Polygenic basis for adaptive morphological variation in a threatened Aotearoa | New Zealand bird, the hihi (*Notiomystis cincta*)

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Abstract

To predict if a threatened species can adapt to changing selective pressures, it is crucial to understand the genetic basis of adaptive traits, especially in species historically affected by severe bottlenecks. We estimated the heritability of three hihi (Notiomystis cincta) morphological traits known to be under selection: nestling tarsus length, body mass and head-bill length, using 523 individuals and 39,699 single nucleotide polymorphisms (SNPs) from a 50K Affymetrix SNP chip. We then examined the genetic architecture of the traits via chromosome partitioning analyses and genome-wide association scans (GWAS). Heritabilities estimated using pedigree relatedness or genomic relatedness were low. For tarsus length, the proportion of genetic variance explained by each chromosome was positively correlated with its size, and more than one chromosome explained significant variation for body mass and head-bill length. Finally, GWAS analyses suggested many loci of small effect contributing to trait variation for all three traits, although one locus (a SNP within an intron of the transcription factor HEY2) was tentatively associated with tarsus length. Our findings suggest a polygenic nature for the morphological traits, with many small effect size loci contributing to the majority of the variation, similar to results from many other wild populations. However, the small effective population size, polygenic architecture and already low heritabilities suggest that both the total response and rate of response to selection are likely to be limited in hihi.

Keywords: genomic relatedness matrix; chromosome partitioning; genome-wide association; polygenic traits; pedigree study; Notiomystis cincta
1 Introduction

In small populations, genetic diversity is eroded over time through genetic drift, including events such as population bottlenecks and founding events (Bouzat, 2010). The continual loss of genetic diversity is of particular interest in species of conservation concern with already low levels of variation. The erosion of genetic diversity may result in the loss of putatively adaptive alleles, leaving the species with low adaptive potential when facing new selection pressures, such as a change in climate or habitat fragmentation (Hoffmann et al., 2017, Chen et al., 2016, de Villemereuil et al., 2019b). However, the complex genomic architecture of traits in non-model organisms is often unknown and, consequently, the underlying evolutionary processes and possible ecological consequences for a population or a species subject to selection cannot be directly assessed (Savolainen et al., 2013). Recently, with the advent of modern and inexpensive sequencing and genotyping technologies, the field of conservation genomics has entered a new era where it is possible to genotype a large number of markers for threatened species with small population sizes (Narum et al., 2013). Now, the challenge at hand is to adequately analyse and interpret these genomic snapshots in a way that we can predict the adaptive potential for threatened species and begin to inform conservation management (Harrisson et al., 2014; Shafer et al., 2015).

Genomic approaches have already proven to be a powerful tool to better understand the structure and evolution of adaptive traits (Pardo-Diaz et al., 2015; Luikart et al., 2018). For example, a closer look at the genetic basis of horn polymorphism in Soay sheep (Ovis aries) revealed that variation is maintained by heterozygote advantage at a single gene (Johnston et al., 2013), which would be impossible to infer from pedigree-based methods alone. Hence, today, many population genomic studies aim to investigate the genetic basis of variation in life-history, fitness and morphological traits in their organism of interest (Luikart et al., 2018). To make assumptions about adaptive potential, it is crucial to understand the proportion of the phenotypic variation observed in a population ($V_p$) that is explained by the additive genetic component ($V_A$), a factor referred to as narrow-sense heritability ($h^2$) (Falconer and Mackay 1996). Traditionally, $h^2$ has been estimated using the expected relatedness between individuals captured by pedigree data. However, those pedigrees may contain errors, not capture all relationships between sampled individuals due to shallow and inconsistent pedigree depth, and only capture expected, rather than realised, genome sharing between two related individuals (Visscher et al., 2006). Therefore, it has recently become common practice to instead use genomic data when estimating trait heritabilities (Gienapp et al., 2017b), for example via inferring the realised genome sharing between individuals described by a genomic
relatedness matrix (GRM). Assuming that marker density is sufficient to accurately capture
genomic relatedness, GRM-based heritabilities can be more precise, providing valuable insight
into the evolutionary potential of the species (Berenos et al., 2014, Gienapp et al., 2017b,
Jensen et al., 2014).

