

# The evolution of ancestral and species-specific adaptations in snowfinches at the Qinghai–Tibet Plateau

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Species in a shared environment tend to evolve similar adaptations under the influence of their phylogenetic context. Using snowfinches, a monophyletic group of passerine birds (Passeridae), we study the relative roles of ancestral and species-specific adaptations to an extreme high-elevation environment, the Qinghai–Tibet Plateau. Our ancestral trait reconstruction shows that the ancestral snowfinch occupied high elevations and had a larger body mass than most nonsnowfinches in Passeridae. Subsequently, this phenotypic adaptation diversified in the descendant species. By comparing high-quality genomes from representatives of the three phylogenetic lineages, we find that about 95% of genes under positive selection in the descendant species are different from those in the ancestor. Consistently, the biological functions enriched for these species differ from those of their ancestor to various degrees (semantic similarity values ranging from 0.27 to 0.5), suggesting that the three descendant species have evolved divergently from the initial adaptation in their common ancestor. Using a functional assay to a highly selective gene, *DTL*, we demonstrate that the nonsynonymous substitutions in the ancestor and descendant species have improved the repair capacity of ultraviolet-induced DNA damage. The repair kinetics of the *DTL* gene shows a twofold to fourfold variation across the ancestor and the descendants. Collectively, this study reveals an exceptional case of adaptive evolution to high-elevation environments, an evolutionary process with an initial adaptation in the common ancestor followed by adaptive diversification of the descendant species.

comparative genomics | high-elevation adaptations | common ancestry | snowfinches | *DTL*

Organisms living at high elevations are exposed to cold temperatures, low levels of oxygen, and strong ultraviolet (UV) radiation. These strong stresses drive drastic phenotypic adaptations, such as the increase of hemoglobin–oxygen affinity and metabolic rates, larger body size, and an enhanced UV tolerance (1–4). Despite the similarity in phenotypic adaptations, recent studies have demonstrated diversified and species-specific adaptations to high elevations at the genomic level (5–15). As the acquisition of similar genetic adaptations in a group of species is shaped by their phylogenetic relationship (16), it is an intriguing question whether a group of species sharing a common ancestry evolve different genetic adaptations to the high elevations.

The snowfinch species complex provides a unique opportunity to study this adaptive diversification within the context of common ancestry, i.e., how the adaptive processes of the descendant species differ from that of their common ancestor. Snowfinch is

one of the few avian clades that have experienced an “in situ” radiation in extreme high-elevation environments, i.e., higher than 3,500 m above sea level (m a.s.l.) (17, 18). The Qinghai–Tibet Plateau (QTP), sometimes described as the “Third Pole” (19), houses six out of the total seven snowfinch species. Compared to most of their closest relatives among the Old World Sparrows (Passeridae), the snowfinches have a larger body size, darker plumage, and increased metabolic rate (20, 21). Previous studies suggested that snowfinches, consisting of the three genera of *Montifringilla*, *Onychostruthus*, and *Pyrgilauda*, form a monophyletic group (22–24). It is therefore possible that the observed similarities in phenotypic adaptations to high elevation are inherited from their common ancestor, assuming that the ancestral species had already adapted to this high-elevation environment. Furthermore, as the extant snowfinches differ somewhat

## Significance

Harsh conditions in high elevations present strong stresses for organisms. Previous studies targeting phylogenetically distinct species revealed cases of diversified adaptations, but it remains largely unknown how common ancestry contributes to evolution of similar adaptations. Our study based on a species complex (snowfinches) living on Qinghai–Tibet Plateau shows that ancestral snowfinches had phenotypically evolved larger body size and genetically an accelerated selection on genes related to development and signaling. From this ancestral state of adaptation three descendants have undergone independent adaptive processes in response to the differences in selective pressures acting on them. A striking example is a DNA repair gene, *DTL*, in which nonsynonymous substitutions evolving in ancestor and descendants have led to different DNA damage repair kinetics.

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The authors declare no competing interest.

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in their morphology, niche utilization, and behavior (20, 21), they may also have evolved the differences in the adaptive characteristics after they split from their common ancestor. Under this adaptive scenario it is likely that the genetic adaptation of the snowfinches to high-elevation environments is an evolutionary process of an initial adaptation in the ancestor and a subsequent adaptive diversification in the descendant species. To our knowledge, this topic has not been explored previously in relation to high-elevation adaptation.

In this study, we generate three de novo genomes from representatives of the three lineages of snowfinches. We integrate phenotypic, genomic, and functional assay data to investigate the impact of the conditions of the ancestor (ancestral adaptation) on the adaptive evolution in the descendant species (species-specific adaptation). Our results show that the snowfinch ancestor has evolved relatively larger body size and an accelerated selection on genes related to development and cellular signaling. From these ancestral adaptive conditions, the three species of snowfinches have evolved species-specific adaptive strategies to high-elevation environments. Using a functional assay on a highly selected gene, denticleless E3 ubiquitin protein ligase homolog (*DTL*), we demonstrate that multiple nonsynonymous substitutions in the ancestor and the descendant species have increased the repair capacity of UV-induced DNA damage. Altogether, our results show an evolutionary process of an initial adaptation in the ancestor followed by adaptive diversification in the descendant species, which have concurrently generated similar but not identical evolutionary routes to reach the high-elevation adaptation.

## Results

**Generation of High-Quality Genomes of the Three Representative Lineages of Snowfinches.** To explore how snowfinches adapt to high-elevation environments, we sequenced the whole genomes of the representatives of three currently recognized major lineages within the group (Fig. 1A), the white-rumped snowfinch, *Onychostruthus taczanowskii* (hereafter referred to as *taczan*), the rufous-necked snowfinch, *Pyrgilauda ruficollis* (*rufico*), and the black-winged snowfinch, *Montifringilla adamsi* (*adamsi*). For each species, we generated more than 110× sequencing data with four to five different insert sizes (Table 1 and *SI Appendix, SI Text 1* and Tables S1 and S2). We achieved high quality assemblies for all three species with scaffold N50 of ~10MB and contig N50 of ~0.5 MB (Table 1 and *SI Appendix, SI Text 1* and Table S3). Combining methods of homology-based modeling, de novo gene prediction, and RNA-sequencing-based assemblies, we annotated around 15,000 protein-coding genes for each of the three species, 99.8% of which have homologs in the public protein databases (Table 1 and *SI Appendix, SI Text 1* and Tables S4–S8). Using benchmarks against universal single-copy orthologs of birds (BUSCO, ref. 25), we estimated that the gene annotation is nearly complete (95%, *SI Appendix, SI Text 1* and Tables S9–S11).

