A Genomewide Scan for Genetic Structure and Demographic History of Two Closely Related Species, *Rhododendron dauricum* and *R. mucronulatum* (Rhododendron, Ericaceae)

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Understanding the processes of divergence and speciation is an important task for evolutionary research, and climate oscillations play a pivotal role. We estimated the genetic structure and demographic history of two closely related species of *Rhododendron*, *R. dauricum*, and *R. mucronulatum*, distributed in northeastern China using 664,406 single nucleotide polymorphic loci of specific-locus amplified fragment sequencing (SLAF-seq) and 4 chloroplast DNA (cpDNA) fragments, sampling 376 individuals from 39 populations of these two species across their geographic distributions. The geographical distribution of cpDNA haplotypes revealed that *R. dauricum* and *R. mucronulatum* have different spatial genetic structures and haplotype diversity. Analysis of molecular variance (AMOVA) results showed that these two species have significant genetic differentiation and that the phylogeny demonstrates that these two species clustered a monophyletic group based on SLAF data, respectively, but not in cpDNA data. The evidence of significant gene flow was also detected from *R. mucronulatum* to *R. dauricum*. A deep divergence between the two species was observed and occurred during the early Oligocene. The niche models showed that the two species have different demographic histories. Thus, our results imply that geography and climate changes played important roles in the evolutionary process of *R. dauricum* and *R. mucronulatum*, and although there was an interspecific gene flow, the divergence was maintained by natural selection.

**Keywords:** specific-locus amplified fragment, nucleotide differentiation, gene flow, niche simulation, climate changes
INTRODUCTION

Past climate oscillations and historical tectonism had a huge impact on the genetic structure and demographic histories of many species, even triggering divergence and speciation (Milne, 2006). Understanding the factors that promote species divergence is of major interest in ecological and evolutionary research (Papadopulos et al., 2011; Butlin et al., 2012). Species divergences are frequently driven by geographic isolation, environmental heterogeneity (ecological speciation) or a combination of both (Orr and Smith, 1998; Schluter, 2001; Nosil, 2012). Geographic isolation is generally considered an allopatric speciation, where gene flow among splitting populations is disrupted by physical barriers, and genetic divergence occurs between taxa by local adaptation, mutation, and genetic drift (Coyne and Orr, 2004). In contrast, under ecological speciation, divergence is driven by divergent natural selection between environments, giving rise to reproductive isolation between subsets of a single population by adaptation to different environments or ecological niches (Schluter, 2000; Schluter, 2001; Rundle and Nosil, 2005).

Gene flow played an important role in the evolution of species, having both positive and negative effects on adaptation (Palme et al., 2004). It is expected due to secondary contact when physical barriers disappear (under allopatric speciation) or range expansions and contractions in response to climatic fluctuation, which is generally regarded as a force counteracting population divergence (Runemark et al., 2012). When the rate of gene flow between species exceeds that among populations within the species, hybridization might occur. These phenomena have been demonstrated not only in animals (Taylor et al., 2005; Seehausen, 2006; Webb et al., 2011), but also in plants (Marczewski et al., 2015; Ma et al., 2016). Interspecies gene flow after speciation can reduce genetic differentiation among species and even achieve homogenization (Vonlanthen et al., 2012; Bhat et al., 2014). However, when the homogenization of interspecific gene flow is weaker than the disproportionation of natural selection, species genetic differentiation may still be maintained (Rundell and Prince, 2009; Ribera et al., 2011).

Rhododendron (Ericaceae L.) is a taxonomically complex genus with approximately 1,000 species (Chamberlain et al., 1996) distributed in Himalayan flora. It is one of the major genera of China and includes approximately 571 species, of which 402 species are endemic to China (Wu et al., 2003; Fang et al., 2005). Northeast Asia is an area with complex topographic and climatic gradients (Lopez-Pujol et al., 2011), and recent research supports a northeastern Asian origin of Rhododendron (Shrestha et al., 2018). Among Rhododendron species, Rhododendron subgen. Rhodorastrum, including four taxa, R. ledebourii, R. dauricum, R. mucronulatum, and R. sichotense, are distributed in this area. The four species exhibit similar phenotypic characteristics, such as leaf, petiole and flower shapes, thus making taxonomy difficult. R. ledebourii, R. mucronulatum, and R. sichotense were regarded as populations, varieties or subspecies of R. dauricum in previous studies (Voroshilov, 1982; Mazurenko and Hohryakov, 1991; Koropachinskii and Vstovskaya, 2002). In contrast, phylogenetic analysis supports the species status of these four species-based cpDNA regions (Tikhonova et al., 2012; Polezhaeva et al., 2018). Additionally, two well-distinguished groups were determined for these four species, including R. ledebourii and R. dauricum as the Siberian group and R. sichotense and R. mucronulatum as the Far Eastern group (Polezhaeva et al., 2018), demonstrating good agreement with the geographical positions of the species collected. However, a main haplotype of R. mucronulatum was clustered with the haplotypes of R. sichotense, and the two species were joined together based on SAMOVA, Furthermore, samples were only from Russia and located in Northeast China (NEC) were lacking. In addition, phylogenetic reconstruction of the genus Rhododendron based on the ITS region showed that R. dauricum and R. mucronulatum were sister species (Kutsev and Karakulov, 2010; Baranova et al., 2014).

