The Genetic Variability of Sicilian Lemon Germplasm Revealed by Molecular Marker Fingerprints

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Abstract. There is a high level of diversity among lemons [Citrus limon (L.) Burm. f. (2n = 2x = 18)] in Sicily, where each growing area has a wide range of landraces mostly derived from bud mutation. Because this variability represents an important resource for future breeding programs and genetic improvement, the relationships among the principal 36 accessions of Sicilian lemon, belonging to three different cultivars (Femminello, Monachello, and Lunario), were examined by intersimple sequence repeat and random amplified polymorphic DNA markers. Three ‘Femminello’ accessions from nearby Italian regions were also examined to study the genetic flow from the continent. The disputed case of the accession ‘Eureka Messina lemon’ was also examined, using ‘Frost Eureka’ as a control. Our results confirmed the extreme polymorphic nature of the three principal Sicilian cultivars and the presence of a wide range of different genotypes. Twenty-two Sicilian genotypes were recognized as unique accessions, reflecting the richness of the lemon germplasm present in Sicily. Each growing area showed the presence of several genetically different landraces, probably preserved by genetic isolation, whereas the continental accessions appeared extremely similar to the island genotypes, showing an exchange of germplasm from the island to the continent.

Lemon is an economically important crop widely grown in southern Italy and especially in Sicily, the first Italian region for lemon production (Cottone et al., 1996; Forte, 1999). While in the other principal producing countries one or few cultivars prevail over the others (‘Eureka’ and ‘Lisbon’ in the United States, ‘Genoa’ and ‘Limeira Lisbon’ in Argentina, ‘Galego’ in Brazil and ‘Berna’ in Spain; Russo and Spina, 1985; Saunt, 2000), in Italy, each lemon-cultivating area (Palermo, Catania, Siracusa, Messina, Reggio Calabria, Salerno, and Naples) has a wide range of landraces mostly derived from bud mutation from the cultivar Femminello (Russo and Spina, 1985).

As the most domesticated fruit crop species over the past century, the Italian lemon has also suffered a dramatic reduction in the gene pool due to the best characteristics (high yields, good fruit size, high juice content, low seededness) of some grown cultivars and the presence of “mal secco” disease [citrus wilt [Phoma tracheiphila (Petri) L.A. Kantachveli and Gikachvili]], which allowed only tolerant cultivars (such as Monachello) to survive (Crescimanno et al., 1992). Therefore, some landraces with interesting breeding characteristics are at risk of disappearance.

In recent years, the collection and the characterization of fruit crop germplasm has become a common concern among geneticists and breeders to identify and preserve the genetic diversity of species and for setting up new breeding programs aimed at genetic improvement.

Currently in Sicily, there are several lemon landraces with interesting agronomic traits, but little information is available about their genetic variability. Until now, Sicilian lemon characterization has been done only through phenotypic analysis (Crescimanno et al., 1992). This characterization requires a large set of data, which is often difficult to access, and in some cases, dependent upon environmental conditions.

Polymerase chain reaction (PCR)-based DNA marker technology has already become a useful tool in characterizing lemon cultivars. In particular, PCR methods using arbitrary primers have become very popular, as they do not require any information about DNA sequences in addition to being cost- and time-effective (Fang and Roose, 1997).

Fang and Roose (1997) used intersimple sequence repeat (ISSR) to discriminate between five genotypes of the cultivars Eureka and Lisbon, whereas Gulsen and Roose (2001a) used ISSR, microsatellites, and isozymes to assess diversity and phylogenetic relationships in several lemon accessions and related taxa. Recently, 10 autochthonous lemon cultivars of the Italian region Campania were identified through ISSR (Capparella et al., 2004) and random amplified polymorphic DNA (RAPD) techniques (Marinelli et al., 2004).

Phylogenetic relationships between lemon and other Citrus L. species and Citrus-related genera have been also investigated by SSR (Barkley et al., 2006), RAPD, and restriction fragment length polymorphism (Federici et al., 1998; Nicolosi et al., 2000).

