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# 1 A multispecies *BCO2* beak color polymorphism in the Darwin's finch radiation

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# 23 Summary

Carotenoid-based polymorphisms are widespread in populations of birds, fish, and reptiles<sup>1</sup>, but 24 25 generally little is known about the factors affecting their maintenance in populations<sup>2</sup>. We report 26 a combined field and molecular-genetic investigation of a nestling beak color polymorphism in 27 Darwin's finches. Beaks are pink or yellow, and yellow is recessive<sup>3</sup>. Here we show that the 28 polymorphism arose in the Galápagos half a million years ago through a mutation associated 29 with regulatory change in the BCO2 gene, and is shared by 14 descendant species. The 30 polymorphism is probably a balanced polymorphism, maintained by ecological selection 31 associated with survival and diet. In cactus finches the frequency of the yellow genotype is 32 correlated with cactus fruit abundance and greater hatching success, and may be altered by 33 introgressive hybridization. Polymorphisms that are hidden as adults, as here, may be far more 34 common than is currently recognized, and contribute to diversification in ways that are yet to be 35 discovered.

# 36 **Results and Discussion**

37 Adaptive radiations are groups of related organisms that have diversified relatively rapidly from 38 a common ancestor<sup>4,5</sup>. A striking feature of some radiations is that polymorphic variation within 39 species is shared among related species (reviewed in<sup>2</sup>). This observation raises the question: how does variation persist in multiple related species<sup>2,6–8</sup>? Shared polymorphisms originate through 40 41 shared ancestral variation, repeated mutation and/or introgression, and are maintained by 42 negative frequency-dependent selection, heterozygous advantage or spatiotemporal fluctuations in selective pressures<sup>2,7,8</sup>. Distinguishing among these alternatives requires an understanding of 43 44 the genetic basis of phenotypic variation, phylogenetic history, and fitness variation in natural 45 populations. This has been rarely achieved, partly because polymorphisms are often associated with large genomic regions containing many genes of uncertain functional importance (e.g.<sup>9-14</sup>), 46 47 and partly because of the difficulties of determining fitness in nature. Here we report a shared 48 color polymorphism in the radiation of Darwin's finches (Thraupidae) on the Galápagos and 49 Cocos islands. We identify its genetic basis and phylogenetic origin, and take advantage of 50 uniquely banded individuals on one island to consider how ecological factors might contribute to 51 the maintenance of the polymorphism.

52

# Identification of the gene responsible for a nestling color polymorphism

A beak color polymorphism in nestlings has been documented in ten species of Darwin's finches (three *Camarhynchus* and seven *Geospiza*<sup>15</sup>). The beaks of nestlings are either pink or yellow (Figure 1A), recognizable at hatching, and similar in appearance among species. The yellow morphs are otherwise indistinguishable. Pedigrees on Daphne Major Island<sup>3</sup> and Genovesa<sup>16</sup> show that the yellow phenotype is recessively inherited and pink and yellow morphs occur in the

same nest, but the causal gene is unknown. The pink beaks appear pink because the blood supply
can be seen. Increasing melanin deposition in the beak obscures phenotypic expression several
weeks after fledging and culminates in complete melanization in breeding birds.

61 In order to identify loci associated with the polymorphism, we focused on two species 62 present on Daphne Major that differ in morph frequencies<sup>3</sup>: *Geospiza scandens* (yellow 63 frequency =  $\sim 30\%$ ) and G. fortis (yellow frequency =  $\sim 20\%$ ). We sequenced whole genomes at 64 low coverage of 456 individuals of known phenotype (mean depth =  $2.2 \pm 1.0$ X) to search for the 65 genetic basis of the polymorphism. We generated genotype likelihoods using the software ANGSD<sup>17</sup> and conducted an association analysis under a generalized logistic regression model 66 that incorporates genotype uncertainty<sup>18</sup> for G. scandens ( $n_{pink} = 98$ ,  $n_{vellow} = 98$ ) and G. fortis 67 68  $(n_{pink} = 130, n_{vellow} = 130)$  separately with nestling beak color as the response variable. We 69 discovered a small region on chromosome 24 harboring a region strongly associated with the 70 yellow phenotype overlapping the carotenoid-cleaving beta-carotene oxygenase 2 gene (BCO2, 71 Figure 1B). Mutations downregulating BCO2 expression or activity are known to increase deposition of carotenoids and pigmented phenotypes in birds, mammals, and reptiles<sup>19–26</sup>. By 72 73 closely inspecting this region in a combined sample of all 456 finches of the two species we 74 identified a single exonic single nucleotide polymorphism (SNP) with a likelihood ratio test 75 statistic (LRT) exceeding 166 (Figure 1C). It is also the only consistently elevated SNP in an 76 analysis of each species alone (Figure S1), is the best fit variant under a recessive model (see 77 note S1), and occurs on multiple haplotypes (Figure S1C). This SNP (chr24:6,166,878; 78 p6166878 hereafter) leads to a synonymous change 32bp into exon 4 of *BCO2*. We used high 79 coverage sequencing data for 16 pink and 8 yellow individuals (Figure S1D), and a larger subset 80 of individuals of unknown phenotype (Figure S1E), to search for SNPs or structural variants

81 linked to p6166878 in the vicinity but found none and confirmed the strong phenotype82 association with p6166878.

83 In order to further confirm the association for p6166878 we designed a TaqMan SNP 84 assay to genotype 1,631 individuals of known phenotype (Table S1) of five species from two 85 Galápagos islands collected during the period 1975-2012. Ninety-eight percent of observed 86 genotypes matched the genotype predicted from phenotype (Figure 1E). The few mismatched 87 pink phenotypes with yellow genotypes could be the result of mis-phenotyping or limited nutrition<sup>27</sup>, but mismatched individuals were notably often clustered in families (Figure S2B, but 88 89 not by species Figure S2A), suggesting a possible unknown genetic or shared environmental 90 contribution. None of the homozygous pink genotypes exhibited the yellow phenotype.

91

# Origin of the polymorphism

92 The vellow allele was not found in warbler finches (*Certhidea olivacea and C. fusca*) or 93 in the vegetarian finch (*Platyspiza crassirostris*) with a combined sample size of 42 individuals 94 (Figure 2), and therefore it probably arose by mutation soon after the split between the vegetarian 95 finch lineage and the ground/tree finch lineage roughly half a million years ago (Figure 2). The 96 polymorphism was retained throughout the radiation except in G. septentrionalis. All individuals 97 of this species have vellow beaks but, uniquely, they also have vellow legs and vellow skin, and 98 all three features are retained into adulthood<sup>15</sup>, strongly suggesting a different genetic basis than 99 for the nestling beak color polymorphism in the other species. We were not able to dissect the 100 genetic basis for yellow color in G. septentrionalis because there is no phenotypic variation 101 within this species.

