PAPER DETAILS

TITLE: Intra-Genetic Variation within Olive Cultivar 'Nabali' in Palestine by Microsatellite and

Random Amplified Polymorphic DNA

AUTHORS: Aziz SALAMEH, Soha GEZAEIL, Alaa LAHLOOH, Dina ARAFAT

PAGES: 30-36

ORIGINAL PDF URL: https://dergipark.org.tr/tr/download/article-file/473975

ANADOLU, J. of AARI ISSN: 1300 - 0225 28 (1) 2018, 30 - 36 MFAL

Intra-Genetic Variation within Olive Cultivar 'Nabali' in Palestine by Microsatellite and Random Amplified Polymorphic DNA

Aziz SALAMEH* Soha GEZAEIL Alaa LAHLOOH Dina ARAFAT

Department of Plant production and Protection, National Agricultural Research Center, PALESTINE

* Corresponding author (Sorumlu yazar): azbarghouthi@hotmail.com

Received (Geliş tarihi): 10.07.2017 Accepted (Kabul tarihi): 30.01.2018

ABSTRACT: Over seventy olive trees (Nabali) from different regions in Palestine were used in this study. Intra-genetic variation within different olive Nabali variants were approved by SSR and RAPD markers. Four SSRs bands were monomorphic revealing a true-to-genotype of Nabali cultivar. Ten RAPD markers produced 60 reproducible bands with an average of 6 bands / marker. Only 24 were polymorphic. The percentage of polymorphic bands was 38% which is relatively high. Similarity matrix for studied populations ranged from moderate (0.610) for Jalkamous and Karawa Bani Zaid (2) to highly genetic similarity or even identity (1.000) in some cases as Aqraba and Aseerah (N). The interaction between different variants trends to be high. The effect of geographic location was absent in this study and has no significant contribution. Dendogram based on Jaccards cofficent revealed three main clusters, the biggest group consisted of the majority of variants including Bieta, Karawa Bani Zaid (1), Salfeet, Salfeet (h), Aqraba, Aseereh (N), Jalkamous, and Alaroub. Second group consisted of two variants Alaar and Nahaleen. The third group is containing only Karawa Bani Zaid (2). The relative high polymorphic bands of RAPD markers (40%) and moderate genetic similarity among different Nabali variants suggested attribution of genetic background. Selection for new traits within Nabali is suggested.

Keywords: Olive, Intra-Genetic Variation, SSR, RAPD.

INTRODUCTION

Olive (Olea europaea L.) is as an important oilproducing crop in the Mediterranean region whose domestication occurred during the Chololithic period (5700-5500 years BC) in the Near-East (Zohary and Hopf, 1994). In Palestine, olive represents the most important fruit trees and growing for hundreds of years. Nabali is the most predominant cultivar and met around 75% of the total planted olive cultivars in Palestine. Nabali is adapted well with the environmental conditions with high productivity of olive oil (22-28%). The chemical parameters of its oil are met the International Olive Council (I.O.C.) standards for extra virgin olive oil except Δ -7stigmastenolw which shows values higher than 0.5% (Qutub et al., 2010). Nabali is susceptible to olive leaf spot disease (Spilocaea oleaginea) and showed alternating bearing.

A high genetic diversity level and the presence of homonyms and synonyms cases observed in olive germplasm (Gomes et al., 2012), therefore, an efficient and rapid discriminatory methods are urgent. Molecular markers have been used successfully in olive characterization (Angiolillo et al., 1999; Khadari et al., 2003; Abdel hamid et al., 2012). Genetic variability among olive cultivars within a country was higher than olive cultivars from different countries (Belaj et al., 2003; Owen et al., 2005). Variability of 27 clones of the Portuguese olive cultivar was investigated by Random Amplified Polymorphic DNA (RAPD), Simple Sequence Repeat (SSR), and Inter Simple Sequence Repeat (ISSR) markers (Gomes et al., 2008), and the intra-variation within an olive cultivar has been approved (Muzzalupo et al., 2010; Ipek et al., 2012).

