

It's a Trap! Escape from an ancient, ancestral sex chromosome system
and implication of *Foxl2* as a primary sex determining
gene in a lizard (Anguimorpha; Shinisauridae)

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Abstract

Although sex determination is ubiquitous in vertebrates, mechanisms of sex determination vary from environmentally- to genetically-influenced. Genetic sex determination is typically accomplished with sex chromosomes via the presence/absence of a specific allele at a genetic locus, which initiates the sex determination signaling network. In vertebrates, well-studied groups like mammals and birds maintain conserved sex chromosome systems (XY and ZW, respectively), while sex chromosomes in many other clades may not be conserved across long evolutionary timeframes. Among squamate reptiles, for example, some groups are relatively stable, like iguanids (Pleurodonta), caenophidian snakes (Caenophidia), and lacertids (Lacertidae) while others are highly variable in their modes of sex determination, such as geckos (Gekkota) and the clade containing chameleons and dragon lizards (Acrodonta). One group inferred to have an evolutionarily stable mode of sex determination is the Anguimorpha, a clade of charismatic taxa, including monitor lizards (*Varanus*), beaded lizards (*Heloderma*), glass lizards (*Ophisaurus*), and the crocodile lizard (*Shinisaurus crocodilurus*). The most recent common ancestor to extant anguimorphs possessed a ZW system that has been retained across the clade. However, determination of sex chromosomes in some anguimorph species has remained elusive including in the endangered, monotypic family of crocodile lizards (Shinisauridae). Here, we use a combination of whole genome re-sequencing and RADseq to demonstrate that *S. crocodilurus* has replaced the ancestral anguimorph ZW system on their LG7 chromosome with a novel ZW system on LG3. The syntenic region that corresponds with this linkage group in chicken is chromosome 9, and this is the first documented use of this syntenic block as a sex chromosome in amniotes. Additionally, this ~1Mb region harbors approximately 10 genes, including a duplication of the well-characterized transcription factor, *Foxl2*—a critical locus for the maintenance of sexual differentiation in vertebrates, and thus a putative primary sex determining gene for *S. crocodilurus*.

Introduction

The evolution of sex determination in vertebrates is impressive in its ability to combine a highly conserved developmental network that can be initiated by quite distinct molecular mechanisms in different species (Bachtrog et al. 2014; Graves, 2008). In vertebrates, sex is commonly determined via either environmental and/or genetic cues at critical points in development. In vertebrate groups that use genetic mechanisms, the most common mechanism is sex chromosomes; either a male or female heterogametic system where the male or female inherits the sex-limited (Y or W) chromosome, respectively (Bachtrog et al. 2014; Gamble et al. 2015). Sex chromosomes have been traditionally identified by comparing male and female karyotypes under the light microscope. The presence of morphological differences between the X and Y (or Z and W) chromosomes (i.e. heteromorphic sex chromosomes) identify a species' sex chromosome system (Stevens, 1905; Bull, 1983). However, many species possess sex chromosomes that cannot be identified via light microscopy because the X and Y (or Z and W) are not morphologically distinguishable from each other (i.e. homomorphic sex chromosomes). Other methods must be employed, such as advanced cytogenetic techniques or high-throughput DNA sequencing technologies, to identify sex chromosome systems in these taxa (Gamble and Zarkower, 2014; Pinto et al. 2022).

Squamate reptiles (lizards and snakes) demonstrate high variability in modes of sex determination: where some clades have conserved, often heteromorphic, sex chromosomes, while others display extraordinary lability in their modes of sex determination and a high incidence of homomorphic sex chromosomes (Gamble et al. 2015, Kratochvíl et al. 2021; Augstenová et al. 2021a). One hypothesis of sex chromosome evolution is that ancient, degenerated sex chromosome systems may act as an “evolutionary trap”, where the existence of highly differentiated (i.e. heteromorphic) sex chromosomes preclude transitions to other sex-determining systems (Bull 1983; Bull and Charnov, 1985; Pokorná and Kratochvíl, 2009). The stability of old sex chromosome systems in mammals, birds, caenophidian snakes, and others, provides anecdotal support for this hypothesis (Bull and Charnov, 1985;