102

# 103 Functional considerations

104 The functional importance of the observed synonymous change is uncertain, and the presence of 105 an unidentified linked causal variant cannot be completely ruled out (see Conclusions). However, a functional explanation is possible because codon usage can be under strong selection<sup>28</sup> and may 106 have functional consequences on translation<sup>28</sup>, RNA stability<sup>29</sup> and transcription<sup>30</sup>. Notably, 107 108 p6166878 changes the highest frequency value codon ( $f_{GTG} = 27.3\%$ ) to the lowest ( $f_{GTA} =$ 109 7.6%) in the reference genome. This is in line with the observed phenotypic effect of the yellow 110 mutation, because a lower abundance codon is expected to be associated with lower protein expression<sup>31</sup>. In this case, less BCO2 activity results in more carotenoid deposition in the yellow 111 112 morph. In fact, we found that yellow homozygotes showed significantly lower BCO2 expression 113 compared to pink homozygotes in the upper beak of developing embryos (Figure 1F) that were 114 sourced from a variety of different species and islands (Table S1): small sample sizes prohibit 115 species-specific analysis. Among the six heterozygous individuals, the pink allele was expressed 116 more than the vellow allele in five samples tested using a droplet-digital PCR (Figure S2C). 117 Differences in expression between the two alleles, and in the absence of alternative splice 118 variants (Methods), raises the possibility that the synonymous change alters transcription factor 119 binding affinity in exon 4. Further research into tissue-specific expression and the specific 120 transcription factors that regulate BCO2 is warranted.

121

# Long-term maintenance of the polymorphism

122 Nucleotide polymorphisms across the genome that are shared among 14 or more species of

123 Darwin's finches make up roughly 5% of all polymorphic sites (Figure S2D). Thus, the *BCO2* 

polymorphism lies in the tail of the distribution of polymorphic sites that show extensive multispecies polymorphism in the phylogeny. Such long-term persistence of a polymorphism (Figure
2) implies some form of balancing selection (reviewed in<sup>2,8</sup>). We next consider possible factors
that contribute to a balance in the short term.

128 Heterozygotes might survive better than homozygotes in their first year, but we found no 129 evidence of heterozygote advantage from the last week in the nest to the year after hatching for individuals captured during nest monitoring between 1978-1998 (G. fortis:  $\chi^2 = 1.2$ , P = 0.5, df = 130 2, n = 964 nestlings; G. scandens:  $\gamma^2 = 0.2$ , P = 0.9, df = 2; n = 326 nestlings). Since color 131 132 polymorphisms in birds are well-known to have signaling functions associated with disassortative mating<sup>32</sup>, predator avoidance<sup>33</sup>, and reproductive parameters<sup>34</sup>, among other 133 factors<sup>35</sup>, the nestling color variation could have a signaling role, allowing parents to feed 134 offspring preferentially<sup>36</sup>. However, a signaling role was rejected in a previous study because 135 136 observations made during half-hour nest watches, and parental feeding of recently fledged juveniles, gave no indication of preferential feeding<sup>15</sup>. 137

138 Alternatively, the significance of the polymorphism might reside in nutritional factors. 139 All species of Darwin's finches obtain carotenoids by feeding on pollen and/or herbivorous 140 insects, mainly Lepidoptera larvae, which parents feed to their nestlings. G. scandens, the species 141 on Daphne Major with the highest frequency of the yellow allele, is a specialist feeder on carotenoid-rich pollen from *Opuntia* cactus (Figure 3A, <sup>37</sup>) capable of feeding nestlings almost 142 143 entirely on a diet of cactus pollen and nectar. In 9 out of 13 years, yellow morph G. scandens 144 experienced higher first-year survival than pink morph G. scandens (Figure S3), reflected in overall higher survival of the yellow morph (34% vs 29%, ,  $\chi^2 = 4.31$ , n = 2065, P = 0.04, 145

146 Pearson's Chi-squared). Strong differences between some years are consistent with fluctuating 147 selection, in addition to random changes (see note S2), but we did not observe negatively 148 covarying trends expected under frequency-dependent selection. Individuals with the yellow 149 genotype survived conspicuously poorly in 1998, a year with el Niño conditions of abundant 150 rain, repeated breeding, but almost zero cactus flower and fruit production (Figure 3B), whereas in 1991, a similar el Niño year<sup>38</sup> except for plentiful cactus production, the yellow genotype 151 152 survived better than pink homozygous G/G or heterozygous G/A individuals. (Figure 4A, Table 153 S2, Generalized linear model: interaction genotype\*year, OR = 0.19, 95% CI = 0.04 - 0.82, P =154 0.03, n = 183 nestlings). As a consequence, the frequency of the yellow genotype G. scandens 155 plummeted with continued high mortality of G. scandens and low cactus fruit abundance into the 156 drought of 1999 (Figure 3C).

157 Although the beak color polymorphism is likely to be balanced, it is not fixed and static. 158 As previously described<sup>39–42</sup>, G. scandens hybridized occasionally with G. fortis and without 159 apparent loss of fitness over a 40-year period. Autosomal genes (alleles) flowed mainly from G. fortis to G. scandens<sup>39,41</sup> and they included BCO2 alleles that are more frequently pink in G. 160 161 fortis than in G. scandens. We evaluated the introgression of pink alleles using whole-genome 162 analysis of 176 individuals homozygous for the pink allele hatched early or late in the study period. Genome-wide divergence prior to 1995 was higher ( $F_{ST} = 0.15$ , n = 88) than after 2008 163  $(F_{ST} = 0.08, n = 88)$ , similar to previous estimates<sup>39</sup>. Four diagnostic G. fortis alleles at SNPs in 164 165 the near vicinity of p6166878 (within 5-kb), rose in frequency by 8-29% in the G. scandens 166 population during this time period (Note S2, Table S4). This is consistent with the convergence 167 in yellow allele frequency shown in Figure 3C, which is possibly a consequence of gene flow of 168 G. fortis-derived pink alleles into the G. scandens population.

