Evolving thermal thresholds explain the distribution of temperature sex reversal in an Australian dragon lizard

Meghan A. Castelli1,2 | Arthur Georges1 | Caitlin Cherryh2,3 | Dan F. Rosauer3 | Stephen D. Sarre1 | Isabella Contador-Kelsall4 | Clare E. Holleley1,2

Abstract

Aim: Species with temperature-dependent sex determination (TSD) are particularly vulnerable to climate change because a resultant skew in population sex ratio can have severe demographic consequences and increase vulnerability to local extinction. The Australian central bearded dragon (Pogona vitticeps) has a thermosensitive ZZ male/ZW female system of genetic sex determination (GSD). High incubation temperatures cause reversal of the ZZ genotype to a viable female phenotype. Nest temperatures in the wild are predicted to vary on a scale likely to produce heterogeneity in the occurrence of sex reversal, and so we predict that sex reversal will correlate positively with inferred incubation conditions.

Location: Mainland Australia.

Methods: Wild-caught specimens of P. vitticeps vouchered in museum collections and collected during targeted field trips were genotypically and phenotypically sexed to determine the distribution of sex reversal across the species range. To determine whether environmental conditions or genetic structure can explain this distribution, we infer the incubation conditions experienced by each individual and apply a multi-model inference approach to determine which conditions associate with sex reversal. Further, we conduct reduced representation sequencing on a subset of specimens to characterize the population structure of this broadly distributed species.

Results: Here we show that sex reversal in this widespread Australian dragon lizard is spatially restricted to the eastern part of the species range. Neither climatic variables during the inferred incubation period nor geographic population genetic structure explain this disjunct distribution of sex reversal. The main source of genetic variation arose from isolation by distance across the species range.

Main conclusions: We propose that local genetic adaptation in the temperature threshold for sex reversal can counteract the sex-reversing influence of high incubation temperatures in P. vitticeps. Our study demonstrates that complex evolutionary processes need to be incorporated into modelling biological responses to future climate scenarios.
1 | INTRODUCTION

In most vertebrates, the outcome of sexual differentiation is binary, with individuals developing phenotypically as male or female following distinct developmental trajectories. Although the outcomes of sexual differentiation are highly conserved among vertebrates, great diversity exists in the genetic and environmental factors that have acquired master or influential roles in the determination of sexual fate (Bachtrog et al., 2014). These signals can be genetic or environmental, but rather than a dichotomy, a continuum occurs from genetic sex determination (GSD) to environmental sex determination (ESD) in that the critical influence can be genetic, environmental or an interaction between the two (Sarre et al., 2004).

Transitions between genetic and environmental sex determination systems have occurred frequently among vertebrates on an evolutionary timescale (Bachtrog et al., 2014; Van Doorn, 2014; Sarre et al., 2011; Warner, 2011), and underlying these transitions is evolution in the threshold temperature for sex reversal (Quinn et al., 2011). The intrinsic and extrinsic factors which are capable of driving evolution towards one or the other extreme of the continuum have been widely modelled computationally and demonstrated experimentally (Sarre et al., 2004). Temperature sex determination (TSD) is the most common mode of ESD and is observed widely among fish, reptiles and amphibians (Bachtrog et al., 2014). Mechanistic models under different climatic scenarios have identified the importance of factors other than temperature alone in generating primary sex ratios, including maternal behaviour, nesting phenology, fecundity, and threshold temperature (Boyle et al., 2014; Mitchell et al., 2010; Schwanz et al., 2020). Species with TSD are particularly vulnerable to climate change because skews in the population sex ratio in response to extreme conditions can have severe demographic consequences, increasing vulnerability to local extinction (Mitchell & Janzen, 2010). Significantly skewed sex ratios caused by climate warming already occurs in contemporary populations of TSD reptiles (Jensen et al., 2018) and fish (Honeycutt et al., 2019). The tipping-point at which skewed sex ratios lead to demographic collapse is not yet known.

