Species–landscape interactions drive divergent population trajectories in four forest-dependent Afromontane forest songbird species within a biodiversity hotspot in South Africa

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Abstract
Species confined to naturally fragmented habitats may exhibit intrinsic population complexity which may challenge interpretations of species response to anthropogenic landscape transformation. In South Africa, where native forests are naturally fragmented, forest-dependent birds have undergone range declines since 1992, most notably among insectivores. These insectivores appear sensitive to the quality of natural matrix habitats, and it is unknown whether transformation of the landscape matrix has disrupted gene flow in these species. We undertook a landscape genetics study of four forest-dependent insectivorous songbirds across southeast South Africa. Microsatellite data were used to conduct a priori optimization of landscape resistance surfaces (land cover, rivers and dams, and elevation) using cost-distances along least-cost pathway (LCP), and resistance distances (IBR). We detected pronounced declines in effective population sizes over the past two centuries for the endemic forest specialist Cossypha dichroa and Batis capensis, alongside recent gene flow disruption in B. capensis, C. dichroa and Pogonocichla stellata. Landscape resistance modelling showed both native forest and dense thicket configuration facilitates gene flow in P. stellata, B. capensis and C. dichroa. Facultative dispersal of P. stellata through dense thicket likely aided resilience against historic landscape transformation, whereas combined forest-thicket degradation adversely affected the forest generalist B. capensis. By contrast, Phylloscopus ruficapilla appears least reliant upon landscape features to maintain gene flow and was least impacted by anthropogenic landscape transformation. Collectively, gene flow in all four species is improved at lower elevations, along river valleys, and riparian corridors—where native forest and dense thicket better persist. Consistent outperformance of LCP over IBR land-cover models for P. stellata, B. capensis and C. dichroa demonstrates the benefits of wildlife corridors for South African forest-dependent bird conservation, to ameliorate the extinction debts from past and present anthropogenic forest exploitation.

KEYWORDS
Afromontane forest, birds, effective population size, gene flow, landscape genetics, microsatellites
1 | INTRODUCTION

Anthropogenic habitat fragmentation is a global threat to biodiversity, yet responses to this phenomenon vary across taxa (Epps & Keyghobadi, 2015; Lowe et al., 2015; Radespiel & Bruford, 2014). Species differ in sensitivity to habitat degradation following fragmentation (Amos et al., 2012; Devictor et al., 2008; Dondina et al., 2017), and landscape configuration change may independently alter local and long-distance dispersal within a species (Freckleton et al., 2005; Richardson et al., 2016). The impacts of these environmental disturbances may take multiple generations to be detected within populations (Epps & Keyghobadi, 2015; Lowe et al., 2015; Samarasinghe et al., 2005; Richardson et al., 2016). The impacts of these environmental disturbances may take multiple generations to be detected within populations (Epps & Keyghobadi, 2015; Lowe et al., 2015; Samarasinghe et al., 2005; Richardson et al., 2016). The impacts of these environmental disturbances may take multiple generations to be detected within populations (Epps & Keyghobadi, 2015; Lowe et al., 2015; Samarasinghe et al., 2005; Richardson et al., 2016).

2 | MATERIALS AND METHODS

2.1 | Study species

We selected four forest-dependent insectivorous songbirds which experienced range declines across South Africa from 1992 to 2014 (Cooper et al., 2017; Freeman et al., 2018). This group is sensitive not only to forest loss and degradation but also to conversion of the natural vegetation matrix—a trait less apparent in other South African forest-dependent birds (Freeman et al., 2018; Neuschulz et al., 2013; Olivier & van Aarde, 2017). Clearance of coastal thicket, a vegetation type resembling low, recovering IOCB forest, is shown to impede the interforest connectivity of these bird species, as well as forest-dependent mammals, both at the local and regional scale (Ehlers-Smith et al., 2017a, 2017b, 2019, 2020). Despite these community-level observations, it is unknown whether elements of the natural landscape matrix facilitate gene flow in forest-dependent insectivorous birds, and the population genetic stability of these species remains unassessed.

Accordingly, we conducted a comparative landscape genetic study of four forest-dependent insectivorous songbird species, focusing on the southeast region of South Africa, where ranges declines of forest-dependent birds between 1992 and 2014 have been most substantial (Cooper et al., 2017). Our study aims were to assess contemporary levels of genetic connectivity between forest metapopulations within each species and to infer whether these species facultatively disperse through the regional landscape matrix. Additionally, we sought to evaluate the historic stability of the effective population sizes within each species. We undertook this study using microsatellite markers and employed an a priori landscape resistance modelling technique developed by Peterman et al. (2014) and Peterman (2018) to conduct our landscape genetics investigation. We hypothesized that (1) connectivity between regional forest fragments would vary between species, depending on known species vagility; (2) each species would exhibit facultative dispersal through well-wooded habitats (thicket) to facilitate gene flow between naturally fragmented forest; and (3) species with greater forest specialization which had experienced greater contemporary range declines would have more rapidly decreasing effective population sizes.
beyond Afromontane forest into mesic and valley subtypes of Albany thicket (Johnson, 1997).

These species are not trans- or intracontinental migrants (Hockey et al., 2005) and would be affected only by regional anthropogenic activity; C. dichroa is endemic to South Africa. Within South Africa, each species may undertake seasonal altitudinal migration between inland and coastal forests (Johnson & Maclean, 1994; Oatley, 2017), especially P. stellata (Craig & Hulley, 2019; Oatley, 1982a), although such movements do not involve whole-population shifts. Ad hoc South African recapture records (Oschadleus & Ranwashe, 2017) suggest higher adult vagility in P. stellata (83 km) and B. capensis (50 km) compared to C. dichroa (8.5 km) and P. ruficapilla (3.4 km).

The former two species more readily traverse open matrix habitats (Aben et al., 2012, 2014; Callens et al., 2011; Dane & Bolton, 2017; Galbusera et al., 2004), although seasonally migrating P. stellata may prefer navigating along riparian forest/thicket corridors (Oatley, 1982a, 2017).

