Evaluation of the genetic variability among a wild *Peganum harmala* L. populations with RAPD-PCR

**Abstract**

Forty wild individuals of wild Peganum harmala (P. harmala) were collected from four sites in Ma’an Governorate and were assessed for genetic variability by the method of random amplified polymorphic DNA (RAPD) PCR. Four primers produced 43 amplified bands and 23 of them were polymorphic. Miniature degree of genetic diversity was detected among studied population of wild P. harmala. Cluster analysis by the un weighted pair-group method with arithmetic averages (UPGMA) divided the wild populations into three main clusters, and one individual was identified in a separated group. It is apparent from this study that individuals of P. harmala species were more closely related. The RAPD markers detected sufficient degree of polymorphism to differentiate among the studied populations, making this technique valuable for wild types management and selectivity for breeding program.

**Keywords:** genetic diversity, jordan, polymorphism, RAPD

**Introduction**

*P. harmala* L. species is well known for its medicinal and traditional uses in folk medicine. Its extract was exposed against two cancer cells line, MDA-MB-231 and Mcf-7, and that showed cell growth inhibition and in higher concentration/longer time, complete cell death occurred Tehrani et al. Assess the diversity of plant genetic resources (germplasm), that have a great chance to occurrence of useful genes, is the first essential prerequisite to plan for the conservation, sustainable use and domestication of them Khoshokhan et al. Molecular markers such as random amplified polymorphic DNA (RAPD) Williams et al. may facilitate more effective diversity analysis of several plants. RAPD-PCR analysis has been used to evaluate the levels of genetic variation among diverse wild cranberry clones, genetic diversity of climbing species of palms Sramah et al., genetic diversity of a medicinal and aromatic plant *Teucrium polium* Bouilla et al. genetic variability among five caraway populations Laribi et al., in *Draba dorneri* vascular plant Catané et al. among cowpea genotypes Patil et al., genetic diversity and related relationships among 22 landrace of *Ficus carica* L. sativa Ali-Shateh et al. and for *Withania somnifera* (L.) Sahu et al. Variability within and among genus, species and families are based on the genetic makeup, the natural selection, human interfering and nature calamities. In the study carried out on analysis of molecular variance of RAPD data of *Aegilops geniculata* populations and *Triticum durum* found that major proportion (80%) of the total variation was existed within populations; in contrast, 20% of the variation was come from among populations Mahjoub et al. However, most RAPD loci are supposed to have only two alleles and separate as dominant markers, foremost to under evaluation of the genetic variation. Little is known about *P. harmala* genetic diversity either in the field or by using RAPD and most of studied concentrated on its pharmacological effects on certain diseases. Moreover, several studies were conducted for evaluation the genetic diversity within and among population of wild types of medicinal plants such as *Teucrium polium* populations, *Lupinus pilosus* L., cultivated tomato, *Artemisia judaica* and estimation of genetic variation within yellow asphodel individuals AlRawashdeh et al. Therefore, the aim of this study was to assess the level of genetic variability and differentiation among population of *Peganum* based on RAPD markers.

