Genetic diversity among *Elaeagnus angustifolia* L. populations based on some morphological traits and Random Amplified Polymorphic DNA Markers

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Abstract

Elaeagnus angustifolia L. is a Eurasian tree that has become naturalized and has various ecological, medicinal and economical uses. In this study, a combination of morphological traits and RAPD markers were used to study the presence or absence of an association between genetic variation and morphological features among five populations of E. angustifolia collected from the East Azarbaijan of Iran. Data analysis of 19 different morphological traits, according to Nei's genetic distance matrix using Nei's in GenAIEx 6.5, showed that genetic distance coefficient ranged from 0.014 (between Jolfa and Ahar populations) to 0.86 (between Jolfa/Marand and Meianeh populations). The cluster analysis based on UPGMA and dendrogram plotted using NTSYSpc 2.02 software, revealed 4 main clusters. RAPD analysis using four random primers generated 29 polymorphic bands. Accordingly, the samples were placed in 4 groups. Based on Nei's genetic distance matrix, a great genetic distance existed between Jolfa and Meianeh populations (0.167) and great genetic similarity existed between Jolfa and Marand populations (0.955). In this research, the results of morphological traits and RAPD markers showed more consistent with each other. Our results showed that RAPD analysis is a suitable method to study genetic diversity and relationships among E. angustifolia populations.

Key words: Elaeagnus angustifolia, Genetic diversity, RAPD, Morphological traits, Iran.

INTRODUCTION

Elaeagnus angustifolia L. that also named Russian olive, is an Eurasian tree, native to southern Europe, central and western Asia. Within this region it occurs primarily on coasts, in riparian areas, and in other relatively moist habitats (Zouhar, 2005). E. angustifolia L. belongs to Elaeagnaceae family with high capacity to grow over a range of environmental conditions (Asadiar et al., 2012a). The small family elaeagnaceae has three genera namely Elaeagnus L., Hippophae L. and Shepherdia Nutt and has 77 species worldwide. E. angustifolia is a deciduous tree, sometime shrub, erect, to 10m tall in cultivation (Sun and Lin, 2010). Various medicinal uses have been shown for E. angustifolia. The ripe fruits of *E. angustifolia* have been used to treat amoebic dysentery. There is general belief that leaves and fruits of the plant have antipyretic effect (Wang et al., 2006). There are large variations in the content of biologically active compounds in the leaves and fruits and the tolerance to drought, salinity and alkalinity stresses among the E. angustifolia populations. There are a variety of molecular markers to identify the valuable *E. angustifolia* resources, and classifying the populations. Some methods such as RP-HPLC biochemical markers (Wang et al., 2006), ISSR genetic markers (Assadiar et al., 2012a) and RAPD molecular markers have been used to study genetic diversity and relationship among E. angustifolia species (Assadiar et al., 2012b). RAPD molecular markers are DNA fragments from polymerase chain reaction (PCR) amplification of random segments of genomic DNA with single primer of arbitrary nucleotide sequences. RAPD markers are able to differentiate between genetically distinct individuals (Taghizad et al., 2010) and are one of the most efficient molecular methods in terms of ability to produce abundant polymorphic markers (Williams et al., 1990). The advantages of RAPD are its rapidity, simplicity and do not require previous knowledge of genome (Rahman, 2006; Tucak et al., 2008). So, RAPD analysis is a valuable tool in studying patterns of gene expression, inter- and intraspecies genetic variations and identification of specific genes using nearly isogenic variants in plant and animal research (Caetano-Anollés et al., 1991; Barker et al., 1999; Kuddus et al., 2002; Bauvet et al., 2004; Arzani and Samei, 2004; Vandemark et al., 2006). In this study, we used morphological traits and RAPD markers to investigate genetic variation among different genotypes of E. angustifolia.