2.2 Field sampling and laboratory procedures

From 2017 to 2018, standardized mist-netting was conducted in six Afromontane forests, two temperate IOCB forests and three forests which are intermediate between both sub-biomes (scarp forests) (Figure 1), visiting each forest for 3 weeks. These 11 forests in the Eastern Cape and southern KwaZulu-Natal Provinces of South Africa fall within the Maputo-Pondoland-Albany biodiversity hotspot (Mittermeier et al., 2004) and were chosen to maximize equal sampling of these scarce species. In total, 114 B. capensis, 94 C. dichroa, 92 P. ruficapilla, and 200 P. stellata were captured (Table 1). Birds were banded to prevent resampling and released at the capture location after sampling of 20–50 μl of blood, collected from the brachial vein using sterile hypodermic needles and heparinized tubes, in conformity with South African legal requirements (see Acknowledgements). Collected blood samples were preserved in 500 μl 95% ethanol, and genomic DNA was extracted using a Nucleospin Tissue DNA extraction kit (Macherey-Nagel). For each species, we screened a separate microsatellite library available from past literature (9–29 loci per species kit (Macherey-Nagel). For each species, we screened a separate microsatellite library available from past literature (9–29 loci per species). We randomized within-species sampling to minimize false-positive discovery from amplification conditions). We randomized within-species samples to minimize false-positive discovery from amplification conditions. We randomized within-species samples to minimize false-positive discovery from amplification conditions.

Amplification errors (large allele dropout, stuttering and null alleles) were checked in MICROCHECKER 2.2.3 (Van Oosterhout et al., 2006) and FREEENA (Kawashima et al., 2009). Forest-level deviations from expectations of Hardy–Weinberg equilibrium (HWE) and linkage disequilibrium (LD) within forests were assessed in GENEPOP4.7 (Rousset, 2008); adjusting significance values using a Benjamini-Hochberg correction (Benjamini & Hochberg, 1995) to control for false discovery rate. Forest-level species genetic diversity was estimated as rarefied allelic richness (AR), and private allelic richness (PrAR) in ADZE1.0 (Szpiech et al., 2008); observed (H̄) and expected (H) heterozygosity; and inbreeding coefficient (F_IS) in GENETIX4.05 (Belkhir et al., 2001). As a precautionary measure against low sample sizes and the limited number of microsatellite loci employed in this study, we used POWSIM 4.1 (Ryman & Palm, 2006) to assess the power of each microsatellite data set to detect population substructures at F_ST = 0.05 (effective population size, N_e = 2000; generations of genetic drift, t = 210), F_ST = 0.01 (N_e = 2000, t = 40), and F_ST = 0.001 (N_e = 2000, t = 4). An N_e of 2000, approaching the upper bounds of the estimated effective population size (see Figure 2; Table S2), was selected as larger N_e are considered more appropriate; the value of t was selected following recommendations by Ryman and Palm (2006) to test for the particular F_ST. Simulations were performed assuming two subpopulations (N = 50 and N = 40) over 1000 replicates, and statistical power was measured as the proportion of significant tests.

Population genetic substructures were investigated through Bayesian clustering using STRUCTURE (Pritchard et al., 2000). The optimal number of genetic clusters per species (K) were tested for K = 1–12 (the number of forest sites +1). Twenty independent runs of 5 × 10^6 Markov chain Monte Carlo (MCMC) iterations and a burn-in period of 5 × 10^5 were performed per K, using the admixture model, correlated allele frequencies, and with LOCPRIOR (grouped by forest). Results of runs averaged in STRUCTURE HARVESTER (Earl & vonHoldt, 2012), and the optimal number of clusters was
determined using the Evanno ΔK statistic derived from posterior probability of each value of K (Evanno et al., 2005). STRUCTURE results were visualized in the Pophelper R package (Francis, 2017).

To further investigate population structure, we performed principal component analysis (PCA) in the adegenet R package (Jombart, 2008) based on individual allele frequencies. Finally, to determine population differentiation, we calculated global and pairwise $F_{ST}$ among forests for each species in ARLEQUIN 3.5 (Excoffier & Lischer, 2010).

### TABLE 1 Sample sizes, estimates of genetic diversity, and inbreeding coefficients within each forest for the four focal bird species

| Forest | N  | AR  | PrAR | $H_o$  | $H_e$  | $F_{IS}$ |
|--------|----|-----|------|--------|--------|----------|
| B. capensis |   |     |      |        |        |          |
| Ngele  | 6  | 4.750 | 0.475 | 0.771  | 0.775  | 0.005    |
| Oribi  | 8  | 4.250 | 0.035 | 0.650  | 0.717  | 0.103    |
| Gomo   | 17 | 6.750 | 0.179 | 0.603  | 0.682  | 0.119    |
| Nqadu  | 8  | 5.625 | 0.243 | 0.607  | 0.806  | 0.260    |
| Baziya | 14 | 6.500 | 0.253 | 0.580  | 0.688  | 0.161    |
| Manubi | 16 | 7.625 | 0.148 | 0.578  | 0.705  | 0.184    |
| Kubusi | 13 | 6.375 | 0.109 | 0.635  | 0.708  | 0.165    |
| Fort Fordyce | 18 | 7.125 | 0.252 | 0.597  | 0.712  | 0.107    |
| The Island | 14 | 5.625 | 0.070 | 0.607  | 0.720  | 0.162    |
| Total  | 114 | 10.625 | 0.196 | 0.625  | 0.709  | 0.151    |
| C. dichroa |   |     |      |        |        |          |
| Ngele  | 6  | 4.000 | 0.228 | 0.688  | 0.653  | −0.058   |
| Oribi  | 5  | 3.125 | 0.157 | 0.500  | 0.536  | 0.077    |
| Baziya | 12 | 5.000 | 0.225 | 0.635  | 0.636  | 0.001    |
| Manubi | 22 | 7.500 | 0.345 | 0.710  | 0.692  | −0.027   |
| Kubusi | 26 | 7.000 | 0.395 | 0.702  | 0.720  | 0.026    |
| Fort Fordyce | 17 | 5.750 | 0.278 | 0.632  | 0.639  | 0.011    |
| Alexandria | 6  | 3.625 | 0.220 | 0.667  | 0.642  | −0.042   |
| Total  | 94 | 8.000 | 0.264 | 0.648  | 0.636  | 0.002    |
| P. ruficapilla |   |     |      |        |        |          |
| Ngele  | 20 | 1.429 | 0.104 | 0.425  | 0.361  | −0.184   |
| Oribi  | 6  | 2.250 | 0.005 | 0.325  | 0.364  | 0.119    |
| Mbotyi | 6  | 2.375 | 0.050 | 0.429  | 0.541  | −0.135   |
| Gomo   | 11 | 2.250 | 0.046 | 0.386  | 0.343  | 0.001    |
| Baziya | 14 | 2.500 | 0.032 | 0.417  | 0.415  | −0.003   |
| Manubi | 15 | 2.875 | 0.100 | 0.367  | 0.401  | 0.087    |
| Kubusi | 9  | 2.375 | 0.003 | 0.375  | 0.354  | −0.063   |
| Fort Fordyce | 11 | 0.286 | 0.002 | 0.403  | 0.390  | −0.035   |
| Total  | 92 | 3.375 | 0.043 | 0.391  | 0.381  | −0.047   |
| P. stellata |   |     |      |        |        |          |
| Ngele  | 30 | 6.625 | 0.212 | 0.667  | 0.673  | 0.009    |
| Oribi  | 13 | 4.875 | 0.122 | 0.625  | 0.683  | 0.088    |
| Gomo   | 26 | 6.125 | 0.188 | 0.705  | 0.679  | −0.038   |
| Baziya | 15 | 5.875 | 0.181 | 0.667  | 0.659  | −0.012   |
| Manubi | 28 | 6.250 | 0.094 | 0.612  | 0.668  | 0.086    |
| Kubusi | 26 | 6.250 | 0.176 | 0.606  | 0.652  | 0.072    |
| Fort Fordyce | 39 | 5.875 | 0.058 | 0.622  | 0.648  | 0.042    |
| Alexandria | 17 | 4.750 | 0.070 | 0.656  | 0.605  | −0.088   |
| The Island | 6  | 4.375 | 0.319 | 0.646  | 0.595  | 0.095    |
| Total  | 200| 9.750 | 0.185 | 0.646  | 0.651  | 0.022    |