Species with GSD are typically assumed to be resilient to sex ratio skew (Bókony et al., 2019). The occurrence of sex reversal (temperature overriding an underlying GSD system) challenges this view. Sex reversal occurs in wild populations of two Australian reptiles: the central bearded dragon (Pogona vitticeps; Holleley et al., 2015), the eastern three-lined skink (Bassiana superrei; Holleley et al., 2016), and is suspected in six more species of lizard and turtle (Holleley et al., 2016; Wiggins et al., 2020). Sex reversal is also a frequent phenomenon in fish (Baroiller & D’Cotta, 2016). Sex reversal provides a mechanism by which increasingly extreme conditions lead to the loss of the heterogametic chromosome (the Y or the W) and a transition from GSD to TSD (Holleley et al., 2015; Schwanz et al., 2020). Persistence of the population then requires one of two things: behavioural modifications that alter the nest environment, or evolution in the temperature threshold for sex reversal (Düsing, 1884; Edwards, 2000; Fisher, 1930). Though maternal behaviours can buffer against the effects of climate, this is not universally true and these behaviours are not always inheritable (Doody et al., 2006; Ewert et al., 2005; Gutzke & Paukstis, 1983; Mitchell & Janzen, 2019;Refsnider et al., 2013; Refsnider & Janzen, 2016; Warner & Shine, 2008). Additionally, evolutionary responses take time and require existing inheritable variation upon which to act (Schwanz et al., 2020). The time lag implicit in these evolutionary forces can leave thermosensitive species (including those with GSD and sex reversal) extremely vulnerable to local extinction under rapid or extreme climate change via demographic skews.

The central bearded dragon (P. vitticeps) displays a sex determination system characterized by a ZZ/ZW GSD system of female heterogamety (Ezaz et al., 2005) and sex reversal of the ZZ male genotype to phenotypically female under the influence of high temperature (Holleley et al., 2015; Quinn et al., 2007). Sex reversal occurs rarely at 31°C but rises in frequency with temperature until 36°C where almost 100% of ZZ genotype animals are feminized (Holleley et al., 2015). This Australian agamid is distributed across a range of climatic types spanning mesic to xeric climatic regions across central, south-eastern and northeastern Australia (Cogger, 2018; Rej & Joyer, 2018), providing a unique opportunity to study the dynamics of genetic and environmental influences on sex determination. Some populations of P. vitticeps may be in the early stages of transition from GSD to TSD via loss of the W chromosome (Holleley et al., 2015).

Here, we examine the distribution of sex reversal across the geographic range of P. vitticeps, using historical museum specimens and contemporary wild-caught individuals. Nest temperatures are predicted to vary across this range (Figure 1). We predict that sex reversal frequency in the wild will be positively correlated with inferred incubation temperatures if there is a common and static threshold for temperature sex reversal. Conversely, if there is local adaptation in the threshold for sex reversal, we predict that the frequency of sex reversal will be independent of incubation temperature across the species range.

2 | METHODS

2.1 | Specimen collection and phenotypic sex identification

A total of 534 wild-caught dragons, collected from field trips (n = 337 dragons) and supplemented by vouchered museum specimens (n = 197) collected over a period of 38 years from 1980 to 2018, were
sampled in this study (File S1). Tissues from dragons collected during field trips were taken as either tail or toe clips, frozen blood or dried blood on DNA-preserving card storage systems (Whatman™ FTA™ Elute Cards, Macherey-Nagel Nucleocards, or PerkinElmer 226 Five Spot cards). Liver samples from museum specimens were sampled by museum staff at the time of collection, and either stored in ethanol at room temperature or frozen (−80°C). Specimens that were wild-caught and released in the field were phenotypically sexed by hemipenial eversion. In this approach, the hemipenes of adult males are reliably evorted by running the thumb up the ventro-lateral surface of the base of the tail, while gently bending the tail to expose the cloacal aperture. Sex identification was confirmed secondarily by examining the animal for gravidity, observing nesting behaviour, and examining male secondary sexual characteristics, such as larger head size and dimorphism in femoral pores. The sex of museum specimen and road-killed animals was identified by dissection of the abdomen and inspection of the gonads. The few juvenile or hatchling animals captured or located during the study were only included if gonadal sex could be determined by dissection.

