Genome divergence and diversification within a geographic mosaic of coevolution

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Abstract

Despite substantial interest in coevolution’s role in diversification, examples of coevolution contributing to speciation have been elusive. Here, we build upon past studies that have shown both coevolution between South Hills crossbills and lodgepole pine (Pinus contorta), and high levels of reproductive isolation between South Hills crossbills and other ecotypes in the North American red crossbill (Loxia curvirostra) complex. We used genotyping by sequencing to generate population genomic data and applied phylogenetic and population genetic analyses to characterize the genetic structure within and among nine of the ecotypes. Although genome-wide divergence was slight between ecotypes ($F_{ST} = 0.011–0.035$), we found evidence of relative genetic differentiation (as measured by $F_{ST}$) between and genetic cohesiveness within many of them. As expected for nomadic and opportunistic breeders, we detected no evidence of isolation by distance. The one sedentary ecotype, the South Hills crossbill, was genetically most distinct because of elevated divergence at a small number of loci rather than pronounced overall genome-wide divergence. These findings suggest that mechanisms related to recent local coevolution between South Hills crossbills and lodgepole pine (e.g. strong resource-based density dependence limiting gene flow) have been associated with genome divergence in the face of gene flow. Our results further characterize a striking example of coevolution driving speciation within perhaps as little as 6000 years.

Keywords: coevolution, divergent selection, ecological speciation, genetic differentiation, Loxia, population genomics

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Introduction

Coevolution, the process of reciprocal adaptation by two or more species in response to reciprocal selection, is thought to be a major driver of biological diversification (Ehrlich & Raven 1964; Thompson 1994, 2005). However, demonstrating coevolution has been challenging (Gomulkiewicz et al. 2007). Moreover, few studies link coevolution directly to speciation and diversification (Althoff et al. 2014; Hembry et al. 2014). Coevolution between the South Hills crossbill (Loxia curvirostra complex) and Rocky Mountain lodgepole pine (Pinus contorta latifolia) is one of the best-documented examples of coevolution (Thompson 2005; Gomulkiewicz et al. 2007), and of coevolution generating reproductive isolation (Althoff et al. 2014; Hembry et al. 2014). In the absence of the red squirrel (Tamiasciurus hudsonicus; a predispersal seed predator), crossbills in the South Hills, Idaho, USA, are resident and much more abundant and exert stronger selection on lodgepole pine cones, causing the evolution of enhanced seed defences directed at crossbills. Where red squirrels occur, they are superior competitors for the lodgepole pine seeds and crossbills are much less abundant. Under these conditions, cones evolve mostly in response to selection

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exerted by red squirrels rather than crossbills and have less crossbill-directed defence, which favours smaller beaked crossbills (Benkman 1999; Benkman et al. 2001, 2003). The result is a geographic mosaic of coevolution, where crossbills coevolve in an arms race with lodgepole pine in the absence of red squirrels but not in their presence (Benkman 1999; Benkman et al. 2001, 2003, 2013).

The South Hills crossbill is one of 10 morphologically and vocally differentiated ecotypes (‘call types’) of the North American red crossbill complex (Groth 1993; Benkman et al. 2009; Irwin 2010) that have evolved in response to selection for specialization on seeds in the cones of different conifer species (Benkman 1993; Benkman et al. 2003). This hypothesis was tested in past studies that quantified feeding performance for five ecotypes, including the South Hills crossbill (Type 9) and the ecotype specialized on lodgepole pine where red squirrels occur (Type 5; Benkman 1993, 2003). Feeding performance varied in relation to beak depth (influences efficiency of seed extraction from conifer cones) and groove width in the horny palate (influences seed husking ability), with each ecotype having beak traits that approximate the predicted optima for foraging on seeds of their ‘key’ conifer (i.e., conifers that reliably produce and hold seeds in cones; Benkman 1993, 2003). The close fit between trait means and both their predicted optima and survival selection strongly implicates resource-based divergent selection in driving this adaptive radiation (Benkman 1993, 2003).

Because divergent selection can reduce gene flow, divergent selection could lead to genetic differentiation, even in the absence of geographic isolation (Endler 1973; Nosil et al. 2008; Shafer & Wolf 2013). In crossbills, reproductive isolation is related to divergent selection among ecotypes, because of expected lower fitness of potentially intermediate (hybrid) phenotypes, habitat isolation, low immigrant reproduction and several forms of behavioural isolation (Smith & Benkman 2007; Snowberg & Benkman 2007, 2009; Smith et al. 2012). In the South Hills, the combination of well-defended lodgepole pine cones (Benkman 1999; Benkman et al. 2001, 2003, 2013), local adaptation by South Hills crossbills and strong density-dependent food limitation prevents all but a few individuals of the nonlocally adapted ecotypes from persisting prior to and during pairing by South Hills crossbills (Smith & Benkman 2007; see Bolnick 2011). Strong density dependence arises because of very stable seed renewal in the South Hills, unlike the episodic abundance of resources that other ecotypes experience (Benkman et al. 2012). This contributes to the high levels of premating reproductive isolation between the South Hills crossbill and the two other ecotypes that breed in the South Hills [0.999 on a scale from 0 (no isolation) to 1 (complete reproductive isolation); Smith & Benkman 2007; Benkman et al. 2009]. However, whether local adaptation and strong contemporary premating reproductive isolation have been of sufficient duration to cause genome divergence is unknown.

