Genetic Diversity and Structure of Walnut Populations in Central and Southwestern China Revealed by Microsatellite Markers

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Abstract. Molecular markers were used to study the genetic diversity, structure, and relationship of Juglans L. with nine populations (five from Juglans regia L. and four from Juglans sigillata Dode) in central and southwestern China. A moderate level of genetic diversity was observed at the population level with the number of effect alleles per locus (Ae) ranging from 1.75 to 3.35 (average 2.29) and the proportion of polymorphic loci (P) equaling 100.0%. The expected heterozygosity (He) within populations ranged from 0.389 to 0.687, and the average was 0.525. The proportion of genetic variation presented among populations accounted for 18.6% of the total genetic diversity. The overall gene flow (Nm) among populations equaled 1.10. The unweighted pair-group method using arithmetic averages (UPGMA) clustering and the Mantel test showed that genetic distances among the nine populations are in a good agreement with their geographic distribution, supporting the viewpoint that J. regia and J. sigillata belong to one species. We suggest that the central area of the southwestern mountain regions of China could be considered as a priority for walnut genetic resource conservation.

The genus Juglans is characterized by a monoecious and heterodichogamous habit. It includes about 20 species (Gleeson, 1982; Manning, 1978), distributed over a wide geographical range, including southern Europe, eastern Asia, and the Americas (Krusman, 1986), and displays differentiation in morphology, particularly in nut characteristics. Patterns of population genetic diversity and structures for some species of this genus have been studied by several molecular marker systems, such as isozymes (Arulsekar et al., 1985; Fornari et al., 2001; Solar et al., 1993), restriction fragment-length polymorphism (RFLP) (Fjellstrom and Parfitt, 1994), randomly amplified polymorphic DNA (RAPD) (Nicese et al., 1998), and intersimple sequence repeat (ISSR) markers (Potter et al., 2002). Simple sequence repeat (SSR) markers exhibit hypervariability and codominance and are highly informative in nature. They exemplify numerous applications in the understanding of the genetic structure of the genus Juglans (Dangl et al., 2005; Foroni et al., 2005; Victory et al., 2006; Woeste et al., 2002).

China is an important place for the resources of genus Juglans. There are 12 walnut species that occur there naturally, including the two dominantly cultivated species, Juglans regia and Juglans sigillata (Kuang and Lu, 1979). Juglans sigillata is found only in southwestern China (Qu and Sun, 1990). Despite the significant global economic value of China’s nut and wood products, China produces about 27.5% of the world’s total nut production (Food and Agriculture Organization of the United Nations, 2005), and little is known about the genetic diversity and structure of populations like the walnut species in China. Southwestern China is a special alpine area with various climate and geographical features and a rich plant diversity (Sun, 2002). The Qinling Mountain range, which stretches from the east to west in central China, is particularly rich in plant diversity. It is noteworthy that populations of J. sigillata and J. regia still grow there, particularly in the canyon regions where human activity, if any, is rare. Moreover, a long-standing debate has arisen about whether J. sigillata and J. regia are regarded as completely different species or as just different ecological types (Kuang and Lu, 1979; Wu et al., 2000; Yang and Xi, 1989). Morphological comparisons allowed L.A. Dode to classify J. sigillata as a new species in 1906 (as cited in Xi and Zhang, 1996). Kuang and Lu (1979) put J. regia and J. sigillata into the section Juglans of the Juglans genus. However, on the basis of peroxidase isozyme analysis, Yang and Xi (1989) suggested that J. sigillata and J. regia are two different ecological types of one species. This appears to be supported by hybridization experiments in which the filial...
generation of *J. regia* and *J. sigillata* was observed to be progestive (Fan and Xi, 2002; Fan et al., 2005; Yang et al., 2007).

The motivation behind this study is to examine the degree of genetic diversity and differentiation of walnuts collected from central and southwestern China and to understand the genetic relatedness of *J. regia* and *J. sigillata* with advanced SSR markers. Our results will provide scientific guidance about effective management, conservation, and improvement of walnut resources.