These four species have different geographical distribution, covering Mongolia, Northeast China, Japan, Korean Peninsula, and Russian (including the southern part of western Siberia, eastern Siberia, and the Russian Far East), and Northeast China (NEC) is in the center of these areas (Figure 1C), among which R. dauricum is widely distributed. R. dauricum and R. mucronulatum have different geographical distribution in NEC, although they both occur along mountain slopes. R. dauricum is distributed along the Great Khingan Mountains (GKM), Lesser Khingan Mountains (LKM) and Changbai Mountain (CBM), while R. mucronulatum is found in the south of the Changbai Mountain and other locations in North China.

Therefore, in this study, to test the genetic divergence and demographic history of R. dauricum and R. mucronulatum driven by possible roles of geological or climate events, we first combined specific-locus amplified fragment sequencing (SLAF-seq) and cpDNA markers. We assessed the genetic diversity, phylogeny, and genetic structures of the two species. The divergence time and ecological niche modeling were also tested to infer the possible population demographic history in light of past climate changes. In addition, gene flow between the species was also detected to further elucidate the role of either climate oscillations or geological processes in shaping nucleotide variation and triggering divergence. Together, these analyses not only provide insights into the evolutionary history of two closely related species but also contribute to understanding biodiversity in northern and northeastern China.

MATERIALS AND METHODS

Plant Materials and Sequencing

On the basis of 23 populations in Jiang et al. (2016), we have newly collected some populations and individuals of R. dauricum. After removing and merging some populations that were close to each other or for which the haplotypes were completely consistent, we newly added 127 R. dauricum individuals in this study. Of these, 88 individuals were from newly added 9 populations and 39 were extensions to the existing populations. Finally, there were a total of 247 R. dauricum...
individuals in 27 populations (Table 1). Moreover, we newly collected 12 populations of 129 individuals with *R. mucronulatum* across all the current relevant regions in China. 5 to 23 individuals for each population, only one individual for LB. We distinguished these two species mainly based on geographical distribution and morphological characteristics. Firstly, *R. dauricum* is mainly distributed in Inner Mongolia, Heilongjiang and Jilin Provinces of China, and *R. mucronulatum* is mainly distributed in Liaoning Province, Hebei Province, Beijing, Shandong Province, and northern Jiangsu Province. Secondly, the main differences in morphology between the two are the characteristic of scales on the leaves abaxial surface and whether there are hairs on the branchlets. The leaves abaxial surface of *R. dauricum* densely covered with scales, imbricate or adjacent to each other, spacing 1/2 or 1.5 times their diameter, and branchlets pilose; the spacing of the scales in *R. mucronulatum* was two to four times its diameter and branchlets glabrous. Sampled individuals were located at least 50 m apart. Additionally, samples of *R. micranthum* and *R. schlippenbachii* were also collected to serve as outgroups for cpDNA phylogenetic analyses. All fresh leaf materials were desiccated in silica gel, and voucher specimens were stored in the Northeast Normal University Herbarium (NENU).

Total genomic DNA was extracted using a modified 4 × CTAB procedure (Doyle and Doyle, 1987). The concentration and quality of genomic DNA were tested by agarose gel electrophoresis (1%) and an ND-1000 spectrophotometer (Thermo Fisher Scientific, USA). Qualified DNA samples were stored at −20°C for polymerase chain reaction (PCR) amplification and high-throughput sequencing.

To obtain the orthologous regions of *R. dauricum* individuals from *R. mucronulatum* samples, we selected four pairs of primers (trnS-trnG, trnS-trnfM, TabE-ndhJ, trnK-matK) used in a previous study (Jiang et al., 2016), which were most informative for *R. dauricum* populations. The reagent concentrations, polymerase chain reaction (PCR) process, and sequencing followed Jiang et al. (2016).

Analyses of cpDNA Sequences

DNA sequence alignment was performed using ClustalX 2.0 (Larkin et al., 2007), and alignments were edited manually in BioEdit 7.0.1 (Hall, 1999) before being concatenated. The number of polymorphic sites (S), nucleotide diversity (π), number of haplotypes (h), and haplotype diversity (Hd) were calculated by DnaSP v5 (Librado and Rozas, 2009). We also calculated Tajimas’ D (Tajima, 1989), Fu and Li’s D and F (Fu and Li, 1993) separately for the two species using this software to test how well the data conformed to the neutral model of evolution. The population differentiation statistic (FST) (Wright, 1984) was calculated by AMOVA in ARLEQUIN 3.5 (Excoffier et al., 2005). Furthermore, the cpDNA haplotype, representing unique DNA sequences or alleles separated by mutational steps, was constructed by TCS v1.2.1 (Clement et al., 2000) with genetic distance and statistical parsimony methods. We chose ‘Gaps = missing’ and set the Fix Connection Limit to 50 steps to ensure that all haplotypes were connected to each other. In order to compare with the phylogenetic tree constructed by SLAF data, we chose the same individuals as SLAF to construct the phylogenetic tree based on cpDNA. A maximum likelihood (ML) tree was constructed in RAxML 8.2.9 software (Stamatakis et al., 2008) under GTRGAMMA selected by

