Phenotypic and iPBS-retrotransposon marker diversity in okra (Abelmoschus esculentus (L.) Moench) germplasm

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Abstract
This study was undertaken to assess genetic and phenotypic diversity of Turkish okra (Abelmoschus esculentus (L.) Moench) germplasm of 26 landraces including three cultivars (Akköy-41, Kabakli-11, and Marmara-1) with 34 phenotypic traits and 74 iPBS-retrotransposon primers. Leaf-blade size, fruit length, fruit diameter, fruit number per plant, petiole length, plant height, stem diameter, number of stem nodes, and plant growth type (degree of branching) were the most important morphological traits contributing to the variation. Comparison of genotypes with 14 iPBS-retrotransposon primers yielded 141 bands, 34 of which (24.1%) were polymorphic, with the primer 2271 producing the highest (6) bands per primer. Cluster analysis based on phenotypic and molecular markers produced two major groups. Phenotypic based unweighted pair group method with arithmetic mean (UPGMA) dendrogram had 12 sub-groups with the highest similarity (0.63) between GAN-19/GAN-21 and MGL-6/Akköy-41 genotypes. The markers, however, produced a dendrogram with eight subgroups, pairwise genetic similarities ranging from 0.43 to 1.00, where MGL-6 singled out with a similarity value of 0.57. Howbeit, the Mantel test between both dendrograms based on the similarity matrix was insignificant.

Introduction
The okra, a member of the Malvaceae family, is taxonomically known as Abelmoschus esculentus (the binominal name is Abelmoschus esculentus (L.) Moench) and by the synonym Hibiscus esculentus L. Although the geographical origin of okra is controversial, it is thought to be originated from a tropical region of Asia or Africa (Tindall, 1983). West Africa was considered okra native land because of the presence of related wild species in the Nile Valley and Ethiopia (De Candolle, 1886; Yildiz et al., 2015a). Genetic studies have shown that okra’s origin may consist of many species from Southeast Asia, India, West Africa, or Ethiopia. It was grown by the ancient Egyptians in the 12th century B.C., and its cultivation spread to the Middle East and North Africa (Lamont, 1999). Okra is cultivated in different tropical, subtropical, and warm temperature regions of the world (Karakoltsidis & Constantinides, 1975). Around 10 million tons of okra are produced worldwide, with most of its production in Asia and Africa. Today, India is the biggest okra producer with 6 million tons, followed by Nigeria, Sudan, and Mali, respectively. In Turkey, okra is annually grown on 5640 ha with a production of 29,111 tons (FAO, 2018). Because of the similarity of the okra germplasm in Turkey to the African germplasm, it is thought that Turkey’s introduction to okra was due to the okras brought to Anatolia from the African continent (Duzyaman, 2009). Production in Turkey is widely with local types. Three commercial cultivars have been recently introduced in Turkey (TTSM, 2020).

In addition to having low essential fatty acid content, okra fruits contain important macrominerals and vitamins (Al-Wandawi, 1983), and...
also another study have shown that okra seed is rich in unsaturated fatty acids (linoleic acid etc.) and high protein quality (Savello, 1980). Okra is not very selective in terms of soil, grown in tropical, subtropical and temperate regions of the world. Although it is a self-pollinated species, cross pollination can also be seen due to bees (Lamont, 1999). And this cross-pollination provides genetic variation that allows local okra genotypes to better adapt to environmental conditions. The chromosome number was reported to be 2n=130 (Joshi & Hardas, 1956) but there are two kinds of okra genotypes diploid (2n = 2x = 60-70) and tetraploids (2n = 4x = 120-130) that were attributed to abnormalities in chromosome movement during the mitotic stage of cell division (Nwangburuka et al., 2011). It is an allopolyploid crop derived from the regular polyploid series with n=12 and has diploid chromosome number ranging between 72 and 144 (Datta & Naug, 1968).