**Materials and Methods**

**Plant materials.** According to the field investigations of other researchers and the information supplied by local forestry agencies, five representative populations of *J. regia* (*N* = 154) and four representative populations of *J. sigillata* (*N* = 132) were sampled in the provinces of Yunnan, Sichuan, Shanxi, and Henan in central and southwestern China (Table 1). The populations were all autochthonous, and their sites were not managed. All sampled trees were estimated to be older than 60 years. The plants of *J. regia* and *J. sigillata* were distinguished according to their phenotypes (Xi and Zhang, 1996). Six populations (SJC-r, QZ-r, LJ-s, ML-s, YB-s, and MN-s) from southwestern China inhabit disjunctive mountainous areas with a narrow latitudinal and longitudinal range (see Fig. 1). The average distance between any pair of populations was about 80 km. Sampled trees within a population were separated by a distance of more than 50 m. Fresh leaves from each plant were collected and stored at −80 °C until DNA extraction for marker analyses.

**DNA extraction.** DNA extraction from young leaves followed the method of Doyle and Doyle (1987), with some modification: 1 g of frozen, ground leaves was added to 10 mL of pre-heated (50 °C) 2× CTAB buffer [2% cetyl trimethyl ammonium bromide, 50 mM 1,4-dithiothreitol, 0.3% β-mercaptoethanol, 1.4 M NaCl, 100 mM Tris, 20 mM ethylenediaminetetraacetic acid (EDTA), pH 8.0] and incubated at 65 °C for 40 min. The aqueous solution was extracted with 10 mL (24:1) of chloroform–isoamyl alcohol and centrifuged at 5000 ×g for 10 min, and the aqueous layer retained; 2/3 volumes of the aqueous layer of 95% ethanol were added (at −20 °C) to precipitate the nucleic acids. The precipitate was washed with 0.2 M ammonium acetate in 75% ethanol and air-dried for 5 min. The pellet was then resuspended in 500 μL of 65 °C pre-heated H2O and treated with 125 μL (10 mg-mL⁻¹) of RNase at 37 °C for 30 min. The DNA was precipitated, washed, dried, and resuspended in 250 μL of H2O, and quantified in a 1% agarose gel against Lambda DNA external standards. All samples were then brought to a working concentration of 10 ng-μL⁻¹.

**PCR amplification and electrophoresis.** DNA was amplified using eight microsatellite primers selected from an earlier study in *Juglans nigra* L. (Woeste et al., 2002) (Table 2). SSR reaction was conducted according to the protocol of Victory et al. (2006) with some modifications. Amplification reaction was performed in a 15-μL volume containing 1× reaction buffer (TaKaRa Biotechnology, Dalian, China), 30 ng of genomic DNA, 0.2 mM of each dNTP (Promega, Beijing, China), 0.4 mM of each primer, and 0.9 unit of *Taq* DNA polymerase (TaKaRa). For DNA amplifications, a GeneAmp PCR System (9700; PerkinElmer, Waltham, MA) was programmed according to the following profile: an initial 3-min incubation at 94 °C; then 30 cycles of 45 s at 94 °C, 30 s at the annealing temperature (Table 2), and 45 s at 72 °C; and a final incubation at 72 °C for 5 min. After amplification, 5 μL of each sample was loaded and electrophoresed on a 2% horizontal agarose gel to control for positive amplification and to determine the approximate amount of product. Then, 2 μL of each sample was electrophoresed on an 8% polyacrylamide gel containing 1× TBE (Tris–borate–EDTA) buffer. After electrophoresis, the gel was silver-stained with the procedure of Panaud et al. (1996). In all cases, PCR reactions were performed at least twice to ensure that absence of bands was not due to a failed reaction.

**Allele scoring and data analysis.** The fragments amplified by microsatellite primers were scored as alleles on the basis of size in a comparison with external standard using a Gel Doc 1000 image analysis system (Bio-Rad, Hercules, CA) and in reference to the expected size in *J. nigra* (Woeste et al., 2002).

The allele data were entered in the form of single-individual genotypes. The following parameters of genetic variation were assessed for each population: mean number of alleles per locus (A), effective number of alleles (Ae) (Kimura and Crow, 1964), percentage of polymorphic loci (99% criterion) (P), expected heterozygosity (Hₑ) (Nei, 1978), and observed heterozygosity (Hₒ). Departures from the Hardy–Weinberg (H-W) equilibrium were assessed at each locus for every population and per locus across all populations using the *F* statistics of Wright (1978).