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**FIGURE 1 |** (A) The Network of haplotypes in *R. dauricum*. The size of haplotype circle represents relative frequency. (B) Geographical distribution of haplotypes within *R. dauricum* and *R. mucronulatum*. The regions of GKM, LKM, and CBM were indicated by dotted ellipses. The populations of *R. dauricum* and *R. mucronulatum* were distinguished by solid line. (C) The distribution of four species in subgen. *Rhodorastrum*. It was drawn based on Polezhaeva (2018) and the sites information collected in ecological niche model simulation. Each color represents a species.
jModeltest 2.0 (Darriba et al., 2012) based on Akaike information criterion (AIC) values and bootstrap supports for clades calculated using 1000 bootstrap replicates.

**SLAF Sequencing, Read Alignment and SNP Calling**

To obtain all haplotypes of cpDNA and represent the distribution of the two species as evenly as possible, we selected 65 individuals (including 40 *R. dauricum* samples and 25 *R. mucronulatum* samples) for subsequent SLAF sequencing (Table 1). Samples of *R. redowskianum* and *R. delavayi* were used as outgroup. The genome of *R. delavayi* (ftp://parrot.genomics.cn/gigadb/pub/10.5524/100001_101000/100331/) was selected as the reference genome for enzyme cutting site prediction. After strict filtering, *Hae*III and *Hpy*166II restriction enzymes were finally used to digest the qualified DNA of the sampled individuals according to the results of prerestriction enzyme digestion. To evaluate the accuracy of the enzyme digestion experiment, *Oryza sativa* subsp. *indica* was

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### TABLE 1 | Sampling information and genetic diversity for SLAF data and cpDNA of *R. dauricum* and *R. mucronulatum*.

| pop | Location | Long (°E) | Lat (°N) | size | S | h | Hd | $\pi \times 10^{-3}$ | voucher |
|-----|----------|-----------|----------|------|---|---|----|-----------------|---------|
| **R. dauricum** | | | | | | | | |
| GKM | | | | | | | | |
| AES* | Aeraha, IM | 120.562 | 47.410 | 20 (2) | 0 | 1 | 0.6610 | 1.8000 (0.2450) | NENU00042001 |
| TP | Taipingling, IM | 120.491 | 47.488 | 10 (1) | 0 | 1 | 0.0000 | 0.0000 | NENU00042002 |
| ZL | Zhilant, IM | 122.700 | 48.063 | 10 | 0 | 1 | 0.0000 | 0.0000 | NENU00042003 |
| BK* | Beikitu, IM | 122.105 | 48.822 | 10 (3) | 0 | 1 | 0.0000 | 0.0000 | NENU00042004 |
| GH | Genhe City, IM | 121.492 | 50.775 | 5 (1) | 0 | 1 | 0.0000 | 0.0000 | NENU00042005 |
| JH | Jinhe Town, IM | 122.34 | 51.041 | 5 | 0 | 1 | 0.0000 | 0.0000 | NENU00042006 |
| AK | Oakley Hill, IM | 122.037 | 48.832 | 10 | 0 | 1 | 0.0000 | 0.0000 | NENU00042007 |
| AL | Ali River, IM | 123.533 | 50.550 | 5 | 0 | 1 | 0.0000 | 0.0000 | NENU00042008 |
| TH* | Tahe County, HLJ | 124.710 | 52.334 | 10 (3) | 1 | 2 | 0.2000 | 0.0001 | NENU00042009 |
| | * | | | | | | | |
| **R. mucronulatum** | | | | | | | | |
| LTD* | Lao Baldingzi, LN | 124.908 | 41.298 | 21 (2) | 0 | 1 | 0.0000 | 0.0000 | NENU00042010 |
| LS* | Sitaizi Forest Farm, JL | 124.702 | 43.189 | 10 | 0 | 1 | 0.0000 | 0.0000 | NENU00042011 |
| FH* | Phoenix Mountain, LN | 120.491 | 41.298 | 21 | 0 | 2 | 0.3250 | 0.0000 | NENU00042012 |
| LD* | Changbai Mountain, JL | 126.335 | 42.856 | 8 (3) | 2 | 3 | 0.4643 | 0.0002 | NENU00042013 |
| LS* | Sitaizi Forest Farm, JL | 124.702 | 43.189 | 10 | 0 | 1 | 0.0000 | 0.0000 | NENU00042014 |
| HP* | Weihuling, HLJ | 127.866 | 43.483 | 10 | 2 | 2 | 0.3560 | 0.0003 | NENU00042015 |
| LT* | Camel Mountain, JL | 129.540 | 43.645 | 10 (2) | 1 | 2 | 0.5660 | 0.0002 | NENU00042016 |
| HS* | Heihe, HLJ | 130.055 | 43.189 | 10 | 2 | 3 | 0.8890 | 0.0003 | NENU00042017 |
| JS* | Triangular Bay, JL | 127.943 | 41.547 | 23 (3) | 0 | 1 | 0.0000 | 0.0000 | NENU00042018 |
| LB* | Old white mountain, JL | 127.919 | 42.424 | 1 | – | – | – | – | NENU00042019 |
| LL* | Tahe County, HLJ | 124.449 | 45.259 | 5 (1) | 0 | 1 | 0.0000 | 0.0000 | NENU00042020 |
| HB* | Heihe, HLJ | 127.452 | 44.247 | 12 (3) | 2 | 3 | 0.4850 | 0.0004 | NENU00042021 |
| S | number of polymorphic sites (SNPs); h, number of haplotypes; Hd, haplotype diversity; $\pi$, nucleotide diversity. *represents the newly added populations of *R. dauricum* and *R. mucronulatum* in this study, +represents the populations of individual expansion based on previous studies (the number of newly added: AES, 10; MLG, 8; CB, 9; LJ, 12). The size in brackets represents the individuals used for SLAF; the $\pi$ in brackets represents the nucleotide diversity based on SLAF. GKM, Great Khingan Mountains; LKM, Lesser Khingan Mountains; CBM, Changbai Mountain; IM, Inner Mongolia; HLJ, Heilongjiang Province; JL, Jilin Province; LN, Liaoning Province; BJ, Beijing; HB, Hebei Province; SD, Shandong Province.}