Interspecific hybridization is possible mechanism in the genus of Abelmoschus (Reddy, 2015), furthermore, the highest variation is seen in A. esculentus species, otherwise called the Saudanien type (Kalloo & Bergh, 1993). It has been determined that there are very few or no distinguishing characteristics in the okra varieties in Turkey, Iran and India (Martin et al., 1981). Although okra in Turkey might differ from each other, it was determined that the original structure is preserved in okra germplasm resources (Duzyaman & Vural, 2002). In the study with genotypes of different origins, it was shown that the morphology of Turkish okra is similar to those originating from India and Africa (Duzyaman & Vural, 2000). Because of high phenotypic similarity among the germplasms of different origins, it becomes crucial to determine genetic variations using DNA markers. In this way, information on the genetic diversity required for the rational use of plant genetic resources can be obtained in the most reasonable way (Chakravarthi & Naravane, 2006).

It is crucial for the agricultural policies of the countries to determine the genetic diversity by using molecular methods to preserve and develop varieties and lines with high yield, quality and adaptability. In this context, molecular markers have proven to be a powerful tool in revealing genetic relationship intra and interspecies. There are different studies assessed the genetic diversity using sequence-related amplified polymorphism (SRAP) (Gulsen et al., 2007), random amplification of polymorphic DNA (RAPD) (Aladele et al., 2008; Prakash et al., 2011), simple sequence repeat (SSR) (Kumar et al., 2017), inter simple sequence repeat (ISSR) (Yuan et al., 2014), inter-primer binding site (iPBS) (Yildiz et al., 2015a; Yaldiz et al., 2018; Barut et al., 2020), microsatellite (Ravishankar et al., 2018) and amplified fragment length polymorphism (AFLP) (Akash et al., 2013) molecular markers in okra.

Retrotransposons are hereditary materials that assume a significant function in the plant genome regarding evolution, which can change the position and number of duplicates in the plant genome (Finnegnan, 1989; Ali et al., 2019). “At the ends of the retrotransposons are long terminal repeats (LTRs) that qualify as a potential tRNA primer binding site (PBS)” (Finnegnan, 1989). The iPBS markers are based on amplification of the PBS of the reverse transcriptase (tRNA complement) in two transposes opposite and close to each other (Kalander et al., 2010). This study aimed to determine the genetic diversity among different local and commercial okra genotypes using both phenotypic and DNA-based iPBS-retrotransposon markers.

Materials and Methods

Genotypes and cultivars

Twenty-six okra genotypes of A. esculentus (L.) collected from different locations in Turkey were evaluated in this study (Figure 1; Table 1). Of these genotypes 20 okra accessions (BLK-1, MGL-2, MGL-3, MGL-4, MGL-5, MGL-6, MGL-7, MGL-8, MGL-9, MGL-10, AYD-11, AYD-12, AYD-13, MGL-14, UIS-15, UIS-16, USK-17, AYD-18, GAN-19, GAN-21) were local genotypes collected by farmers from different locations in Turkey. Two (YLV-22 and YLV-23) were advanced breeding lines developed by Atatürk Central Horticultural Research Institute (ACHRS), Yalova. The list also included 3 commercially registered cultivars (Akköy-41, Kabaki-11 and Marmara-1) obtained from ACHRS and one standard commercial type (STD-20).

Figure 1. Okra genotypes and cultivars investigated in the experiment (numbers refer to row number in Table 1).
Production of plant material

Seeds collected from each genotype were sown at Akdeniz University Faculty of Agriculture Experiment Farm. The soil was off clay loam texture with a slightly alkali pH of 7.62, a high lime content of 17.7%, good organic matter content of 2.1%, total N content of 0.09%, low P content (0.0013%), high K content (0.19%), high Ca content (0.4%), optimum Mg content of (0.09%) and Mn, Zn, Cu and Fe contents respectively 2.67, 0.47, 0.25 and 1.2 mg/kg respectively. Irrigation was applied as required with dripping system, standard fertilizer was applied, and hand weeding was performed during the plant growth period.