In addition to characterising the overall GRM-based heritability, the heritability can be
partitioned across regions of the genome to give further insight into the genomic basis of trait
variation (Yang et al., 2011b). If large chromosomes, containing more genes, explain a more
substantial proportion of the genetic variation compared to smaller chromosomes, the trait is
likely polygenic – i.e. there are many loci, scattered throughout the genome, contributing to
trait differences (Yang et al., 2011b, Robinson et al., 2013). More than one chromosome
explaining significant variation also suggests that many loci contribute to trait variation
(Kemppainen & Husby, 2018a).

Another way of investigating the genetic basis of adaptive traits using genomic data involves
a scan for genome-wide associations (GWAS). This detects specific loci with small to large
effect on the phenotype of the study species. The power to detect association is dependent,
among other factors, on the effect size of the locus. If a trait with significant heritability has no
significant GWAS peaks, it suggests that the trait is polygenic, with many loci of small effect
contributing to variation (Manilo et al., 2009). A further hurdle to finding significant genome-
wide associations is presented when individuals are closely related, which is commonly the
case when sampling from natural populations, particularly those with small population size.
The similarity in genetic background must first be adjusted for before testing for an association,
which may further reduce the power to detect an association (Yang et al., 2014).

It appears that a polygenic architecture underlies many evolutionarily significant traits under
investigation (Jensen et al., 2014; Kardos et al., 2016). Variation in these traits is maintained
by the balance between genetic drift, mutation and selection, with smaller heritabilities
predicted in populations with small effective population size (Hoffman et al., 2017). For these
populations, theoretical and simulation results suggest that polygenic adaptation in response
to selection is likely to proceed via allele frequency changes at a subset of loci with large effect
and/or high allele frequency, rather than frequency shifts across all loci (Höllinger et al., 2019).
Variation at other loci is susceptible to be lost via genetic drift and will, therefore, constrain
both the rate of response and the total response to selection (John & Stephan, 2020). In
addition, the evolution of polygenic traits can be further constrained by linkage disequilibrium
between deleterious and beneficial alleles, and trade-offs with other (potentially unmeasured)
traits (Hill & Robertson 1966; Hospital & Chevalet 1996; Lande 1982). Therefore,
understanding the genetic architecture of adaptive traits is likely to enable better predictions of the adaptive potential of small populations.

The hihi is an endemic passerine of Aotearoa | New Zealand and is becoming a model species for studying the genetics of endangered birds, and species with small population sizes more generally (Brekke et al., 2011; de Villemereuil et al., 2019a; de Villemereuil et al., 2019b). The hihi dataset on the island of Tiritiri Matangi (36°36'S, 174°53'E) contains a multi-generational pedigree and morphological and life history data for every individual born since 1995 when the population was established through reintroduction. Recently, de Villemereuil et al. (2019a, 2019b) predicted little adaptive potential of this species based on signatures of low genetic diversity, low additive genetic variance of fitness, and small pedigree-based heritabilities for traits under selection. All but one life-history trait had very small heritabilities (<0.0001), however, there was good support for low but non-zero heritabilities for beak length, body mass and tarsus length (de Villemereuil et al., 2019b). While previous hihi studies already examined the age-dependent effect of body size on fitness (Low 2006), this large dataset (>3000 birds) for the first time revealed that all three morphological traits had positive linear selection and negative non-linear selection gradients (de Villemereuil et al., 2019b). These gradients suggest that there are optimal values for nestling tarsus length, head-bill length and mass that maximise fitness, but that all trait means are significantly smaller than these optima, and hence are under directional selection to increase body size. Morphological traits are commonly recorded in avian studies, with body size directly associated with fitness in many species (reviewed in Merilä and Sheldon, 2001; Gardner et al., 2011), and are the focus of a growing number of genomic studies (reviewed in Husby et al., 2019).