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used as a control to ensure the effectiveness of the enzyme-cutting scheme. The obtained SLAF tags with an A-tail added to the 3’ ends were ligated with Dual-index sequencing adaptors, amplified by PCR, screened, and then used to construct the SLAF library (for detailed processes of the SLAF library construction, refer to the methods of Sun et al. (2016). The selected tags were then subjected to pair-end sequencing using an Illumina HiSeq 2500 at Biomarker Technologies Corporation in Beijing. Sequencing insert size was 314 to 414 bp and paired-end reads were 126 bp in length.

Clear reads were aligned against the reference genome using Burrows-Wheeler Aligner (BWA) (Li and Durbin, 2009) with parameters defined as missed match = 3; opening gap = 11 and gap extension = 4. False alignment was always detected near Indels; therefore, local alignment was performed. The Genome Analysis Toolkit (GATK) (DePristo et al., 2011) and Sequence Alignment/Map tools (SAMtools) (Li et al., 2009) were used for variant calling under the default setting. Raw SNPs were filtered with a minimum depth (DP) of 3 and minimum mapping quality (MQ) of 30 by our custom Perl scripts, and then PLINK 2 (Purcell et al., 2007) was used to further filter with the minor allele frequency (MAF) of 0.04 and maximum missing rate of 0.5, and SNPs that were out of Hardy-Weinberg equilibrium (HWE) were removed with a p-value threshold < 0.01. Later, considering the linkage disequilibrium (LD) between SNPs, we used the -r2 function in PLINK to quantify pairwise LD between all pairs of SNPs located within 1000 kb of each other, and the results showed that the LD of both species decayed rapidly (Supplementary Figure S1). Thus, we did not further filter SNPs according to LD. All non-biallelic SNPs, which may be caused by sequencing, were filtered using our custom Perl scripts. The final screened SNPs were subjected to subsequent data analyses.

Genetic Diversity, Phylogeny, and Population Structure Analyses

The genetic diversity parameter π and the neutral test of Tajima’s D, Fu and Li’s D and F values were calculated by PopGenome in R (Bastian et al., 2014) based on SNPs. Fs was also calculated by AMOVA in ARLEQUIN 3.5. The ML tree was constructed in the same way as above. The population structure analyses were performed using ADMIXTURE software (Alexander et al., 2009). The number of subgroups (K value) was set as 1 to 10 in advance for clustering, and the clustering results were cross-verified to determine the optimal number of subgroups according to the valley value of the cross-validation error rate. We further performed principal component analysis (PCA) to determine the clustering status of samples using EIGENSOFT (Price et al., 2006).

Divergence Time and Gene Flow

A Bayesian relaxed molecular clock approach was used to estimate species divergence time by MCMCTREE in PAML (Yang, 2007). When using previously published calibration times, the split of Oryza sativa (https://phytozone.jgi.doe.gov/pz/portal.html#info?alias=Org_Osativa) and Arabidopsis thaliana (https://phytozone.jgi.doe.gov/pz/portal.html#info?alias=Org_Athaliana_er) was fixed as 130–200 Mya, and the split of R. delavayi and Actinidia chinensis (http://bioinfo.bti.cornell.edu/cgi-bin/kiwi/download.cgi) was estimated to be in the range of 56.1–120.8 Mya (Zhang et al., 2017). The 150 bp sequences before and after the SNPs were used for BLASTN (Altschul et al., 1990) against the reference genomes with an E-value threshold of 1e-3, and then obtained the sequence with a total length of 12,936,348 bp as an input file. R. dauricum and R. mucronulatum individuals were selected from populations TP and MS, respectively, based on the cpDNA haplotype. The data set was modeled under a correlated rates clock, and the HKY85 nucleotide substitution model was determined by ModelFinder (Kalyaanamoorthy et al., 2017). The first 2000 iterations were discarded as burn-in, and every 10 iterations were sampled until 20000 samples were obtained. Furthermore, to account for gene flow among the groups, we used TREEMIX 1.12 (Pickrell and Pritchard, 2012), which infers patterns of population splitting and mixing accessing the covariance structure of allele frequencies between populations and performing a Gaussian approximation for genetic drift.

