis of studies of recently diverged neosex chromosomes in fruit flies (*Drosophila*) (reviewed in Bachtrog 2013), we expected expression of genes situated near the sex-determining locus to be expressed 1) at a lower total level, 2) higher intensity, 3) lower evenness, and to have 4) higher $dN/dS$ compared with genes in other parts of the genome (i.e., the “non 7p chromosomes”). For the most part, these expectations were not met for genes that were demonstrably very close to the sex-determining region, with the one exception that the intensity of “completely sex-linked” genes were individually significantly higher than the “non 7p chromosomes” ($P < 0.05$, table 3). Evenness of “other orphans” was also significantly lower than “non 7p chromosomes” as was total expression of “other orphans.” $dN/dS$ was significantly higher only in “chromosome 7p” compared with “non-7p chromosomes” but the magnitude of this difference was small. No expression data were recovered from genes on Scaffold 22, which is tightly linked to the sex-determining region (Olmstead et al. 2010), even though it was 1,156,260-bp long. We examined this scaffold using Xenbase (Bowes et al. 2009) and found that it contained a cluster of olfactory receptors, which (not surprisingly) were not highly expressed in any of the EST libraries we examined.

**Discussion**

To explore sex chromosome divergence in an amphibian, we used genotype calls from approximately 20 million positions, information about sex linkage, EST databases, and molecular evolutionary analyses to further characterize the sex chromosomes of the Western tropical frog *S. tropicalis*. Phylogenetic analysis of mitochondrial DNA sequences suggests our samples originated in Nigeria, which is also the source of the female individual from which the genome sequence was generated (Hellsten et al. 2010). Additionally, our analysis also...
suggests that the golden strain analyzed by Olmstead et al. (2010) is from Nigeria.

Known sequences in the \textit{S. tropicalis} genome sequence assembly version 7.1 comprise approximately 80.4\% of the approximately 1.7 Gbp genome, and scaffolds, including “N”s, comprise approximately 84.5\% of the genome. Thus, the RAD tag data could not be compared with 15–20\% of the genome because of gaps in the genome sequence. Because of variation in coverage, high confidence genotype calls were not made on scaffolds that together comprise an additional 2.1\% of the genome. Thus, in this study, we lack information from a nontrivial portion of this genome.

Mindful of these substantial gaps in genome sequence and the uncertainty in linkage relationships among many unassembled (orphan) scaffolds, we leveraged information from a targeted sequencing effort of chromosome arm 7p and also the linkage analysis by Olmstead et al. (2010) to guide our analysis. The dearth of genotypic patterns consistent with divergent sex chromosomes, and particularly patterns that are consistent with female heterogamy (table 2), and the similar level of pairwise nucleotide diversity in males and females throughout the petite arm of chromosome 7 (fig. 3) argues strongly against there being a large sex-specific region of the \textit{S. tropicalis} chromosomes. This inference is consistent with the findings of Uno et al. (2008) who detected no sex differences in C-banded heterochromatin in \textit{S. tropicalis}.

On the basis of studies of fruit flies (reviewed in Bachtrog 2013), we expected genes linked to the sex-determining locus to potentially exhibit lower total expression and higher specificity (i.e., higher intensity and lower evenness as defined in Materials and Methods section). We also expected molecular evolution of these genes to be consistent with relaxed purifying selection. However, on the basis of a small sample size, we only observed a significant increased expression intensity of “completely sex-linked” genes compared with the rest of the genome, with none of these expectations met in “partially sex-linked” genes (table 3). Some of these expectations were also met in orphan scaffolds, which have undetermined linkage relationships with respect to the sex-determining locus, and regions of chromosome arm 7p. It is not clear that these latter observations are related in any way to linkage to the sex-determining region.

Caveats exist in our interpretation of these data. First, non-recombining portions of the genome tend to accumulate repetitive sequences that can be difficult to sequence and map. For this reason, the sex-specific portion of the \textit{S. tropicalis} genome may be under-represented in the current genome assembly and/or our mapped illumina reads. Second, it is conceivable that there is polymorphism in the sex-determining mechanism (Olmstead A, personal communication). Polymorphism in genetic sex determination could occur at a single locus where multiple, differently functioned sex-determining alleles are segregating at a single locus, which have distinct and not necessarily transitive dominance relationships. Polymorphism in genetic sex determination could also occur at multiple loci distributed on the same or different chromosomes. Sex determination in zebrafish, for example, appears to be orchestrated by genes on different chromosomes (Anderson et al. 2012). Genotypic patterns expected with these types of polymorphisms are unclear and could include a dearth or absence of pronounced sex chromosome divergence. A third caveat to our conclusions is that polymorphism among females could also potentially exist in the extent of divergence between the W and Z chromosomes. Under this scenario, it is conceivable that there could be variation among populations in the extent of recombination along the sex chromosomes and consequently the extent and magnitude of divergence between the sex chromosomes. Further exploration of these possibilities will be assisted by the identification of the sex-determining locus in \textit{S. tropicalis}, the completion of high-quality sequencing and assembly of sex-linked regions, and the exploration of variation within and among populations in sex determination and sex chromosome evolution.

