Assessment of Genetic Diversity and Population Genetic Structure of *Corylus mandshurica* in China Using SSR Markers

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Abstract

*Corylus mandshurica*, also known as pilose hazelnut, is an economically and ecologically important species in China. In this study, ten polymorphic simple sequence repeat (SSR) markers were applied to evaluate the genetic diversity and population structure of 348 *C. mandshurica* individuals among 12 populations in China. The SSR markers expressed a relatively high level of genetic diversity (*Na* = 15.3, *Ne* = 5.6604, *I* = 1.8853, *Ho* = 0.6668, and *He* = 0.7777). According to the coefficient of genetic differentiation (*Fst* = 0.1215), genetic variation within the populations (87.85%) were remarkably higher than among populations (12.15%). The average gene flow (*Nm* = 1.8080) significantly impacts the genetic structure of *C. mandshurica* populations. The relatively high gene flow (*Nm* = 1.8080) among wild *C. mandshurica* may be caused by wind-pollinated flowers, highly nutritious seeds and self-incompatible mating system. The UPGMA (unweighted pair group method of arithmetic averages) dendrogram was divided into two main clusters. Moreover, the results of STRUCTURE analysis suggested that *C. mandshurica* populations fell into two main clusters. Comparison of the UPGMA dendrogram and the Bayesian STRUCTURE analysis showed general agreement between the population subdivisions and the genetic relationships among populations of *C. mandshurica*. Group I accessions were located in Northeast China, while Group II accessions were in North China. It is worth noting that a number of genetically similar populations were located in the same geographic region. The results further showed that there was obvious genetic differentiation among populations from Northeast China to North China. Results from the Mantel test showed a weak but still significant positive correlation between Nei’s genetic distance and geographic distance (km) among populations (*r* = 0.419, *P* = 0.005), suggesting that genetic differentiation in the 12 *C. mandshurica* populations might be related to geographic distance. These data provide comprehensive information for the development of conservation strategies of these valuable hazelnut resources.
Introduction

Hazelnut, Corylus mandshurica Maxim. et Rupr (synonym to C. sieboldiana), belongs to the family Betulaceae and is an important species, both economically and ecologically, among nut trees. C. mandshurica is a deciduous shrub of about 2 to 6 m in height with bracts forming a tubular husk [1–3]. The leaves are irregularly serrate and ovate leaves alternate with circular leaves. Their abaxial surface is covered heavily with pubescence. C. mandshurica is widely distributed in Japan, Korea, and Northeast to North China around Beijing. Most of C. mandshurica survived in the mountainous forest belts and deep valleys at high altitude [4]. The shells are thin and thus the ratio of kernel weight to nut weight is high [5], and the nuts are rich in nutrients, mainly including unsaturated fat, protein and vitamins [6]. Compared with C. avellana, the nuts have better taste and fragrance. Its nuts are not only suitable for fresh eating, but also suitable for baking. Its nuts are an important ingredient in cake, and are a favorite of consumers [7]. Moreover, its leaves can be used as fodder to feed domestic silkworms by local farmers. In addition to its economic value, hazelnut is useful for soil and water conservation and sustainable use of local forests [8].

In recent years, molecular markers have shown promise for assessment of genetic diversity, owing to their high discriminatory power and comparatively low cost. In particular, the superiority of molecular markers for the characterization of plants is well recognized. DNA based molecular marker technologies, such as simple sequence repeat (SSR), amplified fragment length polymorphism (AFLP), sequence-related amplified polymorphism (SRAP) and single nucleotide polymorphisms (SNPs), have several advantages including abundant, independent from the environment, suitability for early and rapid evaluation, and having non-tissue specific characteristics. Among them, SSRs, also known as microsatellites or short tandem repeats (STR), are widely present in eukaryotic genomes [9] and very useful for a number of reasons including co-dominant inheritance, high polymorphism, high variability and suitability for automated allele sizing and cross-species transferability.

SSRs have been developed for European hazelnut [10–14], and parentage relationships among the most important hazelnut cultivars were identified [15]. The genetic linkage map has been established [16–18]. Moreover, microsatellite markers are considered as valuable tools in molecular breeding of C. mandshurica [19]. Besides, SSR markers have already been successfully applied to the evaluation of the genetic diversity and population structure of European hazelnut [20–21], comprising eastern filbert blight (EFB)-resistant cultivars [22]. Evaluation of genetic diversity by SSR markers is of importance not only for the breeding of hazelnut, but also for the efficient management and protection of hazelnut germplasm. Characterization of the genetic diversity and genetic differentiation is indispensable for the efficient protection of hazelnut by helping understand its population dynamics, origin and the evolution process [23]. However, limited information has been acquired on the genetic diversity of C. mandshurica resources in China, and its genetic variation and population genetic structure are still unknown.

