BSA and molecular markers screening for salt stress tolerant mutant of *Petunia* obtained in *in vitro* culture

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ABSTRACT: In this study, we performed BSA to identify genetic markers linked to salt tolerance. We tested the genetic diversity among four bulked DNA samples of EMS induced mutant clones and one bulked DNA sample of non-mutated clone of Petunia for salt tolerance in *in vitro* callus cultures using RAPD and ISSR markers. Out of the 36 RAPD and 16 ISSR primers identified, 25 and 13 were effectively used to amplify genomic DNA of all the five bulked samples, respectively. In total, 114 RAPD amplification products were obtained, of which 28% were polymorphic and 2% were genotype-specific bands. Out of the 64 ISSR amplification products obtained, 51% were polymorphic and 1% was genotype-specific bands. Results of this study indicated the existence of two patterns of distorted segregation among the studied markers. The first one indicates the differences between non-mutated clones of *Petunia* and its putative mutants. The second one was observed only between putative mutants and putative mutants tested for salt tolerance in *in vitro* culture. Both RAPD and ISSR analysis successfully detected the association with changes induced by chemical mutagenesis and salinity. Furthermore, our results indicate that BSA method can be useful in the rapid detection of molecular markers for further marker-assisted selection.

Key words: bulk segregant analysis, chemical mutagenesis, ISSR, RAPD, *Petunia × atkinsiana* D. Don, salt stress, tissue culture.

INTRODUCTION

Salinity is one of the abiotic factors that influences growth and vigor of many crops. However, exposure of crops to high salinity and other than abiotic stress factors such as drought, temperature, chemical toxicity, and so on affects its production. Nearly 20% of the cultivated land and about half of the irrigated land are affected by high salinity (PATADE et al., 2006). Therefore, high priority is warranted in breeding programs, which aim salinity tolerant crops (BIDABADI et al., 2011). Tolerance of high salinity of many agricultural crops has been reported (SHALABY & EL-BANNA, 2013). However, floriculture breeding program are not actively selecting new cultivars for salt tolerance. The most effective means that can deal with soil salinity is to grow cultivars that can establish and be productive on such soil (MIRI et al., 2014). *Petunia × atkinsiana* D. Don. belongs to *Solanaceae* family and is often considered a secondary plant model-system, due to various favorable biological features. Cultivar Prism Red has large flowers and made heavy texture largely produced for home-consumption and for gardening (VANDENBUSSCHE et al., 2006; BERENSCOT et al., 2008). Due to the important...
role of the Solanaceae family in agronomic and ornamental crops, salt tolerance in this family was examined. Salt tolerant plants can be derived through somaclonal variation with induced mutations. To maintain high productivity under adverse growing conditions, many breeders search for more tolerant forms using traditional breeding approach or artificial selection; although, much emphasis is currently being laid on molecular techniques. To achieve this researchers aim to obtain crops that demonstrate an increased tolerance to salt stress or drought using in vitro cultures from inter alia, banana (BIDABADI et al., 2011), potatoes (GOPAL & IWAMA, 2007), sweet potatoes (LUAN et al., 2007), and cauliflower (FULLER et al., 2006). Usually, somaclonal variation with or without chemical or physical mutagenesis is used as a source of plants’ new genotype for crop improvement (BAGHWAT & DUNCAN, 1998; JAIN, 2000; BAIRU et al., 2011; MIRI et al., 2014). However, a combination of somaclonal variability along with mutagenesis induced in in vitro cultures may considerably increase the level of variability in the resultant variants that may impart increase tolerance to environmental stress factor. However, several variants derived by this method can be either unstable or non-heritable because of the epigenetic changes in the resultant crop rather than the genetic changes. To make the collection useful for breeders, morphological and molecular characterization of the germplasm is necessary. Application of molecular tools can provide new approaches to increase the salt tolerance of many crops and horticultural species. Genes associated with salt tolerance have been identified in model plants, like Arabidopsis (YOKOI et al., 2002). Detailed knowledge of genome and transcription for most floricultural species is not yet available. Therefore, information about the genes involved, collected from other species, might be used as, markers in order to accelerate the selection process for this property (MUNNS, 2005).