Here, we used genomic data to reveal previously undiscovered genetic patterns underlying these three adaptive morphological phenotypes in the Tiritiri Matangi population. We employed state-of-the-art software and data for 523 individuals genotyped on a recently generated custom SNP array to generate heritability estimates from pedigree relatedness and genomic relatedness, testing whether a genomic snapshot provides comparable results to a pedigree over many generations for the additive genetic variance component of the traits. To explore the genomic architecture of each trait, we further partitioned the variance explained by the genotyped SNPs across chromosomes to test for a possible polygenic architecture. Finally, we performed a genome-wide association study in order to find loci of small, medium and large effect on any of the traits under investigation. For traits with low heritability, we wanted to discern whether a GWAS could detect loci of large effect, or whether there was evidence that
traits are controlled by many loci of small effect, in agreement with studies on many other wild populations. From a conservation management perspective, a low heritability estimate and polygenic makeup of these morphological traits will complement our previous work demonstrating that hihi has little adaptive potential when facing future changes in selection pressures.

2 Methods

Study population and phenotypic data
The hihi was once abundant on the North Island of Aotearoa, until habitat loss and predation by introduced mammals confined its range to only one naturally occurring population on Te Hauturu-o-Toi (Little Barrier Island), in the Hauraki Gulf (36°12’S, 175°05’E) (Brekke et al., 2011). In 1995, the hihi was successfully introduced to the nearby island of Tiritiri Matangi, with a population size of 210 birds prior to the 2019/2020 breeding season (Parlato et al., in review). As a result of extensive monitoring since its translocation, we now have a wealth of environmental, life history, fitness, morphological and pedigree data for the population (de Villemereuil et al., 2019b). Hihi do not migrate between the populations and predominantly nest in nest boxes on Tiritiri Matangi Island. During every breeding season (ranging from October through to February), nestlings are banded and measured for body mass (g), but also for tarsus length (mm) and head-to-bill length (mm) as those two measures stabilize after 21 days. Blood has been routinely taken (around 60µl per individual via brachial vein puncture) from nestlings since 2004.

Genotyping and pedigree data
Due to a very high rate of extra-pair copulations in this species, the paternity is determined via routine genotyping of a panel of 19 microsatellite markers, allowing the construction of a multi-generational pedigree for the population (for details see Brekke et al., 2012; de Villemereuil et al., 2019a). For nestlings without blood samples available (i.e. those born before 2004), fathers are assumed to be unknown in the pedigree. In addition, in 2016, 1,536 birds (from five different hihi populations) were genotyped using a custom 50k Affymetrix single nucleotide polymorphism (SNP) array (manuscript in prep). This array was developed based on the identification of SNPs from de-novo assembly of restriction site-associated DNA sequencing (RAD-seq) of 31 individuals from the Te Hauturu-o-Toi and Tiritiri Matangi populations, and low-coverage whole-genome sequencing of ten of these birds (three of which were from Tiritiri Matangi). The flanking sequences of identified variants were aligned to the Ensemble 86 zebra finch (Taeniopygia guttata) genome, which, given the very high synteny of avian genomes
(Zhang et al., 2014), is used in the following analyses as a proxy for the distribution of markers in the hihi genome (electronic supplementary material, table S1).