**Material and methods**

**Plant material and DNA isolation**

For this investigation, fresh leaves of a total 40 single individuals (10 samples per population) representing four populations were collected in April, 2015 from Ma’an Governorate. DNA was extracted from young leaves of *P. harmala* using CTAB method. Total genomic DNA was extracted from 5g of leaf materials from each sample according to Bader et al. Leaf samples were kept in a fixing solution (95% ethanol) for one hour, kept in oven over night then dried and homogenized in a mortar and pestle. Twenty mg of grinding tissues placed in 1.5ml tubes and mixed with 600µl of freshly prepared and preheated 2x CTAB solution in 2ml tubes then placed at 65°C for 60min. The mixture was added to 600µl of chloroform/isooamyl alcohol (24:1), vortexed for few seconds, and then centrifuged at 14,000g for 15min. DNA was then precipitated with an equal volume of pre cooled ethanol (–20°C), then centrifuged for 15min at 14000g. The purity was measured by the ratio of the absorbance at 260 and 280nm. 40ng of template DNA, 0.25mM dNTPs (Promega), 0.001% (w/v) gelatin), and preheated 2x CTAB solution in 2ml tubes then placed at 65°C for 60min. The mixture was added to 600µl of chloroform/isooamyl alcohol (24:1), vortexed for few seconds, and then centrifuged at 14,000g for 15min. DNA was then precipitated with an equal volume of pre cooled ethanol (–20°C), then centrifuged for 15min at 14000g. The solution was poured in tubes and left to dry, then 600µl of cooled (4°C)70% ethanol was added to the solution and placed in the refrigerator (–20°C) overnight. Next day, ethanol was poured from the dried tubes and 100µl of TE buffer (10mM Tris-HCl, 1mMEDTA, pH 8.0) was added and the whole mixture were placed at 65°C for 60min. Four µl of RNAase (10mg/ml) were added per tube and left for 60min at 37°C. The concentration of DNA was estimated using S2100 UV/VIS DIODE-Array-Spectrophotometer, machine Version 1.7. The purity was measured by the ratio of the absorbance at 260 and 280nm.

**PCR amplification**

PCR reaction was performed as described by Williams et al. with arbitrary oligonucleotides from Operon Technologies (Alameda, Calif.). The final volume of 25µl contained 10 x buffer (50mM KCI, 10 mM Tris-HCl pH 8.3, 1.5mM MgCl2, and 0.001% (w/v) gelatin), 20ng of template DNA, 0.25mM dNTPs (Promega), 0.025mM
of primers, 1.5mM MgCl₂ and 1U of Taq polymerase. Reaction mixtures were amplified in PCT-200 programmable thermo cycler (MJ Research Inc., USA), each reaction was performed using an initial step of 94°C followed by 44 cycles of a denaturation step for 1min at 94°C, followed by an annealing step for 1min at 36°C and an extension step for 2min at 72°C, followed by a further extension step for 5min at 72°C. After the final cycle the samples were holding at 4°C for analysis. Amplified fragments 10µl RAPD-PCR along with 100bp DNA ladder were resolved by electrophoresis on 1.4% a garose gel mixed with 2µl of ethidium bromide in 1X tris-borate-EDTA (TBE) buffer. Primer screening: twenty decamer primers from Operon, Advanced Biotechnologies Inc. Almeda, USA were firstly screened using one individual DNA sample represent a single plant to define the suitability of each primer for this study. After introductory testing on a few samples, four primers were selected for further analysis based on their ability to discover distinguish obviously resolved and bands within the population. The primers generating no and weak amplified fragments were dismissed.

Data analysis

The RAPD fragments were scored as present (1) or absent (0) for each of the markers to approximate the similarity among all populations and then transferred into a binary matrix. Each band was assumed to represent a single locus. All gels were scored for both monomorphic and polymorphic amplified fragments. The matrix of similarity was calculated using the Jaccards’ coefficient and the dendrogram obtained by clustering according to the Unweighted Pair-Group Method with arithmetic averages (UPGMA) using SPSS, V. (11.0), software. Genetic diversity within populations was estimated using the percentage of polymorphic bands [number of polymorphic bands/ number of total bands] x100).

