

GENETICS

Differential intron retention in *Jumonji* chromatin modifier genes is implicated in reptile temperature-dependent sex determination

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In many vertebrates, sex of offspring is determined by external environmental cues rather than by sex chromosomes. In reptiles, for instance, temperature-dependent sex determination (TSD) is common. Despite decades of work, the mechanism by which temperature is converted into a sex-determining signal remains mysterious. This is partly because it is difficult to distinguish the primary molecular events of TSD from the confounding downstream signatures of sexual differentiation. We use the Australian central bearded dragon, in which chromosomal sex determination is overridden at high temperatures to produce sex-reversed female offspring, as a unique model to identify TSD-specific features of the transcriptome. We show that an intron is retained in mature transcripts from each of two *Jumonji* family genes, *JARID2* and *JMJD3*, in female dragons that have been sex-reversed by temperature but not in normal chromosomal females or males. *JARID2* is a component of the master chromatin modifier Polycomb Repressive Complex 2, and the mammalian sex-determining factor *SRY* is directly regulated by an independent but closely related *Jumonji* family member. We propose that the perturbation of *JARID2/JMJD3* function by intron retention alters the epigenetic landscape to override chromosomal sex-determining cues, triggering sex reversal at extreme temperatures. Sex reversal may then facilitate a transition from genetic sex determination to TSD, with *JARID2/JMJD3* intron retention preserved as the decisive regulatory signal. Significantly, we also observe sex-associated differential retention of the equivalent introns in *JARID2/JMJD3* transcripts expressed in embryonic gonads from TSD alligators and turtles, indicative of a reptile-wide mechanism controlling TSD.

INTRODUCTION

Since the initial discovery of vertebrate temperature-dependent sex determination (TSD) in 1966 (1), the mechanism by which temperature exerts its influence on sexual development has been extensively investigated (1–6). Because males and females in a TSD species have the same chromosome complement, sexual differentiation must be accomplished by differential regulation of common genes, rather than by the presence, absence, or dosage of a master sex-determining gene (7). Temperature may exert its influence at any point in the developmental cascade during which the embryo remains sexually labile (7, 8).

The Australian central bearded dragon (*Pogona vitticeps*) is a uniquely powerful model with which to disentangle primary from emergent molecular events associated with TSD. Dragons are exceptional for the fact that genotype (a ZW/ZZ sex chromosome system) interacts with environmental temperature to determine the fate of sexual differentiation (9, 10). ZW embryos develop as female, and ZZ embryos develop as male (11). However, at high incubation temperatures, this system is overridden so that both ZZ and ZW embryos develop as female (Fig. 1A) (9, 10). The mating of a normal ZZ male dragon with a sex-reversed ZZ female necessarily yields ZZ offspring,

whose sex is determined solely by incubation temperature (9). Because female development can be specified independently by temperature or sex chromosome complement, it is possible to distinguish the effects of temperature and genotype on sex determination (Fig. 1A).

Here, we show that sex-reversed female dragons are distinguished transcriptionally from normal chromosomal females and males by (i) up-regulation of the *Proopiomelanocortin* (*POMC*)–mediated environmental stress response and (ii) the preferential retention of introns in mRNA transcribed from each of two *Jumonji* family genes, *JARID2* and *JMJD3*. The retained intron sequences contain premature stop codons and are very likely to alter or abolish the function of these important epigenetic regulators. The same *Jumonji* introns are also differentially retained between embryonic gonads developing at either female-producing temperature (FPT) or male-producing temperature (MPT) in both alligator and turtle, which is suggestive of an ancient mechanism that may control reptile TSD.

RESULTS

To identify changes in gene expression and/or alternative splicing associated with sex reversal and thus TSD, we analyzed RNA sequencing data from adult gonad, brain, and other somatic tissues (liver, kidney, lung, heart, and muscle) in normal female (ZWf), male (ZZm), and sex-reversed female (ZZf) dragons (table S1).

We first generated global gene expression profiles and clustered these according to principal components analysis (Fig. 1B) and Spearman ranked correlation (Fig. 1C). Both methods grouped samples according to the tissue of origin, rather than sex or genotype. ZWf and ZZf ovary samples clustered separately from testis or somatic tissues but were as similar to each other as replicate samples from within