The significance of the deviations was evaluated with a χ² test following the method of Workman and Niswander (1970). Genetic structures were further analyzed with Wright’s analysis of hierarchical *F* statistics (Wright, 1978). Gene flow (*Nₘ*) was estimated from *Nₘ* = 0.25(1 − *F*ₚₛₜ)/*F*ₚₛₜ (Bosart and Prowell, 1998; Whitlock and McCauley, 1999). Nei’s (1978) unbiased genetic distances were calculated for all population

**Table 1. Walnut populations surveyed and their ecological and geographical parameters.**

<table>
<thead>
<tr>
<th>Population</th>
<th>Juglans species</th>
<th>Landform feature</th>
<th>Altitude (m)</th>
<th>Longitude (E)</th>
<th>Latitude (N)</th>
<th>Annual rainfall (mm)</th>
<th>Annual avg. temp. (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FY-r</td>
<td><em>J. regia</em></td>
<td>Plateau and hilly</td>
<td>1,414</td>
<td>111°30’</td>
<td>37°24’</td>
<td>438</td>
<td>10.2</td>
</tr>
<tr>
<td>LC-r</td>
<td><em>J. regia</em></td>
<td>Plateau and hilly</td>
<td>1,130</td>
<td>111°30’</td>
<td>33°42’</td>
<td>672</td>
<td>14.7</td>
</tr>
<tr>
<td>HS-r</td>
<td><em>J. regia</em></td>
<td>Alpine and canyon</td>
<td>2,350</td>
<td>102°59’</td>
<td>32°03’</td>
<td>620</td>
<td>9.5</td>
</tr>
<tr>
<td>SJC-r</td>
<td><em>J. regia</em></td>
<td>Alpine and canyon</td>
<td>2,455</td>
<td>99°26’</td>
<td>28°11’</td>
<td>1,250</td>
<td>5.1</td>
</tr>
<tr>
<td>QZ-r</td>
<td><em>J. regia</em></td>
<td>Alpine and canyon</td>
<td>1,922</td>
<td>99°32’</td>
<td>27°34’</td>
<td>1,050</td>
<td>8.2</td>
</tr>
<tr>
<td>LJ-s</td>
<td><em>J. sigillata</em></td>
<td>Alpine and canyon</td>
<td>2,566</td>
<td>99°45’</td>
<td>26°51’</td>
<td>980</td>
<td>11.0</td>
</tr>
<tr>
<td>ML-s</td>
<td><em>J. sigillata</em></td>
<td>Alpine and canyon</td>
<td>3,120</td>
<td>101°15’</td>
<td>27°54’</td>
<td>818</td>
<td>11.4</td>
</tr>
<tr>
<td>YB-s</td>
<td><em>J. sigillata</em></td>
<td>Alpine and canyon</td>
<td>1,230</td>
<td>101°31’</td>
<td>26°54’</td>
<td>850</td>
<td>20.5</td>
</tr>
<tr>
<td>MN-s</td>
<td><em>J. sigillata</em></td>
<td>Alpine and canyon</td>
<td>1,834</td>
<td>102°09’</td>
<td>28°36’</td>
<td>1,113</td>
<td>13.3</td>
</tr>
</tbody>
</table>

“r” refers to *J. regia*, and “s” to *J. sigillata.*
pairs and used to construct a phylogenetic tree (UPGMA). All of the above calculations were performed using POPGENE software, version 1.32 (Yeh et al., 1997). The proportion of null alleles was calculated as \((H_E - H_O)/(1 + H_E)\) (Brookfield, 1996).

The relationship between latitude and the expected heterozygosity was examined via linear regression using Excel (Office 2003; Microsoft Corp., Redmond, WA). A Mantel test of the association between genetic distance and the geographic distance was performed using GenAlEx6 software (Peakall and Smouse, 2006). Geographic distances were calculated according to the longitude and latitude of every sampled location using CoordTrans software (version 2.3; Franson Technology AB, Johanneshov, Sweden).

**Results**

**Allelic variation at microsatellite loci.** Based on the SSR analysis, a total of 73 alleles were scored with the sizes ranging from 153 to 306 bp (Table 2). The number of alleles per locus varied from six (WGA-79) to 12 (WGA-42), with an average of 9.13 alleles per locus. Nineteen of the alleles had an overall frequency greater than 0.1; the remaining 54 alleles were "rare" with a frequency less than 0.1, but only 4 of the 54 rare alleles were unique to a single population (private), meaning that 92.6% of the rare alleles were found in at least two populations.