### 2.4 Demographic history

To infer contemporary gene flow disruption, we compared pairwise $F_{ST}$ estimated in ARLEQUIN 3.5, to the proportions of shared allele statistic $D_{PS}$, calculated in MSA 4.0 (Dieringer & Schlötterer, 2003). Lag time to detection of new gene flow barriers is shorter when measuring $D_{PS}$ compared to $F_{ST}$ (Landguth et al., 2010; Robin et al., 2015; Savary et al., 2021), and so larger $D_{PS}$:$F_{ST}$ ratios may suggest recent reductions in gene flow (Robin et al., 2015). We
estimated regional contemporary effective population sizes ($CN_e$) for each species, using the LD model (for single sampling events) in NEESTIMATOR 2.1 (Do et al., 2014). We separately assumed random and monogamous mating (typically observed; Hockey et al., 2005). We observed results at 0.02 and 0.01 critical allele frequencies to better accommodate limited data sets (Do et al., 2014) and used a pairwise jackknife approach to assess confidence intervals; within-species samples were pooled to accommodate overlap/inter-breeding among the most recent generations. We further inferred variation in focal species $N_e$ over the most recent 100 generations using the VarEff R package (Nikolic & Chevalet, 2014). Default parameter conditions were kept across species, adjusting maximum distance between alleles ($DMAX = 18 – P. stellata; 17 – C. dichroa; 22 – B. capensis; 10 – P. ruficappila$); number of past $N_e$ changes ($JMAX = 3$); and generations since the most recent common ancestor ($GBAR = 1000; reduced from the default GBAR = 5000 given the low population differentiation observed for each species [Nikolic & Chevalet, 2014]$). Runs were performed under both single-step mutation model (SMM), and 10% single-step two-phase mutation (TPM) to accommodate a broader range of mutation dynamics within natural populations. Mutation models assumed a mutation rate of $\mu = 5 \times 10^{-4}$ per generation (Brohede et al., 2002; Coetzer et al., 2020), with an acceptance ratio of 0.25.

2.5 | Landscape resistance modelling

2.5.1 | Landscape genetics framework

Landscape genetics frameworks provide a means to investigate relationships between genetic distances and features landscape, by modelling resistance surfaces of spatially arranged cost values to gene flow (Mannel & Holderegger, 2013; Mannel et al., 2003; Waits et al., 2015). To investigate the regional landscape influences on the interforest connectivity within each species, we adopted an a priori approach of resistance surface parameterization using RESISTANCEGA 4.1 R package (Peterman, 2018; Peterman et al., 2014). This approach circumvents subjectivity of conflicting expert opinion (Charney, 2012; Zeller et al., 2012) and limited applicability of niche-model derivations towards atypical landscape use (Balkenhol et al., 2015; Keeley et al., 2017; Vasudev et al., 2015; Zhan et al., 2017). The RESISTANCEGA 4.1 R package integrates mixed-effects modelling and stochastic genetic algorithms mimicking natural selection (Scrucca, 2013) specifically to maximize the relationship between pairwise genetic distances of samples and resistance surfaces. Models were fitted using maximum-likelihood population effects (MLPE) parameterization (Clarke et al., 2002) in the LME4 R package (Bates et al., 2014) where fitness was assessed using corrected Akaike information criteria ($AIC_c$). Models with an $AIC_c$ difference ($\Delta AIC_c < 2$ were considered equivalent (Burnham & Anderson, 2004). We modelled two ecological distances for each landscape surface: isolation-by-resistance (IBR) considers cumulative current-flow costs across all possible paths between two points (McRae, 2006), whereas least-cost pathways (LCP) are spatial entities along which the accumulated costs are minimized (Adriaensen et al., 2003; Marrotte & Bowman, 2017). The IBR models were constructed using commute-time resistance distances, an equivalent to circuit theory models for determining flow resistance (Lundgren & Ralph, 2019; Marrotte & Bowman, 2017; McRae et al., 2013), in RESISTANCEGA4.1. The LCP models in turn were made using the ‘cost distance’ function in the gdistance R package (Van Etten, 2015).