In this study, we present the first investigation of genetic diversity of wide-ranging hazelnut (C. mandshurica) in China, with a particular focus on population structure using SSR markers. A total of 12 C. mandshurica populations throughout its present main Chinese distribution range were studied. The aims of this study were to: (1) evaluate the genetic diversity and population structure of C. mandshurica, (2) estimate variance components and partition the within- and among-population variance, (3) obtain comprehensive information for the development of conservation strategies for these valuable C. mandshurica resources.
Materials and Methods

Ethics statement

The collection of plant samples and research activities were conducted with the permission of local forestry departments. No other endangered or protected species were involved in this study.

Population sampling and DNA extraction

A total of 348 trees of *C. mandshurica* were collected from 12 populations, each represented by 20 to 31 individuals (located at least 100 m apart) (Fig 1, Table 1). Fresh young leaf tissues (three to five leaves per tree) of individual plants were collected and promptly dried with silica gel in a sealed manila envelope and then taken back to the laboratory for later DNA extraction. Total DNA was extracted from about 20 mg of leaf tissue per tree, using the modified CTAB procedure [24].

Simple sequence repeat (SSR)

Ten microsatellite loci, CAC-A040, CAC-B005, CAC-B001, CAC-B020, CAC-B028, CAC-B113, CAC-B114, CAC-C003, CAC-C028, and CAC-B105 were analyzed for polymorphism [10, 25].
Polymerase chain reaction (PCR) amplification was performed in a total of 20 μL volume that contained 10–50 ng of plant DNA, 0.2 mM each of forward and reverse primers, 1.5 mM MgCl₂, 50 mM Tris–HCl, 0.2 mM of each dNTP, 1 unit of Taq DNA polymerase (Biotech International) and accompanying buffer. PCR amplification was performed with the following cycling parameters: first a denaturation step at 94°C for 5min, followed by 35 cycles of 94°C for 30 s, annealing at 48°C~55°C for 40 s (different primer annealing temperature, Table 2), 72°C for 40 s and a final extension at 72°C for 3 min. The forward primers were labeled with a fluorochrome (FAM). Amplified fragments of SSRs were analyzed with an ABI 3730XL capillary sequencer (Applied Biosystems) separately along with an internal size standard (GeneScan-500 LIZ, Applied Biosystems). The SSR allele sizes were called with GENEMAPPER software (version 4.0, Applied Biosystems) for all populations and entered in a spreadsheet.

### Table 1. Location of populations, number of individuals sampled in a study of genetic diversity of Corylus mandshurica in China by SSR analyses.

| Population code | Location             | Number of samples | Latitude (N) | Longitude (E) | Altitude (m) |
|-----------------|----------------------|-------------------|--------------|---------------|--------------|
| MJG             | Mengjiagang, Heilongjiang | 30                | 46°26'       | 130°36'       | 326          |
| HR              | Huairou, Beijing     | 26                | 40°19'       | 116°35'       | 180          |
| LH              | Longhua, Hebei       | 28                | 41°18'       | 117°45'       | 643          |
| LQ              | Shangzai, Shanxi     | 31                | 39°19'       | 114°17'       | 1391         |
| FN              | Xiamiao, Hebei       | 29                | 41°12'       | 116°40'       | 723          |
| NA              | Ning’an, Heilongjiang | 30                | 44°18'       | 129°40'       | 534          |
| MS              | Mishan, Heilongjiang | 20                | 45°28'       | 131°56'       | 146          |
| WC              | Weichang, Hebei      | 30                | 41°56'       | 117°44'       | 980          |
| TL              | Fanhe, Liaoning      | 30                | 42°12'       | 123°46'       | 125          |
| CC              | Chicheng, Hebei      | 33                | 40°51'       | 115°44'       | 1302         |
| HG              | Xinhua, Heilongjiang | 31                | 47°23'       | 130°16'       | 214          |
| GZ              | Shagang, Liaoning    | 30                | 40°19'       | 122°27'       | 214          |
| Total           |                      | 348               |              |               |              |

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### Table 2. Genetic diversity at 10 SSR loci in 348 individuals of C. mandshurica.

