Gene flow between two thick-billed grasswren subspecies with low dispersal creates a genomic pattern of isolation-by-distance

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Introgression between grasswren subspecies

Characterising gene flow facilitates conservation management. This study used genomic markers to measure gene flow between thick-billed grasswren subspecies and found results that support taxonomic identification of the two subspecies and suggests grasswrens have low dispersal and may benefit from increased genetic diversity. Recognition of models of divergence with gene flow will be necessary for future conservation management.
Abstract

Context
In the era of the Anthropocene, habitat loss and environmental change threaten the persistence of many species. Genotyping-By-Sequencing (GBS) is a useful molecular tool for understanding how patterns of gene flow are associated with contemporary habitat distributions that may be affected by environmental change. Two parapatric subspecies of the threatened thick-billed grasswren (TBGW; Amytornis modestus) more frequently occur in different plant communities. As such, a preference for plant community type could reduce subspecific introgression and increase genetic diversity at the parapatric boundary.

Aims
We aimed to measure gene flow within and among two TBGW subspecies and tested whether divergent genomic markers were associated with plant community type.

Methods
We sequenced 118 individuals from either of the two TBGW subspecies or in the region of parapatry and identified 7583 SNPs through ddRADseq.

Key results
We found evidence of asymmetric gene flow and a genomic pattern of isolation-by-distance. There were sixteen genomic outliers correlated with plant community type (regardless of location).

Conclusions
These findings show that plant community type does not prevent introgression in one subspecies (A. m. raglessi), but low dispersal and habitat heterogeneity could contribute to the maintenance of distinct subspecific morphotypes. Local adaptation in different plant community types could also provide a mechanism for future divergence.

Implications
We suggest subspecific introgression could increase genetic variation and the adaptive potential of the species, facilitating species persistence under conditions of climate change.

Keywords: genotype by sequencing, dispersal, Maluridae, Amytornis, isolation-by-distance, introgression
Introduction

Habitat loss is the leading cause of reduced species persistence and species extinction (Bradshaw 2012; Newbold et al. 2015; Allan et al. 2019; Thompson et al. 2019). Within Australia, habitat loss has been anthropogenically driven by a multitude of processes that has changed the landscape notably since the late 18th century. These processes include the introduction of invasive species, anthropogenic dispersal of non-local species, redirection/removal of natural water courses, and changes in soil properties due to agricultural practices (Kingsford 2000; Woinarski et al. 2015; Jellinek et al. 2020; Mallen-Cooper and Zampatti 2020). An alarming proportion of extant species are threatened by habitat loss, and, consequently, have reduced population sizes and limited genetic variation on which selection can act (Saccheri et al. 1998; Amos et al. 2012). Molecular tools are important for conservation management practices and species interventions, as they mediate threats to wildlife and ensure long-term success of intervention programs (Elshire et al. 2011; Steiner et al. 2013; Flockhart et al. 2015; Deiner et al. 2017; Forseth et al. 2017). Population genetics can identify populations that may be in greater need of intervention or better suited for conservation management (Dudgeon et al. 2012; Paparella et al. 2015; Whiteley et al. 2015; Willoughby et al. 2015; Rosauer et al. 2018; Mynhardt et al. 2020; Rossetto et al. 2021). Understanding how species respond to habitat changes is relevant for mitigating future threats, especially where further habitat change is predicted to occur.

Populations may be more likely to cope with climate change if they are able to expand their range and move into novel habitats (Hoffman and Blows 1994). There are several evolutionary dynamics that determine whether a species can expand their range or not. These include how much genetic variation there is at the population margin, the strength of genetic swamping of genotypes from central to marginal individuals, and the heritability of adaptive traits at the population margin (Jenkins and Hoffman 1999; Davis et al. 2013; Moerman et al. 2020). Local adaptation into novel environments at the species boundary is one factor that promotes range-expansions, as observed in the European damselfly (*Ischnura elegans*) (Dudaniec et al. 2018). Gene flow can erode local adaption that may favour range expansion, but – if the population is large enough – gene flow could also facilitate local adaptation by...
enhancing genetic variation (Kirkpatrick and Barton 1997; Case and Taper 2000). At
the leading margin of the European lizard (*Zootoca vivipara louislanzi*), low gene
flow has facilitated a range expansion but low genetic diversity throughout the
population could also mean this lizard is susceptible to decline in the face of future
climate change (Dupoué *et al.* 2020). When range-shifts involve secondary contact
between divergent taxa, species persistence could also be affected due to loss of
locally adaptive traits, hybrid swarms or interspecific competition (Case and Taper
2000; Sanchez-Guillen *et al.* 2016). Conservation of threatened species under future
ecological scenarios will depend on the ability to predict range shifts, and an
understanding of the genomics of hybridisation and introgression.