The ecology and evolutionary history of the red crossbill complex have resulted in limited genetic differentiation among the ecotypes. Ecotype diversification probably occurred in the Holocene following the retreat of glaciers and expansion of conifers (Benkman 1993). The distributions of some key conifers relied upon by the ecotypes were so restricted in the late Pleistocene (e.g. coastal Douglas fir Pseudotsuga m. menziesii and Rocky Mountain ponderosa pine P. ponderosa scopulorum; Gugger et al. 2010; Potter et al. 2013) that frequent regional cone crop failures likely prevented earlier specialization by crossbills (Benkman 1993). Such recent divergence is indicated by coalescent analyses of European common (red) crossbill mtDNA haplotypes suggesting that ecotype diversification occurred rapidly over the last 11 000 years (Bjorklund et al. 2013). Patterns of mtDNA divergence and diversity are similar among North American ecotypes (Questiau et al. 1999). In addition, due to regular localized cone crop failures most ecotypes are nomadic and move long distances between natal and breeding locations, and between breeding locations (up to ~3000 km; Newton 2006), often breeding opportunistically and sympatrically in areas with abundant conifer seeds (Groth 1993; Summers et al. 2007). In the South Hills, multiple ecotypes breed (Smith & Benkman 2007). Such conditions and behaviour likely allowed extensive gene flow throughout the young radiation. Indeed, prior analyses using mtDNA and AFLPs found little evidence for genetic differentiation among ecotypes (Questiau et al. 1999; Parchman et al. 2006). Nonetheless, the widely sympatric occurrence of distinct ecotypes suggests that adaptation and reproductive isolation have evolved despite the large potential for homogenizing gene flow.

Recent innovations in DNA sequencing have dramatically increased our ability to address how geography, ecology and history shape genome divergence during the early phases of divergence (Alcaide et al. 2014; Lamichhaney et al. 2015; Martin et al. 2015). Here, we use genotyping by sequencing (GBS) to evaluate the pattern and extent of genome divergence across the North American red crossbill complex. We sampled multiple, geographically dispersed populations within each ecotype (Table S1, Fig. S1, Supporting information) to test the hypothesis that ecotypes are genetically cohesive and differentiated from one another. We tested for isolation by ecology, in the form of the evidence that genetic divergence was associated with divergent
selection (Nosil et al. 2008; Shafer & Wolf 2013), but anticipate little geographic structure (isolation by distance) based on the aforementioned biology of crossbills. Although our focus is on the South Hills crossbill, we include nine ecotypes to provide a broader context of divergence among closely related forms, and because the form of divergent selection and opportunities for geographic isolation are not uniquely different because of coevolution. Finally, the fossil record indicates a large reduction in the amount of lodgepole pine in the South Hills region (Mehringer 1985; Davis et al. 1986) during a several thousand-year warm period centred around 6000 BP (Bartlein et al. 2014). Because this could have prevented the persistence of a local crossbill population (See Siepielski & Benkman 2005), we estimate the distribution of lodgepole pine 6000 BP to characterize the time period over which this coevolutionary interaction could have persisted.

Materials and methods

DNA sequencing, assembly and variant calling

We sequenced DNA from 219 red crossbills representing nine morphologically and vocally differentiated ecotypes, as well as 12 white-winged crossbills (L. l. leucoptera; Table S1, Fig. S1, Supporting information). We utilized a GBS protocol that we have used in previous studies (Gompert et al. 2012; Nosil et al. 2012; Parchman et al. 2012, 2013), and generated three lanes of single-end 100-base sequencing on an Illumina HiSeq 2000 at the National Center for Genome Resources (Santa Fe, NM, USA). We used Perl scripts to remove contaminant DNA, trim barcodes and match barcodes to individual sample information. We first used seqman ngen 3.0.4 (DNASTAR) to perform a de novo assembly for a subset of 30 million reads sampled randomly from the sequencing data for all individuals. The purpose of this step was to produce a consensus GBS reference of the genomic regions represented in our libraries. Because our library preparation method produces reads identical in length from genomic regions beginning with EcoRI cut sites, reads typically align neatly into rectangular contigs, the consensus sequences of which represent a reference of genomic regions sampled by GBS. After removing low-quality or overassembled contigs, we generated a reference of 349 865 contig consensus sequences. We then aligned all reads for each individual onto the GBS reference using bwa 0.7.8 (Li & Durbin 2009). We used samtools 1.19 and bcftools 1.19 (Li et al. 2009) to identify bi-allelic single nucleotide polymorphisms (SNPs), and only called variants when 98% of the individuals had at least one read. We used this high threshold here to obtain SNPs with higher coverage and genotypes having relatively high levels of statistical certainty. While we could have called more SNPs with a lower threshold, we were interested mostly in genome-wide parameter estimates for this study and hope to generate more comprehensive resequencing data for locus-specific analyses in the future. We randomly selected a single variant from each contig to increase the independence of loci, and limited analyses to loci with minor allele frequencies >0.03 for population genetic analyses. Further details on assembly and variant calling are in the Supporting information.

