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






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Genomic erosion across avian lineages in the context of their evolutionary history

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Abstract

Loss of genetic diversity threatens species survival, yet its dynamics and impacts can vary widely across species depending on their evolutionary histories, life-history traits, and demographic trajectories. To investigate these differences, we analyzed the genomes of 3 species that experienced extreme and well-documented population bottlenecks, the Mauritius parakeet, the Mauritius kestrel, and the pink pigeon, and compared them to 36 species spanning the avian phylogeny with varied IUCN Red List statuses. For each species, we assessed nucleotide diversity, genetic load, and inbreeding coefficients based on runs of homozygosity (F_{ROH}). We found a negative correlation between nucleotide diversity and F_{ROH} , but neither metric was a good predictor of the species' Red List status. Rather, the effective population size to census size ratio (N_e/N_c) showed a strong correlation to Red List status. Species with larger historical effective population sizes showed greater heterozygosity but carried a higher heterozygous load, highlighting the importance of historical demography for contextualizing species' vulnerability to genomic erosion. We also found significant differences in genetic load between taxonomic groups (parrots, pigeons, and falcons), possibly due to differences in life-history traits and demographic histories, underscoring the importance of interpreting genomic erosion dynamics in an evolutionary context. By anchoring our study on 3 evolutionarily divergent endangered species from Mauritius, we show how multispecies comparisons can contextualize extreme bottlenecks within a broader evolutionary framework, thereby identifying both general patterns of genomic erosion and species-specific vulnerabilities.

Keywords conservation genomics, genomic erosion, birds, Mauritius

Introduction

Genetic diversity is an essential component of a species' ability to adapt and persist under changing environmental conditions (Spielman et al. 2004; Allendorf et al. 2013; Kardos et al. 2021). For small or isolated populations, maintaining genetic diversity is particularly challenging, as reduced effective population size (N_e) diminishes the efficacy of natural selection, intensifies genetic drift, and leads to increased inbreeding. These processes ultimately lead to genomic erosion, characterized by reduced genetic diversity and the accumulation or fixation of deleterious mutations (van Oosterhout et al. 2022). As inbreeding becomes more frequent, recessive deleterious mutations are increasingly exposed, intensifying negative effects and leading to inbreeding depression (Charlesworth and Willis 2009; Blomqvist et al. 2010;

Hasselgren et al. 2021). These processes collectively reduce fitness and adaptive potential, heightening vulnerability to environmental changes and threatening the long-term persistence of the population (Blomqvist et al. 2010; Hasselgren et al. 2021; Jackson et al. 2022; van Oosterhout et al. 2022; Jeon et al. 2024). Additionally, even populations that have partially recovered demographically retain the genetic legacy of past bottlenecks, known as “drift debt,” which manifests as a time lag between population decline and loss of genome-wide variation (Gilroy et al. 2017; Dussex et al. 2023; Pinto et al. 2024). Recent analyses have shown that species are losing genetic diversity worldwide (Exposito-Alonso et al. 2022; Shaw et al. 2025), highlighting the need for cross-species comparisons to improve our understanding of genomic erosion in biodiversity conservation.

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With the continuous and rapid production of genomic data for wild species worldwide, conservation genomics can now take advantage of high-resolution tools to assess genetic diversity and genetic load (Lewin et al. 2018; Wright et al. 2020; van Oosterhout et al. 2022; Theissinger et al. 2023). For species at risk of extinction, such insights are critical for guiding conservation interventions aimed at reducing genetic load and enhancing population viability (e.g. vaquita (Morin et al. 2021), kākāpō (Dussex et al. 2021), pink pigeon (Speak et al. 2024)). Genome-wide comparisons can also provide insights into an endangered species' susceptibility to introgression from closely related species (Rieseberg 2001; Serrato-Capuchina and Matute 2018; Fontdevila 2019), and opportunities for genetic rescue (Whiteley et al. 2015; Bell et al. 2019). Additionally, using genomic resources across multiple species within a comparative framework may provide valuable insight into how evolutionary and demographic histories shape genomic patterns over both short and long timescales (Grueber 2015). This approach can elucidate how evolutionary history interacts with genomic traits, such as genetic diversity, genetic load, or structural variations, to influence species' long-term viability and extinction risk.

The Mauritius parakeet (*Alexandrinus [Psittacula] eques*), Mauritius kestrel (*Falco punctatus*), and pink pigeon (*Nesoenas mayeri*) exemplify how species can recover demographically from near extinction but remain genetically imperiled. These birds, endemic to Mauritius—a species-rich archipelago in the Indian Ocean that has witnessed over 100 species extinctions in recent centuries (Florens 2013)—experienced some of the most extreme population bottlenecks ever recorded in wild populations. Only 4 Mauritius kestrels remained by 1974, the pink pigeon declined to ~10 individuals by 1990, and the Mauritius parakeet to just ~20 individuals by 1986 (Jones and Swinnerton 1997; Jones 2013; Jones et al. 2013). Intensive conservation management facilitated their demographic recovery to current free-living adult population sizes of approximately 250 Mauritius kestrels, 500 pink pigeons, and 650 Mauritius parakeets (Jones and Swinnerton 1997; Jones 2010; Nicoll et al. 2021) (Fig. 1d). Although these species belong to evolutionarily distant lineages, they underwent similar demographic trajectories of collapse and recovery, providing independent yet comparable case studies to examine the genomic consequences of population decline and recovery across birds.

Despite these recoveries, the legacy of the extreme historical population collapses can jeopardize their long-term viability, as genetic diversity in these species continues to decline due to the accrued drift debt (Tollington et al. 2013; Jackson et al. 2022). These species are also at risk of accumulating an increased genetic load of deleterious mutations, as has been shown in the pink pigeon (Jackson et al. 2022). Beyond genetic challenges, ecological pressures such as habitat loss and degradation persist, compounded by threats like emerging infectious diseases, which can affect individual fitness and population viability (Tollington et al. 2015).

This study investigates the interplay between genome-wide diversity, genetic load, demographic history, and conservation status across a diverse set of avian species. Using recently generated, high-quality chromosome-level reference genomes for 3 focal Mauritian species—the Mauritius parakeet, pink pigeon, and Mauritius kestrel—we compared them to 36 species spanning the avian phylogeny. Placing these species in a broader phylogenetic context allowed us to distinguish general temporal trends from lineage-specific outcomes and to evaluate how such insights can

inform conservation assessments and prioritization. By further focusing on species within the same orders as the Mauritian taxa, we explored potential lineage-specific differences in genomic metrics to contextualize the genetic risk of the Mauritius species against the backdrop of their phylogenetic background and to inform conservation priorities for other vulnerable birds.

Materials and methods

Dataset

The reference genomes from 3 bird species endemic to Mauritius—the Mauritius parakeet (Morales et al. 2024a), the pink pigeon (Morales et al. 2024c), and the Mauritius kestrel (Morales et al. 2024b)—were recently sequenced and reported. All 3 genomes were generated from samples collected postbottleneck, from 2020 to 2021. To build a comparative dataset, 36 additional bird species were selected based on the availability of relatively high-quality reference genomes, prioritizing assemblies with higher scaffold N50, larger average scaffold size, and lower scaffold count within each taxonomic group, while also ensuring a comprehensive representation across the avian phylogeny (Fig. 1b). Although no strict thresholds were applied, the majority of selected genomes had scaffold N50 values >2 Mb and >85% of their genome contained within scaffolds >500 kb; a small number of species with slightly lower metrics were included to fill key phylogenetic gaps. Metadata for all 39 species, including current census population sizes, IUCN conservation statuses, and assembly statistics are compiled in Table S1.

Mapping and variant calling

Raw reads used for assembling the reference genomes were downloaded from NCBI (see Table S1 for Assembly IDs) and aligned to the corresponding genomes. NGS short reads were mapped using BWA (v0.7.17) mem (Li and Durbin 2009) with default parameters. Read duplicates were marked with GATK (4.4.0.0) MarkDuplicates (DePristo et al. 2011). PacBio HiFi reads were mapped and sorted using pbmm2 v1.5.0 (<https://github.com/PacificBiosciences/pbmm2>) with the parameter “-preset HIFI”. GATK HaplotypeCaller was used to call variants for each alignment. Only SNPs were kept for further analyses.

Depth

After mapping the raw reads to each reference genome, we estimated the average genome-wide and per-scaffold depth in each genome using MosDepth v0.3.3 (Pedersen and Quinlan 2018).

Sex chromosome removal

Each reference genome was mapped to the chicken genome (assembly GRCg6a) with minimap2 v2.1 (Li 2018). Any scaffold mapped to chicken sex chromosomes for more than 20% was treated as a potential region from sex chromosomes and removed for further analyses. Any additional scaffolds annotated in the reference genomes as sex chromosomes were also removed.