2.2 Molecular sex identification

Genotypic sex (ZZ/ZW) was determined by extracting DNA from tissue and conducting a PCR sex test (Quinn et al., 2010, modified by Holley et al., 2015). Three DNA extraction methods were performed as appropriate for each tissue type: (a) Qiagen’s Gentra® Puregene® DNA purification kit for tail and toe clips, frozen liver, frozen blood, (b) Macherey-Nagel Nucleospin® Tissue kit for blood on frozen PerkinElmer 226 cards and Macherey-Nagel Nucleocards; (c) Whatman Elute quick extraction protocol for blood on Whatman™ FTA™ Elute Cards. Manufacturer’s instructions were followed in all cases. Purified DNA was quantified using a NanoDrop 1,000 spectrophotometer (Thermo Scientific) and standardized to a concentration of 25 ng/µl prior to PCR. DNA extraction and PCR setup were automated using the epMotion 5,075 platform (Eppendorf, Germany), and included both positive and negative controls following Whiteley et al., (2017). PCR products were visualized on a 1.5% agarose gel. ZZ genotypic individuals displayed one band at 524 bp (both Z fragments) and ZW genotypic individuals displayed two bands, one at 524 bp (Z fragment) and the other at 357bp (W fragment). Internal ZZ and ZW control samples were run on every gel. Individuals with a ZZ genotype and a female phenotype were classified as sex-reversed. Chi-square tests were used to determine if there were sex biases in this dataset between males and females across the entire period of the study. Additionally, chi-square tests were conducted to determine if the proportion of concordant females and sex-reversed females varied across the years in which at least 15 phenotypic females were collected.

2.3 Prediction generation: constant temperature equivalent

The rate of embryonic development of P. vitticeps is dependent on temperatures experienced by the egg within the nest, with embryos developing faster at hotter temperatures (Whiteley et al., 2017). For embryos developing at higher temperatures, this means that
the period during which sex can be determined by temperature is shorter. Constant temperature equivalent (CTE) calculation allows the fluctuations in temperature experienced by a typical buried egg to be converted to a constant temperature which results in the same amount of embryonic development. First, the daily soil temperature was calculated over a single laying season (September to December in year \( y \) and January to February in year \( y + 1 \)) using NicheMapR (M. Kearney, 2020; M. R. Kearney & Porter, 2017) and user-supplied weather input data (SILO climate gridded dataset). Secondly, the daily CTE was calculated using methods described in Georges (1989) and Georges et al. (1994). To generate range-wide long-term trends in CTE, the procedure was repeated over forty consecutive breeding seasons from 1975–2014, and for 1,000 locations randomly selected from within the range of \( P. \) vitticeps.

The microclimate model was driven by the Scientific Information for Land Owners (SILO) climate gridded dataset (Jeffrey et al., 2001). For all the 1,000 randomly selected points, we extracted minimum and maximum temperature, relative humidity at time of minimum and maximum temperature, rainfall, vapour pressure, and solar radiation (used to estimate cloud cover, as outlined in NicheMapR documentation). In addition, we used a daily gridded wind speed dataset at both the minimum and maximum wind temperature (T. McVicar, 2011; T. R. McVicar et al., 2008). Elevation was extracted from the GEODATA 9 s Digital Elevation Model (DEM-9S) Version 3 (Hutchinson, Stein, Stein, Anderson, & Tickle, 2008). For each location, soil properties were extracted from the Soil and Landscape Grid of Australia (Grundy et al., 2015; Viscarra Rossel et al., 2015; Viscarra Rossel et al., 2014a, 2014b, 2014c, 2014d). The proportion of clay, silt and sand at each location was used to determine the soil parameters based on table 9.1 from Campbell and Norman (1998), included in NicheMapR (M. Kearney, 2020). We used a custom R program (R Core Team, 2020) including the packages \texttt{sp} (Bivand et al., 2013; Pebesma & Bivand, 2005), \texttt{RFast} (Papadakis et al., 2020), \texttt{zoo} (Zeileis & Grothendieck, 2005), \texttt{timezone} (Rundel, 2013) and \texttt{raster} (Hijmans, 2020) to collate climate variables into NicheMapR, run the microclimate model and extract the hourly soil temperature output for 10 soil depths between 0 and 200 cm. We assumed a nest depth of 15 cm (Pianka, 2005), and that nests were unshaded.

