Genetic Diversity of Some *Capparis* L. Species Growing in Syria

Bassam Al-Safadi, Hussam Faouri, and Rana Elias

*Department of Molecular Biology and Biotechnology; Atomic Energy Commission of Syria; Damascus - Syria*

**ABSTRACT**

This work investigated the genetic diversity and relationships among *Capparis* species growing in Syria using IRAP and ISSR techniques. Forty-seven samples of three *Capparis* species genotypes were collected from 21 different locations in Syria. The genotypes were morphologically identified based on the descriptions available in the literature. When IRAP technique was used, an average of 71.5% of the amplified fragments were polymorphic compared to 82.04% in ISSR. Morphological characterization along with the cluster and PCoA analyses of the data divided the studied genotypes into three groups. The groups included genotypes identified as *Capparis spinosa* L., *C. sicula* Duh., and *C. aegyptia* Lam. Based on the morphological description, molecular studies and statistical analyses of this study, *C. aegyptia* could be suggested as a separate species and not a varietal rank of *C. spinosa* (*C. spinosa* var. *aegyptia* (Lam.). Two samples (Alep1 and Idl) were not placed in any of the three distinctive groups, despite their closeness morphologically to *C. spinosa*. In PCoA analysis, sample Alep1 came between *C. sicula* and *C. spinosa* and Idl was placed between *C. sicula* and *C. aegyptia*. Although hybridization between *Capparis* species could occur, it was not clear from the present study if these two genotypes were hybrids.

**Key words:** *Capparis*, Diversity, ISSR, IRAP, Syria

**INTRODUCTION**

*Capparis* L. is a genus belonging to the Capparaceae (Cappariidae). It is distributed in the subtropical and tropical regions of the world (Raja et al. 2013) and grows widely in the Mediterranean and Western Asia. It is believed to include more than 250 species (Jacobs 1965; Fici 1993). Zohary (1960) revised the taxa of *Capparis* species, sub-species and varieties of the Mediterranean and Near Eastern countries and divided them into six groups. Two of the groups (*C. spinosa*, *C. ovata* Desf.) were highly variable with at least six varieties in each group and one group (*C. leucophylla* DC) was less variable with only two varieties. The other three groups (*C. cartilaginea* Decne, *C. mucronfolia* Boiss, *C. decidua* (Forsk) Edgew) were not variable. Jacobs (1965) proposed a wider species concept where all *Capparis* species of the Mediterranean region were included in one single species, namely *Capparis spinosa*. Later, Highton and Akeroyd divided *C. spinosa* into two subspecies: subsp. *spinosa* and subsp. *rupestris* (Sm) Nyman (Saadaoui et al. 2013). Inocencio et al. 2005; 2006), using molecular analysis, subdivided the genus *Capparis* into ten species in Central and Western Asia, North Africa and Europe. Five species have been recorded in the Mediterranean region (*C. spinosa* L., *Capparis sicula* Veill, *Capparis aegyptia* (Lam.) Boiss, *Capparis orientalis* Veill, and *Capparis ovata* Desf.).

Genetic diversity among the individuals or populations can be assessed using morphological and molecular markers (Qi and Lindhout 1997; Fang and Roose 1997; Faccioli et al. 1999; Breto...
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et al. 2001; Reddy et al. 2002; Wang et al. 2005; Kumar et al 2006; Jawdat et al. 2010). Molecular tools provide valuable data on genetic diversity through their ability to detect the variation at the DNA level. Identification is very important in biodiversity studies in many different ways. For evaluation of species diversity, it is essential that individuals can be classified accurately. Several types of molecular markers, including random amplified polymorphic DNA (RAPD), simple sequence repeat (SSR) and inter–simple sequence repeat (ISSR) and amplified fragment length polymorphism (AFLP) have been successfully used for germplasm identification and genetic diversity studies (Qi and Lindhout 1997; Simons et al. 1997; Shan et al. 1999; Wang et al. 2005). The inter-simple sequence repeats (ISSRs), is a technique based on DNA polymerase chain reaction (PCR) method. It uses simple sequence repeat primers to amplify the regions between their target sequences. ISSR marker has been used by many researchers to study genetic diversity in the plants (Reddy et al. 2002; Kumar et al. 2006). This technique has the benefit of discriminating between closely related genotypes (Fang and Roose 1997). Inter-retrotransposon amplified polymorphism (IRAP) technique is also based on DNA polymerase chain reaction (PCR) method and has been exploited to study genetic diversity and phylogeny in many plant genera such as Citrus (Breto et al. 2001), Spartina (Baumel et al 2002), Musa (Teo et al. 2005), Diospyros (Guo et al 2006), and Oryza (Branco et al. 2007). Jawdat et al (2010) used IRAP technique to study the genetic relationship among Eryngium species in Syria.