Pokorná and Kratochvíl, 2009; Gamble et al. 2015). As more and more sex chromosome transitions are identified, it remains unclear whether all sex chromosomes are destined to become traps. And examples of taxa transitioning away from ancient, degenerated sex chromosome systems are rare (Acosta et al. 2019; Nielsen et al. 2019; Rovatsos et al. 2019a). Previous phylogenetic studies have supported the trap hypothesis in squamates (Pokorná and Kratochvíl, 2009; Gamble et al. 2015), but also suffered from incomplete taxonomic sampling, which likely biased the conclusions. In other words, testing this hypothesis is contingent upon having sufficient data necessary to identify a transition away from a sex chromosome system which typically requires a reference genome to coordinate linkage groups (which are rare in squamates; Pinto et al. 2023), genome-scale data from both sexes (e.g. Vicoso et al. 2013; Gamble et al. 2015; Pinto et al. 2022), and a robust phylogenetic hypothesis to establish relationships within the focal taxa (Nielsen et al. 2019). Thus, the burden of proof is higher for identifying escapes from these putative “evolutionary traps,” which may be partially responsible for the dearth of examples.

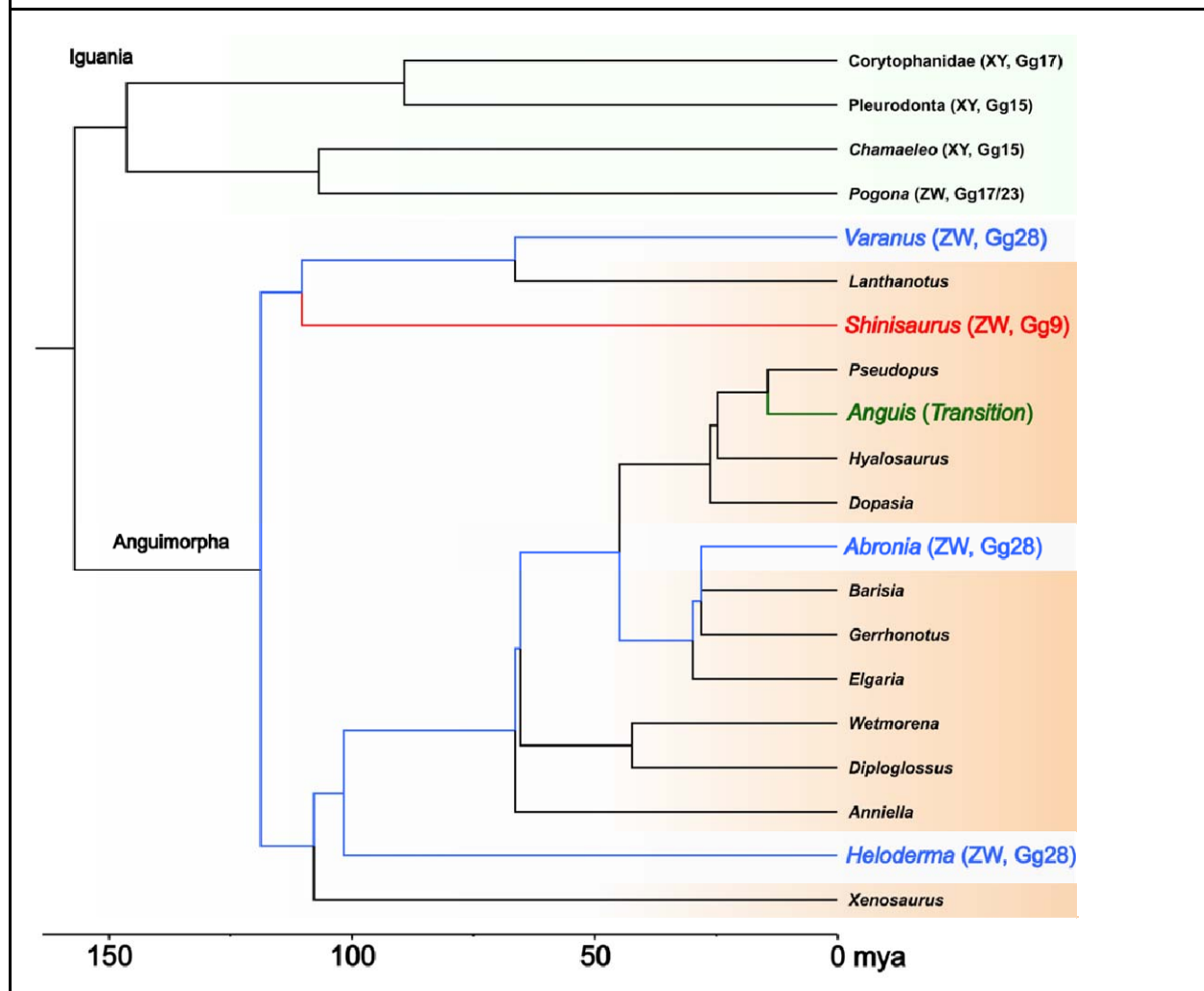
The sex chromosomes of the infraorder Anguimorpha (lizards including monitor lizards, Gila monsters, alligator lizards, and their allies) have been a topic of interest for many years, likely resulting from the paucity of genetic and cytogenetic data for this group. In recent years, advanced cytogenetic techniques (FISH) have facilitated karyotypic analysis and identification of ZW sex chromosomes in the Gila monster (*Heloderma suspectum*; Pokorná et al. 2014) and Komodo dragon (*Varanus komodoensis*; Pokorná et al. 2016) using leading to expanded interest in studying chromosome evolution in this enigmatic group. More recently, RNAseq and qPCR analysis, in conjunction with draft genomes of these same two anguimorph species (Gila monster; Webster et al. 2023, and Komodo dragon; Lind et al. 2019), have provided some additional insights into this system (Rovatsos et al. 2019b). Namely, the homology of the ZW systems in the anguimorph genera *Abronia*, *Heloderma*, and *Varanus* (Rovatsos et al. 2019b; Webster et al. 2023). The presence of a ZW sex chromosome on the same linkage group—syntenic with chromosome 28 in the chicken genome—in these three genera, spanning the phylogenetic breadth of extant

