Full Length Research Paper

Molecular diversification and preliminary evaluations of some satsuma selections' performance under mediterranean conditions

Meral İncesu*, Önder Tuzcu, Turgut Yeşiloğlu, Yıldız Aka Kaçar, Bilge Yıldırım, Melda Boncuk and Berken Çimen

University of Çukurova, Faculty of Agriculture, Department of Horticulture 01330 Adana, Turkey.

Accepted 28 March, 2011

Satsuma (*Citrus unshiu* Marc.) has been cultivated for a long time in Turkey, and therefore variations for agronomical traits are likely among cultivated satsumas due to bud mutations. The objectives of this study were to determine variations for some selected agronomical traits and genetic markers among 21 new satsumas derived from selections. Fruit yield, fruit quality and molecular diversification of these clones were determined. The clones of 62 Adana, 4/2 Izmir and 11/1 Izmir indicated the highest yield. The fruits obtained from all clones were heavier than the control (Owari Satsuma). Molecular analysis, as assessed with 9 random amplification of polymorphic DNA (RAPD) and 14 simple sequence repeats (SSR) primers, indicated that satsuma clones showed a narrow genetic base suggesting that the observed morphological polymorphism within the group must be associated with somatic mutations which were not detected by these molecular markers.

Key words: Mandarins, clonal selection, citrus.

INTRODUCTION

Annual citrus production is approximately 120 million tonnes world-wide (FAO, 2009) and Turkey is one of the important citrus producing countries among the Mediterranean. This region constitutes 17.5% of the world's total citrus production. Total citrus production in Turkey is approximately 3 million tonnes and 25.2% of this was of mandarin (FAO, 2009).

Mandarins exhibit more variation in characters than other citrus species (Reuther et al., 1967), and is the most phenotypically heterogeneous group in *Citrus* (Moore, 2001). Satsuma mandarin is considered to have originated in Japan (Reuther et al., 1967) and also probably as a chance seedling from China (Nishiura, 1964). Chance seedling selections, bud sports and spontaneous mutations are still a major source to find new citrus cultivars. Conventional breeding is slow and difficult due to the complex reproductive biology in citrus (JinPing et al., 2009). Bud mutations arise frequently in citrus, and clonal selection has been used as a traditional breeding method (Uzun et al., 2009).

Although, Turkey is not a natural genetic center of citrus fruits, they have been grown in this country for several years. To find Satsuma varieites which have the features in the direction of the market demands and high yield, the first national citrus budwood selection was carried out between the years 1979-1983 in the Mediterranean and Aegean regions of Turkey. A thorough search of trees in the regions for high yield, trees without pests and diseases and high quality has been repeated every year for 4 years (Özsan et al., 1986). In this project, 21 Satsuma genotypes with high yield and fruit quality were selected from citrus growing areas of Turkey.

RAPD, AFLP, ISSR and SSR markers or isozyme have been used to distinguish between accessions and for investigations of genetic diversity/relatedness and genetic mapping in citrus (Green et al., 1986; Denget al., 1995; Kijas et al., 1995; Machado et al., 1995; Fang et al.,

^{*}Corresponding author: E-mail: meralincesu@gmail.com. Tel: +90 322 338 63 88 or +90 322 338 63 88.

Abbreviations: RAPD, Random amplification of polymorphic; **SSR,** simple sequence repeats; **UPGMA,** unweighted pair group method analysis; **ISSR,** inter-simple sequence repeat.

Table 1. RAPD Primers Used in the study.

| RAPD primer | Primer sequence |
|-------------|-----------------|
| OPA04 | AATCGGGCTG |
| OPA05 | AGGGGTCTTG |
| OPA09 | GGGTAACGCC |
| OPA13 | CAGCACCCAC |
| OPA14 | TCTGTGCTGG |
| OPA15 | TTCCGAACCC |
| OPA16 | AGCCAGCGAA |
| OPA17 | GACCGCTTGT |
| OPA19 | CAAACGTCGG |

1997; Fang and Roose, 1997; Masashi et. al., 1998; Federici et al., 1998; Nicolisi et al., 2000; Abkenar et al., 2004; Oliveira et al., 2004; Aka-Kacar et al., 2005; Lin and Chen, 2006; Pang et al., 2007; Wei, 2007; Hvarleva et al., 2008; Rao et al., 2008; Baig et al., 2009; Uzun et al., 2009). Since the introduction of (RAPD) markers in 1990 (Williams et al., 1990), their use in plant genetic analysis has increased in an exponential manner, principally due to its simplicity, quick and easy assay. RAPD analysis has been used for genotype characerization (Khadari et al., 1995; Lu et al., 1996; Nicese et al.,1998), to assess intra-or interpopulation genetic variability (Huff et al., 1993; Alberto et al., 1997; Baig et al., 2009), for genetic mapping (Chaparro et al., 1994; Oliveira et al., 2004) and to identify molecular markers linked to genes of interest (Nair et al., 1996). SSR markers have several advantages over other molecular markers. They are in fact abundant in most genomes and co-dominant, therefore, their information content is very high; SSRs are PCR based, thus requiring little DNA for the amplification; and every SSR locus is defined by a unique pair of primers, so that information exchange between labotaratories is easy (Lacis et al., 2009).

The objective of the study was to determine the variations of some selected agronomical traits of 21 Satsuma mandarins derived from selection breeding programmes and to characterize them by SSR and RAPD genetic markers.

MATERIALS AND METHODS

Plant materials

21 satsuma varieties collected from the Mediterranean and Aegean regions of Turkey between 1979-1983 were used in this study. Thoroughly searching trees in the regions, selections of trees with high yield and quality have been repeated every year for 4 years. The name of Satsuma clones were 61 Adana, 62 Adana, 89 Adana, 3 Izmir, 22 Izmir, 26 Izmir, 30 Izmir, 4/2 Izmir, 8/3 Izmir, 11/1 Izmir, 13/2 Izmir, 18/1 Izmir, 19/1 Izmir, 24/2 Izmir, 25/1 Izmir, 27/1 Izmir, 27/2 Izmir,23 Mersin,24 Mersin, 25 Mersin, 27 Mersin and Owari Satsuma which was used as the control for the selections.

