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RESEARCH ARTICLE

Molecular Characterization using Directed Amplification of Minisatellite-region DNA (DAMD) Marker in *Ficus Sycomorus* L. (Moraceae)

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Abstract:

Background:

Ficus sycomorus L. species exhibited great importance with various applications in pharmacology and medicine studies. However, little attention has been given to its molecular characterization.

Objective:

The study aimed to assess DNA genetic diversity among 16 genotypes of *F. sycomorus* L. species.

Methods:

Directed Amplification of Minisatellite-region DNA (DAMD) marker has been employed to investigate the genetic relationship among the studied genotypes of *F. sycomorus* L. species based on the estimated Percent Disagreement Values (PDV).

Results:

Twenty-four DAMD primers produced 194 bands, of which, 145 (74.742%) were polymorphic with Polymorphic Information Content (PIC) average of 0.219. DAMD-PCR application highlighted 12 unique markers characteristic for some studied genotypes. Cluster analysis showed that the studied *F. sycomorus* L. genotypes were split into two main distinguished clusters, each one was considered as a subspecies. In this respect, *F. sycomorus14* and *F. sycomorus15* genotypes were considered as subspecies too far from the second one containing the remaining genotypes.

Conclusion:

The DAMD assay successfully highlighted genetic diversity within *F. sycomorus* species. More accurate molecular markers are required to confirm the current data.

Keywords: *Ficus sycomorus* L., DAMD, Molecular marker, Genotype, Genetic diversity, PDV.

Article History

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1. INTRODUCTION

Ficus sycomorus L. (Moraceae) species is characterized by its geographic diversity due to its adaptation to different environmental conditions. Assessment of genetic diversity of *F. sycomorus* L., as in the case of other fruit trees, is important not only in evolutionary studies, but also in the breeding and conservation programs of germplasm. It has been demonstrated that the existence of 400 monoecious and 800 gynodioecious species belonged to the *Ficus* genus [1, 2]. Overall, *F. sycomorus*

L. species taxonomically is classified as a subgenus of the fig, belonging to the Moraceae family.

It was originated in Ethiopia and Central Africa, and planted since antiquity in Egypt, Palestine, Lebanon and Syria. Since a long time ago, Mouterde [3] reported that this species is at risk due to urban development, with little occurrence of this species in Sida and Syrian littoral zones.

This study is part of a series of studies currently underway in Syria and around the world under the project of biodiversity and the inventory of genetic resources, especially those neglected and marginalized to benefit food, medicine and genetic improvement programs to combat diseases, insects and abiotic stresses and a part of the efforts being made to expand

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their cultivation. *F. sycomorus* L. species displayed an important role as a coastal plant, genetically resource tolerant to salinity, as a food (their fruit) and medical treatment [2, 4].

Many DNA-PCR based markers are successfully employed worldwide for investigation of DNA genetic diversity in plants kingdom. Of the available techniques, Directed Amplification of Minisatellite-region DNA (DAMD) technique has been developed by Heath *et al.* [5] and revealed polymorphism due to minisatellites. Since this technique is carried out at higher PCR-stringencies, the patterns produced have a greater reproducibility than Random Amplified Polymorphic DNA (RAPD) [6]. It has been documented that minisatellites called as the Variable Number of Tandem Repeats (VNTR), are tandem repeats of 10-60 bp long DNA sequence motif and are widely distributed throughout the eukaryote genome. The variation in these repeats is proved to be the source of the polymorphism in several organisms [7].

It has been demonstrated that these minisatellites and DNA sequences flanking them are involved in inversions, which result in their distribution on both the strands in opposite orientations. This makes PCR possible using a minisatellite core sequence as a single primer. Moreover, these minisatellites have characteristics of inheritance in Mendelian fashion and extensive genome coverage [5].