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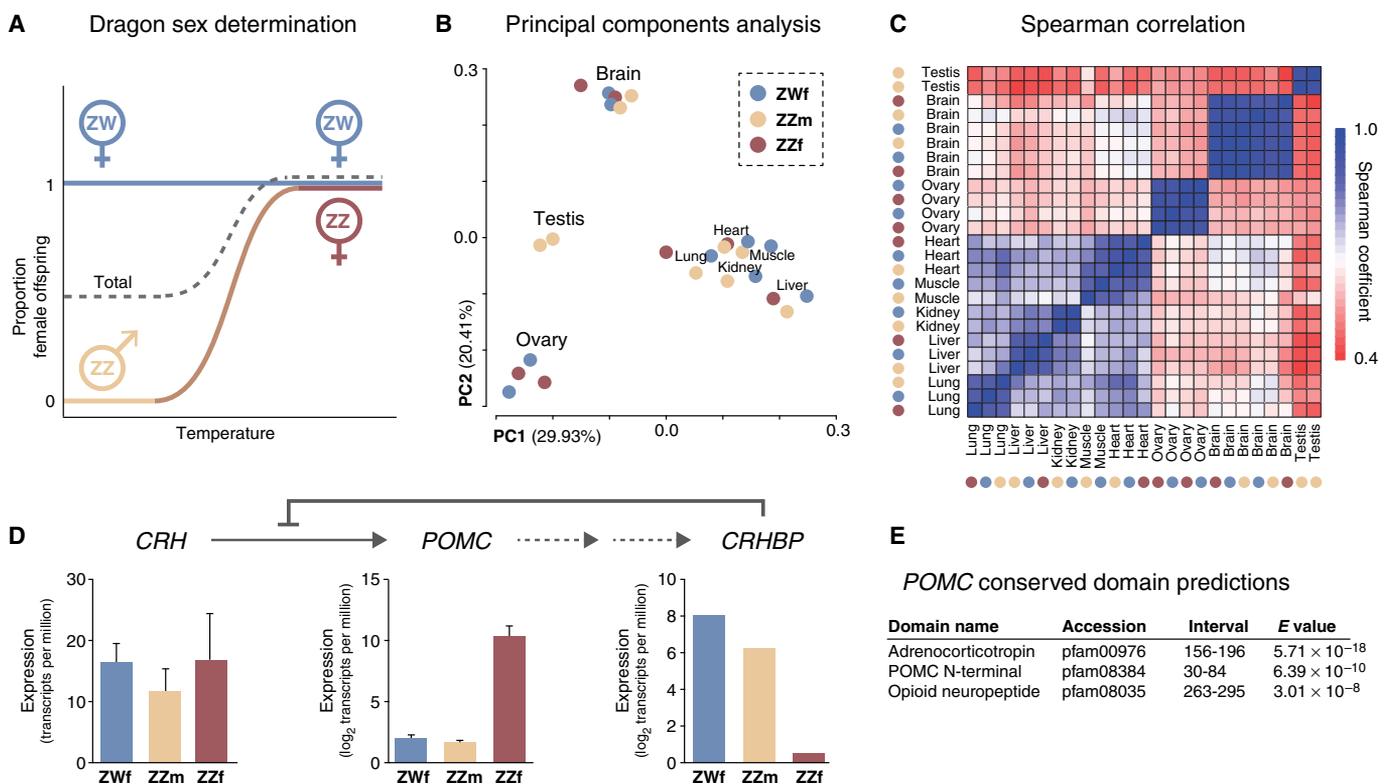


Fig. 1. Comparison of global gene expression profiles for normal and sex-reversed dragons. (A) In dragons, ZW embryos develop as female (ZWf; blue) regardless of environmental temperature. ZZ embryos develop as male (ZZm; yellow) at low temperatures and as sex-reversed females (ZZf; red) at high temperatures. Because female development can be specified independently by temperature or sex chromosome complement, it is possible to distinguish the effects of temperature and genotype on sex determination. (B) Principal components analysis and (C) Spearman ranked correlation clustering of global gene expression profiles for adult tissues from ZWf, ZZm, and ZZf individuals. (D) Top: Expression of *POMC* is activated by *CRH* (13), and *CRH* is inhibited by *CRHBP* (15). Bottom: Normalized expression (transcripts per million; mean \pm SD; $n = 2$) for *CRH* (brain), *POMC* (brain), and *CRHBP* (liver) in ZWf, ZZm, and ZZf tissues. (E) Conserved domain prediction (protein BLAST) for the expressed sequence of *POMC* in dragon.

each genotype (Fig. 1, B and C). Hence, all tissues, including ovaries, in sex-reversed females appear transcriptionally normal.

To find individual genes whose expression deviated reproducibly from these otherwise ordinary profiles, we performed differential gene expression analyses. By first comparing normal ZZm and ZWf dragons, we observed a previously unreported male bias in the expression of canonical muscle genes, including *Actin* (*ACTA1*; 44-fold) and *Myosin* (*MYH1*; 155-fold), in the brain (fig. S1, A to E). However, ZZf individuals exhibited normal female expression of these male-biased genes (fig. S1D). This indicates that, although they exhibit some “male-like” morphological and behavioral traits (12), sex-reversed females are not necessarily male-like in their molecular phenotypes.

Of greater relevance, ZZf brain displayed unique transcriptomic features that could be involved in TSD. Seventeen genes were classified as differentially expressed between ZZf and ZWf brains (fig. S2, A to D). Prominent among these were predicted orthologs for *POMC*, which is an environmental stress gene (13), and chromatin modifiers from the *Jumonji* family (14).

POMC was substantially overexpressed (327-fold) in the brain of ZZf, relative to ZWf or ZZm (Fig. 1D). *POMC* is a pituitary-expressed gene that encodes the precursor of the peptide hormone adrenocorticotropin (ACTH) (13), which is conserved in dragon ($E < 1 \times 10^{-17}$; Fig. 1E). ACTH stimulates synthesis and secretion of glucocorticoids from the adrenal cortex, including cortisol (corticosterone in birds and reptiles), which is the central effector hormone of the vertebrate

stress response (13). *POMC* expression is activated by corticotropin-releasing hormone (*CRH*) (13) and modulated by negative feedback from inhibitory *CRH*-binding protein (*CRHBP*) (15). Although there was no difference in the expression of *CRH* between ZZf and ZWf dragons, *CRHBP* expression was reduced (262-fold) in ZZf liver, the major site of its synthesis (Fig. 1D) (15). This is consistent with a reduction in negative feedback on *POMC* activation, leading to its overexpression in sex-reversed females.