| Locus   | Allele size range(bp) | Annealing temperature(°C) | Na  | Ne  | I   | Ho  | He  |
|---------|-----------------------|---------------------------|-----|-----|-----|-----|-----|
| CAC-A040| 250-300bp             | 50                        | 13  | 3.2180 | 1.6361 | 0.5891 | 0.6902 |
| CAC-B001| 98-120bp              | 50                        | 19  | 7.5308 | 2.2955 | 0.8928 | 0.8685 |
| CAC-B114| 140-160bp             | 55                        | 17  | 5.9692 | 2.0581 | 0.5934 | 0.8337 |
| CAC-C028| 120-150bp             | 51                        | 7   | 2.0804 | 0.9193 | 0.4339 | 0.5201 |
| CAC-C003| 100-130bp             | 48                        | 10  | 3.7291 | 1.5314 | 0.4725 | 0.7329 |
| CAC-B113| 148-180bp             | 51                        | 12  | 6.3050 | 2.0886 | 0.7666 | 0.8426 |
| CAC-B105| 130-160bp             | 51                        | 16  | 5.8002 | 1.9360 | 0.7861 | 0.8288 |
| CAC-B028| 260-280bp             | 51                        | 18  | 6.6402 | 2.1356 | 0.7420 | 0.8506 |
| CAC-B020| 220-270bp             | 50                        | 28  | 12.1102| 2.7178 | 0.8309 | 0.9188 |
| CAC-B005| 270-280bp             | 51                        | 13  | 3.2209 | 1.5347 | 0.5607 | 0.6905 |
| Mean    |                       |                           | 15.3| 5.6604 | 1.8853 | 0.6668 | 0.7777 |
| St. Dev |                       |                           | 5.8128| 2.8877 | 0.4990 | 0.1572 | 0.1188 |

For each locus: Number of alleles observed (Na), Effective number of alleles (Ne), Shannon's Information index (I), Observed heterozygosity (Ho), Expected heterozygosity (He), Gene flow (Nm)

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Data analysis

The presence of null alleles of each locus was estimated by software MICRO-CHECKER 2.2.3 [26]. The number of alleles observed \( (Na) \), effective number of alleles \( (Ne) \) [27], Shannon’s information index \( (I) \), observed heterozygosity \( (Ho) \), expected heterozygosity \( (He) \), genetic differentiation coefficient \( (Fst) \), gene flow \( (Nm) \) and Nei’s genetic distance \( (Gd) \) [28] (S1 Table) were calculated using Popgene32 software. \( Nm \) was estimated as: \( Nm = \frac{0.25(1-Fst)}{Fst} \). Cluster analysis was carried out based on the Nei’s genetic distance matrix with UPGMA (unweighted pair group method of arithmetic averages) and a dendrogram was constructed using MEGA (version 5) [29].

Genetic structure analysis was performed with the program STRUCTURE 2.3.4 with 10 runs and 100,000 Markov Chain Monte Carlo (MCMC) repetitions after a burn-in period of 100,000 interactions for each group number K. The optimum value of K was obtained by calculating the \( \Delta K \) value to determine the most likely number of groups [30]. Analysis of molecular variance (AMOVA) was carried out using the program Arlequin (version 3.5) for estimation of variance components and partition of the within- and among-population variance [31]. In addition, the Mantel test was conducted using the program GenALEX 6.5 for correlation between Nei’s genetic distance [32] and geographic distance (km) [33]. Significance was assessed by conducting 9999 permutations.

Results

Microsatellite variation and genetic diversity

In this study, MICRO-CHECKER analysis indicated the presence of null alleles at a few loci in some populations. At CAC-B114, CAC-C003 and CAC-B028, null alleles were detected in five populations, at CAC-B105, CAC-A040, CAC-B020 and CAC-B005, null alleles were detected in two populations, and at CAC-C028, null alleles were detected in population TL, while at CAC-B001 and CAC-B113, no null alleles were detected. Thus, these 10 SSR loci are a proper set to assess genetic diversity of *C. mandshurica*.

A total of 153 alleles were found at the ten microsatellite loci in the 348 individuals (Table 2). The number of alleles observed \( (Na) \) per locus varied from 7 (CAC-C028) to 28 (CAC-B020) with a mean of 15.3 per locus. At the same time, effective numbers of alleles \( (Ne) \) varied from 2.0804 (CAC-C028) to 12.1102 (CAC-B020) with an average of 5.6604 per locus. Shannon’s Information index \( (I) \) averaged 1.8853 and ranged from 0.9193 (CAC-C028) to 2.7178 (CAC-B020). Observed heterozygosity \( (Ho) \) and expected heterozygosity \( (He) \) ranged from 0.4339 (CAC-C028) and 0.5201 (CAC-C028) to 0.8928 (CAC-B001) and 0.9188 (CAC-B020) respectively.