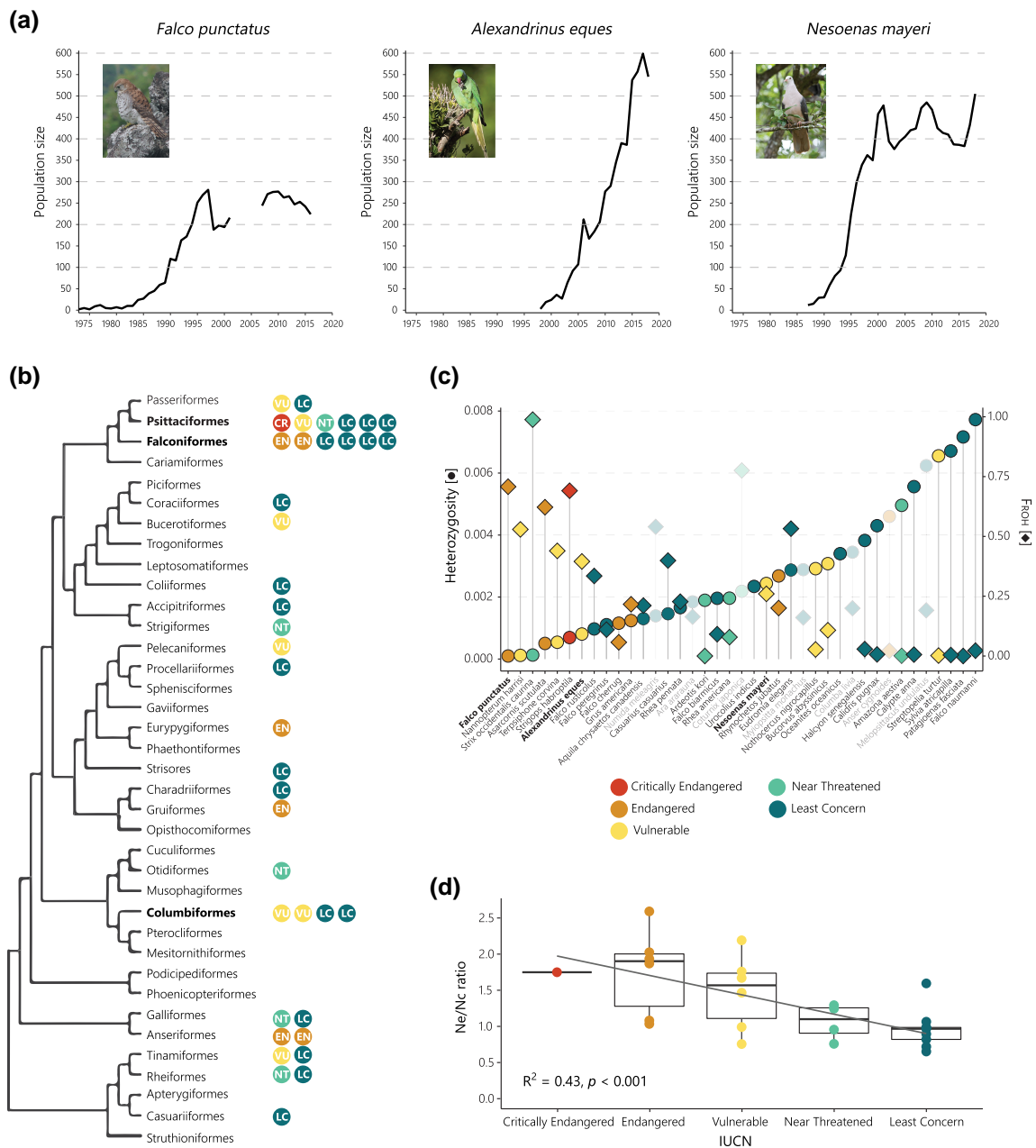


Figure 1 Demographic trajectory of the 3 Mauritius species and phylogenetic, genetic diversity, and Ne/Nc ratio distribution of the 39 species used in this study. a) Demographic trajectory of the wild population derived from field monitoring of adult individuals over time. Note that the 3 species have been monitored in different ways so presented trends are approximations of their total numbers. b) Phylogenetic relationship among the avian orders sampled in this study. Tree topology follows [Stiller et al. \(2024\)](#) and branch lengths are not time-calibrated and do not reflect divergence times. The orders of the 3 Mauritius species are marked in bold. Each circle represents a sampled species within its respective order. The color and initials indicate the IUCN Red List category of each species. c) Genome-wide heterozygosity (circles) and runs of homozygosity-based inbreeding coefficient (F_{ROH} ; diamonds) for each species. Color coding corresponds to IUCN Red List categories. Domestic species are denoted with lighter colors and in gray (outer line in the dots and in their name in x-axis). d) Correlation between the Ne/Nc ratio after log-transform ($\text{Log}(Ne)/\text{Log}(Nc)$) and IUCN Red List categories. The effective population size (Ne) was estimated as the harmonic mean of PSMC values from 10 to 100 kya, whereas the census population size (Nc) was obtained from IUCN Red List data. Photo credits: Samantha Cartwright for the Mauritius kestrel (*Falco punctatus*), Jacques de Speville for the Mauritius parakeet (*Alexandrinus eques*), and Gregory Guida for the pink pigeon (*Nesoenas mayeri*).

Heterozygosity

We estimated genome-wide heterozygosity using ANGSD ([Korneliussen et al. 2014](#)). We first obtained genotype likelihoods on scaffolds larger than 500 kb and only considered sites with a

depth of depth between $\frac{1}{3}$ (-setMinDepth) and 2 times (-setMaxDepth) the average depth for each sample. We assumed that the reference and ancestral states were the same. We applied the following parameters: -uniqueOnly 1 -remove_bads 1 -only_proper_pairs 1 -C 50 -baq 0 -minMapQ 30 -minQ 20

-setMinDepth \$minDP -setMaxDepth \$maxDP -doCounts 1 -GL 2 -doSaf 1. Next, we calculated the folded site frequency spectrum (SFS) with realSFS.

To compare different estimations of heterozygosity, we also estimated genome-wide heterozygosity directly from the VCF files with a custom pipeline. First, we divided scaffolds into sliding windows of size 100 kb (with a slide of 50 kb) using bedtools makewindows v2.30.0 (Quinlan and Hall 2010). Next, we obtained the total number of callable heterozygous and total callable genotypes per window using bcftools v1.20 (Danecek et al. 2021), tabix v1.14 (Li 2011), and vcftestcount from vcflib (Garrison et al. 2022). Genotypes were considered callable if their read depth was between $\frac{1}{3}$ and 2 times the average depth per sample and had a minimum genotype quality (GQ) of 30 or reference genome quality (RGQ) of 10, indels and multiallelic sites were excluded. Only nonmissing and quality-filtered sites (callable) were considered in each window and used as total callable genotypes in the denominator. Windows with less than 50% callable sites relative to the window size were removed for analysis.

Runs of homozygosity (ROH)

As some reference genomes were assembled with only Pacbio long reads, and no short-read data are available for the same individual, commonly used methods like ROHan (Renaud et al. 2019) could not be applied to identify ROHs. To address this limitation, we developed a custom method for this analysis. Given the varying fragmentation levels of the assemblies (Table S1), for each species, we retained only scaffolds with a minimum size of 5 Mb. As a result, 2 species (Red-faced mousebird, *Urocolius indicus* and Wilson's storm petrel, *Oceanites oceanicus*) were excluded. We identified ROH based on per-window heterozygosity estimates (see above). Using the R package bedtoolsr (Patwardhan et al. 2019), we concatenated windows with a heterozygosity lower than $5e^{-4} \text{ bp}^{-1}$ (Fig. S1), except for 2 genomes with exceptionally low average heterozygosity (the flightless cormorant, *Nannopterum harrisi*, and Mauritius kestrel), for which a lower threshold of $1e^{-4} \text{ bp}^{-1}$ was used. Adjacent homozygous regions were merged if separated by a gap shorter than 100 kb, and only ROHs with a minimum size of 500 kb were retained. The inbreeding coefficient (F_{ROH}) was calculated as the ratio of the total length of ROH segments to the total length of the analyzed genome (scaffolds > 5 Mb). We also estimated heterozygosity for the analyzed scaffolds per species, both including and excluding ROHs. To obtain the heterozygosity outside ROHs, we subtracted the windows that fall within a ROH for the estimation.

To validate our custom method, we compared our ROH estimates with those from ROHan for 5 species for which short-read data were available. ROHan was employed using a window size of 100 kb and a rohmu of $5e^{-4}$ to mimic the parameters used in the custom method.

Genetic load

To compare genetic load across species, we used Combined Annotation-Dependent Depletion (CADD) score (Kircher et al. 2014). CADD score integrates multiple genome annotations to rank the deleteriousness of any possible single-nucleotide

variant, including conservation metrics (e.g. GERP, Cooper et al. 2005), regulatory and transcript information, and protein-based scores (e.g. SIFT, Ng and Henikoff 2003). While CADD annotation of deleteriousness of mutations is generally consistent with regular functional annotation based on protein-coding information (Fontseré et al. 2025) such as SnpEff (Cingolani et al. 2012), CADD can also rank mutations in the introns and intergenic regions. We used the CADD score calculated for chicken genome (Groß et al. 2020) and transferred them to evolutionary conserved regions of the genome (ultraconserved elements, UCEs) in our target species. UCEs are highly conserved in vertebrates over a long evolutionary period, especially in birds (Cummins et al. 2024). Though mainly in nonprotein coding regions (introns or lncRNA genes), mutations in UCEs, especially with a high CADD score, are expected to have a strong deleterious effect (Speak et al. 2024). We extracted UCE regions and corresponding flanking regions from each reference genome and from the chicken genome (GRCg6a), and performed a multispecies alignment for each UCE, using the recommended pipeline of Phyluce v1.7.3 (Faircloth 2016) see Fig. S2 for detailed pipeline. Each genome was converted from fasta to 2bit format using UCSC FaToTwoBit (Casper et al. 2018). The UCE probe file was downloaded from <https://github.com/faircloth-lab/uce-probe-sets/tree/master/V1/uce-5k-probe-set>, and used to extract and validate UCE regions per species. The CADD score file of the chicken genome was downloaded from <https://osf.io/c97ez>. For each species, CADD scores were lifted from the chicken genome for homozygous (relative to chicken) and heterozygous sites across UCE regions with customized scripts partially adapted from LoadLift (Speak et al. 2024). Heterozygous sites were extracted from VCF files and filtered using bcftools with the same parameters used for heterozygosity estimations (see above). Only heterozygous sites with one allele equal to the corresponding site in the chicken genome were kept for further analyses.

To quantify genetic load in each genome, we counted the number of heterozygous sites and homozygous substitutions with CADD scores ≥ 20 in UCE regions, representing the top 1% most deleterious sites in the chicken genome. To control for the evolutionary distance to the chicken among species, we rescaled the counts of sites with CADD scores ≥ 20 by dividing them with the counts of substitutions with CADD scores < 3, the latter representing nearly-neutral sites. The rescaled ratios of heterozygous sites and homozygous substitutions were used as an inference of heterozygous and homozygous load. We distinguished these 2 classes because, according to theory (Bertorelle et al. 2022), deleterious homozygous positions are expressed, approximating the realized load, whereas (recessive) heterozygous positions are not expressed, approximating the masked load. However, in the absence of precise estimates of dominance coefficients, these are only approximations, and we therefore prefer to refer to them as the homozygous and heterozygous load.