**Snowfinches as a Monophyletic Group Emerged around 14 Ma.** Previous studies constructed the phylogeny of snowfinches based on a small number of genes (22–24). In this study, we corroborated the phylogenetic relationship of the three aforementioned phylogenetic lineages using whole genomic data. We selected seven phylogenetically representative oscine passerines as outgroups which have relatively well-annotated genomes available. We obtained a robust phylogenomic tree based on single-copy orthologous genes with 100% bootstrap values for all internal branches. This phylogenetic relationship is consistent with previously published results, supporting a monophyletic origin of three snowfinches (Fig. 1B). Given this robust phylogeny, we then estimated the time of divergence between the snowfinches and the other oscine passerines. Using the crown age of Passerida ( $26 \pm 3$  Ma, ref. 26) as a calibration point (Fig. 1B) we found that the ancestral snowfinch (*ancestor*) diverged from the most closely related lowland species, *Passer montanus* (Eurasian tree sparrow, *montan*) around 14 Ma

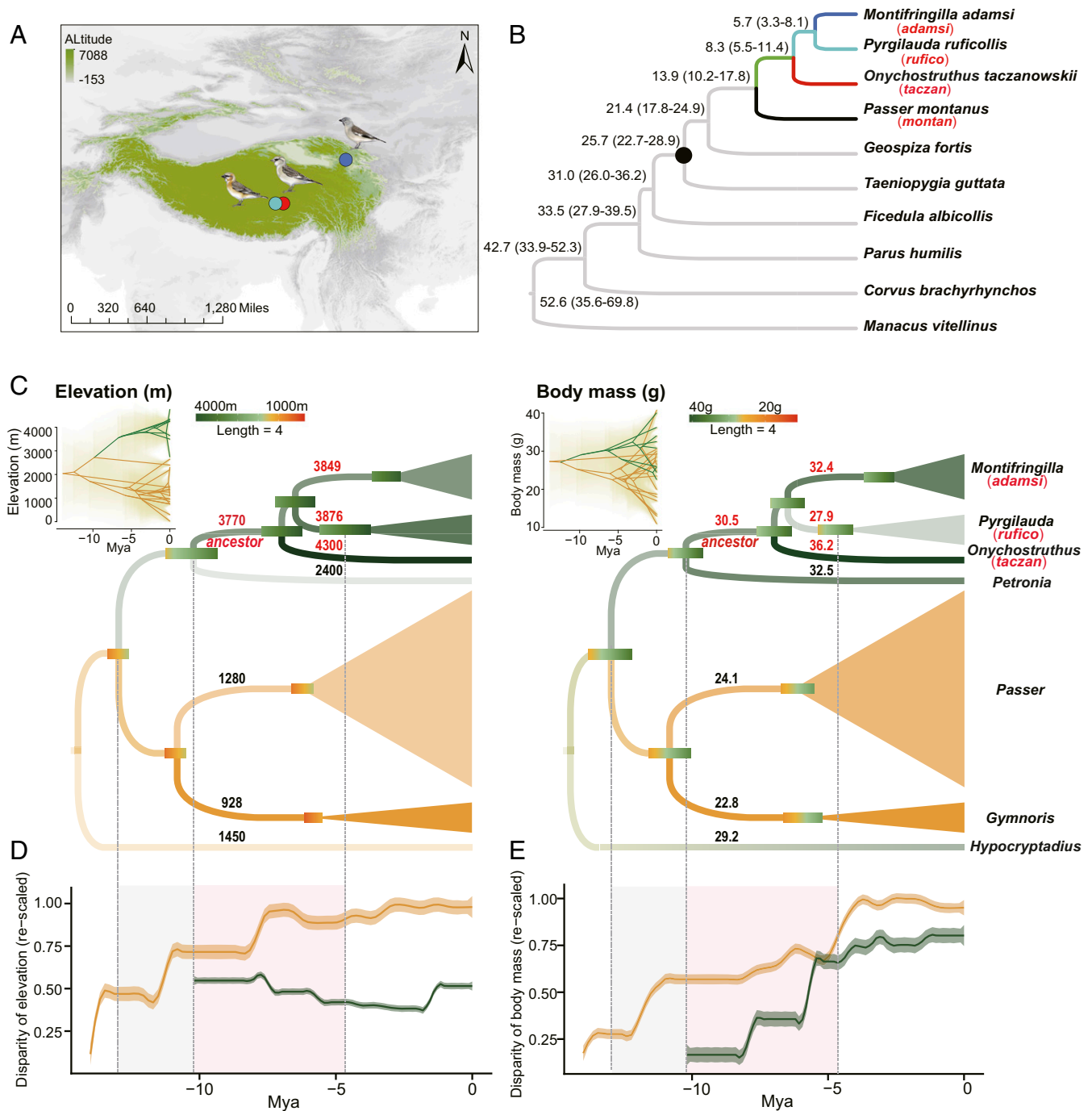
(95% confidence interval 10.3 to 17.8 Ma), and the three major lineages diverged from one another between 5.7 and 8.3 Ma.

**Ancestral Trait Reconstruction of Phenotypic Adaptation to a High-Elevation Environment.** We infer the ancestral elevation distribution of the snowfinches to determine whether the ancestral snowfinch had already inhabited high-elevation environments. To accurately infer the ancestral elevation, we collected mid-elevation records from the 27 extant species across seven genera of Passeridae and performed the ancestral trait reconstruction based on the species tree of Passeridae derived from Päckert et al. (24). Our results showed that the *ancestor* had likely already inhabited the QTP at an elevation of ~3,770 m a.s.l., which is within the elevational range of many extant species of snowfinches (Fig. 1C). Most nonsnowfinch species of Passeridae, however, are distributed at elevation below 2,000 m a.s.l., with the exception of *Petronia petronia* (rock sparrow) at 2,400 m a.s.l. Consistently, we also found a remarkable division in elevational range at the time when the snowfinches split from other Passeridae species and a subsequent decreasing variation in elevation during the diversification of the snowfinches (Fig. 1D).

We then checked if ancestral snowfinch had evolved phenotypic characters specific to high-elevation animals. In general, birds at high elevation evolve a relatively larger body size than their low-elevation relatives (27). We thus examined the variation of body mass across the snowfinch phylogeny. The results of the ancestral trait reconstruction showed that the body mass of the *ancestor* had increased to ~30 g, which is greater than that of most nonsnowfinch species of Passeridae but slightly smaller than that of the rock sparrow (Fig. 1C). During the diversification of snowfinches, both *adamsi* and *taczan* have increased their body mass whereas *rufico* has decreased its body mass compared to the *ancestor* (Fig. 1C). This result indicates an increased variation in body mass among snowfinches after their divergence from the common ancestor (Fig. 1E). Based on the time-scaled phylogeny, the results suggest that the *ancestor* began its adaptation to high-elevation environments around 14 Ma, and the descendant species have evolved divergent phenotypic adaptations during the speciation.