Phenotypic/Agro-morphological traits measured

The phenotypic data of okra (A. esculentus (L.) Moench) were collected according to the criteria in the guidelines for conducting the distinctness, uniformity and stability described by International Union for the Protection of New Varieties of Plants (UPOV, 1999). A total of 34 traits were measured, including 7 agromorphological traits (seed yield per plot (SYP), seed weight per pod (SWPP), seed yield per plant (SYPP), thermal time requirements for flowering (TTRF), thermal time requirements for fruit formation (TTRFF), plant growth type (PGT) and suitability to mechanical harvest (STMH)). TTRF and commercial harvest time were calculated in degree days by average of the daily greatest and least temperatures recorded and compared to a base of temperature, 10°C (Dhankar & Singh, 2013).

DNA extraction

Leaf samples from each genotype were collected from individual healthy green plants for molecular marker analysis. Genomic DNA was extracted from 40–50 mg young fresh leaf tissue of individual genotypes, using 2% CTAB method (Doyle & Doyle, 1987) and all DNA concentration of each genotype was measured using 1% agarose gel.

PCR amplifications with iPBS-retrotransposon markers

PCR amplifications were performed under reaction conditions containing 50 ng DNA template, 0.2 mM of each dNTPs (Thermo Fisher Scientific, USA) 0.2 μM of each iPBS primers, 1X Taq Polimerase Buffer (from 1X stock ammonium buffer with 15 mM MgCl2), 1.25 U/μl Taq DNA polymerase (Ampliqon, Denmark) in the final volume using the following PCR conditions; 5 minute (min) denaturation at 95°C and 35 cycles of 30 second (s) denaturation at 95°C, 45 s annealing at 40-60°C, and 1 min extension at 72°C, followed by a final extension at 72°C for 10 min. The PCR products separated on 1X TBE Buffer (90 mM Tris, pH 8.3, 90 mM boric acid, 0.1 mM EDTA) with 2% agarose gels (Sigma-Aldrich Chemie GmbH, Germany) containing 0.5 μg/ml ethidium bromide (EtBr) and imaged under U.V. imaging system (DNR, Minilumi, Neve Yamin, Israel).

Data analysis

Polymorphic bands obtained by iPBS-retrotransposon primers used in the study were scored as present (1) or absent (0) due to the dominant nature of marker system. A dendrogram was created in “Numerical Taxonomy Multivariate Analysis System (NTSYS-pc)" version 2.1 software package (Rohlf, 1992) using UPGMA method (Michener & Sokal, 1957) to examine genetic relationships between genotypes. To create a dendrogram, DICE coefficient was based on the similarity matrix (Dice, 1945). Principal component analysis (PCA) was performed based on Jaccard similarity coefficient and two and three-dimensional graphs were created using the DCENTER and EIGEN procedures in NTSYS. Analysis was also made on phenotypic/agro-morphological data using SPSS statistical package (IBM, 2015) according to augmented statistical design. Similarity index was calculated and distance cluster dendrogram was created with S.M. coefficient and UPGMA method using NTSYS-pc ver. 2.1 software package. The eigen value of the phenotypic/agro-morphological data was calculated based on the similarity coefficient and of the variation explained. Comparison of marker and agromorphological data based on similarity matrices with Mantel test (Mantel, 1967) was performed.