To calculate the daily CTE, we used the method described in Georges (1989) and Georges et al. (1994). The model assumes that daily soil temperature varies on a cycle about a stationary mean for one day, \( T \) is the temperature at time \( t \) in degrees Celsius, \( M \) is the mean soil temperature, and \( R \) is the maximum deviation from \( M \). \( T_0 \) is the minimum temperature at which embryonic development is possible. In \( P. \) vitticeps, \( T_0 = 16.2^\circ \text{C} \) (unpublished data). The CTE (\( T' \)) occurs at time \( t' \). Equation 2 is solved iteratively for \( t' \) until \( t'_\text{old} - t'_\text{new} < 0.0001 \). If this did not occur within 10,000 iterations, we used the value of \( t' \) that occurred when the difference between iterations was smallest. The value for \( t' \) is then substituted into equation 1 and solved for \( \text{CTE} \).

\[
T = R \cos(t) + M, \quad 0 \leq R \leq M, \quad (1)
\]

\[
t' = \frac{\pi}{2} - \frac{R}{M - T_0} \sin(t') \quad \text{for } \frac{R}{M - T_0} \leq 1. \quad (2)
\]

If soil temperature dropped below \( T_0 \), development ceased for part of that day. In that case, \( t_0 \) was calculated using equation 3 to estimate the time at which soil temperature drops to \( T_0 \) and development ceases. Then, \( t' \) was calculated iteratively using equation 4 until \( t'_\text{old} - t'_\text{new} < 0.0001 \). If this did not occur within 10,000 iterations, we used the value of \( t' \) that occurred when the difference between iterations was smallest. Finally, the CTE was calculated by substituting \( t' \) into equation 1. For further details on the derivation of these equations, please refer to Georges (1989).

\[
t_0 = \cos^{-1}\left(\frac{T_0 - M}{R}\right), \quad 0 < t_0 \leq \pi \quad \text{for } \frac{R}{M - T_0} \leq 1. \quad (3)
\]

\[
t' = t_0 + \frac{R}{2(M - T_0)} \sin(t_0) - \frac{R}{M - T_0} \sin(t'). \quad (4)
\]

For some days, there was little variation in soil temperature and \( M = T_0 \) or \( M = T_0 \) and \( R = 0 \). On these days, the CTE could not be calculated using the method described in Georges (1989) and Georges et al. (1994). When this occurred, we set \( \text{CTE} = T' = M \). We calculated the daily CTE for each day in September to February. The mean CTE for a breeding season is the mean of all daily CTEs from September to February \( (n = 181 \text{ days}) \) for that random location.

The high degree of correlation (Pearson’s Correlation; \( R = 0.890, t = 61.739, df = 998, p < .001 \)) between the average maximum temperature for each location from 1975 to 2014 and the corresponding CTE estimates (derived in part from maximum temperature) led us to use only the extracted SILO data for estimation of inferred incubation conditions for each collected specimen rather than the more derived CTE values.

### 2.4 Geographic correlates of sex reversal

Multinomial logistic regression (R packages: \texttt{nnet} and \texttt{tidy}) was used to determine whether sex-reversed individuals displayed different geographical distributions when compared to the concordant sexes (where genetic sex matches phenotypic sex). A multinomial logistic regression was constructed using latitude, longitude and elevation as the predictor variables, with the sex of each specimen as the response variable, either concordant male (ZZm), concordant female (ZWf) or sex-reversed female (ZZf). The three categorical outcomes were then compared on a pairwise basis, to determine if there was an association between latitude, longitude, or elevation and sex.

Cluster analysis software \texttt{SaTScan v9.6} (2018) was used to identify if sex reversal was clustered in the landscape. A Bernoulli probability model and moving window analysis was applied to identify significant clustering in the spatial distribution of sex-reversed compared to all concordant individuals. The maximum spatial cluster size
was set at 50% of the size of the population, a generally accepted standard (Kim & Jung, 2017), with high abundance clusters defined as containing at least ten instances of sex reversal.

2.5 | Environmental correlates of sex reversal: inferred incubation conditions

Inferred incubation conditions (Table 1) were determined for each individual using its estimated age, location of capture and parameters of the reproductive cycle of *P. vitticeps*. Temperature variables (maximum temperature and diel range) were selected for their direct relationship to sex reversal (Holleley et al., 2015) and rainfall for its effects on temperature. The deviation of these conditions during the inferred incubation period from the long-term average for each site was also included in analysis, to account for the possibility that it may not be absolute temperature which causes sex reversal, but rather deviation from long-term conditions, to which populations in different areas may have become adapted. Age was estimated qualitatively, and by using snout-vent length (SVL). Adult individuals were assumed to be between 3–5 years old, juveniles between 1 and 2 years old and hatchlings less than 6 months old. The breeding season (and therefore time during which incubation conditions could reasonably be experienced by each individual) was determined to be from September-February inclusive based on patterns of gravidity in dissected wild female specimens and patterns of egg laying in a breeding colony (Figure S1). For adults, inferred incubation conditions during the third, fourth and fifth breeding season prior to collection were averaged. For juveniles, inferred incubation conditions during the first and second breeding season prior to collection were averaged. Hatchlings were excluded from analysis.