**Positive Selection in the Common Ancestor and Descendant Species.** As the descendant snowfinches show diversification of phenotypic adaptations following an initial adaptation in the *ancestor*, we expect to find a similar trend at the genetic level. Using the branch-site model in PAML (28) we identified positively selected genes (PSGs) in the *ancestor* (144 PSGs) and the descendant species (355 to 377 PSGs; Fig. 2A and *SI Appendix, Table S12*). We found that ~95% PSGs differ between the *ancestor* and the descendant species. However, we also found an excess of PSGs shared between the *ancestor* and the descendant snowfinches (one-sided binomial tests, false discovery rate [FDR]-adjusted  $P < 0.05$ ; Fig. 2A). For example, the ubiquitin-like modifier activating enzyme 6 (*UBA6*), a gene activating ubiquitin activity (29), shows accelerated selection across snowfinch branches, with the ratio of nonsynonymous substitution rate and synonymous substitution rate ( $d_N/d_S$ ) ranging from 0.23 to 0.44 as compared to ~0.1 in other branches. It is worth noting that the  $d_N/d_S$  values are higher in *adamsi* and *taczan* (~0.44) than in *rufico* (0.25; Fig. 2B), suggesting differences of selective strength among the descendant species. These results suggest that positive selection acting on the descendant species has driven divergent adaptations, demonstrated by the substantially different PSGs across species, although an appreciable number of genes (15 in *adamsi* to 24 in *taczan*; Fig. 2A) are under positive selection in both the ancestor and descendant species.

For the *ancestor*-only PSGs, which are genes showing an increase of  $d_N/d_S$  values only in the ancestral branch but not in the three descendant species, we hypothesize that the three descendant



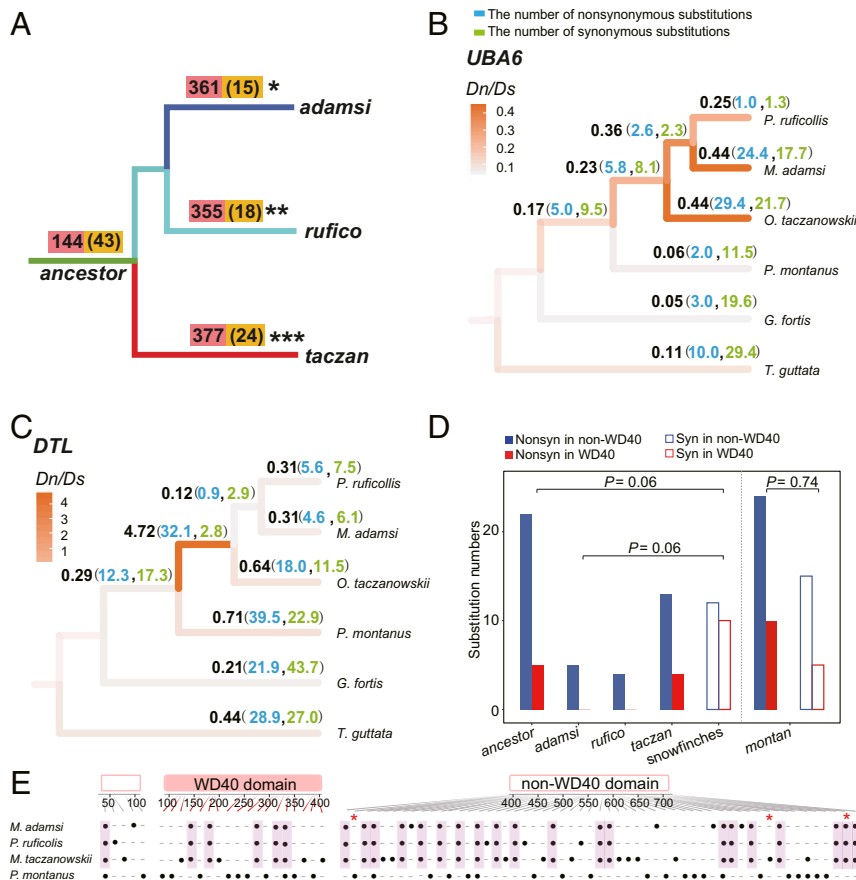
**Fig. 1.** The phylogenomic tree and ancestral trait reconstruction of the snowfinches. (A) Sampling sites of three species of snowfinches used in this study. For (A) and (B), three snowfinch lineages are similarly color-coded. (B) The phylogenomic tree suggests that the snowfinches diverged from *montanus* between 10 and 18 Ma. The black dot marks the calibration point ( $26.0 \pm 3.0$  Ma) in estimating divergence time. (C) Reconstruction of the ancestral traits for 27 species from Passeridae suggests that the ancestor of the snowfinches occupied a high-elevation environment (Left) and evolved a large body size (Right). The ancestral branch (ancestor), the representatives from three main lineages (*adamsi*, *rufico*, and *taczan*), and a lowland relative, the Eurasian tree sparrow (*montanus*), are marked along the tree. Twenty-seven species in Passeridae are collapsed into seven genera. The point estimates and 95% CIs of traits reconstructed for common ancestors of given genera are shown in numerical values and bars. Note only the point estimates are shown for the genera that have a single species (i.e., *Onychostruthus*, *Petronia*, and *Hypocryptadius*). The size of the triangle is proportional to the number of species. Phenograms in the inset show the trait evolution of the 27 species (y axis) over time (x axis) based on ancestral trait reconstruction with 95% CIs shown as shades. (D and E) Accumulation of elevation disparity (D) and body mass disparity (E) through time for all species of Passeridae (yellow curve lines) and snowfinches (green curve lines). The gray and orange shaded areas indicate time ranges when snowfinches split from nonsnowfinches and snowfinches diversified, respectively. Note that the divergence time of snowfinches in Päckert et al. (24) (C, D, and E) slightly differs from our phylogenomic tree in B.

**Table 1. Basic statistics for genome sequencing, assembly, and annotation of snowfinch species**

Species	Common name	Sequence data (G)	Insert size	Scaffold N50, Mb	Contig N50, Mb	Gene no.
<i>O. taczanowskii</i>	White-rumped snowfinch	152.6	170, 500, 800, 2,000, 5,000	9.07	0.67	15,585
<i>P. ruficollis</i>	Rufous-necked snowfinch	118.5	170, 500, 2,000, 5,000	9.60	0.51	15,206
<i>M. adamsi</i>	Black-winged snowfinch	139.7	170, 500, 2,000, 5,000	10.06	0.37	15,136

species may undergo a divergent adaptation by accumulating species-specific nonsynonymous substitutions. To test this hypothesis, we selected *DTL*, a gene related to DNA damage repair (30, 31), because *DTL* exhibits the strongest signal of positive selection in the ancestral branch ( $d_N/d_S$ , 4.72), higher than in any other branches ( $d_N/d_S$ , 0.12~0.71; Fig. 2C). To examine which region is particularly subject to positive selection, we analyzed the spatial position of nonsynonymous substitutions across different functional domains in *DTL* using synonymous substitutions as neutral control. We found that the WD40 domain of *DTL* has been marginally depleted with nonsynonymous substitutions in the ancestor (5 substitutions in WD40 domain vs. 22 in non-WD40 domain region) compared to synonymous substitutions (10 vs. 12, Fisher's exact test, FDR-adjusted  $P = 0.06$ ; Fig. 2D and E). This

pattern differs from the positions of nonsynonymous and synonymous substitutions in WD40 and non-WD40 regions in *montan* (10 vs. 24 as compared to 5 vs. 15, FDR-adjusted  $P = 0.74$ ; Fig. 2D and E). As synonymous substitutions approximate the neutral mutation input, non-WD40 region of ancestor *DTL* seems to be the major target of positive selection. After the three descendant snowfinches split they evolved species-specific nonsynonymous substitutions in the non-WD40 region (4 vs. 22, FDR-adjusted  $P = 0.06$ ; Fig. 2D and E). *DTL* of *taczan* has a relatively large number (4 vs. 13), whereas *adamsi* and *rufico* *DTL*s have small numbers (*rufico*, 0 vs. 4; *adamsi*, 0 vs. 5). Thus, even for a gene that is highly selected in the ancestor, the descendant species have evolved varying numbers of nonsynonymous substitutions that probably lead to different functional dynamics.