In the present study, 41 phenotypically diverse lemon accessions, mostly collected from different parts of Sicily, were analyzed by ISSR and RAPD methods to detect genetic polymorphisms useful for setting up a molecular reference system that would allow precise identification. Data obtained from this study have been also used to provide much information about genetic relationships among the accessions examined. This information, with morphological and phenological descriptors, could be useful in assessing the basis of breeding programs aimed at the genetic improvement of lemons.
Materials and Methods

PLANT MATERIALS AND DNA EXTRACTION. A total of 41 plants of *C. limon*, including 34 ‘Femminello’, 3 ‘Monachello’, 2 ‘Lunario’, and 2 ‘Eureka’ accessions were used in the investigation. The sampling sites, the codes, and the more remarkable quality traits are listed in Table 1. The terms ‘Femminello’, ‘Monachello’, and ‘Lunario’ commonly are used to identify different cultivars, each constituted from a phenotypically diverse, wide range of plants. Thirty-one ‘Femminello’, three ‘Monachello’, and two ‘Lunario’ accessions differing in morphological and physiological traits were collected from several sites located in the major *Citrus* diffusion areas of Sicily. The localization of the sampling sites of Sicilian plants is shown in Fig. 1. Two ‘Femminello’ plants, ‘Femminello a foglia larga’ (FW) and ‘Femminello Favazzina’ (FV), were collected from Calabria region, the closest peninsular Italian region to Sicily, whereas a ‘Femminello common’ (FC) was collected from Salerno (Campania region). Among the two ‘Eureka’ accessions, only ‘Frost Eureka’ (EF) was certified, and it was used as external control. The ‘Eureka Messina lemon’ (EM) was a putative ‘Eureka’ plant collected from a grower in the Siracusa area. All the plants were introduced in the germplasm collection in the Lascari field station (lat. 38°N, long. 14°E).

Genomic DNA was extracted from leaves as described by Doyle and Doyle (1987). The leaves were collected from mature trees present in the germplasm collection and were carefully washed (as advised to avoid insects and fungal contamination; Fang and Roose, 1997), frozen in liquid nitrogen, and stored at −80 °C. The samples were ground in a mortar with liquid nitrogen. DNA was quantified by measuring OD<sub>260</sub> as described by Sambrook et al. (1989).

ISSR ANALYSIS. A total of 20 primers [i.e., (AC)<sub>3</sub>YAC, (AG)<sub>3</sub>YGC, (AC)<sub>2</sub>YAC, (AC)<sub>3</sub>YG, (GT)<sub>3</sub>YAC, (TCC)<sub>3</sub>YAC, (GA)<sub>3</sub>YAC, (CA)<sub>3</sub>YAC, and (GA)<sub>3</sub>YAC, reported by Fang and Roose, 1997, and ENEA13, ENEA14, ENEA21, ENEA34, ENEA36, ENEA45, and ENEA47 (Table 2) given by S. Lucretti of the Biotechnology and Agriculture Division (Ente per le Nuove tecnologie, l’Energia e l’Ambiente, Centro Ricerche Casaccia, Rome) were used to amplify the DNA. The ISSR primers have been selected on the basis of their usefulness in published data (Fang and Roose, 1997; Siragusa et al., 2006, 2007) or previous experience. The primers were purchased from MWG-Biotech AG (Ebersberg, Germany).

Each amplification was performed in a 25-μL reaction volume containing 20 mM Tris-HCl (pH 8.4), 50 mM KCl, 2 mM MgCl<sub>2</sub>, 800 μM dNTP (dATP, dTTP, dCTP, dGTP in 1:1:1:1 ratio), 1.5 μM of each primer, 1 μL of Platinum Taq polymerase (Invitrogen, Carlsbad, CA), and 30 ng of template DNA. The amplification was performed in a 96-well GeneAmp PCR System 9700 (Applied Biosystems, Foster City, CA) equipped with a Hot Bonnet under the following cycle program: initial denaturation step for 4 min at 94 °C, followed by 36 cycles at 94 °C for 30 s (denaturation), 47 °C to 60 °C for 45 s (annealing), and 72 °C for 120 s (extension), followed by a final extension step at 72 °C for 7 min. PCR-amplified DNA fragments were separated on a 1.5% agarose gel containing 1× TBE (45 mM Tris-borate and 1 mM EDTA) and 0.5 μg mL<sup>−1</sup> aqueous solution of ethidium bromide. About 25 μL of reaction products with 5 μL of loading buffer (0.25% BFB, 40%, w/v, sucrose) were loaded onto the gel, which was run for 4 h at 100 V. The gel was then visualized with a ultraviolet transilluminator at 300 nm. To confirm the reproducibility of the banding patterns, the PCR experiments were repeated three times.