Though RAPD provides an inexpensive and reliable method for routine screening of large number of cultivars for olive germplasm collection (Belaj *et al.* 2004) and detecting genetic similarities in olive (Belaj *et al.* 2003). It's reproducibility still under question. Therefore, providing RAPD profile with other molecular technique could be a good option to support the obtained results. Recently, SSRs have become one of the most useful molecular markers in plant breeding, cultivar fingerprinting, and genome mapping genetic (Rallo *et al.*, 2003).

In Palestine, genetic variation for Nabali has been studied and approved previously on eight olive trees from two sites only (Wiesman *et al.*, 1998). No other studies were done to determine the intragenetic variation within Nabali cultivar by using a large number of trees on large scale of area. Identification of genetic variation within Nabali could be a useful tool for breeding program in future in Palestine.

The objectives of this work are;

To identify by RAPD and SSR markers DNA fingerprints of Nabali cultivar, figure out the genetic relationship between them, and define the clones within the studied cultivar.

MATERIALS AND METHODS

Plant material consisted of fresh leaves of 72 trees tagged as Nabali cultivar. Trees were selected from thirty two villages located in eight governorates in Palestine (Table 1). The villages and trees were

selected according to the extension agent's recommendations and knowledge. The samples were stored in paper bags in the field and at cold temperature in the lab until DNA extraction.

Genomic DNA was isolated from 100 mg leaf by ground into fine powder using pistil and mortar in the presence of liquid nitrogen. The leaf powder of each individual sample was then subjected to DNA extraction using DNeasy Plant Mini Kit (Qiagen). DNA quality was determined by visualization on 0.7% ethidium bromide agarose gel. Concentration of DNA for all samples was measured by S-30 spectrophotometer (BOECO, Germany) and uniformed to 20 ng/ μ l. Resulting DNA solutions were stored at -20 °C.

Amplification and genotyping four RAPD primers (opx-09, opj-05, opi-12, and opx-03) were tested (Table 2). All the 72 samples were amplified and analyzed with four RAPD primers. PCRs were performed in a total reaction mixture of 25 µL volumes containing 10 mM Tris-HCl, pH 8.2,50 mM KCl, 1.5 mM MgCl2, 200 mM primer, 0.2 unit of Taq DNA Polymerase (sigma) and 20 ng DNA. Amplifications were performed in PTC 100 thermocycler (MJ Research), initial denaturation at 94 °C for 1min, then 45 cycles at 92 °C for 1 min, 36 °C for 1 min, and 72 °C for 2 min and finally one cycle for 8 min at 72 C for elongation. Amplified products were fractionated bv electrophoresis in 2% ethidium bromide Agarose gels ($1 \times TAE$ buffer) and photographed.

| No | Governorate | Villages |
|----|-------------|--|
| 1 | Jenin | Biet Qad, dier Abu Daaf, Jalkamous, Maithloun, Sier, Alyamoun, Bourqeen, Yaabed, Arabeh, KufarPaa, and Markaa |
| | | Kutet Kaa, aliu Matkaa |
| 2 | Qalqelia | Jayous, BaqatAlhatab, and koufer Qadoum |
| 3 | Nablus | Qabalan, Aqraba, Biet Foureek, and Aseerah(N) |
| 4 | Tulkarem | Dier Ghosoun, Alaar, and Ras Rouman |
| 5 | Salfeet | Alwaraam, Salfeet ¹ , and Dierstya |
| 6 | Ramallah | Karawa Bani Zaid ² , Dier Gassaneh, and Biet Laqya |
| 7 | Bietlahem | Biet Jala, Nahaleen, and Tkouaa |
| 8 | Hebron | Alaroub and Dora, |

Table 1. The Governorates and the villages where the leaves of Nabali olive trees were collected.

¹: In salfeet village two different olive variants were found. Thereafter in the text, salfeet and salfeet (h) used to distinguish between both.