Orchard management

The yield and fruit characteristics of 21 (selections and one control) satsuma mandarin types were examined under ecological conditions of Adana province. The study was conducted at the research area of Çukurova University in Adana. Clones were planted on silty-clay-loam textured soil having a pH of 7.3. Plants were spaced with 7x7 m spacing. Clones were grafted on sour orange (*Citrus aurantium* L.) and the experiment was established using completely randomized design. Five replications were established for each genotype. The experimental area was managed according to standard commercial practces.

Characterization for tree yield performance and fruit quality

Selected Satsuma clones were evaluated as follows.

Yield efficinecy

Yield by tree (kg/tree) were recorded during 3 years of production and according to these data, cumulative yield was calculated.

Fruit quality

Commercial harvest of satsumas in Adana is the middle of November. At harvest, 25 fruits were harvested. All fruit samples were assessed for fruit weight (g), fruit height (mm), fruit diameter (mm), index, rind thickness (mm), juice content (%), brix (%), total acids (%) and brix/acid ratio for 3 years.

Molecular analysis

DNA extraction

Ten young leaves were collected from a single tree for each genotype, and immediately frozen in liquid nitrogen and stored at -80 °C. Total genomic DNA was extracted from leaf tissue following the protocol for minipreps by using CTAB (Dellaporta et al., 1983). DNA concentrations were assessed by spectrophotometer (Nanodrop ND-1000).

RAPD analysis

Nine RAPD primers (OPA04, OPA05, OPA09, OPA13, OPA14, OPA15, OPA16, OPA17 and OPA19) (Table 1) were used for PCR amplifications to determine genetic characterization of Satsuma mandarins. PCR was performed in 12.5 μ l reaction mixture containing the following: 15 ng DNA, 6.25 μ l 2X PCR Master Mix, 0.5 μ l 25 mM MgCl₂, 1.25 μ l primer, 0,05 of *Taq* DNA polymerase and distilled water (1.45 μ l).

Amplified DNA fragments were loaded in a 1.5% agarose gel in 1X TAE buffer and submitted to electrophoresis for 3 h at 70 volt. DNA fragments were visualized under ultra-violet light and gel image was captured by photo documentation system and the images stored for further analysis. A 1 kb ladder was used as the molecular standard in order to confirm the appropriate RAPD markers.

SSR markers

14 SSR primers characterized for *Citrus* were used in this study (Table 2). PCR amplifications were carried out in 20 μ l final reaction

| | SSR Primer |
|----------|------------------------------|
| 458 | F: CCCCCTCTTTTCTCTCCA |
| | R: TTCTGGGCTGGTAGGTTCAG |
| | F: TCGCCCTCCCCCTGAAATTA |
| 571 | R: GAAAGCCTGGTGGGAGCAGA |
| | |
| 197 | F: CGCAATTCAATTCCCTGTCT |
| 407 | R: CGTCGAGCAACAAATCAAGA |
| | |
| 506 | |
| | R. CATATOCCATCACCACCAAA |
| 405 | F: GGCCTTAAACCACCTTGACA |
| 495 | R: TGAGGCTTTTGCTGTTGTTG |
| | |
| 473 | F: CTTGGCGTCGAAAAGAAATC |
| | R: AGCACGGATGTCAAAATTCC |
| | |
| 485 | R: GGTCGAGATTGAGCAGCAGT |
| | |
| 607 | F: TGCATTTTGTGGGTCTTGCTTG |
| 037 | R: GGCCCTGACTGCTGCAAGAT |
| | |
| 498 | |
| | |
| | F: GAGCTCAAAACAATAGCCGC |
| WE51 431 | R: CATACCTCCCCGTCCATCTA |
| | |
| 488 | F: CACGCTCTTGACTTTCTCCC |
| | R: CTTTGCGTGTTTGTGCTGTT |
| | F: GCATAGAATAAGAAATGACAGCAAA |
| 173 | R: ATGCCTGCACCTTTGGTAAG |
| | |
| 54 | F: AACACCTTAAGGCTGCAGGA |
| 04 | R: CGTTGTTGATGATTCTTGATGA |
| | E: TOCOTATOATOGGOAACTTO |
| MEST 121 | R: CAATAATGTTAGGCTGGATGGA |
| | |

Table 2. The Forward and Reverse SSR Primer Information for This Study.

volumes each containing 25 ng of DNA, 2X PCR Master Mix, 5 μ l ddH₂O, 0.5 μ l of 25 mM MgCl₂, 0.05 unit Taq polymerase, 0.025 μ M M13 primer, 3'and 5'end primers (F + R).

The DNA amplifications were carried out in a thermocycler (Eppendorf Mastercyler Gradient). The mixture was initially denatured at 94 °C for 5 min; followed by 35 cycles at 94 °C for 1 min; 55 to 60 °C for 30 sec; 72 °C for 1 min; and the final extension step at 72 °C for 4 min. PCR products were stored at 4 °C before analysis.

In SSR analysis, after amplification, an equal volume of formamide loading buffer was added to each reaction tube containing 95% formamide, 10 mM EDTA (pH 8.0), 0.025% of xylene cyanol and 0.025% of bromophenol blue. The samples were heat denatured for 5 min at 95°C and quickly cooled on ice. After loading 1.0 μ l of each sample, PCR products were separated in a 25-cm, 8% denaturing polyacrylamide gel (Long Ranger, FMC Biozym, Hessisch Oldendorf, Germany) that had been preheated for 30 min. Electrophoresis was conducted in 1.0 Long Ranger TBE buffer at 1500 V, 50 W, 35 mA and 48°C using a Li-Cor DNA Analyzer 4200 (Licor Biosciences, Bad Homburg, Germany). A 50-350 bp DNA ladder mix (MWG Biotech AG, Ebersberg, Germany)