Because the immune and circadian systems are known to be intertwined with stress (16), it was notable that expression levels of prominent immune genes, such as *IgM* and *IRF1*, were significantly lower in ZZf dragons than ZWf or ZZm, whereas the circadian regulator, *CIART*, was overexpressed (fig. S2D). Moreover, canonical stress-related gene ontology terms, such as “defense response” and “response to biotic stimuli,” were enriched among the top 500 genes whose expression varied between ZZf and ZWf dragons (fig. S2F).

There is considerable evidence connecting stress and sex in vertebrates. Cortisol is a regulator of natural sex change in sequential hermaphrodite fish, which change sex in response to environmental and/or social stress (17). In birds, elevated maternal corticosterone skews the sex ratio of offspring (18), as does its presence in the yolk of two Australian TSD lizard species (19). An influence of maternal ACTH levels on sex ratio has been observed in rodents (20), and there is even evidence that maternal stress biases human sex ratios (21). Thus, the strong, potentially constitutive, activation of *POMC*-mediated stress observed in sex-reversed

females may represent a mechanism by which environmental temperature interacts with sexual fate.

The most interesting gene identified in our differential expression analysis was the predicted ortholog for *JARID2*, which was expressed more highly in *ZZf* than *ZWf* tissues (fig. S2D and Fig. 2A). *JARID2* belongs to the *Jumonji* family, whose members have JmjN, JmjC histone demethylase, and AT-rich DNA binding (ARID) domains (14), all of which are conserved in dragon. *Jumonji* genes are a family of epigenetic regulators with diverse developmental roles (22).

In addition to its overexpression, the *JARID2* locus produced a unique alternative transcript in *ZZf* dragons. The 11th intron of *JARID2* was retained in nearly all transcripts from *ZZf* tissues, including the brain and gonad. In contrast, *JARID2* transcripts from *ZWf* or *ZZm* dragons were spliced normally, with no sign of retention of the 11th intron (Fig. 2, A and B, and fig. S3).

Differential intron retention (IR) has recently been established as an important mode of gene regulation in eukaryotes (23). Premature stop codons within a retained intron sequence trigger mRNA degradation, inhibiting or abolishing protein expression independently of transcription levels (23). Multiple stop codons occur within the 11th intron of *JARID2* (Fig. 3, A and B), so its retention should lead to transcript degradation. However, its high relative abundance indicates that degradation does not occur. If translated, the intron-retaining

isoforms would produce truncated products, lacking the zinc finger DNA binding domain that is encoded downstream of intron 11 (Fig. 3C). Alternatively, specific detention of intron-retaining isoforms in the nucleus might prevent translation but protect them from degradation in the cytoplasm (24, 25).

The association of *JARID2* differential IR with sex reversal is particularly interesting because *JARID2* is a component of the master epigenetic regulator Polycomb Repressive Complex 2 (PRC2) (26, 27), in which it fulfills essential roles in tissue transdifferentiation (27) and mammalian X-chromosome inactivation (28). *JARID2* appears to encourage efficient binding of PRC2 to target loci, promoting their silencing (26–28).

Wider comparisons suggest that differential IR in *JARID2/JMJD3* transcripts might be associated with TSD in all reptiles. *JARID2* was recently found to be differentially expressed in embryonic gonad development at either FPT or MPT in alligator (*Alligator mississippiensis*) (4) and turtle (*Trachemys scripta*; both are TSD species) (5). We therefore interrogated RNA sequencing data from these studies (tables S2 and S3).

At FPT in alligator bipotential gonad, we found that the orthologous *JARID2* intron was predominantly retained in mature transcripts, as we observed in sex-reversed *ZZf* dragons (Fig. 2, A and B). The same intron was predominantly spliced at MPT, similarly to normal *ZWf* or *ZZm* dragons (Fig. 2, A and B). This temperature-dependent distinction was