Associations between populations and their habitat develop through ecological
opportunity (Wellborn and Langerhans 2015). For example, morphotypes that give a
population an advantage in their particular habitat type are likely to be retained
(Aiello *et al.* 2021; Grismer 2021). The strength of an ecological association will be
influenced by the amount of gene flow occurring between populations with different
ecological associations, which in turn is dependent on ease of dispersal across the
landscape. Individuals are more likely to disperse to habitats that are similar to their
habitat of origin. This is because individuals that are locally adapted will have lower
fitness outside their original habitat type (Fedorka *et al.* 2012; Berner and Thibert-
Plante 2015). Therefore, populations occurring in linear, unfragmented landscape
arrangements, such as habitat gradients, could have reduced gene flow and in turn
stronger ecological associations (e.g. Cicero 2004). Populations that occur in
landscapes with more diverse patterns of habitat distribution, such as patchy and
heterogeneous landscapes, could have greater gene flow because individuals need to
disperse greater distances to reach particular habitat types and could therefore choose
to remain in an alternate habitat type (Lenormand 2002; Harrisson *et al.* 2012;
Forester *et al.* 2016). It may be less likely for associations between populations and
their habitat to occur in a heterogeneous landscape because gene flow will reduce the
frequency of locally selected alleles. More case studies are needed to complement a
growing body of theoretical modelling, to inform our understanding of the occurrence
of ecological associations and the magnitude of gene flow across different landscape
scenarios, ultimately with a view to better manage extant populations.
The endangered thick-billed grasswren (*Amytornis modestus*, TBGW) is an arid-zone species of the Maluridae family. We adopt the nomenclature of (Black 2011; 2016) which describes seven subspecies of TBGW. There are two extinct and five extant subspecies occurring in parts of the Northern Territory, South Australia and New South Wales (Black et al. 2011; Black and Gower 2017). This taxonomy is a widely accepted (Skroblin and Murphy 2013; Gill and Donsker 2017) however competing taxonomic assignments have been proposed (Christidis et al. 2013; Norman and Christidis 2016). Studies show that *A. m. indulkanna* and *A. m. raglessi* are distinct based on morphology and mitochondrial sequences (Austin et al. 2013). These subspecies share a region of parapatry between the salt lakes, Lake Eyre and Lake Torrens that likely formed due to secondary contact and a possible range expansion (Slender et al. 2017). Outside the region of parapatry, the habitat that each subspecies occupies is characterized by a different and distinct plant community (Slender et al. 2018a). Within the region of parapatry, there is a third ‘sandy’ habitat type where grasswrens were rarely present (Slender et al. 2018a). The Central Australian arid zone is known for its heterogeneous distribution of different plant types (Slatyer 1961; Williams 1982; Brandle 1998). This feature, along with the habitat changes associated with grazing in the arid zone (Jessop 1995; Navarro et al. 2006; Facelli and Springbett 2009), is likely to impact gene flow between populations associated with particular plant communities. In general, the arid zone is predicted to experience greater temperature extremes, less precipitation, and more extreme weather events in the future (Pickup 1998; Lioubimtseva 2004; Lindenmayer and Burgman 2005; Vaghefi et al. 2019). Adaptability through greater genetic diversity will be critical for the persistence of the two parapatric TBGW subspecies.

In this study, we aimed to measure gene flow within and among two TBGW subspecies that have been observed in different plant communities (*A. m. indulkanna* in plant community A, dominated by *Maireana aphylla* [cotton saltbush], and *A. m. raglessi* in plant community B, dominated by *M. astrotricha* [low bluebush] and *M. pyramidata* [blackbush]) (Slender et al. 2018a). The two subspecies may overlap in an area where a third plant community (plant community AB, dominated by *Zygochloa paradoxa* [sandhill canegrass]) occurs but which is not considered suitable foraging habitat for TBGW (Black et al. 2011; Slender et al. 2018a). This area, the parapatric margin, has been proposed as an area of secondary contact. We examine whether
strength of gene flow changes across the three regions that historically were likely to have been demographically different and today contain different plant community types. We test the idea that gene flow is contemporarily higher in the parapatric margin.

**Materials and Methods**

**Samples**

We used DNA from all available TGBW samples which included a combination of 104 contemporary samples and 14 museum samples (Table S1; supplemental material). Contemporary samples were collected in the field by mist-netting birds during the breeding seasons from 2012 to 2015. For further details on the study species and contemporary sample collection methods see Slender et al. (2017).

Museum samples were collected from two time periods; four museum samples were from 1985 (A. m. raglessi) and the remainder were from 2007 to 2009 (A. m. raglessi \[^n = 2\] and A. m. indulkanna \[^n = 8\]) (Austin et al. 2013). Samples were organized into three geographically associated zones described in Slender et al. (2017) in order to compare genetic diversity and gene flow between the subspecies centre’s and their parapatric margin (Figure 1). Zone AB describes the subspecies parapatric margin; zone A describes the geographic centre of A. m. indulkanna and occurs to the west of zone AB and zone B describes the geographic centre of A. m. raglessi and occurs to the east of zone AB. TGBWs in zone A were predominantly found in habitat containing Maireana aphylla (cotton bush) and Atriplex nummularia omissa (Oodnadatta saltbush) (Black et al. 2011; Slender et al. 2018a). While TGBWs in zone B were predominantly found in habitat with M. astrotricha (low bluebush) and M. pyramidata (blackbush) (Black et al. 2011; Slender et al. 2018a). Zone AB contains shrubs typical of TGBW habitat such as M. astrotricha (low bluebush) and A. vesicaria (bladder saltbush), but this was heterogeneously distributed among stands of Zygochloa paradoxa (sandhill caneggrass). The boundary between zone A and zone AB has been extended compared to Slender et al. (2017) so that two museum samples (SAMA B55668 and SAMA B55667) that were formerly included in zone A, now fall within zone AB. This is because the landscape in this area was more like the habitat of zone AB (Slender et al. 2018a).
DNA extraction

Genomic DNA was extracted from tissue and blood in salt solution using a DNeasy Blood and Tissue kit (QIAGEN Pty Ltd, VIC, Australia) or a Gentra Puregene Blood Kit (QIAGEN Pty Ltd, VIC, Australia). Genomic DNA was extracted from FTA samples following Smith and Burgoyne (2004). DNA extractions were carried out in a separate PCR free laboratory in order to minimise DNA contamination. DNA quantity was measured using the Qubit fluorometer (ThermoFisher Scientific Australia Pty Ltd, VIC, Australia). DNA extractions were quality tested using UV-spectrophotometry and agarose gel electrophoresis. Samples were assessed as good quality when they showed 1) a large un-degraded band on an agarose gel and 2) a 260/280 ratio between 1.8 and 2.0 indicating minimal protein and chemical contamination.

Library construction and sequencing

Genotyping-by-sequencing libraries were generated following the protocol in Poland et al. (2012). DNA samples (200 ng) were digested with 8 U of PstI and MspI at 37°C for 2 hrs. Each sample was prepared for multiplexing by ligating a pair of adapters containing a unique barcode to the DNA fragments. We used 96 unique barcodes where the barcodes ranged from 4 to 9 bp (Elshire et al. 2011) to create two pooled libraries. One barcode in each library was assigned as a negative control and seven barcodes in each library were used to duplicate samples within (6 samples) and across (1 sample) libraries. Barcodes were randomly allocated to samples from different geographic locations so that we would detect errors caused by mismatched barcodes that can be made during library preparation or subsequent demultiplexing. We used an adapter mix to DNA ratio of 1:50 ng as this concentration produced libraries with reduced adapter dimer (Elshire et al. 2011). Libraries were then amplified using PCR with the following standard Illumina primers: P5 (5’- AATGATACGGCGACCACCGAGATCTACAC-3’) and P7 (5’- CAAGCAGAAGACGGCATACGAGAT-3’). Sequencing was performed on an Illumina next-seq sequencer that produced single end-reads of 62 bp after adapter trimming. Sequencing data was quality checked using FastQC v10.1 (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/).
SNP calling and filtering