![FIGURE 2 Demographic trends within B. capensis, C. dichroa, P. ruficappila and P. stellata across a region of southeast South Africa: (a) ratios between $F_{ST}$ and $D_{pq}$ genetic distance metrics within each species among sampled forest sites; (b) regional $CN_e$ size of each species measured at 1% and 2% critical allele frequencies, and assuming monogamous mating (with 95% confidence intervals); (c) VarEff plots showing variation in $CN_e$ of each species over the past 100 generations, assuming a single-step mutation model at a constant mutation rate of $\mu = 5 \times 10^{-4}$ per generation. Species demographic trends are inferred from a combination of eight species-specific loci unique to each species.](image-url)
2.5.2 Landscape variables

We assessed the relative influence of three landscape variables on connectivity in each focal species (Figure 3): (i) land cover, as matrix landscape has been shown to affect forest-dependent insectivorous birds; (ii) rivers and dams, as these species may use riparian corridors both for seasonal migration, and dispersal; and (iii) elevation, as species populations may be isolated along an elevation gradient. Land cover and rivers and dams resistance surfaces were based on 20 m categorical land-cover classes taken from the South African National Land-Cover (SANLC) 2018 (Thompson, 2019), while topographic surfaces were based on 7.5 arc-second (250 m) categorical map of mean elevation in metres above sea level (m.a.s.l), taken from the Global Multi-resolution Terrain Elevation Data 2010 (Danielson & Gesch, 2011). We down-scaled the resolution of all resistance surfaces to 250 m × 250 m for computational efficiency without significant loss of landscape-genetic associations (McRae et al., 2008). For riparian corridors, we grouped rivers, estuaries, dams, and herbaceous wetland elements and classified every cell containing these elements as ‘river’: we then added a single-cell buffer around all ‘rivers’ to prevent diagonal side-stepping when modelling LCP and IBR under an 8-connexity parameter. For land cover (see below), we classified cells according to the majority land-cover class within each cell. We prepared resistance surfaces by creating a convex hull around forest sites using the Convex Hull plug-in for QGIS3.10 (QGIS Development Team, 2019), and setting a 50-km buffer as the boundary for downstream analyses. The latter condition was included to accommodate total movement among available forest habitat throughout the study area.

2.5.3 Thematic resolution of landscape matrix

To infer matrix permeability across different land-cover classes, we tested four alternative land-cover surfaces under different classification schemes (Figure 3a). The original 72 land-cover classes were consolidated into seven categories: (a) native forest (≥75% tree canopy cover; ≥6 m canopy height); (b) dense thicket—coastal/mesic/valley thicket (≥75% tree canopy cover; 2.5–6 m canopy height); (c) woodland (35–75% tree canopy cover; ≥2.5 m canopy height); (d) open habitats—grassland, shrubland and savanna (<35% tree canopy cover); (e) artificial plantation (exotic Pinus/Eucalyptus); (f) agriculture; and (g) human infrastructure (urban, suburban, rural, industrial, transportation networks and mining). From this, four resistance surfaces were classified as follows: (1) native forest configuration (the primary habitat of each species); (2) forest and dense thicket configuration, as the latter habitat improves forest bird community connectivity; (3) natural land cover (forest, dense thicket, woodland, open habitats) versus anthropogenically transformed land cover (exotic plantation, agricultural land, human infrastructure); and (4) a comprehensive surface comprising all seven categories. For computational efficiency, land-cover resistances surfaces 1–4 were optimized three times per species, using least-cost distances based on pairwise F<sub>ST</sub>. Given that D<sub>PS</sub> better responds to recent landscape change (Savary et al., 2021), we optimized the these resistance surfaces once using this genetic distance metric, as F<sub>ST</sub> results proved stable across replicates (Figure 4). The best-supported thematic resolution for each species was used in subsequent analyses.

2.5.4 Resistance surface optimizations and landscape distance model comparisons

Univariate optimization was conducted in RESISTANCEGA 4.1, separately on the three landscape resistance surfaces (land cover, riparian corridors and elevation), using both LCP and IBR modelling. Subsequently, we conducted multivariate optimization, wherein the optimizations of all three surfaces are summed to form a composite

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**FIGURE 3** Landscape surfaces at a spatial resolution of 250 × 250 m cell size used to calculate LCP and IBR ecological distance modelling. (a) Four classifications of land-cover surfaces: (i) native forest configuration; (ii) forest and dense thicket configuration; (iii) natural land cover (native forest, dense thicket, woodland and open habitats) and combined anthropogenically transformed land cover (artificial plantation, agricultural land and human infrastructure); (iv) all seven land-cover classes; (b) rivers and dams; (c) elevation (in metres above sea level)
surface over which LCP and IBR distances are modelled. All optimizations (except previously optimized land-cover LCP models) were optimized three times, separately for each species. All IBR and LCP models were compared to null models. Following Cushman et al. (2013) and Khimoun et al. (2017), we used causal modelling to compare species-specific IBR and LCP ecological distance models to isolation-by-distance (IBD), according to the Spearman correlations between genetic and landscape distances. Despite the high rates of type I error alleged for partial Mantel tests (Castellano & Balietto, 2002; Raufaste & Roussel, 2001), use of these tests in a causal model framework to reject the incorrect causal model and to identify the most applicable models driving observed genetic patterns is considered appropriate (Cushman & Landguth, 2010; Cushman et al., 2013; Khimoun et al., 2017). We inferred the most relevant landscape model using Mantel and partial Mantel tests in the ecodist R package (Goslee & Urban, 2015), with 10,000 random permutations, using distance values for LCP and IBR models, and log-transformed Euclidean distances for IBD. Spatially constrained, yet otherwise indiscriminate dispersal should best reflect IBD model. Dispersal which navigates select routes across landscape elements efficiently should reveal LCP, whereas inefficient dispersal, or dispersal across numerous paths through landscape elements should show IBR.