Since carotenoids have an essential role in Vitamin A metabolism<sup>43</sup> and in color vision<sup>44</sup>. 169 170 altered BCO2 expression and carotenoid sequestration may be biochemically advantageous at 171 high intake levels for three reasons. First, deposition in the beak may avoid a toxic accumulation of metabolic breakdown products of circulating carotenoids<sup>43</sup>, so that sequestered carotenoids 172 173 can be metabolized later at a time of lower intake<sup>45</sup>. For example, it is known that excess 174 carotenoid accumulation impairs muscle function in other bird species<sup>46</sup>. Second, the yellow polymorphism could influence maternal investment. Fallahshahroudi et al.<sup>47</sup> found that chicken 175 mothers homozygous for the yellow skin allele, a BCO2 allele silenced in skin tissue<sup>19</sup>, invest 176 more carotenoids in egg yolk than other genotypes<sup>47</sup>. Consistent with this, G. scandens mothers 177 178 with the yellow genotype hatched eggs more successfully than heterozygous individuals (97% vs. 78%: Linear mixed effect model with year-hatched as a random effect,  $\gamma^2 = 12.1$ , P < 0.001, 179 df = 2; n = 138 nests; Figure 4B, Table S2). The pattern was repeated at different times and in 180 181 different age groups in the extended breeding season of 1983 (Table S3). Third, reduced BCO2 182 expression and carotenoid accumulation may alter spectral tuning in the avian retina, where 183 BCO2 is required for the biosynthesis of galloxanthin, a key apocarotenoid involved in shortwavelength vision<sup>44</sup>. In the absence of experimental data, visual perception differences among 184 185 morphs remains unexplored.

186 **Conclusions** 

187 The genetic basis of polymorphic traits in natural populations, and selection pressures acting on 188 them, are generally not known<sup>2</sup>, although there are a few outstanding exceptions involving 189 supergenes<sup>9–14</sup>. Here we have shown that variation at a single locus (*BCO2*) is responsible for a 190 beak color polymorphism in Darwin's finches. This synonymous mutation associated with the

191 vellow morph has an uncertain functional consequence, but it changes the highest frequency 192 valine codon to the least abundant in the finch genome, and it is associated with reduced BCO2 193 expression. We cannot completely exclude the possibility that this synonymous mutation may be 194 linked to one or several unidentified causal variants. However, it occurs on multiple haplotypes 195 (Figure S1C) and a careful analysis of our genome data did not reveal any other candidate 196 mutation or any haplotype pattern consistent with the presence of two independent causal 197 mutations (both associated with the synonymous variant). Identification of a sequence variant 198 linked to the phenotypic polymorphism has enabled us to trace its origin through phylogenetic 199 analysis to a single mutation occurring in the Galápagos archipelago approximately half a 200 million years ago. The polymorphism has been retained in all descendant species except one. The 201 most parsimonious explanation for its occurrence in many species is that the polymorphism is a 202 shared ancestral condition. Persistence across the radiation is notable because the rarer morph 203 may be lost through drift in the founding of new populations by a limited number of individuals during speciation<sup>2,6</sup>. Nonetheless, Darwin's finches satisfy two conditions that are conducive to 204 retention: high speciation rate<sup>48</sup> and presence of several coexisting, and occasionally 205 interbreeding, closely related species<sup>3,16,49</sup>. Loss through drift is likely to be counteracted by 206 reintroduction through introgression<sup>20,50</sup>, especially in the early stages of speciation. 207

Although we do not fully understand the salient selection pressures, we have identified diet as an important factor, because frequencies of the yellow allele in *G. scandens* are associated with changing diet availability in a cactus specialist. In this species, the frequency of the yellow allele decreased abruptly following a resource-induced crash, which may indicate that any advantage to the yellow allele is lost during prolonged periods of high stress lasting for one or more years. The yellow genotype is relatively common in cactus-specialist species feeding on

214 carotenoid-rich pollen and tend to survive better in some years when cactus products are 215 plentiful. The selective advantage of the yellow morph under certain environmental conditions 216 must be counterbalanced by a yet unknown selective advantage for the pink morph under other 217 conditions. A contributing factor may be selection that maintains the essential role of BCO2 in spectral tuning<sup>44</sup> or in carotenoid degradation and detoxification<sup>51,52</sup>. However, BCO2 expression 218 may be normal in the liver of the vellow morph, as in the chicken BCO2 mutation<sup>19</sup>, meaning the 219 220 yellow morph does not become toxified. We found evidence of higher hatching success of eggs 221 produced by females of G. scandens with the homozygous yellow genotype. This raises the 222 intriguing possibility that these females deposit more carotenoids in egg yolks than other genotypes, an important factor for egg quality in birds<sup>47,53,54</sup>, which may influence hatching 223 224 probability. Together, our results suggest that most of the time the yellow polymorphism is 225 approximately neutral, with morph frequencies occasionally perturbed by introgressive 226 hybridization and episodic fluctuations in selection. For other species carrying the yellow allele 227 at high frequency, such as the *Camarhynchus* tree finches, the frequency of the yellow allele has 228 not been studied, but diet may play a role, because all species uptake carotenoids from flowers or 229 caterpillars<sup>55</sup>.

Intraspecies color polymorphisms are exceedingly rare in birds (<3.5% of all birds<sup>35</sup>). Most color polymorphisms that have been studied to date are visible in adults and have signaling functions in contexts of mate choice<sup>11</sup>, social dominance<sup>56</sup>, camouflage<sup>7</sup> or protective mimicry<sup>57</sup>. The yellow beak polymorphism in Darwin's finches differs from all of these, and is more akin to polymorphisms in Major Histocompatibility Complex (MHC) antigens<sup>50</sup>, where fitness advantages are physiological and biochemical. Our study also contributes to recent advances in understanding the genetic basis of carotenoid-based traits<sup>58–65</sup> and a growing appreciation for the

- role of *BCO2* in carotenoid-based phenotypes in birds<sup>19,22-26</sup>, primarily in plumage, suggesting a
- common role for mediating yellow carotenoid-based traits. Since they are largely or entirely out
- of sight in adults, literally, such polymorphisms may be far more common than is currently
- 240 recognized, and contribute to diversification in many ways that are yet to be discovered.