A DAMD marker has been widely used for genetic characterization of many plant species including *Oryza* [8], *Morus* spp [9], *Panax* spp [10], *Shorea curtisii* [11], *Capsicum* [12], *Convolvulus* [13], *Astragalus rhizanthus* complex [14], common bean (*Phaseolus vulgaris* L.) [15], cucumber (*Cucumis sativus* L.) [16], grapevine [17], mango [18], *Citrus* [19], chickpea [20], *Jatropha* [21], *Flavoparmelia caperata* L [22], *Dianthus* L [23]. and recently, in *Citrus* [24] and *Sechium edule* (Jacq.) Sw [25].

The utility of DAMD marker, however, has not been widely reported in *F. sycomorus* L. germplasm characterization. Thereby, the current study aimed to assess the phylogenetic relationship among *F. Sycom* genotypes (scattered in various areas of the Syrian coast) along with the precise identification of the subspecies belonging to the *F. sycomorus* species, especially since there is no precise inventory of the

number of subspecies of this species in Syria and to find specific molecular markers of this species using the DAMD assay.

2. MATERIAL AND METHODS

2.1. Plant Materials

Sixteen *F. sycomorus* leaves samples were collected from the coastal regions of Syria (Table 1). Five to ten leaves from each genotype were bulked as one sample for DNA extraction. Table 1 shows characteristic collection sites in terms of the original site, altitude and annual rainfall. Sample collection has been performed in autumn as previously reported by Saleh [2].

2.2. DNA Extraction

Total genomic DNA of each genotype has been performed by a CTAB (cetyltrimethylammonium bromide) protocol as described by Doyle and Doyle [26]. The concentration of DNA was measured by the DNA fluorimeter and kept at -80°C until use.

2.3. DAMD-PCR Assay

Twenty-four DAMD primers were tested as previously reported in other species [9, 14, 17]. DAMD-PCR amplification reaction was carried out in 25 μl reaction volume containing 1X PCR buffer, 2 mM MgCl_2 , 0.25 mM dNTPs, 25 pmol primer, 1.5 unit of *Taq* DNA polymerase and 50 ng template DNA. PCR amplification was performed in a T-gradient thermal cycler (Bio-Rad; T-Gradient). It was programmed to 35 cycles after an initial denaturation cycle for 4 min at 94°C . Each cycle consisted of a denaturation step for 1 min at 94°C , an annealing step for 2 min at T_m varying according to each examined primer (Table 2), and an extension step at 72°C for 2 min, followed by extension cycle for 7 min at 72°C in the final cycle. PCR products were then separated on a 1.8% ethidium bromide-stained agarose gel (Bio-Rad) in 0.5X TBE buffer. Electrophoresis was performed for 2 h at 100 V and visualized with a UV transilluminator. A standard VC 100bp Plus DNA Ladder (Vivantis) was used to estimate the molecular weight of DAMD-PCR amplification products.

Table 1. Geographical regions in terms of original sites including their altitude (m) and annual rainfall (mm) in which the 16 *F. sycomorus* genotypes were collected.

Genotype Code	Original Site	Altitude (m)	Annual Rainfall (mm)	References
<i>F. sycomor 1</i>	Lattakia	4.5	650-700	[2]
<i>F. sycomor 2</i>	Lattakia	6.5	650-700	Current study
<i>F. sycomor 3</i>	Lattakia	6.3	650-700	[2]
<i>F. sycomor 4</i>	Lattakia	6.5	650-700	Current study
<i>F. sycomor 5</i>	Lattakia	5	650-700	Current study
<i>F. sycomor 6</i>	Lattakia	5	650-700	Current study
<i>F. sycomor 7</i>	Lattakia	5	650-700	Current study
<i>F. sycomor 8</i>	Lattakia	5	650-700	Current study
<i>F. sycomor 9</i>	Lattakia	4	650-700	Current study
<i>F. sycomor 10</i>	Lattakia	6.3	650-700	[2]
<i>F. sycomor 11</i>	Lattakia	5	650-700	Current study
<i>F. sycomor 12</i>	Jableh	11.8	650-700	[2]

(Table 1) contd....