Of the 58,466 SNPs included on the array, 45,553 markers were designated as polymorphic high resolution according to the default quality control metrics in the Axiom Analysis Suite software. PLINK (Purcell et al., 2007) filtering (--maf 0.01, --hwe 1e-20) resulted in the removal of 325 and 69 loci among the autosomes, respectively. 2,734 SNPs appeared as duplicates (i.e. mapped to the same chromosome at the same position) and hence only one SNP from each position was retained. To avoid population structure as a confounding factor, we only included birds from Tiritiri Matangi Island in this study, which shows no structure (electronic supplementary material, figure S1). No individual was removed due to low genotyping rates (-mind), but some were excluded for having incomplete phenotypic information. The final dataset contained 39,699 autosomal genomic loci with a mean call rate of 0.997398 and complete sex and phenotype information on all three traits for 523 birds.

**Trait heritabilities**

We employed several different methods of retrieving heritability estimates for our three traits under investigation, using generalised linear mixed models (an ‘animal model’, Wilson et al., 2009) and either the pedigree relatedness or the relatedness estimated from the genotyped markers (i.e. the genomic relatedness matrix; GRM). First, pedigree- and GRM-based heritabilities ($h^2_{ped}$, $h^2_{GRM1}$) were estimated in a Bayesian framework using the R (R Core Team, 2013c) package MCMCglmm (Hadfield 2010). To estimate $h^2_{ped}$, we fit the relatedness matrix calculated from the full pedigree. To estimate $h^2_{GRM1}$, the GRM was constructed using an approach that scales by the actual variance in relatedness (approach 3; Robinson et al., 2013). Following de Villemereuil (et al., 2019b), the additive genetic variance ($V_A$) for each trait was estimated by fitting sex and clutch size as fixed effects and accounting for the random effects of mother, social father, year and month when the nestling was born. Models were run for 503,000 iterations with a burn-in period of 3,000, sampling every 10th output from the chain, and convergence checked graphically and using the Heidelberger and Welch convergence test. The heritability was then calculated as the ratio of $V_A$ to $V_P$ (total phenotypic variance), where $V_P$ and thus $h^2$ accounts for the variance explained by fixed effects (de Villemereuil et al., 2018).

Furthermore, to be able to compare our results with the heritability of morphological traits reported in other avian studies (reviewed in Husby et al., 2019), we also used the “prefitModel” function in the R package RepeatABEL (Rönnegård et al., 2016). While RepeatABEL is mainly...
known for allowing the inclusion of repeated measures, it was chosen as it allows the inclusion of multiple random effects into the model. The GRM-based heritability ($h^2_{GRM}$) was estimated by fitting the same fixed and random effects as above.

Finally, the package BayesR (Moser et al., 2015) was used to infer the phenotypic variation explained by SNPs in different effect size distributions and to estimate the SNP-based heritability ($h^2_{BayesR}$). The SNP-based heritability differs from the GRM-based heritability as it estimates the proportion of variance accounted for by linear regression on a set of genotyped SNPs (de los Campos et al., 2015). As BayesR does not allow for the inclusion of any fixed or random effects, the raw phenotypes were corrected for those effects in MCMCglmm (same model as for the heritability analysis, but excluding $V_a$) and the residuals then used as phenotypes in BayesR. We set the mean effect sizes of 0.01, 0.001, 0.0001 or 0 for the four mixture distributions and ran the programme for 50,000 iterations, with a burn-in of 20,000, recording every 10th sample in the chain after that (as suggested in the default settings).

Chromosome partitioning of genetic variance

For each chromosome, we estimated the contribution to overall trait variance by fitting a mixed model in MCMCglmm with a GRM estimated from markers on the focal chromosome, a GRM calculated from all other markers in the genome, and accounting for sex, clutch size, mother, social father, year and month of birth. As above, the GRMs were calculated using an approach that scales by the actual variance in relatedness. We ran 203,000 iterations with a burn-in period of 3,000, sampling every 10th output from the chain. Due to convergence issues for chromosomes with fewer than 850 markers, we merged multiple chromosomes based on their size according to the zebra finch karyotypes (electronic supplementary material, table S1). The variance was estimated individually for chromosomes 1-14, 1A and 20 and each merged chromosome group.