Read filtering and SNP calling was performed using STACKS v1.44 (Catchen et al. 2013). Samples were demultiplexed using the `process_radtags` program and reads from sample replicates were merged into one sample (after preliminary SNP calling with separated duplicates was used to determine error rates). Reads were identified if the adapter barcode (with a maximum of 2 mismatches) and the unique barcode (with a maximum of 1 mismatch) were present. Putative alleles were identified from a stack assembly created with the `ustacks` program that was instructed to include loci with a minimum depth of coverage of 5 reads, maximum distance of 2 nucleotides, and maximum number of 50 stacks per locus. The `cstacks` program was used to create a catalog for identifying loci with a maximum of 2 mismatches between putative alleles. SNPs were determined by comparing the output of `ustacks` with the output of `cstacks` using the `sstacks` program. Relaxing the error tolerance rate improves the likelihood of detecting heterozygous calls (Hohenlohe et al. 2010; Lu et al. 2013). We used a bounded model for detecting SNPs with the lower error limit of 0.0001 and an upper error limit of 0.05. Minor alleles with low frequency cause problems in population genetic analyses because they can represent sequencing error and they are not informative population markers (Gonçalves da Silva et al. 2015). We removed loci (1) that were missing calls in more than 80% of all individuals, or (2) if the minor allele frequency was < 0.05. An individual was considered heterozygous at a locus if there was a proportion of <0.75 reads per allele. We checked that the dyadic likelihood of relatedness did not exceed 0.4 between any individual within zone A and zone AB or zone B and zone AB using the program COANCESTRY v1.0.1.2 (Wang 2011). A related individual of a pair or group of related individuals was excluded if they were related to more individuals and if they had more missing data.

The output from STACKS consisted of 16,569 loci that we applied additional filtering steps to with a custom script implemented in R STUDIO v1.0.136 (R Core Development Team 2008). Loci were removed if they appeared in the negative control and were observed in less than 85% of samples. We used a Principal Component Analysis (PCA) in the R package adegenet v2.0.1 (Jombart 2008) to explore preliminary population structure. The putative clusters without admixed individuals were each analysed for loci out of Hardy-Weinberg Equilibrium (HWE) in
the R package *pegas* v0.9 (Paradis 2010). We removed loci from further analysis that did not conform to HWE in (1) both putative clusters or (2) one putative cluster when a SNP was only present in one cluster. We identified linked loci in each putative cluster excluding potentially admixed individuals, using PLINK v1.07 (Purcell *et al.* 2007). We removed loci from further analysis that were highly correlated ($r^2 > 0.1$) and had a p-value <0.01 in (1) both putative clusters or (2) one putative cluster when a SNP was only present in one cluster. Within a linkage pair, we removed the locus with the most linkage pairs. When both loci had even numbers of linkage pairs, we removed the locus with the most missing data.

**Differences between putative genetic clusters**

$F_{ST}$ outlier loci between putatively non-admixed individuals in zone A and zone B were identified using two programs. We ran BAYESCAN v2.1 (Foll and Gaggiotti 2008) with default settings after data format conversion with PGDSPIDER v2.1.1.0 (Lischer and Excoffier 2012) and the R package *OutFLANK* v0.1 (Whitlock and Lotterhos 2015). $F_{ST}$ outlier loci were defined as having a $q$-value and corresponding false discovery rate of < 0.1. Using a consensus list of $F_{ST}$ outlier loci from both analyses, the dataset was separated into three versions, one with neutral loci (n-SNP), one with only outliers putatively under selection (o-SNP), and a third with both neutral and outlier loci (n+o-SNP). The closest known species relative with an available whole genome sequence is the zebra finch (*Taeniopygia guttata*) (Warren *et al.* 2010). We performed a discontiguous megablast search that looked for sequence similarities between TBGW o-SNPs and the zebra finch GenBank and refseq assemblies using the blastn and blastx functions respectively with an evalue threshold of 1e-6.

To further understand the distribution of shared and distinct genetic variation, we performed an Analysis of Molecular Variance (AMOVA) and calculated the significance of pairwise $F_{ST}$ between zones using GENODIVE v2.0b27 (Meirmans and Van Tienderen 2004) with 10,000 permutations. We tested differences between genetic clusters in three separate analyses; one where the region of parapatry was merged with zone A, one where the region of parapatry was merged with zone B and the last where zone AB was excluded. We repeated these analyses with the n-SNP dataset and n+o-SNP dataset. Expected heterozygosity ($H_e$) is a measure of genomic
diversity when the dataset consists of SNPs (Fischer et al. 2017). \( \hat{H}_e \) was calculated for each zone separately using the n+o-SNP dataset.

Isolation-By-Distance

We tested for Isolation-By-Distance (IBD) among eleven sampling localities by calculating geographic and genetic distance matrices that excluded the locality MTB (zone A) as it contained only one individual (Figure 1). The Euclidean distance between localities (km) was first calculated in GENALEX v6.5 (Peakall and Smouse 2006; Peakall and Smouse 2012). Any paths between localities that passed through Lake Eyre or Lake Torrens (e.g., MUL and WIT) were corrected so that it did not pass through the salt lake. This was done by calculating the Euclidean distance from the first sampling location to a point in the middle of the space between Lake Eyre and Lake Torrens and then calculating the Euclidean distance between that point and the second sampling location and adding the distances together. All geographic distances between sampling localities were then log transformed to account for individuals moving in two dimensions. We calculated a pairwise \( F_{ST} \) genetic distance matrix with n-SNPs using GENODIVE (Meirmans 2020) and also transformed the genetic data \( (F_{ST}/1 – F_{ST}) \) (Nei 1977). Tests for IBD are easily biased by hierarchical population structure where allele frequencies are sharply divided geographically (Meirmans 2012) as well as uneven sample sizes and the spatial patterns between sampling localities (Balkenhol et al. 2009; Guillot and Rousset 2013; Kierepka and Latch 2015). We therefore performed a series of tests for IBD using three different methods; (1) Mantel and partial Mantel tests, (2) Decomposed Pairwise Regression (DPR), and (3) spatial autocorrelation. To test for limitations in gene flow that might prevent genetic swamping at marginal locations we performed two Mantel tests: (1) across locations within zone A (\( A. m. indulkanna \)) and zone AB, and (2) across locations within zone B (\( A. m. raglessi \)) and zone AB. We included zone AB in an analysis with either zone because this area appears to be the population margin for both subspecies (the region of parapatry) (Slender et al. 2017). We then performed a partial Mantel test across all zones to test for gene flow among subspecies while accounting for potential population structure across these regions. We used a binary matrix that compared zone B versus zone AB and A combined or zone A versus zone AB and B combined. We used GENODIVE (Meirmans 2020) to perform mantel and
partial mantel tests with 1000 permutations. DPR is useful for detecting outlier populations that may be associated with weak geographic barriers such as heterogeneous landscapes (Koizumi et al. 2006). We performed a DPR using the R package DPR v1.0 (Reynolds 2011).