| Clone | 10 year old | 11 year old | 12 year old | Cumulative yield |
|---------------|-----------------------|---------------------|-----------------------|-----------------------|
| 61 Adana | 37.25 ^{b-e+} | 81.00 ^a | 94.70 ^{bcd} | 218.40 ^{efg} |
| 62 Adana | 24.07 ^e | 116.33 ^a | 147.30 ^a | 289.60 ^{a-d} |
| 89 Adana | 37.58 ^{b-e} | 89.12 ^a | 107.70 ^{abc} | 311.00 ^{ab} |
| 3 Izmir | 32.37 ^{cde} | 54.00 ^a | 83.13 ^{cd} | 197.65 ^g |
| 22 Izmir | 39.63 ^{b-e} | 102.60 ^a | 108.73 ^{abc} | 272.90 ^{b-e} |
| 26 Izmir | 36.23 ^{b-e} | 93.67 ^a | 122.47 ^{ab} | 281.30 ^{a-e} |
| 30 Izmir | 44.37 ^{a-d} | 122.67 ^a | 115.13 ^{ab} | 282.16 ^{a-e} |
| 4/2 Izmir | 53.47 ^{ab} | 126.28 ^a | 120.80 ^{ab} | 337.95 ^a |
| 8/3 Izmir | 31.17 ^{cde} | 62.30 ^a | 93.97 ^{bcd} | 232.90 ^{d-g} |
| 11/1 Izmir | 41.65 ^{b-e} | 119.74 ^a | 130.17 ^{ab} | 316.13 ^{ab} |
| 13/2 Izmir | 39.93 ^{b-e} | 87.60 ^a | 126.73 ^{ab} | 255.50 ^{b-g} |
| 18/1 Izmir | 40.63 ^{b-e} | 108.67 ^a | 89.30 ^{cd} | 231.56 ^{d-g} |
| 19/1 Izmir | 40.00 ^{b-e} | 97.00 ^a | 138.47 ^{ab} | 301.10 ^{abc} |
| 24/2 Izmir | 24.63 ^e | 126.00 ^a | 118.63 ^{ab} | 269.26 ^{b-f} |
| 25/1 Izmir | 58.80 ^a | 91.78 ^ª | 129.35 ^{ab} | 269.40 ^{b-f} |
| 27/1 Izmir | 40.63 ^{ab} | 108.67 ^a | 89.30 ^{cd} | 273.85 ^{a-e} |
| 27/2 Izmir | 48.27 ^{abc} | 105.00 ^a | 94.90 ^{bcd} | 219.03 ^{efg} |
| 23 Mersin | 41.93 ^{b-e} | 87.08 ^a | 111.87 ^{ab} | 242.95 ^{c-g} |
| 24 Mersin | 38.70 ^{b-e} | 91.00 ^ª | 107.93 ^{bcd} | 286.25 ^{a-d} |
| 25 Mersin | 27.25 ^{de} | 92.63 ^a | 63.43 ^{de} | 205.60 ^{fg} |
| 27 Mersin | 33.07 ^{cde} | 80.25 ^a | 105.43 ^{abc} | 279.70 ^{a-e} |
| Owari Satsuma | 41.93 ^{a-e} | 45.00 ^a | 37.00 ^e | 129.86 ^h |
| Signifinance | ** | NS | ** | ** |

Table 3. Yield of new Satsuma mandarin clones on sour orange (kg/tree).

**Significant at P<0.01; NS, not significant.

Values followed by a common letter in a column for each clone are not significantly different at P<0.01 by Duncan's Multiple Range test.

was run alongside the amplified PCR products to determine their approx. sizes.

Data analysis

The experiment was carried out during a three year period on 10-11 and 12 year old trees. All data were analyzed using analysis of variance (ANOVA) with the SPSS 17.0 (SPSS, Chicago, USA) program. Mean seperations were determined using Duncan Multiple Range test at a 0.05 significance level.

For molecular analysis, amplified fragments were classified as present (represented with 1) or absent (represented with 0) for both RAPD and SSR analysis. Genetic similarity values (Nei and Li, 1979) were calculated, and Unweighted Pair Group Method analysis (UPGMA) cluster analysis was performed to generate dendrogram with NTSYS-PC version 2.02i programme.

RESULTS AND DISCUSSION

Yield efficiency

Statistically, significant differences were recovered among the clones for most of the traits evaluated. The clones of 62 Adana (101.88 kg/tree), 4/2 lzmir (103.33 kg/tree) and 11/1 lzmir (96.57 kg /tree) gave the highest yield. These means were much higher than Owari Satsuma with 40.49 kg/tree, commonly grown up in the conditions of Çukurova, was used as control (Table 3). Observations of all clones indicated that, 4/2 Izmir clone was the highest (337.95 kg/tree), regarding cumulative yield. In contrast, the lowest cumulative yield (129.86 kg/tree) was determined from Owari Satsuma, as control, among all the selected clones. Also 89 Adana, 11/1 Izmir and 19/1 Izmir were significantly prominent with their high cumulative yields, 311 kg/tree, 316.13 kg/tree and 301,10 kg/tree respectively.

Fruit quality

The heaviest fruits were obtained from the clones of 62 Adana (144.58 g/fruit), 18/1 Izmir (141.28 g/fruit) and 30 Izmir (135.14 g/fruit). The control, Owari Satsuma, had fruit averaging 103.67 g. All clones gave heavier fruits than the control. Köse (2000) analyzed some Satsuma clones in the conditions of Çukurova and determined the fruit weight of Satsuma Sato (145.70 g), Satsuma Suzuki Wase (98.20 g) and Satsuma Owari Sra 12 (75.80 g). Most clones whose performance was tested in this study showed better performance than the cultivars aforementioned.