Genetic differentiation

The inbreeding coefficient \( (Fis) \) per locus ranged from -0.1167 (CAC-B001) to 0.1143 (CAC-B114), with an average of 0.0215 alleles per locus. Furthermore, genetic differentiation \( (Fst) \) of individual loci ranged from 0.0689 at CAC-A040 to 0.2016 at CAC-B114, and averaged at a value of 0.1215 alleles per individual locus, suggesting low genetic differentiation among the populations. Moreover, the genetic diversity within populations (87.85%) was significantly higher than that between populations (12.15%). In addition, gene flow \( (Nm) \) ranged from 0.9903 at CAC-B114 to 3.3798 at CAC-A040 and averaged 1.8080 (Table 3). Similarly, the results of AMOVA analysis indicated that variation within populations was 88.50% while variation among populations was 11.50% (Table 4), possibly because of the relatively high gene flow \( (Nm = 1.8080) \) between *C. mandshurica* populations.
Genetic relationships and population structure analysis

A dendrogram was drawn based on the SSR data depicting the genetic relationships among the 12 populations (Fig 2). The populations were differentiated into two main clusters, and each cluster was further separated into smaller clusters. Group I included populations from Heilongjiang Province (MJG, NA, MS and HG) and Liaoning Province (GZ and TL). Group II contained populations from Hebei Province (LH, FN, WC and CC), Shanxi Province (LQ) and Beijing City (HR), and was separated into two sub clusters. The results showed that distinct genetic differentiation among populations from Northeast China and North China.

The 348 hazelnut individuals were further assessed for population stratification using the STRUCTURE program. SSR data were analyzed with possible cluster number (K-value) ranging from 1 to 12. The ΔK showed a clear maximum for K = 2 (ΔK = 185.0287), indicating among 12 populations the existence of two groups (S1 Fig). Group I included populations from MJG, NA, MS, TL, HG and GZ. Group II included populations from HR, LH, LQ, FN, WC and CC (Fig 3). Structure analysis suggested differentiation between the two groups, and grouped them approximately in line with the geographic area. These results indicated that there were different degrees of introgression in the populations, detected as differences in allele frequencies among the populations.

The Mantel test revealed a weak but positive correlation between Nei’s genetic distance and geographic distance (km) (r = 0.419, P = 0.005; Fig 4), suggesting that genetic differentiation in the 12 populations might be caused by geographic isolation due to distance.

Discussion

SSR polymorphism and genetic diversity among populations

In recent years, SSR marker techniques have been extensively applied in the detection of genetic variation in hazelnut populations, thereby estimating their genetic diversity [34–39]. In this study, SSR polymorphism was examined using 10 microsatellite loci to analyze the genetic diversity of 348 hazelnut individuals from 12 populations in China. The results showed that genetic diversity was significantly higher in the populations from Northeast China than in those from North China. The genetic diversity was also positively correlated with geographic distance, suggesting that isolation-by-distance played a major role in the genetic differentiation of these populations.

Table 3. Summary of F statistics and gene flow for the 10 loci.

| Locus      | Fis        | Fst        | Nm        |
|------------|------------|------------|-----------|
| CAC-A040   | 0.0935     | 0.0689     | 3.3798    |
| CAC-B001   | -0.1167    | 0.0770     | 2.9960    |
| CAC-B114   | 0.1143     | 0.2016     | 0.9903    |
| CAC-C028   | 0.0749     | 0.0878     | 2.5972    |
| CAC-C003   | 0.2052     | 0.1772     | 1.1607    |
| CAC-B113   | -0.0616    | 0.1413     | 1.5195    |
| CAC-B105   | -0.0718    | 0.1113     | 1.9968    |
| CAC-B028   | 0.0275     | 0.1055     | 2.1196    |
| CAC-B020   | 0.0119     | 0.0834     | 2.7469    |
| CAC-B005   | 0.0200     | 0.1579     | 1.3336    |
| Mean       | 0.0215     | 0.1215     | 1.8080    |

For each locus: Fis, coefficient of inbreeding; Fst, Genetic differentiation coefficient; Nm, Gene flow

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Table 4. Analysis of molecular variance (AMOVA) of genetic diversity of C. mandshurica populations.