Environmental data during the inferred incubation period of each individual were extracted from the Scientific Information for Land Owners (SILO) database of Australian climate data, in gridded format (~25 km²). We needed to estimate the home range size to ensure that we could reasonably assume that each specimen was incubated within the grid in which it was collected. There are no published data on the home range and post-hatching dispersal of *P. vitticeps*, but the data available in other Australian terrestrial reptiles indicates that home ranges are likely to be less than 25 km² (Piza-Roca et al., 2018; Shine & Lambeck, 1989). Thus, we have extracted and analysed inferred incubation conditions from only the SILO grid which contains the collection location of the specimen, assuming that this grid also contained the location in which the individual developed in the egg. All data analyses were conducted in R v.4.0.2 (R Core Team, 2020), and significance was set at *p* = .05 for all tests.

To identify environmental variables associated with the occurrence of sex reversal, we used multiple model inference analysis [R package MuMIn v.1.42.14: (Barton, 2019)]. A binomial logistic regression model containing all of the model predictors (Table 1) was generated, and the MuMIn package then used to generate models with all possible combinations of the predictor variables. Akaike information criteria (AICc, corrected for small sample sizes) values assigned to each model were then used to determine the model/s which best explained variation in the response variable. Both minimum and maximum air temperature were considered for inclusion in analysis, but owing to the high degree of correlation displayed between these two variables and the more direct link between high temperature and sex reversal, we decided to include only maximum daily temperature in analysis. The inferred incubation conditions of each ZZ individual (excluding hatchlings) were entered into a single binomial logistic regression model and MuMIn used to perform model averaging and to determine which variables best explain the binomial regression model.

| Variable | Description |
|----------|-------------|
| Average daily maximum temperature during incubation | Average maximum temperature (°C) for days during the inferred incubation period |
| Average daily rainfall during incubation | Average daily rainfall (mm) for days during the inferred incubation period |
| Average diel temperature range during incubation | Average diel temperature range (°C) for days during the inferred incubation period |
| Deviation from long-term average daily maximum | Average deviation of the *daily maximum temperature* from the long-term mean of daily maximums during the breeding season for that site from 1910 until 5 years before specimen collection |
| Deviation from long-term average rainfall | Average deviation of the *daily rainfall* from the long-term mean of daily rainfall during the breeding season for that site from 1910 until 5 years before specimen collection |
| Deviation from long-term average diel temperature | Average deviation of the *diel temperature range* from the long-term mean of diel range during the breeding season for that site from 1910 until 5 years before specimen collection |
To determine whether population structure across the species geographic range may explain patterns in sex reversal, reduced representation sequencing was conducted on a sample of 218 individuals (13 ZZf, 73 ZWf and 132 ZZm; Figure S2). Each individual was genotyped by reduced representation Illumina short-read sequencing using commercial provider Diversity Arrays Technology (DArT Pty Ltd, Canberra, ACT, Australia). Briefly, SNP genotyping was performed using a combination of complexity reduction by using two restriction enzymes, implicit fragment size selection and next-generation sequencing (Kilian et al., 2012). A pair of restriction enzymes PstI (recognition sequence 5’-CTGCA∥G-3’) and SphI (5’-GCATG∥C-3’) was used for complexity reduction by double digestion. Sequences were processed using proprietary DArT analytical pipelines (Kilian et al., 2012) to yield SNP markers polymorphic within the set of samples. Calling quality was assured by high average read depth per locus (medium coverage, 10X). In addition, approximately one-third of the samples were processed twice from DNA to allelic calls as technical replicates. Scoring consistency (repeatability) was used as the main selection criteria for high-quality and low error-rate markers. Refer to Georges et al. (2018) for further detail. Additional filtering of loci and individuals and preliminary analysis was undertaken using the R package dartR (v.1.1.11; Georges et al., 2018). Loci with a call rate of less than 95%, repeatability of less than 99%, monomorphic loci and secondary SNPs were removed prior to analysis. Individuals with a call rate of less than 75% (n = 2) were also removed from analysis. Structure across the landscape was visualized using principal coordinates analysis (PCoA) in the dartR package, and isolation by distance was assessed using Mantel’s test in the R package adegenet (Jombart, 2008).