**Fig. 2. Positive selection analyses of snowfinches.** (A) Numbers of PSGs in ancestor, *taczan*, *rufico*, and *adamsi*. Numbers of all PSGs are marked in red, and those shared between ancestor and all three descendant snowfinches (list on the ancestral branch) as well as between ancestor and each of the descendant snowfinches (list on the respective snowfinch branches) are marked in yellow and in brackets. We calculated the observed ratios of shared PSGs between ancestor and *adamsi* (0.04, 15/361), ancestor and *rufico* (0.05, 18/355), or ancestor and *taczan* (0.06, 24/377). We then tested these ratios against the expected ratio (0.02, 144/6,572 genes) using a binomial test. \*\*\* $P < 0.001$ ; \*\* $P < 0.01$ ; \* $P < 0.05$ . (B and C) Evolutionary rates of two representative genes (B, *UBA6* and C, *DTL*).  $dN/dS$ ,  $dN$  and  $dS$  in each branch are shown by black, blue, and green, respectively. (D) Statistics of nonsynonymous and synonymous substitutions in the WD40 and non-WD40 regions of the ancestor and three descendant snowfinch *DTL*s. The reason that we merged all three descendant species is because each species consists of only a few synonymous substitutions. Nonsyn, nonsynonymous substitutions; syn, synonymous substitutions. (E) Distribution of the nonsynonymous substitutions of the snowfinch and *montan* *DTL*s in the WD40 and non-WD40 regions, which are marked by black dots. The nonsynonymous substitutions of ancestor *DTL*, i.e., those shared by three snowfinches, are highlighted by purple boxes. Asterisks show sites containing two nonsynonymous substitutions, which could not be drawn separately due to the space limit.



**Impact of DTL Nonsynonymous Substitutions in the Ancestral and Descendant Snowfinches on UV-Induced DNA Damage Repair.** Because *DTL* is involved in DNA damage repair (30, 31), we examined whether the nonsynonymous substitutions in snowfinches affect repair capability of UV-induced DNA damage. We first predicted the functional effects of nonsynonymous substitutions using Provean (32) and Sift (33). Compared to those in *montan*, 6% (Provean) and 21% (Sift) substitutions potentially cause functional change and only one substitution (P506L) is predicted by both programs to be functionally relevant (*SI Appendix, Table S13*). Because the two programs give different results, we hypothesized that a cumulative effect of snowfinch-specific nonsynonymous substitutions could lead to more radical functional change. To examine this, we chemically synthesized the full-length protein-coding sequences of *DTLs* for the three descendant snowfinches (*DTL<sub>taczan</sub>*, *DTL<sub>rufico</sub>*, and *DTL<sub>adamsi</sub>*) and their ancestor (*DTL<sub>ancestor</sub>*). We also synthesized *montan DTL* (*DTL<sub>montan</sub>*) as a control. We then cloned these *DTL* sequences into lentiviral vectors (Fig. 3 and *SI Appendix, Text 2*). We used our previously established embryo fibroblast cell line (GEF; Fig. 3) from a wild great tit (34) to test the repair function of *DTL*. The great tit is the closest relative to snowfinches with cell lines available. After knocking down the endogenous *DTL* expression in GEF (*SI Appendix, Fig. S1*), we overexpressed *DTL<sub>taczan</sub>*, *DTL<sub>rufico</sub>*, *DTL<sub>adamsi</sub>*, *DTL<sub>ancestor</sub>*, and *DTL<sub>montan</sub>* via the lentiviral vectors. We exposed all five cell lines to UV irradiation and subsequently examined the repair kinetics of *DTL* (proportion of repair to damage) by measuring two representative UV-induced DNA lesions, pyrimidine (6-4) pyrimidone photoproduct (6-4PP) and cyclobutane pyrimidine dimers (CPD) (35).

In a two-way repeated measures ANOVA the two assays showed that snowfinch *DTLs* had a significant improvement of repair effects (6-4PP,  $P < 0.001$ ; CPD,  $P < 0.001$ ) and repair kinetics through time (6-4PP,  $P < 0.001$ ; CPD,  $P < 0.001$ ) as compared to *montan DTL* (Fig. 3). Specifically, at the time point of 1.5 h we found that 6-4PP repair capacity was significantly higher in the ancestral and descendant snowfinch *DTLs* (*DTL<sub>taczan</sub>*, *DTL<sub>rufico</sub>*, *DTL<sub>adamsi</sub>*, and *DTL<sub>ancestor</sub>*) than in *DTL<sub>montan</sub>* (post hoc  $t$  test, FDR-adjusted  $P < 0.01$ ; Fig. 3). The descendant and ancestral snowfinch *DTLs* differ in their repair kinetics (FDR-adjusted  $P < 0.05$ ; Fig. 3 and *SI Appendix, Table S14*). *DTL<sub>adamsi</sub>* shows the greatest repair capability by repairing ~65% DNA damage, whereas *DTL<sub>rufico</sub>* exhibits the smallest repair capability and only repairs ~15% DNA damage. Both *DTL<sub>ancestor</sub>* and *DTL<sub>taczan</sub>* have an intermediate level of repair capability by repairing ~45% DNA damage (FDR-adjusted  $P < 0.05$ ). At the time point of 6 h all snowfinch *DTLs* had reached the repair levels of ~80%, which are significantly higher than that of *montan* (~60%, FDR-adjusted  $P < 0.05$ ). *DTL<sub>ancestor</sub>* and *DTL<sub>taczan</sub>* show a similar repair kinetics (FDR-adjusted  $P$  value, nonsignificant), which significantly differs from that of *DTL<sub>rufico</sub>* and *DTL<sub>adamsi</sub>* (FDR-adjusted  $P < 0.05$ ).