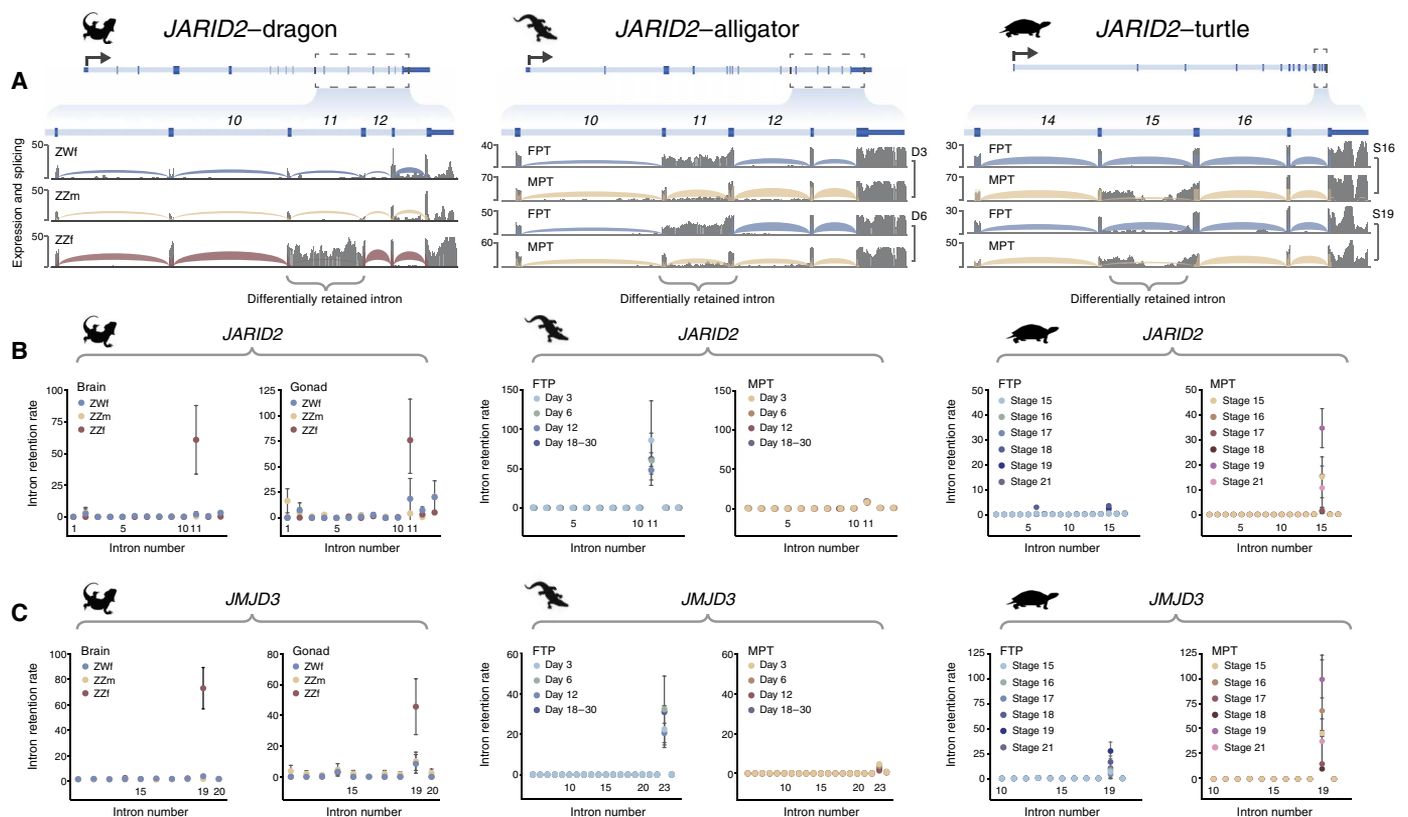


Fig. 2. Association of *JARID2/JMJD3* differential IR with temperature-dependent sex. (A) Annotated gene models for the predicted ortholog of *JARID2* in dragon (left), alligator (middle), and turtle (right). For dragon, normalized coverage by mapped RNA sequencing reads (gray) and density of spliced read junctions spanning annotated introns are shown for a single replicate of *ZWf* (blue), *ZZm* (yellow), and *ZZf* brain (red). For alligator, a single replicate of embryonic gonad after 3 and 6 days of incubation at FPT (blue) or MPT (yellow) is shown. For turtle, a single replicate of embryonic gonad at developmental stages 16 and 19 at FPT (blue) or MPT (yellow) is shown. Note that the section of zero coverage in the center of the retained intron for turtle is a string of undefined (N) bases in the genome, to which reads cannot be mapped. (B and C) Normalized rates of IR (retained transcript fragments/kilobase (kb)/spliced intron junctions; mean \pm SD) are shown for individual introns in *JARID2* (B) and *JMJD3* (C) in *ZWf*, *ZZm*, and *ZZf* dragons (brain and gonad; $n = 2$), alligator embryonic gonad (FPT and MPT; days 3, 6, 12, and 18 to 30; $n = 3$), and turtle embryonic gonad (FPT and MPT; stages 15 to 19 and 21; $n = 2$).