To account for potential lineage-specific adaptive substitutions that have occurred since the divergence with chicken, homozygous substitutions that were shared by 20 or more species were excluded, as these are more likely to reflect long-term adaptive changes rather than harmful mutations. To further confirm that we were correctly retaining sites that have not changed since the divergence with chicken, we compared 3 species (saker falcon, *Falco cherrug*; budgerigar, *Melopsittacus undulatus*; and rock pigeon, *Columba livia*) included in the phylogenetic tree

by Feng et al. (2020), to their closest reconstructed ancestral nodes in the tree, which correspond to genus or family level. The command “hal2fasta” from HAL tools v2.3 (Hickey et al. 2013) was used to extract the sequences of the ancestral nodes from the genome-level alignment from this phylogenetic tree of 363 bird species. We extracted the reconstructed full sequences from the closest ancestral nodes and cut them into 200-bp short sequences with 20-bp step sliding windows. The reads were mapped to the chicken genome with BWA. For each species, we confirmed that homozygous substitutions had at least one ancestral read mapped to the chicken genome, and the ancestral state matched the chicken sequence. Homozygous substitutions with CADD scores above 20 are concentrated at the terminal branches of the phylogenetic tree of birds (Fig. S3), indicating that they are more likely to be deleterious substitutions than lineage-specific adaptations. Although we cannot rule out lineage-specific adaptive substitutions when measuring homozygous load, focusing on sites with CADD > 20 within the most conserved regions provides a higher likelihood of targeting truly deleterious variation (Rentzsch et al. 2019; Speak et al. 2024; Fontsero et al. 2025) (Fig. S3). As above, the counts of substitutions with CADD scores < 3, considered as nearly-neutral sites, were used to rescale the counts of sites with CADD scores ≥ 20 .

Demographic history

We inferred historical fluctuations of effective population size (N_e) for each species using PSMC (Li and Durbin 2011) with the parameters “-N30 -t5 -r5 -p 1+1+1+1+30*2+4+6+10” (Hilgers et al. 2025) and estimated the harmonic N_e mean from 10 to 100 kya for further analyses. The generation times and mutation rates used for PSMC can be found in Table S1. Generation times were retrieved from IUCN Red List (IUCN 2024), and the source of mutation rates were listed in Table S1.

Statistical and phylogenetic comparative analyses

To explore the relationships among genome-wide heterozygosity, F_{ROH} , N_e , and genetic load across species while accounting for phylogenetic signals, we reconstructed the phylogenetic relationships among the studied species and incorporated this inference into 2 statistical frameworks. The phylogeny was reconstructed from a subset of the UCE dataset described above (see Genetic Load section). Using the software AMAS (Borowiec 2016), we selected and concatenated 1,526 UCE sequences, retaining only loci with $\leq 2\%$ missing data and $> 30\%$ parsimony-informative sites. As previously published phylogenies did not include the target Mauritius species, we used IQ-TREE (Minh et al. 2020) to infer a maximum-likelihood tree using the edge-linked partition model (Chernomor et al. 2016) constraining deeper relationships based on the topology from Stiller et al. (2024). This approach allowed us to integrate our focal species into a well-supported, coalescent-informed avian phylogeny. Using this inferred phylogenetic tree, we assessed different univariate models with a phylogenetic generalized least squares (PGLS) approach (Freckleton et al. 2002). These analyses were conducted using the R packages *ape* (Paradis et al. 2004;

Paradis and Schliep 2019), *caper* (Orme et al. 2023) and *nlme* (Pinheiro et al. 2017). When PGLS indicated a nonzero lambda (λ) value, suggesting a significant phylogenetic signal, we further examined these effects using a Bayesian phylogenetic generalized linear mixed model (pGLMM) (Hadfield and Nakagawa 2010). In this framework, the phylogenetic relationship among species was modeled as a random effect. pGLMM analyses were performed using the R packages *ape* and *MCMCglmm* (Hadfield 2010).

Genomic synteny

We inferred multigenome synteny for chromosome-level reference genomes for pigeons (3 species in Columbidae), parrots (5 species in Psittaciformes), and falcons (6 species in *Falco*) separately using ntSynt v1.0.2 (Coombe et al. 2024) with the divergence range (-d) set to 10. Synteny results were visualized using scripts from ntSynt based on the R package *gggnomes* (Hackl et al. 2024).

Identification of transposable elements

To annotate repetitive elements (RE) in the genomes of the pink pigeon, Mauritius kestrel, and Mauritius parakeet, we produced de novo libraries of RE for each species using RepeatModeler2 (Flynn et al. 2020). We combined the de novo libraries with previously published manually curated libraries of RE from the Collared flycatcher (*Ficedula albicollis*) and Blue-capped cordon-bleu (*Uraeginthus cyanocephalus*) from Storer et al. (2021), and from the Emu (*Dromaius novaehollandiae*), Anna’s hummingbird (*Calypte anna*), and Kākāpō (*Strigops habroptilus*) from Peona et al. (2021). Using the resulting custom libraries, we annotated the RE from the genomes using RepeatMasker version 4.0.8 (Smit et al. 2015). We repeated this process for 3, 5, and 4 additional species of Columbiformes, Falconiformes, and Psittaciformes, respectively, to enable comparisons of proportions of RE across species.

Results

Genome-wide diversity and inbreeding

The sequencing depth across the dataset ranged between $15 \times$ and $96 \times$ (mean = 46, SD = 19). We estimated genome-wide heterozygosity with both genotype likelihoods in ANGSD and by SNP-calling, resulting in very similar estimates (adjusted $R^2 = 0.78$; Tables S1 and S2, Fig. S4). We used heterozygosity estimates from ANGSD for all subsequent analyses, except for the estimation of ROHs (see Material and Methods). Neither heterozygosity or F_{ROH} estimates showed significant correlation with the quality of the genomes (e.g. N50) or depth (Fig. S5). Our in-house method produced consistent ROH results to those from ROHan (Fig. S6), validating our approach.

Consistent with recent demographic change, genome-wide heterozygosity showed a strong negative correlation with inbreeding coefficient F_{ROH} (Fig. 1c; PGLS: $\lambda = 0$, $R^2 = 0.46$, $F_{1,28} = 26.56$, $P < 0.001$). Samples with domestic or pet origins (Fig. 1c, Table S1) are outliers with higher F_{ROH} compared to wild samples with similar levels of heterozygosity (Fig. 1c), and higher heterozygosity outside

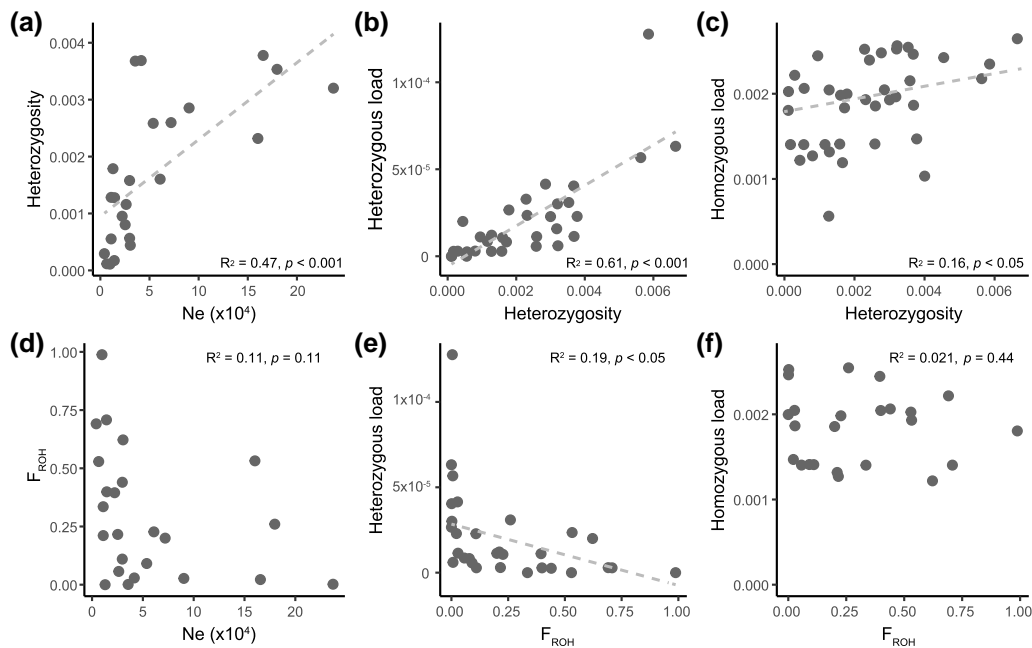


Figure 2 Comparison between genetic diversity metrics, genetic load, and effective population sizes. The dashed line represents the linear correlation when $P < 0.05$. a) Correlation between genome-wide heterozygosity and effective population size (Ne), with Ne estimated as the harmonic mean of PSMC values between 10 and 100 kya. b) Correlation between heterozygous load and genome-wide heterozygosity. Heterozygous load is defined as the ratio of heterozygous substitutions with CADD > 20 to homozygous substitutions with CADD < 3 . c) Correlation between homozygous load and genome-wide heterozygosity. Homozygous load is defined as the ratio of counts of filtered homozygous substitutions with CADD > 20 and the number of filtered homozygous substitutions with CADD < 3 . d) Correlation between inbreeding coefficient (based on runs of homozygosity; F_{ROH}) and Ne. e) Correlation between heterozygous load and inbreeding coefficient. f) Correlation between estimated homozygous load and inbreeding coefficient.

of ROH regions compared to genome-wide heterozygosity (Fig. S7); we therefore excluded them from further analyses. IUCN Red List status did not correlate with genome-wide heterozygosity, F_{ROH} (Fig. S8), or genetic load (Fig. S9).

Comparing historical Ne (between 10 and 100 kya estimated from PSMC) to current census size (Nc) reveals the magnitude of recent demographic change, with elevated values indicating more abrupt declines. The Ne/Nc ratio ($\text{Log}(\text{Ne})/\text{Log}(\text{Nc})$) was significantly lower (Wilcoxon two-sample test $P < 0.001$) in non-threatened species (least concern, mean = 0.91, SD = 0.23) compared to threatened species (remaining IUCN status categories, mean = 1.49), yet more varied (SD = 0.52). Ne/Nc ratio showed a strong linear correlation with IUCN status (Fig. 1d, GLM: $R^2 = 0.43$, $F_{1-24} = 12.49$, $P < 0.001$), when numerically coded from 0 critically endangered to 4 least concern. To disentangle the contribution of each component, we tested log-transformed Nc and Ne separately (Fig. S8). In these tests, only Nc was significant ($P = 0.0186$, $R^2 = 0.27$), whereas Ne was not. However, the Ne/Nc ratio explained more variance than either component alone ($P < 0.001$, $R^2 = 0.43$), indicating that the ratio captures additional, meaningful variation. Moreover, the Ne/Nc ratio reveals discrepancies that are not evident from Nc alone. For example, the pink pigeon and the whooping crane (*Grus americana*) both exhibit unusually high Ne/Nc ratios (2.19 and 2.59, respectively) compared to other species within the same IUCN categories (Vulnerable and Endangered, respectively). These elevated ratios point to a history of intense demographic decline despite currently recovered census sizes, underscoring that the severity of underlying genomic erosion is not apparent with Nc or Ne alone.

