Viviparous Reptile Regarded to Have Temperature-Dependent Sex Determination Has Old XY Chromosomes

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Data deposition: This project has been deposited at NCBI-SRA database (www.ncbi.nlm.nih.gov/sra, last accessed May 25, 2020) under the accession BioProject PRJNA573688. Y-linked sequences found in Eulamprus heatwolei are available in Supplementary Material online. Shapefiles of geographical ranges of reptiles are available in the figshare platform at the following link https://figshare.com/articles/Reptile_shapefiles/7416638 (last accessed May 25, 2020). The program to map climatic data onto geographical ranges is available in the GitHub platform at the following link https://github.com/annakrystalli/IUCNextractR (last accessed May 25, 2020).

Abstract

The water skinks Eulamprus tympanum and Eulamprus heatwolei show thermally induced sex determination where elevated temperatures give rise to male offspring. Paradoxically, Eulamprus species reproduce in temperatures of 12–15 °C making them outliers when compared with reptiles that use temperature as a cue for sex determination. Moreover, these two species are among the very few viviparous reptiles reported to have thermally induced sex determination. Thus, we tested whether these skinks possess undetected sex chromosomes with thermal override. We produced transcriptome and genome data for E. heatwolei. We found that E. heatwolei presents XY chromosomes that include 14 gametologs with regulatory functions. The Y chromosomal region is 79–116 Myr old and shared between water and spotted skinks. Our work provides clear evidence that climate could be useful to predict the type of sex determination systems in reptiles and it also indicates that viviparity is strictly associated with sex chromosomes.

Key words: temperature-dependent sex determination, viviparous reptiles, genetic sex determination systems, water skinks, Eulamprus heatwolei.
Introduction

Eulamprus tympanum and Eulamprus heatwolei Reproduce in Colder Conditions Compared with Other Species with Temperature-Dependent Sex Determination

Vertebrates exhibit two major classes of sex determination systems. Genotypic sex determination (GSD), where genetic components guide the development of the gonads, and temperature-dependent sex determination (TSD), where specific incubation temperatures define the sex of the embryos (Bachtrog et al. 2014). TSD in reptiles is thought to have evolved when external conditions that enhance either male or female offspring fitness could influence the sex of the embryos (Charnov and Bull 1977; Shine 1999). For this reason, the discovery of TSD in a viviparous skink was particularly notable (Robert and Thompson 2001). In viviparous species, the external conditions have little effect because embryonic development and hatching occur inside the mother’s womb in a relatively stable environment.

The viviparous water skinks Eulamprus tympanum and E. heatwolei (family Scincidae) are classified as TSD species (Tree of Sex 2014) because cytogenetic analyses found no evidence of heteromorphic sex chromosomes and female Eulamprus skinks give rise to male offspring when they are kept at warm temperatures (32°C) during pregnancy (Robert and Thompson 2001). Three features, however, make this classification of Eulamprus as TSD suspect: 1) These two species inhabit alpine habitats in southeastern Australia (Cogger 2000), whereas most reptiles with TSD systems inhabit lowland areas; 2) Uniquely, although all known viviparous reptiles have genetic sex determination systems, E. tympanum and E. heatwolei are the only known viviparous reptiles classified as TSD; and 3) Several studies have found 1:1 sex ratios in E. heatwolei at mild temperatures, both in the laboratory and in the field (Schwarzkopf and Shine 1991; Robert and Thompson 2001; Allsop et al. 2006). Taken together, these features implied either a GSD system with thermal override or, although less likely, an atypical TSD system.

We first examined whether ambient temperatures in areas inhabited by E. tympanum and E. heatwolei during breeding seasons were unusual compared with reptile species with TSD or GSD. For this, we mapped 30 years of ambient temperatures onto the geographic ranges of 101 species with TSD and 99 species with GSD during their breeding season (fig. 1). Average ambient temperatures for E. heatwolei and E. tympanum during their breeding seasons are 15 and 12.4°C, respectively (fig. 1). Thus, E. heatwolei and E. tympanum are clear outliers when considered as TSD species, located at 3 and 4 SDs away from the mean of the distribution, respectively (fig. 1). In contrast, Eulamprus species are found within the distribution of species with GSD (fig. 1). These results are suggestive of the presence of previously undetected sex chromosomes in these two species.