| Source of variation | d.f. | Sum of squares | Variance components | Percentage of variation | P      |
|---------------------|------|----------------|---------------------|------------------------|--------|
| Among populations   | 11   | 317.662        | 0.43999             | 11.50%                 | <0.001 |
| Within populations  | 684  | 2316.455       | 3.38663             | 88.50%                 | <0.001 |
| Total               | 765  | 2947.051       | 3.89756             |                        |        |

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study, the SSR marker analysis showed a high degree of genetic diversity *C. mandshurica*. The average number of alleles observed per locus (15.3) was significantly higher than other *Corylus* species such as *C. heterophylla* (4.18) [40] and *C. avellana* (7.16 [10], 7.1 [41] and 11.9 [25]). *Ho* and *He* values were 0.6709 and 0.7954 respectively. These results were similar to the values previously reported by Bassil et al [10] for *C. avellana* and Wang et al [40] for *C. heterophylla*.

The high genetic diversity in *C. mandshurica* populations may relate to the biological characteristics and living environment of this species. *C. mandshurica*, as a widely distributed, perennial woody plant species, can preserve its genetic diversity for quite a long period of time. Moreover, *C. mandshurica* is monoeocious, wind-pollinated and self-incompatible mating system of species. In a long term, outcrossing will decrease differentiation among individuals in a diverse population. In addition, *C. mandshurica* is an ancient species capable of clonal propagation due to the formation of adventitious buds on rhizomes. Asexual reproduction and sexual reproduction of plants will produce a life-long effect. Clonal and sexual preproduction can result in many generations coexisting in a population. Such populations are less susceptible to genetic drift [42] and help to preserve genetic variation.

**Genetic structure, gene flow and differentiation among populations**

AMOVA revealed that 11.50% of variation is among populations and the remaining 88.50% of variation is within populations (*P*<0.001) (Table 4). Moreover, the results of AMOVA analysis
Fig 3. Bayesian STRUCTURE bar plot of membership for 12 *C. mandshurica* populations (K = 2). Red represents Groups I and green represents Groups II. For details of locations abbreviations and locations, see Table 1 and Fig 1.

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Fig 4. Mantel test for matrix correlation between Nei’s genetic distance and geographic distance for 12 *C. mandshurica* populations.

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were consistent with the mean value of $F_{st}$, further suggesting the existence of low genetic differentiation among different populations. Similar results were recently reported by Di et al [38], who analyzed 8 populations of *C. heterophylla* and showed that 16% of the genetic variation existing among populations.

Comparison of these results with the UPGMA dendrogram and the Bayesian STRUCTURE ($K = 2$) bar plot showed general agreement between the population subdivisions and the genetic relationships among the populations. The populations were divided into two main clusters. Similar results were previously reported on Chinese walnut (*Juglans mandshurica*) [43], two refuges in the last ice age have been inferred: The Taihang and Qinling Mountain in North China and Changbai Mountain in Northeast China. Another possible reason relates to the existence of the Yanshan Mountains. It is worth noting that a number of genetically similar populations were located in the same geographic region. The Mantel test revealed a significant correlation between the genetic distance and geographic distance among populations ($r = 0.419, P = 0.005$). Therefore, it’s speculated that the genetic structure of *C. mandshurica* population in China can be affected by geographic distance.

Gene flow is defined as the transfer of alleles from one population of a species to another. Study of gene flow is critical for the understanding of population processes within and among species [44]. The carrying of alleles to a population where they did not previously exist can be a very important origin of genetic variation [45]. The most important factor impacting the rate of gene flow between different populations is mobility [46]. Slatkin [47] considered that if $Nm > 1$, the gene exchange among population can prevent the impact of genetic drift and reduce the genetic variance among populations. In this study, the average of gene flow of *C. mandshurica* at each locus was 1.8080, indicating that the gene flow was one of important factors influencing the genetic structure of *C. mandshurica* populations. The relatively high level of gene flow likely prevents genetic differentiation, which is the reason for the observed low genetic differentiation. That is the reason why the variance within populations was significantly higher than that between populations. Therefore, the lack of differentiation is attributed to gene flow.