| Clone | Fruit weight | Fruit height | Fruit diameter | Index | Rind thickness |
|---------------|---------------------|-----------------------|----------------------|----------------------|---------------------|
| 61 Adana | 126.35 ^a | 54.11 ^{b-d+} | 65.87 ^{abc} | 1.224 ^{abc} | 3.10 ^{a-f} |
| 62 Adana | 144.58 ^a | 53.52 ^{cde} | 68.14 ^{ab} | 1.282 ^{ab} | 3.15 ^{a-e} |
| 89 Adana | 131.67 ^a | 54.69 ^{ab} | 67.09 ^{ab} | 1.234 ^{abc} | 3.12 ^{a-f} |
| 3 Izmir | 134.99 ^a | 54.71 ^{ab} | 66.55 ^{ab} | 1.218 ^{abc} | 2.92 ^{efg} |
| 22 Izmir | 121.21 ^a | 52.13 ^e | 66.75 ^{ab} | 1.288 ^a | 2.86 ^{fg} |
| 26 Izmir | 126.09 ^a | 52.85 ^{de} | 67.41 ^{ab} | 1.286 ^a | 3.34 ^a |
| 30 Izmir | 135.14 ^a | 53.97 ^{cd} | 68.24 ^{ab} | 1.266 ^{abc} | 3.22 ^{a-d} |
| 4/2 Izmir | 121.26 ^a | 52.65 ^{de} | 66.83 ^{ab} | 1.272 ^{ab} | 2.87 ^{fg} |
| 8/3 Izmir | 120.34 ^a | 52.20 ^e | 66.72 ^{ab} | 1.284 ^a | 2.97 ^{d-g} |
| 11/1 Izmir | 131.72 ^ª | 54.06 ^{b-d} | 66.79 ^{ab} | 1.240 ^{abc} | 3.07 ^{b-f} |
| 13/2 Izmir | 125.74 ^a | 56.17 ^a | 67.09 ^{ab} | 1.200 ^c | 2.82 ^g |
| 18/1 Izmir | 141.28 ^a | 55.73 ^{ab} | 68.97 ^a | 1.242 ^{abc} | 2.96 ^{d-g} |
| 19/1 Izmir | 129.89 ^a | 53.82 ^{cde} | 66.76 ^{ab} | 1.248 ^{abc} | 3.27 ^{ab} |
| 24/2 Izmir | 129.07 ^a | 54.93 ^{ab} | 66.36 ^{abc} | 1.214 ^{bc} | 3.36 ^a |
| 25/1 Izmir | 132.28 ^a | 54.30 ^{b-d} | 69.02 ^{ab} | 1.276 ^{ab} | 3.18 ^{a-e} |
| 27/1 Izmir | 119.36 ^a | 53.62 ^{cde} | 64.97 ^{bc} | 1.212 ^{bc} | 2.99 ^{c-g} |
| 27/2 Izmir | 122.08 ^a | 54.07 ^{b-d} | 65.29 ^{bc} | 1.212 ^{bc} | 2.98 ^{d-g} |
| 23 Mersin | 131.57 ^a | 53.77 ^{cde} | 66.70 ^{ab} | 1.244 ^{abc} | 3.05 ^{b-f} |
| 24 Mersin | 131.95 ^a | 56.36 ^a | 63.12 ° | 1.124 ^d | 3.30 ^{ab} |
| 25 Mersin | 127.10 ^a | 53.42 ^{cde} | 66.22 ^{abc} | 1.250 ^{abc} | 3.25 ^{abc} |
| 27 Mersin | 130.65 ^a | 54.88 ^{ab} | 67.26 ^{ab} | 1.228 ^{abc} | 2.83 ^g |
| Owari Satsuma | 103.67 ^a | 48.97 ^f | 59.09 ^d | 1.218 ^{abc} | 2.83 ^g |
| Significance | NS. | ** | ** | ** | ** |

Table 4. Fruit Quality of New Satsuma Mandarin Clones on Sour Orange Rootstock.

**Significant at P<0.01; NS, not significant.

Values followed by a common letter in a column for each clone are not significantly different at P<0.01 by Duncan's Multiple Range test.

All clones gave bigger fruits than the control when they were analyzed in terms of fruit diameter and fruit height. Fruit shape, evaluated by fruit index, was almost similar among the clones tested.

The thinnest peeled ones were from 13/ 2 Izmir (2.82 mm) and the control plants (2,83 mm), while the thickest peeled fruits were being observed in the clones of 24/2 Izmir (3.36 mm), 26 Izmir (3.37 mm) and 24 Mersin (3.30 mm) (Table 4).

The highest amount of fruit juice was obtained from the clone of 19/1 Izmir (46.27 %), 8/3 Izmir (46.20%) and 61A (45.69%) whereas the lowest one was obtained from the clone of the control (39.78%) and 27/1 Izmir (40.37%) (Table 5). Similarly, Urgun (1997) and Köse (2000) notified that the amount of fruit juice in the same Satsuma types (Sugiyama, Silverhill, Ikeda, Hayashi) that they analyzed, showed changes between 40.49 - 45.75%. According to these results, the clones selected show superiorities in terms of fruit juice amount.

22 Izmir (11.38%) clone had the highest total soluble solids, in contrast the lowest was determined from 24 Mersin (10.09%) clone. In terms of total acidity, the highest and the lowest were obtained from 27/1 Izmir

(1.47%) and Owari Satsuma (1.16%) respectively (Table 5).

The highest amount of soluble solids/acidity was obtained from the clones of 13/2 Izmir (9.33), control (8.53), 4/2 Izmir (8,78) and 30 Izmir (8,57) but the lowest amount was obtained from the clones of 24 Mersin (7.40) and 27/1 Izmir (7.30) (Table 5).

Molecular analysis

RAPD analysis

The nine 10-mer RAPD primers generated 75 amplification products; the total number of marker ranged from 3 (OPA15) to 12 (OPA16). The sizes of the bands ranged from 250 to 2200 bp (Table 6). In average, each primer gave 8.33 scorable markers per amplification. The similarity value among 22 genotypes ranged from 0.79 to 1. This may imply low genetic variability among the genotypes studied.

Based on RAPD analysis, a dendrogram is illustrated in Figure 1. The results reveal that, 21 genotypes grouped