Eulamprus heatwolei Has XY Chromosomes

To test for the presence of previously unidentified sex chromosomes in skinks, RNAseq data were generated from brain,
liver, and gonads of one adult male and one adult female
E. heatwolei. We then applied a subtraction approach
(Cortez et al. 2014; Marin et al. 2017) to the male and female
transcriptomic data of E. heatwolei. Specifically, we assem-
bled a male-restricted transcriptome and used male and fe-
male genomic reads to uncover Y-linked transcripts (see
Materials and Methods). We identified Y-linked transcripts
from 14 protein-coding genes with known orthologous genes
located on a single syntenic block on chromosome 5 in
Anolis
carolinensis
and chromosome 1 in chicken (fig. 2
a and supple-
mentary table 3, Supplementary Material online).
Additionally, we performed a male and female genomic
read coverage analysis of six chromosomes of E. heatwolei
(see Materials and Methods). We found a region on chromo-
some 5 where the male shows only half of the coverage (i.e.,
one genomic copy, fig. 2b, and supplementary fig. 1,
Supplementary Material online). XY gametologs map to this
specific region on chromosome 5 (fig. 2a and b) and analysis
of their genomic coverage is consistent with two X gameto-
logs in females but one X and one Y gametolog in males
(supplementary fig. 2, Supplementary Material online).
Lastly, we screened the genomes of seven males and seven
females using standard PCRs and found that we could only
amplify Y-linked sequences in males (fig. 2c and supplemen-
tary fig. 3, Supplementary Material online). In summary, the
results reveal the presence of sex chromosomes in E. heatwolei.

Fig. 2.—(a) Synteny of the 14 XY gametologs in other species. (b) Male (blue) and female (red) genomic coverage along the chromosome 5 of
Eulamprus heatwolei. A syntenic region shows half of the coverage in males (one copy) but regular coverage in females (two copies). XY gametologs map
to this region. Blue arrows show the matching locations of Y-linked markers from Niveoscincus ocellatus. (c) PCR screenings of two males and two females
using primers designed to amplify three Y-linked genes (seven males and seven females were screened in total; see supplementary fig. 3, Supplementary
Material online). (d) Time-calibrated synonymous substitution tree used to estimate the age of the XY chromosomes in E. heatwolei. Branch lengths represent
millions of years.
Viviparous Reptile Has Old XY Chromosomes

**Materials and Methods**

Data Generation

One adult male (Euhea_18_05) and one adult female individual (Euhea_18_03) of *E. heatwolei* species were captured from a population that inhabits Woods Reserve, Corin Road, ACT, Australia (–35.480751, 148.940398). Both individuals were sacrificed by intraperitoneal injection of pentobarbitone following the standard operating procedures specified by the animal ethics committee of the University of Canberra. We generated DNA-seq libraries for a male and female *E. heatwolei* from liver tissue using the Illumina TruSeq DNA protocol for short insert size (400–450 nt). We generated strand-specific RNA-seq libraries (using the Illumina TruSeq Stranded mRNA Library protocol) for a total of six samples obtained from brain, liver, and gonads for a male and female *E. heatwolei*. All libraries were sequenced on Illumina HiSeq 2500 sequencers at the University of Canberra. We generated 262–269 million 150-nt paired-end DNAseq reads. We generated 82–95 million 125-nt paired-end RNAseq reads. Further details in supplementary table 4, Supplementary Material online. Quality of the reads...
was verified using FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc, last accessed May 25, 2020) and the remaining adaptors were removed with Trimmomatic (Bolger et al. 2014).

Assembly of Y-Linked Transcripts

To assemble Y-linked transcripts in *E. heatwolei*, we used a subtraction approach based on male and female RNAseq data (Cortez et al. 2014; Marin et al. 2017; Acosta et al. 2019). Briefly, male RNA-seq reads were aligned onto the de novo reconstructed female transcriptome from *E. heatwolei* using Hisat2 (v2.0.2) (Kim et al. 2015); no mismatches allowed; reads not mapping were selected. We also removed male RNA-seq reads sharing k-mers with the female transcriptome (Akagi et al. 2014). The selected reads were passed to Trinity (v2.0.2, default k-mer of 25 bp) (Grabherr et al. 2011) to assemble transcripts that were only present in male tissues. We obtained 21,249 transcripts that were subsequently aligned to the male and female genomic reads using BlastN (Altschul et al. 1990); at a 100–99% identity threshold. We selected those transcripts showing 4x–14x of averaged coverage of male genomic reads and zero averaged coverage of female genomic reads (supplementary table 3, Supplementary Material online). To establish Y gene identity, we searched NCBI GenBank (Reptile taxa only; http://www.ncbi.nlm.nih.gov/genbank, last accessed May 25, 2020) with BlastN and BlastX for the closest homologs and identified transcripts that coded for 14 proteins (supplementary table 3, Supplementary Material online). BlastX searches also allowed the identification of CDS regions. For these 14 Y-linked protein-coding genes, we performed BlastN searches against the de novo reconstructed female transcriptome from *E. heatwolei* to find the X gametologs (best match over the entire sequence; 95–97% identity). We verified the X gametologs identity using coverage analyses of male and female genomic reads and GenBank searches (same gene identity as Y gametologs). XY gametologs in *E. heatwolei* were searched against the *A. carolinensis* and chicken genomes using the sequence search engine at the ENSEMBL webpage (https://www.ensembl.org/Multi/Tools/Blast, last accessed May 25, 2020) to establish whether they formed a syntenic block. We validated the presence of a Y chromosome by PCR screenings using genomic DNA obtained from tails snips of seven males and seven females. Additional information can be found in the extended Materials and Methods section in the Supplementary Material online.