3 | RESULTS

3.1 | Microsatellite characteristics and genetic diversity

All individuals were successfully genotyped for all loci (Table 1). The eight informative microsatellite markers retained per species exhibited no large allele dropout, or stuttering, displayed null allele frequencies <5% across populations, and had limited deviations from expectations of LD and HWE (Appendix S1–S3). At the forest level, loci were in HWE after Benjamini–Hochberg corrections, except two expectations of LD and HWE (Appendix S1–S3). At the forest level, P. ruficapilla species-specific sets of loci (in corrections, while no significant linkage was evident within the other species in 2/252 (0.79%) LD tests concerning different loci/population combinations showed it was possible to detect genetic divergence as low as $F_{ST} = 0.01 (t = 40)$ with 94.4% certainty in B. capensis, 92.4% in C. dichroa and 91.1% in P. stellata, but only 55.7% certainty for P. ruficapilla, although $F_{ST} = 0.02 (t = 80, n = 2000)$ could be detected with 84.7% certainty for this species. All four microsatellite data sets had low certainties (6.5%–12.0%) to detect $F_{ST} = 0.001 (t = 4)$.

3.2 | Genetic population structure

Global genetic differentiation was significant for C. dichroa ($F_{ST} = 0.036, p < 0.001$) and P. stellata ($F_{ST} = 0.016, p < 0.001$), but not for B. capensis ($F_{ST} = 0.013, p = 0.100$) and P. ruficapilla ($F_{ST} = 0.006, p = 0.261$). STRUCTURE identified a maximum of two genetic clusters within the regional populations of each species, respectively, according to Evanno $\Delta K$: B. capensis ($\Delta K = 7.275$), C. dichroa ($\Delta K = 1.184$), P. ruficapilla ($\Delta K = 1.099$) and P. stellata ($\Delta K = 3.034$) (Figure 5a–d, Figure S1.1–S1.4). In both B. capensis and P. stellata, there is a subtle north–south gradient, wherein individuals from the southern IOCB forests (Alexandria and The Island) appear partially differentiated from individuals within Ngele to Fort Fordyce (Figure 5a,d). This gradient is less apparent in P. stellata where individuals from Ngele to Fordyce showed higher mixed ancestry compared to B. capensis (Figure 5a,d). In C. dichroa, there is also a subtle north–south gradient, although individuals from Kubusi appear particularly differentiated from other regional forests (Figure 5b). Finally, in P. ruficapilla, the population structure appears uniform across the study area (Figure 5c). The PCA results suggested that all four species, individuals across regional forests broadly comprise genetically homogenous groups (Figure 6a–d). However, in C. dichroa, individuals from Kubusi show large variability and partially cluster away from other forest groups (Figure 6b), and this is similarly observed in P. ruficapilla for individuals from Mbotyi (Figure 6c).

3.3 | Demographic history

Pairwise $D_{PT}^{*}:F_{ST}$ ratios were highest for B. capensis, P. stellata and C. dichroa, and lowest for P. ruficapilla (Figure 2a). Overall $CN_e$ appears lowest in C. dichroa and highest in P. ruficapilla, although both have larger 95% CI compared to P. ruficapilla and P. stellata (Figure 2b; Table S2). Species $CN_e$ assuming monogamous mating (typically observed in each species [Hockey et al., 2005]) were twice as high compared to assuming random mating (Table S2). Disparities between $CN_e$ at 1% and 2% critical allele frequencies were minimal in B. capensis, 18% in P. stellata, 33% in C. dichroa and 150% for P. ruficapilla, reflecting lower rare allele frequencies...
in the last three species (Do et al., 2014). Fluctuations in $N_e$ over the past 100 generations varied across the four songbirds, consistent across single-step (Figure 2c), and two-phase (Figure S2) mutation models. Historically, $B. capensis$ and $C. dichroa$ had the largest $N_e$, but declined to levels comparable to $P. ruficapilla$ and $P. stellata$, which both appear more temporally stable, but still in decline (Figure 2c). Assuming a two-year generation time (Bird et al., 2020), or three years for $P. stellata$ (Oatley, 1982b), these events relate to the past three centuries, with most declines beginning <100 years (~20–60 generations) ago.
3.4 | Landscape genetics

3.4.1 | Land-cover thematic resolution

Land-cover thematic surface evaluation (Figure 4) indicated that genetic distances in *P. stellata* were best explained by the native forest and dense thicket configuration model, which outranked the geographic distance model by >17 AICc units. For *B. capensis* and *C. dichroa*, both the native forest configuration, and the native forest and dense thicket configuration model were comparable to the geographic distance model (<2 AICc units) (Figure 4). However, in *P. ruficapilla*, the native forest configuration model ranked second to the geographic distance model (<4 AICc units) (Figure 4). For each species, land-cover models ranked similarly across pairwise *F*<sub>ST</sub> and *D*<sub>PS</sub> (Figure 4, Figure S3), except for *C. dichroa*, in which the null model outranked all other models when using pairwise *F*<sub>ST</sub> (Figure 4), but ranked below the geographic distance model, and native forest and dense thicket configuration model by <4 AICc units (Figure S3). According to both pairwise *F*<sub>ST</sub> and *D*<sub>PS</sub>, detailed land-cover configurations were inadequate to explain genetic distances observed in the four species (Figure 4; Figure S3).

3.4.2 | Landscape resistance surfaces

For *B. capensis*, only the resistance distance model from elevation outranked the geographic distance model (>8 AICc units), although least-cost distance models from both land-cover (native forest and dense thicket configuration), and rivers and dams were comparable to the geographic distance model (Figure 4). For *C. dichroa*, the null model outranked all landscape models (Figure 4). For *P. ruficapilla*, the least-cost distance model from elevation, and the resistance distance models from elevation, and rivers and dams were comparable to the geographic model (Figure 4). Only in *P. stellata* did univariate least-cost and resistance distance models from each univariate landscape surface consistently outranked the geographic distance model (Figure 4), of which the least-cost distance model from land cover (native forest and dense thicket configuration) ranked the highest (>17 AICc). Across all four species, the least-cost and resistance distance models from the composite landscape surface ranked far lower than the geographic distance model (Figure 4).