| Clone | Juice content (%) | Total soluble solid (TSS) (%) | Total acidity (TA) (%) | TSS/TA |
|---------------|-----------------------|----------------------------------|---------------------------|---------------------|
| 61 Adana | 45.69 ^{abc+} | 11.14 ^{abc} | 1.39 ^{abc} | 7.99 ^{bcd} |
| 62 Adana | 44.30 ^{a-d} | 10.92 ^{a-e} | 1.38 ^{bcd} | 8.05 ^{bcd} |
| 89 Adana | 43.43 ^{c-f} | 10.79 ^{be} | 1.29 ^{d-h} | 7.77 ^{bcd} |
| 3 Izmir | 43.70 ^{c-f} | 10.60 ^{de} | 1.21 ^{ghi} | 8.28 ^{a-d} |
| 22 Izmir | 43.35 ^{c-f} | 11.38 ^a | 1.36 ^{cde} | 8.46 ^{abc} |
| 26 Izmir | 43.10 ^{def} | 10.54 ^{def} | 1.30 ^{def} | 8.13 ^{bcd} |
| 30 Izmir | 45.59 ^{abc} | 10.98 ^{a-e} | 1.22 ^{f-i} | 8.57 ^{ab} |
| 4/2 Izmir | 42.37 ^{d-g} | 11.02 ^{a-d} | 1.28 ^{e-h} | 8.78 ^{ab} |
| 8/3 Izmir | 46.20 ^{ab} | 10.76 ^{b-e} | 1.33 ^{cde} | 8.18 ^{bcd} |
| 11/1 Izmir | 43.88 ^{b-e} | 10.91 ^{a-e} | 1.37 ^{bcd} | 8.07 bcd |
| 13/2 Izmir | 44.22 ^{a-d} | 10.99 ^{a-d} | 1.19 ⁱ | 9.33 ^a |
| 18/1 Izmir | 41.39 ^{e-h} | 10.74 ^{cde} | 1.21 ^{hi} | 8.34 ^{a-d} |
| 19/1 Izmir | 46.27 ^a | 10.47 ^{ef} | 1.29 ^{d-g} | 8.16 ^{bcd} |
| 24/2 Izmir | 41.70 ^{e-h} | 10.98 ^{a-d} | 1.40 ^{abc} | 7.96 ^{bcd} |
| 25/1 Izmir | 42.91 ^{d-f} | 11.35 ^a | 1.46 ^{ab} | 7.86 ^{bcd} |
| 27/1 Izmir | 40.37 ^{gh} | 11.27 ^{ab} | 1.47 ^a | 7.30 ^d |
| 27/2 Izmir | 42.37 ^{d-g} | 11.04 ^{a-d} | 1.38 ^{bcd} | 8.04 ^{bcd} |
| 23 Mersin | 42.06 ^{d-h} | 11.19 ^{abc} | 1.36 ^{cde} | 8.26 ^{a-d} |
| 24 Mersin | 42.48 ^{d-g} | 10.09 ^f | 1.37 ^{cd} | 7.40 ^{cd} |
| 25 Mersin | 44.37 ^{a-d} | 11.02 ^{a-d} | 1.34 ^{cde} | 8.28 ^{a-d} |
| 27 Mersin | 41.38 ^{fgh} | 10.80 ^{b-e} | 1.23 ^{f-i} | 8.23 ^{bcd} |
| Owari Satsuma | 39.78 ^h | 10.55 ^{def} | 1.16 ⁱ | 8.53 ^{ab} |
| Significance | ** | ** | ** | * |

Table 5. Average fruit quality of new Satsuma mandarin clones on sour orange.

**Significant at P<0.01; *significant at P<0.05.

Values followed by a common letter in a column for each clone are not significantly different at P<0.05 by Duncan's multiple range test.

Table 6. Band size and band number of primers used for RAPD analysis.

| S/N | Primer | Band size | Band number |
|-----|--------|--|-------------|
| 1 | OPA04 | 250, 350, 500, 510, 750, 760, 1000, 1250, 1500, 1510, 2000 | 11 |
| 2 | OPA 05 | 250, 450, 500, 600, 750, 800,1000,1250,1500, 2000 | 10 |
| 3 | OPA09 | 250, 350, 500, 510, 750, 850, 1000, 1450 | 8 |
| 4 | OPA13 | 300, 480, 600, 730, 800, 1100 | 6 |
| 5 | OPA14 | 730, 850, 1000, 1400,1500 | 5 |
| 6 | OPA15 | 500, 750, 1000 | 3 |
| 7 | OPA16 | 250, 480, 500, 550, 750, 800, 1000, 1450, 1500, 2000, 2100 | 12 |
| 8 | OPA 17 | 250, 400, 500, 250, 350, 480, 498, 500, 600, 750, 1250, 1500, 2000, 2200 | 11 |
| 9 | OPA 19 | 250, 400, 500, 600, 750, 1000, 1100, 1500, 2000 | 9 |

in two clusters was based on cutoff value of the average similarity at 0.79. Cluster A included 4/2 Izmir, 25 Mersin, 24/2 Izmir, 62 Adana, 19/1 Izmir genotypes. Cluster B comprised of the other satsumas.

Coletta et al. (1998) used RAPD markers to evaluate genetic similarity among 35 mandarin accessions, including 10 species and 7 hybrids. The genetic similarity within the mandarin group was found to be high, and suggests that cultivated mandarins have a narrow genetic base. Researchers propose that somatic mutations accounts for additional diversity within groups of cultivars, such as 'Satsuma. In their study, satsuma miyagawa wase, Owari, Wase and Unshiu Wase showed close genetic similarity in RAPD analysis.

Machado et al. (1995), random amplified polymorphic DNA (RAPD) analysis was carried out to evaluate polymorphism and genetic similarity between 39 Mediterranean mandarin genotypes. UPGMA cluster analysis



Figure 1. UPGMA dendrogram of 22 satsuma genotypes from RAPD data. Similarity values and matrix corellation are shown at the bottom of the dendrogram. (Matrix correlation: r = 0.82, 1:4/2 Izmir; 2: 26-Izmir; 3: 13/2 Izmir; 4: 27/1 Izmir; 5: 23Mersin; 6: 24 Mersin; 7: 3-Izmir; 8: 25Mersin; 9: 27/2 Izmir; 10: 8/3 Izmir; 11: 22-Izmir; 12: 61Adana; 13: 89 Adana; 14: 19/1 Izmir; 15: 11/1 Izmir; 16: Owari Satsuma; 17: 30-Izmir; 18: 18/1 Izmir; 19: 24/2 Izmir; 20: 62Adana; 21: 27Mersin; 22: 25/1 Izmir).

revealed the low level of genetic variation between accessions of Mediterranean mandarins, whereas their hybrids with other *Citrus* species showed greater genetic dissimilarity. Researchers claimed that twenty accessions yielded very similar patterns, suggesting either that they could be a single clone, or that the technique was not able to detect genomic variation. However, for the other specimens genetic polymorphism can easily be detected by RAPD, although the genetic variation between accessions was quite low.

SSR analysis

The fourteen SSR primer pairs used to characterize the Satsuma clones produced discrete reproducible fragments for genotypes tested. The number of fragments amplified from an individual mandarin clones with each primer pair ranged from one to two. Primer pair 497 was the most informative, producing four different sized fragments (Table 7).