²: In Karawa Bani Zaid village two different olive variants were found. Thereafter in the text, Karawa Bani Zaid (1), and Karawa Bani Zaid (2), used to distinguish between both.

Later on, in order to minimize the size of population (72 samples), only samples which showed at least a different RAPD profile bands or more were selected for regenotyping again with ten RAPD and six SSR primers in order to assess the genetic diversity (Table 2). Out of seventy two, eleven samples of Nabali were selected for second cycle of genotyping. SSR primers were described and used by Cipriani et al. (2002). SSR primers were amplified by PTC 100 thermocycler (MJ Research), initial denaturation at 94 °C for 2 min, then 35 cycles at 92 °C for 45s., 57 °C for 45s., and 72 °C for 45s., and finally one cycle for 8 min at 72 °C for elongation. Amplified SSR bands were fractionated by electrophoresis in 4% ethidium bromide garose gels (1×TAE buffer).

DATA analysis

Each SSR and RAPD fragments was treated as a unit character and was scored presence or absence of the band (1 or 0). The 1/0 matrix was prepared for all fragments scored and the data were used to generate Jaccard's similarity coefficients (1908) for RAPD bands depending on the following formula:

Table 2. RAPD and Olive SSR primers characterized in the study.

Sij = a / (a+b+c)

Where Sij: standard Jaccard between two individuals i and j;

- a = bands shared by both individuals;
- b = bands present in i but not in j; and
- c = bands present in j but not in I

Jaccard's coefficients were subjected to unweighted pair-group method using arithmetical averages (UPGMA) to generate a dendrogram using linkage procedure. The RAPD data were analyzed using fingerprint analysis missing DATA (FAMD version1.25) (Schlüter and Harris, 2006).

RESULTS

The aim of this work is determining the genetic diversity within Nabali cultivar in Palestine. Seventy two variants of Nabali from different areas were scored with four RAPD markers. All produced RAPD profiles were selected and analysed in next step with SSR and RAPD markers. Eleven Nabali variants were selected as a result of first RAPD running.

| Type of primers | primer | Primer sequences | Size of sequenced alleles | | |
|-----------------|------------|-------------------------------------|---------------------------|--|--|
| | - | (5' to 3') | (bp) | | |
| | OPA-19 | CAA ACG TCG G | | | |
| | OPK-16 | GAG CGT CGS A | | | |
| | OPX 09 | GGT CTG GTT G | | | |
| | OPJ-06 | CTC AGT CGC A | | | |
| RAPD | OPZ-11 | GGG AAT TCG G | | | |
| | OPF-06 | CCA GGA GGA C | | | |
| | OPJ 05 | CTC CAT GGG G | | | |
| | OPZ-07 | TCG TTC CGC A | | | |
| | OPI 12 | AGA GGG CAC A | | | |
| | OPX 03 | TGG CGC AGT G | | | |
| | | TCA GTT TGT TGC CTT TAG TGG A | 172 | | |
| | UD099-006 | TTG TAA TAT GCC ATG TAA CTC GAT | 1/2 | | |
| | | AAA AAC ACA ACC CGT GCA AT | 159 | | |
| | UD099-008 | AAA TTC CTC CAA GCC GAT CT | | | |
| | 110000.004 | GGA TTT ATT AAA AGC AAA ACA TAC AAA | 188 | | |
| SSR | 0D099-024 | CAA TAA CAA ATG AGC ATG ATA AGA CA | 100 | | |
| | 110000 021 | TAT CCT CTA TGT GGC GAT | 151 | | |
| | 00099-031 | TTG GTT AAA AGC ATT GAT ACA | | | |
| | 110000.042 | TCG GCT TTA CAA CCC ATT TC | 174 | | |
| | 0D099-043 | TGC CAA TTA TGG GGC TAA CT | | | |
| | 110000 020 | AAT TAC CAT GGG CAG AGG AG | 170 | | |
| | 00099-039 | CCC CAA AAG CTC CAT TAT TGT | | | |

Only four SSR markers produced readable bands (UDO99-006, UDO99-024, UDO99-039, and UDO99-043). All SSR markers were monomorphic with the eleven Nabali variants. This outcome confirmed that all studied and selected olive trees were Nabali and any presented genetic variation due to intra-variation within Nabali rather than mis–selection with other cultivar. Ten RAPD primers produced 60 reproducible bands with an average of 6 bands / marker. Out 60 bands only 24 were polymorphic. The percentage of polymorphic bands was 40% which is relatively high.