Historical demographic trends (Ne) have a significant positive correlation with genome-wide heterozygosity (Fig. 2a; PGLS: $\lambda = 0$, $R^2 = 0.47$, $F_{1-22} = 19.65$, $P = 2.1e^{-4}$). In contrast, F_{ROH} showed no significant correlation with historical Ne (Fig. 2d; PGLS: $\lambda = 0$, $R^2 = 0.11$, $F_{1-22} = 2.82$, $P = 0.11$). This lack of correlation is not surprising as the inbreeding coefficient is expected to have a nonlinear relationship with Ne (Reed et al. 2003), because inbreeding increases rapidly in small populations but approaches equilibrium asymptotically as Ne grows. Furthermore, our analysis focused only on long ROHs (> 500 Kb), which reflect population history within tens to hundreds of generations ago.

Species with higher genetic diversity or a lower inbreeding coefficient tended to carry a higher heterozygous load, calculated as the corrected ratio of heterozygous sites with a CADD score above 20 (Fig. 2b; PGLS: $\lambda = 0$, $R^2 = 0.61$, $F_{1-30} = 45.99$, $P = 1.6e^{-7}$, and Fig. 2e; PGLS: $\lambda = 0$, $R^2 = 0.19$, $F_{1-28} = 6.74$, $P = 0.015$). In contrast, estimated homozygous load, calculated as the corrected ratio of homozygous substitutions with a CADD score above 20, showed a statistically significant but weak association with heterozygosity (Fig. 2c; PGLS: $\lambda = 0.81$, $R^2 = 0.16$, $F_{1-30} = 5.61$, $P = 0.0245$), and no association with F_{ROH} (Fig. 2f; PGLS: $\lambda = 0.86$, $R^2 = 0.02$, $F_{1-28} = 0.61$, $P = 0.4424$). Because $\lambda \approx 0.8$ indicated phylogenetic signal in the residuals, we refitted these models using a phylogenetic GLMM (pGLMM) with a random effect for shared ancestry. This model recovered the same fixed-effect pattern, as heterozygosity remained significant and F_{ROH} nonsignificant, but revealed that phylogenetic variance was negligible compared to residual variance ($\sigma^2_{\text{phylo}} \approx 4.1 \times 10^{-14}$; $\sigma^2_{\text{species}} \approx 1.0 \times 10^{-14}$, Table S3). Thus, although residuals appeared phylogenetically structured in the PGLS, shared ancestry explained little additional variation once predictors were included, indicating that phylogenetic

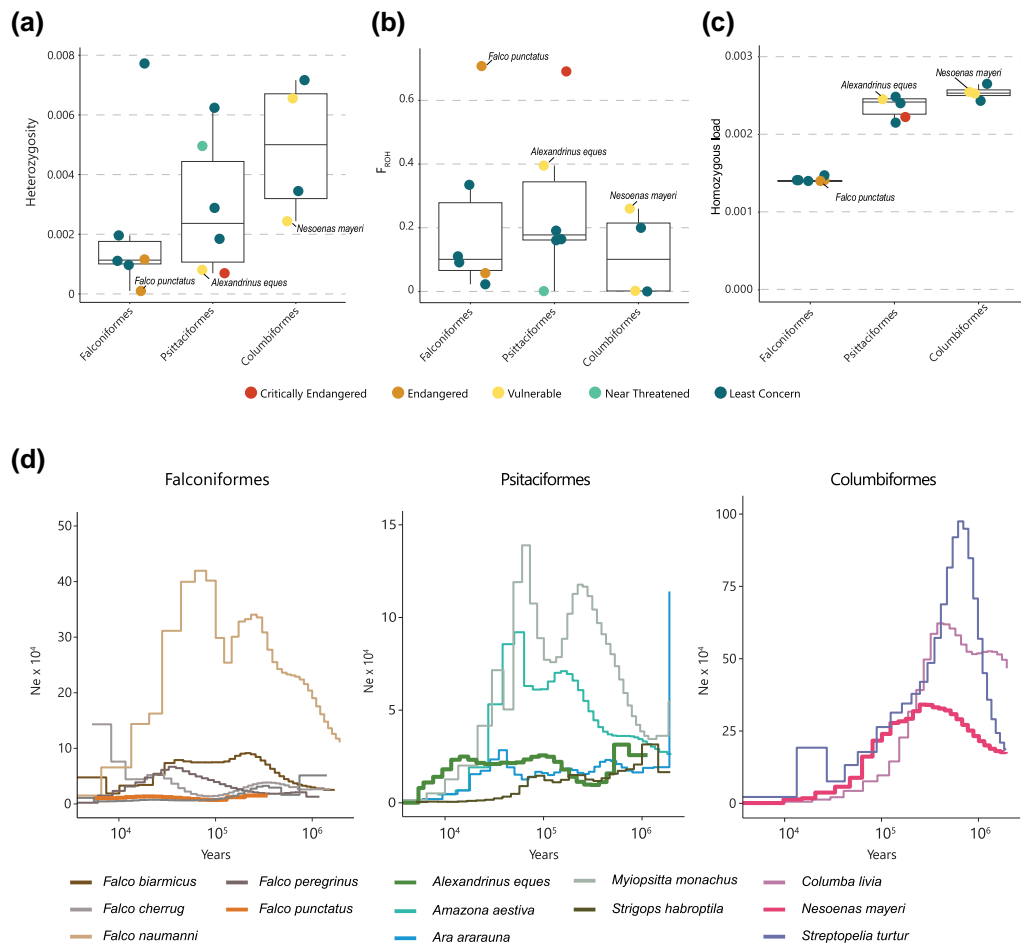


Figure 3 Genetic diversity, genetic load, and demographic history of Falconiformes, Psittaciformes, and Columbiformes. a) Genome-wide heterozygosity in $\text{het} \times \text{bp}^{-1}$, b) inbreeding coefficient, and c) homozygous load distribution across the orders of the 3 target species. The inbreeding coefficient (F_{ROH}) was estimated using runs of homozygosity, and homozygous load was based on the ratio of homozygous substitutions with CADD scores above 20 to those with CADD scores below 3. d) Demographic histories are shown as variation in N_e (effective population size) inferred with PSMC, and a log-scaled x-axis shows years before present. The falcons had the lowest historical N_e (5.0×10^4 , harmonic mean for 10 to 100 kya) on average, though *Falco naumanni* is higher. The parrots show more variable, generally higher N_e (7.6×10^4), with *Myiopsitta monachus* and *Amazona aestiva* peaking. The pigeons had the highest N_e (2.3×10^5), led by *Streptopelia turtur*. Thick lines refer to Mauritius species. Only species with chromosomal-level assemblies were included.

relatedness is not a major determinant of homozygous load variation. Although statistically significant, the association between homozygous load and heterozygosity remained weak, and the absence of a positive correlation between homozygous load and F_{ROH} was unexpected (Kyriazis et al. 2025). This pattern may reflect the influence of deep-time factors such as lineage-specific mutation rates, adaptive substitutions, or life-history differences. Moreover, because only one individual per species was analyzed, it remains difficult to distinguish fixed substitutions accumulated through deep phylogenetic time from genuinely deleterious homozygous alleles (see Discussion).

Genomic features within and across taxonomic groups

Despite similar distribution ranges and histories of population decline in the past decades (Fig. 1a), the 3 Mauritius species

showed contrasting levels of genetic heterozygosity, inbreeding coefficients (Figs. 1c, 3a, b), and homozygous load (Fig. 3c). The Mauritius kestrel had the lowest heterozygosity of all samples ($8.33 \times 10^{-5} \text{ het} \times \text{bp}^{-1}$) and the second highest F_{ROH} (0.71) among the wild species included in this study, with 50% of its genome in very long ROHs ($>10 \text{ Mb}$), as evidence of sustained recent inbreeding after recovering demographically from a bottleneck of only 4 individuals. The Mauritius parakeet had the second lowest heterozygosity ($8.07 \times 10^{-4} \text{ het} \times \text{bp}^{-1}$) among parrots, closely following another extremely bottlenecked species, the critically endangered Kākāpō (*Strigops habroptilus*). However, the Mauritius parakeet's F_{ROH} , while high (0.40), is lower than that of the Kākāpō (0.69), with 24.7% of the Mauritius parakeet's genome in ROHs longer than 10 Mb, evidence of their extreme bottleneck of ~ 12 individuals. The pink pigeon exhibits a heterozygosity of $2.38 \times 10^{-3} \text{ het} \times \text{bp}^{-1}$, the lowest among the analyzed pigeons, but higher than more than half of the species included in the study and nearly 30 times greater than the

Mauritius kestrel. Additionally, the pink pigeon showed an F_{ROH} of 0.26, with 12.3% of its genome in ROHs of a length longer than 10 Mb (Fig. S7).

Reflecting a deeper evolutionary process, the differences in genetic diversity of Mauritius species were associated with the differences between their taxonomic groups (Fig. 3). Falcons exhibit lower heterozygosity (one-sided Wilcoxon test $P=0.07$) than the other 2 taxonomic groups. Likewise, falcons carry 23.7% less homozygous load than the parrots ($P=0.001$) and 29.7% less than the pigeons ($P=0.005$). No significant difference in F_{ROH} was found between taxonomic groups. Within their respective taxonomic groups, the 3 Mauritius species showed the lowest genome-wide heterozygosity (Fig. 3a) and the highest F_{ROH} (Fig. 3b). Genetic diversity estimates carry the signal of ancestral population size, as the 3 Mauritius species had relatively low population sizes within their respective taxonomic groups (Fig. 3d). However, the pink pigeon had a larger historical population than most studied species, including falcons and parrots, which is reflected in its higher heterozygosity compared to the average levels in the other taxonomic groups (Fig. 3a). This reveals the importance of considering genetic diversity within the context of a species' long-term evolutionary history and taxonomic group. Note that all species that we have included from Falconiformes are from the same genus, *Falco*, with a divergence time of 12 million years (Kumar et al. 2022), which could explain the small deviations for heterozygosity and homozygous load, whereas the divergence time of the study Columbiformes and Psittaciformes species was roughly 16 and 55.6 million years, respectively (Fig. S10).