MATERIALS AND METHODS

Plant material and DNA extraction

The leaves of *E. angustifolia* were collected from 5 different locations of East Azarbaijan province of Iran (Figure 1) in May 2012 (Table 1). Genomic DNA was extracted from dried leaves. Approximately 100 mg of samples was pulverized in a mortar and then extracted by cetyltrimethylammonium bromide (CTAB) method

(Doyle and Doyle, 1987). The quantity and quality of each DNA bulk sample was determined spectrophotometrically at 260 nm (nanodrop 1000, Thermo Scientific) and 1% agarose gel electrophoresis.

RAPD-PCR amplification

In this study, seven RAPD primers, which were taken

Table 1. *E. angustifolia* genotypes collected from EastAzarbaijan province of Iran.

Sample	Location	Latitude	Longitude	Habitat (m)
Campio	200041011		0	
Ea1	Jolfa	42º 75'	56° 09'	965
Ea2	Jolfa	43º 04'	55º 45'	833
Ea3	Jolfa	43º11'	55° 60'	703
Ea4	Jolfa	43º 01'	57º 43'	981
Ea5	Marand	42º 53'	57º 13'	1314
Ea6	Marand	42º 46'	56º 18'	1743
Ea7	Marand	42º 72'	56º 31'	1392
Ea8	Marand	42º 43'	57º 53'	1695
Ea9	Meianeh	41º 56'	70º 75'	1486
Ea10	Meianeh	41º 64'	69º 23'	1605
Ea11	Meianeh	41º 66'	71º 25'	1696
Ea12	Meianeh	41º 63'	71º 16'	1684
Ea13	Ahar	42º 55'	66º 75'	1453
Ea14	Ahar	42º 56'	66º 77'	1445
Ea15	Ahar	42º 59'	67º 86'	1492
Ea16	Ahar	42º 64'	67º 84'	1420
Ea17	Tabriz	42º 16'	61º 75'	1512
Ea18	Tabriz	42º 22'	60° 63'	1356
Ea19	Tabriz	42º 16'	63º 08'	1525
Ea20	Tabriz	42º 18'	63º 09'	1529

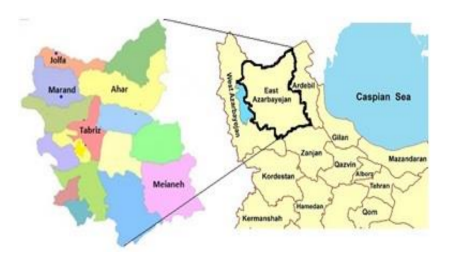


Figure 1. Geographical distribution and collection sites of *E. angustifolia* genotypes in East Azerbayejan province of Iran.

Primer	Primer sequence	GC (%)	ТМ	Number of amplified bonds	Number of polymorphic bonds	Polymorphic/amplified bondnds (%)
RP ₁	5'TGCCCGTCGT 3'	70	34	10	10	100
RP ₂	5'ACAACGCCTC 3'	60	32			
RP ₃	5'TGCCGAGCTG 3'	70	34	8	8	100
RP ₄	5'GGGTAACGCC 3'	70	34	9	8	88.88
RP ₅	5'GGTGAACGCT 3'	60	32			
RP_6	5'GGACCCAACC 3'	70	34	5	5	100
RP ₇	5'TGCGCCCTTC 3'	70	34			

Table 2. Primer sequences and RAPD products generated by primers in *E. angustifolia* genotypes.