Anguimorpha, is strong evidence that this is the ancestral sex chromosome system in the clade. Ancient sex chromosome systems, like those ancestral to anguimorphs (115–180 million years old), fit the criteria that should render them as an evolutionary trap (Pokorná and Kratochvíl, 2009; Rovatsos et al. 2019b). However, the sex chromosomes of many anguimorph taxa remain unknown, including the monotypic family Shinisauridae, which is nested deep within the anguimorph phylogeny (Figure 1).

The crocodile lizard (*Shinisaurus crocodilurus*) is the sole living member of the family Shinisauridae and native to small disjunct regions of southeastern China and northern Vietnam (Le and Ziegler, 2003; Huang et al. 2008; Nguyen et al. 2014). It is one of the rarest lizard species in the world and is listed as Endangered in the IUCN Red List (Nguyen et al. 2014). Due to anthropogenic pressures, populations have experienced severe declines in recent years with less than 1000 individuals in the wild in China and less than 100 adults in Vietnam (Huang et al. 2008; van Schingen et al. 2014a). They are semi-aquatic habitat specialists and depend upon clean streams in broadleaf evergreen forest (Ning et al. 2006; van Schingen et al. 2016a) and their restricted ecological niche is predicted to all but disappear due to climate change by the end of this century (Li et al. 2013; van Schingen et al. 2014a; see also van Schingen-Khan et al. 2022). Habitat destruction threatens remaining populations, as well as being collected for food and the international pet trade (Huang et al. 2008; van Schingen et al. 2014b; van Schingen et al. 2016a). Although still recognized as a single species, there exist several conservation units, with *S. crocodilurus vietnamensis* currently only known from Vietnam and the nominate subspecies from China consisting of several distinct lineages (van Schingen et al. 2016b; Ngo et al. 2020; Nguyen et al. 2022). Crocodile lizards do not have a clear sexual dimorphism. While morphological traits, such as coloration or body morphometry may provide an indication of the sex, a robust method to reliably identify the sex of individuals is lacking (van Schingen et al. 2016b). Relevant to the present study, examination of male and female *S. crocodilurus* karyotypes have revealed no heteromorphic sex chromosomes (Zhang et al. 1996; Augstenová et al. 2021b). To identify sex chromosomes in this species, we analyzed whole-genome re-sequencing data for more than 50 individual crocodile lizards (held animals originating

from China; Xie et al. 2022) using whole-genome re-sequencing and, in parallel, restriction site-associated DNA sequencing (RADseq) data for 13 additional individuals (held animals originating from both China and Vietnam) to show that the sex determining system in *S. crocodilurus* is a novel ZW system that has eluded previous analyses, at least in part, due to the small size (<1Mb) of its sex determining region (SDR).