We examined synteny and repetitive element content in the 3 Mauritius species to ensure that their differences did not confound interspecific comparisons. All exhibited conventional avian genome structures, with conserved synteny to their closest relatives (Fig. S10) and typical repetitive element proportions (15% to 20%; Fig. S11; Hughes and Piontkivska 2005), consistent with previous reports of lower repeat content in birds relative to mammals.

Discussion

By analyzing 39 avian genomes spanning diverse taxonomic groups, we show that the dynamics of genetic diversity, genetic load, and inbreeding are shaped by both recent demographic changes and deep evolutionary history. While the genomic effects of recent population decline have been well documented (van der Valk, Diez-del-Molino, et al. 2019; Hasselgren et al. 2021; Khan et al. 2021; Cavill et al. 2024), we show how deep demographic history also exerts a long-lasting influence on patterns of genomic erosion. As a result, species from different taxonomic groups exhibit distinct levels of heterozygosity and genetic load, underscoring the need to account for both demographic history and phylogenetic context when comparing genomic metrics. Using 3 Mauritian species as focal case studies, we illustrate how comparative analyses can uncover conservation-relevant patterns that would be overlooked in single-species studies. Despite challenges in standardizing genomic metrics across diverse taxa, our findings demonstrate how evolutionary history constrains the interpretation of genomic metrics widely used in conservation, highlighting the value of comparative frameworks for advancing biodiversity conservation.

Conservation status is shaped by recent and long-term history

Genetic diversity has been considered a classical indicator for population resilience and risk of extinction (Breed et al. 2019; DeWoody et al. 2021; Teixeira and Huber 2021; Jeon et al. 2024). Consistent with this expectation, we observed a negative correlation between genome-wide heterozygosity (a proxy of genetic diversity) and F_{ROH} (a proxy of inbreeding) (Fig. 1c), reflecting intensified inbreeding and genetic drift in recently small populations (Brüniche-Olsen et al. 2018; Grossen et al. 2020). Reference genomes sequenced from domestic or pet samples had elevated F_{ROH} , despite relatively high heterozygosity outside ROHs, highlighting how recent inbreeding can bias genome-wide heterozygosity estimates and the importance of checking sample origin when using public genomic data (Fig. S7).

Over longer evolutionary timescales (10,000 to 100,000 years ago), genetic diversity was strongly correlated with historical population size (Fig. 2a), indicating that present-day genomic variation continues to reflect ancient demographic history. This suggests that species' genetic susceptibility to future habitat and environmental change is associated with its ancient demographic history, even prior to the accelerated recent environmental changes induced by human activity (Tan et al. 2023). F_{ROH} , on the other hand, did not show a strong correlation with the historical population size (Fig. 2d), as it should instead reflect the effect of recent demographic change (Ceballos et al. 2018). Consequently, joint monitoring of demographic, environmental, and genetic change—using complementary genomic metrics—is essential for evaluating short- and long-term risks. This is particularly important for species with historically low genetic diversity, as they may remain more vulnerable to future environmental shifts even if current population sizes appear stable (Ellstrand and Elam 1993; van der Valk, de Manuel et al. 2019; Brüniche-Olsen et al. 2021; Liu et al. 2025; Willi et al. 2006).

Data on modern census population sizes (N_c) are undoubtedly crucial in conservation assessments (Shaffer 1981; Lande 1988; Willi et al. 2006; Frankham et al. 2014), with the rate of demographic decline being one of the major factors considered by the IUCN Red List rating (McNeely et al. 1990; Frankham et al. 2014). However, the N_e/N_c ratio, reflecting the balance between long-term genetic diversity and current population size, can also have a prominent role in conservation assessments (Frankham 1995; Kalinowski and Waples 2002). Elevated N_e/N_c ratios based on current N_e likely reflect genetic erosion that has already occurred, while elevated ratios based on historical N_e reflect recent demographic declines that have not yet resulted in a proportional loss of genome-wide diversity. The latter ratio reflects the time-lag between population decline and genetic diversity loss (Gargiulo et al. 2025), resulting from the drift debt (Gilroy et al. 2017; Dussex et al. 2023; Pinto et al. 2024; Liu et al. 2025), and serves as an early warning sign of an imminent population collapse (Amos and Balmford 2001; Wilder et al. 2023). In this study, we use historical N_e estimated via PSMC, enabling a comparison across many species without the need for population-level sampling. We find that elevated N_e/N_c ratios are associated with higher IUCN risk categories (Fig. 1d). Although the correlation between IUCN status and N_e/N_c ratio is mainly driven by N_c (Fig. S8), as IUCN rating is partly based on current population sizes,

the N_e/N_c ratio has a stronger correlation to IUCN status than N_c alone (Fig. S8), suggesting that it captures additional demographic-genetic dimensions. Notably, species such as the pink pigeon and whooping crane exhibit unusually high N_e/N_c ratios relative to their IUCN category, revealing discrepancies that may signal severe ongoing genomic erosion, as a result of a sharp historical demographic decline despite demographic recovery. This highlights the value of integrating genomic data into conservation assessments to better capture long-term genetic threats, as previously discussed for these species (Jackson et al. 2022; Fontseré et al. 2025). One potential caveat is the uncertain sampling time of the biological materials used to develop the reference genomes, which could bias our analyses of IUCN status, since 9 out of 39 species had status changes within the past 20 years. Our findings indicate that the N_e/N_c ratio, even when based on historical N_e , is not only a meaningful indicator of conservation status (Fig. 1d) but also a potential flag for discrepancies between genomic erosion and current IUCN status. This underscores the importance of understanding and incorporating long-term demographic history into conservation assessments to better capture latent genetic risks.

Interpreting genetic load patterns across species

Our results highlight that the relationship between diversity and load must be interpreted in light of evolutionary timescale and demographic context. We found a strong positive correlation between the relative number of heterozygous deleterious sites—a proxy for masked load—and genome-wide heterozygosity (Fig. 2b). Therefore, similarly to genetic diversity, masked load is also shaped by demographic history. Population decline often leads to the expression of masked genetic load, driven by genetic drift and reduced purging (van der Valk, Diez-del-Molino, et al. 2019; Dussex et al. 2023). With habitat loss predicted to intensify in the near future, resulting in accelerated population declines and loss of genetic diversity (Exposito-Alonso et al. 2022), species with currently higher diversity may face rapid exposure of these deleterious mutations before effective purging can occur, increasing the risk of fitness reductions and jeopardizing population viability (van Oosterhout et al. 2022). Therefore, estimating heterozygous load remains highly informative, as it integrates demographic and selective history and may ultimately help predict population fitness once empirical links are established.

Our results reveal that species with lower genetic diversity exhibit reduced homozygous load (Fig. 2c). This contrasts population-genetics expectations that small, low-diversity populations accumulate more deleterious mutations through genetic drift (Kimura et al. 1963; Bertorelle et al. 2022; Robinson et al. 2023), and with empirical studies (Khan et al. 2021; Wang et al. 2023), based on standing variation under constant selection coefficients. Several factors may explain this apparent discrepancy.

First, our estimates rely on single individuals per species, which limits the resolution of polymorphic deleterious alleles and sensitivity to recent demographic effects. As we are comparing single genomes per species spanning >60 million years of avian evolution, our estimates of homozygous load primarily reflect long-term evolutionary accumulation of substitutions in conserved regions, a mixture of mildly deleterious and

lineage-specific adaptive changes (Grossen et al. 2020; Dussex et al. 2023; Kardos et al. 2023; Wang et al. 2023). The difficulty of distinguishing between these categories likely weakens cross-species correlations with diversity and inbreeding. Improved ancestral state inference and population-level sampling will be required to disentangle fixed deleterious substitutions from adaptive changes and to accurately quantify load within species.

Moreover, differences in life-history traits may modulate the rate of deleterious allele accumulation. Species with long generation times experience fewer generations over comparable evolutionary periods, potentially slowing the accumulation of weakly deleterious alleles, whereas species with short generation times but large population sizes may maintain more efficient purifying selection. However, these relationships are complex, as generation time often covaries with population size and other ecological factors (Chao and Carr 1993; Sæther et al. 2005; Plough 2016).

Finally, because our lifted-over CADD annotations are based on noncoding regions ultraconserved across birds, they provide reliable indicators of evolutionary constraint, but the direct functional consequences of mutations in these regions remain incompletely understood. Therefore, our estimates should be interpreted as comparative indicators of evolutionary constraint rather than as absolute measures of fitness impact.

Lineage-specific patterns across taxonomic groups

The 3 Mauritian species exhibit low genetic diversity and high inbreeding (F_{ROH}) within their respective taxonomic groups, consistent with their severe population bottlenecks. These metrics highlight an ongoing risk of inbreeding depression despite their successful conservation management (Jones et al. 1995; Jones and Swinnerton 1997; Tollington et al. 2013; Nicoll et al. 2021). Although absolute levels of heterozygosity and homozygous load differ among the 3 species, the similarly high F_{ROH} values indicate comparable levels of recent inbreeding across the Mauritian taxa (Fig. 3b), suggesting the need for future population monitoring and management focused on reducing the risk of inbreeding.

Placing these patterns in a broader evolutionary context, the contrasts among taxonomic groups reveal the interplay between long-term demographic history and deep-time evolutionary processes. Falcons as a group show consistently low heterozygosity and homozygous load, whereas parrots and pigeons display higher overall genomic diversity. This pattern mirrors variation in ancestral effective population size inferred from PSMC trajectories (Fig. 3d), indicating that ancient demography has left a strong imprint on present-day diversity. By contrast, F_{ROH} estimates based on long ROHs show similar values across taxonomic groups (Fig. 3b, pairwise Wilcoxon tests were not significant), as these capture recent inbreeding rather than long-term demographic history.

Life-history traits likely mediate these taxonomic differences. Parrots have long generation times, low reproductive rates, and high parental investment (Jones and Swinnerton 1997; Jones 2010; Jones et al. 2013), making them particularly vulnerable to genomic erosion. These traits slow the recovery of genetic diversity after bottlenecks and exacerbate the accumulation of homozygous load. In contrast, pigeons have shorter generation

times and higher reproductive rates (Jones 2013), facilitating faster recovery and preserving higher genetic diversity despite similar population collapses. Falcons exhibit intermediate traits, with low reproductive rates but shorter generation times and the ability to disperse to new environments (Jones et al. 1995; Cartwright et al. 2014; Nicoll et al. 2021), which can limit genetic drift and inbreeding but may not fully mitigate the effects of historically small population sizes.