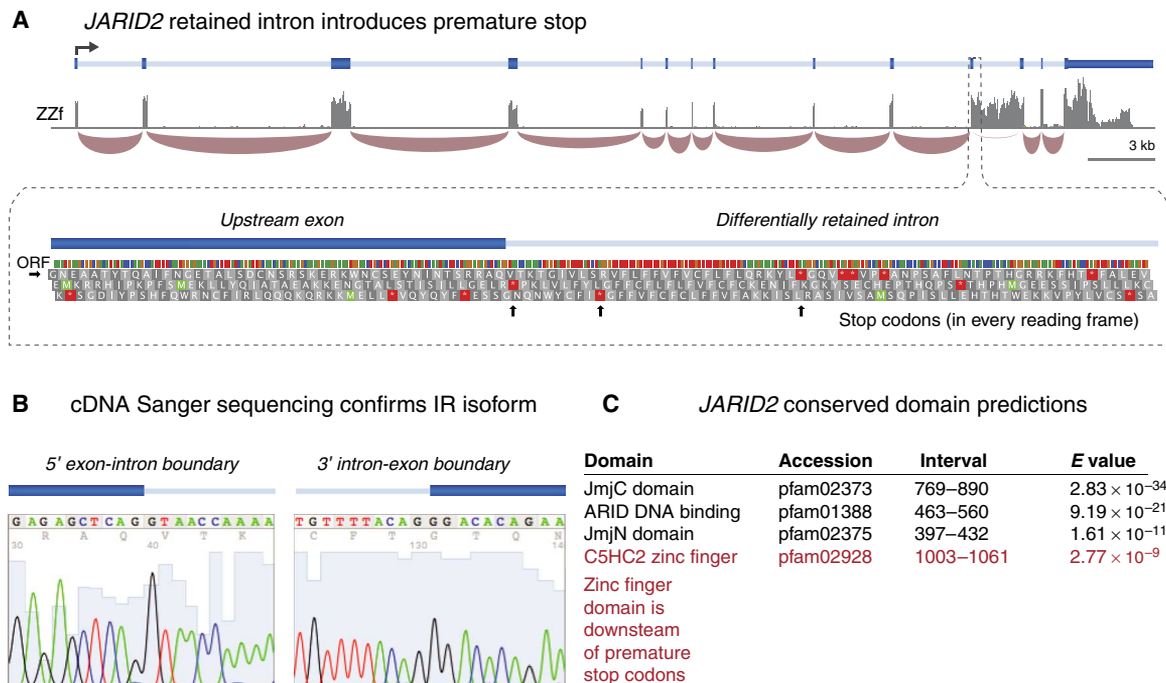


Fig. 3. *JARID2* retained intron creates premature stop. (A) Differential IR may regulate gene expression/function via the inclusion of premature stop codons within the retained intron sequence. The browser graphic shows multiple stop codons (in every reading frame) encoded within the sequence of *JARID2* intron 11, which is retained in Zzf but not ZWf or ZZm dragons. ORF, open reading frame. (B) Intron-retaining isoform was validated by Sanger sequencing on complementary DNA (cDNA) using primer pairs that spanned each exon-intron boundary. (C) Conserved domain predictions (BLASTp) for the expressed *JARID2* transcript from dragon included Mjmn, Mjmc histone demethylase, and ARID domains that characterize *Jumonji* family genes. The predicted zinc finger DNA binding domain is encoded downstream of intron 11 and will be deleted from a possible protein product.

apparent from day 3 of the experimental regime, wherein a subset of eggs was switched from FPT to MPT at day 0 (fig. S4A), and persisted throughout the entire 30-day period of comparison (Fig. 2B). *JARID2* IR was also predominant at FPT at day 0, which corresponds to Ferguson developmental stage 19 (fig. S4, B and C). A change in splicing of *JARID2* therefore occurred within 3 days of the temperature shift from FPT to MPT, roughly coinciding with the start of the temperature-sensitive period of development for alligator (29). *JARID2* was also more highly expressed at FPT than MPT throughout the period of comparison (fig. S4, B and C).

Similarly, the orthologous *JARID2* intron was differentially retained between FPT and MPT in turtle bipotential gonad. However, the directionality of the association was reversed; the intron was predominantly retained at MPT and excluded at FPT (Fig. 2, A and B). Differential IR was apparent from developmental stages 15 to 21 (Fig. 2B), encompassing the temperature-sensitive period for turtle (5). *JARID2* was also more highly expressed at turtle MPT than FPT (fig. S4, B and C), again the opposite relationship to that observed in alligator and dragon. Whole embryos sampled at developmental stage 12, before the initiation of gonad development (fig. S4A), showed no difference in *JARID2* splicing or expression between FPT and MPT (fig. S4, B and C).

Another member of the *Jumonji* family, *JMJD3* (*KDM6B*) (14), was also reported to be up-regulated at alligator FPT relative to MPT (4) and vice versa in turtle (5). *JMJD3* is an active histone demethylase and positively regulates gene expression (14). Strikingly, we found that *JMJD3*, just like *JARID2*, contained an intron that was preferentially retained at FPT in alligator and MPT in turtle (Fig. 2C). We therefore examined the predicted *JMJD3* ortholog in dragon, revealing that this gene also exhibits sex reversal-specific IR (Fig. 2C) and is more highly expressed in Zzf individuals than ZWf or ZZm (fig. S5, A to C). The introns

retained in *JARID2* and *JMJD3* are not paralogous to each other (fig. S6), implying that their capacity for differential IR does not derive from a shared sequence element.

We have therefore identified two sex reversal-specific and TSD-associated differential IR events that are expected to reduce, alter, or abolish the functionality of *JARID2* and *JMJD3* in at least three divergent reptile lineages (Fig. 4A). Although neither gene has known roles in sex determination, a closely related *Jumonji* family member, *JMJD1A*, is a direct regulator of the mammalian sex-determining factor *SRY*, and *JMJD1A* dysfunction causes male-to-female sex reversal in mice (30). Thus, we have identified two compelling candidate genes with the capacity to influence sex gene dosage and fulfill a sex-determining role.