Figure 1: Summary of current anguimorph sex chromosome knowledge summarized from Rovatsos et al. (2019) indicated by blue and green tips/branches); information identified in this study indicated by red tips/branches and what remains unknown across the phylogeny indicated by black tips/branches. Phylogeny from TimeTree using a representative species from each clade (Kumar et al. 2017) and visualized using Figtree [v1.4.4] (<http://tree.bio.ed.ac.uk/software/figtree/>). “Gg” stands for chicken (*Gallus gallus*) linkage group.



Methods

WGS analysis

We downloaded whole genome resequencing reads from NCBI SRA for multiple male and female individuals (see *Data Availability* for accessions). We constructed a Snakemake [v6.10.0] (Mölder et al. 2021) workflow in an isolated conda environment [v4.11.0] (<https://docs.anaconda.com/>) containing relevant packages: BBmap [v38.93] (Bushnell, 2014), FastQC [v0.11.9] (Andrews, 2010), Freebayes [v1.3.5] (Garrison and Marth, 2012), GFF utilities [v0.10.1] (Pertea and Pertea, 2020), Minimap2 [v2.22] (Li, 2018), Mosdepth [v0.3.2] (Pedersen and Quinlan, 2018), MultiQC [v1.11] (Ewels et al. 2016), Parallel [v20211022] (Tange, 2018), pixy [v1.2.5.beta1] (Korunes and Samuk, 2021), Sambamba [v0.8.1] (Tarasov et al. 2015), Samtools [v1.12] (Li et al. 2009), seqkit [v0.11.0] (Shen et al. 2016), STACKS [v2.6.0] (Catchen et al. 2013), Trim Galore! [v0.6.7] (<https://doi.org/10.5281/zenodo.5127899>), and vcftools [v0.1.15] (Danecek et al. 2011). To process the raw sequencing data, we trimmed adapters and low-quality regions using Trim Galore!, then removed PCR duplicates using BBmap. Quality assessment using FastQC and MultiQC was conducted at each step and we subsequently removed samples with fewer than 5 million PE reads after filtering PCR duplicates. The final dataset possessed 28 male and 23 female individuals. We proceeded to map reads for each individual to the female reference genome (Xie et al. 2022) with minimap2 and calculated read depth and read mapping statistics using mosdepth and samtools, respectively. Then generated an all-sites vcf file with freebayes-parallel. Lastly, we calculated Weir and Cockerham (1984) F_{ST} and nucleotide diversity statistics using pixy at 500kb resolution and, for LG3 only, also at 100kb resolution.

RADseq analysis

In a similar fashion, we analyzed a modest dataset of restriction site-associated DNA sequencing (RADseq) for four males and six females (Supplemental Table 1). We followed a modified protocol from Etter et al. (2011) as outlined in Gamble et al. (2015). Libraries were size-selected into 300- to 600-bp fragments using magnetic beads in a PEG/NaCl buffer (Rohland and Reich, 2012), then pooled and sequenced using paired-

end 150-bp reads on an Illumina® HiSeqX at Psomagen®. Similarly to the WGS data, raw illumina reads were trimmed using Trim Galore!, PCR duplicates removed using BBmap, and mapped to the reference genome using minimap2 with quality assessment using FastQC and MultiQC at each step. Diverging from the WGS pipeline, we processed the RADseq data herein using the `refmap.pl` script included in the STACKS suite and calculated Weir and Cockerham (1984) F_{ST} using VCFtools.

Validation of the putative ZW system in Shinisaurus crocodilurus

We generated a dataset of ‘*in silico* poolseq’ reads by subsampling each read file to 10 million reads (20 million per sample) using seqkit and combined into male and female pools. We analyzed the pools using Pooled Sequencing Analyses for Sex Signal [PSASS; v3.1.0] (<https://doi.org/10.5281/zenodo.3702337>). We then generated PCR primers targeting the annotated version of *Foxl2*’s second exon [FOXL2-ex2-F2 5’ – CAGAGCTCGTCCCATTCACTT – 3’ and FOXL2-ex2-R2 5’ – GAGAGATGTACCACCGGGAG – 3’] and sequenced the resultant amplicon using Sanger sequencing (Psomagen).