Ecological Niche Modeling

MaxEnt software (Phillips et al., 2006) was used to predict the distribution models for R. dauricum and R. mucronulatum in the last interglacial (LIG) (0.14–0.12 Mya), the last maximum glacial (LGM) (0.02–0.018 Mya) and current time based on modern distribution records and bioclimatic variables. The occurrence data of species were mainly collected from the following three sources: Chinese Virtual Herbarium (CVH) (http://www.cvh.ac.cn/) and relevant literature; Global Biodiversity Information Facility (GBIF) (http://www.gbif.org); and sampling point information obtained from our field survey. 316 sites of R. dauricum and 214 sites of R. mucronulatum worldwide were obtained. Then, the distribution data obtained in these three ways were manually checked and proofread, and duplicate records within each 2.5-arc-minute cell and artificially planted species distribution points were removed. Finally, 238sites of R. dauricum and 214 sites of R. mucronulatum worldwide were obtained.

The 19 bioclimatic environmental variables were downloaded from the world climate data website (http://www.worldclim.org) with a 30 arc-second resolution for the present and LIG and 2.5 arc-minute for LGM. Then, ArcGIS software (Johnston et al., 2001) was used to perform pairwise correlations among the 19 variables. If r > 0.8, only one of the variables was selected based on the relative contribution to the model to minimize biased fitting of niche models. Accordingly, seven environmental variables were used to simulate species distribution areas for R. dauricum and R. mucronulatum. Model evaluation statistics were produced from 15 replicate runs with 60% of the data used for training and 40% for model testing. Finally, the accuracy of the predicted distribution results was tested by the ROC curve analysis method, and the AUC value (the area under the ROC curve) was obtained. The closer the AUC value is to 1, the farther it is from a random distribution, the greater the correlation.
between environmental variables and the predicted geographical distribution of species, and the more accurate the model prediction results will be.

RESULTS

Genetic Diversity and Neutrality Test

The concatenated cpDNA sequences (including indels) had an aligned length of 2,711 bp, in which 15 SNPs and 10 haplotypes were identified (Table 1). Excluding the 5 haplotypes published by Jiang et al. (2016), the new 5 haplotypes sequences were deposited in the GenBank database (MT603719–MT603738). In 376 individuals from 39 populations, the haplotype diversity (Hd) and nucleotide diversity (π) ranged from 0.5018 and 0.0014, respectively. Independently, 15 SNPs and 10 haplotypes were identified in R. dauricum populations, and the Hd was 0.6610 and π was 0.0018. The results of the neutrality test showed no significant positive or negative value (Tajima’s D = 0.1348, Fu and Li’s D = −0.1766, Fu and Li’s F = 0.3798; P > 0.1). In contrast, only one haplotype (H6) was detected in R. mucronulatum populations and shared with R. dauricum; that is, there was no variation. The pairwise FST values (0.3541) showed significant genetic differentiation between R. dauricum and R. mucronulatum.

After SLAF library construction and high-throughput sequencing, a total of 735,337 SLAF tags was developed, and the average depth was 16.33x, which generated 328.74 Mb reads, with a mean Q30 of 92.02% and a GC content of approximately 41.53%. The numbers of SLAF markers in each individual ranged from 95,609 to 282,183. Among the SLAFs that were detected in total, 79,696 SLAFs were mapped to the genome and were distributed equally on each chromosome (Supplementary Figure S2A). Then, a total of 6,514,510 SNPs across the 65 accessions were identified by both GATK and SAMTools, which were considered to be reliable and equally well spread across all chromosomes (Supplementary Figure S2B). After various filtering, 664,406 SNPs were finally selected for downstream analyses. The SLAF data have been submitted to the Sequence Read Archive (SRA) database in the NCBI and the accession number was PRJNA589346. The average nucleotide diversity of R. dauricum was 0.2450 × 10^−3, which was lower than that of the cpDNA (Table 2). In contrast, the average nucleotide diversity of R. mucronulatum (0.1850 × 10^−3) was higher than that of cpDNA. The results of the neutrality test showed a significant positive value (Tajima’s D = 0.8196, Fu and Li’s D = 1.5824, Fu and Li’s F = 1.5494 in R. dauricum; Tajima’s D = 0.6661, Fu and Li’s D = 0.9743, Fu and Li’s F = 0.9989 in R. mucronulatum; P < 0.1), which might indicate that these species experienced a bottleneck in their evolutionary history or balancing selection of these loci.