Finally, we used spatial autocorrelation (Smouse and Peakall 1999) in GenAlEx v6.5 to further evaluate spatial structure in the genetic data at an individual level. A pairwise matrix with Rousset’s $a$ genetic distance (Rousset 1997; Rousset 2000) between all individuals with the n-SNP dataset was calculated using SPAGeDi v1.4b (Hardy and Vekemans 2002). Geographic distances between individuals were calculated in GenAlEx using the same method to create the geographic distance matrix for the mantel tests. Distance classes were sufficiently small enough to evaluate any non-linear correlations with the autocorrelation coefficient ($r$) where the sample size within each distance class was relatively even. We looked for the presence of IBD within each distance class as well as the detectability of IBD across multiple distance classes (Diniz-Filho and Pires de Campos Telles 2002). Significance was assessed for both tests using 95% confidence intervals for the null hypothesis of no spatial structure using 999 random permutations, and for estimates of $r$ by bootstrapping 999 pairwise comparisons for each distance class.

Gene flow

We investigated population structure and admixture using the n-SNP dataset with two methods: (1) Discriminant Analysis of Principal Components (DAPC) (Jombart et al. 2010) implemented in the R package adegenet v2.0.1 (Jombart 2008), which assigns individuals to genetic clusters following a PCA while accounting for within-population variation; and (2) Bayesian clustering with the program STRUCTURE v2.3.4 (Pritchard et al. 2000; Falush et al. 2003) that determines genetic clustering based on HWE. All methods are useful for detecting admixed individuals. For the DAPC, we retained one principal component, as this returned the optimum $a$-score, which is the difference between the proportion of successfully reassigned individuals compared to the number of principal components retained. The optimum number for $K$ was inferred from the retained principal component by identifying $K$ where the Bayesian Information Criterion (BIC) produced an elbow in the curve of BIC values as a function of $K$. Admixture was inferred if the proportion of population assignment
was <0.9 or >0.1 in any individual. For the STRUCTURE analysis, three replicate runs for each $K$ were analysed (as Standard Deviation of $\ln P(K)$ was small) using default settings, unless stated. We used the admixture model with correlated allele frequencies and an MCMC chain of 1,000,000 iterations with a burnin of 10,000 iterations to test $K$ between 1 and 5. To estimate the probability of mixed ancestry for each individual, the option ANCESTDIST was used. Admixture was inferred if the confidence intervals of the individual population assignment did not include 1 or 0 in all three replicate runs. STRUCTURE HARVESTER (Earl and vonHoldt 2011) was used to estimate the best fitting value for $K$. When the highest $\ln P(K)$ was not $K = 1$, then the most likely $K$ was determined using Delta $K$ (Evanno et al. 2005). Cluster assignments were merged in CLUMPP v1.1.2 (Jakobsson and Rosenberg 2007) and results were visualized with DISTRACT v1.1 (Rosenberg 2004). Population assignments of individuals were compared to their mtDNA haplotype (Slender et al. 2017). Further hierarchical population structure was investigated by repeating the analysis on individual populations detected in the initial run (Evanno et al. 2005).

Selection

Previous comparisons of habitat within the three zones identified three predominant plant communities represented by three principal components (Slender et al. 2018a). We used Latent Factor Mixed Models (LFMMs) (Frichot et al. 2013) to test for associations between genotype (n+o-SNPs) and the environmental variables defined by the principal components. The LFMM test was performed in the R package lfmm v0.0.

Migration

We tested the proportion of migrants between the three zones with a reduced dataset of 200 loci in BAYESASS v3.0.4 (Wilson and Rannala 2003). We performed a PCA with all individuals in the n-SNP dataset with the R package adegenet v2.0.1 (Jombart 2008) and selected loci for use that had the highest loading in the PCA. Three independent MCMC runs were performed with 1,000,000 iterations and a burn-in of 10,000 iterations. The alpha (allele frequency) and delta (inbreeding coefficient) values were adjusted to 0.6 and 0.4 respectively so that the acceptance rates were between 20% and 60%. Iterations were sampled every 100 intervals to determine the
posterior distribution of the parameters. Convergence of the MCMC run was assessed by inspecting the trace file in the program TRACER v1.6.0 (Rambaut et al. 2015).

**Results**

**DNA extraction and sequencing statistics**

We used samples from across three zones: zone A (n = 44), zone B (n = 61), and zone AB (n = 13) to assess gene flow between the two parapatric TBGW subspecies. Following DNA extractions, samples stored on FTA® cards produced considerably lower quantities of DNA (<500 ng) compared to blood stored in salt solution (>1,000 ng). Following illumina sequencing, the average number of reads per sample (before filtering) was 2,539,005 with a coefficient of variation of 24.6%. The average between run reproducibility, calculated by determining when the genotype was the same in duplicates on different plates, was 95.7% (n = 12,192 loci, excluding missing genotypes). The average within run reproducibility, calculated by determining when the genotype was the same in duplicates on the same plate, was 90.5% (n = 146,304 loci, excluding missing genotypes). The average genotyping error rate, calculated from the number of allelic mismatches across duplicates, was 0.31% (n = 316,992 loci). There were no individuals that exceeded > 30% missing data; overall the dataset contained 5.56% missing data. Following SNP calling in STACKS, we removed 5 loci that appeared in the negative control and 2929 loci that had low coverage across samples. An initial PCA showed two putative genetic clusters with individuals from zone A forming one cluster, individuals from zone B forming the second cluster and 19 potentially admixed individuals from zone AB and zone B (Figure S1). We observed similar amounts of missing data between the clusters, excluding potentially admixed individuals (cluster 1 [zone A]: 6.41%, cluster 2 [zone B]: 6.58%). We removed a further 625 loci from further analysis that did not conform to HWE and 5428 loci that could potentially introduce linkage disequilibrium.