Despite its medical, nutritional and ecological importance, there have been no molecular studies in Syria that cover all the geographical areas and all the species. Morphological and taxonomical studies are also rare and old (Post 1932; Mouterede 1966; 1984). Most of the previous studies have indicated that only two Capparis species in Syria (Al-Oudat 2008; Chikhali 2013). These studies concentrated on the distribution and morphological characterization of C. spinosa without any investigation on the genetic relationships. Babojian (2007) conducted a taxonomic study on C. spinosa in Syria using morphological characterization in an attempt to classify C. spinosa plants into sub-species. Inocenio et al. (2006), although not working specifically on Capparis species in Syria, collected the samples from some parts of Syria and identified them as C. aegyptia.

This study aimed at investigating the distribution of Capparis species in Syria and their genetic relationships using morphological and DNA molecular analysis.

MATERIALS AND METHODS

Plant Material

Seeds, immature fruits, and stem cuttings of the genus Capparis were collected from different geographical and climatic regions in Syria. A total of 47 samples were collected from 21 different sites, out of which 24 samples were chosen (based on the morphological characteristics) for the molecular study. Collection sites of Capparis L. species along with information regarding geography, altitude, rainfall are shown in Table 1.

Identification of Plants

The morphological characterization and identification of the plants were based on available literature (Zohary 1973; Inocencio et al 2005; 2006; Danin 2010) and covered the following traits: general shape of the plant (Erect, procumbent, pendulous), leaf shape (ovate or elongated), leaf color (dark green or light green), leaf surface (hairy or glabrous), anther filament color (white or colored), fruit shape (round or elongated), pulp color (red or white), and stem color (light pink or green).

Molecular Analyses

DNA extraction

Total genomic DNA was extracted from the leaves of collected Capparis L. DNA extraction was conducted following a modified protocol of Leach et al (1986). Total DNA concentration was detected by the Spectrophotometer (Gene quant, Amersham Biosciences, USA) and the concentration was adjusted to 20 ng/µL.
Table 1 - Collection sites of Capparis L. species along with information regarding geography, altitude, rain-fall.

| Collection site | Code | Geographical Location | Altitude (m) | Rainfall (mm) | Species |
|-----------------|------|-----------------------|--------------|---------------|---------|
| 1               | Alep | North                 | 450          | 365           | Spi, Sic |
| 2               | Idl  | North                 | 350          | 500           | Spi     |
| 3               | Ras  | North Center          | 295          | 450           | Sic     |
| 4               | Tai  | North Center          | 290          | 450           | Sic     |
| 5               | Tha  | North Center          | 275          | 450           | Sic     |
| 6               | Slu  | Coastal Mountains     | 1100         | 1000          | Spi     |
| 7               | Tar  | Coast                 | 50           | 830           | Spi     |
| 8               | Lat  | Coast                 | 50           | 750           | Spi     |
| 9               | Pal  | Center                | 750          | 170           | Sic, Aeg |
| 10              | Rust | Center                | 450          | 360           | Spi, Sic |
| 11              | Zein | Center                | 500          | 360           | Spi, Sic |
| 12              | Sham | Center                | 700          | 120           | Spi, Aeg |
| 13              | Jin  | South West            | 1200         | 650           | Spi     |
| 14              | Mad  | South West            | 1400         | 600           | Spi     |
| 15              | Tal  | South West            | 1050         | 250           | Spi, Aeg |
| 16              | Er   | South West            | 1400         | 650           | Spi     |
| 17              | Qiq  | South West            | 1100         | 500           | Spi, Aeg |
| 18              | Adra | South West            | 750          | 120           | Sic, Aeg |
| 19              | Bus  | South                 | 800          | 230           | Spi, Sic |
| 20              | Izra | South                 | 570          | 290           | Spi, Sic |
| 21              | Swei | South                 | 1800         | 450           | Spi, Sic |