DISCUSSION

Here, we propose a unified model for the molecular basis and evolution of TSD. We suggest that *JARID2/JMJD3* differential IR forms the mechanistic link between environmental temperature and epigenetic state underpinning sex reversal and reptile TSD.

Because retention was favored at (i) FPT in alligator (in contrast to MPT in turtle) and (ii) high temperatures in dragon (in contrast to low temperatures in alligator/turtle), *JARID2/JMJD3* IR is not simply associated with high or low incubation temperatures or with either the male or female phenotype. What occurs instead must be a complex interaction with the genetic background of each evolutionarily independent system of sex determination. As postulated previously (8, 31), we argue that some TSD species have evolved from female heterogametic sex determination systems (ZZ/ZW) and others have evolved from male heterogametic systems (XX/XY). Sex reversal can rapidly trigger the loss of

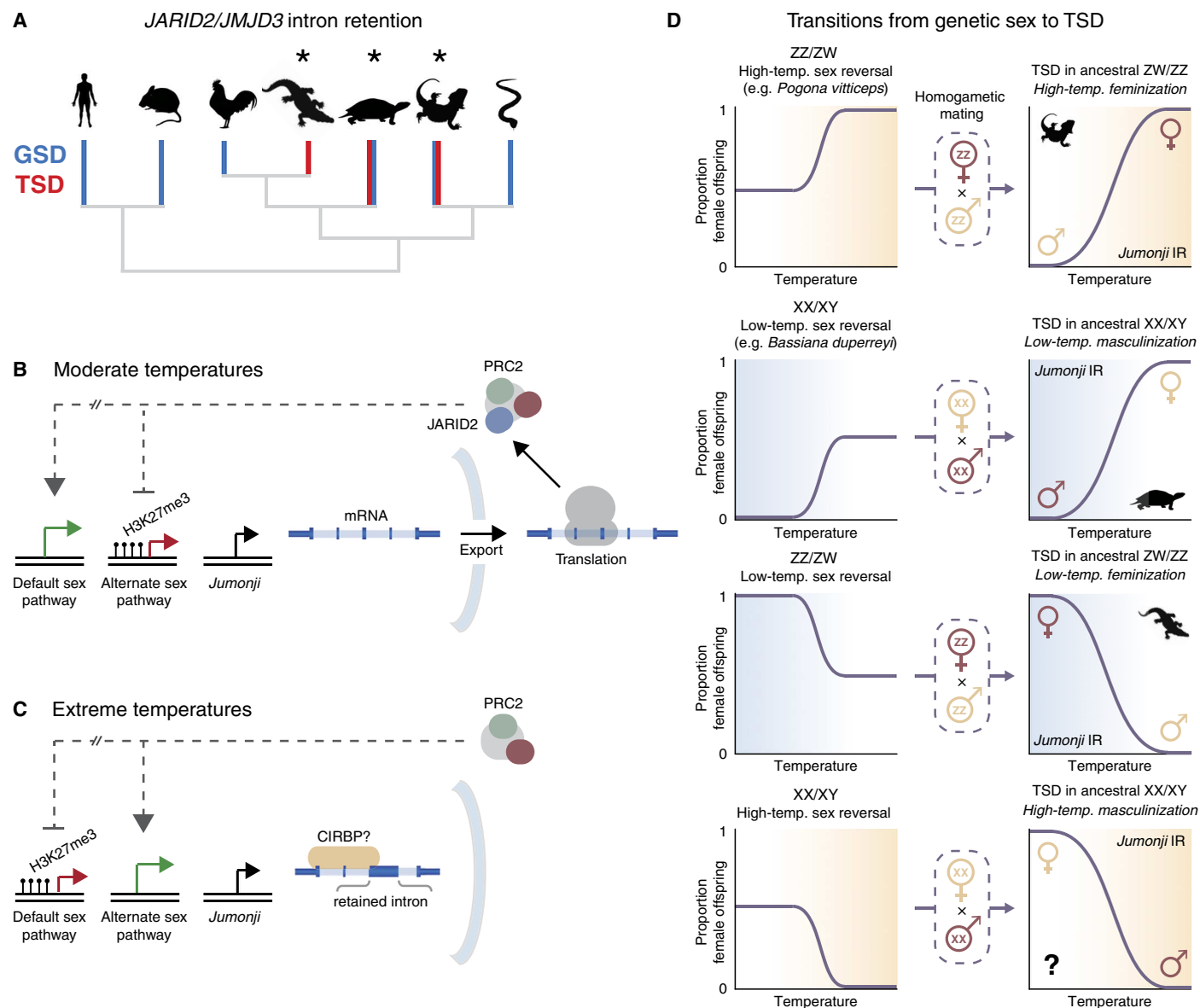


Fig. 4. Model for role of *JARID2/JMJD3* differential IR in temperature-dependent sex. (A) Phylogenetic relationships of reptile lineages, coded for genetic sex determination (GSD; blue) and/or TSD (red), and occurrence of sex-associated *JARID2/JMJD3* IR (asterisk). (B to D) The sex-determining action of *JARID2/JMJD3* IR is to divert sexual differentiation from the ancestral homogametic state (the default path) to the alternate developmental pathway. (B) At moderate temperatures, *JARID2/JMJD3* mRNA is spliced, exported, and translated. *JARID2* facilitates recruitment of PRC2 to target genes, which are repressed via H3K27 trimethylation (26–28). *JARID2* and/or *JMJD3* may also directly activate genes via histone demethylation (14). (C) At extreme temperatures, introns are retained in *JARID2/JMJD3* mRNA. IR may be induced by binding of CIRBP (6, 37) to *JARID2/JMJD3* transcripts. IR isoforms are not exported or translated (24, 25). *JARID2* is depleted from PRC2, reducing recruitment to target genes. Direct activation of genes by *JARID2/JMJD3*-catalyzed histone demethylation is also perturbed. The epigenetic landscape is thereby altered, repressing genes that maintain the default sex pathway and/or activating sex reversal genes. This can lead to feminization or masculinization, depending on the ancestral GSD system in a given species (8, 31, 32). (D) Four possible sex-associated patterns of IR may occur. Left: Ancestral GSD states (ZZ/ZW or XX/XY), with sex reversal at either high (beige) or low (blue) temperatures. Mating of sex-reversed and wild-type homogametic individuals causes transition to TSD (9, 30), with *JARID2/JMJD3* IR maintained as the regulatory signal controlling differentiation. Right: Four TSD patterns emerge: female-specific IR at high temperatures, male-specific IR at low temperatures, female-specific IR at low temperatures, and male-specific IR at high temperatures.

the heterogametic sex chromosome and, thereby, a shift from genetic sex to TSD (9). This phenomenon has been observed in wild populations for lizard species with both ZZ/ZW (*P. vitticeps*) (9) and XX/XY (*Bassiana duperreyi*) (32) systems, triggered by high and low sex-reversing temperatures, respectively.

We suggest that the sex-determining action of *JARID2/JMJD3* IR is to divert sexual differentiation from the ancestral homogametic state to

the alternate developmental path, in response to extreme temperatures (Fig. 4, B and C). This model explains the variable relationships we observed between *JARID2/JMJD3* IR, temperature, and sex among turtle, alligator, and dragon (Fig. 4D).