**Subspecies variation**

The proportion of heterozygous SNPs per sample varied from 0.234 in a sample from zone A to 0.301 in samples from both zone A and zone B. Mean ± SE estimates of heterozygosity (He) were slightly higher for zone A and zone B compared to zone AB (zone A = 0.303 ± 0.002, zone B = 0.304 ± 0.001; zone AB = 0.288 ± 0.002). The
number of private alleles within zone B was greater (n = 16) than for either zone A (n = 1) or zone AB (n = 0). We identified 39 loci as potential $F_{ST}$ outliers under selection which left 7543 loci that were treated as neutral loci not under selection. Therefore, the dataset o-SNP contained 39 loci and the dataset n-SNP contained 7543 loci. Of the 39 outlier loci, nine loci were monomorphic in zone A; three loci were monomorphic in zone B and four were monomorphic in zone AB. Of the four monomorphic loci in zone AB, three were shared with the monomorphic outliers of zone A and one was shared with the monomorphic outliers of zone B. Four outliers had hits to nucleotide sequences from the zebra finch GenBank assembly (Table S2) but there were no matches to protein sequences from the refseq assembly. These blast hits did not reveal why there could be associations between outlier loci and plant community type.

Zone B had slightly more polymorphic loci in the n-SNP dataset (99.9%) compared to zone A (99.2%). Using the n-SNP dataset, the proportion of total genetic variance was shared among individuals and populations similarly when the region of parapatry (zone AB) was combined with either zone A or zone B, or even when it was excluded (Table 1). The proportion of variance in the case of n-SNP was greater among individuals (mean 0.080%, $p < 0.001$) than among populations (mean 0.008%, $p < 0.001$; Table 1). Using the n+o-SNP dataset, the proportion of total genetic variance explained by population was greater than that explained among individuals for all three tests (A+AB v B, B+AB v A, A v B). This difference was greatest when zone AB was excluded. When zone AB was not excluded the difference was greater when combined with zone B (among individuals = 0.185%, $p < 0.001$; among individuals = 0.094%, $p < 0.001$). Using the n-SNP dataset, there was no difference in pairwise estimates of $F_{ST}$ when the region of parapatry was combined with either zone A or zone B (Table 2). The pairwise estimates of $F_{ST}$ using n+o-SNP was higher when zone AB was combined with zone B (0.202, $p < 0.001$) compared to when zone AB was combined with zone A (0.165, $p < 0.001$; Table 2).

Isolation-By-Distance

IBD was detected in only one Mantel test that included localities from zone A and zone AB ($R^2 = 0.112$, $R_{xy} = 0.335$, $p = 0.029$) (Figure S2). There was no correlation between genetic and geographic distance across localities from zone B and zone AB ($R^2 = 0.012$, $R_{xy} = 0.110$, $p = 0.435$). However, this result may have been affected by
the small number of localities used in this test (Figure S2). Partial Mantel tests across all zones where zone AB was in the same cluster as A or B were significant (zone A+AB vs B; $R^2 = 0.317$, $R_{xy}$ (Spearman’s $r$) = 0.514, $p = 0.001$ and zone B+AB vs A; $R_{xy} = 0.385$, $p = 0.015$) (Figure 2). The sample sizes for the spatial autocorrelation were skewed for the lowest distance class (0-20 kms) but for all other distance classes the sample size was on average ($\pm$ SD) 293 $\pm$ 132. This analysis showed that at an individual level there was positive spatial autocorrelation for the first two distance classes (0-20 and 20-40 kms) (Figure 3). When plotting $r$ as a function of increasing distance classes, the curve intercepted the x-axis at 123.6 kms (Figure 3). IBD was detectible from 0 - 60 km and between 80-100 and 140-160 km ($p < 0.01$). This suggests that spatial autocorrelation is linear up to 60 kms and non-linear at other intervals, which may indicate a pattern of low habitat connectivity. Initial results of the DPR analysis suggested that there were no populations that had greater divergence than what was expected based on distance alone. The model with the smallest AIC$_C$ where $R^2$ was also the highest and where $\Delta$AIC$_C < 2$ was for 3 sub-populations (OOW, MTL, and MUR) to be potential outliers however this was not significant (Table 3). Regression of all sub-populations with all other sub-populations also suggested genetic drift and gene flow were in equilibrium and no population structure was present.

Geneflow

A PCA showed limited population structure between zone A and B along the first component (1.7% of variation) as there was no separation of individuals into clusters (Figure 4). Despite this, STRUCTURE identified two major genetic clusters (Table S3) corresponding to eastern and western populations. Two genetic clusters were also identified by the DAPC analysis albeit with weaker support (Figure S3). Using $K = 2$, results from both STRUCTURE and the DAPC were concordant in that both analyses showed that 1) zone AB contained the highest proportion of admixed individuals 2) there were greater proportions of admixture in individuals in zone AB than either zone A or zone B, and 3) there were greater proportions of admixture in individuals in zone B than in zone A (Figure 5). Comparison of the two methods showed there were discrepancies in the identity of admixed individuals as well as in the proportions of admixture. The DAPC method compromises the power for detecting admixture with
the assignment of individuals to populations, therefore we have limited the discussion
of admixture below to the STRUCTURE results. In zone A, 2.3% of individuals were
admixed and these individuals had a relatively low proportion of assignment
probability from the eastern genetic cluster (< 18%). In zone B, 18% of individuals
were admixed and these individuals had low to high proportions of assignment
probability from the western genetic cluster (18.7 – 52.0%). Two of the admixed
individuals in zone B came from museum samples that were either collected in 1985
or 2007 and were from localities furthest from the region of parapatry (MUR and
MTL). In zone AB, all individuals were admixed and had low to high levels of
assignment probability from both the eastern (17.5 – 71.4%) and western genetic
clusters (28.6 – 82.5%). To look at hierarchical substructure within the identified
populations, individuals in zone B and then zone A were excluded from two separate
STRUCTURE analyses. For zone A and zone AB, $K = 1$ was the most likely using
mean LnP($K$) and for zone B and zone AB, $K = 3$ was most likely using Delta $K$
(Figure S4). Two of smaller clusters from the zone B and zone AB analysis comprised
of groups of individuals that were from the same or neighbouring territories and had
slightly higher levels of relatedness. An earlier analysis with COANCESTRY showed
that the Dyadic likelihood and the 95% confidence intervals for those groups were: $r$
= 0.28 (0.26,0.30) – 0.30 (0.28,0.32) for three individuals in the first cluster and $r$
= 0.14 (0.12,0.16) – 0.28 (0.25,0.30) for seven individuals in the second cluster. The
three individuals in first cluster were also separated along component two (PC2; 1.4%
variance) of the PCA (Figure 4).