**Genetic variation within populations.** Genetic diversity parameters based on allelic frequencies are shown in Table 3. In individual population, \(A\) varied from 3.25 to 6.00, with an average of 4.60, while \(A_E\) varied from 1.75 to 3.35, with an average of 2.39. \(H_O\) ranged from 0.281 to 0.492, with an average of 0.414. \(H_E\) was 0.525 and varied from 0.389 to 0.687. For entire populations, \(H_O\) and \(H_E\) values were 0.411 and 0.632, respectively, with \(A_E = 2.91\).

A comparison of the genetic diversity was performed among the nine populations. The highest level of diversity existed in population QZ-r with \(H_E\) values of 0.687, while the lowest values existed in population HS-r, with \(H_E = 0.389\). The correlation coefficient between the latitude and the expected heterozygosity in all the populations was significantly negative, \(-0.577 (P = 0.011)\).

**Genetic structure.** Genetic analyses revealed a high level of differentiation within the populations. The coefficient of hierarchical \(F_{ST}\) (Table 4), estimated according to Wright (Office 2003; Microsoft Corp., Redmond, WA). A Mantel test of the association between genetic distance and the geographic distance was performed using GenAlEx6 software (Peakall and Smouse, 2006). Geographic distances were calculated according to the longitude and latitude of every sampled location using CoordTrans software (version 2.3; Franson Technology AB, Johanneshov, Sweden).

**Table 2.** Characterization of the eight SSR markers based on nine walnut populations.

| Locus | Repeat type | Alleles (no.) \(^a\) | Ranges of allele sizes (bp) \(^b\) | Annealing temp. (°C) | Primer sequence (5’ → 3’)
|-------|-------------|---------------------|-------------------------------|---------------------|---------------------|
| WGA-1 | (AG)_6C(AG)_3 | 10 | 189–221 | 60 | F : ATTGGAGAGGAGGGAATTAG
|       |             |       | 189–207 | R : CGCCGCACATCAATGCACAC
| WGA-42 | (GA)_14 | 12 | 210–260 | 55 | F : GTGGGTTCGACCGTGAC
|       |             |       | 210–260 | R : AACTTTGCACCACATCCACA
| WGA-70 | (GA)_14 | 10 | 161–177 | 55 | F : TGTAATTGGGGAATGTTCAA
|       |             |       | 159–187 | R : TGGGAGACACAATGATCGA
| WGA-71 | (GA)_d(G)_12 | 8 | 212–226 | 57 | F : ACCCGAGAGATTCTGAG
|       |             |       | 212–228 | R : GGAGCCGATCTCCTCCTCTCT
| WGA-72 | (CT)_14 | 8 | 153–173 | 58 | F : AAAAACTAATTTTTGCA
|       |             |       | 153–173 | R : ACCATCCAATATGCTTCA
| WGA-76 | (GA)_12 | 10 | 242–306 | 55 | F : AGGGCCTCCCTCTTTAGG
|       |             |       | 246–302 | R : CAGTCATCCCCTCCTT
| WGA-79 | (GA)_12 | 6 | 204–220 | 55 | F : CACTTGCGAGCTGCA
|       |             |       | 204–220 | R : TTGGAGCTGAGAC
| WGA-89 | (TG)_d(GA)_21 | 9 | 216–240 | 56 | F : ACCCATCTTCCGAC
|       |             |       | 214–232 | R : TGCCATAGCCATAC

\(^a\)Number of alleles observed for each marker.

\(^b\)Top ranges are for *J. regia*; bottom ranges are for *J. sigillata.*
(1978), ranged from 0.140 for locus WGA-70 to 0.248 for locus WGA-79, with the average value equaling 0.186. This indicates that, on average, 18.6% of the total genetic diversity present is present within a given population. All loci possessed a close FST value except locus WGA-79. The overall gene flow (N_m) among populations was 1.097, which gave an estimate of the average number of migrants between all studied populations per generation. The observed value indicated that gene exchange between populations is low.