Phylogenetic analyses

Phylogenetic analyses included the 219 red crossbills, and 12 white-winged crossbills for use as an outgroup. We first generated a second data set of SNPs present in alignments of both crossbill species using bwa, samtools and bcftools. While calling variants, we disregarded insertions and deletions, and only considered SNPs when 95% of the individuals had at least one read at that locus. We used a custom Perl script to produce a multiple alignment by defining the DNA state of each individual and variant as the genotype with the highest likelihood. Heterozygotes were coded using IUPAC ambiguities (i.e. M for A/C, R for A/G, W for A/T, S for C/G, Y for C/T and K for G/T), and loci with too much uncertainty (i.e. equal likelihoods for the three genotypes) were encoded as missing data. This resulted in a multiple alignment of 238 615 positions and 231 individuals. We inferred maximum-likelihood (ML) trees using examl 2.0.4 (Stamatakis & Aberer 2013), and executed 25 independent ML searches using as starting points 25 parsimony trees inferred using parsimony 1.0.3 (Stamatakis 2014). We conducted ML inferences using a GTR + Γ substitution model and performed 500 bootstrap replicates using examl with a GTR substitution model using the CAT approximation. We produced the bootstrapped ML analysis with raxml 8.0.20 (Stamatakis 2014) and used parsimonator as before to obtain starting trees from every alignment. We summarized the ML analyses using raxml in two different ways: (i) drawing bootstrap support values onto the best-supported ML tree and (ii) computing a bootstrap consensus tree using the majority rule extended criterion.

Population genetic analyses

We used analyses based on allele frequencies and genotype probabilities to quantify patterns of genetic structure within and among the red crossbill ecotypes using the 18 385 high-coverage SNPs described above. We estimated population allele frequencies and genotype
probabilities based on genotype likelihoods estimated with 

SCFTools using a hierarchical Bayesian model (Gompert et al. 2012). This model treats population allele frequencies and individual genotypes as unknown model parameters and utilizes Markov Chain Monte Carlo (MCMC). We used this model to estimate allele frequencies across populations within ecotypes for significant differentiation among populations (Gompert et al. 2013). We ran MCMC chains for 20 000 steps, discarded 5000 as burn-in and recorded every fifth step. We first summarized genotypic variation across all individuals of the red crossbill using principal components analysis (PCA). We generated a genetic covariance matrix based on the genotype point estimates for each bird and performed the PCA on this genetic covariance matrix using the PRCOMP function in R (Core Team, 2013). We tested for significant differentiation between ecotypes and for significant differentiation among populations within ecotypes using permutational multivariate analysis of variance (PERMANOVA; Anderson 2001) based on Euclidian distances of the first two principal components using the vegan package in R (Oksanen et al. 2013).

We used allele frequency estimates to calculate Nei’s genetic distance (Nei’s D; Nei 1972) among ecotypes and among all samples (populations) within ecotypes. We calculated pairwise Hudson’s $F_{ST}$ (Hudson et al. 1992) based on estimated allele frequencies at all loci for each ecotype and each population, using code written in R. We generated ML estimates of the folded-site allele frequency spectrum, nucleotide diversity ($\pi$) and expected heterozygosity as indicators of genetic variation within each ecotype using the expectation maximization algorithm of Li (2011) as implemented in SAMTOOLS and ran the algorithm for 20 iterations for each population.

We further investigated hierarchical patterns of genetic structure across the ecotypes and populations within ecotypes using a hierarchical Bayesian model that is similar to the correlated allele frequency model of STRUCTURE (Pritchard et al. 2000; Falush et al. 2003). We used this model (hereafter ENTROPY, described in Gompert et al. 2014) to characterize population structure and estimate admixture proportions for individuals in the absence of information on sample origin. Importantly, ENTROPY allows for stochastic variation in sequence coverage across individuals and loci and estimates allele frequency and genotype probability parameters along with admixture proportions. Similar to the admixture model in STRUCTURE, ENTROPY assumes that the genome of each individual consists of loci with ancestry from one of $k$ ancestral populations and makes no a priori assumptions about the population or cluster origin of individual samples. Admixture proportions, which represent the fraction of an individual’s genome inherited from each of the $k$ clusters, are estimated for each individual. In addition, ENTROPY generates estimates of deviance information criterion (DIC) as a metric for model choice and comparison; models with lower DIC values are those that fit the data better (Gompert et al. 2014).