### Table 3

| Region                  | Number of Genes (Expression) | Total | Intensity | Evenness | Number of Genes (dN/dS) | dN/dS |
|-------------------------|------------------------------|-------|-----------|----------|-------------------------|-------|
| "Non-7p chromosomes"   | 35,135                       | 0.00065 | 0.00015  | 3.19655  | 9183                    | 0.2702 |
| "Chromosome 7p"        | 2,114                        | 0.00079 | 0.00019  | 3.17054  | 546                     | 0.2850*|
| "Other orphans"        | 260                          | 0.00040* | 0.00012  | 2.59443* | 55                      | 0.2670 |
| "Partially sex-linked" | 246                          | 0.00090 | 0.00018  | 3.37658  | 71                      | 0.2751 |
| "Completely sex-linked"| 35                           | 0.00101 | 0.00049* | 2.56020  | 8                       | 0.2615 |

*Values that are individually significantly different from the "Non-7p chromosomes" (\(P < 0.05\), one-sided permutation tests).
degenerated because, after duplication, the degenerate ancestral sex chromosome would segregate as a new autosomal chromosome, and the resulting homozygous null genotypes could be detrimental (Evans et al. 2012). Sex chromosome degeneration also creates imbalances in allelic copy number between the sexes, which can lead to the evolution of dosage compensation—a factor that is also potentially relevant to polyploid speciation (Orr 1990). Dosage compensation is a process that equalizes expression levels in each sex of a gene that has a different number of alleles in each sex. This could evolve in a species with female heterogamy, for example, through inactivation of one of the Z alleles in males or through upregulation of the Z allele in females. Orr (1990) proposed that dosage compensation in species with a degenerate sex chromosome could act as a barrier to polyploid speciation because dosage compensation would be disrupted when a newly formed triploid individual backcrosses with a diploid parental individual. Our analyses suggest that the sex-specific region of \textit{S. tropicalis} is small, that sex chromosome divergence is minimal, and therefore that dosage compensation associated with degeneration of the sex-specific sex chromosome would have evolved in very few genes or none at all. Together these features of the sex chromosomes may have facilitated (or at least not impeded) polyploidization in \textit{Silurana}, which occurred at least once (reviewed in Evans 2008). Interestingly, the sister genus \textit{Xenopus} has a newly evolved sex-determining gene called \textit{DM-W} (Yoshimoto et al. 2008; Bewick et al. 2011). Species in this group also probably have minimally diverged sex chromosomes and have undergone polyploid speciation multiple times (Evans 2008). Clearly, however, this is not the only consideration in the ability of species to tolerate polyploidization because many amphibian groups that have homomorphic sex chromosomes lack polyploid species.

The extent of sex chromosome degeneration is also relevant to the chances a species experiences future sex chromosome turnover—a change in which pair of chromosomes carries the trigger for sex determination (Charlesworth and Mank 2010). If sex chromosome turnover occurs in a species with a diverged and degenerate sex chromosome, the ancestral degenerate chromosome could segregate autosomally, and some individuals could inherit two copies and be homozygous for degenerate alleles (Charlesworth and Mank 2010). Thus, sex chromosome turnover may be more likely in species that have sex chromosomes that are not substantially degenerated.

If sex chromosome turnover were common, this could maintain homomorphy of sex chromosomes. Recent work on sex chromosomes in African clawed frogs has established nonhomology between the sex chromosomes of \textit{X. laevis} and \textit{S. tropicalis} (Uno et al., submitted) and the recent appearance of a novel sex-determining locus in \textit{X. laevis} (Yoshimoto et al. 2008; Bewick et al. 2011). Thus, it appears that the origin of a new sex-determining gene in \textit{X. laevis} was associated with a reassignment of sex chromosomes without necessarily involving a change in heterogamy, and that recent sex chromosome turnover can account for sex chromosome homomorphy in this species (Tymowska 1991). In other species, including \textit{S. tropicalis}, it is also possible that the sex-determining mechanism of nondiverged sex chromosomes could be old, but that divergence is prevented by periodic recombination, which possibly could be facilitated by breeding individuals that are phenotypically sex reversed (the “fountain of youth” hypothesis; Perrin 2009). Because we do not yet know the sex-determining gene(s) of \textit{S. tropicalis} or other frogs that might have inherited this sex determination system from a recent common ancestor (e.g., genera \textit{Hymenochirus}, \textit{Pseudohymenochirus}, or \textit{Pipa}), we cannot determine at this time whether turnover or recombination best accounts for the apparent homomorphy of the sex chromosomes of this species. Additional identification of sex-determining genes in amphibians, and analysis of their evolutionary histories and genomic context, is thus an exciting direction for future research.

**Supplementary Material**

Supplementary figures S1–S3 and table S1 are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org/).

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