**Genotypic structure.** In individual populations, tests for the departure from H-W equilibrium showed significant deviations for at least one locus in every population. The χ² test (P < 0.05) indicated that significant departures from the H-W equilibrium appeared in many cases (≥60%). The deviations were primarily attributed to a deficit of heterozygotes.

F_{IT} is the overall inbreeding coefficient of an individual relative to the whole set of populations, while F_{IS} is the inbreeding coefficient of an individual relative to its own population. The mean F_{IT} value of 0.348 indicates that, overall, nine populations have a deficiency of heterozygotes. Within populations, all loci but one (WGA-42, F_{IS} = -0.106) contained a deficiency of heterozygotes (F_{IS} ranging from 0.047 to 0.417), with an average F_{IS} value of 0.199, which is significantly different from zero (Table 4). This suggested that there was a regular tendency toward heterozygote deficiency and indicated the presence of inbreeding within the populations. In addition, the proportion of null alleles was estimated, and they were found to occur at a moderate frequency: for eight loci, the estimated values varied from 0.04 to 0.22 (Table 4) and the mean value equaled 0.13.

**Genetic relationships.** Genetic distances were calculated for each pair of populations to estimate the extent of their divergence (Table 5). In the nine populations, the average genetic distance among populations equaled 0.311. The lowest genetic distance (0.049) was found between populations YB-s and MN-s, and the greatest genetic distance (0.845) was found between populations LC-r and MN-s. In addition, The average genetic distances between *J. regia* populations is 0.222, that between *J. sigillata* populations is 0.122, and that for *J. regia*-to-*J. sigillata* is 0.423.

The UPGMA cluster analyses based on Nei’s unbiased genetic distances was performed to further show the genetic relationships among the populations (Fig. 2). The dendrogram separates the nine populations into two main groups. The first group consists of populations SJC-r, QZ-r, LC-r, MN-s, YB-s, and ML-s. In this group, population QZ-r differs considerably from the others. The second group consists of populations FY-r, LC-r, and HS-r. In this group, populations FY-r and LC-r cluster together while population HS-r was more distant. A Mantel test with 1000 random permutations revealed a significant correlation between pairwise genetic distance and geographic distances among the nine sampled populations (R² = 0.354, P < 0.005).

**Discussion**

We observed a moderate genetic diversity in nine walnut populations in central and southwestern China based on 73 loci alleles. The expected heterozygosity (H_E) of the populations ranged from 0.389 to 0.687, and their mean H_E was 0.525. In comparison, the average H_E of those nine populations was lower than that observed in other *Juglans* species, e.g., in *J. nigra* (mean H_E = 0.807) (Vicent et al., 2006) and *Juglans mandshurica* Max. (mean H_E = 0.806) (W.N. Bai, unpublished data), and in some temperate forest tree species (Derory et al.,

<table>
<thead>
<tr>
<th>Loci</th>
<th>F_{IS}</th>
<th>F_{IT}</th>
<th>F_{ST}</th>
<th>N_m</th>
<th>H_O</th>
<th>H_E</th>
<th>Proportion of null alleles</th>
</tr>
</thead>
<tbody>
<tr>
<td>WGA-01</td>
<td>0.406</td>
<td>0.522</td>
<td>0.196</td>
<td>1.029</td>
<td>0.350</td>
<td>0.731</td>
<td>0.22</td>
</tr>
<tr>
<td>WGA-42</td>
<td>-0.106</td>
<td>0.103</td>
<td>0.189</td>
<td>1.071</td>
<td>0.578</td>
<td>0.648</td>
<td>0.04</td>
</tr>
<tr>
<td>WGA-70</td>
<td>0.417</td>
<td>0.499</td>
<td>0.140</td>
<td>1.532</td>
<td>0.372</td>
<td>0.740</td>
<td>0.21</td>
</tr>
<tr>
<td>WGA-71</td>
<td>0.089</td>
<td>0.278</td>
<td>0.208</td>
<td>0.952</td>
<td>0.382</td>
<td>0.540</td>
<td>0.10</td>
</tr>
<tr>
<td>WGA-72</td>
<td>0.198</td>
<td>0.315</td>
<td>0.146</td>
<td>1.469</td>
<td>0.345</td>
<td>0.498</td>
<td>0.10</td>
</tr>
<tr>
<td>WGA-76</td>
<td>0.327</td>
<td>0.422</td>
<td>0.141</td>
<td>1.525</td>
<td>0.278</td>
<td>0.480</td>
<td>0.14</td>
</tr>
<tr>
<td>WGA-79</td>
<td>0.047</td>
<td>0.283</td>
<td>0.248</td>
<td>0.761</td>
<td>0.489</td>
<td>0.679</td>
<td>0.11</td>
</tr>
<tr>
<td>WGA-89</td>
<td>0.152</td>
<td>0.322</td>
<td>0.201</td>
<td>0.993</td>
<td>0.491</td>
<td>0.729</td>
<td>0.14</td>
</tr>
<tr>
<td>Mean</td>
<td>0.199</td>
<td>0.348</td>
<td>0.186</td>
<td>1.097</td>
<td>0.411</td>
<td>0.631</td>
<td>0.13</td>
</tr>
</tbody>
</table>