In our model (Fig. 4, B and C), *JARID2/JMJD3* IR is favored at extreme temperatures. *JARID2* is depleted from PRC2, reducing its recruitment to target genes. Direct activation of genes by *JARID2/JMJD3*-catalyzed

histone demethylation (33, 34) is also perturbed. Under these conditions, we propose that the epigenetic landscape is modified, disturbing developmental canalization toward the ancestral homogametic sex (the “default” state) and thereby enabling sex reversal. This can lead to feminization or masculinization of the developing gonad, depending on the ancestral homogametic sex in a given species (Fig. 4D).

How might *JARID2/JMJD3* differential IR relate to stress? Given the strong up-regulation of *POMC*-mediated stress that we observed in sex-reversed dragons, it is noteworthy that, in two other Australian TSD lizards, *B. duperreyi* and *Amphibolurus muricatus*, it is possible to interfere with sex ratio by manipulating yolk corticosterone levels (emulating maternal stress) during embryonic development (18). For these species, which have evolved from XX/XY and ZW/ZZ systems, respectively, corticosterone skewed sex ratio in opposing directions, consistent with the predictions of our model that accounts for different genetic sex backgrounds (Fig. 4D).

We therefore propose that *POMC*-mediated stress provides the temperature-sensitive signal that regulates *JMJD3/JARID2* differential IR and thereby sex reversal. Consistent with this, expression of *JARID2* and *JMJD3* is regulated by stress in mammalian systems (34, 35). Cellular stresses, including heat shock, also directly modulate the efficiency of transcript splicing and can cause general or specific IR (24, 36). It is possible that stress-activated *JARID2/JMJD3* IR, which appears to occur systemically, could have additional consequences, unrelated to sex determination, in other tissues or, alternatively, that this phenomenon simply lacks functional relevance outside the developing gonad. The response may also involve a gene previously implicated in TSD in turtles, *CIRBP* (6), which encodes a temperature-inducible RNA binding protein with imputed function in mRNA stability and translational regulation (37). It is plausible that IR could be induced by the binding of *CIRBP* to *JARID2/JMJD3* mRNA during thermal stress.

Our identification of differential IR in genes critical for epigenetic regulation therefore provides another compelling link between the environmental stress and sex determination pathways. Definitive causal evidence for our proposed model will require the development of techniques for targeted gene manipulation and/or transcriptional perturbation, such as Tol2 constructs or CRISPR (clustered regularly interspaced short palindromic repeats)–Cas9 (CRISPR-associated protein 9), which have yet to be established in reptiles. Nevertheless, our observations open up new directions for investigations into the molecular mechanisms by which genes and environment interact to control reptile sex determination.

MATERIALS AND METHODS

Bearded dragon RNA sequencing libraries

We used paired-end RNA sequencing libraries from *P. vitticeps*, as published previously (38). Polyadenylation-selected RNA libraries were derived from adult male (ZZm), female (ZWf), and sex-reversed female (ZZf) dragon tissues, including gonad ($n = 2$), brain ($n = 2$), and additional somatic tissues (liver, kidney, lung, heart, and muscle; $n = 1$). Raw sequencing reads were trimmed using TrimGalore with default parameters. After trimming and quality control (QC), a total of 386 million read pairs were retrieved (table S1). Raw libraries can be accessed under the European Nucleotide Archive study accession number PRJEB5206.