Bulked segregant analysis (BSA) is a method that can be used for the rapid screening and verification of useful molecular markers during the selection and breeding process. It can be used to identify individual markers associated with a specific phenotype. BSA can efficiently and rapidly identify genetic markers associated with a particular trait (MASOJČ et al., 2009). In addition to BSA, random amplified polymorphic DNA (RAPD) and inter simple sequences repeat (ISSR) markers used in this study provide greater opportunities in determining the exact relationship between a specific DNA sequence and the examined trait and also in determining the exact relationship between a specific DNA sequence and the examined trait and also in determining a plant’s identity using particular genes. Both of these methods are popular DNA fingerprinting methods because they are inexpensive, highly polymorphic, quick and easy to perform.

In this study, we tested genetic diversity among four bulked DNA samples of ethyl methane-sulphonate (EMS)–induced mutant clones and one bulked DNA sample of non–mutated clone of Petunia × atkinsiana D. Don cv. Prism Red to identify genetic markers linked to salt tolerance in in vitro callus cultures using RAPD and ISSR markers.

MATERIALS AND METHODS

Plant material

The samples used in this study were leaves of Petunia × atkinsiana D. Don cv. Prism Red. Clones were obtained from callus culture, treated with 0.5mM EMS and screening for salt tolerance using 50, 100 and 150mM NaCl in in vitro culture (KRUPA-MAŁKIEWICZ et al., 2017). Genetics analysis based on molecular markers using BSA method in bulked samples as follows: (1) control plants derived from callus not treated with chemical mutagen; (2) plants derived from callus treated with 0.5mM EMS; (3) plants derived from callus treated with 0.5mM EMS and cultured on MS medium (MURASHIGE & SKOOG, 1962) supplemented with 50mM NaCl; (4) plants derived from callus treated with 0.5mM EMS and cultured on MS medium supplemented with 100mM NaCl; and (5) plants derived from callus treated by 0.5mM EMS and cultured on MS medium supplemented with 150mM NaCl. Each bulk contained 12 individual samples selected to have identical phenotypes and increased reaction to salt stress. Explants that did not indicate tolerance to salt stress were dying out and were not taken for further research, due to the lack of a sufficient amount of plant material. Bulks were prepared by mixing equal proportions of DNA from each individual in each bulk. All bulks were subjected to genetic analysis using RAPD and ISSR techniques to search for differentiating polymorphisms. Only DNA fragments that differentiated between the control bulk and putative mutants’ bulk were considered mutation linked markers.

Plant DNA extraction

Total genomic DNA was isolated separately from lyophilized leaves (20mg) of Petunia × atkinsiana D. Don in -56°C using the standard
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protocol of DNeasy Plant Mini Kit (Qiagen). Qualitative and quantitative DNA isolates were made using a spectrophotometer Epoch (BioTek) by spectroscopy method UV-Vis and visually checked on 1% agarose gel.

**RAPD amplification**

PCR amplification was carried out with a set of 36 oligonucleotides, arbitrary primers (Biomers) according to the method described by WILLIAMS et al. (1990). The PCR-RAPD reaction and temperature profiles of each step were optimized and standardized to laboratory condition and type of chemical reagents, especially polymerase used in this study. Modifications included amount of polymerase, number of reaction cycles, annealing temperature and elongation time to ensure optimal conditions and obtain stable and reproducible amplification products. Amplifications were performed in 15µl reaction volume containing 10×PCR buffer with (NH₄)₂SO₄, 2.0mM of dNTPs, 2.0mM MgCl₂, 0.5µM of primer, 1.0U Taq DNA polymerase (Thermo Scientific) and ~10ng/µl of template DNA. PCR-RAPD reactions were carried out in a total volume of 25µl containing 50ng of template DNA, 1U Taq DNA polymerase (Thermo Scientific), 2.5µl 10×PCR buffer, 2.0mM dNTP and 2.0mM MgCl₂. 16 primers (Metabion, Germany) were used for the PCR. DNA was amplified using a DNA Engine Dyad Thermal Cycler (Biorad) and using the following program: initial denaturation step at 94°C for 3min, 10 cycles - denaturation 94°C 1min, annealing - 37°C 30 s, elongation 72°C 30s, 35 cycles - 94°C 1min, annealing 37°C 30s, elongation 72°C 1min with final extension step at 72°C 5min.