Inter-Retrotransposon Amplified Polymorphism (IRAP) Procedure

Eleven IRAP primer combinations (Tables 2) were synthesized using the PolyGen DNA synthesizer (PolyGen DNA-Synthesizer, Germany) and the iCycler PCR machine (BIO-RAD, USA) was used for the amplification of total genomic DNA. The amplification reactions of total genomic DNA were performed in 25µL reaction volume, containing 2.5 µL of 10 X Taq polymerase reaction buffer, 2.0 µL of 10 mM dNTPs mix, 0.05 µL of 5.0 U/µL Taq DNA polymerase, 3.4 µL of 50 mM, 2.0 µL of each of the 50 pmol/µL of forward and reverse primer and 40 ng DNA.

Table 2 - Primer combinations used for IRAP with annealing temperature.

| No. | Primer Combination | Annealing Temperature °C |
|-----|--------------------|--------------------------|
| 1   | D26-D26            | 41.5                     |
| 2   | D26-D27            | 43.5                     |
| 3   | D16-D17            |                          |
| 4   | D18-D19            |                          |
| 5   | D20-21             |                          |
| 6   | D22-23             |                          |
| 7   | D24-25             | 45.5                     |
| 8   | D29-D29            |                          |
| 9   | D30-D31            |                          |
| 10  | D32-D32            |                          |
| 11  | D33-D33            | 48.5                     |

Inter-simple sequence repeats (ISSR) procedure

Thirteen ISSR primers (Table 3) were used for the amplification of total genomic DNA. The amplification reactions of total genomic DNA were performed in 20 µL reaction volumes, containing 2.5 µL of 10 X Taq polymerase reaction buffer, including 0.8 µL of 10 mM dNTPs mix, 0.05 µL of 5 U/µL Taq DNA polymerase, 2.5 µL of 10 pmol/µL of each primer, and 25 ng of DNA. For IRAP, fragments were separated on 2.5% agarose and for ISSR, they were separated on 1.5% agarose gels in 1X TAE buffer and stained with ethidium bromide (10 mg/mL).

Table 3 - ISSR primers used for the amplification.

| No. | Primer | Sequence |
|-----|--------|----------|
| 1   | ISSR-1 | (CAC)_n T |
| 2   | ISSR-2 | (GA)_n C |
| 3   | ISSR-7 | (CA)_n G |
| 4   | ISSR-8 | (CT)_n G |
| 5   | ISSR-10| (TCC)_n |
| 6   | ISSR-13| (AG)_n G |
| 7   | ISSR-14| (GA)_n T |
| 8   | ISSR-15| (GA)_n C |
| 9   | ISSR-16| (GA)_n A |
| 10  | ISSR-19| (CT)_n T |
| 11  | ISSR-22| (GT)_n A |
| 12  | ISSR-43| (AC)_n CTA |
| 13  | ISSR-47| (ACC)_n |
Data Analysis
IRAP and ISSR bands were transformed into a binary matrix, where the presence of reproducible polymorphic DNA band at a particular position on the gel was manually scored as 1 (present) and 0 (absent). The genetic similarity matrices were constructed using the correlation coefficient. Dendrograms were generated by the un-weighted pair-group method using the correlation similarity measure. Principal coordinate analysis (PCoA) was performed to provide the spatial representation of the relative genetic distances among the individuals and to determine the consistency of differentiation among the populations defined by the cluster analysis. Data analyses were performed using PAST program (Hammer et al. 2001).