For the CPD assay, the three *DTLs* of the descendant snowfinches showed a similar repair kinetics and reached the peak of repair capability at the time point of 6 h (recovering ~90% DNA damage, FDR-adjusted  $P$  values, nonsignificant). Their repair kinetics was significantly and marginally higher than that of *DTL<sub>montan</sub>* (~57%, FDR-adjusted  $P < 0.05$  and  $P = 0.08$ ). In contrast, *DTL<sub>ancestor</sub>* only repairs about 40% damage, and its kinetics differs significantly from those of the descendant snowfinch *DTLs* (FDR-adjusted  $P < 0.05$ ) and that of *DTL<sub>montan</sub>* (FDR-adjusted  $P = 0.05$ ). At the time point of 24 h, the repair kinetics of all snowfinch *DTLs* is higher than that of *DTL<sub>montan</sub>* (~95% vs. ~74%), although the difference is not statistically significant after the multiple correction (Fig. 3 and *SI Appendix, Table S14*). Notably, *DTL<sub>ancestor</sub>* shows lower efficiency of CPD repair at 6 h but higher efficiency at 24 h when compared to *DTL<sub>montan</sub>*, and *DTL<sub>ancestor</sub>* also has a constant higher efficiency of

6-4PP repair compared to *DTL<sub>montan</sub>*. This pattern is likely a trade-off due to functional antagonism between 6-4PP and CPD repair or between early and late repair dynamics. Taken together, these results suggest that all snowfinch *DTLs* exhibit an improved repair capability of DNA damage, and the repair kinetics is different across the *ancestor* and the descendant species.

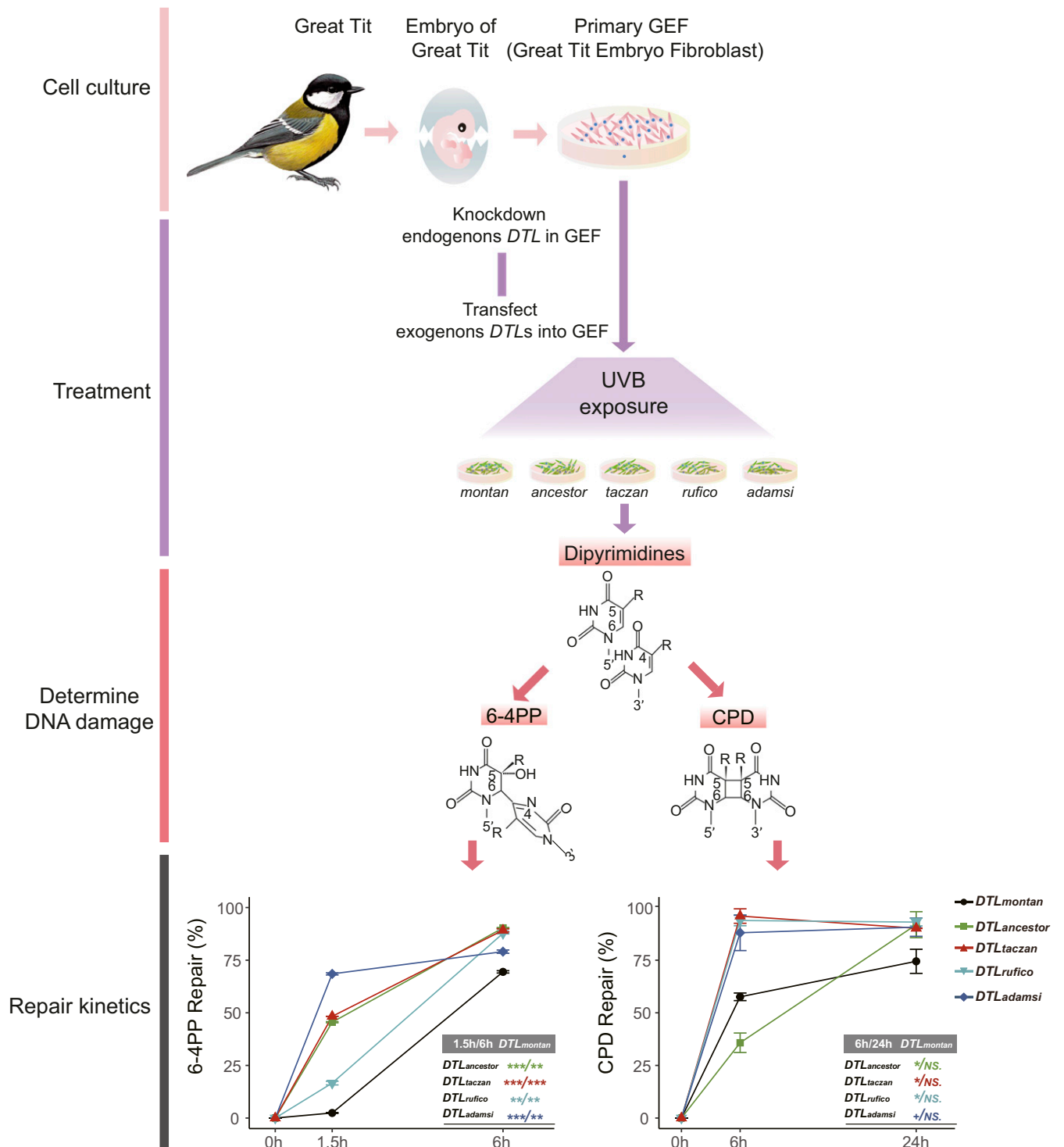
**Comparison of Enriched Biological Processes between Descendant Species and Their Ancestor.** We tested whether the adaptive divergence observed in PSGs and *DTL* function in the descendant species also occurs in biological processes represented by Gene Ontology (GO) terms. We utilized GO annotations to analyze the functional enrichments of PSGs in the *ancestor* and the descendant species. We found that PSGs are generally related to development, signaling, and cellular processes (Fig. 4A). To quantify the differences between *ancestor* and descendant snowfinches in functional enrichments we used GOSemSim (36) to compute pairwise semantic similarity (SS) of GO annotations. We implemented two measurements (RCMAX and BMA) to examine the robustness of the results. A greater SS value indicates more similarity between the two sets of GO terms (37, 38). Our results show that the pairwise SS values are the highest between the *ancestor* and *adamsi* (0.547 and 0.446), intermediate between the *ancestor* and *rufico* (0.46 and 0.394), and lowest between the *ancestor* and *taczan* (0.274 and 0.273; Fig. 4B). This result indicates that the three descendant species exhibit different levels of similarity of GO terms to the *ancestor*.

To test whether the observed SS values are statistically significant, we benchmarked the observed values against two permutations of the SS values calculated from 100 repeated samplings. We found that the observed SS values between *adamsi* and *ancestor* were higher than the 95th percentile value of the permutations (Fig. 4B), while the SS values between *rufico* and *ancestor* were higher than, or nearly equal to, the 90th percentile value. In contrast, the SS values between *taczan* and *ancestor* fall in the middle of the permutations, suggesting that they could be explained by random chance. We then generated a new permutation by sampling the same numbers of GO terms randomly. Using the new permutations, we obtained a similar trend (Fig. 4C). These results suggest that positive selection has driven different extents of adaptive divergence among the descendant species.