3.4.3 | Comparative performance of landscape models

Both *B. capensis* and *P. stellata* showed significant IBD according to pairwise *F*<sub>ST</sub> (Table 2), whereas *C. dichroa* and *P. stellata* showed significant IBD according to pairwise *D*<sub>PS</sub> (Table 3). Partial Mantel tests of either LCP or IBR models controlling for IBD suggest that for *B. capensis*, *P. ruficapilla* and *P. stellata*, genetic distances (pairwise *F*<sub>ST</sub>) better correlated with certain landscape elements than geographic distance (Table 2). In *B. capensis*, genetic distances correlated significantly with the IBD-controlled LCP (LPC|IBD) models for native forest and dense thicket, rivers and dams, and landscape elevation (Table 2). In this species, landscape elevation appears especially pertinent to gene flow, with both LCP|IBD and IBR|IBD models for elevation strongly correlating to genetic distances (Table 2). In *P. ruficapilla*, genetic distances correlated strongly with IBR|IBD models with native forest land-cover, rivers and dams, and especially with landscape elevation (Table 2). In this species, causal modelling, however, did not support one ecological distance model of the other (Figure 7). In *P. ruficapilla*, genetic distances correlated strongly with IBR|IBD models with native forest land-cover, rivers and dams, and especially with landscape elevation (Table 2). In this species, causal modelling showed that the IBR model of landscape elevation remained significant even after controlling for LCP (Figure 7). In *P. stellata*, genetic distances correlated with all tested ecological distance models (after controlling for IBD), but especially so for the LCP|IBD model for native forest and dense thicket land cover (Table 2). The association between genetic distances within *P. stellata* and these land-cover classes remained significant across both pairwise *F*<sub>ST</sub> (Table 2), and pairwise *D*<sub>PS</sub> (Table 3). In this species,
causal modelling further corroborated the LCP model over the IBR model for these land-cover classes, further showing that IBD may better explain genetic distances than this land-cover IBR model (Figure 5). Although no landscape model pairwise $F_{ST}$ genetic distances in *C. dichroa* (Table 2), the LCP model for both native forest, and native forest and dense thicket remained significant after controlling for IBD (Table 3).

### 4 | DISCUSSION

The four forest-dependent insectivorous songbirds in this study adequately maintain gene flow among spatially isolated metapopulations confined to the fragmented native forests of southeast South Africa (Figure 5). Gene flow disruptions are however notable in *B. capensis*, and to a lesser extent in *P. stellata* and *C. dichroa* (Figure 2a). Two species, *B. capensis* and *C. dichroa*, show pronounced declines in effective population sizes (Figure 2c), signifying vulnerability of these avian insectivores to increasing anthropogenic habitat changes. The endemic forest-specialist *C. dichroa* has experienced the most drastic population declines over the past two centuries, which have continued well after the widespread cessation of commercial selective logging and deforestation 80 years ago.

#### 4.1 | Population genetic structures

The South African endemic *C. dichroa* displayed the highest population structuring (Figure 5a), and the large genetic variability unique to Kubusi (Figure 6b) affirms the climatic refugial importance of the eastern Amatole forest complex (Dalton et al., 2015; Kushata et al., 2020; Madisha et al., 2018). The higher population complexity of this forest specialist contrasts with that of the forest generalist *B. capensis*, which is near endemic to South Africa, and more genetically diverse than *C. dichroa* (Table 1). Higher historic availability of suitable habitat for *B. capensis* likely afforded larger populations that were more buffered against Paleoclimatic fluctuations, although both native forest (Ivory et al., 2018; Lawes, Eeley, et al., 2007) and Albany thicket (Potts et al., 2013) biomes were susceptible to contractions during periodic aridity. Consistently higher genetic diversity at Manubi across species (Table 1) corroborates the refugial significance of intermediate (scarp) forests within South Africa (Grass et al., 2015; Lawes, Eeley, et al., 2007; Moir et al., 2020; Moir et al., 2021), which are close to the coast, buffering them from paleoclimatic extremes. The lower diversity at Oribi Gorge may reflect greater proximity to subtropical IOCB forests (Mucina, 2018; Mucina et al., 2006), which re-established only ~8 kya (Huntley et al., 2016) and are generally avoided by these four songbird species. The low genetic diversity observed in *P. ruficapilla* suggests more recent, or perhaps constrained, colonization of South Africa than does *P. stellata*, and the unexpectedly low regional complexity within *P. ruficapilla* contrasts with the strong population insularity observed in East Africa (Callens et al., 2011). This alludes to dispersal behaviour...
of this species changing from sedentary in tropical Africa, to vagile in higher latitude forests of South Africa (Martin & Tewksbury, 2008; Moore et al., 2008; Salisbury et al., 2012).

### 4.2 Population viability and effective population size

Our study findings suggest that historic deforestation and commercial selective logging likely had a large negative impact on the viability of these four species (Figure 2c), more so than contemporary informal forest resource harvesting, despite it being largely unregulated (Leaver & Cherry, 2020a, 2020b; Leaver et al., 2019). The contemporary effective population sizes of all four species (Figure 2b; Table S2) are likely underestimates resulting from pooled generations (Luikart et al., 2010), although true numbers are likely to remain low. Surprisingly, B. capensis exhibited the most restricted gene flow and highest signs of inbreeding (Table 1), despite showing the lowest range contraction (~1.3%), and broadest habitat tolerance, extending into dense thicket. The Albany thicket biome has experienced minimal loss (~8.9%) between 1750 and 2014 (Skowno et al., 2019), and total thicket vegetation has steadily increased across the Eastern Cape since 1950 (Njwaxu & Shackleton, 2019; Skowno et al., 2019; Stickler & Shackleton, 2014). But 63% of Albany thicket is severely degraded (14.0–25.4% valley thicket, and 12.8% mesic thicket) (Lloyd et al., 2002), and the extent of coastal thicket degradation is unknown. The cumulative effect of both native forest and dense thicket transformation has likely affected B. capensis more adversely than either P. stellata or C. dichroa, which probably use dense thicket only for facultative dispersal. Recuperation of the Albany thicket sub-biome augurs well for the population recovery of B. capensis, due to greater habitat availability, as well as the improved viability of P. stellata and C. dichroa. Improved landscape resistance modelling performance according to pairwise $D_{ps}$ for C. dichroa (Table 3) may reflect species recovery.

In east Africa, post-fragmentation sensitivity is evident for P. ruficapilla and P. stellata (Callens et al., 2011; Githiru & Lens, 2006;...
Sirén et al., 2018), yet the populations of both species in the Eastern Cape appear to have been largely resilient to historic forest exploitation. Afrotropical forest-dependent species have been observed to initially display stable effective population sizes following forest fragmentation (Husemann et al., 2015; Lens et al., 2002), but continued forest degradation eventually undermines population viability (Habel et al., 2014; Korfanta et al., 2012; Lens & Van Dongen, 1999). In South Africa, this is observable in both *P. ruficapilla* and *P. stellata* (Figure 2c).