RESULTS

Morphological characterization
*Capparis* L. species investigated in the current study were characterized using several morphological traits as previously described. Three different *Capparis* species were identified to grow in Syria. The species and some of their morphological characters were:

1- *C. spinosa* L.: Shrubs were erect, and glabrous; the leaves were dark green and generally ovate large (3-3.5 cm) with glabrous texture somewhat fleshy; the stem was green or colored (mostly light to dark pink). The flowers were large zygomorphic and abaxial sepal was slightly galeat shaped. The fruits were oblong with inner white wall. Stamen filaments were colored (mostly light to dark pink) (Fig. 1A).

2- *C. sicula* Duh.: Shrubs were procumbent and gray green, densely covered by white hair on the stems, leaves and buds. Leaves were round to ovate (2-2.3 cm) green with dense pubescence not fleshy (herbaceous). Flowers were zygomorphic but smaller than *C. spinosa*. Sepal was galeat. The fruit was obovate to oblong with inner red wall. Filaments were white (Fig. 1B).

3- *C. aegyptia* Lam.: Shrubs were somewhat erect (pendulous) glabrous and green or blue-green. Leaves were obovate to ovate and fleshy glabrous (2.3-2.7 cm); adult leaves became later in the season gray to blue thicker and covered with wax. Flowers were slightly zygomorphic similar in size to *C. spinosa* or a bit smaller; sepal was slightly galeat. Fruit was oblong and the pulp was white to yellow. Stamen filaments were colored (mostly light to dark pink) (Fig. 1C).

Figure 1 - Photos showing: A: *Capparis spinosa* (Beit Jin), B: *C sicula* (Izraa), C: *C. aegyptia* (Abou Shamat area)

Molecular characterization
IRAP Data
Eleven primer combinations generated multiple bands from the genomic DNA of all the studied *Capparis* L. species (Fig. 2). The banding pattern of the amplification using IRAP primers is summarized in Table 4. A total of 153 fragments were amplified; 109 (71.5%) of them were polymorphic. The number of polymorphic fragments varied from 6 (D18-D19) to 14 (D26-D27) with an average of 9.9 per primer. Two unique bands were scored in the samples collected from Sweida and Latakia (Table 4).
Table 4 - The banding pattern of IRAP amplification products.

| Primer Combination | Total bands | Polymorphic bands | % polymorphism | unique bands |
|--------------------|-------------|-------------------|----------------|--------------|
| nikita              | 18          | 13                | 72.2           | 0            |
| sukkula             | 11          | 8                 | 72.7           | 0            |
| D30-D31             | 9           | 8                 | 88.8           | 0            |
| D22-D23             | 8           | 7                 | 87.5           | 0            |
| D26-D27             | 15          | 14                | 93.3           | 0            |
| D20-D21             | 16          | 8                 | 50             | 0            |
| D26-D26             | 13          | 9                 | 69.2           | 0            |
| D29-D29             | 17          | 13                | 76.4           |              |
| D24-D25             | 14          | 9                 | 64.2           | 0            |
| D16-D17             | 18          | 13                | 72.2           | 0            |
| D18-D19             | 15          | 6                 | 40             | 0            |
| **Total**           | **153**     | **109**           | **71.5**       | **2**        |

The dendrogram resulting from the cluster analysis of the data showed that the genotypes were separated into two main groups (Fig. 3). The first group was separated into several sub-clusters and included the samples morphologically identified as *C. spinosa*, collected mainly from the coastal and southern regions. The second group was separated into two sub-clusters. The first sub-cluster included the samples identified as *C. aegyptia* and the second one included the samples of *C. sicula*.

**ISSR data**

The 13 selected ISSR primers produced various numbers of DNA fragments (Fig. 4), depending on their simple sequence repeat motifs (Table 3). In total, 84 fragments were produced, out of which 69 fragments (82.04%) were polymorphic. The number of polymorphic fragments varied from three (ISSR-8, ISSR-10, ISSR-43) to 11 (ISSR-16) with an average of 5.3 per primer; five unique bands were scored in the samples collected from Sweida, Erna, Taiba (2) and Palmyra (Table 5).