Genome Annotation

We used previously lifted over annotations (Pinto et al. 2023; <https://doi.org/10.6084/m9.figshare.20201099.v1>) from the draft genome of a male *S. crocodilurus* (Gao et al. 2017) to the new, unannotated female reference genome (Xie et al. 2022; GCA_021292165.1). We pulled coding transcripts from the genome using GFF Utilities. We used the 10 genes within the putative ~1Mb SDR on LG3 to perform a high-stringency tBLASTx query (Altschul et al. 1990) to the chicken genome on Ensembl (Howe et al. 2020) with a word size of 3, maximum of 10 hits, e-value cutoff of $1e^{-50}$, using BLOSUM62 scoring matrix. These queries received hits on 7 of the 10 total genes (Table 1).

Results

Across WGS experiments, read mapping efficiency ranged from 80.60% (for SRR5019740) to 99.40% (for SRR14583318). In RADseq experiments, read mapping efficiency ranged from 96.03% to 98.67%. Across both experiments, only one F_{ST} peak appeared in both datasets (Supplemental Figure 1). The poor resolution of the RADseq dataset, compared with the whole genome resequencing data, likely results from the small number of individuals used and the relatively small size of the non-recombining portion of the sex chromosome. Comparing M/F F_{ST} values does not necessarily diagnose which sex is heterogametic (i.e. XY vs. ZW). Therefore, we composed a dataset of ‘*in silico* poolseq’ reads to identify an excess of female-associated SNPs aligning to the previously identified region of high M/F F_{ST} (Supplemental Figure 2). Taken together, these data suggest that *S. crocodilurus* possesses a female heterogametic system (ZW) with an SDR located in a ~900kb region on LG3.

Upon further investigation of the SDR, we identified a total of 10 genes annotated within this region. To better characterize these genes, we BLAST-ed each to the chicken genome. We recovered high-quality BLAST hits for seven of the 10 annotated *S. crocodilurus* SDR genes in chicken (Table 1). Six out of the seven queries hit genes located on chicken chromosome 9, while the other landed on a chicken chromosome 30 (Table 1). In our poolseq analysis, one of these genes possessed half the read depth in females relative to males and, upon closer inspection, we identified a duplicated, unannotated copy of that gene Forkhead Box L2 (*Foxl2*) also located within the SDR. We included this *Foxl2* copy in a BLAST search against chicken, where it was again identified as a *Foxl2* homolog (Table 1). We also BLAST-ed *Foxl2* to the earlier male *S. crocodilurus* draft genome (Gao et al. 2017) and found only a single copy of *Foxl2* in this genome matching one copy in the updated reference genome with 100% sequence identity, consistent with both (1) the duplicated version being W-specific and (2) the female reference being chimeric for Z and W alleles (Xie et al. 2022). Lastly, we generated a gene tree using *Foxl2* copies from across reptiles to confirm its duplicated origination was within Shinisauridae (Supplemental Figure 3). Thus, in the chimeric female reference genome, this putative W-linked *Foxl2* copy was located approximately

70kb upstream of the annotated Z-linked copy of *Foxl2* on the other side of an assembly gap.

Our WGS data used in poolseq and other analyses were restricted to only individuals from Chinese populations to reduce the influence of population-specific demographic processes (Xie et al. 2022). To include the less-numerous Vietnam samples, we generated PCR primers for a segment of *Foxl2*'s second exon and Sanger sequenced multiple females (Vietnam) and males (China and Vietnam) (Supplemental Figure 4). We identified one SNP in the female Vietnamese samples in this region and tested its association with sex using Fisher's exact test (p-value = 0.0047***). Thus, the ZW SDR containing *Foxl2* appears to be conserved between populations of *S. crocodilurus* from both China and Vietnam.

Table 1: Top tBLASTx hits in chicken for each of the genes present in the *Shinisaurus* 900kb-SDR. The duplicated *Foxl2* copy is dubbed ENSGALP00000033127-W.