We also used GCTA (V1.24, Yang et al., 2011a) to estimate the variance explained by each chromosome. GRMs were generated with the --make-grm option for each individual chromosome and all but the focal chromosome. Because the model failed to converge when fitting all autosomes simultaneously, we fit the GRM pairs (e.g. GRM for Chr 1 and GRM for all but Chr 1, using the --mgrm option) in the default REML model one by one to estimate the additive genetic variance explained by each chromosome. Both pathways, however, should theoretically result in the same heritability estimate per autosomal chromosome (Yang et al., 2011a). Fixed effects included sex and clutch size, and we adjusted for population stratification using the --qcovar function on the top ten principle components.
For both methods, we used the cor.test function in R to test the significance of a positive Pearson’s correlation between chromosome size (based on the zebra finch genome) and variance explained. To account for p-value inflation as a result of heteroscedasticity and censoring, we used 100,000 permutations to calculate corrected p-values following the approach of Kemppainen and Husby (2018b).

GWAS

To test for association between trait and genotype, we used the RepeatABEL package. The “prefitModel” function first fits a model without the SNP-effects and computes the variance components for the trait, using the hglm package. SNPs are then tested for association by accounting for these random effects in the “rGLS” function in RepeatABEL, using the same fixed and random effects as in the heritability analysis. P-values for each SNP are computed using Wald statistics, together with the SNP effect for each marker. We used a Bonferroni correction to control for family-wise error rate from multiple testing (significance value of 0.05 divided by the number of markers, giving a critical p-value of 1.26x10^{-6}). All genomic SNP were tested for association, including those that mapped to random or the Unknown zebra finch chromosomes, or were not mapped to a zebra finch chromosome (which we combine as ‘chromosome 77’, electronic supplementary material, table S1). SNPs of interest were further investigated based on the zebra finch genome annotation in the Ensembl database (Version 100.12: bTaeGut1_v1.p; Cunningham et al., 2019). In addition, we calculated the genomic inflation factor (λ, computed using lower 90 per cent of the distribution), to determine whether population structure was likely to have led to inflation of the test statistics.

3 Results

i) Trait heritability

The pedigree-based estimates for the heritability of nestling morphological traits tended to be higher than the GRM-based heritabilities, and MCMCglm GRM-based estimates \( h^2_{GRM} \) were lower than corresponding RepeatABEL heritabilities (table 1; electronic supplementary material, table S2, table S3, figure S2). The SNP-based heritabilities from BayesR were 0.117, 0.079 and 0.076, respectively (electronic supplementary material, table S5). All methods agreed on tarsus length being the ‘most heritable’ of the three morphological traits.

Table 1: Summary table of the outputs (electronic supplementary material, table S2, table S3) for the trait mean, total phenotypic variance \( V_P \), pedigree-based and GRM-based heritability \( h^2 \) estimates and additive genetic variance \( V_A \) for tarsus length (mm), body mass (g) and head-bill length (mm). We report the mean \( h^2 \) and \( V_A \) for
the MCMCglmm (ped, GRM1) and RepeatABEL (GRM2) estimates for 523 hihi (268 males and 255 females). Credible / confidence intervals (MCMCglmm) or standard error (RepeatABEL) are reported in square brackets.