Table 3. Morphological characters and their code for preparing matrix data.
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No.	Traits
1	Length of leaves: ≤ 6 cm (0); 6 to 7 cm (1); > 7cm (2)
2	Width of leaves: ≤ 1.3 cm (0); 1.3 to 1.7 cm (1); > 1.7 cm (2)
3	Length of leaves pedicel: ≤ 9 mm (0); 9 to 11 mm (1); > 11 (2)
4	Speckles on leaves: absent (0); present (1)
5	Color of abaxial leaf surface: yellowish white (0); silvery (1); rust- colored or ferruginous (2)
6	Color of adaxial leaf surface: yellowish white or silvery (0); rust- colored or ferruginous (1)
7	Shape of leaf blade: round or ovate (0); oblong or elliptic (1); lanceolate (2)
8	Shape of leaf apex: round or obtuse (0); acute or acuminate (1)
9	Shape of leaf margin: revolute (0); entire (1)
10	Ripe fruit color: red (0); yellowish brown or yellow to orange (1); yellowish gray (2)
11	Fruit surface type: hairy (0); scaly (1)
12	Type of fruit pedicel: erect, robust (0); slender, nodding (1)
13	Presence of wings on fruit surface: absent (0); present (1)
14	Shape of fruit: nearly globose or obovoid (0); ellipsoid (1); ovate (2)
15	Length of fruit pedicel: ≤ 2 mm (0); 2 to 6mm (1); > 6 mm (2)
16	Fruit length diameter: ≤ 4.6cm (0); 4.6 to 6cm (1); > 6 (2)
17	Fruit width diameter: \leq 3.9cm (0); 3.9 to 4.4 (1); > 4.4cm (2)
18	Shape of seed: narrow and long (0); ovate (1)
19	Seed size: ≤ 1.5cm (0); 1.5 to 1.8cm (1); > 1.8 cm (2)

from various previous studies, were used (Table 2). The reaction mixture for RAPD amplification assay had a total volume of 20 μ l, which contained 40 ng genomic DNA, 10 μ l Master mix (from BIORON company, containing 1 unit *Taq DNA polymerase*, 0.1 mM of each dNTPs, 2.5 mM MgCl₂, 0.01% Tween 20, 65 mM Tris-HCl and 16 mM (NH₄)₂SO₄ and 70 pmol primer. The amplification was carried out on a gradient thermo cycler (LabCycler/SensoQuest, Germany), with an initial step of 5 min denaturation at 94°C, followed by 45 cycles of 45 s at 94°C, 1 min at 37°C and 90 s at 72°C, and a final extension step for 7 min at 72°C. The PCR amplified products were run on a 2% agarose gel containing safe dye stain in 1× TBE buffer for 2h. Then gels were digitally photographed under ultraviolet light in a transilluminator documentation system (Gerix 1000, Biostep).

RAPD data analysis

DNA banding patterns generated were scored for the presence (1) or absence (0) of each amplified band to create binary data matrices. To assess the genetic relationships between populations based on Nei's genetic distance coefficients (Nei, 1973), NTSYS- pc 2.02 software was used to construct UPGMA (Unweighted Pair Group Method of Cluster Analysis) – dendrogram.

Morphological traits

Nineteen morphological characteristics were considered to evaluate genetic diversity as described before by

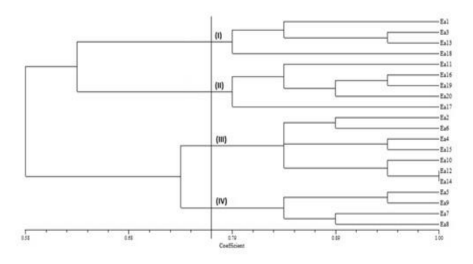


Figure 2. Dendrogram generated using UPGMA method showing relationships between populations of *E. angustifolia* using morphological characters data.

Table 4. Nineteen morphological characters used forconstruction of cluster analysis for *E. angustifolia*genotypes (C: character).

Sample	C: 1 2 3 4 5 6 7 8 9 0 1 2 3 4 5 6 7 8 9
E.a1	0101100011000011010
E.a2	0101101110000011011
E.a3	0101100011000011000
E.a4	0101101111000011010
E.a5	0101101111000011000
E.a6	0101101110000111010
E.a7	0101101111000111000
E.a8	0101101110000111000
E.a9	0101101111000011010
E.a10	0101101110000011000
E.a11	0101101111000011001
E.a12	0101101110000011000
E.a13	0101100011000011000
E.a14	0101101110000011000
E.a15	0101101111000011010
E.a16	0101101111000011000
E.a17	1101101111110011000
E.a18	0101101011010011000
E.a19	0101101111010011000
E.a20	1101101111000011000

Asadiar *et al.* (2012) (Table 3). Some traits that were identical among genotypes were not considered for further analysis (Tables 4 and 5).