To facilitate the convergence and stabilization of MCMC chains, we initialized individual admixture proportions in the chains using probabilities of cluster membership based on $k$-means clustering of the principal component scores (equivalent to a no-admixture model; Falush et al. 2003). Specifically, we used $k$-means clustering (kMEANS package in R) based on the principal components estimated from genotypes in a linear discriminant analysis (LDA package in R; Jombart et al. 2010). This provided reasonable starting values of $q$ to initialize MCMC and ensured proper mixing and convergence of MCMC chains. Importantly, this approach uses genotypic data without reference to sample origin and does not constrain posterior sampling. We ran ENTROPY separately for predefined values of $k = 1$–9 and ran five independent chains for each $k$. Each chain used the probability of cluster membership as mean expectation for the admixture proportion $q$, but random deviates with a precision scalar of 20 were drawn from a Dirichlet distribution to initialize $q$ for each chain. We used an upper value of 9 for $k$, representing the number of ecotypes included in our analysis. We ran each MCMC chain for 80 000 steps following 60 000 steps that were discarded as burn-in and saved every 10th step. We estimated posterior medians, and 95% credible intervals for parameters of interest. We checked for mixing and convergence of posterior parameter estimates by plotting MCMC steps for different parameter sets and inspected mixing during the burn-in period and convergence among chains.

The localities sampled are geographically separated and could have allele frequencies that differ due to genetic drift, with population homogenization due to migration declining with distance, leading to isolation by distance. Likewise, divergent selection could reduce gene flow and potentially lead to differences in population allele frequencies (Endler 1973; Nosil et al. 2008; Shafer & Wolf 2013). To investigate the extent to which allele frequency differences can be attributed to geographic and phenotypic distances between populations, we modelled pairwise genetic distances (Nei’s $D$) between populations as a function of geographic and phenotypic distances (difference in mean beak depth) between populations. Beak depth data (sample
sizes for males and females in parentheses) were from Groth (1993) for ecotypes 1 (39 and 33) and 7 (5 and 1), unpublished measurements of live birds and museum specimens by CWB for ecotypes 2 (226 and 149), 3 (47 and 34), 4 (25 and 15), 5 (61 and 31) and 6 (150 and 71), Benkman et al. (2013) for ecotype 9 (471 and 335) and Irwin (2010) for ecotype 10 (54 and 35). Phenotypic distances were calculated based on differences between ecotype means. The use of mean trait values should be conservative, because it provides less power to detect patterns than analyses based on measurements from each individual. Geographic distances were calculated based on Haversine distances, as implemented in the R package FOSSIL (Vavrek 2011). Geographic and phenotypic distances were normalized (transformed to Z-scores) so that their coefficients would be on the same scale. Genetic distances were logit-transformed and centred on the mean so as to not be bounded by zero and one. We used a Bayesian linear model that did not require all observations in the response variable to be independent, but instead modelled random effects for all population pairs (Clarke et al. 2002; Gompert et al. 2014) and over all coefficients for geographic and genetic distances, and separately for phenotypic and genetic distances. The model was specified in JAGS (version 3.4; Plummer 2003), and samples were gathered from the R interface to JAGS (RJAGS; R core Team 2013). After discarding 2000 steps as burn-in, we obtained 2000 samples of the posterior distributions from each of three chains, by retaining every fifth iteration of 10 000 MCMC steps. All chains were inspected graphically for adequate convergence and mixing.

Past forest distribution estimation

Random Forests (Breiman 2001) is a nonparametric classification and regression tree approach that we used to model the distribution of lodgepole pine, because of its past success when true absence data are available, and its ability to identify nonlinear relationships and interaction terms (Cutler et al. 2007). We used the USFS Forest Inventory and Analysis data (O’Connell et al. 2014) for 3406 presences and 10 855 absences of lodgepole pine in Idaho, Montana, Wyoming, Colorado and Utah to estimate the distribution of Rocky Mountain lodgepole pine. This was augmented with 50 000 ‘likely-absence’ points randomly placed within cells classified as ‘unforested’ in the LANDFIRE Forest Canopy Cover data set; this likely-absence data set was reduced by 59 points by eliminating all points within 1 km of lodgepole pine presence points to allow for error or lack of precision in the canopy cover layer, resulting in a total of 60 796 absences.

Covariates used for modelling were the BIOCLIM set (Hijmans et al. 2005) and elevation (Gesch et al. 2002). BIOCLIM values are similar to climate predictors used to model the distribution of tree species in other studies, including lodgepole pine, by other researchers (Boucher–Lalonde et al. 2012; Bell et al. 2014), but have the advantage of having been modelled for the mid-Holocene (6000 BP). Covariates were resampled to 1 km spatial resolution for modelling. We iteratively generated 100 models, each of which used a subsample of the absence data so that there were three times the number of absence points as presence points. This was done because Random Forests performs poorly when classes are highly imbalanced (Chen et al. 2004). The 100 models were combined into a single classification model. Summary statistics and graphs of covariance convergence for the subsampled absence data were used to evaluate the stability of the model. The out-of-bag (Breiman 2001) error rate in predicting known presences and absences was 9%. This model was applied to the historical distribution of climate variables in the BIOCLIM data sets to predict the past distribution of lodgepole pine. We estimated the relative amount of lodgepole pine forest 6000 BP compared with the current amount based on the combined area and relative probabilities of occurrence during the two time periods.