## Discussion

High-elevation environments exert a strong selective pressure on organisms living there and thus provide an ideal natural setting for the study of adaptation. Using snowfinches, a species complex living at the QTP, we investigated the contribution of ancestral adaptation to the adaptive evolution in the descendant species. Our results show that some key features for the high-elevation adaptation, including the developmental process, cellular signaling, and DNA repair capacity, have already evolved in the common ancestor. After the initial adaptation in the ancestor, the descendant species have adapted divergently in response to local selective pressures and microhabitats unique to each species (20–22, 39), leading to a deviation of adaptations between the ancestor and each of its descendants. Altogether, this study demonstrates a case of high-elevation adaptation, in which the descendant species have evolved diversified and species-specific adaptations after a common adaptation established in the ancestor.

The snowfinches have evolved a large body size compared to their lowland relatives. This phenotypic change likely began in their common ancestor. In addition, we identify strong signals of positive selection on genes related to development, signaling and DNA repair in the ancestor. We further demonstrate the ancestral adaptation using a functional assay on a highly selected gene, *DTL*, where nonsynonymous substitutions of ancestral snowfinch have improved the repair capacity of UV-induced DNA damage.

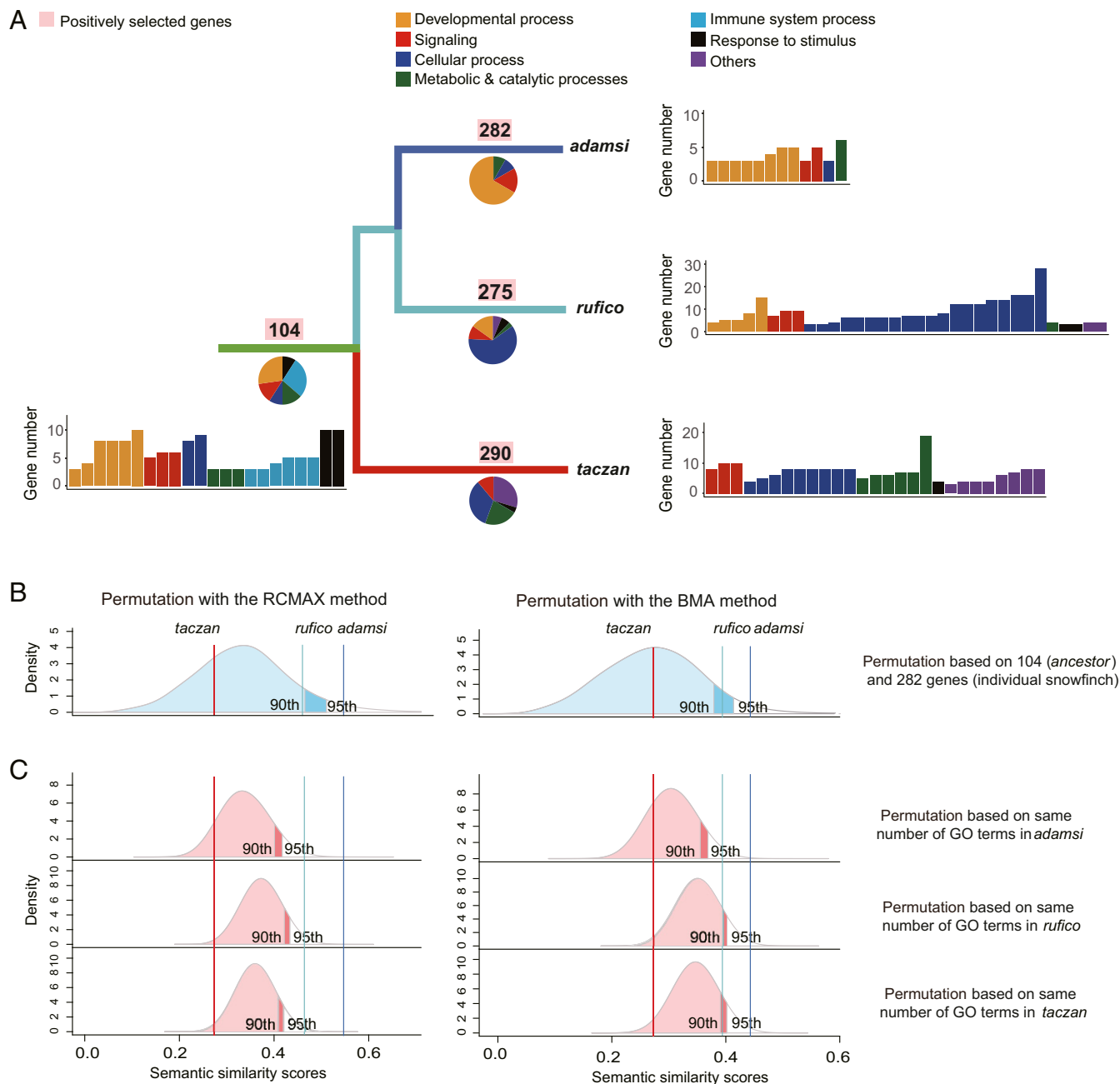


**Fig. 3.** A schematic representation of *DTL*'s functional experiment. The embryo fibroblast cells from the great tit (named GEF) were used. After knocking down endogenous *DTL* and transfecting exogenous *DTL<sub>taczan</sub>*, *DTL<sub>rufico</sub>*, *DTL<sub>adamsi</sub>*, *DTL<sub>ancestor</sub>*, and *DTL<sub>montan</sub>* (SI Appendix, Fig. S1), we exposed GEFs to UVB irradiation and checked repair kinetics (percent repair) by measuring two UV-induced photoproducts, 6-4PP and CPD. For each curve, the bar of SE is shown based on three biological replicates. Only post hoc *t* tests between each of snowfinch *DTL*s and *DTL<sub>montan</sub>* are shown. \*\*\**P* < 0.001; \*\**P* < 0.01; \**P* ≤ 0.05; +*P* = 0.08; NS, not statistically significant. Statistical results of pairwise comparisons of snowfinch *DTL*s are shown in SI Appendix, Table S14.

These adaptive characteristics are consistent with previous studies (5–14). Given the functional benefits of these characteristics, for example, a large body size can reduce heat loss with low surface-area-to-volume ratio (40), they are likely to be several common adaptive strategies for organisms to survive in a high-elevation

environment with low oxygen levels, low temperatures, and strong UV radiation.