<table>
<thead>
<tr>
<th>Trait</th>
<th>Mean</th>
<th>$V_P$</th>
<th>$h^2_{ped}$</th>
<th>$V_{A, ped}$</th>
<th>$h^2_{GRM1}$</th>
<th>$V_{A, GRM1}$</th>
<th>$h^2_{GRM2}$</th>
<th>$V_{A, GRM2}$</th>
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</thead>
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<tr>
<td>Tarsus length</td>
<td>27.576</td>
<td>1.440</td>
<td>0.133</td>
<td>0.190</td>
<td>0.077</td>
<td>0.111</td>
<td>0.081</td>
<td>0.082</td>
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<td></td>
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<td>[1.39e-8,</td>
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<td></td>
<td></td>
<td></td>
<td>0.253]</td>
<td>0.362]</td>
<td>0.154]</td>
<td>0.219]</td>
<td>0.182]</td>
<td>0.126]</td>
</tr>
<tr>
<td>Body mass</td>
<td>37.011</td>
<td>43.662</td>
<td>0.086</td>
<td>2.880</td>
<td>0.042</td>
<td>1.812</td>
<td>0.065</td>
<td>2.050</td>
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<td></td>
<td>0.151]</td>
<td>6.646]</td>
<td>0.105]</td>
<td>4.548]</td>
<td>0.161]</td>
<td>3.276]</td>
</tr>
<tr>
<td>Head-bill length</td>
<td>38.793</td>
<td>2.845</td>
<td>0.120</td>
<td>0.343</td>
<td>0.032</td>
<td>0.091</td>
<td>0.070</td>
<td>0.159</td>
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<td>[0.101,</td>
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<td></td>
<td></td>
<td>0.252]</td>
<td>0.727]</td>
<td>0.094]</td>
<td>0.270]</td>
<td>0.168]</td>
<td>0.251]</td>
</tr>
</tbody>
</table>

The correlation between off-diagonal elements of the pedigree-based and the GRM-based relatedness matrices was 0.749 (electronic supplementary material, figure S3). A correlation of less than one is expected due to factors such as Mendelian sampling and agrees with other studies (e.g. Pryce et al., 2012; Bérénos et al., 2014; Galla et al., 2020).

ii) Chromosome partitioning of genetic variance

There was evidence that multiple loci contributed to the variance in all three morphological traits (figure 1; electronic supplementary material, figure S4). For tarsus length, we found a positive but non-significant correlation between the proportion of additive genetic variance explained by each chromosome and the chromosome size (HC corrected p-value = 0.113). Chromosome 1, 1A and 6 explain the most variation, although the largest chromosome (2) explained only moderate variation.

For body mass, we found a weak and non-significant positive correlation between chromosome size and phenotypic variance explained (HC corrected p-value = 0.422). Here, chromosomes 14 and 9 explain the most variation.
Figure 1: The variation explained by each chromosome or chromosome composite (in yellow circles, excluding 'Chr 77') for all three traits (tarsus length, body mass and head-bill length) as calculated in MCMCglmm. Plotted is additive genetic variance explained by each chromosome ($V_A$) against the length of the chromosome (in Mbp) based on the zebra finch reference genome. After HC correction, all p-values were non-significant (0.113, 0.422 and 0.574). a = chromosomes 15 and 4A, b = chromosomes 17-19, c = chromosomes 21-28 and 1B (electronic supplementary material, table S1).

Finally, for head-bill length, we found a weak negative correlation between the variation and the size of the genomic region. The three largest chromosomes explain relatively little of the overall variation, while chromosome 14, and micro-chromosome composites 'b' and 'c' each explain the most variation. Chromosome partitioning in GCTA resulted in a comparable picture for all three traits (electronic supplementary material, figure S5).

iii) GWAS

No SNP was significantly associated with the morphological traits after testing for association in RepeatABEL (figure 2). According to the well-curated zebra finch genome (which the hihi chromosome assignment is derived from), the SNP with the highest support for association with tarsus length is located in one of the intronic regions of the HEY2 transcription factor (electronic supplementary material, figure S6), which is involved in cardiac morphogenesis, neurogenesis and somitogenesis (Ihara et al., 2020).