Ecological associations and migration

A unique plant community was previously identified in each of the three zones using
a PCA reported in Slender et al. (2018a). PC1 was associated with low abundance of
Atriplex vesicaria and high abundance of Zygochloa paradoxa and was predominant
in Zone AB. PC2 was associated with high abundance of Maireana aphylla and low
abundance of M. astrotricha and M. pyramidata and was predominant in Zone A. PC3
was associated with low abundance of A. nummularia omissa and high abundance of
Acacia spp and Rhagodia spinescens and was predominant in Zone B (Table S4).
Using $K = 2$ output from structure, the LFMM analysis identified 328, 333 and 419
loci associated with PC1, PC2 and PC3 respectively. Of the 39 $F_{ST}$ outliers, there
were 12 loci that correlated with PC2 (two of these also correlated with PC3) and six loci that correlated with PC3. No loci were found to correlate to PC1. The results from BAYESASS suggested that zone AB received more migrants per generation than zone A or zone B (Figure 6). Zone AB received more migrants per generation from zone A than zone B; the mean ± SD migration from zone A = 21.0 ± 4.7% and from zone B = 10.3 ± 4.6%. Zone B received some migration per generation from zone A (4.5 ± 1.6 %), but zone A received < 1% migration per generation from either zone B or zone AB.

**Discussion**

This study aimed to measure patterns of genetic diversity between the parapatric margin (zone AB) of two TBGW subspecies, and their population centre's (A. m. raglessi = zone B and A. m. indulkanna = zone A). Greater genetic variation at the margin could increase the potential for local adaptation to occur in different vegetation types at the margin. We detected gene flow occurring between the subspecies that was not restricted to zone AB as was previously thought and observed no evidence for greater diversity in zone AB compared to other zones. We discovered a pattern of IBD across the subspecies, low genotypic evenness and low genetic differentiation at neutral SNPs based on FST values indicating the subspecies have introgressed considerably. Spatial autocorrelation at short distances suggests that IBD is likely caused by short-range dispersal. We detected more migration between the parapatric margin and the population centre of A. m. raglessi suggesting introgression was asymmetric towards A. m. raglessi. There was evidence of local adaptation in both subspecies to different plant communities, which suggests selection could lead to future differentiation of the subspecies.

IBD increases genetic variation because it occurs when there is low gene flow between distant locations. The presence of IBD indicates that individuals within a population only disperse short distances (Aguillon et al. 2017). Grasswrens are thought to have poor dispersal ability due to their small size and short wings and have highly localized taxonomies (Christidis et al. 2010; Austin et al. 2013). This study found evidence for IBD across the TBGW subspecies, A. m. raglessi and A. m. indulkanna, which have been geographically isolated in the past and have subsequently made secondary contact (Austin et al. 2013; Slender et al. 2017). The
population structure demonstrated in this study is likely biased by the presence of IBD, as limited sampling across large areas replicates patterns of population structure (Perez et al. 2018). Poor dispersal is likely to be one mechanism that has created IBD between these subspecies; however, we also detected patterns suggesting landscape heterogeneity could influence gene flow strength. Further work could assess landscape effects on gene flow strength (van Strien et al. 2015).

IBD in this study indicates considerable nuclear introgression between the subspecies and a low risk of outbreeding depression (Frankham 2010). Introgression may have ensued over a long period of time if secondary contact between A. m. indulkanna and A. m. raglessi occurred a long time ago. Alternatively, there may be a preference for heterospecific mates which could also have led to increased introgression. We previously found that A. m. indulkanna more often and more intensely responded to hetero-subspecific song than con-subspecific song (Slender et al. 2018b). While we know little about the function of grasswren song, it is plausible that greater response to song could indicate mating preferences (Nowicki and Searcy 2005). Introgression of taxonomically young lineages such as subspecies could increase their genetic diversity and the adaptive potential (Grant and Grant 2019). Acknowledging populations that interbreed for conservation planning is a useful component of biodiversity management strategy that is gaining traction in conservation programs (Chan et al. 2019). This stands in contrast to previous concerns that introgression is a threat to biodiversity such as when anthropogenic interference creates conditions that promote species collapse via hybridisation (Allendorf et al. 2001). Conservation approaches need to evaluate the role of hybridisation between populations and species as increased genetic variation may be supported by introgression (Bohling 2016).

Subspecies classifications have a major impact on the allocation of conservation resources (Zink 2004). Both A. m. raglessi and A. m. indulkanna are currently classified as subspecies based on plumage and morphological differences, and a mitochondrial divergence of 1.7% at ND2 (Black 2011; Austin et al. 2013). However, these subspecies are also known to have a continuous distribution and mitochondrial paraphyly (Slender et al. 2017). The lack of genetic differentiation and high level of gene flow between A. m. raglessi and A. m. indulkanna suggests these subspecies could be lumped into one Evolutionarily Significant Unit (ESU) for conservation.
purposes (Moritz 1994; Zink 2004). However, clinal genetic variation caused by IBD
indicates subspecies classifications that require separate management approaches
tailored specifically to A. m. raglessi or A. m. indulgekanna. Other studies show that
phenotypic variation moderately correlates with genotypic variation in natural
populations (Wood et al. 2021). This supports the argument that morphologically
divergent populations make a significant contribution to biodiversity. Managing the
genetic variation captured by each of these subspecies will enable greater adaptive
potential in the future (Fraser and Bernatchez 2001; Coates et al. 2018). Gene flow
and genetic variation have an integral role in conservation management of subspecies,
and sometimes units defined by morphotype is appropriate.