<i>Shinisaurus</i> Gene ID	Chicken Gene Id	Location (Chicken)	E-value
ENSACAP00000003394-D1	ENSGALG00000026187	9:6115461-6115832	1.42E-95
ENSGALP00000008531-D1	<i>RBP1</i>	9:6110534-6110713	7.54E-54
ENSACAP00000003392-D1	<i>No hits.</i>		
ENSACAP00000003371-D1	ENSGALG00000034575	30:1402166-1402369	2.51E-78
ENSACAP00000003355-D1	ENSGALG00000005357	9:6041607-6041822	0
ENSACAP00000003221-D1	ENSGALG00000005367	9:6037018-6037164	1.03E-80
ENSGALP00000033127-W	<i>FOXL2W</i>	9:5875297-5875587	2.44E-76

ENSGALP00000033127-D1	<i>FOXL2</i>	9:5875297-5875587	6.78E-77
ENSACAP00000003172-D1	<i>PIK3CB</i>	9:5800560-5800790	0
ENSGALP00000040175-D1		<i>No hits.</i>	
ENSACAP00000002765-D1		<i>No hits.</i>	

Discussion

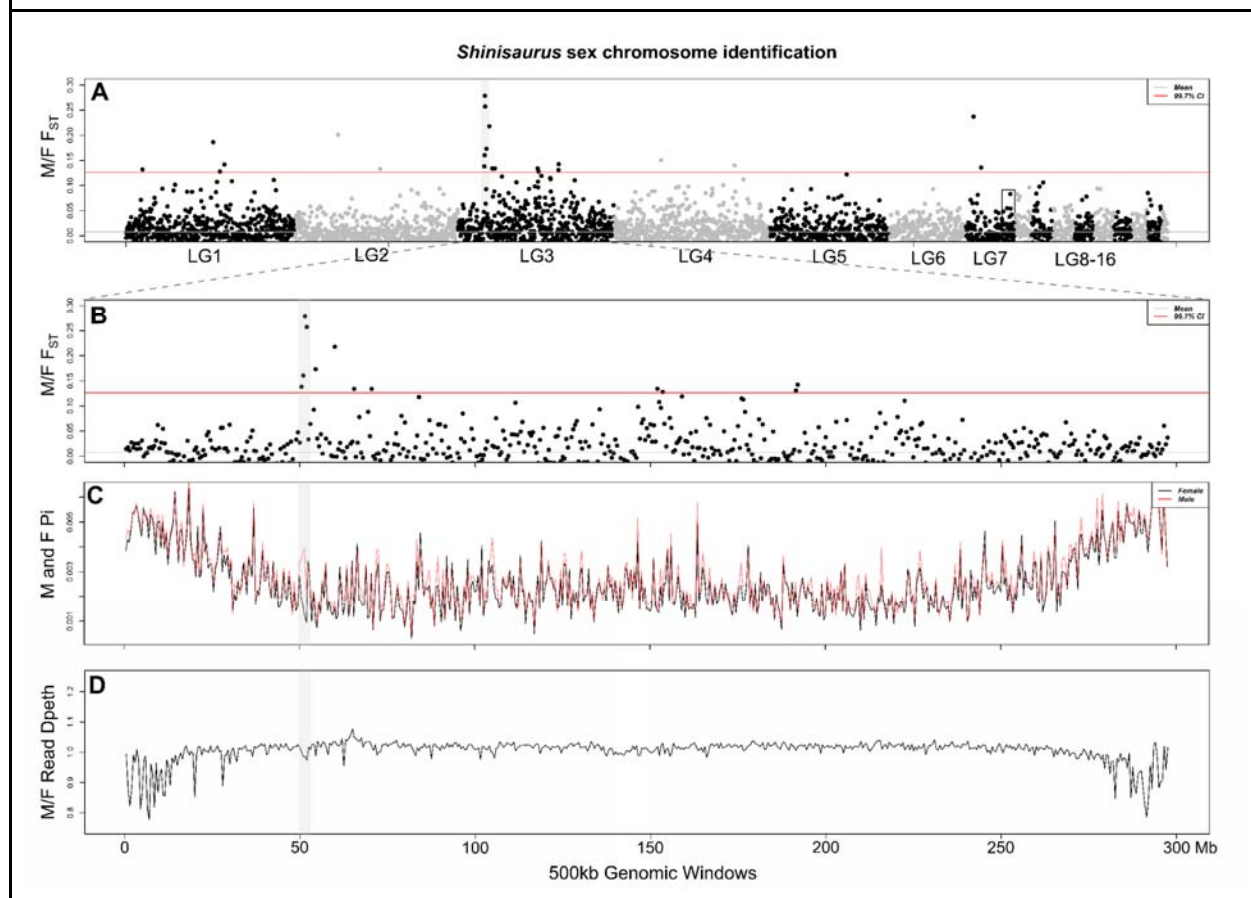
Escaping the “Evolutionary Trap”