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- 250

Author contributions: LA, PRG, BRG, and EDE conceived and designed the study. PRG and
 BRG collected blood samples and field observations. EDE conducted all bioinformatic and
 ecological analyses with input from PRG and BRG. CGS was responsible for all genomic

- 254 material preparation, sequencing, and pedigree analysis. CGS and HB conducted genotyping.
- 255 MD, OO, and AA collected samples and generated expression data. CJR generated the genome
- assembly. EDE wrote the first draft of the manuscript with input from LA, PRG, and BRG and
- all authors commented on the final manuscript. All authors approved the manuscript before
- 258 submission.259
- 260 Declaration of interests: Authors declare no competing interests.
- 261

262

# 263 Figure Legends

264

# 265 Figure 1: Genetic basis for a beak polymorphism in Darwin's finches.

266 (A) G. magnirostris nestlings with the yellow beak phenotype (left) or pink beak phenotype

(right). Images by P.R. Grant. (B) Association test result for nestling beak color showing the
 likelihood ratio test (LRT) statistic per-site genome-wide. The top plot shows *G. fortis* (n<sub>pink</sub> =

- $130, n_{vellow} = 130)$  and bottom *G. scandens* ( $n_{pink} = 98, n_{vellow} = 98$ ). (C) SNP association test
- results for *G. scandens* and *G. fortis* combined showing the region of strong association on
- 271 chr24. An exonic SNP in BCO2 is highlighted (LRT = 166.6). (D) Alignment of the first 68bp in
- exon 4 of *BCO2* from the yellow and pink allele, with amino acids indicated between
- alignments. The high LRT SNP from (C) is highlighted. (E) Phenotype to genotype matching
- using 1,631 individuals of known phenotype. Sample sizes for each group are marked above bars
- and per-species summaries are shown in Figure S2A. (F) Fragments Per Kilobase per Million
- 276 mapped reads (FPKM) of *BCO2* embryos of several finch species (Table S1), contrasted across 277 known genotypes (n = 35; AA = 3, GA = 6, GG = 26). *P*-values for Tukey's post-hoc analysis is
- known genotypes (n 55; AA 5; GA 6; GG = 26). *P*-values for Tukey's post-noc analysis 1 278 noted above each individual comparison and boxplot hinges correspond to the first and third
- 278 noted above each individual comparison and boxplot ninges correspond to the first and third 279 quantiles, center line is median, and whiskers mark 1.5x the interquartile range. Raw data are
- 219 quantiles, center line is median, and whiskers mark 1.5x the interquartile range. Raw dat 280 shown as points and jittered. See also Figure S1 and Figure S2.
- 280

# Figure 2: Frequency of the yellow allele across the Darwin's finch radiation.

Left, a species chronogram for Darwin's finches (reproduced from<sup>66</sup>), colored by genera, with

- the parsimonious origin of the yellow allele marked with a yellow star (previously dated at 546
- 285 KYA<sup>48</sup>). A yellow asterisk marks all species where the yellow nestling phenotype has been
- observed, a black asterisk indicates that the yellow nestling phenotype has not been observed,
- and no asterisk indicates that the nestling phenotype has not yet been studied in that species. *G. septentrionalis* has a different yellow phenotype (see text). Right, the frequency of the yellow
- allele (*BCO2* SNP p6166878, allele A) in all finch species with the number of individuals
- 290 genotyped marked along the vertical axis (including samples from<sup>48,66</sup>). See also Figure S2D.
- 291

# Figure 3: Changes in yellow genotype frequency over time in relation to cactus abundance in two Darwin's finch species on Daphne Major.

- 294 (A) Proportion of adult diet during early breeding in G. scandens and G. fortis (reproduced
- 295 from<sup>37</sup>). (B) Annual maximum *Opuntia* cactus representing fruit availability on Daphne Major.
- No data were collected in 2005. (C) Frequency of the yellow (A/A) genotype over 26 years for G.
- 297 fortis (blue) and G. scandens (red); points are scaled by sample size. Frequencies are plotted
- beginning in 1987, one year before blood sampling began. See also Figure S3A and Figure S4.
- 299

# Figure 4: Survival and hatching success in relation to genotype in *G. scandens*, the common cactus finch.

- 302 (A) Differences in survival to 1-year after hatching between the 1991 cohort and 1998 cohort.
- 303 Yellow genotype individuals survived better in 1991 than 1998, corresponding to years of high
- and low cactus production, respectively. (B) Lifetime hatching success for 22 mothers and 138
- 305 nests, colored by genotype. Yellow genotype mothers experienced greater hatching success than
- 306 heterozygotes. We cannot evaluate homozygous pink hatching success, for which we have only

- two mothers. 95% confidence intervals are shown and only comparisons with significance P < 0.05 are marked (see summaries in Table S2). See also Figure S3.

309

### 310 **STAR METHODS**

311

### 312 **RESOURCE AVAILABILITY**

313

314 Further information and requests for resources and reagents should be directed to and will be

315 fulfilled by the lead contacts, Erik D. Enbody (erik.enbody@gmail.com) and Leif Andersson 316 (leif.andersson@imbim.uu.se).

317

### 318 Materials availability

- 319 This study did not generate new unique reagents.
- 320

### 321 Data and code availability

- 322 Resequencing data is deposited at NCBI# PRJNA678752. The genome assembly for
- 323 Camarhynchus parvulus V1.1 can be found at NCBI# GCA 902806625.1. Data tables for
- 324 TaqMan genotypes, list of samples used for whole-genome resequencing, RNAseq sample
- 325 metadata, and code for bioinformatic analyses are uploaded to the GitHub page of EDE during
- 326 review, and will be archived at time of publication:
- 327 (https://github.com/erikenbody/Finch beak color polymorphism).
- 328

### 329 **METHODS DETAILS**

330

### 331 **Sample collection**

332 Blood was collected as part of a long-term monitoring of finches on Daphne Major and other

- 333 islands beginning with samples first collected in 1988. Sampling was conducted in accordance
- 334 with protocols of Princeton University's Animal Welfare Committee, and stored on EDTA-
- 335 soaked filter paper in Drierite to preserve red blood cells for DNA extraction later. Additional
- 336 details on sample collection can be found elsewhere<sup>3</sup>, as well as for samples collected on other islands and used in the TaqMan assays<sup>48,66</sup>. Nestlings were phenotyped for the beak color 337
- 338 polymorphism by eye and scored for the presence or absence of having extensive yellow on the
- 339 lower mandible. Although individual phenotypes change with age (the beak is eventually
- 340 covered by melanin), the beak color of nestlings is dichotomous in variation and binned visually
- 341 for the presence or absence of yellow. Among Daphne individuals, we selected all 98 G.
- 342 scandens and all 130 G. fortis carrying the yellow phenotype for which we have collected
- 343 genetic material for. We then selected an equal number of pink individuals per species (pink is
- 344 the more common phenotype, so there are more samples collected of the pink phenotype), for a
- 345 total of 456 samples used for low-coverage sequencing and genome-wide association analysis.
- 346 Later, we sequenced an additional 151 pink individuals to test for introgression (see section on
- 347 introgression below). This study includes in total 607 Darwin's finch whole-genome samples 348 sequenced at low coverage. Embryos (n = 35, Table S1) were collected on Santa Cruz and Pinta
- 349 according to<sup>67</sup>. Tissues were stored in RNAlater (ThermoFisher, CA) until further use.
- 350