Speciation was traditionally thought to be more commonly associated with population
divergence in allopatry, which has affected how we define species and subspecies
(particularly those not in allopatry) (De Queiroz 2007; Marie Curie Speciation
Network 2012). Examples where populations have undergone divergence with gene
flow are now becoming more common since genomic techniques to assess gene flow
are more accessible (Sousa and Hey 2013; Seehausen et al. 2014; Toews et al. 2016).
This study shows that A. m. raglessi and A. m. indulkanna display patterns of
morphological divergence that are in congruence with outlier loci associated with the
subspecies occurrence in different vegetation types. This pattern is similar to other
models of divergence with gene flow such as the little greenbul (Andropadus virens)
where morphological divergence is more likely explained by the birds occurrence in
different habitat types (savanna versus forest or mountain versus forest) than their
allopatric history (Smith et al. 2005). In another model of divergence with gene flow
(Littorina saxatilis), outlier loci have been genomically linked with loci that control
phenotype, which are selected for according to ecotype (Hollander et al. 2015).
Lastly, Haenel et al. (2021) show how populations of the threespine stickleback fish
(Gasterosteus aculeatus) that possess phenotypes associated with either stream or lake
environments have developed reproductive isolation without any form of geographic
barrier. This research outlines a mechanism for divergence with gene flow and
suggests that the two parapatric TBGW subspecies in this study could be a model of
divergence with gene flow that may continue to diverge in the future.
This study detected two interesting genomic patterns, the cause of which remain unresolved. Mitochondrial paraphyly at ND2 detected by Slender et al. (2017) predicted a low rate of genomic introgression as only 10% of A. m. raglessi individuals had an A. m. indulkanna haplotype. The contradictory results of this study based on nuclear markers could be explained by a number of processes, for example, selection for particular mtDNA haplotypes (Toews and Brelsford 2012; Morales et al. 2015; Morales et al. 2018), or greater dispersal of males compared to females. Other malurid species are known to display female sex-biased dispersal (Cockburn et al. 2003). However, the adult sample size per sex in this study was too small to investigate sex-biased dispersal. Intriguingly, this study also detected a third cluster of individuals in the middle of the A. m. raglessi range that displayed unique genomic variation. We can only hypothesize why these individuals were identified as distinct, but one possible scenario may be that limited gene flow between A. m. raglessi and another TBGW subspecies, such as A. m. curnamona, could be occurring or has more likely occurred in the past. The location of the nearest A. m. curnamona sighting is less than 100 km southeast from an A. m. raglessi sighting (Black et al. 2010). Further sampling of adult grasswrens and the inclusion of samples from other grasswren subspecies may reveal patterns of sex-biased dispersal and explain the source of distinct genomic variation detected within the A. m. raglessi population.

TBGW subspecies show asymmetric gene flow from A. m. indulkanna to A. m. raglessi. The dune field that runs between Lake Eyre and Lake Torrens demarcates the boundary of the asymmetry (Slender et al. 2017). Asymmetric gene flow could occur as the result of several processes, such as greater niche breadth in A. m. indulkanna, demographic or ecological differences on either side of the dune field that promote greater geneflow from A. m. indulkanna to A. m. raglessi (e.g. Oswald et al. 2017), or a mating advantage for A. m. indulkanna (e.g. Baldassarre and Webster 2013; Baldassarre et al. 2014; Slender et al. 2018b). The more frequent and intense response of A. m. indulkanna towards hetero-subspecific song compared to con-subspecific song could suggest A. m. indulkanna is more competitive than A. m. raglessi. A. m. raglessi did not show the same strength of response to hetero-subspecific song compared to con-subspecific song (Slender et al. 2018b). Further work is needed to test hypotheses regarding subspecies behaviour and habitat...
This study shows that two parapatric TBGW subspecies introgressed and that gene flow is asymmetric towards *A. m. raglessi*. Gene flow between the subspecies is limited by distance probably due to the low dispersing ability of the species as well as landscape heterogeneity. We suggest that these subspecies should be taxonomically (and administratively) managed as distinct units despite considerable introgression. Plant community type does not appear to limit geneflow nor does it provide a mechanism for increased genetic diversity at the parapatric margin as was predicted. However, adaptation to different plant community types suggests divergence with gene flow could be a pathway towards increased genomic variation in the future. This study provides an Australian arid zone example to show that gene flow between subspecies can increase genetic variation within a species. Increased gene flow is expected to facilitate persistence of the species through enhanced adaptive capacity. Populations that contain distinct genomic variation should be managed separately particularly in environments that are likely to be affected by future climate change.

*Data availability statement*

The data that support this study will be shared upon reasonable request to the corresponding author.

*Conflicts of Interest*

The authors declare no conflicts of interest

*Declaration of Funding*

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Figure 1. South Australian collection localities for samples of two thick-billed grasswren (TBGW) subspecies used for nuclear genomic sequencing. Localities are grouped into three zones. Sand dunes (grey shade) that run between Lake Eyre and Lake Torrens demarcate a novel habitat type where TBGWs are rarely observed; localities in zone A (solid circle) occur to the west of the sand dunes, localities in zone AB (grey square) occur immediately east of the sand dunes in a region of parapatry (referred to as the contact zone), and localities in zone B occur to the east of zone AB. Locality abbreviations are listed in Table S1. Numbers indicate sample size at that locality.

Figure 2. The pairwise genetic ($F_{ST}/(1 – F_{ST})$) and geographic (log(1 + km)) relationship between localities by zone (zone A: $n = 6$, zone B: $n = 3$, and zone AB: $n = 2$) using a Mantel test ($R^2 = 0.317$). There was only one sample collected at the locality MTB, therefore this locality was excluded. The solid line is the line of best fit and the broken lines are the 95% confidence intervals.

Figure 3. Correlogram showing the spatial autocorrelation coefficient $r$ as a function of distance (km) indicated by distance class (end point). Dotted lines are the 95% CI about the null hypothesis of a random distribution of genotypes and error bars are 95% CI of $r$.