### 4.3 Gene flow among regional forests

Metapopulation dynamics of these four songbird species do not appear to wholly contingent upon observed adult mobility. Observed regional adult vagility in *B. capensis* and *P. stellata* (Oschadleus & Ranwashe, 2017) did not preclude long-term (*F_{ST}*) isolation-by-distance in these species (Table 2). Local seasonal migration observed elsewhere in *C. dichroa* (Johnson & Maclean, 1994; Oatley et al., 2017; Oatley, 1966, 1969) is contested within southeast South Africa (Craig & Hulley, 2019; Wolmarans, 2015). Stronger population differentiation (Figure 5) and more recent (*D_{ps}* regional isolation-by-distance in this species indicates that regional populations of this species are likely sedentary. (Table 3). The apparent panmixia (Figure 5) and lack of isolation-by-distance (Table 2) observed within *P. ruficapilla* strongly indicates underestimated regional dispersal ability of this species.

However, philopatry—the tendency for vagile organisms to return to the same breeding location—has been observed in *P. stellata* (Dowsett, 1985; Oatley, 1982a) and *C. dichroa* (Oatley, 2017); strong adult site fidelity is observed in all four species (Callens et al., 2011; Habel et al., 2016; Oatley, 1982a; Oschadleus & Ranwashe, 2017) and attested to by recapture records (Oschadleus & Ranwashe, 2017). Adult philopatry (Habel et al., 2016; Oatley, 1982a) could mean that gene flow is mostly contingent upon natal dispersal of young birds (Gerrard et al., 2012). Natal dispersal of birds is poorly documented in South Africa, but has been observed in *P. stellata* (Oatley, 1982a), with 2- to 3-month-old individuals having been observed moving through plantations, woodland and riparian thicket. Intuitively, young *P. stellata* should seek to minimize exposure and attempt cost-efficient navigation of hospitable matrix vegetation, explaining the high performance of the LCP model of forest and coastal/mesic thicket configuration.

### 4.4 Species–landscape interactions

Landscape resistance modelling suggests that the configuration of both native forest, and dense (coastal/mesic/valley) thicket is important to gene flow in *B. capensis*, *C. dichroa* and *P. stellata* (Figures 4 and 7; Tables 2 and 3). *Pogonocichla stellata* demonstrates higher gene flow resistance through these thicket habitats compared to *B. capensis* and *C. dichroa* (Figure 7), potentially indicating that *P. stellata* disperses facultatively through select thicket habitats, while *B. capensis* and *C. dichroa* are more inclined to inhabit these habitats (especially mesic Albany thicket adjacent to native forests) in southeast South Africa (Johnson, 1997; Oatley, 1997a). Dispersing *P. ruficapilla* appear not to discriminate land-cover beyond forest configuration (Figure 4). This weak land-cover association could be due to a type 1 error derived from low sample size (Winarski et al., 2020), but the near panmixia within *P. ruficapilla* (Figure 5), and equilibrium between historic and contemporary gene flow (Figure 2b) supports the notion of high dispersal within this species, and tolerance towards anthropogenic landscape transformation.

The significant influence of regional elevation on the population structure of *B. capensis*, *P. ruficapilla* and *P. stellata* (Figure 7; Table 2) could indicate an elevation gradient to gene flow, potentially supporting altitudinal migration in *P. stellata*. For these three species, lower elevations appear more conducive to dispersal (Figure 7), with ravines, valleys and gorges serving as conduits into interior Afromontane forests. Outperformance of IBR over LCP elevation models respectively demonstrate inefficient navigation of landscape elevation, or interference by other landscape features. Rivers and dams appear to impact the gene flow of *B. capensis*, *P. ruficapilla* and *P. stellata* (Table 2). This landscape genetic association is similarly observed in a forest-associated pipistrelle (Moir et al., 2020).

### 4.5 Implications for regional Afromontane forest bird conservation

Stronger performance of the LCP models of native forest and dense thicket configuration over respective IBR models for *P. stellata* (Figure 7e), as well as *B. capensis* and *C. dichroa* (Tables 2 and 3) demonstrate the potential utility of conservation corridors in the Eastern Cape and southern KwaZulu-Natal to preserve genetic integrity within regional Afromontane forest birds. Such corridors should promote resilience anthropogenic climate change, as recommended by Colyn et al. (2020). The highest priority forests for conservation are the scarp forests present along the Wild Coast, as well the eastern Amatole Afromontane forests, and Afromontane (eastern mistbelt) forests in southern KwaZulu-Natal. These forests harbour the largest overall and unique genetic diversity (Table 1) and are therefore the most likely to serve as future climatic refugia. Effective creation of conservation corridors could incorporate forest and coastal/mesic thicket vegetation at lower elevations, particularly where these two land-cover classes coincide with rivers and dams, to ensure the preservation of optimal dispersal pathways beneficial for these four species. The utility of coastal thicket in facilitating movement of forest-dependent taxa is already recognized, and many authorities regard coastal thicket as secondary forest (Ehlers-Smith et al., 2018a, 2019; Ehlers-Smith, Ehlers-Smith, Ramesh, et al., 2017; Olivier et al., 2013). Beyond forest and dense thicket configuration, matrix land cover was not shown to impact the gene flow of these four songbird species significantly. Avian connectivity between
IOCB forests in KwaZulu-Natal can remain high across anthropogenically transformed areas, but minimally in forest-dependent insectivores (Neuschulz et al., 2013). Matrix transformation, however, can diminish regional South African forest ecological integrity (Botzat et al., 2015; Ehlers-Smith et al., 2019, 2020; Ehlers-Smith, Si, Ehlers-Smith, Kalle, et al., 2018; Freeman et al., 2018) and undermine the population viability of forest-dependent birds within forests.