### 3 RESULTS

#### 3.1 Geographic distribution of sex reversal

Among the 534 wild-caught individuals sampled across the species range, 294 (55%) were ZZ males, and 240 (45%) were phenotypic females. Of those females, 28 (12%) possessed a ZZ genotype and were therefore sex-reversed. The bias towards males in our data was significant ($\chi^2 = 5.46, df = 1, p = .019$) possibly due to capture-bias and the higher visibility of males which display prominently. Sex-reversed females were collected in 12 non-consecutive years of the 38 years sampled. The earliest case of sex reversal was in South Australia in 1983 and the most recent in New South Wales in 2018 (Figure 2). There was no range-wide temporal trend in sex reversal. The proportion of females which were sex-reversed varied from 6% to 27% but without significant difference between years ($\chi^2 = 6.031, df = 5, p = .303$) among the six years in which at least 15 phenotypic females were collected.

Sex reversal was detected across a substantial proportion of the range of *P. vitticeps* but was not randomly distributed (Figure 2). Spatial cluster analysis using SaTScan identified a single large cluster in the south-eastern part of the sampling area where sex reversal occurred at a rate higher than would be expected if sex-reversed individuals were distributed randomly throughout the study range. The radius of the cluster was 410.83 km, containing 23 out of the 28 cases of sex reversal (Expected cases: 12.9, Log-Likelihood Ratio: 8.182, $p = 0.039$). We found that the latitudinal, longitudinal and elevational distributions of concordant males (ZZm) and concordant females (ZWf) did not differ (Table 2). Sex-reversed individuals (ZZf) were found not to differ in terms of latitudinal or elevational distribution compared to both ZZm and ZZf.
ZWf, but were significantly different in their longitudinal distributions, being largely absent from lower longitudes (the western part of the species range; Table 2; Figure 2).

In a previous report detailing the extent of wild sex reversal (Holleley et al., 2015), the area examined spanned a minimum convex polygon (MCP) of ~260,000 km² (12.2% of the known species range), and sex-reversed females were found to occupy an MCP of ~26,000 km² (9.9% of the sampled range). Here we report a substantial increase in the sampling area, with an MCP of ~1.7 million km² (79.5% of the species range) sampled, of which sex-reversed females were found in a ~400,000 km² MCP representing approximately 24.2% of the sampled range.

3.2 | Environmental correlates of sex reversal

Overall, the binomial logistic regression comparing ZZ males and ZZ females using all six inferred incubation condition predictor variables had exceptionally poor predictive ability ($R^2 = 0.032$) (Table 3). An unbiased model inference approach using MuMIn extracted nine “best” models to explain the binomial logistic regression. These models differed by less than two AICc values from the best model with the lowest AICc value and are therefore indistinguishable in terms of their explanatory power (Burnham & Anderson, 2002). Five of these were single-variable models, three contained two variables and one was the null model. That the null model was also included indicates that no combination of inferred incubation conditions was able to explain the binomial regression model of the frequency of sex reversal (ratio of ZZ females to ZZ males) more parsimoniously than the null model. This suggests that of the inferred incubation conditions included in the model, none are capable of adequately explaining the occurrence of sex reversal.

3.3 | Population structure

Pre-filtering, this dataset contained 197,890 SNP loci for 218 individuals with a missing data rate of 34.29%. After filtering and quality control, the dataset was reduced to 11,128 SNPs for 216 individuals with a missing data rate of 2.1%.
We were unable to detect any broadscale geographic population structure. In a PCoA analysis, the first and second axes explain only 2.4% and 1.3% of variation respectively (Figure 3), indicating that despite the large sampling area, there is limited population differentiation. A relatedness matrix constructed using dartR identified a higher degree of relatedness in all specimens collected within a geographically isolated area south of the Murray River in South Australia. In the PCoA analysis, these individuals do cluster and appear to be genetically divergent from the rest of the sampled specimens, but low variation explained by the PCoA indicates that this divergence is minor. There was significant evidence of weak isolation by distance (Wright, 1943) across the entire range of the species (Mantel’s test; $r = .571, p = .001$, Figure 3) suggesting a stepping-stone model of dispersal for this species (Kimura, 1953).