RAPD ANALYSIS. Twenty-two arbitrary decamer primers i.e., OP01, OPAT 04, OPAT14, OP04, OPH15, OPK04, OP04, OPM06, OPN07, OPN14, OPQ14, and P130 (reported by Coletta Filho et al., 1998) and UBC219, UBC234, UBC237, UBC239, UBC247, UBC251, UBC264, UBC266, and UBC272 (reported by Wang et al., 1999) were used for the amplification of DNA sequences. The primers were purchased from MWG-Biotech AG.

DNA amplification reactions were performed in a volume of 25 μL with 20 mM Tris-HCl (pH 8.4), 50 mM KCl, 3 mM MgCl<sub>2</sub>, 800 μM dNTP, 0.4 μM of each primer, 1 U of Platinum Taq polymerase (Life Technologies, Grand Island, NY), and 30 ng of template DNA. The amplification was performed in a 98-Well GeneAmp PCR System 9700 thermocycler (Applied Biosystems) equipped with a Hot Bonnet under the following cycle program: initial denaturation step for 90 s at 94 °C, followed by 40 cycles at 94 °C for 1 min (denaturation), 35 °C for 2 min (annealing), and 72 °C for 2 min (extension), followed by a final extension step at 72 °C for 10 min. PCR-amplified DNA fragments were visualized as described above. To confirm the reproducibility of the banding patterns, all analyses were repeated three times.

DATA ANALYSIS. Amplified bands from each primer were scored as present (1) or absent (0) for all cultivars studied. Only those bands showing consistent amplification were considered; smeared and weak bands were excluded from the analysis. Dice’s (1945) coefficient of similarity (*D<sub>D</sub>*) was determined between each pair of accessions. Dice’s coefficient has been recommended for the evaluation of genetic similarities when using RAPD data (Lamboy, 1994). The genetic distance (*GD*) between two samples was calculated as: *GD* = 1 − *D<sub>D</sub>*. The estimates of similarity between cultivars were then used for cluster analysis by unweighted pair group method of arithmetic average (UPGMA; Sneath and Sokal, 1973) using NTSYS, version 2.02, for Windows (Rohlf, 1994). The Mantel test of genetic and geographic (Euclidian) distances was carried out using 250 permutations.

Additional statistics were computed to estimate the grade of polymorphism among cultivars studied. The average marker allele frequency (*p*) for each primer and among all primers was calculated for each single accession and among all accessions. Genetic diversity (*H*) of Nei (1973) and Shannon index (*S*) (Lewontin, 1972) were used to summarize the data for molecular markers, and their standard deviations (SD) were indicated. The percentage of polymorphisms (*Pp*) was given as number of polymorphic loci/number of total loci, regardless of allele frequencies. The fixation index (*G<sub>ST</sub>*) was computed as a proportion of genetic diversity expressed between accessions coming from different geographic regions. From this, the gene flow (*Nm*) was derived and interpreted according to McDermott and McDonald (1993). All calculations and analyses were conducted using POPGENE, version 1.31 (Yeh et al., 1999).

Results

A total of 558 reproducible and well-resolved band classes were observed in all lemon plants; 120 loci of these (*Pp* = 21.5%) appeared polymorphic. When the analysis was carried out only for Sicilian accessions (excluding also the disputed
Table 1. Lemon accessions analyzed and relatively more remarkable quality traits (Crescimanno et al., 1992).