Gene flow among populations was influenced by a number of factors including the mating system, geographic distribution, mechanisms of seed dispersal, and the stage of succession and colonization [48, 49]. An earlier study found that the pollen and seed dispersal was the foremost mechanism for gene exchange among plants [50]. As pointed out in a more recent study, gene flow mainly arises from two diffusion processes: seed, the extent and relative importance of which largely influence the landscape of neutral genetic diversity across a geographical area [44]. *C. mandshurica* is wind-pollinated and self-incompatible; its pollen is easily dispersed over medium and long distances. In addition, the nuts of *C. mandshurica* are nutritious and thus serve as an important food source for small mammals and birds, who can transport them in small numbers over long distances. All of these factors can contribute to the observed high gene flow between different populations.

**Genetic resource conservation strategies**

In recent years, it has become increasingly important to adopt a holistic view of biodiversity, which includes agricultural biodiversity, conservation and sustainable development. An in-depth understanding of hazelnut genetic diversity and ecological distribution is indispensable for the protection and utilization of this species [35]. Analyzing the genetic diversity and genetic structure of *C. mandshurica* populations can provide a theoretical basis for the conservationist to formulate future conservation strategies of *C. mandshurica*.

In this work, the results based on SSR techniques discovered that wild *C. mandshurica* populations have a relatively high level of genetic diversity. However, in the past several decades,
excessive deforestation and over-exploitation have damaged the ecological environment of many wild *C. mandshurica* living on natural hills. And local farmers have devastated the natural habitats of some populations in order to obtain the economic value. If not properly preserved, this species will likely become endangered because of both high demand and adverse climatic conditions, and even become extinct in the future. We suggest *in situ* conservation method: conservation to prevent the decrease in population sizes and loss of genetic diversity. Moreover, in order to achieve effective conservation of *C. mandshurica* germplasm resources, efforts are needed to carefully plan and construct pollen banks and gene banks for hazelnut. In the meantime, protected areas can be established to conserve and restore the habitat and the populations. In addition, it is important to develop a core collection of *C. mandshurica* in greater breadth and depth, which would not only mitigate the pressure of excessive use of wild resources, but also help achieve more effective management and use of hazelnut germplasm.

**Supporting Information**

S1 Fig. Delta K value as a function of K based on 10 runs, indicating the most likely number of two genetic clusters. DeltaK = mean (|L''(K)|) / sd(L(K)).

(TIF)

S1 Table. Nei's genetic distance among *C. mandshurica* populations.

(DOCX)

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**Author Contributions**

Conceived and designed the experiments: GXW TTZ JWZ QHM. Performed the experiments: JWZ TTZ GXW QHM LSL. Analyzed the data: JWZ TTZ GXW. Contributed reagents/materials/analysis tools: JWZ TTZ GXW. Wrote the paper: JWZ TTZ GXW.