4 | DISCUSSION

Here, we present the most detailed in situ account of naturally occurring sex reversal in a terrestrial vertebrate. Over a 38-year period encompassing a study area of 1.7 million km$^2$ (almost 80% of the species range), we show that sex reversal in *P. vitticeps* occurred across a quarter of the species range but was not directly explained by inferred incubation conditions. Sex-reversed ZZ females comprise 12% of all phenotypic females collected, demonstrating that the sex-reversed phenotype is a substantial demographic in this species.

Counter to our predictions based on the established relationship between high temperature and sex reversal in a laboratory setting (Holleley et al., 2015; Quinn et al., 2007), we did not observe an association between environmental temperatures and sex in the wild. There is a noticeable lack of sex reversal in the northern and north-western part of the species range where nest CTE and ambient temperatures are the hottest (Figure 1). Instead, we observed a significant cluster of sex reversal in the south-eastern portion of the species range, where inferred incubation conditions are relatively cooler than in the north, and where long-term CTE’s do not approach the 32°C sex reversal threshold. It must be noted that while we cannot conclude definitively that sex reversal is absent from the north of the species range, owing to lower sample size in this area, our clustering methods do account statistically for sampling heterogeneity and still identified a significant cluster. The occurrence of sex reversal in wild populations is clearly more complex than the simple relationship with temperature observed under controlled laboratory conditions, even after correcting for diel variation in temperature in natural nests (Georges, 1989; Georges et al., 1994), and could be explained by differences in the thermal threshold for sex reversal.

We argue that the geographically disjunct distribution of sex reversal is most likely explained by local genetic adaptation to environmental conditions and evolution in the thermal threshold for reversal. Such a mechanism is involved in transitions between GSD and TSD (Quinn et al., 2011), and our population genetic data demonstrate that isolation by distance is the primary pattern of genetic differentiation in this species, providing a pattern of underlying genetic variation. In contrast to other reptiles with very large species distributions, *P. vitticeps* has no obvious barriers to dispersal and no significant population structure (Atkins et al., 2019; Melville et al., 2001, 2017; Sovic et al., 2016). The Murray River in South Australia was the only weak and porous barrier to dispersal that we detected. We propose that the thermal threshold for sex reversal varies in a gradient

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**FIGURE 3** Genetic structure in the central bearded dragon (*Pogona vitticeps*). The structure displays a pattern of isolation by distance, but no strong genetic differentiation between populations throughout the species range. Slight genetic differentiation is apparent among specimens collected south of the Murray River, a geographical barrier to gene flow. (a) Principal coordinates analysis indicates that there is little genetic differentiation between individuals from across the species range. Axes 1 and 2 explain only 2.4% and 1.3% of variation, respectively, and this combined with the lack of clear delineation of groups of individuals, indicates that gene flow is occurring between groups of individuals across the species range, resulting in a geographical gradient of genetic differentiation. (b) Significant and relatively strong isolation by distance was detected (Mantel’s test: $r = 0.571, p = 0.001$), indicating that across the species range, *P. vitticeps* genetic similarity decreases with increasing distance. PCoA and isolation by distance analysis were generated from a filtered SNP dataset of 11,128 binary SNPs and 216 individuals selected from across the species range [Correction added on 7 December 2020, after first online publication: the figure has been modified.]
pattern across the landscape, matching the pattern of isolation by distance. We predict that threshold temperatures will be higher in the northern part of the species range, where incubation temperatures are the hottest.

For evolution of the thermal threshold to occur, heritable variation in the propensity to reverse must exist. Preliminary observations suggest that this is the case in *P. vitticeps*. Offspring of sex-reversed *P. vitticeps* have a lower threshold for sex reversal than the offspring of ZW females (Holley et al., 2015). Additionally, individual females can produce offspring that are completely resistant to sex reversal at all temperatures (Holley et al., 2015). The exact molecular mechanisms of high temperature sex reversal are not known, but a genetic component to the threshold for reversal could be controlled by a single gene of major influence [e.g. CIRBP in the snapping turtle, *Chelydra serpentina* (Schroeder et al., 2016)] or have a polygenic basis [e.g. European sea bass, *Dicentrarchus labrax* (Faggion et al., 2019)]. There is also likely to be an epigenetic component to the sex reversal threshold. Epigenetic mechanisms already implicated in sex reversal include differential expression and temperature-specific splicing of chromatin modifying genes JARID2 and JMJD3/KDM6B (Deveson et al., 2017; Ge et al., 2018).