Results

After removing barcodes from the raw reads, and discarding contaminant reads, we retained 321 627 388 reads representing all 231 individuals. Initial de novo assembly placed 24 352 918 reads into 403 678 contigs; the 349 865 highest quality contigs from this assembly were used as a GBS reference. We subsequently aligned reads from all individuals to this reference using BWA. After using SAMTOOLS and BCFTOOLS to call variant sites, discarding loci with minor allele frequency <0.03, and randomly sampling a single SNP per contig, we retained a final set of 18 385 SNPs (mean coverage per individual per locus of 7.2×) for population genetic analyses across the red crossbill complex. Phylogenetic analyses were based on a set of 238 615 SNPs that were called in the alignments of red crossbills and white-winged crossbills, as described above and in the Supporting information.

Phylogenetic analyses revealed topologies with white-winged crossbills and red crossbills each forming strongly supported monophyletic groups (Fig. 1), consistent with previous studies (Questiau et al. 1999; Parchman et al. 2006). In contrast to past studies (Parchman et al. 2006), South Hills crossbills formed a strongly supported monophyletic group and were the only monophyletic red crossbill lineage (Fig. 1). Individuals
of the remaining ecotypes were dispersed throughout the tree and showed no evidence of clustering, with the exception of Type 6, for which bootstrap support was weak (Fig. 1). Type 6 is the only ecotype that could be considered allopatric to the other ecotypes, as it is confined mainly to Mexico south of the other ecotypes (Groth 1993). Consistent with phylogenetic analyses, population genetic analyses based on 18 385 SNPs revealed low levels of genome-wide genetic differentiation between the different ecotypes, as indicated by small Nei’s D and F_{ST} estimates (mean F_{ST} = 0.021, range: 0.011–0.035; Table S2, Supporting information).

Despite low levels of divergence, genetic differentiation among the ecotypes and similarity among geographically dispersed samples within individual ecotypes was evident in the PCA (Fig. 2), where the first two principal components differed significantly among ecotypes (PERMANOVA, F_{R, 210} = 1220; R^2 = 0.98; P < 0.001). South Hills crossbills were the most distinct in these analyses and were separated from other ecotypes along PC1, while the remaining ecotypes were separated mostly along PC2 (Fig. 2). The four smallest ecotypes (types 1, 3, 4, and 10) have the highest PC2 scores (Fig. 2), three of which are found mostly in the Pacific Northwest (types 3, 4, and 10), whereas Type 1 is the one ecotype found mostly in eastern North America (Fig. S1, Supporting information; Groth 1993). Intermediate-sized ecotypes are found in the middle cluster, including the two most abundant ecotypes in the Rocky Mountain region (types 2 and 5) and Type 7, which is uncommon but found within the geographic ranges of types 2 and 5 in the northern Rocky Mountains and west to the Cascades (C. W. Benkman, personal observations; Groth 1993). The largest ecotype is Type 6, which has the smallest PC2 scores (Fig. 2) and occurs mostly in Mexico, allopatric to the other ecotypes. Geographically dispersed samples from within the same ecotype overlapped extensively in PC space, indicating genetic cohesiveness within ecotypes.

Support for the distinctiveness of the South Hills crossbill and genetic similarity of geographically dispersed samples within each of the ecotypes was also found in Bayesian clustering analyses (ENTROPY; Gompert et al. 2014). DIC values were similar from k = 2 through k = 6 (Table S3, Supporting information), and all five models led to conclusions consistent with PCAs above. Inspection of MCMC chains indicated sufficient mixing and convergence only for k ≤ 6 models. Subtle allele frequency differences among some of the clusters likely caused problems with the mixing of the MCMC chains for k > 6 models. We highlight results from k = 2, 3 and 5. In the k = 2 model, South Hills crossbills were assigned to one cluster, whereas all other individuals were assigned to the other (Fig. 3A). The k = 3 model also assigned South Hills crossbills to a single cluster and assigned types 1, 3, 4 and 10 to a second cluster, and types 2, 5, 6 and 7 to a third (Fig. 3B). The k = 5 model assigned individuals to clusters that largely reflect the four nonoverlapping groups of ecotypes in the PCA (Fig. 2), with types 5, 6 and 9 each assigned to their own clusters, types 1, 3, 4 and 10 assigned to one cluster, and types 2 and 7 assigned to the fifth cluster (Fig. 3C).