Results

Variation in phenotypic/agro-morphological traits

Considerable variation was recorded for all the qualitative and quantitative plant traits investigated.
Higher variation was observed in quantitative traits compared with qualitative traits (Table S1). Variance was the highest in SYP (26273.7 g²) followed by TTRFF (2572.6-degree day²), and TTRF (2421.4-degree day²), SYPP (193.2 g²), petiole length (PL) (47.8 cm²), diameter of young fruit (FD) (29.2 cm²) and length of mature fruit (LMF) (25.3 cm²). There were 3-fold difference in plant height (PH), ranging from 40.0 cm (GAN-19, MGL-7) to 120 cm (MGL-9) whereas stem diameter (SD) ranged from 4.58 cm (AYD-13) to 9.43 cm (MGL-9) (Table S1). FD ranged from 10.57 mm (BKL-1) to 26.69 mm (AYD-18) while diameter of mature fruit (DFM) ranged from 16.86 mm (Akköy-41) to 32.18 mm (UIS-16). Time of flowering (TF) was the earliest in MGL-9 (55 and 58 days) and the latest in USK-17 and GAN-21 (68 vs 72 days, respectively). The earliest harvest was obtained from MGL-9 (61 days), YLV-23 (62 days) and BLK-1 (63 days). The highest number of fruits was obtained from Marmara-1 (16) and MGL-8 (14). The fruit number per plant (FNP) ranged from 3 (AYD-11, AYD-12, MGL-6, MGL-7 and GAN-21) to 16 (Marmara-1) with a mean of 6.23 (Table S1).

PCA with 34 traits produced 9 components with eigen values explaining 85.8% of total variation where the first 2 explaining 23.4% and 17.1% of variance, respectively (data not presented). The first three components explained 52.9% of cumulative variance. Leaf blade size (LBS) (0.793), SYPP (0.756), flower size (FS) (0.754), (FNP) (0.695), PH (0.657), LMF (0.653), SD (0.639) and fruit thickness of carpel (TFC) (0.603) had the highest contributions in the first component whereas SWPP (0.658), FD (0.608), stem number of nodes (NSN) (0.603) and seed yield per plant (SYPP) (0.602) had the highest values in the second component as opposed to PGT (plant growth type) (0.721) having the highest value in the third component (data not presented).

The biplot for PCA of the first two components discriminated 4 groups of traits that distinctly comprised plant for quantitative (PH, SD etc.), color (fruit color (FC), stem color (SC) etc.), shape (fruit shape of apex (FSI), fruit constriction of basal part (CFBP) etc.) and phenological traits (TF, TFH etc.) (Figure 2). Shape and color traits were closely related while phenological and quantitative traits were more clearly separated (Figure 2). Shape and color traits are grouped on the negative part of X and Y axis while quantitative traits were positioned on the positive scale of X and Y axis. Time related traits were positioned on Y axis (Figure 2). NSN, STMH, DFM, TFC, FNP, PL, FS, LBS, plant growth type (PDB), petiole diameter (PD) and number of fruit locules (NFL) appeared to be the most important agromorphological traits positively relating to yield (SWPP, SYPP and SYP).

Cluster analysis based on qualitative and quantitative phenotypic traits
In line with cluster analysis of 27 phenotypic data described by UPOV, the dendrogram was created according to the UPGMA method and as a result, the existence of two major groups and different sub-groups under these major groups were determined. The highest similarity (0.63) was seen between GAN-19/GAN-21 genotypes collected from Gaziantep region and MGL-6/Akköy-41 genotypes. MGL-2 and MGL-9 genotypes formed a separate group in the dendrogram (Figure 3).

The first two eigen values explained 21.05% of the total variation. In addition, 2D/3D plots of agromorphological data were obtained by PCA analysis (Figure 4; Figure 5).
Comparison of genotypes with iPBS-retrotransposon primers

Initially, 74 iPBS-retrotransposon primers were used to screen the bulk DNAs of okra genotypes for amplification. Then 14 iPBS-retrotransposon primers were determined yielding polymorphic and highly reproducible bands. In this study, a total of 26 okra genotypes collected from different locations in Turkey were analysed with 14 different polymorphic iPBS-retrotransposon primers to determine the genetic relationship between genotypes.

At the same time, characterization study was carried out according to various criteria, and the data obtained were compared with iPBS marker and the correlation between the two dendograms was determined by the Mantel test. The 14 primers (2271, 2379, 2095, 2393, 2238, 2388, 2423, 2424, 2087, 2270, 2390, 2272, 2382, 2384) yielded 141 bands, 34 of which (24.1%) were polymorphic. One to six polymorphic band was obtained per primer. The primer 2271 yielded the highest (6) while the primers 2390, 2272, 2382, 2384 produced the lowest number of polymorphic bands (1) per primer. Polymorphic band sizes ranged from 230 to 2200 bp in size while the average number of bands and polymorphic markers per primer was 10.07 and 2.42, respectively (Table 2).