Compensatory maternal choice in nesting phenology, depth and location cannot fully explain the distribution of sex reversal in *P. vitticeps*. While we acknowledge that uncertainty in nest depth, location and specimen age may contribute to low power in this analysis, another study in a TSD reptile found significant environmental associations at this scale with a similar sampling and analytical approach (Ewert et al., 2005). Additionally, studies in thermally sensitive reptiles are mixed in terms of whether maternal choices are capable of buffering the effects of climate (Doody et al., 2006; Ewert et al., 2005; Gutzke & Paukstis, 1983; Mitchell & Janzen, 2019;Refsnider et al., 2013; Refsnider & Janzen, 2016; Warner & Shine, 2008). We consider it very unlikely that behavioural repertoires to avoid sex reversal would be 100% effective for *P. vitticeps* in the hottest parts of the species range and yet less effective in the relatively cooler regions in which sex reversal was detected (Figure 2). Given the underlying genetic variation in this species (Figure 3) and role of threshold evolution in mediating transitions between GSD and TSD (Quinn et al., 2011), we consider it likely that the sex reversal threshold varies across the species range, buffering against the sex-reversing conditions of northern Australian nesting habitats (Figure 1).

Understanding the capacity of the threshold for sex reversal to evolve is essential for predicting demographic outcomes under altered climate scenarios, including the likelihood of transitions in sex determining modes. Contemporary climate change could accelerate the transition to TSD through greater production of ZZ females and local/widespread loss of the W chromosome. Conversely, conferral of a reproductive advantage to high temperature males could decelerate the transition to TSD (Schwanz et al., 2020). Here, a poor relationship between temperature and the incidence of sex reversal suggests that local genetic adaptation to extreme conditions has already occurred as a sex ratio balancing mechanism. Populations that resist sex reversal then become a source of ZW immigrants, and even small rates of ZW immigration (1% p.a.) significantly decelerate the transition to TSD (Schwanz et al., 2020). Thus, local thermal adaptation and an influx of ZW immigrants could allow *P. vitticeps* to remain stably as a mixed sex determination system in the long term. Importantly, the existence of multiple evolutionary processes working in opposition means that the outcome of transitions between sex determining modes cannot be predicted based on climatic data alone (Schwanz et al., 2020).

The capacity of thermal thresholds to evolve is a critical component of the biological response to climate change, particularly in ectothermic animals. We demonstrate here that threshold evolution may have already occurred across the geographic range of a species with temperature sex reversal. Understanding this component of resilience to changing thermal regimes is essential to future conservation efforts.

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**PEER REVIEW**

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DATA AVAILABILITY STATEMENT
The Scientific Information for Land Owners data used in this paper can be accessed at https://www.longpaddock.qld.gov.au/silo/, and other sources of data were indicated. The associated collection and inferred incubation data for each specimen and CTE calculations for Figure 1 are provided in supplementary files and archived on Dryad. Sequencing data and code for analyses are also provided on Dryad. https://doi.org/10.5061/dryad.98sf7m0h7.

ORCID
Meghan A. Castelli https://orcid.org/0000-0003-1041-0719
Arthur Georges https://orcid.org/0000-0003-2428-0361
Caitlin Cheryb https://orcid.org/0000-0001-6146-4376
Dan F. Rosauer https://orcid.org/0000-0002-2371-1767
Stephen D. Sarre https://orcid.org/0000-0002-7158-2517
Clare E. Holleley https://orcid.org/0000-0002-5257-0019

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BIOSKETCH

Team Pogona is based at the Institute for Applied Ecology (University of Canberra). The team applies a multidisciplinary approach to uncover the molecular basis of genetic and temperature-dependent sex determination in the central bearded dragon and to determine the ecological and evolutionary consequences of sex reversal in the wild. M.A.C. conducted laboratory work and analysis and led the writing of the manuscript. A.G., C.C., D.F.R and S.D.S. conducted and advised on incubation or genetic analysis. I.C-K. assisted with laboratory work. C.E.H. conceived the study and advised on all aspects. All authors contributed to manuscript writing.

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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