We detected no evidence for isolation by distance. Pairwise genetic distances were unrelated to geographic distances (Fig. 4A), both for pairs of samples within the same ecotype and between all 22 geographically separate samples [the credible interval for the slope for the relationship between genetic and geographic distance included zero; slope for full analysis: 1.5 × 10^{-6}; 95% credible or equal-tail probability interval (ETPI): −1.8 × 10^{-5} to 4.2 × 10^{-5}]. In contrast to the lack of isolation by distance, ecotype beak depth tended to decrease with increasing PC2 values (Fig. 2), and these groupings of ecotypes were detected using ENTROPY (Fig. 3). In addition, genetic distance tended to increase with increasing beak depth divergence among ecotypes (Fig. 4B), but this pattern was not statistically significant (slope: 0.212; 95% ETPI: −0.008 to 0.439).

Although South Hills crossbills stand out in the phylogenetic and population genetic analyses (Figs 1–3), point estimates of genome-wide differentiation were not greater than in pairwise comparisons among all of the other ecotypes (Fig. 5A). Instead, the upper tails of the F_{ST} distributions for the South Hills crossbill had higher
densities (more loci with especially high $F_{ST}$) (Fig 5B), and these loci differentiated South Hills crossbills in PCA and Bayesian clustering analyses. For example, in pairwise comparisons between South Hills crossbills and other ecotypes, there was a strong relationship between locus-specific $F_{ST}$ and the strength of PC1 loading (Fig. 5C), a pattern that does not exist in comparisons among the other ecotypes (Fig. 5D). Similarly, the 0.8 quantiles of the genome-wide $F_{ST}$ distributions were higher for pairwise analyses involving South Hills crossbills (Fig. 5B). Thus, elevated divergence in a restricted number of genomic regions, rather than mean genome-wide genetic divergence, distinguished South Hills crossbills in PCA and ENTROPY analyses.

Estimates of nucleotide diversity indicate that South Hills crossbills harbour lower levels of genetic diversity.
than the other ecotypes (Fig. 7), which could reflect the fact that South Hills crossbills reside in only \(\sim 70 \text{ km}^2\) of lodgepole pine forest (Fig. 6) and likely have a much smaller effective population size than other ecotypes. Our historical reconstruction of lodgepole pine distribution in the region where South Hills crossbills occur suggests that there was little pine forest available only 6000 years ago during a period of warming (Fig. 6B, D). Unless the South Hills crossbill and pine began coevolving elsewhere and subsequently codispersed to the South Hills, these results suggest that coevolution and genome divergence occurred within the last 6000 years.

**Discussion**

Although levels of genetic differentiation were low, many ecotypes correspond to genetically cohesive groups that are differentiated from other such groups (Fig. 2). The low levels of genetic differentiation in our results and those of previous studies (Parchman et al. 2006; Bjorklund et al. 2013) are consistent with the ecotype diversification occurring recently, in the face of gene flow, or both. Glacial advances during the Pleistocene caused severe reductions in habitat that likely eroded ecotype diversity before glacial retreats allowed a vast expansion of conifers and an ensuing diversification of ecotypes (Benkman 1993; Dynesius & Jansson 2000). The absence of isolation by distance (Fig. 4A), consistent with ecotype nomadism and opportunistic breeding, indicates that genetic differentiation was not dependent on geographic distance or isolation. Instead, our results highlight the importance of adaptation to alternative conifer species (Benkman 1993, 2003) in contributing to reproductive isolation and genetic differentiation. These results contrast with the evidence that divergence without geographic isolation appears uncommon in birds (Price 2008). Although this difference might be attributable to the use of much smaller sets of genetic markers in past studies (but see Poelstra et al. 2014; Mason & Taylor 2015), strong reproductive isolation as a by-product of adaptation to alternative resources (Smith et al. 1999, 2012; Smith & Benkman 2007) distinguishes crossbills from most bird species (Price 2008).

Population genomic analyses indicate that the South Hills crossbill (Type 9) was the most genetically distinct ecotype (Figs 2 and 3A), and it was the only monophyletic ecotype in phylogenetic analyses (Fig. 1). This pattern of genetic differentiation indicates that previously documented patterns of divergent selection, adaptation (Benkman 1999; Benkman et al. 2003), and reproductive isolation (Smith & Benkman 2007; Benkman et al. 2009) associated with a local coevolutionary arms race have contributed to genome divergence. Our results show that, rather than overall genome-wide divergence, elevated genetic differentiation in a small number of genomic regions characterizes divergence in the South Hills crossbill (Fig. 5). This pattern is expected when adaptive divergence occurs in the face of gene flow (Peccoud et al. 2009; Feder et al. 2012), although such a pattern could also arise from selective sweeps in the absence of reproductive isolation (Cruickshank & Hahn 2014). Our findings are consistent with recent studies of other vertebrate taxa with phenotypic differentiation and reproductive isolation (Poelstra et al. 2014; Malinsky et al. 2015; Mason & Taylor 2015), where differentiation is restricted to few genomic regions across a background of genomic homogeneity. While GBS data offer a coarse assessment of patterns of ecotype genome divergence, whole genome resequencing