### 351 **DNA extraction library preparation**

- 352 We extracted DNA from blood on filter paper using a custom salt preparation protocol. Briefly,
- 353 we submerged clipped blood on filter paper in a buffer containing 400mM NaCl, 2mM EDTA
- (pH 8.0), 10mM TrisHCl (pH 8.0), and dH20. Next, we added a freshly prepared buffer 354

containing 5% SDS, proteinase K (2mg/mL), and dH20. Samples were incubated overnight at
55°C, the filter paper removed, and 135µL of saturated NaCl was added to the mixture. The
sample tube was vortexed and spun down at 4,000 rpm for 15min at 4°C and the supernatant
transferred to a new 2mL tube. DNA was precipitated using 2 volumes 95% EtOH and mixed by

- inverting the tube. Finally, samples were spun at 13.3rpm for 3min to pellet the DNA, EtOH
- removed, and 50-200 LTE added to elute the DNA. DNA concentration was measured on a
- 361 Nanodrop (ThermoFisher, CA).
- 362

We generated fragment libraries for whole-genome sequencing using a custom Tn5-based tagmentation protocol based on<sup>68</sup>. Briefly, we assembled the Tn5 transposon construct using the stock Tn5 (prepared by Karolinska Institutet Protein Science Facility) and primers described in<sup>68</sup>:

366

367 Tn5MErev: 5;-[phos]CTG TCTCTTATACACATCT-3' 368

369 Tn5ME-A (Illumina FC-121-1030): 5'-TCGTCGGCAGCGTCAGATGTGTATAAGAGACAG-3' 370

371 Tn5ME- B (Illumina FC-121-1031): 5'-GTCTCGTGGGGCTCGGAGATGTGTA TAAGAGACAG-3'

372

373 Samples were tagmented by adding to a solution containing the Tn5 construct and H<sub>2</sub>0, 5x TAPs,

and 40% PEG. The mixture was incubated on a Bio-Rad thermocycler (Bio-Rad, CA) for 10min

at 55°C. Next, genomic libraries were PCR enriched using Kapa Biosystems HiFi HotStart

- 376 (Wilmington, MA) PCR kit (annealing temperature 63°C). DNA libraries were size selected
- 377 using .38X and .16X Ampure XP beads (Brea, CA) for a target insert size of 350bp, and

378 resulting product quantified on a Tecan microplate reader (Tecan Life Sciences, Switzerland)

using Qubit (ThermoFisher, CA) reagents. Samples were pooled in equimolar concentrations and

a final size selection performed using .45X and .3X Ampure beads on the resulting pool. Pools

were sequenced on an Illumina NovaSeq S4 flow cell (Illumina, CA) with a target sequencingdepth of 2x.

383

# 384 **TaqMan genotyping assay**

385 Custom SNP genotyping TaqMan assay (ThermoFisher, CA) were applied to perform genotypic

analysis of the SNP of interest in exon 4 of *BCO2*. We designed primers (BCO2\_F: 5'-

387 TGTTTCAGAACCCAGTGACAACT-3'; BCO2\_R:5'-TTCCAGTGTCTCTGGGTCCA-3') and

388 probes (BCO2\_VIC:5'-ATGTGAACTACGTGCTGTAC-3'; BCO2\_FAM:5'-

- 389 ATGTGAACTACGTACTGTAC-3') for the SNP of interest on chr24:6166878 (A/G). We used
- this assay to genotype 2,859 individuals, for which 1,631 had a known nestling beak phenotype.
- 391

# 392 RNA sample preparation and sequencing

393 We dissected the upper beak primordia of 35 Darwin's finch embryos (6 Geospiza magnirostris,

394 7 G. fortis, 7 G. fuliginosa, 8 Camarhynchus psittacula, 1 C. parvulus, and 6 Platyspiza

- 395 *crassirostris*) and extracted RNA with E.Z.N.A. Total RNA Kit I (Omega Bio-tek, GA). We
- 396 prepared cDNA libraries with the NEBNext Ultra RNA Library Prep Kit for Illumina
- 397 (NewEngland Biolabs, MA) with poly(A) selection. The libraries were then sequenced on HiSeq

398 4000 (Illumina, CA).

- 400 In order to search for splice variations, RT-PCR was applied to amplify the regions around
- 401 p6166878 in cDNA by the use of forward 5'- CCCATCCCAGCCAAGATCAA-3' and reverse

- 5'- CGTAGTGGGGATGAGCTGTG-3' primers under the following conditions: 95°C for 5 402 403 min, followed by 35 cycles of 95°C for 30 s, 60°C for 30 s and 72°C for 1 min. The amplified 404 fragments were subjected to Sanger sequencing where splice variants were searched for by eve 405 and none were detected.
- 406

### 407 ddPCR for allele specific expression in heterozygotes

- 408 Droplet digital PCR was performed using Bio-Rad QX100 Droplet Digital PCR system (Bio-
- 409 Rad, CA) to analyze allele-specific RNA expression. The reaction mix was prepared by using 11
- 410 ul of 2X ddPCR Supermix for Probes (no dUTP)(Bio-Rad, CA), 1.1 ul 20× Primer and Probe
- 411 Mix (final concentration of 800 and 300nM, respectively), 7.9 µl Nuclease free water, and 2 µl 412 reverse transcriptase product. Probes were reused from the TagMan analysis described above and
- 413 primers crossing exon-intron junctions are listed below. Twenty microliters of the prepared
- 414 mixture were loaded into a disposable droplet generator cartridge (Bio-Rad, CA), along with 70
- 415 µL of droplet generation oil for probe (Bio-Rad, CA) to generate droplet with the QX100 droplet
- 416 generator (Bio-Rad, CA). After generation, samples were transferred to a 96-well plate and
- 417 cycled in a C1000 touch Thermal Cycler (Bio-Rad, CA) under the following cycling protocol:
- 418 95°C for 10min, then 42 cycles of 95°C for 30s (denaturation) and 58°C for 120s (annealing),
- 419 followed by post-cycling steps of 98°C for 10min and an infinite 10°C hold. The ramp rate
- 420 among the steps of the amplification was adjusted to 1°C/sec. The cycled plate was then read in
- 421 the QX100 Droplet Reader (Bio-Rad. CA) and the data was analyzed with QuantaSoft software (Bio-Rad, CA). Primers:
- 422
- 423
- 424 BCO2 F2 - GCCACAACCCAGTGACAACT
- 425 BCO2 R1 - TTCCAGTGTCTCTGGGTCCA
- 426