| Accession                      | Sampling site | Code | Remarkable quality traits                      |
|--------------------------------|---------------|------|------------------------------------------------|
| Femminello comune              | Palermo       | FC1  | High juice production (33.8%) (7.6.7.-7)        |
| Femminello comune              | Palermo       | FC2  | High juice production (38.3%) (7.6.7.-7)        |
| Femminello Lo Porto            | Palermo       | FL   |                                                 |
| Femminello comune              | Santa Flavia (PA) | FC3  | High juice production (43.7%) (7.6.7.-7)        |
| Femminello verdellifero        | Santa Flavia (PA) | FV1  | High juice production (33.3%) (7.6.7.-7)        |
| Femminello femminino           | Santa Flavia (PA) | FF   | Low seeds number per fruit (4.3) (7.7.1.-2); high juice production (34.4%) (7.6.7.-7) |
| Femminello masculino           | Santa Flavia (PA) | FM   | High juice production (39.1%) (7.6.7.-7)        |
| Femminello Pozzetti            | Lascari (PA)  | FP   | Citrus wilt tolerance (10.12.11.1)              |
| Femminello comune              | Mandanici (ME) | FC4  | High acidity juice (7.80) (8.5.1)               |
| Femminello comune              | Roccalumera (ME) | FC5  | High acidity juice (7.30) (8.5.1)               |
| Femminello paddaru             | Roccalumera (ME) | FD   | High juice production (35.2%) (7.6.7.-7)        |
| Femminello cuduzzaro           | Roccalumera (ME) | FU1  | High acidity juice (7.17) (8.5.1)               |
| Femminello palermitano         | Roccalumera (ME) | FH   |                                                 |
| Femminello familario a foglia larga | Roccalumera (ME) | FF1  | High acidity juice (40.6%) (7.6.7.-7); high acidity juice (7.23) (8.5.1) |
| Femminello familario a foglia stretta | Roccalumera (ME) | FF2  | High juice production (34.7%) (7.6.7.-7); high acidity juice (7.63) (8.5.1) |
| Femminello Germanà             | Capo d’Orlando (ME) | FG   | Citrus wilt tolerance (10.12.11.1)              |
| Femminello Pettineo            | Pettineo (ME) | FO   | Low seeds number per fruit (2.2) (7.7.1.-1); high juice production (41.8%) (7.6.7.-7); high acidity juice (7.27) (8.5.1) |
| Femminello famularo             | Rometta (ME)  | FF3  | Citrus wilt tolerance (10.12.11.1)              |
| Femminello cucuzzaro           | Brolo (ME)    | FU2  | Citrus wilt tolerance (10.12.11.1)              |
| Femminello famularo            | Brolo (ME)    | FF4  | Citrus wilt tolerance (10.12.11.1)              |
| Femminello precoce             | Brolo (ME)    | FE   |                                                 |
| Femminello sfusato             | Brolo (ME)    | FN   | High juice production (40.6%) (7.6.7.-7); high acidity juice (7.23) (8.5.1) |
| Femminello verdellifero        | Brolo (ME)    | FV2  | High acidity juice (7.10) (8.5.1)               |
| Femminello tondo precoce       | Brolo (ME)    | FT   |                                                 |
| Femminello verdellifero        | Giarre (CT)   | FV3  |                                                 |
| Femminello Conti               | Riposto (CT)  | FC   |                                                 |
| Femminello innesto palermitano | Mascalci (CT) | F1   | Low seeds number per fruit (2.2) (7.7.1.-1); high juice production (37.4%) (7.6.7.-7); high acidity juice (7.13) (8.5.1) |
| Femminello comune              | Catania (CT)  | FC6  | High juice production (40.1%) (7.6.7.-7)        |
| Femminello fior d’arancio      | Catania (CT)  | FA   | High juice production (33.2%) (7.6.7.-7); citrus wilt tolerance (10.12.11.1) |
| (or zagara bianca)             | Siracusa (SR) | FS   | Low seeds number per fruit (4.6) (7.7.1.-2); high juice production (43.9%) (7.6.7.-7); high acidity juice (7.03) (8.5.1) |
| Monachello non rifiorente      | Fiumefreddo (ME) | MN   | Low seeds number per fruit (4.2) (7.7.1.-2); citrus wilt tolerance (10.12.11.1) |
| Monachello Sant’Antonio        | Messina       | MS   | Low seeds number per fruit (4.8); citrus wilt tolerance (10.12.11.1) |
| Feinmellino - Monachello quattrocchi | Catania        | MQ   | Citrus wilt tolerance (10.12.11.1)              |
| Lunario                        | Villagrazia (PA) | LV   | Low seeds number per fruit (3.9) (7.7.1.-1); high acidity juice (7.20) (8.5.1) |
| Lunario spinoso                | Fiumefreddo (ME) | LT   |                                                 |
| Femminello Favazzina           | Reggio Calabria | FV   | High juice production (34.0%) (7.6.7.-7); high acidity juice (7.50) (8.5.1) |
| Femminello a foglia larga      | Reggio Calabria | FW   |                                                 |
| Femminello comune              | Salerno       | FC7  |                                                 |
| Frost Eureka                   | Riverside (U.S.A.) | EF   |                                                 |
| Eureka Messina lemon           | Messina       | EM   |                                                 |