*Inbreeding coefficient at the population level.

*Inbreeding coefficient at the sample level.

*Proportion of differentiation among populations.

*Proportion estimated from N_m = 0.25 (1 – F_{ST})/F_{ST}.

*Proportion of null alleles = (H_E – H_O)/(1 + H_E).
2002; Heuertz et al., 2004; Jones et al., 2002; Muir et al., 2004). As an economically important species, walnut plants were inevitably disturbed by human activities, such as deforestation and selection (Lee and Lee, 1997), particularly over the past 100 years. Consequently, many walnut genetic resources have been fragmented or lost entirely. Those that remain are located only along canyons and besides streams and rivers. The *Juglans* species are monoecious woody plants with separate male and female inflorescences and a wind-pollinated out-crossing mating system (Gleeson, 1982); its pollen has at most only 300 m of transmitting distance by wind (Han et al., 1996). Therefore, fragmentation can induce a certain level of genetic drift. Moreover, the *J. regia* and *J. sigillata* species have higher percentages of self-pollination and parthenogenesis (Fan and Xi, 2002; Wang, D.L., et al., 1999; Wang, G.A., et al., 2003). Self-pollination is generally considered a more important factor of a population’s genetic diversity level (Charlesworth and Charlesworth, 1987). Therefore, human disturbance, mating system, and self-pollination could be the major causes for the lower level of genetic diversity in the nine walnut populations compared with other *Juglans* species and temperate forest tree species.

The genetic analyses of the nine populations showed a high level of differentiation among populations. The coefficient of hierarchical $F_{ST}$ among populations ranged from 0.140 to 0.248. Its mean $F_{ST}$ was 0.186, indicating that 18.6% of the total genetic diversity existed among populations and 81.4% within populations. This result was consistent with the variation models reported in anemophilous pines (Yeh and Layton, 1997). The high level of interpopulation differentiation might result from the differentiation of habitats, for instance, with respect to climate, temperature, annual rainfall, or landform features. Southwestern China is located in the center area of Himalayan–Hengduan Mountains and is characterized by rolling hills, high peaks, and criss-crossing valleys and canyons. It contains various geographical features and microenvironments. Heterogeneous environments promote maintenance of high genetic diversity due to developmental homeostasis (Lerner, 1954). Various climate and ecological conditions in these areas have led to a high species diversity and genetic diversity (Peng et al., 2005). The Himalayan–Hengduan Mountains and surrounding areas are considered to be a new concentration area of temperate and alpine plants due to their geographic and environmental characteristics (Ming et al., 2006; Sun, 2002). Suitable habitats and little disturbance by human activities might be the main reasons for the high diversity within these walnut populations.

The genetic diversity of the nine populations presented a geographic gradient. Six populations distributed in the southwest mountains of China (LJ-s, ML-s, YB-s, MN-s, SJC-r, and QZ-r) had a higher level of genetic diversity within populations than those in the northward areas. A negative correlation between latitude and $H_E$ in the nine populations was detected. This result seems to be consistent with the viewpoint proposed by Lu and Zhang (1990) that walnuts were indigenous to the tropical mountains in southwestern China, but—because of the seasonal drought that persists there—they evolved their sustainability from a humid environment in torrid zones to a drier environment. Populations of SJC-r, QZ-r, and LJ-s, distributed in the central area of the southwest mountains of China (in northwestern Yunnan Province), had a higher genetic diversity than other populations, and those three populations captured more than 95% of the genetic variation presented in all populations under investigation. Therefore, we could consider this region as an important hotspot of walnut diversity.