The common ancestry may partially constrain genetic adaptations in the descendant species, evidenced by the finding that a significant excess of PSGs (4 to 6%) shared between each of the



**Fig. 4.** Functional enrichment analyses of PSGs. (A) Distribution of overrepresented GO terms is shown as a pie chart below the branch. Numbers of PSGs applicable for enrichment analyses are shown above branches. Note that the number is smaller than the counterpart in Fig. 2A because some genes do not have a human ortholog and thus could not be included in the analyses. GO terms are grouped into seven major categories including developmental process (yellow), signaling (red), cellular process (dark blue), metabolic and catalytic processes (green), immune system process (light blue), response to stimulus (black), and others (purple). (B and C) The observed SS scores and permuted ones between *ancestor* and the descendant snowfinches based on genes (B) and GO terms (C). Two algorithms of summarization including RCMAX (Left) and BMA (Right) are used. Dark blue, light blue, and red vertical lines show the observed pairwise SS values between *ancestor* and each of the snowfinches (*adamsi*, *rufico*, and *taczan*, respectively). Blue (permutations based on same number of genes) and red (permutations based on same number of GO terms) shades show 10,000 permuted SS values with 95th and 90th percentiles labeled.

descendant species and the common ancestor, and GO semantic similarity between ancestor and two descendants. This pattern could be explained by the evolution of a certain genetic makeup in the ancestor that had contributed to the acquirement of adaptive characteristics in the descendant species later on. This finding differs from what has been reported in previous genomic studies in the high-elevation animals, such as the Yak (*Bos grunniens*), Tibetan humans, ground tit (*Parus humilis*), and great

tit (*Parus major*), as genes under positive selection have been rarely found to be shared across those species (5, 7, 41, 42). It is worthwhile to investigate how the common ancestry influences the adaptive evolution in more animal systems in the future.

Besides the inheritance of adaptive characteristics for high elevation from the common ancestor, the descendant species exhibit various extents of adaptive divergence compared to that of their common ancestor. These results suggest that the descendant

species have evolved species-specific adaptive characteristics after they diversify. Since we find the GO terms enriched in *adamsi* and *rufico* show greater SS values than *taczan*, as compared to their common ancestor, this similarity seems evolved from their phylogenetic relatedness. However, we think this is less likely because of several lines of evidence. For example, the SS value between *adamsi* and *rufico* (0.28) is intermediate compared to the values between *adamsi* and *taczan* (0.20) and between *taczan* and *rufico* (0.34). Furthermore, *adamsi* and *rufico* do not exhibit more similarity when compared to *taczan* in terms of body mass (Fig. 1C) and *DTL* repair dynamics (Fig. 3). Moreover, our phylogenetic signal test does not find evidence for the influence of phylogenetic dependence on the evolution of the adaptive characteristics in the three descendant species (*SI Appendix*, Table S15). We therefore conclude that adaptive diversification of the three descendant species has evolved as a result of selection acting separately on each of them (20–22, 39).

## Materials and Methods

**Genome Sequencing, Assembly, and Annotation.** We collected samples of *taczan* and *rufico* from Naqu, Tibet (4,200 m a.s.l.) and of *adamsi* from Geermu, Qinghai (4,100 m a.s.l.; Fig. 1A). Our collection complies with the National Wildlife Conservation Law of China. Four to five paired-end short-read ( $2 \times 150$  bp) sequencing libraries with insert sizes of 170 bp, 500 bp, 800 bp, 2 kb, and 5 kb were constructed and sequenced on the Illumina HiSeq 2000 sequencing platform at BGI-Shenzhen. After the quality filtration, we assembled clean reads via SOAPdenovo (43) and SSPACE (44). We constructed gene sets by integrating de novo gene predictions, homology-based methods, and RNA-sequencing data-based methods. We annotated gene functions using Blastp based on their highest match to proteins in the UniProt database (release 2011-01), GO (45) and Kyoto Encyclopedia of Genes and Genomes database (release 58) (46). See *SI Appendix*, Text 1 for a detailed description of genome assembly and annotation.

**Phylogenomic Tree Reconstruction.** We reconstructed a phylogenomic tree including three snowfinches and seven other oscine species. Both phylogenetic representativeness and genome assembly quality were considered to select the outgroups. We blasted all the protein-coding sequences from these species using TreeFam (47) with an E-value threshold of  $10^{-7}$ . We used Solar to concatenate high-scoring pair segments of each protein pair and used H-scores to evaluate the similarity among genes with a custom script. We identified gene families by clustering homologous gene sequences using *Hcluster\_sg* (Version 0.5.0, <https://github.com/douglasgscfield/hcluster>). We identified the 784 best-to-best single-copy orthologous genes across all 10 species and used them for phylogenetic reconstruction. We calibrated the tree with the date of the clade Passerida (Fig. 1) (26),  $26.0 \pm 3.0$  Ma. This clade includes three species of snowfinches, *montan*, the medium ground finch, *Geospiza fortis*, and the zebra finch, *Taeniopygia guttata*. We then estimated the divergence time by implementing a Bayesian relaxed clock model in Beast v.2 (48). We ran Markov chain Monte Carlo chains for 100 million generations (sampling once every 1,000 generations) with a relaxed lognormal distribution for the molecular clock model and assuming a birth–death speciation process for the tree prior. The gamma substitution model was applied. We checked for convergence and performance using Tracer v.1.5 (49) and accepted the results if the values of the estimated sample size were larger than 200, suggesting little autocorrelation between samples. We combined resulting trees in TreeAnnotator v.1.7.5 and visualized the consensus tree with the divergence dates in FigTree v.1.4.3 (50).

**Ancestral Trait Reconstruction of Elevation and Body Mass.** We performed an ancestral trait reconstruction using a well-established phylogeny of Passeridae (24), which includes all 27 species from seven genera: *Montifringilla*, *Onychostruthus*, *Pyrgilauda*, *Petronia*, *Gymnoris*, *Passer*, and *Hypocryptadius*. We collected the elevation records from eBird (<https://ebird.org>) and China Bird Reports (<http://www.birdreport.cn>). Body masses were curated from the National Zoological Museum, Institute of Zoology, Chinese Academy of Sciences, and from the *Handbook of the Birds of the World* (51). We inferred ancestral traits for midelevation and body mass using BayesTraits v.3.0.2 (52). We used a variable-rates model to account for variation in evolutionary rates across phylogenetic trees. We ran 110 million iterations to ensure parameter convergence by discarding the first 10 million iterations in the burn-in phase. We sampled parameters every 1,000 iterations and final parameters were estimated based on all samples. We summarized the results by calculating a

mean rate-transformed tree based on all the trees scaled by the rate of change in the posterior samples. We estimated ancestral values for each trait using a maximum likelihood approach using the package Phytools (53). To account for unequal rates of trait evolution across the tree, we used the mean rate-transformed trees for each trait to estimate ancestral trait as described in Cooney et al. (54) and Venditti et al. (55). We used a 1-My time slice to estimate ancestral disparity through time using functional dispersion (FDISP) in package FD (56), which is a measurement of the average distance of all branches in the phylogeny to the centroid in a principal coordinates analysis (PCoA) of the given trait.