5 | CONCLUSION

Our results show that reported forest-dependent range contractions in four insectivorous birds do not closely reflect species genetic responses to anthropogenic activity within the study area of the Eastern Cape and southern KwaZulu-Natal provinces of South Africa. The forest generalist B. capensis underwent the lowest range contraction (−1.3%), yet showed the most substantial gene flow restrictions, alongside pronounced declines in effective population size. More extensive range contractions observed in P. ruficapilla (−20.7%) and P. stellata (−23.0%) do not correspond to the comparatively stable effective population sizes observed, although gene flow restriction is evident in the latter species. Only the South African endemic forest specialist C. dichroa showed simultaneous declines in distribution (−19.5%) and effective population size, alongside gene flow disruption, and thus appears especially vulnerable to forest degradation.

In all four species, landscape resistance modelling suggested that regional gene flow within each of the four species is likely affected by landscape features. Native forest and dense thicket configuration is important to gene flow in P. stellata, B. capensis and C. dichroa, with B. capensis seeming most averse to thicket degradation. Beyond dense thicket, all four species, but particularly P. ruficapilla, do not facultatively disperse through matrix land cover. Finally, we propose that by conserving optimal dispersal routes through the two landcover classes, predominantly within low-elevation regions and coinciding with prominent river systems, should effectively ameliorate gene flow disruption and mitigate extinction debts culminating from historic forest exploitation.

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CONFLICT OF INTEREST

The authors declare no conflicts of interest.

DATA AVAILABILITY STATEMENT

Data for this study will be available at the Dryad Digital Repository: to be completed after manuscript is accepted for publication. Microsatellite genotyping: https://doi.org/10.5061/dryad.r4xgd7b.

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REFERENCES

Aben, J., Adriaensen, F., Thijs, K. W., Pellikka, P., Sijjander, M., Lens, L., & Matthysen, E. (2012). Effects of matrix composition and configuration on forest bird movements in a fragmented Afrotropical biodiversity hot spot. Animal Conservation, 15(6), 658–668. https://doi.org/10.1111/j.1469-1795.2012.00562.x
Aben, J., Strubbe, D., Adriaensen, F., Palmer, S. C. F., Travis, J. M. J., Lens, L., & Matthysen, E. (2014). Simple individual-based models effectively represent Afrotropical forest bird movement in complex landscapes. Journal of Applied Ecology, 51(3), 693–702. https://doi.org/10.1111/1365-2664.12224
Adie, H., Rushworth, I., & Lawes, M. J. (2013). Pervasive, long-lasting impact of historical logging on composition, diversity and above ground carbon stocks in Afrotropical forest. Forest Ecology and Management, 310, 887–895. https://doi.org/10.1016/j.foreco.2013.09.037
Adriaensen, F., Chardon, J. P., De Blust, G., Swinnen, E., Villalba, S., Guinick, H., & Matthysen, E. (2003). The application of "least-cost" modelling as a functional landscape model. Landscape and Urban Planning, 64(4), 233–247. https://doi.org/10.1016/S0169-2046(02)00242-6
Amos, J. N., Bennett, A. F., Mac Nally, R., Newell, G., Pavlova, A., Radford, J. Q., Thomson, J. R., White, M., & Snnucks, P. (2012). Predicting landscape-genetic consequences of habitat loss, fragmentation and mobility for multiple species of woodland birds. PLoS One, 7(2), e30888. https://doi.org/10.1371/journal.pone.0030888
Amos, J. N., Harrison, K. A., Radford, J. Q., White, M., Amos, J. N., Harrisson, K. A., Radford, J. Q., White, M., Newell, G., Nally, R. M., Sunnucks, P., & Pavlova, A. (2014). Species- and sex-specific connectivity effects of habitat fragmentation in a suite of woodland birds. Ecology, 95(6), 1556–1568. https://doi.org/10.1890/13-1328.1
Balkenhol, N., Cushman, S. A., Storfer, A. T., & Waits, L. P. (2015). Landscape genetics: Concepts, methods, applications. John Wiley and Sons Ltd.
Bates, D., Sarkar, D., Bates, M. D., & Matrix, L. (2014). lme4: Linear mixed-effects models using Eigen and 54 (R package version 1.1-6). http://cran.r-project.org/package=lme4
Belkhir, K., Borsa, P., Goudet, J., Chikhi, L., & Bonhomme, F. (2001). GENETIX 4.02, logiciel sous Windows TM pour la genetique des populations. Laboratoire Genome, Populations, Interactions, CNRS UMR 5000: Universite de Montpellier II, Montpellier (France). http://www.univ-montp2.fr/~genetix/genetix/genetix.htm
Benjamini, Y., & Hochberg, Y. (1995). Controlling the false discovery rate: A practical and powerful approach to multiple testing. Journal of the Royal Statistical Society: Series B (Methodological), 57(1), 289–300.
Berliner, D. (2009). Systematic conservation planning for South Africa’s forest biome: An assessment of the conservation status of South Africa’s forests and recommendations for conservation. University of Cape Town.
Berruti, A. (1997). Yellowthroated Warbler Geelkelsanger Phylloscopus ruficapilla. In J. A. Harrison, D. G. Allan, L. G. Underhill, M. Herremans, A. J. Tree, V. Parker & C. J. Brown (Eds.), The atlas of southern African birds. Vol. 2: Passerines (Vol. 2, pp. 256–257). BirdLife South Africa.
Winiarski, K. J., Peterman, W. E., & McGarigal, K. (2020). Evaluation of the R package 'resistancega': A promising approach towards the accurate optimization of landscape resistance surfaces. *Molecular Ecology Resources, 20*(6), 1583-1596. https://doi.org/10.1111/1755-0998.13217

Wolmarans, M. H. L. (2015). A flight of fancy in the Chorister Robin-Chat (*Cossypha dichroa*): An isotopic standpoint. Rhodes University. https://core.ac.uk/download/pdf/145047054.pdf

Zeller, K. A., McGarigal, K., & Whiteley, A. R. (2012). Estimating landscape resistance to movement: A review. *Landscape Ecology, 27*(6), 777-797. https://doi.org/10.1007/s10980-012-9737-0

Zhan, L., Paterson, I. G., Fraser, B. A., Watson, B., Bradbury, I. R., Nadukkalam Ravindran, P., Reznick, D., Beiko, R. G., & Bentzen, P. (2017). Megasat: Automated inference of microsatellite genotypes from sequence data. *Molecular Ecology Resources, 17*(2), 247-256. https://doi.org/10.1111/1755-0998.12561

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