*The province of the sampling site is showed in brackets; CT = Catania, ME = Messina, PA = Palermo, SR = Siracusa.
*International Plant Genetic Resources Institute code of the quality traits is signed in bold type.
The 20 ISSR primers used in this analysis gave rise to 251 well-resolved band classes, ranging from 200 bp [primer \((GA)_{8}YG\)] to 2.5 Kb [primer \((CA)_{8}RG\)] in size. The number of ISSR bands obtained for each primer varied from 8 [primer \((AG)_{8}YG\)] to 18 [primer \((AC)_{8}YT\)], with an average of 12.1 bands per primer. Fifty-two polymorphic band classes (9.3% of the total amplified band classes) were identified with ISSR primers, with an average of 2.6 polymorphic band classes per primer. Some polymorphisms identified with the ISSR primer ENEA34 are shown in Fig. 2.

The RAPD primers produced 307 reproducible band classes, ranging from 300 bp (primer OPN14) to 2.5 kb in size (primer OPM04). The number of RAPD bands obtained for each primer varied from 9 (with the primer OPO14) to 18 (with the primer OPM04), with an average of 13.4 bands per primer. Sixty-eight polymorphic band classes (12.2% of the total amplified band classes) were identified with RAPD primers, with an average of 3.1 polymorphic band classes per primer. Some polymorphisms identified with the RAPD primer OPM04 are shown in Fig. 3. 

Fig. 1. Geographic localization of sampling sites of Sicilian lemon accessions analyzed.

Table 2. Sequences and annealing temperatures of ENEA primers used for ISSR analysis.

| Primer name | Primer sequence (5'-3') | Annealing temp. (°C) |
|-------------|-------------------------|---------------------|
| ENEA7       | GGTC(CA)_{7}            | 56                  |
| ENEA8       | CAGC(AC)_{7}            | 56                  |
| ENEA12      | CCAT(GT)_{7}            | 53                  |
| ENEA13      | GCA(AC)_{7}             | 53                  |
| ENEA14      | GGG(AC)_{7}             | 53                  |
| ENEA21      | (GA)_{6}GG              | 54                  |
| ENEA34      | (ACC)_{6}CC             | 56                  |
| ENEA36      | CC (ATG)_{6}            | 56                  |
| ENEA45      | AA(CT)_{8}              | 52                  |
| ENEA47      | AG(CA)_{8}              | 53                  |

‘Eureka Messina lemon’), the percentage of polymorphic loci was reduced to 17.9%.

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The RAPD primers produced 307 reproducible band classes, ranging from 300 bp (primer OPN14) to 2.5 kb in size (primer OPM04). The number of RAPD bands obtained for each primer varied from 9 (with the primer OPO14) to 18 (with the primer OPM04), with an average of 13.4 bands per primer. Sixty-eight polymorphic band classes (12.2% of the total amplified band classes) were identified with RAPD primers, with an average of 3.1 polymorphic band classes per primer. Some polymorphisms identified with the RAPD primer OPM04 are shown in Fig. 3.

Dice’s similarity coefficient was used to carry out cluster analysis and to generate a dendrogram showing the relationships among the selected accessions (Fig. 4). The dendrogram showed two main groups, one including most ‘Femminello’ accessions (group 1) and another including the accessions ‘Monachello Sant’ Antonio’ (MS), ‘Lunario’ (LV), ‘Femminello femminino’ (FF), and ‘Femminello fumaluro a foglia larga’ (FF1; group 2), having a genetic similarity of 0.97. In group 1, accessions ‘Frost Eureka’ and ‘Eureka Messina lemon’ appeared different and stood alone, close to the root of the relationship group. Subgroup 1A, having a genetic similarity higher than 0.99, included only the ‘Femminello’ accessions and contained all ‘Femminello’ from other Italian regions, ‘Femminello comune’ from Campania (FC7), ‘Femminello a foglia larga’ (FW), and ‘Femminello Favazzina’ (FV) from Calabria. In this subgroup, 10 accessions, including ‘Femminello comune’ from Salerno (FC7), were not distinguished, showing the typical Sicilian ‘Femminello’ profile. Similarly, ‘Femminello comune’ from Mandanici (FC4) with ‘Femminello comune’ from Roccalumera (FC5), ‘Femminello verdellifero’ from Brolo (FV2) with ‘Femminello verdellifero’ from Giarre (FV3), and ‘Femminello Pettineo’ (FO) with ‘Femminello Favazzina’ (FV) appeared indistinguishable. The remaining accessions were separated from each other, clustering in groups that cannot be explained by their likely area of origin. The Mantel test was carried out to confirm this hypothesis. A very low normalized Mantel statistic Z value was obtained (matrix correlation r = 0.11), with a probability random $Z_{\text{observed}} = 0.50$.