For head-bill length, there are two SNPs that, while not significant, show a stronger association with the trait than all others. One of them is placed in an intron of an uncharacterized protein-coding gene (ENSTGUG00000006844) on chromosome 1, the other in between two genes associated with cat eye syndrome (CECR1, CECR2) on chromosome 1A. The third most associated SNP for head-bill length was found on chromosome 20, a chromosome previously associated with bill length (Knief et al., 2012; Silva et al., 2017).
The top ten SNPs with the lowest p-values across all traits can be found in electronic supplementary material, tables S4. We refrained from calculating the additive genetic variance explained by each of the most associated SNPs, as p-values were generally high. The genomic inflation factor values ranged from 0.96 to 1.05 for all three phenotypic traits under investigation (electronic supplementary material, figure S7), suggesting that population structure has been adequately accounted for.

**Figure 2**: Manhattan plots for the association between single nucleotide polymorphisms and (a) tarsus length, (b) body mass and (c) head–bill length in 523 hihi nestlings of Tiritiri Matangi. Top black dashed line: Bonferroni significance threshold, bottom dashed line: nominal significance (p-value = 0.05). A description of the chromosome order and naming can be found in the electronic supplementary material, table S1.
The *BayesR* analyses confirmed that all traits under investigation appear polygenic, with less than 9% of the SNPs affecting trait variation, the majority of which contributed to the smallest effect size distribution (electronic supplementary material, table S5). A comparison of the SNPs with the lowest p-values from the GWAS showed reasonable agreement with those inferred to have a high probability of inclusion in the largest effect size distribution from *BayesR* (electronic supplementary material, tables S4).

### 4 Discussion

Adaptive response requires that there is genetic variation in traits linked to fitness, and further, that response to selection is unconstrained by trade-offs with other traits and loci. Here, we examined the genetic basis of variation in nestling tarsus length, body mass and head-bill length in the hihi, a threatened yet well-monitored bird species of Aotearoa | New Zealand, in order to understand whether these traits can respond to current and future selection pressures.

Our analyses confirmed low heritability estimates, and support a polygenic basis for all three traits, with no loci of large effect contributing to trait variation. Given the low effective population size and the polygenic architecture, such that adaptive variation lost via drift is unlikely to be replenished by mutation in the short-term, we conclude that evolutionary potential in hihi is likely constrained even further than the already low heritabilities suggest.

We have three reasons to believe that a polygenic architecture underlies all three morphological traits under investigation: (i) for tarsus length, the variance explained by each chromosome scaled with chromosome size, and for all traits more than one chromosome contributed to variation, (ii) there were no significant SNP associations in the GWAS and (iii) the majority of loci explaining variation are of small effect size. This study is in good agreement with studies in other passerines, (e.g. great tits (Santure et al., 2013; 2015; Gienapp et al., 2017a; Bosse et al. 2017; Kim et al., 2018), collared flycatchers (Husby et al., 2015; Kardos et al., 2016; Silva et al., 2017), house sparrows (Silva et al., 2017; Lundregan et al., 2018), great reed warblers (Hansson et al. 2018), reviewed in Husby (et al., 2019)) as well as studies from other wild animal populations (e.g. Bérénos et al., 2015), where many loci with small effect are responsible for variation in adaptive phenotypes, rather than a few large-effect loci.