RESULTS AND DISCUSSION

Matrix data was prepared according to the morphological states (Table 4). Cluster analysis of 19

morphological traits showed that the genotypes of E. angustifolia divided into 4 main clusters (Figure 2). Each of I, II and IV clusters was divided into two sub clusters while cluster III was subdivided into three sub clusters. In cluster III, E.a12 and E.a14 were common in all morphological traits. According to Nei's genetic distance matrix using Nei in GenAlEx 6.5, genetic distance coefficient ranged from 0.014 (between Jolfa and Ahar populations) to 0.086 (between Jolfa/Marand and Meianeh populations). Out of seven RAPD primers used in this study, four primers produced reproducible bands. A total of 29 polymorphic bands were identified ranging from 100 to 1500 bp. Three primers (RP1, RP3 and RP6) produced 100% polymorphic bands (Table 2). The Cluster analysis, using RAPD data showed that the genotypes of E. angustifolia were divided into 4 main clusters (Figure 3).

According to the previous plant classification based on morphological traits, each plant depends on morphological properties based on one classification unit. In this study, combination of morphological and genetically parameters were used for analyzing the relationship between the populations. Maximum calculated genetic distance based on Nei genetic distance matrix using Gen Alex 6.5 software was 0.086 between Jolfa and Meianeh populations; and maximum genetic similarity matrix was 0.986 between Ahar and Jolfa populations, according to morphological data (Tables 5 and 6). Data analysis using NTsyspc 2.02 showed that genetic distance dendrogram was including four major groups. Also, according to RAPD data analysis, Maximum calculated genetic distance was

	E.a1	E.a2	E.a3	E.a4	E.a5	E.a6	E.a7	E.a8	E.a9	E.a10	E.a11	E.a12	E.a13	E.a14	4	14 E.a15		E.a15	E.a15 E.a16	E.a15 E.a16 E.a17
E.a1	1.000																			
E.a2	0.736	1.000																		
E.a3	0.842	0.684	1.000																	
E.a4	0.842	0.894	0.789	1.000																
E.a5	0.786	0.736	0.842	0.842	1.000															
E.a6	0.736	0.894	0.684	0.894	0.736	1.000														
E.a7	0.684	0.736	0.842	0.842		0.842	1.000													
E.a8	0.684	0.736	0.736	0.736		0.842	0.894	1.000												
E.a9	0.842	0.789	0.789	0.894	0.947	0.789	0.842	0.842	1.000											
E.a10	0.684	0.842	0.842	0.842	0.894	0.842	0.894	0.894	0.842	1.000										
E.a11	0.736	0.894	0.789	0.894	0.842	0.789	0.842	0.736	0.789	0.842	1.000									
E.a12	0.736	0.894	0.789	0.894	0.842	0.894	0.842	0.842	0.789	0.947	0.894	1.000	8	00	00	00	00	00	00	00
E.a13	0.894	0.736	0.947	0.842	0.789	0.736	0.789	0.684	0.736	0.789	0.842	0.842	42 2							
E.a14	0.736	0.894	0.789	0.894	0.842	0.894	0.842	0.842	0.789	0.947	0.894	1.0	1.000	00 0.842	0.842	0.842	0.842	0.842	0.842	0.842
E.a15	0.789	0.842	0.842	0.947	0.894	0.842	0.894	0.789	0.947	0.894	0.842	0.842	42		0.789	0.789 0.842	0.789 0.842	0.789 0.842	0.789 0.842	0.789 0.842
E.a16	0.842	0.789	0.789	0.894	0.947	0.789	0.842	0.842	0.894	0.842	0.894	0.894	94		0.842	0.842 0.894 0.842	0.842 0.894 0.842	0.842 0.894 0.842	0.842 0.894 0.842	0.842 0.894 0.842
E.a17	0.631	0.684	0.684	0.789	0.736	0.684	0.736	0.631	0.684	0.736	0.789	0.789	89		0.736	0.736 0.789 0.736	0.736 0.789 0.736 0.789	0.736 0.789 0.736 0.789	0.736 0.789 0.736 0.789	0.736 0.789 0.736 0.789
E.a18	0.789	0.578	0.842	0.684	0.842	0.578	0.736	0.736	0.789	0.736	0.684	0.684	84 4			0.789 0.684 0.736	0.789 0.684 0.736 0.789	0.789 0.684 0.736 0.789 0.684	0.789 0.684 0.736 0.789 0.684	0.789 0.684 0.736 0.789 0.684
E.a19	0.789	0.736	0.736	0.842	0.894	0.736	0.789	0.789	0.842	0.789	0.842	0	0.842		0.789	0.789 0.842 0.789	0.789 0.842 0.789 0.947	0.789 0.842 0.789 0.947 0.842	0.789 0.842 0.789 0.947 0.842 0.842	0.789 0.842 0.789 0.947 0.842
E.a20	0 280			0010	0 00/	364 0	0 789	0 7 8 0	0 842	0 789	0 842	5	840		0 789	0 789 0 842 0 789	0 789 0 842 0 789 0 947	0 789 0 842 0 789 0 947 0 842	0.789 0.842 0.789 0.947 0.842 0.736	0.842 0.789 0.947 0.842 0.736