Population Structure

The topology obtained from TCS based on cpDNA was used to infer relationships among the 10 cpDNA haplotypes (Figure 1A), of which H1, H6, H8 were the most frequent haplotypes and were located in the center of the haplotype network. However, these three major haplotypes are geographically segregated: H1 from the Changbai Mountains populations of R. dauricum, H6 from R. mucronulatum populations and H8 exclusive to the Great/Lesser Khingan Mountains populations of R. dauricum. Eight mutational steps differentiate H6 from H8, while only two steps separate haplotype H1 from H6 (Figure 1A). In addition, some low-frequency haplotypes were found in a few populations of R. dauricum.

To further understand the evolutionary history of the two species, we used a Bayesian clustering algorithm to estimate the population genetic structure. The results of ADMIXTURE showed that K=2 was the optimal value when the cross-validation (CV) errors were applied (Supplementary Figure S3). When K=2, R. dauricum and R. mucronulatum clustered into distinct groups and showed majority populations collected from the Changbai Mountains (MES, SL, LTS, MH, HC, LJ, CB, WT) of R. dauricum with some admixture from R. mucronulatum (Figure 2A). When K=3, the admixture R. dauricum samples were separate. PCA results also showed that samples from R. dauricum and R. mucronulatum clustered into different groups by PC1, while the samples of R. dauricum from Great/Lesser Khingan Mountains and Changbai Mountains were clustered into distinct groups by PC2 (Figure 2B), consistent with the result of the ADMIXTURE analysis.

Phylogeny and Divergence Time

The maximum-likelihood (ML) phylogenetic tree of all samples based on the 664,406 SNPs showed that R. dauricum and R. mucronulatum formed a well-supported clade, respectively. (Figure 3A). In the R. dauricum clade, the individuals from Changbai Mountain formed a subclade, and others formed another clade, with a high support value (bootstrap values > 90%). However, the ML tree based on cpDNA (Figure 3B, note that it represents a bootstrap consensus tree and therefore does not show all individual cpDNA haplotypes) presented multiple incongruences with the phylogenetic tree based on SLAF data. The cpDNA phylogeny identified three main lineages: R. mucronulatum, R. dauricum from Changbai Mountain and R. dauricum from elsewhere. R. dauricum did not constitute a monophyletic group.

Based on the two previously published calibration time points, the divergence time was calculated based on the maximum-likelihood (ML) tree using MCMCTREE (Figure 4), which pointed toward the divergence time of outgroup R. redowskianum from the most recent common ancestor of R. dauricum, and R. mucronulatum was dated to have occurred.

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TABLE 2 | Summary results of AMOVA and FST.

| Locus                  | cpDNA           | SLAF            |
|-----------------------|-----------------|-----------------|
|                       | % of variation | % of variation |
| Among species         | 28.57           | 57.98           |
| Among populations     | 68.06           | 18.59           |
| within species        |                 |                 |
| Within populations    | 3.37            | 23.43           |
| FST                   | 0.3541          | 0.2886          |

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68.65 Mya (95% HPD interval: 71.42–120.48 Mya), while the divergence time of *R. dauricum* and *R. mucronulatum* was estimated at 32.41 Mya (95% HPD interval: 17.54–53.11 Mya) during the early Oligocene. This was in effect also the divergence time between the Siberian and the Far Eastern clades of subgen. *Rhodorastrum*.

**Genetic Differentiation and Gene Flow**

Our hierarchical AMOVA showed that the percentage of variation among species was 28.57% and was significant in terms of cpDNA (P < 0.01) (Table 2). In general, the variations among species in SLAF were 57.98% higher than cpDNA and were the main source of variation, while the main source of variation in cpDNA was found among populations within species (68.06%). The pairwise F<sub>ST</sub> values between *R. dauricum* and *R. mucronulatum* showed significant genetic differentiation among species (F<sub>ST</sub> = 0.3541 in cpDNA; F<sub>ST</sub> = 0.2886 in SLAF). The results of TREEMIX suggested a higher proportion of gene flow in populations of *R. dauricum* and *R. mucronulatum* from Changbai Mountain (Figure 2C), which may be a major reason for the lack of complete species divergence and the shared haplotype (H6) between *R. dauricum* (SJ, SL, and WHL) and *R. mucronulatum* (Figure 1B).