The total Nei’s genetic diversity ($H_T$) for Sicilian accessions was $0.037 \pm 0.002$. It was lower if calculated only for ‘Femminello’ accessions ($0.019 \pm 0.006$). An additional measure for genetic variation was obtained from the Shannon index, a genetic diversity index suitable when dominant markers are used (Dawson et al., 1995). $S$ value was $0.060 \pm 0.005$ for all Sicilian accessions analyzed, and it was $0.034 \pm 0.008$ when only ‘Femminello’ accessions were considered.

An analysis of gene diversity in Sicilian ‘Femminello’ accessions was carried out in the three principal growing areas (Palermo, Messina, and Catania). The accession ‘Femminello siringo’ (FS), unique to the Siracusa zone, was excluded from this analysis.

Low gene diversity was found in all three regions analyzed, with the greatest value in the area of Palermo ($h = 0.020 \pm 0.087$), followed by Messina ($h = 0.016 \pm 0.069$) and Catania ($h = 0.011 \pm 0.059$). The greatest number of polymorphic loci was identified in Messina accessions ($Pp = 8.5\%$), whereas Palermo
...and Catania areas showed lower values (Pp = 5.8% and 4.5%, respectively). The average variation within samples H̄ was 0.012 ± 0.001.

Genetic distances between accessions coming from Palermo and Messina was the lowest (0.003), whereas the GDs between Catania accessions and Palermo and Messina accessions were 0.006 and 0.004, respectively.

The fixation index (GST = 0.42) showed that the observed genetic variability could be attributed slightly more to within-area accession differences than to differences among the three areas. As expected, the estimated gene flow was extremely low (Nm = 0.69).

Some markers were found to be associated with the different areas. The 750-bp fragment amplified from the RAPD primer OPH15 was principally present in the Messina area accessions, whereas an 1850-bp fragment amplified from the RAPD primer OPH04 was absent only in Messina accessions. Three fragments were commonly amplified in the accessions growth on the North coast of the island: the 550-bp fragment amplified from the RAPD primer OPH04, and the 650- and 1150-bp fragments from the ISSR primers (AC)8YG and (GA)8YC, respectively. The 1150-bp fragment was also amplified in all accessions from Reggio Calabria, demonstrating the most probable route of gene flow from the island to the continent.

Discussion

Lemon characterization studies have already been carried out to examine the genetic diversity among the most important cultivars in some growing areas and to detect phylogenetic relationships between lemon and other Citrus species and Citrus-related genera.

The use of molecular markers surely gave important guidance to these studies. Microsatellites, ISSR, and RAPD were successfully used to show relationships among individuals of the same or closely related species and also among more distantly related taxa.

In all the studies of phylogenetic relationships within the genus Citrus, lemon cultivars always showed high levels of heterozygosisty (Barkley et al., 2006; Federici et al., 1998; Gulsen and Roose, 2001b; Nicolosi et al., 2000; Torres et al., 1978). Furthermore, Fang and Roose (1997) found a much higher proportion of unique PCR-amplified fragments in lemon cultivars than sweet orange and grapefruit cultivars. On the contrary, when lemon accessions were analyzed, a low level of genetic variability was found (Gulsen and Roose, 2001a). This was also confirmed when Campania lemon germplasm was analyzed, showing high genetic similarity levels even if the analyzed cultivars appeared phenotypically different.
In this study, we carried out the analysis of the genetic diversity of Sicilian lemon germplasm using two multilocus PCR-based techniques, ISSR and RAPD. The usefulness and reliability of these molecular markers has already been proven in several other lemon fingerprinting studies (Capparelli et al., 2004; Fang and Roose, 1997). The data obtained have been used to find genetic relationships among different cultivars and landraces commonly cultivated in this region and to assess the genetic status of lemon germplasm on the island.