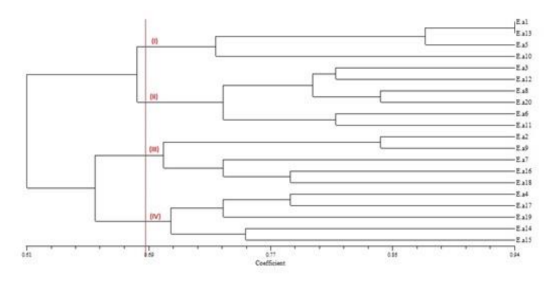


Figure 3. Dendrogram generated using UPGMA method showing relationships between populations of *E. angustifolia* using RAPD marker data.

Table 6. Pairwise popula	ation matrix o	of Nei's Ge	netic identity
based on morphological of	haracters.		

Table 8. Pairwise	population mat	rix of Nei's Ger	etic distance
based RAPD data			

Population	Jolfa	Marand	Tabriz	Ahar	Mianeh	Population	Jolfa	Marand	Tabriz	Ahar	Meianeh
Jolfa	0.000					Jolfa	0.000				
Marand	0.085	0.000				Marand	0.046	0.000			
Tabriz	0.069	0.015	0.000			Tabriz	0.075	0.094	0.000		
Ahar	0.014	0.046	0.032	0.000		Ahar	0.068	0.052	0.111	0.000	
Meianeh	0.086	0.081	0.066	0.052	0.000	Mianeh	0.167	0.146	0.150	0.152	0.000

0.167 between Jolfa and Meianeh populations; and maximum genetic similarity matrix was 0.955 between Ahar and Jolfa populations (Tables 7 and 8). In this study, the morphological data and RAPD data are consistent with each other. Existing high similarity between *E. angustifolia* populations could be due to ecological conditions in regions.

Since *E. angustifolia* is tolerant to severe drought, high salinity and alkalinity in soils, it is said to play a very important role in maintaining ecosystem function in the hyper arid areas. It is also used for various medicinal purposes (Zhang and Zhao, 1996). Therfore, various types of biochemical, morphological and molecular markers have been widely used for the analysis of genetic diversity among and between *E. angustifolia* populations worldwide. Wang *et al.* (2006) used RP-HPLC (reversed-phase high-performance liquid chromatography) for the classification and analysis of intra-specific genetic relationships of seventeen populations of *E. angustifolia*. Asadiar *et al.* (2012b) evaluated genetic relationships and polymorphism among genotypes of *E. angustifolia* collected from different locations of West Azarbaijan province using a combination of morphological traits and molecular RAPD marker. Uzun *et al.* (2015) used a combination of fruit characteristics, RAPD and ISSR markers for the evaluation of genetic variation among 56 *E. angustifolia* accessions collected from the Central Anatolian region.