Cluster Analysis Based on iPBS-retrotransposon Markers

As a result of the dendrogram created, the existence of 2 different major groups was determined. Pairwise genetic similarities ranged from 0.43 to 1.00 and maximum similarity value of 1.00 were observed between 4 genotypes (MGL-3, MGL-9, AYD-11, AYD-13). iPBS markers failed to separate the four genotypes from each other. MGL-7 genotypes created a different group, separating from the rest in dendrogram with 0.57 genetic similarity. The remaining 25 genotypes were distinguished from one another with similarity values ranging from 0.57 to 1.00. Among the 25 genotypes, the MGL-6 genotype differed with a similarity value of 0.57 (Figure 6). The first two eigen values (2.63 and 1.86, respectively) obtained with PCA explained 36.1% of the total variation. The 2D and 3D graphs showed a similar result with dendrogram and most of the genotypes clustered together, especially MGL-3, MGL-9, AYD-11, AYD-13 genotypes with similarity values of 1.00 were clustered in a close area with STD-20 cultivars in this.

Table 2. Genetic analysis of 14 polymorphic iPBS-retrotransposons markers with 26 Okra genotypes

<table>
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<tr>
<th>Primers</th>
<th>Sequence</th>
<th>Tm (°C)</th>
<th>Total Band Numbers</th>
<th>Polymorphic Band Numbers</th>
<th>Product Sizes for Polymorphic Bands (bp)</th>
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<tr>
<td>2271</td>
<td>GCCTCGGATGCCA</td>
<td>54.3</td>
<td>11</td>
<td>6</td>
<td>500-2000</td>
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<tr>
<td>2379</td>
<td>TCCAGAGATCCA</td>
<td>41.5</td>
<td>9</td>
<td>4</td>
<td>400-1500</td>
</tr>
<tr>
<td>2095</td>
<td>GCTGGGATACCA</td>
<td>44.8</td>
<td>11</td>
<td>3</td>
<td>280-600</td>
</tr>
<tr>
<td>2393</td>
<td>TACCGTAGGCCA</td>
<td>47.1</td>
<td>10</td>
<td>3</td>
<td>400-650</td>
</tr>
<tr>
<td>2238</td>
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<td>55.5</td>
<td>10</td>
<td>3</td>
<td>260-1250</td>
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<tr>
<td>2388</td>
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<td>43.4</td>
<td>10</td>
<td>3</td>
<td>300-1250</td>
</tr>
<tr>
<td>2243</td>
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<td>54.9</td>
<td>11</td>
<td>2</td>
<td>1900-2200</td>
</tr>
<tr>
<td>2249</td>
<td>AACCGACCTCTGACCCA</td>
<td>54.7</td>
<td>11</td>
<td>2</td>
<td>500-2000</td>
</tr>
<tr>
<td>2087</td>
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<td>10</td>
<td>2</td>
<td>230-250</td>
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<tr>
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<tr>
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<td></td>
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<td>10.07</td>
<td>2.42</td>
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Figure 6. Dendrogram of 26 okra genotypes constructed by UPGMA based on iPBS-retrotransposon markers.

Figure 7. Two-dimensional plot of PCA of 26 okra genotypes based on iPBS-retrotransposon markers. Dim-1 and Dim-2 explain 21.12% and 14.98% of variation, respectively.

Figure 8. Three-dimensional plot of the principal component analysis (PCA) of 26 okra genotypes based on iPBS-retrotransposon markers. Dim-1, Dim-2, and Dim-3 explain 21.12% 14.98% and 9.86% of variation, respectively.