**Distribution Model Analysis**

Based on the model fitting of the *R. mucronulatum* and *R. dauricum* distribution areas by MaxEnt software, the high ROC values (*R. dauricum*: 0.896, 0.881, 0.881, 0.909; *R. mucronulatum*: 0.910, 0.945, 0.959, 0.948) indicate the good accuracy of our model predictions. *R. mucronulatum* in China is mainly distributed in Liaoning, Beijing, Hebei, and Shandong, and the simulated result of the current potential distribution areas is basically consistent with the current distribution areas (Figure 5H), although some areas of southern China were also predicted to be suitable. Based on the paleoclimate information simulation, this species had a relatively large distribution range in the LIG period (Figure 5E), while in the subsequent LGM period (Figures 5F, G), both the CCSM and MIROC models showed that the distribution areas shrunk and migrated to the southeast. From LGM until now (Figures 5F–H), the distribution range had neither obvious shrinkage or expansion. In contrast, *R. dauricum* populations showed the opposite trend, with a relatively small distribution range in the LIG (Figure 5A) and expansion during the LGM (Figures 5B, C), with subsequent shrinkage when the climate was warm (Figure 5D). This is probably related to the fact that *R. dauricum* is a cold-tolerant plant.
DISCUSSION

Differences in Genetic Diversity Between R. dauricum and R. mucronulatum

Based on the genetic diversity values obtained in the present study, the total nucleotide diversity ($\pi = 0.0014$, Table 1) is relatively high at the subgen. *Rhodorastrum* level, and the haplotype diversity ($Hd = 0.5198$) is at a medium level. These two species showed different levels of nucleotide diversity based on SLAF and cpDNA data. The analysis results of SLAF data showed that *R. dauricum* and *R. mucronulatum* have similar nucleotide diversity, but a higher level of species-wide nucleotide diversity was found in *R. dauricum*, consistent with a previous report (Jiang et al., 2016) and that of *R. mucronulatum* was zero in cpDNA. The nucleotide diversity of *R. dauricum* was higher than that of other species of *Rhododendron*, such as *R. simii* ($\pi = 1.06 \times 10^{-3}$) (Li et al., 2012). In addition, *R. dauricum* contained all 10 haplotypes, while *R. mucronulatum* had only one haplotype (H6) and shared it with *R. dauricum*. Compared to other temperate species, such as *R. delavayi* ($Hd = 0.5570$) (Sharma et al., 2014), *R. simii* ($Hd = 0.749$) (Li et al., 2012), and *Platycrater arguta* ($Hd = 0.8820$) (Qiu et al., 2009), *R. dauricum* was in the middle ($Hd = 0.6610$), but fewer than 170 angiosperms indicate cpDNA haplotype diversity ($Hd = 0.6700$) (Petit et al., 2005).

As noted by Wright and Gaut (2005), a series of factors may contribute to genetic diversity in plant species. The niche simulation analysis showed that *R. dauricum* and *R. mucronulatum* occupy different niches (Figures 5D, H), so climate may be an important driver that causes genetic diversity between the two species (Wang et al., 2015). For the extreme situation where *R. mucronulatum* has only one cpDNA haplotype, similarly low cpDNA diversity has been found in *Acer mono* (Guo et al., 2014) and *Juglans mandshurica* (Bai et al., 2016) distributed in NEC. These provide evidence that many of these northern populations may have experienced a bottleneck during LGM. Also, the limitation of marker sampling cannot be excluded. Furthermore, due to the ornamental value of this species, the influence of human factors on diversity cannot be ignored.

Oligocene Divergent Event and Demographic Histories of Two Species

Climate changes and tectonic events have had a profound influence on the evolution of different lineages and have often been associated
with rapid divergence or speciation (Milne and Abbott, 2002; Milne, 2006; Mao et al., 2012; Zou et al., 2013). Previous studies showed drastic global cooling after the early Eocene Climatic Optimum (EECO; 52–50 Ma) (Zachos et al., 2001). Our study inferred that the divergence of *R. dauricum* and *R. mucronulatum* occurred at 32.41 Mya (i.e., the divergence between the Siberian and the Far Eastern clades, Figure 4). This may be driven by the global dramatic climate cooling and reduced sea level (Ivany et al., 2000; Katz et al., 2008; Liu et al., 2009; Buerki et al., 2013) during the late Eocene to the early Oligocene (40–30 Mya). In addition, the separation of most continents and the collision between the plates have a direct or indirect impact on speciation (Hall, 2009). During the North China tectonic period of the Middle Eocene-Oligocene (52–23.3 Mya), the Pacific plate displaced in the NWW (West-Northwest) direction. At approximately 34 Mya, the direction of motion suddenly changed from NNW (West-Northwest) to NWW, causing severe deformation

**FIGURE 5** | Ecological niche models (ENMs) for *R. dauricum* and *R. mucronulatum*. (A–D) represents the distribution of *R. dauricum*; (E–H) represents the distribution of *R. mucronulatum*. The black dots in D and H represent the sites of *R. dauricum* and *R. mucronulatum* used for simulation.
in the eastern part of the Eurasian plate, and mainland China was squeezed in an east-west direction, simultaneously resulting in a series of near-south-north folds (Wan and Cao, 1992). Thus, the divergence between Siberian and the Far Eastern clades was likely to be directly impacted by geomorphological changes.