Results

The P. harmala populations were collected to assess the genetic variability at the DNA level. Random amplified polymorphic DNA (RAPD) analysis has been demonstrated as a good tool for this aim. Initially, RAPD analysis was passed out using asset of more than 25 random primers of Opern Kits A, B, F, W and Z, for amplification during PCR. Among them, four RAPD primers produced polymorphic amplified fragments except one primer did not yield any polymorphic bands (Table 1). A representative banding pattern is shown in (Figure 1A), which demonstrates a total of 43 diverse bands from four populations of P. harmala. The primer (ACAACGCCTC) and (CCGTGACTCA) showed the highest percentage of polymorphism (75%) and (63%), respectively (Table 1). The primer P1 shows the highest number (8) of polymorphic bands followed by P2, which shows 6 bands. Most of similarity among testing samples were ranged between 1.00-0.58 (Table 2). The dendrograms depended on the Jaccard’s coefficients values (Figure 1B) and is investigative of low to medium level of polymorphism among the four populations at the DNA level. The samples of P. harmala were grouped into two major clusters. The first cluster (A) was divided into 5 sub-clusters (A1-A5) (Figure 2). The first sub-cluster was then divided into three main sub-sub clusters. The first sub-sub cluster contained 5 samples from Alshoubak, one sample of Adruh, one Wadimusa sample (Figure 2). Among these samples (0.93) similarity was formed between five samples 33, 36 Alshoubak regions, 31, 33 Alshoubak site, Adruh sample 13 and Wadimusa sample 23 had 1.00 similarities. Also two Adruh samples (14 and 15) showed 1.00 similarity values (Figure 2). The second sub-sub cluster consists of two samples of Alshoubak, one samples of Ma’an and one sample Wadimusa. The third sub-sub cluster formed 4 Wadimusa sample, 8 Adruh and 3 of Ma’an site. The second sub-sub cluster appeared two Ma’an samples, one of Adruh individuals and one sample Wadimusa. The third sub-sub cluster integrated three Wadimusa. The forth sub-sub cluster enclosed three of Ma’an individuals and two of Alshoubak samples. The fifth sub-sub cluster had only one samples of Alshoubak region. The second main cluster formed a separate cluster and showed only one individual (no.2) collected from Ma’an site (Figure 2), formed diverse cluster compared to the rest of individuals. Variability among populations was existed through distribution the sites samples overall the clusters. Variation within sites was observed but with a low level.

### Table 1 Primer sequence and monomorphic and polymerphic bands obtained by using four RAPD primers

<table>
<thead>
<tr>
<th>Primer name</th>
<th>Primer sequence</th>
<th>Band total</th>
<th>Monomorphic</th>
<th>Polymorphic</th>
<th>% Polymorphism</th>
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<tr>
<td>OPA-14</td>
<td>TCTGTGCTGG</td>
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<td>4</td>
<td>4</td>
<td>50</td>
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<tr>
<td>OPM-02</td>
<td>ACAACGCCTC</td>
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<td>2</td>
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<tr>
<td>OPM-07</td>
<td>CCGTGACTCA</td>
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<td>3</td>
<td>5</td>
<td>63</td>
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<tr>
<td>OPM-06</td>
<td>CTGGGCAACT</td>
<td>6</td>
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</table>

Figure 1 A representative RAPD banding profiling using OPM-02primer among different populations of Peganum harmala species. M: 100bp DNA ladder (Gene Direx).

conducted at the three hierarchical levels (population, ecological, and ploidal group) showed that most of the total variation was found among individuals within populations or within populations in their corresponding group. This suggests that mating occurs mainly among individuals within a subpopulation, thus favoring the divergence between populations. Sarmah et al. reported that the RAPD markers grouped Rattan species collected in their specific species level with some exceptions and discovered intra-specific diversity. Our results showed that the similarity represented by the polymorphism of RAPD bands between individuals from the four sites of Ma’an governorate is approximately 63% (sample 1, 3). In spite of the small size of the population, which usually reveals lower levels of genetic diversity, our findings based on the RAPD technique showed that the genetic diversity of this species is not low. However, this study showed that the groupings of populations were mostly in agreement with their geographical localities assigned the low level of genetic variation within populations. The occurrences of polymorphism within or among studied P. harmala populations point to potential of using RAPDs in category the genotypes. This study provides leadership for the prospective analysis of genetic diversity in Peganum species with even more consistent molecular markers such as AFLP and ISSR. To conclude, this is the first report on the evaluation of genetic variability and population discrimination analysis in four P. harmala species, mainly originating from Jordan using RAPD. We successfully provided deep insights in the genetic background of the studied populations. Considerable genetic variability has been detected at low level within or between populations. Further studies are needed to cover all science areas associated with this specie in the future.

Acknowledgements

None.

Conflict of interest

The author declares no conflict of interest.

References