Discussion

The okra germplasm including 3 commercial cultivars showed considerable variation in the qualitative and quantitative plant growth descriptive traits investigated. LBS, FL, FD, PL, FNP, PH, SD, NSN and PDB were the most important quantitative characteristics contributing to variation within the germplasm. Yildiz et al. (2015b) working with 30 okra genotype and 35 morphological traits reported that fruit, leaf, stem color, leaf shape, flower size, fruit-surface angularity, petal color were important characteristics to distinguish the okra accessions.

Duzyaman (2005) working with 11 Turkish genotypes and 25 morphological, horticultural and physiological characteristics found that number of locules, mature pod length and internode length, pod yield/plant, pod number/plant, pedicel length and number of branches were main traits contributing the variability. Days to first flowering, nodes per plant, length of internode, fruit weight and fruits per plant were the most important traits contributing to heterosis in okra breeding (Bhatt et al., 2016). The higher number of accessions and descriptors were suggested in order to characterise better the representative regional morphotypes (Martin et al., 1981). Earliness and fruit yield as well fruit size and fruit color are important traits for consumers and producers. The germplasm we investigated had considerable variability for the traits that may serve as genetic material for a breeding program. The germplasm included landraces with thinner fruits (STD-20), short but thicker fruits, late maturing (GAN-21), determinate growth type (AYD-13), shortest plant height (MGL-7), thinner fruit diameter (MGL-6), shortest plants and fruits (GAN-19), highest seed yield (MGL-2), late, highest seed yield (USK-17), low fruit diameter (UIS-15), wider fruit diameter, easy harvest brittle fruit stems (UIS-16), dark green fruits (AYD-18), early, taller plants, long fruits (MGL-9), wide fruit diameter (YLV-22), taller plants and fruits (BKL-1), high seed yield (MGL-14) and earliness, higher fruit number (MGL-8). Although it was not a planned descriptive trait, brittleness of fruit stems discerned during harvest of UIS-16 could be an important quality for developing easily harvestable cultivars while dark green fruits of AYD-18 and very early flowering MGL-9 may contribute to genetic variation. Small okra pods of 2-4 days (<4 cm) are preferred largely in Turkey (Duzyaman & Vural, 2003b) whereas larger pods (>7-8 cm)
cm) are marketed in North America and India (Duzyaman & Vural, 2003a). Slower fruit fibre development in some cultivars (Kyriakopoulou et al., 2014) may be suitable for marketing long pods. Further work is required for the investigation of late fibre development properties of genotypes in Turkish germplasm in order to extend marketability of okra. This quality may compensate disadvantages of early harvest failure and exploit fruit yield potential of okra genotypes. Cultivars with earliness, easily harvestable traits, higher yield and superior market value may boost commercial potential of okra. Assessing genetic diversity of germplasm is crucial before establishing successful breeding programs.

Although the difference between okras in Turkey is usually determined by morphological properties, we have shown that iPBS markers can contribute to differentiate okra genotypes. Hence, both molecular and qualitative-quantitative agro-morphological traits were used to genotype and phenotype the 26 okra genotypes collected from different locations in Turkey. iPBS-retrotransposon marker systems, previously used to define the relationship between genotypes in different plant species (Barut et al., 2020; Yaldiz et al., 2018). Polymorphism was obtained for 26 okra genotypes using 14 of the 74 iPBS-retrotransposon primers used in the study. The polymorphism rate was 24.11%. At the same time, significant phenotypic variations were identified in terms of different characters handled within the local okra genotypes from different locations. Dendrograms created using molecular and agro-morphological data showed that two major groups exist among the genotypes. As a result of marker-data analysis, MGL-3, MGL-9, AYD-11, AYD-13 and also STD-20 genotypes are similar in terms of both dendrogram and 2D/3D graph. The first two eigen values explained 36.1% of the total variation. Based on agromorphological clustering, genotypes collected from similar location were determined to be located close to each other in the dendrogram. The highest similarity (0.63) was obtained with the GAN-19 and GAN-21 genotypes collected from the same location. Akköy-41 and Marmara-1 varieties were in a separate group than Kabakli-11. However, in the dendrogram and 2D/3D graphs created by iPBS-retrotransposon molecular marker system, Kabakli-11 and Akköy-41, developed from Sultani, were in the main group with the Sultani type standard variety (STD-20), while Marmara-1, a plump variety, was in the other group. The first two eigen values obtained as a result of analysis of agromorphological data explained 21.05% of the total variation. Correlation between marker-based and agromorphological-based dendrogram created according to similarity matrix was determined with Mantel test, was, however, insignificant (r = -0.11).