Based on Jaccards coefficient, similarity matrix for studied populations was done. Similarity ranged from moderate (0.610) for Jalkamous and Karawa Bani Zaid (2) to highly genetic similarity or even identity in some cases such as Aqraba and Aseerah (N) (1.000) (Table 3). Interaction between different variants showed high trends. For example the interaction between Nabali variants from Jalkamous and Alaroub was high (0.925) the same trend was found between variants Salfeet and Salfeet (h) (0.953), Agraba and Aseerah (N) (0.952). On other side some exceptions were found, the interaction for Nabali variants from Karawa Bani Zaid, Bieta and Alaroub was (0.675).

The effect of geographic location was absent in this study and has no significant contribution. For instance for two variants from Aqraba and Aseerah (N) the similarity was 1.000. As well variant from Nahaleen (in the south of Palestine) with variants from Akarab, Aseerah (N), and Alaar (North of Palestine) were 0.914, 0.914, and 0.912, respectively. On other side, Karawa Bani Zaid (1) and Karawa Bani Zaid (2) are two variants from the same village showed low genetic similarity (0.667). The high similarity in this study suggested a genetic similarity or even identity between different variants from different locations. This could be due to fact that these variants come from the same gene pool.

Dendogram based on Jaccards cofficent revealed three main groups (Figure 1), the biggest group consisted of the majority of variants including eight Nabali varints; Bieta, Karawa Bani Zaid (1), Salfeet, Salfeet (h), Aqraba, Aseerah (N), Jalkamous, and Alaroub. Second group consisted of two variants Alaar and Nahaleen. While the third group is contains only Karawa Bani Zaid (2).

DISCUSSION AND CONCLUSIONS

Two types of molecular markers were used in this study, SSR and RAPD markers. SSR markers were monomrphic and produced the same profile for all Nabali variants. In opposite, microsatellites were effective tool to discriminate on intra-varietal genotypes level (Muzzalupo *et al.*, 2010; Ipek *et al.*, 2012). The negative result of SSR may be due to the low number of used markers and the small size of the population. The same copies of profile of microsatellite for all samples confirmed that, all selected trees belong to the same olive cultivar Nabali. As well, any existed genetic variation within the olive population were due to the

Table3. Similarity matrix of Jaccard's coefficient of 11 olive trees (variants) in different sites in Palestine.

| | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | |
|----|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|--|
| 1 | 1.000 | | | | | | | | | | | |
| 2 | 0.868 | 1.000 | | | | | | | | | | |
| 3 | 0.610 | 0.667 | 1.000 | | | | | | | | | |
| 4 | 0.770 | 0.829 | 0.788 | 1.000 | | | | | | | | |
| 5 | 0.925 | 0.892 | 0.675 | 0.842 | 1.000 | | | | | | | |
| 6 | 0.814 | 0.893 | 0.675 | 0.842 | 0.864 | 1.000 | | | | | | |
| 7 | 0.878 | 0.917 | 0.692 | 0.865 | 0.864 | 0.826 | 1.000 | | | | | |
| 8 | 0.878 | 0.865 | 0.692 | 0.865 | 0.910 | 0.867 | 0.953 | 1.000 | | | | |
| 9 | 0.829 | 0.914 | 0.730 | 0.914 | 0.860 | 0.864 | 0.952 | 0.952 | 1.000 | | | |
| 10 | 0.829 | 0.914 | 0.730 | 0.914 | 0.860 | 0.864 | 0.952 | 0.952 | 1.000 | 1.000 | | |
| 11 | 0 780 | 0.757 | 0 722 | 0.912 | 0 773 | 0 778 | 0.860 | 0.860 | 0.857 | 0.857 | 1 000 | |

Abberiviation: 1: Jalkamous, 2: Karawa bani zied (1), 3: Karaw abani zied (2), 4: Nahaleen, 5: Alaroub, 6: Bieta, 7:Salfeet, 8: Salfeet (h), 9: Aqraba, 10: Aseerah(N), and 11: Alaar.