An open question within sex chromosome evolution is whether ancient, degenerated sex chromosomes act as evolutionary traps (Pokorná and Kratochvíl, 2009; Nielsen et al. 2019; Pinto et al. 2023). The most recent common ancestor of extant anguimorphs is thought to have possessed a ZW system on the linkage group syntenic with chicken chromosome 28, which is located on the distal region of LG7 in the *S. crocodilurus* reference genome (Rovatsos et al. 2019b; Webster et al. 2023). The sex determining region (SDR) in *S. crocodilurus* is located on LG3, a region syntenic with chicken chromosome 9. To our knowledge, this is the first demonstration in a tetrapod of the syntenic region of chicken chromosome 9 being recruited in a sex determining role (Kratochvíl et al. 2021), lending further support to the idea that all chromosomes will likely be recruited into a sex determining role given thorough enough phylogenetic sampling (Graves and Peichel, 2010; Hodgkin, 2002; O’Meally et al. 2012; Pinto et al. 2022).

It is clear from these genomic data that *S. crocodilurus* possesses a distinct sex chromosome system from all other known anguimorphs. Unlike the case of Corytophanidae and other pleurodonts, where phylogenetic relationships among taxa were inconclusive (Nielsen et al. 2019), the relationship of *S. crocodilurus* to all other anguimorphs is far less divisive. Indeed, *S. crocodilurus* is well-supported as nested within Anguimorpha—either sister to Varanidae as a member of the “Paleoanguimorpha” (Burbrink et al. 2020) or as sister to a clade containing Varanidae and Lanthanotidae (Singhal et al. 2021), depending on taxonomic sampling. Thus, assuming the hypothesis that an ancient origin of the ZW sex chromosome system possessed by extant *Varanus*, *Heloderma*, and *Abronia* is correct, then *S. crocodilurus* has successfully escaped the evolutionary trap of their ancestral, degenerated sex chromosome system—a system nearly as ancient as those systems found in both mammals and birds (Rovatsos et al. 2019b; Webster et al. 2023). It is worth noting that there remains another putative escape from the ancestral anguimorph sex chromosome system in *Anguis* that has yet to be explored further (Rovatsos et al. 2019b) and more

recent phylogenetic work has implicated that Corytophanidae is likely nested somewhere within other pleurodonts, rather than being sister to all other species (Burbrink et al. 2020; Singhal et al. 2021). This suggests that there are a minimum of two evolutionary escapes within Toxicofera (snakes, iguanians, and anguimorphs)—and perhaps even two within the infraorder Anguimorpha alone.

Figure 2: Identification of the ZW sex chromosome system in *Shinisaurus crocodilurus*. (A) Whole genome F_{ST} scan with a clear peak in a ~1Mb region on LG3. The square block on LG7 is syntenic with the sex-determining region in *Varanus* and *Heloderma* (Webster et al. 2023). (B) Isolation and magnification of LG3 F_{ST} peak. (C) Modest increase in male, relative to female, nucleotide diversity and (D) decrease in male/female read depth in the region corresponding to the F_{ST} peak on LG3.



Primary Sex Determination in Shinisauridae

In many vertebrate groups where the primary sex determiner (PSD) is known, a relatively short list of commonly-recruited PSDs have been identified (i.e. the 'usual

suspects'; Adolphi et al. 2021; Dor et al. 2019; Herpin and Schartl, 2015). Indeed, the same genes, or paralogs, have been independently co-opted to function as the PSD in many taxa, including *Sox3* in placental mammals and some medaka (members of the *Oryzias celebensis* and *O. javanicus* groups); *Amh* in tilapia, northern pike, and potentially other anguimorphs (Li et al. 2015; Myosho et al. 2015; Pan et al. 2019; Rovatsos et al. 2019; Webster et al. 2023; and see Pan et al. 2021 for recent review); and *Dmrt1* in birds, a frog (*Xenopus laevis*), tongue sole, and other medaka fish (members of the *Oryzias latipes* group) (Chen et al. 2014; Ioannidis et al. 2021; Matsuda et al. 2002; Nanda et al. 2002; Smith et al. 2009).