Marker-assisted molecular studies on okra are rare in the world. Previously, there were only two molecular marker studies with okra genotypes found in Turkey where SRAP, iPBS and SSR molecular markers were used (Gulsen et al., 2007; Yildiz et al., 2015a). At the end of a study using 66 okra genotypes and 13 polymorphic IPBS-retrotransposon primers, 88 bands and 40.2% total polymorphism rates were obtained as a result (Yildiz et al., 2015a). Higher total polymorphism rates of Yildiz et al. (2015a) probably originated from higher number of genotypes tested. Gulsen et al. (2007) using 31 Turkish and 2 randomly selected U.S. genotypes and applying 39 RAPD primer combinations obtained approximately 50% of the 97 bands obtained which were polymorphic for 23 genotypes. It was determined that 17 of these 23 genotypes were separated from each other with an average of 0.93 similarities. However, although the UPGMA dendrogram based on 33 phenotypic markers distinguished all genotypes, the geographic relationship between okra genotypes was not detected.

In another study with 44 okra genotypes collected from different locations in India, a total of 104 bands were obtained with 14 RAPD primers randomly amplifying the genome, and 74.03% polymorphism was obtained. These 14 primers were found to make a clear distinction between genotypes (Prakash et al., 2011). The reason for the higher rate of polymorphism obtained in their study was the fact that the centre of origin of okra is Indian subcontinent with considerable variation. Another reason for the higher polymorphism rates obtained is thought to be the use of the RAPD molecular marker system, which probably provides random amplification to produce a higher number of bands. Work on the assessment of okra germplasm is not sufficient and hence further studies are required with higher number of landraces, accessions, lines and cultivars representing major okra growing areas and dominant types in Turkey. Many of the previous work cited varied in the traits investigated. Further studies should apply wider number of marker systems and commercial cultivars as well as morphological descriptors universally used by breeders of okra plant. It was reported that genetic base of Turkish okra germplasm is narrow requiring inclusion of external diversity to national collections to increase variability (Yildiz et al., 2015b). Because of this narrow genetic base, polymorphism detected by limited number of molecular marker systems was low and hence further work is required with the inclusion of the external germplasm having the qualitative and quantitative traits scarce in Turkish national gene pool and testing new marker systems and phenotypic markers.

Conclusion

LBS, LMF, FD/DMF, PL, FNP, PH, SD, NSN, and PDB were the most important characteristics contributing to the variation in the okra germplasm. The 14 iPBS primers yielded 141 markers, 34 of which (24.1%) were polymorphic. Cluster analysis based on phenotypic and iPBS-retrotransposon data produced two major groups with 12 and 8 subgroups respectively without any
significant correlation between the two in Mantel test. Selection of parents in okra breeding program by phenotypic descriptors in conjunction with iPBS-retrotransposon marker system may aid better selection for conservation and breeding studies.

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Author Contribution

FK: Conceptualization, Data Curation, Formal Analysis, Methodology, Project Administration, Resources, Supervision, Validation, Visualization, Writing – review and editing; SNY: Formal Analysis; CB: Formal Analysis, Investigation, Visualization, Writing – Original Draft Preparation; NY: Formal Analysis, Data Curation; and NM: Conceptualization, Methodology, Project Administration, Supervision, Validation, Writing – review and editing.

Additional Information


References