### **OUANTIFICATION AND STATISTICAL ANALYSIS** 427

428

### 429 **Population genomics**

- 430 All Illumina short reads were mapped to the chromosome-scale Camarhynchus parvulus V1.1 genome assembly (GCA 902806625.1) using BWA mem v0.7.17<sup>69</sup> and the resulting BAMs 431 432 were sorted using SAMTOOLS v1.10 (http://www.htslib.org/). Sequencing coverage was
- 433 estimated for chromosome 4 (a large, representative subset) using the SAMTOOLS coverage
- 434 command. We used ANGSD v0.935-33-g79d9455 (Analysis of Next Generation Sequencing
- Data<sup>17</sup>) to estimate genotype likelihoods used for running association tests. We ran the following 435 436 commands for each species (G. scandens and G. fortis; n = 2 runs total), which outputs a beagle
- 437 file of genotype likelihoods:
  - 438

```
439
      $ANGSD PATH/angsd -b $BAMLIST1 -ref $REFGENOME -anc $REFGENOME -r $INTERVAL \
440
                    -out Results gwas/${POP1} ${POP2} ${INTERVAL}.ref \
441
                    -uniqueOnly 1 -remove bads 1 -only proper pairs 0 -trim 0 \
442
                    -minMapO 20 -minO 20 -doCounts 1 \
443
                    -domajorminor 1 -domaf 1 \setminus
444
                    -GL 1 -P 5 -doGlf 2 -SNP pval 1e-6 -minMaf 0.05 \
445
                    -dumpCounts 1
```

446

447 We next generated an estimate of relatedness by calculating principal components using all sites on the chromosome being run using PCAngsd v $1.01^{70}$ , and processed output using a custom R 448 449 script:

```
450
451
       python $PCANGSD/pcangsd/pcangsd.py -beagle
452
       Results_gwas/${POP1}_${POP2}_${INTERVAL}.ref.beagle.gz -o
453
       Results gwas/${POP1} ${POP2} ${INTERVAL}.ref.pcangsd -threads 5
454
455
       Rscript ~/bc/gwas angsd/get PCs.R
456
       Results gwas/${POP1} ${POP2} ${INTERVAL}.ref.pcangsd.cov
457
458
       The first two principal components were used as an estimate of relatedness to run the following
459
       association test in ANGSD, which performs the logistic regression score test described in<sup>18</sup> under
460
       a recessive model with pink coded as 0 and yellow coded as 1:
461
462
       $ANGSD PATH/angsd -doMaf 4 \
463
       -beagle Results gwas/${POP1} ${POP2} ${INTERVAL}.ref.beagle.gz \
464
       -fai $FAIFILE -yBin scandens bin pheno.txt -doAsso 2 -model recessive -cov
465
       Results gwas/${POP1} ${POP2} ${INTERVAL}.ref.PC1 PC2.txt \
466
       -out Results gwas/${POP1} ${POP2} ${INTERVAL}.ref.lrt.2.rec
467
468
       We extracted allele frequencies at each site used in the association test on chromosome 24 (n =
469
       111,890) by running ANGSD for G. fortis yellow, G. fortis pink, G. scandens yellow, and G.
470
       scandens pink (n = 4 \text{ runs}) with the following settings:
471
472
       $ANGSD PATH/angsd -b $BAMLIST1 -ref $REFGENOME -anc $ANCESTRAL \
473
                       -r $INTERVAL -sites $SITES \
                       -out Results_af/${POP1} ${INTERVAL} BALANCED.all sites \
474
475
                       -uniqueOnly \overline{1} -remove bads 1 -only proper pairs \overline{0} -trim 0 \setminus
476
                       -minMapQ 20 -minQ 20 -doCounts 1 \
477
                       -domajorminor 5 -domaf 2 \setminus
478
                       -GL 1 -P 5 -SNP pval 1e-6
479
480
       All subsequent analysis of association test results were performed using custom scripts in R
       v4.0.3<sup>71</sup> and various Tidvverse packages<sup>72</sup>.
481
482
       We created a neighbor-joining tree using PCAngsd v1.01<sup>70</sup> for all homozygous AA (n = 176)
483
484
       and GG individuals (n = 80) for the region 5-kb up and downstream of the p6166878 variant
485
       using a beagle file generated as described above for the association analysis. PCAngsd generates
486
       neighbor-joining trees based on a covariance matrix of individual allele frequencies.
487
488
       Bioinformatic analysis of introgression
489
490
       After the identification of the BCO2 variant of interest at p6166878, we selected an additional
491
       151 individuals to sequence at low-coverage whole-genome sequencing (mean depth = 1.9 \pm
492
       1.2X) to test the hypothesis that the frequency of the pink allele changes in the G. scandens
493
       population due to introgressive hybridization with G. fortis. We selected all the G. scandens
494
       samples homozygous for the pink allele and were hatched after the year 2008 as representative of
495
       samples collected "late" in the study (n = 44). We next randomly selected an equal number of G.
496
      fortis samples that were homozygous for the pink allele and were hatched after the year 2008 (n
497
       = 44). In order compare samples to those collected early in the study, we sequenced an equal
498
       number of randomly selected samples hatched between 1988 and 1995 for G. fortis (n = 44) and
```