This is the first time Forkhead Box L2 (*Foxl2*) has been implicated as a PSD in a vertebrate, although it has been predicted to be one (Ma et al. 2022). *Foxl2* is a direct transcriptional activator of aromatase, involved in development of the ovaries, and its loss in mice during embryogenesis leads to abnormal ovarian development and infertility (Fleming et al. 2010; Pannetier et al. 2006; Schmidt et al. 2004; Uda et al. 2004). After primary sex determination and sexual development has concluded, *Dmrt1* and *Foxl2* antagonize each other transcriptionally in gonadal tissue, where sustained *Dmrt1* and *Foxl2* expression is required for adult maintenance of testis and ovary tissue, respectively (Garcia-Ortiz et al. 2009; Matson et al. 2011; Uhlenhaut et al. 2009). Importantly, *Dmrt1* has been recruited to act as a primary sex determining gene in multiple taxa, while *Foxl2* has remained mysteriously absent from this list—with the singular putative exception being recently described in some species of bivalve mollusks (Han et al. 2022). Thus, the identification of both *Foxl2* and a duplicated *Foxl2* copy in the W-limited region of the *Shinisaurus* genome confirms the expanded list of the “usual suspects” that can act as the PSD in vertebrates.

Pragmatically, the identification of a novel ZW system in *S. crocodilurus* may present an important juncture in the conservation efforts of this endangered lizard species, that are urgently needed (Nguyen et al. 2014). Body morphometrics in mature specimens may provide an indication of the sex, i.e. males tend to have a relatively larger head, relative to abdomen length than females (van Schingen et al. 2016b).

However, definitive sexually dimorphic characters are lacking in the species, especially in hatchlings, juveniles, and subadults. Therefore, a molecular genetic sex test could assist in well-managed captive breeding efforts in this species (Ziegler et al. 2019). This is vital as it's estimated only ~1,000 individuals remain in the wild populations in China and Vietnam during the last census (Huang et al. 2008; van Schingen et al. 2016a), while loss of remaining habitats as well as poaching is considered ongoing.

In conclusion, using a combination of sequencing and validation techniques we identified the elusive ZW system in the endangered crocodile lizard, *Shinisaurus crocodilurus*. This ZW system is located on LG3 and, although interpretation inherits strong reference bias, the SDR appears to be <1Mb in size and contains approximately 10 genes. One of these genes, *Foxl2*, possesses a duplicated copy and is important in ovarian development and fertility in vertebrates. Because of its sequence conservation (either strictly age-related or via gene conversion) and proximity to the original Z copy of *Foxl2*, we hypothesize that if *Foxl2* is the PSD in this system, it's likely a gene dosage-dependent mechanism, where ZW females possess three copies of *Foxl2* instead of the two copies of ZZ males. Our proposed mechanism would essentially be the inverse of *Dmrt1* sex determination in birds, where a lack of *Dmrt1* on the W decreases *DMRT1* expression in females, allowing for *Foxl2* to proceed with ovarian development (Ioannidis et al. 2021; Smith et al. 2009). Here, extra gene copies of *Foxl2* increase *FOXL2* expression to initiate ovarian development in the developing gonad. This information may play a vital role in conservation efforts of this species and this information should be adopted in ongoing captive breeding work (Ziegler et al. 2019).

Data Availability:

The data used in this study is available on NCBI, SRA accessions for WGS data are: SRR14583317, SRR14583321, SRR14583324-26, SRR14583330, SRR14583333, SRR14583340-49, SRR14583351, SRR14583353-54, SRR14583356, SRR14583360-66, SRR5019733-45, SRR14583318-20, SRR14583322-23, SRR14583331, SRR14583334-39, SRR14583346, SRR14583350, SRR14583352, SRR14583355, SRR14583357-59. RADseq and HiC data generated in this study are available on SRA under BioProject PRJNA975696 (SAMN35342808-17).

Code used in the WGS analysis is available on GitHub:

https://github.com/DrPintoThe2nd/Shinisaurus_ZW.

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