**References**

1. Wu ZY. Vegetation in China. Beijing: Science and Technology Press; 1995.
2. Zhang Y, Li F, Tao R, Li Z, Liang Y. An investigation of wild Corylus resource at Changbai Mountains. J Jilin Agric Sci. 2007; 32:56–57.
3. Editorial Board of the Flora of China Academy of Sciences. Flora of China. Beijing: Science Press; 1979.
4. Zhang PF, Zhang FH, Zhang R, Liu YL, Wang ZG. Genetic diversity of hazel populations in Shanxi province based on SSR markers. Plant Sci J. 2014; 32: 131–138.
5. Zhao D, Su S, Ni B, Wang W, Meng X, Liu W. Germplasm resources investigation and utilization prospects of hazel in Small Xing'an Ridge region. Chin Agric Sci Bull. 2012; 28: 87–94.
6. Liu HJ, Liu GC, Liu GY, Han CR. Analysis on Properties of Woody Diesel Plants Corylus mandshurica Maxim.el Rupr. J Chin Cere Oils Assoc. 2013; 28: 41–45.
7. Wang GX. Shanxi forest. Beijing: China Forestry Press; 1992.
8. Liu H. Exploring the utilization of Corylus. Farm Prod Proc. 2010; 1: 24–25.
9. Sharma PC, Grover A, Kahi G. Mining microsatellites in eukaryotic genomes. Trends Biotechnol. 2007; 25: 490–498. PMID: 17945369
10. Bassil NV, Botta R, Mehlenbacher SA. Microsatellite markers in the hazelnut: isolation, characterization and cross-species amplification in Corylus. J Am Soc Hort Sci. 2005; 130: 543–549.
11. Bassil NV, Botta R, Mehlenbacher SA. Additional microsatellites of the European hazelnut. In VI International Congress on Hazelnut. 2005; 686: 105–110.
12. Boccacci P, Akkak A, Bassil NV, Mehlenbacher SA, Botta R. Characterization and evaluation of microsatellite loci in European hazelnut (Corylus avellana L.) and their transferability to other Corylus species. Mol Ecol Notes. 2005; 5: 934–937.
13. Gürcan K, Mehlenbacher SA. Transferability of microsatellite markers in the Betulaceae. J Amer Soc Hort Sci. 2010; 135: 159–173.
14. Gürcan K, Mehlenbacher SA. Development of microsatellite marker loci for European hazelnut (Corylus avellana L.) from ISSR fragments. Mol Breeding. 2010; 26: 551–559.
15. Boccacci P, Akkak A, Botta R. DNA-typing and genetic relationships among European hazelnut (Corylus avellana L.) cultivars using microsatellite markers. Genome. 2006; 49: 598–611. PMID: 16936839
16. Mehlenbacher SA, Brown RN, Nouhra EN, Gökirmak T, Bassil NV, Kubisiak TL. A genetic linkage map for hazelnut (Corylus avellana L.) based on RAPD and SSR markers. Genome. 2006; 49: 122–133. PMID: 16498462
17. Sathuvalli VR, Chen H, Mehlenbacher SA, Smith DC. DNA markers linked to eastern filbert blight resistance in “Ratoli” hazelnut (Corylus avellana L.). Tree Genet Genomes. 2011; 7: 337–345.
18. Rowland LJ, Ogden EL, Bassil N, Buck EJ, McCallum S, Graham J, et al. Construction of a genetic linkage map of an interspecific diploid blueberry population and identification of QTL for chilling requirement and cold hardiness. Mol Breeding. 2014; 34: 2033–2048.
19. Cheng L, Huang W, Zhou Z, Liu J, Wang Y, Su S, et al. Genetic diversity of six Corylus species in China detected with microsatellite isolated from Corylus avellana. Sci Silvae Sin. 2009; 45: 22–26.
20. Gürcan K, Mehlenbacher SA, Erdoğan V. Genetic diversity in hazelnut (Corylus avellana L.) cultivars from Black Sea countries assessed using SSR markers. Plant Breeding. 2010; 129: 422–434.
21. Boccacci P, Botta R. Microsatellite variability and genetic structure in hazelnut (Corylus avellana L.) cultivars from different growing regions. Sci Hortic-Amsterdam. 2010; 124: 128–133.
22. Muehlbauer MF, Honig JA, Capik JM, Vaichunas JN, Molnar TJ. Characterization of eastern filbert blight-resistant hazelnut germplasm using microsatellite markers. J Amer Soc Hort Sci. 2014; 139: 399–432.
23. Fritsch P, Rieseberg LH. The use of random amplified polymorphic DNA (RAPD) in conservation genetics. Mol Genet Approaches Conser. 1996; 1996: 54–73.
24. Doyle JJ, Doyle JL. A rapid DNA isolation procedure for small quantities of fresh leaf tissue. Phytochemical Bulletin. 1987; 19: 11–15.
25. Gökirmak T, Mehlenbacher SA, Bassil NV. Investigation of genetic diversity among European hazelnut (Corylus avellana) cultivars using SSR markers. Acta Hortic. 2005; 686: 141–147.
26. Van Oosterhout C, Hutchinson WF, Wills DP, Shipley P. MICRO-CHECKER: software for identifying and correcting genotyping errors in microsatellite data. Mol Ecol Notes. 2004; 4: 535–538.
27. Kimura M, Crow JF. The number of alleles that can be maintained in a finite population. Genetics. 1964; 49: 725. PMID: 14156929
28. Slatkin M, Barton NH. A comparison of three indirect methods for estimating average levels of gene flow. Evolution. 1989; 43: 1349–1368.
29. Tamura K, Peterson D, Peterson N, Stecher G, Nei M, Kumar S. MEGAS: Molecular Evolutionary Genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. Mol Biol Evol. 2011; 28: 2731–2739. doi: 10.1093/molbev/msr121 PMID: 21946353
30. Evanno G, Regnaut S, Goudet J. Detecting the number of clusters of individuals using the software STRUCTURE: a simulation study. Mol Ecol. 2005; 14: 2611–2620. PMID: 15969739
31. Excoffier L, Lischer HEL. ARIEQUIN suite ver 3.5: a new series of programs to perform population genetics analyses under Linux and Windows. Mol Ecol Resour. 2010; 10: 564–567. doi: 10.1111/j.1755-0998.2010.02847.x PMID: 21565059
32. Nei M. Genetic distance between populations. Am Nat. 1972; 106, 283–292.
33. Peakall R, Smouse PE. GenAIEx 6.5: genetic analysis in Excel. Population genetic software for teaching and research—an update. Bioinformatics. 2012; 28: 2537–2539. PMID: 22820204
34. Erfatpour M, Hamidogli Y, Kaviani B, Fatahi R, Falahati M, Javadi D, et al. Assessment of genetic diversity among some Iranian hazelnut genotypes using SSR markers. Aust J Crop Sci. 2011; 5: 1286.
35. Sathuvalli VR, Mehlenbacher SA. Characterization of American hazelnut (*Corylus americana*) accessions and *Corylus americana* × *Corylus avellana* hybrids using microsatellite markers. Genet Resour Crop Ev. 2012; 59: 1055–1075.