Figure 1. Dendogram based on Jaccards coefficient illustrating genetic similarity among eleven Nabali variants in Palestine based on 10 RAPD markers.

intra-genetic rather than inter-genetic variation level. In this study, RAPD were able to distinguish the genetic variation among different Nabali variants. Over seventy samples were screened with RAPD in first time. RAPD marker provides a cheap and fast tool for screening large population, which offers a good choice for researchers in developing countries (Belaj *et al.*, 2003; Belaj *et al.*, 2004). Combining two types of Markers SSR and RAPD in this study was good strategy to overcome the low reproducibility of RAPD. Out of fifty eight, twenty two bands were polymorphic, the relative high polymorphic bands (40%) and moderate genetic similarity among different Nabali variants suggested attribution of genetic background. In Palestine, Nabali known as difficult rooting olive cultivar and grafting is the only and common vegetative propagation method. As well, sexual propagation is not used. Therefore, the variation due to somatic mutation within Nabli cultivar is possible since somatic mutations considered as an important source of intra-plant genetic variation (Salomonson, 1996). Genetic contribution within Nabali variants or other olive cultivars was indicated before. (Wiesman *et al.*, 1998; Muzzalupo *et al.*, 2010; and Ipek *et al.*, 2012). The low-moderate genetic variation obtained in this study could be due to two main reasons. First, the predominant propagation method for olive in Palestine is grafting. Second, Nabali is self-pollinated cultivar. Therefore, the maintenance of cultivar is expected and could lead to minimize the genetic diversity within Nabali olive trees in Palestine.

In this study, two Nabali olive trees from Akarab and Aseerah (N) were completely genetically identity (1.00). This may due to the fact that Aseerah is known as a high quality olive oil producer village in Palestine and containing very old olive orchards which back to the Roman period. Therefore, trees from Aseerah (N) are expected to be used as a good source for scions which are used in grafting in many olive orchards. Oppositely, the similarity for two Nabali trees from Karawa Bani Zaid was moderate (0.667). In other cases, the genetic similarity was high for two Nabali variants from too far distance areas, Nahaleen (south of Palestine) and Aqraba (North of Palestine) (0.914). Therefore, the geographic

REFERENCES

- Abdelhamid, S., N. Grat-Kamoun, F. Marra, and T. Caruso. 2012. Genetic similarity among Tunisian cultivated olive estimated through SSR markers. Scientia Agricola 70: 33-38.
- Angiolillo, A., M. Mencuccini, L. Baldoni. 1999. Olive genetic diversity assessed using amplified fragment length polymorphisms. Theoretical and Applied Genetics 98: 411-421.
- Belaj, A., Z. Satovic, G. Cipriani, L. Baldoni, R. Testolin, L. Rallo, I. Trujillo. 2003. Comparative study of the discriminating capacity of RAPD AFLP and SSR markers and of their effectiveness in establishing genetic relationships in olive. Theoretical and Applied Genetics 107: 736-744.
- Belaj, A., Z. Satovic, L. Rallo, I. Trujillo. 2004. Optimal use of RAPD markers for identifying varieties in olive (*Olea europaea* L.) germplasm collections. Journal of the American Society for Horticultural Science 129: 266-270.

contribution in genetic variation within Nabali variants is absent and eliminated.

The intra- genetic variation within Nabali variants is approved in this study. Few groups or lines of Nabali were found. The main group contains the majority of studied lines (eight variants). These group representatives the predominant Nabali variant in Palestine. Other lines of Nabali were found in this study. The available of these lines or genotypes is less.