Genotype Code	Original Site	Altitude (m)	Annual Rainfall (mm)	References
<i>F. sycomor 13</i>	Banyas	10	700-750	[2]
<i>F. sycomor 14</i>	Banyas	220	~850	[2]
<i>F. sycomor 15</i>	Banyas	12	700-750	Current study
<i>F. sycomor 16</i>	Banyas	250	~850	[2]

Table 2. DAMD primers tested in the current study in terms of primer name, primer sequence and temperature annealing (°C).

Primer Number	Primer Name	Primer Sequence 5'-3'	Source Species	Ta (°C)	References
1	URP1F	ATCCAAGGTCCGAGACAACC	Rice	55	[16]
2	URP2R	CCCAGCAACTGATCGCACAC	Rice	61	[16]
3	URP4R	AGGACTCGATAACAGGCTCC	Rice	58	[16]
4	URP9F	ATGTGTGCGATCAGTTGCTG	Rice	60	[16]
5	URP25F	GATGTGTTCTTGGAGCCTGT	Rice	58	[16]
6	HVR(-)	CCTCCTCCCTCCT	Rice	50	[16]
7	OGRB01	AGGGCTGGAGGAGGGC	Rice	55	[16]
8	FVIIex8C	CCTGTGTGTGTGCAT	Human	47	[16]
9	HBV3	GGTGAAGCACAGGTG	Human	53	[16]
10	HBV5	GGTGTAGAGAGGGGT	Human	56	[16]
11	YNZ22	CTCTGGGTGTGGTGC	Human	56	[16]
12	14C2	GGCAGGATTGAAGC	Human	53	[16]
13	33.6	GGAGGTGGGCA	Human	52	[16]
14	6.2H(+)	AGGAGGAGGGGAAGG	Human	56	[16]
15	M13	GAGGGTGGCGGCTCT	Phage M13	57	[16]
16	HBVb	GGTGTAGAGAGAGGGGT	Rice	52	[9]
17	HVRc	CCTCCTCCCTCCT	Rice	52	[9]
18	URP2F	GTGTGCGATCAGTTGCTGGG	Rice	50	[17]
19	URP6R	GGCAAGCTGGTGGGAGGTAC	Rice	50	[17]
20	URP13R	TACATCGCAAGTGACACAGG	Rice	50	[17]
21	URP17R	AATGTGGGCAAGCTGGTGGT	Rice	50	[17]
22	M13	GAGGGTGGCGGTTTCCT	Rice	55	[14]
23	HVA	AGGATGGAAAGGAGGC	Rice	55	[14]
24	HVY	GCCTTTCCCGAG	Rice	55	[14]

2.4. DAMD Analysis

The presence or absence of each size class was scored as 1 or 0, respectively. The Percent Disagreement Values (PDV) found were used to generate a matrix *via* the Unweighted Pair Group Mean Arithmetic average (UPGMA) using Statistica program [27]. Then the previous matrix was used to estimate genetic similarity [28]. Polymorphic Information Content (PIC) value was estimated for each DAMD primer according to the formula:

$$PIC = 1 - \sum(P_{ij})^2$$

Where P_{ij} is the frequency of the i th pattern revealed by the j th primer summed across all the patterns revealed by the primers [29].

3. RESULTS

The size of DAMD-PCR amplification products ranged between 100-1500 bp. Fig. (1) shows polymorphic patterns within *F. sycomorus* L. species produced by 14C2 (a) and URP17R (b) DAMD primers.

The DAMD marker applied in the molecular study within *F. sycomorus* L. species, produced total bands number (TB) of 194 bands, of which, 145 were polymorphic (PB) representing percentage polymorphism (P%) of 74.742% (Table 3). Data showed that TB ranged between 4 [URP1F & 6.2h(+)] and 19 (URP4R) with a mean average of 8.083 bands/ primer. Whereas, PB ranged between 2 [URP9F, HVR(-), 6.2H(+)] and 19 (URP4R) with a mean average of 6.042 bands/ primer (Table 3). As for estimated PIC, it ranged between 0.098 (URP6R) and 0.369 (M13) with a mean average of 0.219 (Table 3).