**ISSR amplification**

PCR reactions were carried out in a total volume of 25µl containing 50ng of template DNA, 1U Taq DNA polymerase (Thermo Scientific), 2.5µl 10×PCR buffer, 2.0mM dNTP and 2.0mM MgCl₂. 16 primers (Metabion, Germany) were used for the PCR. DNA was amplified using a DNA Engine Dyad Thermal Cycler (Biorad) and using the following program: initial denaturation step at 94°C for 3min, 10 cycles - denaturation 94°C 1min, annealing - 37°C 30 s, elongation 72°C 30s, 35 cycles - 94°C 1min, annealing 37°C 30s, elongation 72°C 1min with final extension step at 72°C 5min.

**RESULTS**

Genome screening using RAPD and ISSR analysis revealed genetic polymorphism between the bulked DNA samples of *Petunia × atkinsiana* D. Don cv. Prism Red treated with mutagen from the control bulk samples. Two types of segregation patterns in the markers were observed in this study. First polymorphism was observed in control bulk and bulks from the explants treated with chemical mutagen (EMS) (Figure 1). The second polymorphism differentiated within putative mutants of *Petunia* and putative mutants tested for salt stress (Figure 1).

**RAPD analysis**

Of the 36 RAPD primers initially screened, 25 were effectively used to amplify genomic DNA of all five bulked samples. Length of bands varied from 380 to 1820bp in size. On an average, each primer produced four bands. In total, 114 RAPD products were obtained, of which 64 (65%) were monomorphic, 41 (28%) were polymorphic, and 9 (2%) - genotype-specific bands (Table 1). The primer pr29 produced the maximum number of bands (nine bands), of which four were polymorphic bands. Only five primers (pr879, pr611, pr110, pr310 and pr483) generated monomorphic bands. The RAPD profile of the selected putative mutant, putative mutants tested for salinity and lines from the control plants revealed genetic polymorphism. The pr539, pr429 and pr797 primers generated four polymorphic bands characteristic only for all EMS putative mutants and three (pr88, pr1100 and 1049) were only for the control (Table 2). In this study, we recorded an interesting observation in case of RAPD profile obtained from sixteen primers (pr176, pr269, pr447, pr598, pr29, pr66, pr88, pr99, pr107, pr139, pr815, pr875, pr144, pr519, pr920, pr1049); we obtained specific bands only for EMS putative mutants tested for salt tolerance (Table 2).

**ISSR analysis**

Five bulked samples were fingerprinted using 13 of the 16 ISSR primers generating 64 amplicons ranging from 410 to 1560bp. Of these, 23 (45%) were monomorphic and 38 (51%) were...
polymorphic bands (Table 1). On an average, each primer produced five bands. More number of bands (eight bands) was generated by primer 826, and less number of bands (two bands) by primer 846. Only one primer (808) generated monomorphic bands. The polymorphic bands were exhibited by twelve primers, and out of these five primers (843, 853, 813, 829, 846) generated an amplification product in

Table 1 – Characteristics of fragments amplified. Number of monomorphic, polymorphic and genotype-specific loci generated in the reaction with RAPD and ISSR primers for control and putative mutants of Petunia × atkinsiana D. Don, containing different EMS and NaCl treatments.

<table>
<thead>
<tr>
<th>Technique</th>
<th>Primer No.</th>
<th>Fragment size range (bp)</th>
<th>Loci</th>
<th>Genotypes of Petunia</th>
<th>Mutant 50mM NaCl</th>
<th>Mutant + 100mM NaCl</th>
<th>Mutant + 150mM NaCl</th>
<th>Total generated amplicons</th>
</tr>
</thead>
<tbody>
<tr>
<td>RAPD</td>
<td>pr176, pr269, pr429, pr447, pr539, pr598, pr99, pr107, pr139, pr310, pr797, pr815, pr879, pr875, pr828, pr1049</td>
<td>1820-380</td>
<td>320/64 (65%) 133/41 (28%) 9/9 (2%)</td>
<td>69 79 104 101 109</td>
<td>462/114</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ISSR</td>
<td>859, 843, 810, 819, 853, 849, 840, 808, 813, 823, 826, 829, 846</td>
<td>1560-410</td>
<td>115/23 (45%) 121/38 (51%) 3/3 (1%)</td>
<td>25 32 60 60 62</td>
<td>239/64</td>
<td></td>
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</tbody>
</table>
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Table 2 - Specific products from RAPD and ISSR primers for EMS induced putative mutants, putative mutants tested for salt tolerance medium (MS supplemented with 50, 100 and 150mM NaCl) and for control.