Together, these results illustrate that genomic erosion cannot be interpreted outside its phylogenetic and life-history context, and they highlight the need to deepen our understanding of how life-history traits influence genetic diversity (Germain et al. 2023) and load.

Future directions to benefit conservation genomics

As whole-genome data become more accessible (Feng et al. 2020; Stiller et al. 2024), the integration of genomic-derived metrics (e.g. demographic reconstructions, heterozygosity, F_{ROH} , N_e/N_c) with demographic, ecological, and environmental data will substantially improve conservation assessments and planning. Expanding the availability and taxonomic breadth of reference genomes will further increase the utility of genomic resources in conservation biology (Grueber 2015; Supple and Shapiro 2018; Mc Cartney et al. 2024). Given the substantial differences of genome-wide features observed between groups (Fig. 3, Figs. S10, S11), it is advantageous, even in the absence of a species-specific reference genome, to identify a closely related reference genome to minimize mapping bias and improve inference accuracy (Prasad et al. 2022). However, population-level data remain essential, as a single individual cannot fully represent the genetic diversity of an entire species. Incorporating population-level data also allows for more robust estimates of realized genetic load by leveraging site frequency distributions (Grossen et al. 2020; Bertorelle et al. 2022).

Although the importance of genetic diversity is well established, understanding the fitness effects of deleterious alleles is central to predicting species' adaptive potential and persistence (Kardos et al. 2021). Future progress will require combining genomic data with direct fitness measures, temporal datasets tracking load dynamics, and improved methods to identify deleterious mutations (Bosse et al. 2019; van der Valk, de Manuel, et al. 2019; Bertorelle et al. 2022; Fontseré et al. 2025).

Integrating demographic history changes is also critical for interpreting genetic diversity trends. While genomic inference with modern samples reveals population history in both the long term (N_e) and short term (recent inbreeding with F_{ROH}), historical genetic data provides critical predecline information in accurately assessing trends in population size and genetic load in a more recent, crucial time scale (van der Valk, de Manuel, et al. 2019; Femerling et al. 2023; Cavill et al. 2024; Dehasque et al. 2024; Silver et al. 2024; Fontseré et al. 2025). Such data are invaluable for identifying rapid losses of genetic diversity and increases in load, which may otherwise go undetected until demographic impacts become severe (Diez-del-Molino et al. 2018).

Although the relationships among genetic diversity, demographic history, and deleterious variation are well established

in theory, our findings underscore the importance of considering evolutionary history, demographic processes, and life-history traits when applying genetic measures in conservation. Our comparative analyses indicate that low genetic diversity or high load does not carry equivalent implications across taxa, and that evolutionary context is essential for interpreting genomic risk. Comparative analyses across lineages provide a unique chance to evaluate how recent demographic collapse interacts with deep-time history, helping to refine expectations for genetic recovery and to better contextualize genomic data for guiding future conservation strategies in an era of rapid environmental change.

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Supplementary material

Supplementary material is available at *Molecular Biology and Evolution* online.

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Conflicts of interest

None declared.

Data availability

The scripts used in this study are available on GitHub: [PachecoMC/CompConGen](https://github.com/PachecoMC/CompConGen).

References

Allendorf FW, Luikart G, Aitken SN, Antunes A. *Conservation and the genetics of populations*. 2nd ed. John Wiley & Sons; 2013.

- Amos W, Balmford A. When does conservation genetics matter? *Heredity (Edinb)*. 2001;87:257–265. <https://doi.org/10.1046/j.1365-2540.2001.00940.x>.
- Bell DA *et al*. The exciting potential and remaining uncertainties of genetic rescue. *Trends Ecol Evol*. 2019;34:1070–1079. <https://doi.org/10.1016/j.tree.2019.06.006>.
- Bertorelle G *et al*. Genetic load: genomic estimates and applications in non-model animals. *Nat Rev Genet*. 2022;23:492–503. <https://doi.org/10.1038/s41576-022-00448-x>.
- Blomqvist D, Pauliny A, Larsson M, Flodin L-Å. Trapped in the extinction vortex? Strong genetic effects in a declining vertebrate population. *BMC Evol Biol*. 2010;10:33. <https://doi.org/10.1186/1471-2148-10-33>.
- Borowiec ML. AMAS: a fast tool for alignment manipulation and computing of summary statistics. *PeerJ*. 2016;4:e1660. <https://doi.org/10.7717/peerj.1660>.
- Bosse M, Megens H-J, Derks MFL, de Cara Ángeles MR, Groenen MAM. Deleterious alleles in the context of domestication, inbreeding, and selection. *Evol Appl*. 2019;12:6–17. <https://doi.org/10.1111/eva.12691>.
- Breed MF *et al*. The potential of genomics for restoring ecosystems and biodiversity. *Nat Rev Genet*. 2019;20:615–628. <https://doi.org/10.1038/s41576-019-0152-0>.
- Brüniche-Olsen A, Kellner KF, Anderson CJ, DeWoody JA. Runs of homozygosity have utility in mammalian conservation and evolutionary studies. *Conserv Genet*. 2018;19:1295–1307. <https://doi.org/10.1007/s10592-018-1099-y>.
- Brüniche-Olsen A, Kellner KF, Belant JL, DeWoody JA. Life-history traits and habitat availability shape genomic diversity in birds: implications for conservation. *Proc R Soc B Biol Sci*. 2021;288:20211441. <https://doi.org/10.1098/rspb.2021.1441>.
- Cartwright SJ, Nicoll MAC, Jones CG, Tatayah V, Norris K. Anthropogenic natal environmental effects on life histories in a wild bird population. *Curr Biol*. 2014;24:536–540. <https://doi.org/10.1016/j.cub.2014.01.040>.
- Casper J *et al*. The UCSC Genome Browser database: 2018 update. *Nucleic Acids Res*. 2018;46:D762–D769. <https://doi.org/10.1093/nar/gkx1020>.
- Cavill EL *et al*. When birds of a feather flock together: severe genomic erosion and the implications for genetic rescue in an endangered island passerine. *Evol Appl*. 2024;17:e13739. <https://doi.org/10.1111/eva.13739>.
- Ceballos FC, Joshi PK, Clark DW, Ramsay M, Wilson JF. Runs of homozygosity: windows into population history and trait architecture. *Nat Rev Genet*. 2018;19:220–234. <https://doi.org/10.1038/nrg.2017.109>.
- Chao L, Carr DE. The molecular clock and the relationship between population size and generation time. *Evolution*. 1993;47:688–690. <https://doi.org/10.1111/j.1558-5646.1993.tb02124.x>.
- Charlesworth D, Willis JH. The genetics of inbreeding depression. *Nat Rev Genet*. 2009;10:783–796. <https://doi.org/10.1038/nrg2664>.
- Chernomor O, von Haeseler A, Minh BQ. Terrace aware data structure for phylogenomic inference from supermatrices. *Syst Biol*. 2016;65:997–1008. <https://doi.org/10.1093/sysbio/syw037>.
- Cingolani P *et al*. A program for annotating and predicting the effects of single nucleotide polymorphisms, SnpEff. *Fly (Austin)*. 2012;6:80–92. <https://doi.org/10.4161/fly.19695>.
- Coombe L, Kazemi P, Wong J, Birol I, Warren RL. 2024 February 13. Multi-genome synteny detection using minimizer graph mappings [preprint]. bioRxiv 579356. <https://doi.org/10.1101/2024.02.07.579356>
- Cooper GM *et al*. Distribution and intensity of constraint in mammalian genomic sequence. *Genome Res*. 2005;15:901–913. <https://doi.org/10.1101/gr.3577405>.
- Cummins M, Watson C, Edwards RJ, Mattick JS. The evolution of ultraconserved elements in vertebrates. *Mol Biol Evol*. 2024;41:msae146. <https://doi.org/10.1093/molbev/msae146>.
- Danecek P *et al*. Twelve years of SAMtools and BCFtools. *Gigascience*. 2021;10:giab008. <https://doi.org/10.1093/gigascience/giab008>.
- Dehasque M *et al*. Temporal dynamics of woolly mammoth genome erosion prior to extinction. *Cell*. 2024;187:3531–3540.e13. <https://doi.org/10.1016/j.cell.2024.05.033>.
- DePristo MA *et al*. A framework for variation discovery and genotyping using next-generation DNA sequencing data. *Nat Genet*. 2011;43:491–498. <https://doi.org/10.1038/ng.806>.
- DeWoody JA, Harder AM, Mathur S, Willoughby JR. The long-standing significance of genetic diversity in conservation. *Mol Ecol*. 2021;30:4147–4154. <https://doi.org/10.1111/mec.16051>.
- Díez-del-Molino D, Sánchez-Barreiro F, Barnes I, Gilbert MTP, Dalén L. Quantifying temporal genomic erosion in endangered species. *Trends Ecol Evol*. 2018;33:176–185. <https://doi.org/10.1016/j.tree.2017.12.002>.
- Dussex N *et al*. Population genomics of the critically endangered kākāpō. *Cell Genom*. 2021;1:100002. <https://doi.org/10.1016/j.xgen.2021.100002>.
- Dussex N, Morales HE, Gossen C, Dalén L, van Oosterhout C. Purging and accumulation of genetic load in conservation. *Trends Ecol Evol*. 2023;38:961–969. <https://doi.org/10.1016/j.tree.2023.05.008>.
- Ellstrand NC, Elam DR. Population genetic consequences of small population size: implications for plant conservation. *Annu Rev Ecol Evol Syst*. 1993;24:217–242. <https://doi.org/10.1146/annurev.es.24.110193.001245>.
- Exposito-Alonso M *et al*. Genetic diversity loss in the anthropocene. *Science*. 2022;377:1431–1435. <https://doi.org/10.1126/science.abn5642>.
- Faircloth BC. PHYLUCS is a software package for the analysis of conserved genomic loci. *Bioinformatics*. 2016;32:786–788. <https://doi.org/10.1093/bioinformatics/btv646>.
- Femerling G *et al*. Genetic load and adaptive potential of a recovered avian species that narrowly avoided extinction. *Mol Biol Evol*. 2023;40:msad256. <https://doi.org/10.1093/molbev/msad256>.
- Feng S *et al*. Dense sampling of bird diversity increases power of comparative genomics. *Nature*. 2020;587:252–257. <https://doi.org/10.1038/s41586-020-2873-9>.
- Florens FBV. Conservation in Mauritius and Rodrigues: challenges and achievements from two ecologically devastated oceanic islands. In: Raven PH, Sodhi NS, Gibson L, editors. *Conservation biology*. 1st ed Wiley; 2013. p. 40–50.
- Flynn JM *et al*. RepeatModeler2 for automated genomic discovery of transposable element families. *Proc Natl Acad Sci U S A*. 2020;117:9451–9457. <https://doi.org/10.1073/pnas.1921046117>.