**Positive Selection Analysis of Protein-Coding Genes.** Since the previous species tree including ten species only covers 784 orthologous genes, we performed positive selection analysis with six closely related species including *taczan*, *rufico*, *adamsi*, *montan*, medium ground finch, and zebra finch. Using Blast (57) and TreeFam (47), we obtained 6,572 single-copy best-to-best hits as orthologous genes, which were shared by all six species and used for positive selection analysis (*SI Appendix*, Fig. S2). We aligned the orthologous proteins using Mafft v.7.310 (58), which were reversely translated as codon-level alignment using Pal2nal v.14 (59). We used branch-site likelihood ratio tests in Codeml of the PAML package (v.4.9e) (28) to identify PSGs for ancestral branch and the branches leading to *adamsi*, *rufico*, and *taczan*, respectively. The branch-site model allows  $\omega$  to vary both among sites in the protein and across branches on the tree and aims to detect positive selection affecting a few sites along particular lineages. Following <http://abacus.gene.ucl.ac.uk/software/pamlDOC.pdf>, we performed a branch-site model A test by comparing two models: the null model (using the settings model = 2, NSites = 2,  $\omega = 1$ ) and the alternative model (model = 2, NSites = 2). The likelihood ratio test has degrees of freedom = 1. We used the F3  $\times$  4 codon model of Goldman and Yang (60) to calculate the equilibrium codon frequencies from the average nucleotide frequencies at the three codon positions (CodonFreq = 2). We identified PSGs with multiple comparisons with Bonferroni–Holm correction at  $P < 0.05$ . We counted the shared PSGs between the ancestor and each of the descendant snowfinches and tested if the proportions of shared genes are significantly different from expected proportion (144/6,572) using a binomial test. FDR correction was used to control for multiple comparisons. For a few genes of interest, such as *UBA6* and *DTL*, we used free-ratio model in Codeml to calculate  $d_N/d_S$  value on every branch. We tested whether the spatial position of nonsynonymous substitutions as compared to synonymous substitutions between WD40 and non-WD40 domain regions differs between snowfinch *DTLs* and *montan DTL* used Fisher's exact test and FDR correction.

**Functional Assays for Repair Capacity of DNA Damage to UV Irradiation.** The coding sequences were synthesized for ancestral snowfinch *DTL* (*DTL<sub>ancestor</sub>*), descendant snowfinch *DTLs* (*DTL<sub>taczan</sub>*, *DTL<sub>rufico</sub>*, and *DTL<sub>adamsi</sub>*), and *montan DTL* (*DTL<sub>montan</sub>*). For each case, we cloned the *DTL* sequence into the pCDH-CMV-MCS-EF1-copGFP vector and packaged the vector into lentiviruses (Huaaobio). We then took the lentivirus to infect the great tit GEFs, whose endogenous *DTL* has been knocked down with short hairpin RNA targeting a conserved region (5'-GCACCAGCAAGCTCATCTTTA-3'; *SI Appendix*, Fig. S1). Applying a slot blot assay as described in Shah et al. (61), we treated GEF cells with UVB irradiation at 100 J/m<sup>2</sup> for 2 min and subsequently harvested cells at 0, 1.5, and 6 h afterward for 6-4PP quantitation or at 0, 6, and 24 h post-UVB irradiation for CPD determination. To determine DNA repair kinetics, we calculated the percentage of repair as 1 minus the ratio between densities of the bands at specified times and that at the 0 time point (100% damage after 2 min of UVB exposure). We used two-way repeated measures ANOVA to investigate if time points of repair and different *DTLs* significantly influenced repair capacity of DNA damage. For each factor, we conducted a post hoc *t* test comparison with FDR correction.

We inferred ancestral *DTL* sequences using phylogenetic model of coding sequence in a maximum likelihood framework in PAML. The inferred ancestral sequences show alternative sequences during the inference process. For example, a topology with tree sparrow as outgroup was used to infer the ancestral sequence. The inferred sequence with the high likelihood (*DTL<sub>ancestor\_1</sub>*) differs from the sequence inferred from alternative topology using the three species of snowfinches (at two sites, L566P and H586N, *DTL<sub>ancestor</sub>*). The ancestral sequence inference also identified two other sites with possible alternative variants (showing a probability of 0.3 to 0.4). Together the most likely (*DTL<sub>ancestor\_1</sub>*) and the least likely sequences (*DTL<sub>ancestor\_2</sub>*) differ at three sites (L566P, T472R, and Y605C; *SI Appendix*, Table S16).

We examined the functional difference among *DTL<sub>ancestor\_1</sub>*, *DTL<sub>ancestor\_2</sub>* sequences using Provean (32) and PolyPhen-2 (62). We



found that all these amino acid changes have negligible functional effects (SI Appendix, Table S16). We then carried out a functional experiment to test the repair capacities of the DNA damage of the three ancestral sequences using the 293T cell line. Both the CPD and 6-4PP functional assays show that the three ancestral sequences display similar repair capacities of DNA damage at all time points examined, except that  $DTL_{\text{ancestor}_2}$  (the least likely sequence) shows a significantly lower repair capacity at the 24-h time point of CPD (SI Appendix, Fig. S3). Taken together, our computational and functional assay analyses show that the functionality of the three ancestral  $DTL$  sequences is roughly similar and stable to alternative amino acid state during the inference process.

**Functional Enrichment Analyses and Semantic Similarity Permutation.** We performed functional enrichment analyses using Metascape (63), which has integrated the latest GO annotation. Specifically, we took advantage of the aforementioned 6,572 orthologs and compared snowfinch genes to zebra finch genes and then to their corresponding human orthologous based on Ensembl ortholog annotation (64). For 6,572 genes, 4,679 had orthologs in the human genome, which further reduced PSG numbers to 104 in *ancestor* and 276 to 290 in descendant species. We identified overrepresented biological process terms ( $P < 0.01$  and enrichment fold  $> 2$ ) in each group of these orthologs using 4,697 orthologs as a background gene set.

To evaluate the overall similarity of the functional enrichment patterns between the *ancestor* and each descendant species, we calculated pairwise SS values of the GO terms and examined SS values relative to random samples. We implemented the R package GoSemSim 3.10 (36) to calculate individual SS values. We applied two different algorithms, BMA and RCMA, to check if results are robust when summarizing SS values of individual GO terms. To determine if the observed SS values significantly deviated from a random expectation, we performed two different permutations. First, we randomly repeated 100 times by sampling genes from the same gene pools (104 for *ancestor* and 282 for descendant species, median value of 276 to 290 PSGs of the three descendant species) from the pool of all 4,697 genes and

performed analogous functional enrichment analysis for each random gene set followed by calculation of SS values. Second, we sampled 100 times of the same numbers of GO terms for *ancestor* and each species of snowfinches and calculate SS values. For both permutations, values greater than or equal to the 95th percentile values of the random samples were considered to be significant. Major conclusion is reproducible when more strict cutoffs ( $P < 0.005$  and enrichment fold  $> 2$ ) were used (SI Appendix, Fig. S4).

**Phylogenetic Signal Test of the Adaptive Characteristics.** We tested whether there are significant phylogenetic signals on the adaptive characteristics of the three descendant species using Blomberg's  $K$  (65) in Phytools (53), including the SS values, the numbers of the species-specific nonsynonymous substitutions of  $DTL$ , and the DNA damage repair kinetics.

**Data Availability.** Sequencing data for the three species of the snowfinches have been deposited in the Sequence Read Archive under project number PRJNA417520.

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