L.A. Dode considered *J. sigillata* as a different species from *J. regia* mainly based on leaf morphological difference. For instance, *J. regia* grows elliptic to ovate leaflets, has 5–9 leaflets in a compound leaf, and has less than 15 pairs of lateral veins in a leaflet, whereas *J. sigillata* grows elliptic to

### Table 5. Nei’s unbiased measures of genetic identities (above the diagonal) and genetic distances (below the diagonal) among nine *Juglans* populations (Nei, 1978).

<table>
<thead>
<tr>
<th>Population</th>
<th>FY-r</th>
<th>LC-r</th>
<th>HS-r</th>
<th>SJC-r</th>
<th>QZ-r</th>
<th>LJ-s</th>
<th>ML-s</th>
<th>YB-s</th>
<th>MN-s</th>
</tr>
</thead>
<tbody>
<tr>
<td>FY-r</td>
<td>****</td>
<td>0.937</td>
<td>0.890</td>
<td>0.819</td>
<td>0.720</td>
<td>0.624</td>
<td>0.728</td>
<td>0.515</td>
<td>0.495</td>
</tr>
<tr>
<td>LC-r</td>
<td>0.065***</td>
<td>0.791</td>
<td>0.759</td>
<td>0.617</td>
<td>0.580</td>
<td>0.693</td>
<td>0.463</td>
<td>0.430</td>
<td></td>
</tr>
<tr>
<td>HS-r</td>
<td>0.116</td>
<td>0.234</td>
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<td>0.845</td>
<td>0.780</td>
<td>0.664</td>
<td>0.738</td>
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<tr>
<td>SJC-r</td>
<td>0.200</td>
<td>0.276</td>
<td>0.169</td>
<td>****</td>
<td>0.902</td>
<td>0.836</td>
<td>0.926</td>
<td>0.793</td>
<td>0.744</td>
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<td>QZ-r</td>
<td>0.328</td>
<td>0.482</td>
<td>0.248</td>
<td>0.103</td>
<td>****</td>
<td>0.788</td>
<td>0.807</td>
<td>0.759</td>
<td>0.734</td>
</tr>
<tr>
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<td>0.544</td>
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<td>0.179</td>
<td>0.238</td>
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<td>0.894</td>
<td>0.856</td>
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<tr>
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<td>0.215</td>
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<tr>
<td>YB-s</td>
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<td>0.515</td>
<td>0.232</td>
<td>0.276</td>
<td>0.156</td>
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<tr>
<td>MN-s</td>
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<td>0.296</td>
<td>0.310</td>
<td>0.151</td>
<td>0.152</td>
<td>0.049</td>
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**Fig. 2.** UPGMA dendrogram of nine *Juglans* populations in central and southwestern China based on Nei’s unbiased genetic distances. The letter of “s” at the end of population names refers to *J. sigillata*, and “r” to *J. regia*. 
ovate–lanceolate leaflets, has 9–11 leaflets in a compound leaf, and has 15–33 pairs of lateral veins in a leaflet (Kuang and Lu, 1979). However, plant morphology, particularly leaf morphology, is influenced strongly by the environment. The genetic diversity of these two species based on isozymes markers showed no significant difference between them (Yang and Xi, 1989). Kuang and Lu (1979) classified J. regia and J. sigillata into the section Juglans under the Juglans genus on the basis of plant morphologic and geographic characteristics. A crossing result of J. regia × J. sigillata showed that their filial generation was progenitive (Fan and Xi, 2002; Fan et al., 2005). Some offspring of J. regia generated by cross-fertilization with J. sigillata or self-fertilization in the scope of J. regia distribution exhibited morphological characteristics of J. sigillata (Fan et al., 2005). In this research, UPGMA clustering showed that the nine populations considered were congregated according to their geographic distribution rather than the definition of traditional taxonomy, suggesting that the genetic differentiation of J. regia and J. sigillata was small. A Mantel test with 1000 random permutations revealed a significant correlation between pairwise genetic distance and geographic distances among the nine sampled populations ($R^2 = 0.354, P < 0.005$). This result supports the viewpoint that J. regia and J. sigillata are different ecological types of the same species. Their differentiation might result from geographic migration and reproductive isolation.

**Literature Cited**