The dendrogram resulting from the cluster analysis of the data showed that genotypes were separated into two main groups (Fig. 5). The first group, which in turn was divided into sub-clusters, included only *C. spinosa* samples collected from different parts of Syria (similar to those revealed by IRAP). The second group was also divided into two sub-clusters with the first sub-cluster including only *C. aegyptia* and the other including all *C. sicula* samples and two *C. spinosa* samples.

![Figure 2 - Amplification of genomic DNA for the 24 Capparis samples using IRAP primer combination D29-D29. The arrow indicates a unique band in the sample collected from Latakia using the same primer combination.](image)

![Figure 3 - Dendrogram of 24 Capparis genotypes generated by the paired group method using correlation matrix, based on data of IRAP. The numbers on the tree branches indicate the percentage of replicates included in a given node calculated out of 1000 bootstrap replicates.](image)
Figure 4 - Amplification of genomic DNA for the 24 *Capparis* samples using ISSR primer 16 (GA)8 A. The arrow indicates a unique band in the sample collected from Sweida using the same primer.

Table 5 - The banding pattern of ISSR amplification products.

| Primer  | Total bands | Polymorphic bands | % polymorphism | unique bands |
|---------|-------------|-------------------|----------------|-------------|
| ISSR-1  | 7           | 6                 | 85.7           | 0           |
| ISSR-2  | 7           | 6                 | 85.7           | Erna (Spi)  |
| ISSR-7  | 6           | 5                 | 83.3           | Palmyra (Sic) |
| ISSR-8  | 4           | 3                 | 75             | Taiba (Sic) |
| ISSR-10 | 6           | 3                 | 50             | 0           |
| ISSR-13 | 11          | 9                 | 81.8           | Taiba (Sic) |
| ISSR-14 | 5           | 5                 | 100            | 0           |
| ISSR-15 | 7           | 6                 | 85.7           | 0           |
| ISSR-16 | 11          | 10                | 90.9           | Sweida (Spi) |
| ISSR-19 | 5           | 5                 | 100            | 0           |
| ISSR-22 | 6           | 5                 | 83.3           | 0           |

Figure 5 - Dendrogram of 24 *Capparis* genotypes generated by the paired group method using correlation matrix, based on data of ISSR. The numbers on the tree branches indicate the percentage of replicates included in a given node calculated out of 1000 bootstrap replicates.

**Combined (IRAP and ISSR) data**

The data obtained from both the marker techniques were combined. A total of 237 DNA fragments were scored, of which 178 (75.1%) were polymorphic with an average of 7.4 polymorphic fragments per primer. The genetic relationships among the studied genotypes through IRAP and ISSR markers, showed similar dendrograms as obtained from each single technique with some exceptions. However, the general dendrogram constructed using the similarity coefficient values of combined molecular data (IRAP+ISSR) revealed a more realistic representation of the relationship than from the individual markers. The genotypes were grouped into two main clusters (Fig. 6). The first cluster was divided into sub-clusters, which included *C. spinosa* samples collected from the coast-line and coastal mountains (Latakia, Tartus, Sulunfa), the mountains of south-west (Erna, Madaia, Beit Jin) and southern region of Syria (Sweida, Izraa). *C. spinosa* samples collected from Latakia and Tartus (similar geography and climate) showed a close genetic relationship (0.83) as was revealed by the similarity analysis (data not shown) with very high bootstrap value (96). Genotypes identified as *C. sicula* were also very closely related as they too came in one cluster with very high bootstrap value (97) with similarity coefficient values between 0.68 and 0.89 (data not shown). These genotypes were collected mostly from the areas in the center of Syria with altitude...
between 275-800 m above the sea level and rainfall between 170-450 mm.