The fourteen SSR primer pairs used on all Satsuma

| | SSR | Fragment size | Band number |
|---------|-----|--------------------|-------------|
| 458 | | 233, 238 | 2 |
| 571 | | 280, 285 | 2 |
| 497 | | 355, 357, 358, 362 | 4 |
| 506 | | 165, 167, 170 | 3 |
| 473 | | 220, 234 | 2 |
| 485 | | 216, 219 | 2 |
| 498 | | 362, 370 | 2 |
| MEST431 | | 346, 357 | 2 |
| 488 | | 160 | 1 |
| 173 | | 204 | 1 |
| 54 | | 173 | 1 |
| 121 | | 198 | 1 |
| 495 | | 272 | 1 |
| 697 | | 393 | 1 |

Table 7. Band size and band number of primers used for SSR analysis.

clones produced 25 fragments. Among 25 bands generated by 14 selected primer pairs, 68% were monomorphic that is present in all individuals. The mean number of putative alleles detected per primer pair in this study ranged from 1 to 4 (Table 2) with a mean value of 1.79 alleles per primer pair. This value is very low when compared with other SSR studies on plants (Aka-Kacar et al., 2006).

A dendrogram was constructed from the SSR data (Figure 2). The average similarity value 0.87 was used as a cutoff value for defining the clusters. Two major clusters were formed. Cluster B consisted of genotype "26 Izmir". Cluster A, the largest group, included the 21 genotypes and this group could be divided into two subgroups. 11/1 Izmir, control, 25/1 Izmir, 18/1 Izmir shared cluster A2 whereas the others take place cluster A1. Genetic similarity within the mandarin genotypes used in this study have a narrow genetic base. The dendrogram indicates that the genotypes can be separated into two major groups with a similarity value of 0.87 (A and B).

Based on 25 fragments, a similarity matrix was generated from the 22 Satsuma selections using the Dice coefficient of Nei and Li (1979). The data obtain from RAPD and SSR analyses were combined to perform genetic similarity. The dendogram constructed by UPGMA cluster analysis separated the Satsuma selections into two groups with a similarity value of 0.84 (Figure 3).

The group B occurred as a big group and divided into two groups in itself (Group B1 and Group B2). Group B2 was divided into two groups. The subgroup B.2.2. clustered into two groups. The subgroup B.2.2. contains only genotype 27 Mersin; the other subgroup B2.1. contains genotypes 13/2 Izmir, 30 Izmir, 18/1 Izmir, 3 Izmir, 61 Adana, 25/1 Izmir, 11/1 Izmir, 27/1 Izmir, 23 Mersin, 22 Izmir, 8/3 Izmir. The other subgroup A1 and A2 contains 4/2 Izmir ,25 Mersin, 24/2 Izmir, 62 Adana, 19/1 Izmir which are all similar patterns (Figure 3).

Group A also divided into two group and the A2 is consists of only "genotype 19/1 Izmir" and the other cluster (A1) has 4/2 Izmir, 25 Mersin, 24/2 Izmir, 62 Adana (Figure 3).

SSR markers could not distinguish between accessions within a few groups in which cultivars have arisen by apparent spontaneous mutation, such as sweet oranges (*Camellia sinensis*), Clementines (*Camellia reticulata*), Satsumas (*Camellia reticulata*), and small-fruited acid limes (*Cycloptera aurantifolia*) (Barkley et al., 2006).

Selection of new citrus genotypes collected as natural mutants from citrus orchards was previously reported (Becerra 1979; Robles-González et al., 1993; Uzun et al., 2009). Bud mutations is generally detected in horticultural traits and fruit characteristics in citrus. Contrasting with diversity for agronomic traits, vey low genetic variability has been found in cultivated citrus by use of molecular markers (Uzun et al., 2009).

Previous studies also have found few molecular polymorphisms within groups like these, consisting of cultivars developed by spontaneous mutation (Fang and Roose 1997; Bretó et al., 2001).

Even if mandarins exhibit much variation in characters (Reuther et al., 1967), they have narrow genetic variability. The genetic similarities between mandarins were found close together in other studies. Santos et al. (2003), reported in their molecular characterization studies with 34 mandarin genotypes that the genetic distance between the maximum of 0.32 (0.68 similarity level). Fang and Roose (1997), identified in their study with ISSR markers that five satsuma cultivars were in the same genetic structure. Uzun (2009), studied on the same selected satsuma clones to determine genetic similarity with SRAP markers, reported that polymorphism



Figure 2. Dendrogram of the genetic relationship of 22 Satsuma clones based on 22 SSR primer pairs. (1:4/2 Izmir; 2: 26-Izmir; 3: 13/2 Izmir; 4: 27/1 Izmir; 5: 23Mersin; 6: 24 Mersin; 7: 3-Izmir; 8: 25Mersin; 9: 27/2 Izmir; 10: 8/3 Izmir; 11: 22-Izmir; 12: 61Adana; 13: 89 Adana; 14: 19/1 Izmir; 15: 11/1 Izmir; 16: Owari Satsuma; 17: 30-Izmir; 18: 18/1 Izmir; 19: 24/2 Izmir; 20: 62Adana; 21: 27Mersin; 22: 25/1 Izmir).

value of the satsuma clones were 0,97-1,00.

RAPD amplification and SSR loci analyses revealed low genetic polymorphism in the grapefruit accessions studied. Varieties of great economic importance and distinct morphological characteristics (for example, pigmented pulp, pale yellow pulp, seeded and seedless fruits) such as 'Marsh Seedless', 'Duncan', 'Thompson Pink', 'Foster' and 'Red Blush' showed complete genetic similarity. These results suggest that these accessions represent variations of a single clone with different names or that they are in fact different varieties that were derived from somatic mutations that were not detected by the molecular markers used (Corarza-Nunes et al.,2002).

Biotechnological methods are offering several opportunities that can be used in breeding Citrus and relatives, the first step of improving genotypes is to assess the genetic structure of genotypes (Aka-Kacar et al., 2005). Though, morphological and isozyme markers have been employed in assessing the underlying genetic

variation of a genotype, the accuracy of the assessment is questionable. The availability of a low number of morphological and biochemical markers, their poor or unknown genetic control, environmental influence on the phenotypic expression, stage specific identification and procedural difficulties are known impediments in using these as genetic markers in genetic diversity analysis. Considering the problems associated with morphological and isozyme markers, researchers searched for alternative tools (Ravi et al., 2003). This study employed SSR and RAPD markers to evaluate genetic polymorphisms between the Satsuma genotypes which were obtained from our selection breeding program.