The DAMD assay successfully produced twelve unique markers, characterized *F. sycomorus* genotypes (Table 4).

Based on UPGMA, Fig. (2) shows that the studied *F. sycomorus* genotypes were clustered into two main clusters. The first cluster contained *F. sycomorus14* and *F. sycomorus15* genotypes, whereas, the second one contained three sub-clusters. The first subcluster contained *F. sycomorus1*, *F. sycomorus2*, *F. sycomorus3*, *F. sycomorus4*, *F. sycomorus5* and *F. sycomorus11* genotypes (Fig. 2), where, *F. sycomorus2*

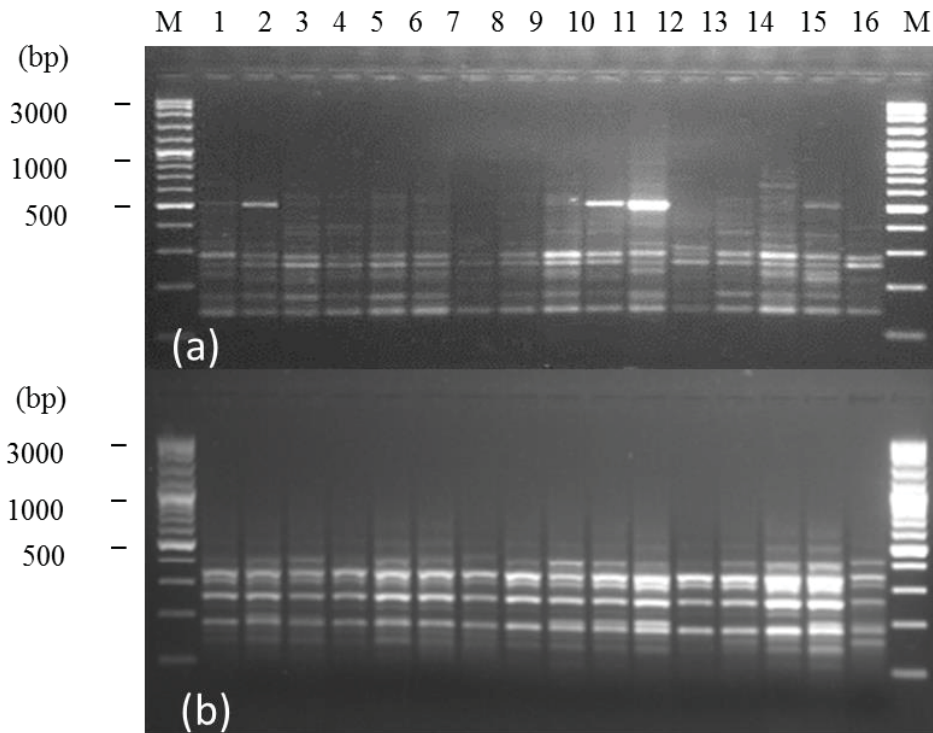


Fig. (1). DAMD polymorphism produced by 14C2 (a), and URP17R (b) tested primers among the 16 *F. sycomorus* studied genotypes. M: A VC 100bp Plus DNA Ladder (Vivantis) ladder standard.