499 G. scandens (n = 44), all homozygous for the pink allele. 25 of these samples were already

```
500
       sequenced for the genome-wide association study described earlier (i.e. 176 samples were used
501
       in this analysis, 151 of them uniquely generated for introgression analysis). We only selected
502
       homozygous pink alleles in order to evaluate if the frequency of the pink allele in the G.
503
       scandens population rose in frequency as a consequence of introgression from G. scandens.
504
505
       We used ANGSD v0.935-33-g79d9455 to generate allele frequency estimates for each of the
506
       four groups (early G. fortis, early G. scandens, late G. fortis, late G. scandens) separately.
507
508
       $ANGSD PATH/angsd -b $BAMLIST1 -ref $REFGENOME -anc $REFGENOME -r $INTERVAL \
509
                        -out Results fortis scandens/${POP1} ${INTERVAL}.ref \
510
                        -uniqueOnly 1 -remove bads 1 -only proper pairs 0 -trim 0 \
511
                        -minMapQ 20 -minQ 20 -doCounts 1 \
512
                        -GL 1 -P 8 \
513
                        -doSaf 1
514
515
       We next used the realSFS command in ANGSD to generate the 2d SFS (site frequency
516
       spectrum) which was used as input to the realSFS `fst index` command to calculate pairwise
517
       genomic divergence (i.e. F_{ST}) between the two species at both the early and late time points. F_{ST}
518
       was then summarized in 10,000-kb, non-overlapping, windows using the realSFS `fst stats2`
519
       command. We calculated mean genome-wide F_{ST} between G. fortis and G. scandens at both the
520
       early (pre-1995) and late (post-2008) time periods using these windowed-values. Per-site F_{ST}
521
       values were extracted for the region 5-kb upstream and downstream (Table S4) using the realSFS
522
       'fst print' command. We identified five SNPs exceeding the 99.95% percentile of early F_{ST}
523
       values in the vicinity of p6166878 using custom R-scripts. Allele frequencies within each of the
524
       four groups were additionally extracted using the `-domajorminor 5 -domaf 2` for the BCO2
525
       region in order to calculate the change of allele frequency between early and late G. scandens
526
       samples (Table S4). For this analysis, conducted using custom R-scripts, we set the major allele
       as the most common allele in early G. fortis samples.
527
528
529
       Analysis of high coverage data
530
       Short read sequencing data for 293 samples from 20 Darwin's finch species and two outgroup
531
       species was accessed from NCBI sequence read archive (www.ncbi.nlm.nih.gov/sra) BioProjects
       PRJNA392917<sup>66</sup>, PRJNA263122<sup>48</sup> and PRJNA301892<sup>73</sup>. Two homozygous vellow (A/A) from
532
533
       G. scandens and G. fortis (n = 4 total) that were sequenced at low coverage were also sequenced
534
       to a target coverage of 15x (this study, PRJNA678752). All short-reads were aligned to
       Camarhynchus parvulus V1.1 using BWA mem v0.7.17<sup>69</sup>. SNPs were called using GATK's
535
       HaplotypeCaller and joint genotyping using GenotypeGCVFs (v4.1.4.1<sup>74</sup>). Filtering was done for
536
537
       SNPs using filter-expressions in VariantFiltration and only biallelic SNPs were retained:
538
539
       "QUAL < 100 || MQ < 40.0 || MQ > 80.0 || MQRankSum < -4.0 || MQRankSum > 4.0
540
       || ReadPosRankSum < -4.0 || ReadPosRankSum > 4.0 || QD < 5.0 || FS > 30.0 ||
541
       DP < 50 || DP > 29300"
542
543
       And removing genotypes with low depth and low genotype quality using -G-filter:
544
545
       "DP < 1 || DP > 200 || GQ < 10"
546
547
       We searched for indels and SNPs that might be linked to p6166878 by searching the unfiltered
548
       joint-genotyped VCF for all SNPs and indels 200-kb upstream and downstream of p6166878.
```

- 549 We calculated allele frequency for homozygous alternative individuals (A/A and G/G) at this 550 position and removed variants with a minor allele frequency < 0.05. Delta allele frequency was 551 calculated as the difference in frequency of non-reference allele in individuals genotyped as
- homozygous yellow or homozygous pink based on SNP p6166878 (Figure S1E).
- 553

559

```
We extracted genotypes at the SNP position of interest (p6366878) using BCFtools v1.10 (http://www.htslib.org/):
```

```
556
557 bcftools query -r chr24:5966878-6366878 -f '[%GT\t]\n' $VCF > p
558 6366878_raw.genos
```

560 We calculated the frequency of the yellow allele (A) by counting the number of alternate alleles 561 per species and dividing by 2n (n = number of individuals per population). Allele frequencies 562 depicted in Figure 2 include a combination of high coverage samples (n = 293) and samples that 563 were individually genotyped (see below, n = 2,859), but we omitted samples that were 564 determined from field observations to be of hybrid origin in Figure 2.

565

566 In order to approximate the timing of the appearance of the yellow allele (A), we placed the

origin of the polymorphism on an existing Darwin's finch phylogeny. We downloaded a recent
 phylogenetic hypothesis for all species that used a maximum-likelihood approach on a

569 concatenated SNP matrix to generate a tree (<sup>66</sup>, <u>https://treebase.org/treebase-</u>

- 570 <u>web/search/study/summary.html?id=21803</u>). We pruned this tree to one branch per species and
- 571 converted the tree to an ultrametric tree using makeChronosCalib to set root time to 1 MYA

based on previous estimates<sup>48</sup> with the R package ape<sup>75</sup>. We then converted the original tree to a
chronos time tree using the following command:

```
575 timetree <- chronos(finch.pruned.tre, lambda = 1, model = "correlated",
576 calibration = mycalibration, control = chronos.control() )
```

577

578 The final plot was produced using ggtree for the base tree<sup>76</sup> and adding the bar plots of allele 579 frequency<sup>77</sup>. The estimated divergence time between *P. crassirostris* (lacking the A allele) and 580 all *Geospiza, Camarhynchus,* and *Pinaroloxias* species was estimated using this method as 456 581 KYA compared to  $546\pm74$  reported in a previous analysis<sup>48</sup>.

582

For all species in the dataset that carry the yellow allele (A, n = 15), we tabulated the number of SNPs that are shared between  $n=\{1..15\}$  species. Specifically, for each SNP called (n = 26,056,248) in the dataset, we summed the number of species carrying at least 1 alternate allele at that position. Allele frequencies per species were tallied using bcftools, summed using custom

- 587 bash scripts (allele\_sharing.sh), and plotted using custom R scripts (plot\_af.R).
- 588

# 589 Allele specific expression

- 590 For 29 individuals for which we have RNAseq data, we also had genomic DNA available. For
- these individuals we used the same TaqMan assay to determine genotype. For all other
- individuals we inferred genotype based on RNA sequencing depth. This includes 1 AA, 4 GG,
- and 1 GA individuals. Before mapping RNAseq data, Illumina adaptor and primer sequences
- 594 were removed with CutAdapt v.1.9<sup>78</sup> and low-quality bases (PHRED < 20) were removed using
- 595 Trim Galore (v.0.4.1, available at

596 http://www.bioinformatics.babraham.ac.uk/projects/trim galore/) with default settings. Cleaned 597 RNAseq reads were then mapped to the genome and Fragments Per Kilobase of transcript per 598 Million mapped reads (FPKM) was calculated using the HISAT2 (v. 2.1.0) - StringTie (v. 1.3.6) - Ballgown (v. 2.20.0) pipeline<sup>79</sup>. For the 6 individuals we genotyped using RNAseq data, 599 genotypes were called using SAMTOOLS and BCFtools v1. 9: 600

601

```
602
      samtools mpileup -Q 20 -q 20 -t DP4,DP -vuf ${ref} *.bam | bcftools call -M -
603
      f GQ -mq 3 -Ov > snps.vcf
```

604

605 We compared the number of Fragments Per Kilobase of transcript per Million mapped reads 606 using a three-way ANOVA with the aov command and post-hoc comparisons using the 607 TukevHSD command in R (v4.0.3).