Alligator RNA sequencing libraries

We used paired-end RNA sequencing libraries from *A. mississippiensis*, as published previously (4). Total RNA libraries were derived from al-

ligator embryonic gonad developing at either FPT (30°C) or MPT (33.5°C) over a 30-day time course ($n = 3$) (4). Raw sequencing reads were trimmed using TrimGalore with default parameters. After trimming and QC, a total of 324 million read pairs were retrieved (table S2). Raw libraries can be accessed using the DRA (DNA Databank of Japan Sequence Read Archive) accession number DRA004128-41.

Turtle RNA sequencing libraries

We used paired-end RNA sequencing libraries from *T. scripta*, as published previously (5). Total RNA libraries were derived from turtle embryonic gonad developing at either FPT (31°C) or MPT (26°C) at developmental stages 15 to 19 and 21 ($n = 2$) (5). Raw sequencing reads were trimmed using TrimGalore with default parameters. After trimming and QC, a total of 697 million sequenced read pairs were retrieved (table S3). Raw libraries can be accessed at the Sequence Read Archive under BioProject PRJNA331105.

Gene expression profiling

We used the recently assembled *P. vitticeps* genome (38) and accompanying gene annotation (<http://gigadb.org/dataset/100166>) to perform gene expression profiling, with expression values derived using Kallisto (39) alignment-free quantification over annotated coding sequences. Kallisto was used with a fixed k -mer length of 30 nucleotides. We excluded genes from downstream analyses that did not have a minimum measured abundance of ≥ 3 transcripts per million in multiple replicates from at least one tissue. A total of 17,818 genes fulfilled these criteria.

Comparisons of global expression profiles were performed in R. Spearman correlation matrices were generated with the `corrplot` package (<https://cran.r-project.org/web/packages/corrplot/>) and principal components analysis with the `EDASeq` package from Bioconductor (<http://bioconductor.org/>).

Differential gene expression analysis

The `edgeR` package from Bioconductor (<http://bioconductor.org/>) was used for differential gene expression analysis. A threshold of false discovery rate (FDR) < 0.01 was required for candidacy as a differentially expressed gene.

To verify that these candidates were robust to bioinformatics variables, we repeated gene quantification using a traditional alignment-based approach (as opposed to Kallisto alignment-free quantification; see above). We used a STAR-RSEM (40, 41) workflow with the following STAR-mapping parameters: `-outSAMstrandField intronMotif`, `-outFilterIntronMotifs RemoveNoncanonicalUnannotated`, `-outReadsUnmapped None`, `-alignMatesGapMax 200000`, and `alignIntronMax 5000000`. We also removed reads with a mapping quality score < 10 , as these represent potential multi-mappers.

Only genes that were concordant in their differential expression (FDR < 0.01) between both independent methods of quantification (Kallisto/STAR-RSEM) were reported as differentially expressed.

Intron retention

Normalized IR rates were assessed for all introns within *JARID2* and *JMJD3* using custom scripts. Our approach was analogous to that reported by Braunschweig *et al.* (42). Specifically, for a given intron in a given sample, the rate of retention was calculated as follows. (i) All split-mapped reads spanning the intron, with ends mapping exactly to its annotated splice junctions, were retrieved (`spliced_junctions`). (ii) Unspliced read fragments mapping within or overlapping (by at least 5 base pairs) the intron sequence were retrieved (`retained_fragments`).

(iii) The IR rate was calculated as retained_fragments/intron_length (kb)/spliced_junctions. Retention rates were averaged between replicate samples and are reported as mean \pm SD.

For alligator (*A. mississippiensis*) and turtle (*T. scripta*), IR was assessed in the same manner as described above. However, because no reference genome was available for *T. scripta*, we used the closely related *Chrysemys picta* draft genome and accompanying gene annotation (43).

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at <http://advances.sciencemag.org/cgi/content/full/3/6/e1700731/DC1>

- fig. S1. Comparison of gene expression in male and female dragons.
 fig. S2. Comparison of gene expression in normal and sex-reversed female dragons.
 fig. S3. Differential *JARID2* IR in normal and sex-reversed dragons.
 fig. S4. Temporal dynamics of *JARID2/JMJD3* expression and splicing in alligator and turtle embryo.
 fig. S5. Expression and splicing of *JMJD3* in the brain and gonad from normal and sex-reversed dragons.
 fig. S6. Differentially retained introns in *JARID2/JMJD3* are nonparalogous.
 table S1. RNA sequencing libraries for *P. vitticeps*.
 table S2. RNA sequencing libraries for *A. mississippiensis*.
 table S3. RNA sequencing libraries for *T. scripta*.

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the evolutionary model. I.W.D. prepared the manuscript with contributions from C.E.H., J.B., J.A.M.G., P.D.W., J.S.M., and A.G. **Competing interests:** The authors declare that they have no competing interests. **Data and materials availability:** All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from the authors. All data have been published previously and are publicly available (tables S1 to S3).

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