Genetic similarity analysis among the studied Capparis species showed that the genotypes identified as C. aegyptia were genetically much closer to C. sicula than to C. spinosa. This was apparent from the dendrogram and similarity coefficient values. Those values ranged from 0.11-0.34 when comparing C. aegyptia values collected from Addra, Abou Shamat, and Kafr Quq to C. spinosa collected from Latakia, Tartus, and Slunfa. On the other hand, the comparison of the samples of C. sicula with C. aegyptia showed these values to range from 0.44-0.56. The closest genetic relationship (0.89) was observed between the two C. sicula samples, Ras2 and Ras3 (Rasafa), and the farthest genetic relationship (0.09) was observed between C spinosa (Sweida) and C aegyptia (Addra).

The Principle Coordinate Analysis (PCoA) based on genetic similarity a matrix was used to visualize the genetic relationships among Capparis genotypes (Fig. 7).

**Figure 6** - Dendrogram of 24 Capparis genotypes generated by the paired group method using correlation matrix, based on combined data of IRAP and ISSR. The numbers on the tree branches indicate the percentage of replicates included in a given node calculated out of 1000 bootstrap replicates.

**Figure 7** - Relationships among the Capparis genotypes revealed by principal coordinate analysis based on combined IRAP and ISSR genetic similarity.
The first coordinate (horizontal axis) accounted for 48.7% of the variation and the second coordinate (vertical axis) accounted for 15.2% of the variation. Three groups were observed: \textit{C. sicula} and \textit{C. aegyptia} appeared on the left side of the plot, forming two independent groups that were separated by the horizontal axis but close to each other. On the right side of the plot, \textit{C. spinosa} was isolated from the other two groups. Two genotypes (Spi Alep1 and Spi Idl) were isolated from the other three major groups. This confirmed the results of cluster analysis where \textit{C. spinosa}, \textit{C. aegyptia}, and \textit{C. sicula} samples were separated in three isolated groups.

**DISCUSSION**

It has become obvious these days that the morphological characterization of plants is not sufficient to make definitive discrimination among the species, sub-species, and varieties as morphological markers are influenced by the environmental conditions and developmental stages, which make their use limited in genetic diversity studies. For example, in Sicily, \textit{C. spinosa} showed remarkable variability with regard to growth forms and other vegetative characters. In typical individuals of \textit{C. spinosa} observed on clay soil, each year from renewal buds at or below ground level, only a few fast-growing, multiramified shoots were produced. In contrast, individuals of the same species growing under atypical pedological conditions (rocky habits) usually showed a lignified shoot system like \textit{C. spinosa} \textit{L.var. intermis} Turra (Fici 2001). DNA molecular markers, on the other hand, are independent of environmental conditions or developmental stage and show high level of polymorphism.

Genetic studies on \textit{Capparis} in Syria are rare and limited to one species or limited to one region. This study was the first study on \textit{Capparis} species, which covered most of the geographical and ecological areas and the various \textit{Capparis} species growing in Syria (Fig. 8).

![Distribution map of Capparis species in Syria](image)

**Figure 8** - Distribution of the three \textit{Capparis} species found to grow in Syria.

Molecular analysis in this study confirmed the outcomes of morphological and taxonomic analyses done on the collected samples. The combined IRAP and ISSR data were utilized to correlate the genetic makeup with the morphological description and geographical distribution of \textit{Capparis} species. The clustering of genotypes within the groups was almost similar in both the techniques with some exceptions. ISSR dendrogram was much similar to the combined
dendrogram than IRAP dendrogram due to the higher percentage of polymorphic bands obtained by ISSR was higher than that obtained by IRAP technique (82% vs 71.5%). This made the use of ISSR technique in this study more efficient and more informative than IRAP. This finding was in line with the findings of Bublyk et al. (2013) who investigated the efficiency of different PCR-based marker systems (ISSR, IRAP, REMAP, RGAP and LP-PCR) for the assessment of Iris pumila genetic diversity and found that ISSR-markers were the most efficient, followed by LP-PCR-markers. A similar result was achieved by Žiarovská et al. (2012) studying variability of Linum usitatissimum L. using several molecular marker techniques.