The fourteen SSR and nine RAPD primer pairs were not able to differentiate all Satsuma selections. These selections have slight morphological differences and may represent sports caused by somatic mutations selected during vegetative propagation from an original cultivar. The genotypes yielded similar patterns, suggesting either



Figure 3. Dendrogram of the genetic relationship of 22 Satsuma clones based on 9 RAPD and 22 SSR primer pairs. (1:4/2 Izmir; 2: 26-Izmir; 3: 13/2 Izmir; 4: 27/1 Izmir; 5: 23Mersin; 6: 24 Mersin; 7: 3-Izmir; 8: 25Mersin; 9: 27/2 Izmir; 10: 8/3 Izmir; 11: 22-Izmir; 12: 61Adana; 13: 89 Adana; 14: 19/1 Izmir; 15: 11/1 Izmir; 16: Owari Satsuma; 17: 30-Izmir; 18: 18/1 Izmir; 19: 24/2 Izmir; 20: 62Adana; 21: 27Mersin; 22: 25/1 Izmir).

they are identical genotypes or we did not have enough markers to detect variation between these genotypes. Our preliminary result indicates that SSR and RAPD markers are useful tools for Satsuma clone varietal identification. But this marker system was not enough to distinguish the genotypes.

In conclusion, these results demonstrated the importance of the evaluation of new natural selections of Satsuma mandarins in order to know their field performance. Among the new 21 satsuma mandarin selections several tree growth, fruit yield, fruit quality, and molecular profile differences were observed under ecological conditions of Adana along three years. According to the results; 30 Izmir, 4/2 Izmir, 62 Adana and 13/2 Izmir clones were found to have better fruit quality and the higher yield than the control. The yield differences among selections began to be more evident since eleven year after planting. Since genotypes showed a real performance in field due to juvenility. As a result, these selections (30 Izmir, 4/2 Izmir, 62 Adana and 13/2 Izmir clones) can be suggested as new satsumas in citrus growing area.

REFERENCES

- Abkenar AA, Isshiki S, Tashiro Y (2004). Phylogenetic relationships in the true citrus fruit trees revealed by PCR-RFLP analysis of cpDNA. Scientia Horticulturae, 102(2): 233-242.
- Aka-Kacar Y, Demirel A, Tuzcu O, Yesiloglu T, Ulas M, Yildirim B (2005). Preliminary results on Fingerprinting of Lemon genotypes tolerant to Mal Secco Disease by RAPD Markers, Biologia, 60: 295-300.
- Aka-Kacar Y, Cetiner S, Cantini C, Iezzoni AF (2006). Simple sequence repeat (SSR) markers differentiate Turkish sour cherry germplasm, J. Am. Pomological Soc. 60(3): 136-143.
- Alberto F, Santos R, Leitao JM (1997). DNA extraction and RAPD markers to assess the genetic similarity among Gelidium sesquipedale (Rhodophyta) polpulations. J. Phycol. 33(4): 706-710.

Baig MN, Yu A, Guo W, Deng X (2009). Construction and characterization of two Citrus BAC libraries and identification of clones containing the phytoene synthase gene. Genome, May, 52(5):

484-489.

- Barkley NA, Mikeal L, Roose ML, Krueger C, Federici T (2006). Assessing Genetic Diversity and Population Structure in a Citrus Germplasm Collection Utilizing Simple Sequence Repeat Markers (SSRs). Theor. Appl. Genet. 112: 1519-1531.
- Becerra RS (1979). Un clon sin espinas de limón mexicano (*Citrus aurantifolia* Swingle). Agricultura Técnica en México, pp. 65-71.
- Bretó M, Ruiz P, Pina A, Asíns MJ (2001). The diversification of Citrus clementina Hort. ex Tan. a vegetatively propagated crop species. Mol. Phylog. Evol. 21: 285-293.
- Chaparro JX, Werner DJ, O'Malley D, Sederoff RR (1994). Targeted mapping and linkage analysis of morphological, isozyme and RAPD markers in peach. Theor. Appl. Genet. 87: 805-815.
- Coletta HD, Machado MA, Targon MLPN, Moreira MCPQDG (1998). Analysis of the genetic diversity among mandarins (*Citrus* spp.) using RAPD markers. Euphytica, 102(1): 133-139.
- Corarza-Nunes MJ, Machado MA, Nunes WMC, Cristofani M, Targon MLPN (2002). Assessment of genetic variability in grapefruits (*Citrus paradisi* Macf.) and pummelos (*C. maxima (Burm.) Merr.*) using RAPD and SSR markers. Euphytica, 126: 169-176.
- Dellaporta SL, Wood J, Hicks JB (1983). A plant DNA minipreparation: Version II Plant Mol. Biol. Rep. Vol. 1, Number 4, Springer Netherlands.
- Deng ZN, Gentile A, Nicolosi E, Domina F, Vardi A, Tribulato E (1995). Identification of in vivo and in vitro lemon mutants by RAPD markers. J. Hortic. Sci. 70: 117-125.
- Fang DQ, Roose ML (1997). Identification of closely related citrus cultivars with inter-simple sequence repeat markers. Theor. Appl. Genet. 95(3): 408-417.
- Fang DQ, Roose ML, Krueger RR, Federici CT (1997). Fingerprinting trifoliate orange germ plasm accessions with isozymes, RFLPs, and inter-simple sequence repeat markers. Theor. Appl. Genet. 95(1-2): 211-219.
- Federici CT, Fang DQ, Scora RW, Roose ML (1998). Phylogenetic relationships within the genus Citrus (Rutaceae) and related genera as revealed by RFLP and RAPD analysis. Theor. Appl. Genet. 96(6-7): 812-822.
- Green RM, Vardi A, Galun E (1986). The plastome of Citrus. Physical map, variation among Citrus cultivars and species, and comparison with related genera. Theor. Appl. Genet. 72: 170-177.

http://www.faostat.org, 2010. Last Access on July 2010.