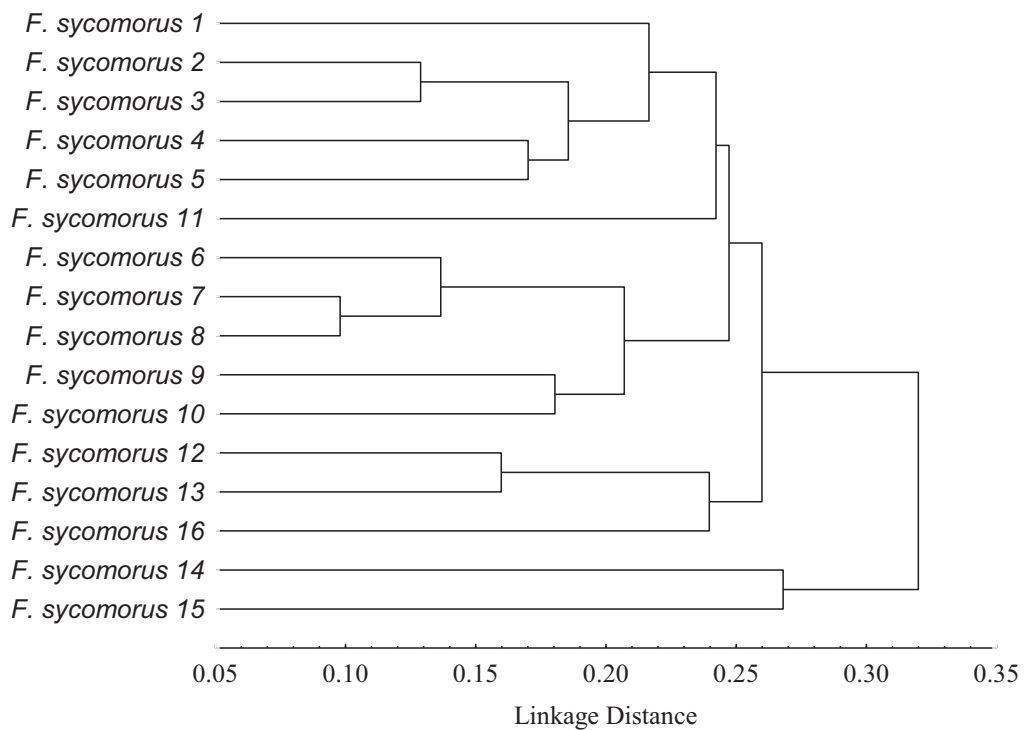


Fig. (2). Cluster analysis generated using UPGMA analysis, showing relationships among the 16 *F. sycomorus* studied genotypes based on DAMD data.

Table 3. Banding pattern of DAMD amplified fragments scored.

Primer Number	Primer Name	TB	PB	P%	PIC
1	URP1F	4	3	75.000	0.178
2	URP2R	11	11	100.000	0.363
3	URP4R	19	19	100.000	0.366
4	URP9F	5	2	40.000	0.173
5	URP25F	7	4	57.143	0.141
6	HVR(-)	6	2	33.333	0.099
7	OGRB01	9	6	66.667	0.22
8	FVIIex8C	10	8	80.000	0.191
9	HBV3	14	10	71.429	0.235
10	HBV5	6	4	66.667	0.202
11	YNZ22	9	4	44.444	0.145
12	14C2	7	5	71.429	0.166
13	33.6	7	5	71.429	0.166
14	6.2H(+)	4	2	50.000	0.152
15	M13	9	8	88.889	0.297
16	HBVb	6	5	83.333	0.323
17	HVRc	6	4	66.667	0.178
18	URP2F	7	5	71.429	0.246
19	URP6R	5	2	40.000	0.098
20	URP13R	9	7	77.778	0.202
21	URP17R	9	6	66.667	0.179
22	M13	11	11	100.000	0.369
23	HVA	6	4	66.667	0.268
24	HVY	8	8	100.000	0.287
Total		194	145	–	–
average		8.083	6.042	70.374	0.219

TB – Total Bands; PB – Polymorphic Bands; P (%) – Polymorphic % and PIC – Polymorphic Information Content.

Table 4. Molecular weights of unique fragments based on DAMD data.