Our analyses showed a low grade of polymorphism among all the analyzed genotypes, even though the pool represented by the unique accessions was broad enough (22 of 36 Sicilian genotypes). Also, if the observed morphological differentiation of Sicilian lemon landraces was higher than the identified genetic diversity, these data reflect the richness of lemon germplasm present in Sicily at the moment and suggest that it originated from a common ancestor, probably a ‘Femminello’ plant, from bud mutation. The reproductive characteristics of the lemon (nucellar embryony), the conservative selection criteria of Sicilian traditional agriculture, and genetic isolation (shown by the low Nm and by a very low correlation between genetic and geographical relationships) explain the finding of low levels of heterozygosity. On the contrary, plants belonging to group 2 showed a genetic diversity (GD = 0.03) with the ‘Femminello’ group (group 1) comparable to the diversity identified by Federici et al. (1998) between the cultivars Frost Lisbon and Eureka. Effectively, ‘Monachello Sant’Antonio’ (MS) and ‘Lunario’ (LV) belong to two groups with many phenotypic traits different from ‘Femminello’ (for example, citrus wilt disease tolerance). The other two plants belonging to group 2, ‘Femminello femminino’ (FF) and ‘Femminello famularo a foglia larga’ (FF1), until now considered belonging to the cultivar Femminello, appeared instead genetically very different.

‘Monachello non rifiorente’ (MN) and ‘Lunario spinoso’ (LT) also appeared separated from subgroup 1A, even though more strictly related. ‘Monachello’ and ‘Lunario’ accessions appeared scattered in the UPGMA dendrogram. It warrants consideration that these cultivars could be constituted from genetically very different genotypes, originated several times from different mutational or hybridization events. It explains the wide range of morphotypes constituting these cultivars, but only partially agrees with the hypothesis that Monachello cultivar is a lemon × citron (Citrus medica L.) hybrid (Morton, 1987). ‘Monachello quattroccoli’ (MQ) was not discriminated. This was expected for the accession ‘quattroccoli,’ which is now considered a ‘Femminello’ accession, even though in some areas of Catania it is still called “monachello” because of its high tolerance to citrus wilt disease. The certified ‘Eureka’ (EF) and the disputed ‘Eureka Messina lemon’ (EM) also appeared different from the ‘Femminello’ group and slightly different from each other.

Our analyses also confirmed three other important facts shown by previous phenotypic analysis (Crescimanno et al., 1992). First, accessions coming from the southeast area of Sicily were shown to be more different from other areas (GDs = 0.006 and 0.004 between Catania accessions and Palermo and Messina accessions, respectively). This could be explained with some genetic pool isolation of local accessions. Second, the accession ‘Lunario spinoso’ (LT) appeared tightly associated with the ‘Femminello’ group and, especially, with ‘Femminello verdellifero’ (FV1). Third, the accession ‘Femminello innesto palermitano’ (FI) appeared very different from other ‘Femminello’ accessions. On the contrary, the accessions ‘Femminello famularo a foglia stretta’ (FF2) and ‘Femminello comune’ from Mandanici (FC4), which appeared identical by phenotypic analysis reported from Crescimanno et al. (1992), were differentiated by molecular markers.

Furthermore, ‘Femminello’ accessions from other Italian regions appeared similar to the typical Sicilian ‘Femminello’ profile, providing evidence for movement and exchange of germplasm from the island to the continent.

In conclusion, the molecular investigation carried out in this study allowed an evaluation of the degree of genetic differentiation seen in probably one of the most important clusters of C. limon germplasm in the world. ‘Femminello’, ‘Monachello’, and ‘Lunario’ groups appeared to be effectively different cultivars, each composed of a phenotypically and genotypically diverse range of plants. Much data (GST and Nm) showed that genetic isolation preserved several genetically different landraces, especially in the southeastern area of Sicily, but the situation of the Sicilian lemon cultivation suggest that they could quickly decrease. Actually, Citrus cultivation in Sicily is going through a long crisis period, and the reduced level of gene flow found among populations may decrease the potential for species persistence in face of biotic and abiotic environmental changes (Soulé, 1980). In this situation, collecting local accessions is of great importance for ecological (maintaining rare genes) and cultural (saving landraces) reasons.

Analysis of the genetic variability of lemon landraces and organization of a molecular characterization system could form the basis for future germplasm conservation programs and may represent a useful tool in planning genetic breeding aimed at developing new and improved lemon cultivars.

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