Based on the cpDNA haplotype geographical distribution (Figure 1), we found that the two populations of MLG and WD of *R. dauricum* located in LKM have the ancient haplotype H8, special haplotypes H9, H10, and a higher haplotype diversity. The populations of *R. dauricum* in CBM also have special haplotypes and higher haplotype diversity. In addition, the results of the niche model showed that the populations of *R. dauricum* clustered to the LKM and CBM in the LIG, with subsequent local expansion during the LGM. Therefore, it is speculated that multiple refugias were maintained in *R. dauricum*. Li et al. (2012) and Bao et al. (2015) also found similar situations for *R. simsii* and *Pinus koraiensis*. In contrast, *R. mucronulatum* is currently mainly distributed in North China. The niche model showed that its distribution shrank back to the Korean Peninsula during the LGM; thus, the Korean Peninsula may be a glacial refugia for *R. mucronulatum* (Watanabe et al., 2016).

### The Relative Effectiveness of Natural Selection and Gene Flow in the Evolution of the Two Species

Natural selection and gene flow are two important influential factors in the evolution of species. Gene flow generally occurs within a species, but there are numerous examples of interspecific gene flow. Gene flow can achieve homogenizing effects between gene pools (Coyne and Orr, 2004), while natural selection can weaken this effect. When natural selection overcomes the homogenizing effects of gene flow, populations or species are able to evolve separately. Our results of the species tree estimated by TREEMIX suggested a higher proportion of gene flow in populations of *R. dauricum* from *R. mucronulatum* (Figure 2C), and the main hybrid zone was observed in the CBM region. Higher levels of migration were suggested from populations of *R. mucronulatum* to populations of *R. dauricum* located in the CBM. The divergence time analysis shows a pre-LGM divergence of the two species (Figure 4), and ecological niche modeling revealed the distribution from LGM to now of the two species overlapped in the CBM region (Figure 5). Therefore, gene flow may occur due to the overlap of regions (Li et al., 2010; Abbott et al., 2013; Sun et al., 2016).

In the haplotype geographical distribution, the range-edge populations (SJ, SL, and WHL) of *R. dauricum* contained the haplotype (H6) from *R. mucronulatum* (Figure 1B). Consistent with this, the phylogeny tree based on cpDNA showed that the *R. dauricum* individuals from CBM did not cluster with others that was assigned to the *R. mucronulatum* clade (Figure 3B). This suggests that there may be seed-mediated unidirectional gene flow from *R. mucronulatum* to the populations of *R. dauricum* in the CBM. However, the phylogeny tree based on markers sampled across the SLAF data present the correct topology that *R. dauricum* and *R. mucronulatum* form a monophyly, respectively. In addition, the results of clustering analysis based on the SLAF data (K=2) showed that individuals from the CBM experienced introgression from *R. mucronulatum*, and that group was separate when K=3 (Figure 2), but the optimal K value was 2. The AMOVA results showed that the main source of variation occurred among species in the SLAF data. In general, the main source of variation in cpDNA was among populations within species (Table 2), which may be due to the gene flow from *R. mucronulatum* increasing genetic differences between populations of *R. dauricum*. Furthermore, the FST value showed that the genetic differentiation between *R. dauricum* and *R. mucronulatum* was generally significant (Table 2). Therefore, all our results suggest that interspecies gene flow exists, but the role of gene flow is not sufficient to offset the disproportionate of natural selection, and the status of the two species has been maintained (Sun et al., 2016). Therefore, we support Tikhonova’s point mentioned above regarding the relationship between the two species that acted as two independent species.

On the other hand, the inconsistency between SLAF and cpDNA also indirectly illustrates the limitations of cpDNA for studying the genetic differentiation and speciation of closely related species, especially the presence of gene flow. Previous research has also demonstrated low species discrimination (24%) for *Rhododendron* based on cpDNA for both coding and noncoding gene regions when used singly or in combination (Li et al., 2011).

### CONCLUSION

Numerous studies have shown that speciation and diversification cannot be adequately explained only by geographic isolation. Factors such as local selection and gene flow may likely and prevalently influence the divergence and diversification of closely related species. Our study investigated the genetic structure and demographic history of two closely related species distributed in northeastern China and revealed that divergence was strongly influenced by early the Oligocene climatic shift. Despite the deep differentiation of the two species, gene flow was not completely interrupted. The significantly different biogeographic structures indicated that the two species experienced divergent selection since their differentiation and overcame the homogenizing effect of gene flow. Our study provides an additional case of the evolutionary history of two closely related species and contributes to understanding the factors affecting this process.

### DATA AVAILABILITY STATEMENT

The original contributions presented in the study are publicly available. The SLAF data can be found in NCBI: [PRJNA589346], the cpDNA data can be found here: [MT603719-MT603738]. Publicly available datasets were analyzed in this study. This data can be found here: [https://www.ncbi.nlm.nih.gov/genbank/ KT696944-KT697615].
AUTHOR CONTRIBUTIONS

BY and HX designed the research and wrote the manuscript. GZ conducted data analysis and figures made. FG performed the experiments. GZ, FG, MW, and HW collected experimental samples.

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SUPPLEMENTARY MATERIAL

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**Conflict of Interest:** Author Baiming Yang was employed by company Changchun Guoxin Modern Agricultural Technology Development Co., Ltd., The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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