Combining the data of both the techniques was successful in sorting the Capparis genotypes into three somewhat distinctive groups matching the morphological analysis. Although C. spinosa and C. sicula were relatively similar morphologically and had often been confused taxonomically (Inocencio et al. 2005; 2006), careful examination of the plants (especially color of filaments, color of leaves, presence of hair, shape of fruit and color of inside wall) as well as DNA analysis could sort them out. The same could be said with regard to C. aegyptia, which could be confused visually with C. spinosa. A recent study on Egyptian taxa of Capparis and related genera (Capparaceae) conducted by Moubasher et al. (2011), using RAPD technique revealed that C. spinosa var. deserti (C. aegyptia), C. spinosa var. canescens (C. sicula), and C. spinosa var. spinosa were more related to each other than to C. decidua, and C. sinaica.

Inocencio et al. (2006) reported that C. spinosa was found rarely in the wild and that most known populations were cultivated and it possessed intermediate morphological characters between C. orientalis and C. sicula, suggesting that C. spinosa could be of hybrid origin. In the same study, Inocencio et al. (2006) did not indicate the existence of C. spinosa in Syria. However, several other studies have reported the existence C. spinosa in Lebanon and Syria (Post 1932; Mouterede 1966; 1984; Chalak et al. 2001; Giuliani and Buerli 2006; Babojian 2007; Al-Oudat 2008; Al-Safadi and Elias 2011; Chikhali 2013). The present study found C. spinosa to grow commonly in the wild, and in fact it grew in diverse geographical and climatic regions in Syria (Table 1 and Fig. 8). On the other hand, there were a few places in Syria where C. spinosa was cultivated (Giuliani 2004).

Previous studies considered C. aegyptia as a varietal rank of C. spinosa (C. spinosa var. aegyptia (Lam.) Boiss) (Zohary 1960; 1962; Boulos 1999; Özbek and Kara. 2013). This was rejected by Danin (2010) who disputed this notion based on the morphological description of the plants and their geographical and climatic distribution. Based on the results of morphological description, molecular studies and statistical analyses of the present study, C. aegyptia, C. sicula and C. spinosa were considered as separate species. This was also in line with the findings of Inocencio et al. (2005; 2006) who studied C. spinosa, C. orientalis, C. ovata, C. aegyptia and C. sicula using morphological and molecular analyses and found these to be separate species. They also suggested that C. spinosa might be a hybrid between C. orientalis and C. sicula.

Capparis orientalis was not found in the present study. It might, however, exist hence further studies are needed in the future. Two samples (Alep1 and Idl) were not placed in any of the three distinctive groups, despite their closeness morphologically to C. spinosa. In PCoA analysis (Fig. 7), Alep1 came between C. sicula and C. spinosa, indicating its close relationship to both of the species. This was confirmed by the analysis of genetic relationship (correlation’s coefficients) where these values ranged from 0.46-0.57 when comparing Alep1 with C. sicula genotypes and from 0.42-0.59 when comparing it with C. spinosa genotypes. The same argument could be extended to Idl where PCoA analysis placed Idl between C. sicula and C. aegyptia with genetic relationship coefficient values ranging from 0.63-0.70 when comparing Idl with C. sicula genotypes and from 0.43-0.57 when comparing it with C. aegyptia genotypes. Although hybridization between Capparis species might occur (Inocencio et al. 2006), it was not clear from the present study if these two genotypes were hybrids (Alep1 between C. spinosa with C. sicula, and Idl between C. sicula with C. aegyptia) or other Capparis species that were not investigated thoroughly in this study.

**CONCLUSIONS**

Based on the morphological and molecular study, three Capparis species were identified to grow in Syria, which were C. spinosa L, C. sicula Duh and
C. aegyptia Lam. There was also a possibility of the existence of one or two more species due to the fact that two genotypes collected from Aleppo and Idlep were not placed in any of the three distinctive groups of the species. This suggested the necessity to conduct further studies to investigate the existence of these species and perhaps to work at the subspecies level to learn more about these important plants.

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