Genotype Code	Original SiteS	Primer	Unique Positive Marker Molecular Weight (bp)
<i>F. sycomor 1</i>	Lattakia	FVIIex8C	~150
<i>F. sycomor 1</i>	Lattakia	33.6	500
<i>F. sycomor 1</i>	Lattakia	URP2F	~290
<i>F. sycomor 4</i>	Lattakia	URP25F	~150
<i>F. sycomor 6</i>	Lattakia	URP13R	200
<i>F. sycomor 8</i>	Lattakia	URP4R	~450
<i>F. sycomor 9</i>	Lattakia	HVRc	200
<i>F. sycomor 9</i>	Lattakia	HVY	200
<i>F. sycomor 13</i>	Banyas	URP4R	~150
<i>F. sycomor 13</i>	Banyas	FVIIex8C	~1100
<i>F. sycomor 14</i>	Banyas	URP4R	900
<i>F. sycomor 14</i>	Banyas	HBV3	1500
<i>F. sycomor 15</i>	Banyas	33.6	1000
<i>F. sycomor 15</i>	Banyas	URP2F	100
<i>F. sycomor 15</i>	Banyas	URP6R	800
<i>F. sycomor 15</i>	Banyas	URP13R	~450
<i>F. sycomor 15</i>	Banyas	M13	~150
<i>F. sycomor 16</i>	Banyas	URP1F	~190
<i>F. sycomor 16</i>	Banyas	6.2H(+)	200

Table 5. Percent disagreement values (PDV) generated by DAMD data based on UPGMA routine in statistical program.

Genotype	<i>F. syc1</i>	<i>F. syc2</i>	<i>F. syc3</i>	<i>F. syc4</i>	<i>F. syc5</i>	<i>F. syc6</i>	<i>F. syc7</i>	<i>F. syc8</i>	<i>F. syc9</i>	<i>F. syc10</i>	<i>F. syc11</i>	<i>F. syc12</i>	<i>F. syc13</i>	<i>F. syc14</i>	<i>F. syc15</i>	<i>F. syc16</i>
<i>F. syc1</i>	0.00															
<i>F. syc2</i>	0.19	0.00														
<i>F. syc3</i>	0.23	0.13	0.00													
<i>F. syc4</i>	0.22	0.17	0.21	0.00												
<i>F. syc5</i>	0.23	0.19	0.18	0.17	0.00											
<i>F. syc6</i>	0.29	0.21	0.27	0.21	0.21	0.00										
<i>F. syc7</i>	0.29	0.25	0.28	0.22	0.24	0.10	0.00									
<i>F. syc8</i>	0.27	0.23	0.27	0.19	0.24	0.17	0.10	0.00								
<i>F. syc9</i>	0.29	0.24	0.24	0.25	0.24	0.22	0.22	0.16	0.00							
<i>F. syc10</i>	0.24	0.22	0.24	0.22	0.24	0.23	0.22	0.20	0.18	0.00						
<i>F. syc11</i>	0.23	0.25	0.26	0.23	0.24	0.32	0.28	0.26	0.26	0.21	0.00					
<i>F. syc12</i>	0.25	0.29	0.29	0.27	0.29	0.30	0.26	0.25	0.28	0.18	0.22	0.00				
<i>F. syc13</i>	0.27	0.24	0.31	0.24	0.28	0.28	0.27	0.24	0.30	0.19	0.27	0.16	0.00			
<i>F. syc14</i>	0.31	0.25	0.26	0.27	0.28	0.32	0.30	0.32	0.35	0.27	0.28	0.31	0.25	0.00		
<i>F. syc15</i>	0.37	0.31	0.37	0.35	0.36	0.32	0.38	0.38	0.35	0.34	0.34	0.38	0.30	0.27	0.00	
<i>F. syc16</i>	0.26	0.23	0.26	0.25	0.29	0.28	0.23	0.23	0.26	0.24	0.27	0.23	0.25	0.29	0.35	0.00