Figure 4. PCA of 7543 loci where individuals from different zones are indicated with different shapes; zone AB (region of parapatry) are white triangles, zone A (A. m. indulkanna) are black diamonds, and zone B (A. m. raglessi) are grey circles.

Figure 5. Population assignment tests using 7543 n-SNP loci where $K = 2$ for (a) DAPC and (b) STRUCTURE or using (c) mitochondrial haplotype for ND2 across three zones (Figure 1). Individuals are ordered by latitude in the order listed in Table S1. The proportion of each colour shows the posterior mean proportion of ancestry from the subspecies A. m. indulkanna or western haplotype (dark grey) and A. m. raglessi or eastern haplotype (light grey). Individuals marked with an asterisk were identified as admixed.

Figure 6. The proportion of migrants (average and standard deviation) assessed between each zone (zone A, zone B, and zone AB) with BAYESASS. Migration from zone A is in black, migration from zone B is in light gray, and migration from zone AB is in dark gray. The analysis was performed following a PCA to identify 200 loci with the highest loading that were used in the BAYESASS analysis.
Table 1. Partitioning of the molecular variance among (1) individuals within zone A and zone B and (2) between zone A and zone B using AMOVA. Zone AB was merged with zone A or zone B in two separate analyses, excluded in a third, or analysed as a separate population. Variance was compared for both n-SNP and o-SNP datasets. Significant $p$-values (< 0.05) are shown in bold.

| Zones included | Dataset | Source of variation | Nested in | % variance | SS          | F-stat | F-value | $P$-value |
|----------------|---------|---------------------|-----------|------------|-------------|--------|---------|-----------|
| A+AB v B       | n-SNP   | Among individuals   | Population| 0.081      | 145109.166  | Fis    | 0.082   | <0.001    |
|                |         | Among populations   | --        | 0.007      | 2197.339    | Fst    | 0.007   | <0.001    |
| A+AB v B       | n+o-SNP | Among individuals   | Population| 0.105      | 806.051     | Fis    | 0.124   | <0.001    |
|                |         | Among populations   | --        | 0.150      | 135.610     | Fst    | 0.150   | <0.001    |
| B+AB v A       | n-SNP   | Among individuals   | Population| 0.082      | 145111.592  | Fis    | 0.082   | <0.001    |
|                |         | Among populations   | --        | 0.007      | 2194.607    | Fst    | 0.007   | <0.001    |
| B+AB v A       | n+o-SNP | Among individuals   | Population| 0.094      | 788.943     | Fis    | 0.115   | <0.001    |
|                |         | Among populations   | --        | 0.185      | 159.104     | Fst    | 0.185   | <0.001    |
| A v B          | n-SNP   | Among individuals   | Population| 0.078      | 128405.213  | Fis    | 0.079   | <0.001    |
|                |         | Among populations   | --        | 0.011      | 2591.425    | Fst    | 0.011   | <0.001    |
|                |         | Among individuals   | Population| 0.075      | 668.666     | Fis    | 0.097   | <0.001    |
|                |         | Among populations   | --        | 0.227      | 184.020     | Fst    | 0.227   | <0.001    |
| A v B v AB     | n-SNP   | Among individuals   | Population| 0.080      | 143301.521  | Fis    | 0.081   | <0.001    |
| Test Type          | Population | Fst     | p-value |
|-------------------|------------|---------|---------|
| Among populations | --         | 0.011   | <0.001  |
| Among individuals | Population | 0.087   | 136065.534 | 0.088 | <0.001 |
|                  | --         | 0.010   | 4008.841 | 0.010 | <0.001 |
Table 2. Pairwise differentiation when zone AB is merged with zone A, zone AB is merged with zone B or zone AB is excluded for both the n-SNP and n+o-SNP datasets. Cells show $F_{ST}$ and $p$-values in parentheses. $P$-values were calculated after 10,000 permutations. Significant $p$-values ($< 0.05$) are shown in bold.

| Zone   | n-SNP  | n-SNP  | n+o-SNP   | n+o-SNP   |
|--------|--------|--------|-----------|-----------|
|        | A      | B      | A         | B         |
| B+AB   | 0.008 ($<0.001$) | --     | 0.202 ($<0.001$) | --         |
| A+AB   | --     | 0.008 ($<0.001$) | --     | 0.165 ($<0.001$) |
| A      | --     | 0.010 ($<0.001$) | --     | 0.227 ($<0.001$) |
Table 3. Fit of alternative isolation by distance models with and without putative outlier populations (see Figure 1 for population codes) $n$ shows the number of populations in the model, $\text{AIC}_C$ shows the corrected Akaike’s information criteria, $\Delta\text{AIC}_C$ shows the difference in $\text{AIC}_C$ between alternative models. These values are used to assess the most likely model. Models are ranked from highest to lowest.

| Population excluded                        | $n$ | $R^2$ | $P$   | $\text{AIC}_C$ | $\Delta\text{AIC}_C$ |
|-------------------------------------------|-----|-------|-------|----------------|-------------------|
| MUL, COP, OOE, WIC, WIT, COS, STC, PEK   | 3   | 0.92  | 0.182 | -26.635        | 0                 |
| MUL, COP, OOE, WIC, WIT, COS, STC        | 4   | 0.883 | 0.005 | -39.507        | -12.872           |
| MUL, COP, OOE, WIC, WIT                  | 5   | 0.825 | <0.001| -48.503        | -21.868           |
| MUL, COP, OOE, WIC                       | 6   | 0.821 | <0.001| -59.135        | -32.500           |
| MUL, COP, OOE                            | 7   | 0.795 | <0.001| -67.610        | -40.975           |
| MUL, COP OOE                             | 8   | 0.742 | <0.001| -76.613        | -49.978           |
| MUL, COP                                 | 9   | 0.575 | <0.001| -82.782        | -56.148           |
| MUL                                      | 10  | 0.434 | <0.001| -89.224        | -62.589           |
| None                                     | 11  | 0.317 | <0.001| -94.404        | -67.770           |
Genetic Distance ($F_{ST}/(1-F_{ST})$) vs. Log($1+$Corrected Geographic Distance)
Proportion of migration per generation

Geographic Origin

- Zone A
- Zone AB
- Zone B

[Graph showing the proportion of migration per generation for different geographic zones]