Genomic Coverage Analyses

We followed a methodology previously published (Vicoso et al. 2013). Briefly, the male and female genomic reads were assembled into contigs. The contigs were subsequently aligned and ordered based on the *A. carolinensis* reference genome. We used bowtie2 (Langmead and Salzberg 2012) to align the DNA-seq reads from the male and female *E. heatwolei* onto the reconstructed chromosomes. Coverage along the chromosomes was calculated using BEDTools (Quinlan and Hall 2010), bins of 100,000 nucleotides. Additional information can be found in the extended Materials and Methods section in the Supplementary Material online.

Data Collection

Full list of reptiles with known TSD system was obtained from the Tree of Sex database (Tree of Sex 2014) and literature searches. We searched the literature and dedicated databases for the duration and month intervals of the breeding seasons. We collected information for 101 species with TSD (supplementary tables 1 and 2, Supplementary Material online). Temperature data from the entire surface of the planet were downloaded from the Climatic Research Unit (http://catalogue.ceda.ac.uk/uuid/3df7562727314bab963282e6a0284f24, last accessed May 25, 2020; version 3.24.01). Additional information can be found in the extended Materials and Methods section in the Supplementary Material online.

Geographical Ranges

Shapefiles for 29 species where downloaded from the RedList database (http://www.iucnredlist.org/, last accessed May 25, 2020; version 3; supplementary table 1, Supplementary Material online). For 72 additional species (supplementary table 2, Supplementary Material online) we generated geographic ranges using the ecological niche modeling routines applying the maximum entropy algorithm in Maxent (Phillips et al. 2006) using the R package kuemn (Cobos et al. 2019). Additional information can be found in the extended Materials and Methods section in the Supplementary Material online.

Mapping Climate Data to the Species Distribution

We matched the climate data with the species shapefiles using a dedicated R package built by Dr Anna Krystalli as part of the Newton Advanced Fellowship program (https://github.com/annakrystalli/UCNexttractR, last accessed May 25, 2020). We recovered the median temperature (ambient temperature) of all months comprised in the breeding season. Additional information can be found in the extended Materials and Methods section in the Supplementary Material online.
Synonymous Substitution Analyses

To assess the age at which the XY system was originated in *E. heatwolei*, we followed a previous procedure (Cortez et al. 2014; Marin et al. 2017; Acosta et al. 2019). Briefly, we aligned using PRANK (Loytynoja and Goldman 2005) the coding sequences of XY gametologs in *E. heatwolei* and coding sequences of 1–1 orthologous in other reptiles, mammalian and Xenopus species downloaded from the Ensembl database (https://www.ensembl.org/, last accessed May 25, 2020). We obtained the species’ tree from the TimeTree database (http://www.timetree.org/, last accessed May 25, 2020). We concatenated the alignments and calculated synonymous substitution rates (*dS*) using codeml (Yang 1997) and a bootstrap approach. Branch lengths on the species’ tree were used to obtain an ultrametric, time-calibrated, tree using the chronos library (ape package in R, v5.0) (Paradis and Schliep 2019). The age of the sex chromosomes was obtained from the calibrated branch lengths just before and after the split of the XY gametologs and the time since *E. heatwolei* diverged from the Snake–Pogona–Anolis lineage (divergence data retrieved from TimeTree; http://www.timetree.org/, last accessed May 25, 2020). We also calculated the age of the sex chromosomes using BEAST v1.10.4 (http://beast.bio.ed.ac.uk/), which resulted in an age estimate of 93 Ma. We used the relaxed clock and calibrated the tree based on the reptile/mammalian divergence time. We ran the analyses two independent times for 100,000,000 generations, sampling every 1,000 generations. Additional information can be found in the extended Materials and Methods section in the Supplementary Material online.

Supplementary Material

Supplementary data are available at Genome Biology and Evolution online.

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Author Contributions

D.C., A.O.U., T.S., and A.G. designed the study. D.S.B.D. and A.G. performed the fieldwork, tissue collection, DNA and RNA extractions, and PCR screenings. P.C.-P., D.S.B.D., A.L.N., M.L.M.P., A.A., and D.C. performed the analyses. C.R.-S. performed additional analyses. F.R.M.C. contributed to the analyses, discussion, and ecological data collection. All authors contributed to the interpretation of the results. A.O.U., A.G., T.S., and D.C. wrote the article. All authors read and approved the final article.

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