and *F. sycomorus3* were too closed with the PDV value of 0.13 (Table 5). Indeed, *F. sycomorus4* and *F. sycomorus5* were also too closed with the PDV value of 0.17. Whereas, the second subcluster contained *F. sycomorus6*, *F. sycomorus7*, *F. sycomorus8*, *F. sycomorus9* and *F. sycomorus10* genotypes, where, *F. sycomorus7* and *F. sycomorus8* were the closets genotypes with the lowest PDV of 0.10. Indeed, *F. sycomorus9* and *F. sycomorus10* were also too closed with the PDV value of 0.18, while, the third subcluster contained *F. sycomorus12*, *F. sycomorus13* and *F. sycomorus16* genotypes, where, *F. sycomorus12* and *F. sycomorus13* were too closed with the PDV value of 0.16 (Table 5). The DAMD marker revealed that the lowest PDV was recorded to be 0.10 between *F. sycomorus6* & *F. sycomorus7* and between *F. sycomorus7* & *F. sycomorus8* genotypes with the highest genetic similarity of 0.831 and 0.830, respectively. Whereas, the highest one was recorded to be 0.38 between *F. sycomorus8* & *F. sycomorus15* (with a similarity of 0.526); *F. sycomorus7* & *F. sycomorus15* (with a similarity of 0.523) and *F. sycomorus12* & *F. sycomorus15* (with a similarity of 0.500) genotypes. It worth noting that the latest genotypes were genetically too far, exhibiting a high PDV of 0.38 than the mean PDV of 0.24.

DISCUSSION AND CONCLUSION

DNA genetic diversity among 16 genotypes of *F. sycomorus* L. species has been assessed using 24 DAMD primers. Data showed that TB ranged between 4-19 bands with a mean average of 8.083 bands/ primer, whereas, PB ranged between 2-19 bands with a mean average of 6.042 bands/ primer. Another investigation by Saleh [2] reported DNA genetic diversity among 10 *F. sycomorus* genotypes using RAPD and Inter-retrotransposon Amplified Polymorphism (IRAP) markers. The previous study revealed that the RAPD marker produced TB ranging between 5-21 bands with an average of 9.778 bands/primer and PB ranging between 2-16 bands with an average of 7 bands/primer. Whereas, in IRAP marker case, TB ranged between 7-17 bands with an average of

11.125 bands/primer and PB ranged between 3-15 bands with an average of 9.438 bands/primer.

Previously, Zhou *et al.* [8] reported high levels of variation between different species and little variation between different cultivars of *O. sativa* using DAMD. Whereas, Ince *et al.* [12] reported that TB varied from 1 to 12 fragments combined with a total of 38 accession-specific DNA markers in *Capsicum* using 22 DAMD primers.

Moreover, Seyedimoradi *et al.* [17] reported a DAMD marker for characterizing 21 grapevine accessions from Iran. The previous study revealed that P% average was recorded to be 66% with a PIC average value of 0.44. Indeed, genetic dissimilarity of genotypes ranged from 0.12 to 0.67. Srivastava *et al.* [18] applied 4 DAMD primers for genetic diversity studying among 46 mango varieties. They reported 110 (with P%=87.3%) fragments with a PIC value of 0.245. In addition, Kumar and Nair [19] applied 4 DAMD primers to study phylo genetic relationships among 50 wild and cultivated accessions of 19 Indian *Citrus* genotypes. They reported a total of 45 bands, of which 35 (78%) were polymorphic with a mean genetic similarity of 0.68, indicating moderate genetic divergence among the studied accessions. Moreover, Pakseresht *et al.* [20] reported a DAMD marker for studying phylogenetic relationships of 40 landraces chickpea genotypes in Iran. The previous study revealed the PIC average 0.232, whereas, Murty *et al.* [21] applied 4 DAMD primers for studying genetic diversity in 19 *Jatropha* accessions. The previous study revealed that P% average was 91.02% with a mean PIC value 0.873. Indeed, Ince and karaca [23] applied 22 DAMD primers to study genetic diversity in *Dianthus* L. genus. The previous study revealed that TB varied between 36-168 bands and PB varied between 19-120 bands with P% ranging between 37.3-100% and PIC values ranging between 0.120-0.341.

Recently, Pinar *et al.* [24] applied 20 DAMD primers for *Citrus* molecular characterization. The previous study revealed that DAMD marker gave 296 total bands, of which, 296

(100%) were polymorphic with an average of 14.8 bands/primer. Whereas, Jain *et al.* [25] applied 12 DAMD primers for *Sechium edule* (Jacq.) Sw. molecular study. They reported a total of 97 bands (average of 8.033 bands/primer) of which 92 (average of 7.667 bands/primer) were polymorphic with P% of 94.845%.