- Huff DR, Peakall R, Smouse PE (1993). RAPD variation within and among natural populations of outcrossing buffalo grass [Buchloë dactyloides (Nutt.) Engelm.]. 1993. Theor. Appl. Genet. 86: 927-934.
- Hvarleva T, Kapari-Isaia T, Papayiannis I, Atanassov A, Hadjinicoli A, Kyriakou A (2008). Characterization of Citrus Cultivars and Clones in Cyprus through Microsatellite and RAPD Analysis. Biotechnol. Biotechnolological Equipment, 22: 787-794.
- JinPing X, LiGeng C, Ming X, HaiLin L, WeiQi Y (2009). Identification of AFLP fragments linked to seedlessness in Ponkan mandarin (Citrus reticulata Blanco) and conversion to SCAR marker. Scientia Horticulturae, 121(4): 505-510.
- Khadari B, Lashermes PH, Kjellberg F (1995). RAPD fingerprints for identification and genetic characterization of fig (*Ficus carica* L.) genotypes. J. Genet. Breed. 49: 77-86.
- Kijas JMH, Fowler JCS, Thomas MR (1995). An evaluation of sequence tagged microsatellite site markers for genetic analysis within Citrus and related species. Genome, 38: 349-355.
- Köse H (2000). Features of Some important satsuma clones under Adana ecological conditions in 1998. University of Çukurova, Faculty of Agriculture, Department of Horticulturae. Bachelor thesis. P. 35. (in press).
- Lacis G, Rashal I, Ruisa S, Trajkovski V, lezzoni AF (2009). Assessment of genetic diversity of Latvian and Swedish sweet cherry (*Prunus avium* L.) genetic resources collections by using SSR (microsatellite) markers. Scientia Horticulturae, 121(4): 451-457.

Lin SY, Chen IZ (2006). Chen Rapd profiling of four native citrus species in taiwan (). XXVII International Horticultural Congress-IHC: International Symposium on Citrus and Other Tropical and Subtropical Fruit Crops. ISHS Acta Horticulturae, p. 773.

- Lu Z, Reighard GI, Baird WV, Abbott AG, Rajapakse S (1996). Identification of peach rootstock cultivars by RAPD markers. Hort. Sci. 31: 127-129
- Machado MA, Coletta-Filho HD, Targon MLPN, Pompeu JJR (1996). Genetic relationship of Mediterranean mandarins (*C. deliciosa* Ten.) using RAPD markers. Euphytica, 92: 321-326.
- Moore GA (2001). Oranges and lemons: clues to the taxonomy of Citrus from molecular markers. Trends Genet. 17(9): 536-540.
- Nei M, Li WH (1979). Mathematical models for studying genetic variation in terms of restriction endonucleases. Proc. Natl. Acad. Sci. USA, 76: 5269-5273.
- Nair S, Kumar A, Srivastava MN, Mohan M (1996). PCR-based DNA markers linked to a gall midge resistance gene, Gm4t, has potential for marker-aided selection in rice. Theor. Appl. Genet. 92(6): 660-665.
- Nicese FP, Hormaza JI, McGranahan GH (1998). Molecular characterization and genetic relatedness among walnut (*Juglans regia* L.) genotypes based on RAPD markers. Euphytica, 101: 199-206.
- Nicolisi ZN, Deng A, Gentil S, Lamalfa G, Tribulato E (2000). Citrus phylogeny and genetic origin of important species as investigated by molecular markers, Theor. Appl. Genet. 100(2000): 1155-1166.
- Nishiura M (1964). Citrus Breeding And Bud Selection In Japan. http://www.fshs.org/Proceedings/Password%20Protected/1964%20V ol.%2077/79-83%20(NISHIURA).pdf
- Oliveira RP, Cristofani M, Vildoso CIA, Machado MA (2004). Genetic linkage maps of 'Pêra' sweet orange and 'Cravo' mandarin with RAPD markers. Pesquisa Agropecuária Brasileira, 39: 159-165.
- Özsan M, Tuzcu O, Akteke SA, Inci HB, Çelikel K, Özdemir E, Çimen I (1986). Budwood selection -certification and breeding in citrus. Derim 3(4): 147-156.
- Pang XM, Hu CG, Deng XX (2007). Phylogenetic relationships within Citrus and its related genera as inferred from AFLP markers. Genet. Resour. Crop Evol. 54(2): 429-436.
- Rao MN, Soneji JR, Chen CX, Huang S, Gmitter FG (2008). Characterization of zygotic and nucellar seedlings from sour orangelike citrus rootstock candidates using RAPD and EST-SSR markers. Tree Genet. Genomes. 4(1): 113-124.
- Ravi M, Geethanjali S, Sameeyafarheen F, Maheswaran M (2003). Molecular Marker based genetic diversity analysis in rice (*Oryza sativa* L.) using RAPD and SSR markers. Euphytica, 133: 243-252.
- Reuther W, Batchelor LD, Webber HJ (1967). The Citrus Industry. 1: 500-505.
- Robles-Gonzales MM, Medina-Urrutia VM, Velazquez-Monreal JJ, Simpson J (2008). Field performance and molecular profiles of Mexican lime selections. Euphytica, 161: 401-411.
- Santos PK, Cunha-Dornelles AL, Loreta Brandão de Freitas L (2003). Characterization of mandarin citrus germplasm from Southern Brazil by morphological and molecular analyses. Pesq. agropec. Bras. vol. 38, no. 7.
- Urgun Ş (1997). Pomological characteristics of some mandarin varieties under Adana ecological conditions. Master Thesis. University of Çukurova, Institute of Natural and Applied Sciencies. p. 253. (in pres).
- Uzun A (2009). Characterization of genetic diversity of citrus by SRAP markers. Ph.D. Thesis. University of Çukurova, Institute of Natural and Applied Sciencies. p. 365.
- Uzun A, Gulsen O, Kafa G, Seday U (2009). Field performance and molecular diversification of lemon selections. Source: Scientia Horticulturae, 120(4): 473-478.
- Wei J (2007). Characterization of retrotransposon elements and development of related molecular markers in citrus. PhD Thesis, Huazhong Agricultural University, Wuhan, China
- Williams JGK, Kubelik AR, Livak KJ, Rafalski JA, Tingey SV (1990). DNA polymorphisms amplified by arbitrary primers are useful as genetic markers. Nucleic Acids Res. 18: 6531-6535.