- Fontdevila A. Hybrid genome evolution by transposition: an update. *J Hered.* 2019;110:124–136. <https://doi.org/10.1093/jhered/esy040>.
- Fontserè C *et al.* Persistent genomic erosion in whooping cranes despite demographic recovery. *Mol Ecol.* 2025;34:e70088. <https://doi.org/10.1111/mec.70088>.
- Frankham R. Effective population size/adult population size ratios in wildlife: a review. *Genet Res.* 1995;66:95–107. <https://doi.org/10.1017/S0016672300034455>.
- Frankham R, Bradshaw CJA, Brook BW. Genetics in conservation management: revised recommendations for the 50/500 rules, Red List criteria and population viability analyses. *Biol Conserv.* 2014;170:56–63. <https://doi.org/10.1016/j.biocon.2013.12.036>.
- Freckleton RP, Harvey PH, Pagel M. Phylogenetic analysis and comparative data: a test and review of evidence. *Am Nat.* 2002;160:712–726. <https://doi.org/10.1086/343873>.
- Gargiulo R, Budde KB, Heuertz M. Mind the lag: understanding genetic extinction debt for conservation. *Trends Ecol Evol.* 2025;40:228–237. <https://doi.org/10.1016/j.tree.2024.10.008>.
- Garrison E, Kronenberg ZN, Dawson ET, Pedersen BS, Prins P. A spectrum of free software tools for processing the VCF variant call format: vcflib, bio-vcf, cyvcf2, hts-nim and slivar. *PLoS Comput Biol.* 2022;18:e1009123. <https://doi.org/10.1371/journal.pcbi.1009123>.
- Germain RR *et al.* Species-specific traits mediate avian demographic responses under past climate change. *Nat Ecol Evol.* 2023;7:862–872. <https://doi.org/10.1038/s41559-023-02055-3>.
- Gilroy DL, Phillips KP, Richardson DS, van Oosterhout C. Toll-like receptor variation in the bottlenecked population of the Seychelles warbler: computer simulations see the ‘ghost of selection past’ and quantify the ‘drift debt’. *J Evol Biol.* 2017;30:1276–1287. <https://doi.org/10.1111/jeb.13077>.
- Groß C *et al.* 2020. Prioritizing sequence variants in conserved non-coding elements in the chicken genome using chCADD. *PLoS Genet.* 16:e1009027. <https://doi.org/10.1371/journal.pgen.1009027>.
- Grossen C, Guillaume F, Keller LF, Croll D. Purging of highly deleterious mutations through severe bottlenecks in Alpine ibex. *Nat Commun.* 2020;11:1–12. <https://doi.org/10.1038/s41467-020-14803-1>.
- Grueber CE. Comparative genomics for biodiversity conservation. *Comput Struct Biotechnol J.* 2015;13:370–375. <https://doi.org/10.1016/j.csbj.2015.05.003>.
- Hackl T, Ankenbrand MJ, van Adrichem B. gggenomes: a Grammar of Graphics for Comparative Genomics, 2024. Available from: <https://github.com/thackl/gggenomes>
- Hadfield JD. MCMC methods for multi-response generalized linear mixed models: the MCMCglmm R package. *J Stat Softw.* 2010;33:1–22. <https://doi.org/10.18637/jss.v033.i02>.
- Hadfield JD, Nakagawa S. General quantitative genetic methods for comparative biology: phylogenies, taxonomies and multi-trait models for continuous and categorical characters. *J Evol Biol.* 2010;23:494–508. <https://doi.org/10.1111/j.1420-9101.2009.01915.x>.
- Hasselgren M *et al.* Genomic and fitness consequences of inbreeding in an endangered carnivore. *Mol Ecol.* 2021;30:2790–2799. <https://doi.org/10.1111/mec.15943>.
- Hickey G, Paten B, Earl D, Zerbino D, Haussler D. HAL: a hierarchical format for storing and analyzing multiple genome alignments. *Bioinformatics.* 2013;29:1341–1342. <https://doi.org/10.1093/bioinformatics/btt128>.
- Hilgers L *et al.* Avoidable false PSMC population size peaks occur across numerous studies. *Curr Biol.* 2025;35:927–930.e3. <https://doi.org/10.1016/j.cub.2024.09.028>.
- Hughes AL, Piontkivska H. DNA repeat arrays in chicken and human genomes and the adaptive evolution of avian genome size. *BMC Evol Bio.* 2005;5. <https://doi.org/10.1186/1471-2148-5-12>.
- IUCN. The IUCN red list of threatened species. Version 2024-1, 2024. Available from: <https://www.iucnredlist.org>
- Jackson HA *et al.* Genomic erosion in a demographically recovered bird species during conservation rescue. *Conserv Biol.* 2022;36:e13918. <https://doi.org/10.1111/cobi.13918>.
- Jeon JY *et al.* Genomic diversity as a key conservation criterion: proof-of-concept from mammalian whole-genome resequencing data. *Evol Appl.* 2024;17:e70000. <https://doi.org/10.1111/eva.70000>.
- Jones CG *et al.* The restoration of the Mauritius Kestrel *Falco punctatus* population. *Ibis (Lond 1859).* 1995;137:S173–S180. <https://doi.org/10.1111/j.1474-919X.1995.tb08439.x>.
- Jones CG. Back from the brink: the echo parakeet story. *PsittaScene.* 2010;22:3–5.
- Jones CG. Pink Pigeon *Nesoenas mayeri*. In: Safford R, Hawkins F, editors. *The birds of Africa: volume VIII: the Malagasy region: Madagascar, Seychelles, Comoros, Mascarenes.* Christopher Helm; 2013. p. 484–489.
- Jones CG *et al.* Echo parakeet *Psittacula eques*. In: Safford RJ, Hawkins AFA, editors. *The birds of Africa. Vol. VIII: the Malagasy region.* Christopher Helm; 2013. p. 517–522.
- Jones CG, Swinnerton KJ. A summary of conservation status and research for the Mauritius kestrel *Falco punctatus*, Pink Pigeon *Columba mayeri* and echo parakeet *Psittacula eques*. *Dodo.* 1997;33:72–75.
- Kalinowski ST, Waples RS. Relationship of effective to census size in fluctuating populations. *Conserv Biol.* 2002;16:129–136. <https://doi.org/10.1046/j.1523-1739.2002.00134.x>.
- Kardos M *et al.* The crucial role of genome-wide genetic variation in conservation. *Proc Natl Acad Sci U S A.* 2021;118:e2104642118. <https://doi.org/10.1073/pnas.2104642118>.
- Kardos M *et al.* Inbreeding depression explains killer whale population dynamics. *Nat Ecol Evol.* 2023;7:675–686. <https://doi.org/10.1038/s41559-023-01995-0>.
- Khan A *et al.* Genomic evidence for inbreeding depression and purging of deleterious genetic variation in Indian tigers. *Proc Natl Acad Sci U S A.* 2021;118:e2023018118. <https://doi.org/10.1073/pnas.2023018118>.
- Kimura M, Maruyama T, Crow JF. The mutation load in small populations. *Genetics.* 1963;48:1303–1312. <https://doi.org/10.1093/genetics/48.10.1303>.
- Kircher M *et al.* A general framework for estimating the relative pathogenicity of human genetic variants. *Nat Genet.* 2014;46:310–315. <https://doi.org/10.1038/ng.2892>.
- Korneliussen TS, Albrechtsen A, Nielsen R. ANGSD: analysis of next generation sequencing data. *BMC Bioinformatics.* 2014;15:356. <https://doi.org/10.1186/s12859-014-0356-4>.
- Kumar S *et al.* TimeTree 5: an expanded resource for species divergence times. *Mol Biol Evol.* 2022;39:msac174. <https://doi.org/10.1093/molbev/msac174>.