The current study revealed that DAMD assay produced 194 bands, of which, 145 (74.742%) were polymorphic with a PIC average value of 0.219. Saleh [2] however, reported the molecular characterization of 10 *F. sycomorus* genotypes using 36 RAPD and 22 IRAP primers. The previous study revealed that RAPD assay produced 352 distinguished bands, of which 252 (71.59%) were polymorphic. As for IRAP assay, there were 178 bands, of which, 151 (84.83%) were polymorphic. Moreover, the mean PIC value was recorded to be 0.215 and 0.299 for RAPD and IRAP assays, respectively.

It worth noting that our data presented herein were comparable with that reported by Saleh [2]. Since, the effectiveness of a molecular marker for detection of DNA genetic diversity is based on P% and PIC values resulting from the application of the given marker, in this respect, in our case study, the DAMD marker was more effective than RAPD by showing the highest P% (74.742%) and PIC (0.219) values compared to RAPD with the lowest P% (71.59%) and PIC (0.215) values. This observation was in agreement with Karaca and Ince [6] who reported that the DAMD marker was more effective than the RAPD one. While, the DAMD marker was less effective than the IRAP one in the study by Saleh [2], where, IRAP marker exhibited the highest P% (84.83%) and PIC (0.299) values than that those reported in the current study.

Overall, based on data previously reported by Saleh [2] and herein regarding P% and PIC values, we could suggest that moderate genetic divergence was observed among the tested *F. sycomorus* genotypes.

From data presented herein, the current study could suggest that the 16 studied *F. sycomorus* genotypes belonged to two separate distinguished subspecies; the first subspecies contained *F. sycomorus14* and *F. sycomorus15* genotypes. Whereas, the second subspecies contained the remaining studied genotypes.

Overall, the DNA genetic diversity detected among the studied *F. sycomorus* genotypes was not observed to be related to their geographical distribution, but could be related to their common origin as previously reported by Saleh [2] who stated similar observation within *F. sycomorus* species. This finding was coherent with those reported by Machado *et al.* [30], showing the evolution of *Ficus* monoecious origin into gynodioecious types. Other studies, however, reported similar findings related to Tunisian fig based on the RAPD assay application [31].

In conclusion, data presented herein using the DAMD maker and in the previous study [2] using RAPD and IRAP markers, highlighted moderate genetic divergence among the studied *F. sycomorus* genotypes. Overall, the current study suggests that the studied *F. sycomorus* genotypes were split into two main subspecies. Indeed, DAMD-PCR assay successfully highlighted 12 unique markers characterized some

studied genotypes. Thereby, due to the importance of *F. sycomorus* as a coastal species and to confirm the actual results obtained regarding the mentioned species for DNA genetic diversity; further research works like *e.g.* Simple Sequence Repeats (SSRs) are required to confirm the number of subspecies to which *F. sycomorus* trees growing in Syria belong; especially, there is no precise inventory of the number of subspecies of this species in Syria.

LIST OF ABBREVIATIONS

CTAB	= Cetyltrimethylammonium Bromide
DAMD	= Directed Amplification of Minisatellite-region DNA
IRAP	= Inter-retrotransposon Amplified Polymorphism
P%	= Percentage Polymorphism
PB	= Polymorphic Bands
PIC	= Polymorphic Information Content
PDV	= Percent Disagreement Values
RAPD	= Random Amplified Polymorphic DNA
SSRs	= Simple Sequence Repeats
TB	= Total Bands
UPGMA	= Unweighted Pair Group Mean Arithmetic Average
VNTR	= Variable Number of Tandem Repeats

ETHICS APPROVAL AND CONSENT TO PARTICIPATE

Not applicable.

HUMAN AND ANIMAL RIGHTS

No animals/humans were used for studies that are the basis of this research.

CONSENT FOR PUBLICATION

Not applicable.

AVAILABILITY OF DATA AND MATERIALS

Not applicable.

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CONFLICT OF INTEREST

The author declares no conflict of interest, financial or otherwise.

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