608

### 609 Codon usage bias

610 We queried the Codon and Codon Pair Usage Tables (CoCoPUTs<sup>80</sup>) for the reference genome

assembly build for Camarhynchus parvulus V1.1. The CoCoPUTs tool pulls from the GenBak 611

612 refseq database to compute codon usage for publically available data. The frequency of the four

- 613 valine codons in the C. parvulus genome are as follows:
- 614
- 615 GTT = 13.50 %
- 616 GTC = 12.61 %
- 617 GTA = 7.63 %
- 618 GTG = 27.25 %
- 619

### 620 **Cactus fruit abundance**

621 We counted the number of flowers and fruits on 10 marked *Opuntia* bushes at 7 to 10-day 622 intervals during each annual visit to Daphne Major Island. We focus on fruits as a measure of

623 annual food availability because the sample of fruits represents all flowers produced up to that

624 time. For an analysis of survival from hatching to the beginning of the following year we focus

625 on flowers as a source of pollen and nectar at the time that nestlings are fed. Flower numbers for

626 1991 and 1998 are shown in Figure S3B. Flowering typically begins in October or November.

- 627 Flowers and fruits were counted in January, February and occasionally later at the end of the
- 628 flowering season.
- 629

### 630 Survival analysis and hatching success

631 We used a 2-sided Pearson's chi-squared test in R (v4.0.3) to test for a difference in first-year

- 632 survival among genotypes for all G. scandens (n = 326) and G. fortis (n = 964) nestlings
- 633 captured with genetic samples during regular nest monitoring that took place between 1978 and
- 634 1998. When analyzing survival of the two morphs in G. scandens across all years, we used a 2-
- 635 sided Pearson's chi-squared test with a Yate's continuity correction in R (v4.0.3) to test for
- 636 differences in morph survival for 2065 nestlings with known phenotypes (see note S3). The
- 637 difference in sample size between these two approaches is due to the larger number of
- 638 individuals with phenotypic data than were available for genetic analysis (i.e. not all nestlings
- 639 had samples for genetic analysis collected).
- 640
- 641 We modeled first year survival in 1991 and 1998 using a generalized linear model (GLM) in R

- 643 *BCO2* genotype, year, and the interaction between the two to test the hypothesis that first year
- 644 survival differed between 1991 and 1998. We used the tab\_model function from sjPlot to 645 summarize models<sup>81</sup> and report odds ratios.
- 646

647 We analyzed lifetime mean female hatching success in *G. scandens* using linear mixed models in

648 the package  $ImerTest^{82}$ . Mean hatching rate was calculated per nest as the ratio of number of 649 hatched divided by the total number of eggs laid. Consequently, hatching rate was only tabulated

650 for individuals where the number of eggs laid and hatched were known. We removed nests

651 where no eggs hatched (which could have been the result of other factors, such as nest predation)

and birds with only one nest (which prevented a reliable mean rate across nests). One yellow

653 phenotype individual was included as A/A who failed to amplify using the TaqMan assay. Each

654 individual was given a single value for lifetime average hatching success, which was used as the

response variable in the LMM. We included cohort (year hatched) of each female as a random

effect in the model. We used the Anova command (type = "II") from the car package<sup>83</sup> to test

657 significance of predictor effects and tab\_model from sjPlot to summarize models<sup>81</sup>.

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# KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Biological Samples		
Darwin's finch blood samples	This paper	https://github.com/erikenbody/Finch_beak_colo
Chemicals, Peptides, and Recombinant Proteins		• • • • • • • • • • • • • • • • • • •
Tn5 Transposon	Karolinska Institutet Protein Science Facility	Addgene #79107
Critical Commercial Assays		
TaqMan Assay	ThermoFisher Scientific	Cat#: 4316034
Kapa Biosystems HiFi HotStart	Roche	Cat#: 7958927001
NEBNext Ultra RNA Library Prep Kit	New England Biolabs	Cat#: E7530L
E.Z.N.A. Total RNA Kit I	Omega Bio-Tek	R6834-01
Deposited Data		
Darwin's finch resequencing data	This project	PRJNA678752
Darwin's finch resequencing data (high depth)	64	PRJNA392917
Darwin's finch resequencing data (high depth)	46	PRJNA263122
Darwin's finch resequencing data (high depth)	71	PRJNA301892
Camarhynchus_parvulus_V1.1 reference genome	Rubin and Enbody et al. <i>in prep</i>	GCA_902806625.1
Darwin's finch phylogeny	46	21803
Oligonucleotides		
Tn5MErev, 59-[phos]CTG TCTCTTATACACATCT- 39	66	
Tn5ME-A (Illumina FC-121-1030), 59- TCGTCGGCAGCGTCAGATGTGTATAAGAGACAG- 39	66	
Tn5ME- B (Illumina FC-121-1031), 59- GTCTCGTGGGCTCGGAGATGTGTA TAAGAGACAG-39	66	
BCO2_F: 5'-TGTTTCAGAACCCAGTGACAACT-3'	This study	
BCO2_R:5'-TTCCAGTGTCTCTGGGTCCA-3'	This study	
BCO2_VIC:5'-ATGTGAACTACGTGCTGTAC-3'	This study	
BCO2_FAM:5'-ATGTGAACTACGTACTGTAC-3'	This study	
5'- CCCATCCCAGCCAAGATCAA-3'	This study	
5'- CGTAGTGGGGATGAGCTGTG-3'	This study	
BCO2_F2 - GCCACAACCCAGTGACAACT	This study	
BCO2_R1 - TTCCAGTGTCTCTGGGTCCA	This study	
Software and Algorithms		
BWA mem v0.7.17	67	
	1	1

SAMTOOLS v1.10	http://www.htslib.or	
	<u>g/</u>	
ANGSD v0.935-33-g79d9455	17	
PCAngsd v1.01	68	
R v4.0.3	69	
Tidyverse R packages	70	
GATK v4.1.4.1	72	
BCFtools v1.10	http://www.htslib.or	
Ape R package	73	
ggtree R package	74	
CutAdapt v.1.9	76	
Trim Galore v.0.4.1	http://www.bioinfor matics.babraham.a c.uk/projects/trim_g alore/	
HISAT2 v. 2.1.0	77	
StringTie v1.3.6	77	
Ballgown v2.20.0	77	
ImerTest R package	80	
car R package	81	
sjPlot	79	