36. Bacchetta L, Rovira M, Tronci C, Aramini M, Drogoudi P, Silva AP, et al. A multidisciplinary approach to enhance the conservation and use of hazelnut (*Corylus avellana*) genetic resources. Genet Resour Crop Ev. 2014; doi: 10.1007/s10722-014-0173-7

37. Valentini N, Calizzano F, Boccacci P, Botta R. Investigation on clonal variants within the hazelnut (*Corylus avellana*) cultivar ‘Tonda Gentile delle Langhe’. Sci Hortic-Amsterdam. 2014; 165: 303–310.

38. Di XY, Liu KW, Hou SQ, Ji PL, Wang YL. Genetic variation of hazel (*Corylus heterophylla*) populations at different altitudes in Xingtangsi forest park in Huoshan. Plant Omics J. 2014; 7: 213–220.

39. Mohammadzedeh M, Fattahi R, Zamani Z, Khadivi-Khub A. Genetic identity and relationships of hazelnut (*Corylus avellana*) landraces as revealed by morphological characteristics and molecular markers. Sci Hortic-Amsterdam. 2014; 167: 17–26.

40. Wang YM. Study on the phylogenetic relationship of *corylus* species and genetic diversity of *C. heterophylla*. M.Sc. Thesis, Beijing Forestry University. 2008. Available: http://www.cnki.net/KCMS/detail/detail.aspx?dbcode=CDFD&QueryID=6&CurRec=6&dbname=CDFD9908&filename=2008085055.nh&urlid=&yx=&v=MzAzMzMzczVRyV0oxRnJDVJMK2ZadVZ2RnlYbFZidkJWMT3RnJPd0c5SEpxEVIEJOGVYMUp1eFITN0RoMVQ=

41. Boccacci P, Botta R, Rovira M. Genetic diversity of hazelnut (*Corylus avellana*) germplasm in northeastern Spain. Sci Hortic-Amsterdam. 2008; 43: 667–672.

42. Loveless MD, Hamrick JL. Ecological determinants of genetic structure in plant populations. Annu Rev Ecol Evol Syst. 1984; 15: 65–95.

43. Bai W, Liao W, Zhang D. Nuclear and chloroplast DNA phylogeography reveal two refuge areas with asymmetrical gene flow in a temperate walnut tree from East Asia. New Phytol. 2010; 188: 892–901. doi: 10.1111/j.1469-8137.2010.03407.x PMID: 20723077

44. Gerber S, Chadoeuf J, Gugerli F, Lascoux M, Buiteveld J, Cottrell J, et al. High Rates of Gene Flow by Pollen and Seed in Oak Populations across Europe. Plos One. 2014; 9: e85130. doi: 10.1371/journal.pone.0085130 PMID: 24454802

45. Futuyma DJ. Evolutionary Biology. 3rd ed. Sunderland MA: Sinauer Assoc; 1998.

46. Mayr E. Evolution and the Diversity of Life. Selected essays, Cambridge: Harvard University Press; 1997.

47. Slatkin M. Rare alleles as indicators of gene flow. Evolution. 1985; 39: 53–65.

48. Hamrick JL, Godt MJW, Brown AHD, Clegg MT, Kahler AL, Weir BS. Allozyme diversity in plant species. Plant population genetics, breeding, and genetic resources: Sinauer Assoc Inc; 1990.

49. Levin DA, Kerster HW. Gene flow in seed plants. In Evolutionary biology. Springer US; 1974, pp. 139–220.

50. Whitlock MC, McCauley DE. Indirect measures of gene flow and migration: $F_{st}$ doesn't equal 1/ (4Nm+1). Heredity. 1999; 82: 117–125. PMID: 10098262