Today, genetic markers are widely used for genetic diversity studies. In addition, comparison between the molecular markers is inevitable.

The present study showed that RAPD markers provide some useful information about relationship between *E. angustifolia* populations, the distribution

Table	Table 7. Similarity matrix for <i>E. angustifolia</i> populations based on RAPD data	larity ma	atrix for	. E. ang	tustifoli	a populi	ations b	based o	n RAPI	D data.										
	E.a1	E.a2	E.a3	E.a4	E.a5	E.a6	E.a7	E.a8	E.a9	E.a10	E.a11	E.a12	E.a13	E.a14	E.a15	E.a16	E.a17	E.a18	E.a19	E.a20
E.a1	1.000																			
E.a2	0.515	1.000																		
E.a3	0.666	0.606	1.000																	
E.a4	0.363	0.727	0.636	1.000																
E.a5	0.848	0.606	0.818	0.515	1.000															
E.a6	0.636	0.757	0.666	0.666	0.666	1.000														
E.a7	0.545	0.666	0.575	0.757	0.515	0.787	1.000													
E.a8	0.666	0.606	0.818	0.575	0.696	0.727	0.696	1.000												
E.a9	0.606	0.848	0.696	0.575	0.696	0.666	0.636	0.757	1.000											
E.a10		0.606	0.696	0.454	0.696	0.727	0.575	0.636	0.696	1.000										
E.a11		0.636	0.787	0.727	0.606	0.818	0.666	0.787	0.606	0.606	1.000									
E.a12		0.666	0.818	0.636	0.757	0.787	0.636	0.757	0.696	0.757	0.787	1.000								
E.a13	0.939	0.575	0.727	0.424	0.909	0.696	0.606	0.727	0.666	0.727	0.575	0.727	1.000							
E.a14		0.666	0.575	0.696	0.454	0.606	0.575	0.575	0.636	0.515	0.666	0.575	0.363	1.000						
E.a15		0.545	0.636	0.696	0.515	0.606	0.575	0.575	0.575	0.575	0.666	0.575	0.424	0.757	1.000					
E.a16	0.575	0.757	0.666	0.606	0.606	0.696	0.727	0.606	0.727	0.606	0.636	0.666	0.636	0.666	0.606	1.000				
E.a17	0.515	0.636	0.727	0.787	0.606	0.757	0.727	0.666	0.606	0.606	0.818	0.787	0.575	0.666	0.787	0.696	1.000			
E.a18	0.484	0.666	0.696	0.696	0.575	0.606	0.757	0.757	0.757	0.515	0.727	0.696	0.545	0.696	0.696	0.787	0.787	1.000		
E.a19	0.454	0.636	0.545	0.727	0.484	0.696	0.727	0.666	0.606	0.545	0.636	0.606	0.515	0.727	0.666	0.636	0.757	0.727	1.000	
E.a20	0.696	0.696	0.787	0.666	0.727	0.696	0.666	0.848	0.727	0.666	0.696	0.848	0.757	0.484	0.545	0.636	0.757	0.727	0.696	1.000

pattern, genetic variation and the geographical and ecological factors. They might also provide data for the taxonomy of the species or intraspecific relationship patterns and for the evaluation of the ecological adaptation of *E. angustifolia*.

In conclusion, molecular variation assessed in this study in combination with morphological characters of *E. angustifolia* can be useful in conventional and molecular breeding programs for this medicinal plant. But it is suggested that: (1) More samples of this plant should be collected for further genetic variation studies; (2) In addition to other molecular markers, pollen micromorphological studeis can also be used for genetic diversity assessments.

CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

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