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**Fig. 4** (A) Genetic distances were unrelated to geographic distances between geographically separate samples within ecotypes (black symbols) and between all 22 samples, and (B) genetic distances between ecotypes did not increase significantly with increasing beak depth divergence among ecotypes (B includes only between-ecotype comparisons). Estimates of the effect of geographic distance or beak depth on genetic distance are for appropriately transformed variables (see 'Materials and methods'), rather than the untransformed values in the plots.
could eventually resolve the size and organization of genomic regions involved in divergence and speciation. Genome divergence in the South Hills crossbill could be influenced by geographic isolation, strength of reproductive isolation and effective population size. South Hills crossbills are not geographically isolated, as other nomadic ecotypes regularly move through and breed in the South Hills annually (Smith & Benkman 2007; Benkman et al. 2009). Moreover, the large scale over which we were unable to detect isolation by distance (Fig. 4A), suggests that 150 km of forestless area separating the South Hills from the vast forests to the north is unlikely to affect the opportunity for gene flow. Alternatively, reproductive isolation is potentially stronger in the South Hills than elsewhere, because strong density-dependent food limitation (Benkman et al. 2012) and cones with elevated defences against crossbills make it more difficult for nonlocally adapted ecotypes to persist and breed (Smith & Benkman 2007). Our test of isolation by ecology was not statistically significant (Fig. 4B), indicating that increasing divergent selection alone did not result in an increase in genetic differentiation. However, as implied above, our measure of divergent selection (beak depth divergence) does not capture certain

Fig. 5 Ecotypes 6 and 9, shown on the right in A and B, do not have particularly large pairwise $F_{ST}$ estimates relative to those including only the other ecotypes (A), but they do have more numerous locus-specific $F_{ST}$ estimates $>0.8$ (of 18 385 loci) (B) in comparison with pairwise estimates between the other ecotypes (ecotype 6 vs. all others: Wilcoxon pairwise test, $Z = 2.48$, $P = 0.013$; ecotype 9 vs. all others: $Z = 3.57$, $P = 0.0004$; ecotype 6 vs. 9: $Z = 0.16$, $P = 0.16$; similar patterns were found for $F_{ST} > 0.9$, but are not shown). Analyses of PC1 loadings in relation to locus-specific $F_{ST}$ estimates for pairs of ecotypes revealed much stronger relationships when ecotype 9 was included (C: ecotypes 2 and 9 shown; $P < 2.2 \times 10^{-16}$) than when it was excluded (D: ecotypes 2 and 10 shown; $P < 2.2 \times 10^{-16}$). Box plots (A, B) show minimum, first quartile, median, third quartile and maximum. Dashed lines (C, D) represent least-squares linear regressions.
elements of ecology such as strength of density dependence that likely affect reproductive isolation (Bolnick 2011). Furthermore, given the heterogeneity of differentiation across the genome (Nosil et al. 2009), average genome-wide divergence is a poor measure for such analyses when divergence occurs with gene flow. An additional factor that could contribute to the greater genetic distinctiveness of the South Hills crossbill is its small effective population size, as genetic drift could increase relative genetic differentiation ($F_{ST}$). Indeed, ecotype-level estimates of heterozygosity ($\pi$) are lowest for the South Hills crossbill (Fig. 7), consistent with a small effective population size [our current (October–November 2015) total population estimate is $N_e \approx 4000$ birds; N. Bohl and C. W. Benkman, unpublished data]. Small population size, in addition to geographic isolation, has likewise been suggested to contribute to the relatively elevated levels of genetic divergence of the crossbill endemic to the Aleppo pine forests ($P. halepensis$) on the island of Mallorca ($L. c. balearica$; Bjorklund et al. 2013).

Reconstructions of historical lodgepole pine distribution using classification models (Random Forests) based on BioClim data were consistent with palaeobotanical
studies (Mehringer 1985; Davis et al. 1986) indicating that pine forest was sparse during several thousand years of warming centred on 6000 yr BP in the region where South Hills crossbills currently reside (Fig. 6). Given the current South Hills crossbill population size, such a large (~86%) reduction in the amount of lodgepole pine would likely have prevented a distinct crossbill population from persisting in this region (Siepielski & Benkman 2005). This conclusion is supported by the evidence for exceptionally high temperatures 5000–7000 yr BP (Bartlein et al. 2014) conducive to lodgepole pine experiencing frequent catastrophic fires (Westerling et al. 2011), which would further reduce habitat for crossbills.