<table>
<thead>
<tr>
<th>Technique</th>
<th>Primers and length size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Putative mutants tested for salt tolerance medium</td>
<td>RAPD</td>
</tr>
<tr>
<td></td>
<td>ISSR</td>
</tr>
<tr>
<td>Putative mutants</td>
<td>RAPD</td>
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<tr>
<td></td>
<td>ISSR</td>
</tr>
<tr>
<td>Control</td>
<td>RAPD</td>
</tr>
<tr>
<td></td>
<td>ISSR</td>
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</table>

100% of the accession tested. An intense non-control 28 specific bands in PCR reactions with eleven primers was obtained among the selected salt tolerant lines putative mutants (Table 2). Furthermore, in our experiment, we noticed that three ISSR primers (819, 849 and 823) generated one band that was characteristic of only all EMS putative mutants of *Petunia* (Table 2). Moreover, one genotype-specific band for control bulked sample was generated by primer 823 (Table 2).

**DISCUSSION**

In this study, molecular characterization was performed to detect mutation and somaclonal variation among *Petunia × atkinsiana* D. Don cv. Prism Red explants obtained in *in vitro* callus culture. We used two types of molecular techniques to amplify different genomic regions of *Petunia*: RAPD and ISSR. Many authors (LAKSHMANAN et al., 2007; MIRI et al., 2009; SHALABY & EL-BANNA, 2013; FAROKHZADEH & ALIFAKHERI, 2014) have indicated that these techniques, based on PCR reaction are a powerful and reliable tool in genetic studies for improvement in tolerance to environmental stresses in many crop plants. These techniques are a simple starting point for comparative analysis and screening that indicate the loci directly related to the observed variability. However, genotyping a large population using these technique is time-consuming and expensive. QUARRIE et al. (1999) have suggested grouping of plants according to the expression of a particular trait and extracting DNA from these bulks. BSA has been successfully used to rapidly identify markers linked to any specific locus with different traits, such as changes in *Coffea arabica* explants regenerated by indirect and direct somatic embryogenesis (SANCHEZ-TEYER et al., 2003), sex in jojoba (SHARMA et al., 2008), resistance to pre-harvest sprouting in rye (MASOJČ et al., 2009), and DNA markers linked to salt tolerance traits in wheat (BHUTTA & HANIF, 2013), indicating their wide use in genetic analysis. Differences in results obtained from various studies on levels of polymorphism can be attributed to differences among the genotypes chosen for the studies. However, polymorphisms usually result in the presence or absence of an amplification product from a single locus. The presence of products that can be polymorphic can be useful as genetic markers. When markers linked to traits of interest are available, a marker-assisted selection can be performed. For this purpose, highly saturated linkage maps can provide a choice of markers closely linked to a specific trait (MUNNS, 2005).

BSA using RAPD and ISSR markers in this study provided information on genetic diversity caused by environmental stress factor such as chemical mutagen (EMS) or salt (NaCl) in *Petunia × atkinsiana* D. Don cv. Prism Red in *in vitro* callus culture. We performed molecular analysis by assuming the selection of genetic markers that may differentiate the control sample and four bulked samples of the plants treated with the chemical mutagen with an indication of their correlation with the trait - increased tolerance to salinity. After analyzing the results of molecular markers’ segregation, 133 RAPD and 121 ISSR polymorphic loci were selected. Furthermore, analysis of electrophoregrams revealed two patterns in segregation of molecular markers within the
examined bulked groups. The first pattern pointed to the differences between the plants’ constituting control (non–mutated) and putative mutants and was observed in all variants treated with EMS. The use of RAPD markers for the genetic analysis allowed to obtain 17% of the differentiating products and 14% ISSR markers. Such strong bidirectional effect of the disruptive selection on the segregation of alleles was detected for six RAPD markers (pr539, pr429, pr797, pr88, pr1100, pr1049) and for three ISSR markers (819, 849, 823). Thus, it can be concluded from electrophoretic separation images that the use of a chemical mutagen (EMS) caused numerous mutation in the Petunia genome, and the number of identified markers differentiating the examined samples indicates their dispersal throughout the entire genome of the analyzed species.