Because Nabali variant is adapted well to the local conditions and producing oil with high quality, Nabali still the preferred olive variant for farmers in Palestine. Nabali is known as susceptible to peacock disease and showed an alternative bearing phenomena. The fact that there intra-genetic variation is existed within Nabali variants, justifies the selection for new lines with promising traits. In this study, several new lines (variants) of Nabali were identified. These variants will be subject for more research in order to evaluate its performance against important agronomic traits such as, alternative bearing phenomena and resistant to peacock disease, consequently, could be integrated in future in national breeding program.

- Cipriani, G., M. T. Marrazzo, R. Marconi, A. Cimato, R. Testolin. 2002. Microsatellite markers isolated in olive (*Olea europaea* L.) are suitable for individual fingerprinting and reveal polymorphism within ancient cultivars. Theoretical and Applied Genetics 104 (2-3): 223-228.
- Gomes, S., P. Martins-Lopes, J. Lima-Brito, J. Meirinhos, J. Lopes, A. Martins, H. Guedes-Pinto. 2008. Evidence of clonal variation in olive 'Verdeal-Transmontana' cultivar using RAPD, ISSR and SSR markers. Journal of Horticultural Science and Biotechnology 83: 395-400.
- Gomes, S., P. Martins-Lopes., H. Guedes-Pinto. 2012. Olive tree genetic resources characterization through molecular markers. *In:* Mahmut Caliskan (Ed.), Genetic Diversity in Plants, ISBN: 978-953-51-0185-7, InTech, DOI: 10.5772/32973. Available from: http://www.intechopen.com/books/genetic-diversityin-plants/olive-tree-genetic-resources-characterizationthrough-molecular-markers.
- Ipek, A., E. Barut, H. Gulen, M. Ipek. 2012. Assessment of inter and intra-cultivar variations in olive using SSR markers. Sci. Agric. 69: 327-335.

- Jaccard P. 1908. Nouvelles recherches sur la distribution florale. Bulletin de la Société Vaudoise des Sciences Naturelles 44: 223-270.
- Khadari, B., C. Breton, N. Moutier, J. Roger, G. Besnard, A. Berville, F. Dosba. 2003. The use of molecular markers for germplasm management in a French olive collection. Theoretical and Applied Genetics 106: 521-529.
- Muzzalupo, I., A. Chiappettac, C. Benincasaa, E. Perri. 2010. Intra-cultivar variability of three major olive cultivars grown in different areas of central-southern Italy and studied using microsatellite markers. Scientia Horticulturae 126: 324-329.
- Owen, C., A. Carolyn, L. Bita, G. Banilas, S. Hajjar, V. Sellianakis, U. Aksoy, S. Hepaksoy, R. Chamoun R., S. Talhook., I. Metzidakis, P. Hatzopoulos, P. Kalaitzis. 2005. AFLP reveals structural details of genetic diversity within cultivated olive germplasm from the Eastern Mediterranean. Theoretical and Applied Genetics 110: 1169-1176.

- Qutub, M., S. Ali, M. Mutawea, M. Abed, T. Arabasi, F. Pierini, E. Lodolini. 2010. Characterization of the main Palestinian olive cultivars and olive oil.
- Salomonson, A. 1996. Interactions between somatic mutations and plant development. Vegetatio 127: 71-75.
- Schlüter, P. M., S. A. Harris. 2006. Analysis of multilocus fingerprinting data sets containing missing data. Mol. Ecol. Notes. 6: 569-572.
- Wiesman, Z., A. Avidan, S. Lavee, B. Quebedaux.1998. Molecular characterization of common olive varieties in Israel and the West bank using randomly amplified polymorphic DNA (RAPD) markers. J. Amer. Soc. Hort. Sci. 123: 837-841.
- Zohary, D., and M. Hopf. 1994. Domestication of plants in the Old World: The origin and spread of cultivated plants in west Asia, Europe, and the Nile Valley. 2nd ed. Oxford Clarendon Press.