Thus, the South Hills crossbill either diverged as it coevolved with lodgepole pine in the South Hills during the last 6000 years, or it diverged elsewhere then subsequently colonized the expanded lodgepole pine forests of the South Hills. The latter is unlikely, as our reconstructions of historical lodgepole pine distributions for 22 000 yr BP (not shown) and palaeobotanical studies (Mehringer 1985) provide no indication that large forested areas occurred in this region at an earlier time. Furthermore, coevolving crossbills and pines were unlikely to have moved to the South Hills from elsewhere, because to the east and north red squirrels are widespread and lodgepole pine cones there reflect strong selection exerted by red squirrels suggesting a history of interaction (Benkman 1999; see Arbogast et al. 2001). To the west and south, lodgepole pine does not and has not occurred (Wells 1983). Finally, the high level of reproductive isolation in the South Hills crossbill is related to both the local resource characteristics that have evolved in response to the absence of red squirrels (Benkman & Siepielski 2004; Benkman et al. 2012) and crossbill-pine coevolution resulting in strong density dependence and local adaptation (Smith & Benkman 2007).

Conclusions

Coevolution has often been invoked to explain patterns of macroevolutionary diversification (Ehrlich & Raven 1964; Thompson 2005; Jablonski 2008), and some components of coevolution have been documented in numerous natural populations (Thompson 1994, 2005). However, clear examples of reciprocal selection and adaptation driving speciation (the link between coevolution as a micro- and macroevolutionary process) are largely lacking (Althoff et al. 2014; Hembry et al. 2014). Past studies on the South Hills crossbill have provided strong evidence for the role of coevolution in driving morphological divergence and reproductive isolation (Benkman 1999, 2003; Benkman et al. 2001, 2013; Smith & Benkman 2007). Our results indicate that the high contemporary measures of premating reproductive isolation (Smith & Benkman 2007; Benkman et al. 2009) reflect a longer term barrier to gene flow. Moreover, the nomadic behaviour of crossbills, their common sympatric occurrence and the absence of isolation by distance across all ecotypes (Fig. 4A) suggest that local coevolution rather than geographic isolation per se is responsible for the high levels of reproductive isolation for the South Hills crossbill. Model-based reconstructions of the past lodgepole pine distribution in the South Hills region (Fig. 6) and other lines of the evidence indicate that the ancestors of South Hills crossbills became resident and began coevolving with lodgepole pine more recently than 6000 yr BP. If the South Hills crossbill evolved so recently, it could represent one of the fastest examples of speciation in birds (Price 2008). Unfortunately, it is likely that this most genetically differentiated New World red crossbill lineage will go extinct within this century due to climate change and loss of suitable habitat (Santisteban et al. 2012; Benkman in press).

The evidence presented here for genetic differentiation associated with resource specialization in the absence of clear geographic isolation is rare, but similar to that seen in host races of insects (Mallet 2008; Pecquod et al. 2009; Nosil 2012; Powell et al. 2013). More generally, there is growing evidence that reproductive isolation can arise in part as a by-product of adaptation to alternative resources (i.e. ecological speciation; Schluter 2000; Nosil 2012). Crossbills might be unusual among birds, which diverge primarily in allopatry

Fig. 7 Ecotype 9 (South Hills crossbills) exhibits much lower genetic diversity (likelihood estimates of heterozygosity) than the other ecotypes. The dashed line represents the overall mean.
(Price 2008), but the mechanisms underlying their diversification could prove general considering the tremendous diversity of host-specific insects (Mallet 2008) and that coevolution is thought to be a major driver of diversification (Thompson 2005; Althoff et al. 2014; Hembry et al. 2014). In particular, geographic variation in the coevolutionary process has been documented as an important source of divergent selection for many interactions (Thompson 2005, 2009). In cases where such divergent selection generates reproductive isolation, the geographic mosaic of coevolution could contribute prominently to ecological speciation.

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T.L.P., C.A.B. and C.W.B. designed research; T.L.P. generated and assembled data; T.L.P., C.A.B. and V.S.C. analysed data; and T.L.P. and C.W.B. wrote paper with contributions from C.A.B. and V.S.C.

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**Data accessibility**

A detailed description of our genotyping by sequencing protocol, a matrix of genotype probabilities and. bam files from **bwa** assemblies are available at the Dryad digital repository (http://dx.doi.org/10.5061/dryad.65d97).

**Supporting information**

Additional supporting information may be found in the online version of this article.

**Appendix S1 Methods.**

Table S1 The number of individuals sampled of *Loxia curvirostra* ecotypes, including geographic locations of separate samples within each ecotype, as well as white-winged crossbills *Loxia leucoptera*.

Table S2 Pairwise estimates of Nei’s *D* (upper triangle) and *FST* (lower triangle) among *Loxia curvirostra* ecotypes (call types).

Table S3 Deviance information criterion (DIC) estimates for entropy models run for *k* = 2 through *k* = 9. Lower estimates of DIC reflect better model fit.

**Fig. S1** The map illustrates sampling localities for red crossbill (*Loxia curvirostra* complex). Individual points refer to geographically separate collection locations and correspond with Table S1. Circles of the same number and color represent different geographical samples from a given ecotype.

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