Our pedigree- and GRM-based heritability estimates support a small but non-zero heritability for all traits and agree with previous pedigree heritabilities estimated from a much larger number (>2,000) of individuals (de Villemereuil et al., 2019b). Interestingly, the GRM-based heritabilities were lower than the pedigree-based estimates, although the credible intervals are overlapping in all cases. One reason for this may be that our SNP array is not dense enough...
to have captured true genome sharing more accurately than the pedigree has done. Given the number of SNPs genotyped, this seems unlikely (Berenos et al., 2014), although we note that we have not successfully genotyped SNPs from a number of microchromosomes, including chromosome 16. We acknowledge that, while 523 individuals is a large dataset for a threatened species, our sample size is likely to be somewhat limiting and has certainly contributed to the large credible intervals and standard errors around both the pedigree- and GRM-based heritability estimates, which may also explain some of the differences between estimates. Interestingly, a study of Corsican blue tits (Cyanistes caeruleus; Perrier et al., 2018) suggests that a sample size of around 500 birds and data from a 50K SNP chip can outperform heritability estimates from a seven-generation pedigree. The ability to estimate heritabilities without long-term and near-complete monitoring of a population offers exciting opportunity to infer adaptive potential in other wild populations and will be particularly valuable in assessing the vulnerability of endangered species (Gienapp et al., 2017b). The sample size, number of markers and low heritabilities is also likely to have limited power to detect association in the GWAS analyses. Linkage disequilibrium between neighbouring markers is moderate (mean of 0.518, calculated with all 523 individuals using plink --r2 and reporting all pairwise values with --ld-window-r2 0.0) and suggests that power to detect a locus explaining all of the trait heritability ranges from 0.306 to 0.516 (Wang and Xu 2019; electronic supplementary material, table S6). As expected, nestling mass, with the lowest heritability due to substantial maternal and month effects compared to the other traits (de Villemereuil et al. 2019b), has the lowest power to detect association.

Morphological traits are often the subject of wild bird studies, as they can reveal evolutionary patterns over time and in turn shed light on the adaptive response of traits known to correlate with fitness (Merilä and Sheldon, 2001; Gardner et al., 2011). The three morphological traits studied here are all under selection as previous analyses suggest that the population mean of all three traits is substantially smaller than the fitness optimum (de Villemereuil et al. 2019b) and that there is a trade-off between body size and agility mediated during the highly energetic breeding season of the hihi (Low 2006). Although there was no significant association detected in the GWAS, it is interesting to note the regions that were most highly associated with tarsus length and head-bill length. The tentative association of a locus on chromosome 3 with tarsus length and of variants on chromosomes 1 and 1A with head-bill length does not appear to be replicated in some studies of passerine body size (Silva et al., 2017; Bosse et al., 2017), while Lawson and Petren (2017) highlight a region on chromosome 1A in increased linkage with regard to beak shape in Darwin finches (Geospiza). Additionally, the third-highest association for head-bill, a SNP on chromosome 20, stands within an intron of NFS1 cysteine desulfurase
and is 7-12Mb distant from QTLs previously associated with beak morphology (Knief et al., 2012; Silva et al., 2017). The challenges in identifying significant loci in a genome-wide association analysis include low SNP density and small sample sizes (Santure & Garant 2018), both of which are likely to apply to the hihi study system to some degree. All the above-mentioned loci are interesting candidates for further study if sample size and SNP density in these regions can be increased. Creating a larger sample size through combining all monitored populations could improve the precision of estimates, and shed more light on the effects of translocations on trait variation and responses to selection.

Despite the low heritabilities of the traits, our genomic analyses have provided important further insights for hihi, and threatened species more generally. A more nuanced understanding of the genomic architecture of a trait under selection helps assess the likely speed of the adaptive response in a species and the longer-term total response to selection (John & Stephan, 2020; Höllinger et al., 2019). We have demonstrated that many loci of small effect are likely to contribute the majority of variation to all three morphological traits. Given the small effective population size of all hihi populations, many adaptive variants are likely to be lost to drift at a rate that is faster than the input of new variation via mutation (reviewed in Johnson & Barton, 2005; Willi & Buskirk, 2006). This is likely to slow the speed of adaptation and lead to further constraint on the adaptive potential of the species. Most importantly, other wild populations may be facing similar threats as the hihi, highlighting the need for continued collection and use of genomic resources to aid conservation measures. We therefore encourage further characterisation of the heritability and genomic architecture for species that face anthropogenic threats including habitat decimation and climate change, in order to better predict their evolutionary potential to adapt to such challenges (Gienapp & Brommer, 2014). Finally, we acknowledge that future studies will need to go beyond describing quantitative parameters in order to successfully combine genomic insights with active species conservation, including informing management actions that can buffer species with small population sizes from the impacts of low adaptive potential.

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