The second pattern was observed within the putative mutants. The polymorphism of markers involved an existence in the presence or absence of bands between the putative mutants tested on MS media with different concentrations of salt (NaCl) and putative mutants propagated on MS media with no salt addition. We observed 22% of this kind of pattern of distorted segregation that was characteristic for sixteen RAPD markers (pr176, pr269, pr447, pr598, pr29, pr66, pr88, pr99, pr107, pr139, pr815, pr875, pr144, pr519, pr920, pr1049) and 44% for eleven ISSR markers (859, 843, 810, 819, 853, 849, 840, 813, 826, 829, 846). This type of segregation thus indicates the presence of an additional genetic variability caused by abiotic stress factor, which is salinity.

Low genetic diversity observed in our study may be because the genomic regions conferred to EMS and NaCl salt stress were tagged; this is because the primers generated polymorphism between non–mutants, putative mutants and putative mutants tested for salt tolerance in bulked DNAs used to characterize these 12 genotypes.

Identification of many genetic markers that differentiate the examined bulked samples confirms that the occurrence of increased tolerance to salinity as a stressor is controlled by many differentiating loci located on many sites in the genome. This confirms that many different genes and interactions between them control this trait.

According to YOKOI et al. (2002) it is possible to transfer a single gene in a plant to increase its tolerance by many folds. However, inter– or intra–species hybridization induces mutations or somaclonal variation in cell and tissue cultures to produce a salt-tolerant cultivar is very slow with limited success. The literature on obtaining a new variety with an increased tolerance through conventional breeding approach is scarce (TAHIRA et al., 2006). Traditional approaches are limited by the complexity of stress tolerance traits, low genetic variation and field components under stress, and lack of efficient selection techniques (BHUTTA & HANIF, 2013; HADI & FULLER, 2013). Moreover, it happens that the resistance to salt stress is subject to changes during plant growth. Therefore, conventional breeding programs can be more productive if they are integrated with newer technologies of plant molecular biology.

CONCLUSION

Our study detected two RAPD and ISSR markers significantly associated with changes induced by chemical mutagenesis and salt stress factor (NaCl). Till date, there are few scientific reports on the use of BSA method to determine genetic variability between the putative mutants obtained in in vitro culture or putative mutants characterized by an increased tolerance to environmental stress. Indication of molecular markers that differentiate the analyzed bulked group is just a starting point to verify the usefulness of these markers in the selection work on a broader genetic material. Hence, an attempt to identify the marker loci determining the observed variability of the trait was undertaken. Results of research confirmed the hypothesis that part of the marker loci within which genetic changes were noted, with the participation of chemical agent or stressor, will be differentiating (polymorphic loci) in the analyzed pooled groups. These preliminary investigations using BSA method could be useful in marker–assisted breeding programs that can be used as a candidate marker for further gene mapping.

Use of RAPD and ISSR markers in this study has given newer opportunities for plant biologists to study the relationship between traits and their genetic control. The BSA is a valuable alternative approach that overcomes the necessity to genotype every member of a population. BSA and their genetic control. The BSA is a valuable alternative approach that overcomes the necessity to genotype every member of a population. BSA allows a faster validation of molecular markers for an identification of genotypes with differentiated allele frequency between the lines selected for the examined trait. In this study, we have shown that BSA coupled with physiological studies can help to identify traits important in determining salt stress resistance in Petunia × atkinsiana D. Don cv. Prism Red and provide molecular markers for this trait. The BSA should be a valuable aid for plant biologists in the future. These results suggest the importance...
of studying the salt response of ornamental potted plants to help the growers and gardeners to select the species, which are more tolerant to salt stress.

DECLARATION OF CONFLICTING INTERESTS

The authors declare no conflict of interest. The founding sponsors had no role in the design of the study, in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results.

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AUTHORS’ CONTRIBUTIONS

MKM conceived and designed experiments. MKM and AB performed the experiments and carried out the lab analyses. MKM prepared the draft of the manuscript. All authors critically revised the manuscript and approved of the final version.

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