- Kyriazis CC, Robinson JA, Lohmueller KE. Long runs of homozygosity are reliable genomic markers of inbreeding depression. *Trends Ecol Evol*. 2025;40:874–884. <https://doi.org/10.1016/j.tree.2025.06.013>.
- Lande R. Genetics and demography in biological conservation. *Science*. 1988;241:1455–1460. <https://doi.org/10.1126/science.3420403>.
- Lewin HA *et al*. Earth BioGenome Project: sequencing life for the future of life. *Proc Natl Acad Sci U S A*. 2018;115:4325–4333. <https://doi.org/10.1073/pnas.1720115115>.
- Li H. Tabix: fast retrieval of sequence features from generic TAB-delimited files. *Bioinformatics*. 2011;27:718–719. <https://doi.org/10.1093/bioinformatics/btq671>.
- Li H. Minimap2: pairwise alignment for nucleotide sequences. *Bioinformatics*. 2018;34:3094–3100. <https://doi.org/10.1093/bioinformatics/bty191>.
- Li H, Durbin R. Fast and accurate short read alignment with Burrows–Wheeler transform. *Bioinformatics*. 2009;25:1754–1760. <https://doi.org/10.1093/bioinformatics/btp324>.
- Li H, Durbin R. Inference of human population history from individual whole-genome sequences. *Nature*. 2011;475:493–496. <https://doi.org/10.1038/nature10231>.
- Liu X *et al*. Time-lagged genomic erosion and future environmental risks in a bird on the brink of extinction. *Proc R Soc B Biol Sci*. 2025;292:20242480. <https://doi.org/10.1098/rspb.2024.2480>.
- Mc Cartney AM *et al*. The European Reference Genome Atlas: piloting a decentralised approach to equitable biodiversity genomics. *npj Biodivers*. 2024;3:28. <https://doi.org/10.1038/s44185-024-00054-6>.
- McNeely JA, Miller KR, Reid WV, Mittermeier RA, Werner TB. *Conserving the world's Biological diversity*. IUCN; 1990.
- Minh BQ *et al*. IQ-TREE 2: new models and efficient methods for phylogenetic inference in the genomic era. *Mol Biol Evol*. 2020;37:1530–1534. <https://doi.org/10.1093/molbev/msaa015>.
- Morales HE *et al*. The genome sequence of the Mauritius parakeet, *Alexandrinus eques* (formerly *Psittacula eques*) (A. Newton & E. Newton, 1876). *Wellcome Open Res*. 2024a;9:378. <https://doi.org/10.12688/wellcomeopenres.22583.1>.
- Morales HE *et al*. The genome sequence of the Mauritius kestrel, *Falco punctatus* (Temminck, 1821). *Wellcome Open Res*. 2024b;9:312. <https://doi.org/10.12688/wellcomeopenres.22452.1>.
- Morales HE *et al*. The genome sequence of the Pink Pigeon, *Nesoenas mayeri* (Prévost, 1843). *Wellcome Open Res*. 2024c;9:336. <https://doi.org/10.12688/wellcomeopenres.22471.1>.
- Morin PA *et al*. Reference genome and demographic history of the most endangered marine mammal, the vaquita. *Mol Ecol Resour*. 2021;21:1008–1020. <https://doi.org/10.1111/1755-0998.13284>.
- Ng PC, Henikoff S. SIFT: predicting amino acid changes that affect protein function. *Nucleic Acids Res*. 2003;31:3812–3814. <https://doi.org/10.1093/nar/gkg509>.
- Nicoll MAC *et al*. Contrasting recovery trajectories of four reintroduced populations of the Endangered Mauritius Kestrel (*Falco punctatus*). *Ibis (Lond 1859)*. 2021;163:1294–1309. <https://doi.org/10.1111/ibi.12987>.
- Orme D *et al*. caper: comparative analyses of phylogenetics and evolution in R, 2023. Available from: <https://github.com/davidorme/caper>
- Paradis E, Claude J, Strimmer K. APE: analyses of phylogenetics and evolution in R language. *Bioinformatics*. 2004;20:289–290. <https://doi.org/10.1093/bioinformatics/btg412>.
- Paradis E, Schliep K. Ape 5.0: an environment for modern phylogenetics and evolutionary analyses in R. *Bioinformatics*. 2019;35:526–528. <https://doi.org/10.1093/bioinformatics/bty633>.
- Patwardhan M, Wenger C, Davis E, Phanstiel D. Bedtools: An R package for genomic data analysis and manipulation. *J Open Source Softw*. 2019;4:1742. <https://doi.org/10.21105/joss.01742>.
- Pedersen BS, Quinlan AR. Mosdepth: quick coverage calculation for genomes and exomes. *Bioinformatics*. 2018;34:867–868. <https://doi.org/10.1093/bioinformatics/btx699>.
- Peona V *et al*. The avian W chromosome is a refugium for endogenous retroviruses with likely effects on female-biased mutational load and genetic incompatibilities. *Philos Trans R Soc B Biol Sci*. 2021;376:20200186. <https://doi.org/10.1098/rstb.2020.0186>.
- Pinheiro J *et al*. Package ‘nlme’. Linear and nonlinear mixed effects models, version 3, 2017;no. 1:274.
- Pinto AV, Hansson B, Patramanis I, Morales HE, van Oosterhout C. The impact of habitat loss and population fragmentation on genomic erosion. *Conserv Genet*. 2024;25:49–57. <https://doi.org/10.1007/s10592-023-01548-9>.
- Plough LV. Genetic load in marine animals: a review. *Curr Zool*. 2016;62:567–579. <https://doi.org/10.1093/cz/zow096>.
- Prasad A, Lorenzen ED, Westbury MV. Evaluating the role of reference-genome phylogenetic distance on evolutionary inference. *Mol Ecol Resour*. 2022;22:45–55. <https://doi.org/10.1111/1755-0998.13457>.
- Quinlan AR, Hall IM. BEDTools: a flexible suite of utilities for comparing genomic features. *Bioinform*. 2010;26:841–842. <https://doi.org/10.1093/bioinformatics/btq033>.
- Reed DH, Lowe EH, Briscoe DA, Frankham R. Inbreeding and extinction: effects of rate of inbreeding. *Conserv Genet*. 2003;4:405–410. <https://doi.org/10.1023/A:1024081416729>.
- Renaud G, Hanghøj K, Korneliusen TS, Willerslev E, Orlando L. Joint estimates of heterozygosity and runs of homozygosity for modern and ancient samples. *Genetics*. 2019;212:587–614. <https://doi.org/10.1534/genetics.119.302057>.
- Rentsch P, Witten D, Cooper GM, Shendure J, Kircher M. CADD: predicting the deleteriousness of variants throughout the human genome. *Nucleic Acids Res*. 2019;47:D886–D894. <https://doi.org/10.1093/nar/gky1016>.
- Rieseberg LH. Chromosomal rearrangements and speciation. *Trends Ecol Evol*. 2001;16:351–358. [https://doi.org/10.1016/S0169-5347\(01\)02187-5](https://doi.org/10.1016/S0169-5347(01)02187-5).
- Robinson J, Kyriazis CC, Yuan SC, Lohmueller KE. Deleterious variation in natural populations and implications for conservation genetics. *Annu Rev Anim Biosci*. 2023;11:93–114. <https://doi.org/10.1146/annurev-animal-080522-093311>.
- Serrato-Capuchina A, Matute DR. The role of transposable elements in speciation. *Genes (Basel)*. 2018;9:254. <https://doi.org/10.3390/genes9050254>.
- Shaffer ML. Minimum population sizes for species conservation. *BioScience*. 1981;31:131–134. <https://doi.org/10.2307/1308256>.
- Shaw RE *et al*. Global meta-analysis shows action is needed to halt genetic diversity loss. *Nature*. 2025;638:704–710. <https://doi.org/10.1038/s41586-024-08458-x>.

- Silver LW *et al.* 2024 November 10. Temporal loss of genome-wide and immunogenetic diversity in a near-extinct parrot. 2024.11.10.622863. <https://www.biorxiv.org/content/10.1101/2024.11.10.622863v1>
- Smit AFA, Hubley R, Green P. RepeatMasker open-4.0, 2015. Available from: <http://www.repeatmasker.org>
- Speak SA *et al.* Genomics-informed captive breeding can reduce inbreeding depression and the genetic load in zoo populations. *Mol Ecol Resour.* 2024;24:e13967. <https://doi.org/10.1111/1755-0998.13967>.
- Spielman D, Brook BW, Frankham R. Most species are not driven to extinction before genetic factors impact them. *Proc Natl Acad Sci U S A.* 2004;101:15261–15264. <https://doi.org/10.1073/pnas.0403809101>.
- Sæther B-E *et al.* Generation time and temporal scaling of bird population dynamics. *Nature.* 2005;436:99–102. <https://doi.org/10.1038/nature03666>.
- Stiller J *et al.* Complexity of avian evolution revealed by family-level genomes. *Nature.* 2024;629:851–860. <https://doi.org/10.1038/s41586-024-07323-1>.
- Storer J, Hubley R, Rosen J, Wheeler TJ, Smit AF. The Dfam community resource of transposable element families, sequence models, and genome annotations. *Mob DNA.* 2021;12:2. <https://doi.org/10.1186/s13100-020-00230-y>.
- Supple MA, Shapiro B. Conservation of biodiversity in the genomics era. *Genome Biol.* 2018;19:131. <https://doi.org/10.1186/s13059-018-1520-3>.
- Tan HZ *et al.* Megafaunal extinctions, not climate change, may explain Holocene genetic diversity declines in Numenius shorebirds. *eLife.* 2023;12:e85422. <https://doi.org/10.7554/eLife.85422>.
- Teixeira JC, Huber CD. The inflated significance of neutral genetic diversity in conservation genetics. *Proc Natl Acad Sci U S A.* 2021;118:e2015096118. [10.1073/pnas.2015096118](https://doi.org/10.1073/pnas.2015096118).
- Theissinger K *et al.* How genomics can help biodiversity conservation. *Trends Genet.* 2023;39:545–559. <https://doi.org/10.1016/j.tig.2023.01.005>.
- Tollington S *et al.* Long-term, fine-scale temporal patterns of genetic diversity in the restored Mauritius parakeet reveal genetic impacts of management and associated demographic effects on reintroduction programmes. *Biol Conserv.* 2013;161:28–38. <https://doi.org/10.1016/j.biocon.2013.02.013>.
- Tollington S *et al.* Detailed monitoring of a small but recovering population reveals sublethal effects of disease and unexpected interactions with supplemental feeding. *J Anim Ecol.* 2015;84:969–977. <https://doi.org/10.1111/1365-2656.12348>.
- van der Valk T, de Manuel M, Marques-Bonet T, Guschanski K. 2019 April 22. Estimates of genetic load in small populations suggest extensive purging of deleterious alleles. bioRxiv 696831. <https://doi.org/10.1101/696831>
- van der Valk T, Diez-del-Molino D, Marques-Bonet T, Guschanski K, Dalén L. Historical genomes reveal the genomic consequences of recent population decline in eastern Gorillas. *Curr Biol.* 2019;29:165–170.e6. <https://doi.org/10.1016/j.cub.2018.11.055>.
- van Oosterhout C *et al.* 2022 September 15. Genomic erosion in the assessment of species extinction risk and recovery potential [preprint]. bioRxiv:507768. <https://doi.org/10.1101/2022.09.13.507768>
- Wang X, Peischl S, Heckel G. Demographic history and genomic consequences of 10,000 generations of isolation in a wild mammal. *Curr Biol.* 2023;33:2051–2062.e4. <https://doi.org/10.1016/j.cub.2023.04.042>.
- Whiteley AR, Fitzpatrick SW, Funk WC, Tallmon DA. Genetic rescue to the rescue. *Trends Ecol Evol.* 2015;30:42–49. <https://doi.org/10.1016/j.tree.2014.10.009>.
- Wilder AP *et al.* The contribution of historical processes to contemporary extinction risk in placental mammals. *Science.* 2023;380:eabn5856. <https://doi.org/10.1126/science.abn5856>.
- Willi Y, Buskirk JV, Hoffmann AA. Limits to the adaptive potential of small populations. *Annu Rev Ecol Syst.* 2006;37:433–458. <https://doi.org/10.1146/annurev.ecolsys.37.091305.110145>.
- Wright BR *et al.* A demonstration of conservation genomics for threatened species management. *Mol Ecol Resour.* 2020;20:1526–1541. <https://doi.org/10.1111/1755-0998.13211>.