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R Anandan

Assistant Professor (Biotechnology), Department of Genetics & Plant Breeding, Faculty of Agriculture, Annamalai University, Annamalai Nagar, Tamil Nadu, India

Rohini Srivastava

Senior Scientist, NRC Plant Biotechnology, IARI, Pusa Campus, New Delhi, India

S Murugan

Department of Genetics & Plant Breeding, Faculty of Agriculture, Annamalai University, Annamalai Nagar, Tamil Nadu, India

M Prakash

Department of Genetics & Plant Breeding, Faculty of Agriculture, Annamalai University, Annamalai Nagar, Tamil Nadu, India

Correspondence R Anandan

Assistant Professor (Biotechnology), Department of Genetics & Plant Breeding, Faculty of Agriculture, Annamalai University, Annamalai Nagar, Tamil Nadu, India

Sonication and vacuum infiltration assisted genetic transformation in bhendi [Abelmoschus esculentus (L.) Moench] using cry2Aa gene for pest resistance

R Anandan, Rohini Srivastava, S Murugan and M Prakash

Abstract

Somaclonal variation and recalcitrance nature were the major bottlenecks of *agrobacterium* mediated genetic transformation in Bhendi (*Abelmoschus esculentus* (L.). Therefore, *Agrobacterium* mediated in planta transformation protocols was standardized using sprouted seed explants of bhendi cv. MDU-1, an important vegetable. *agrobacterium tumifaciens*-mediated transformation was performed using EHA105 strain harboring the binary vector pBinAR carrying *cry2Aa* gene under the control of CaMV35s promoter and *npt II* gene as a selectable marker. The transformation event consisted of sonicating the explants for 3 minutes, vacuum infiltration (750 mm of Hg) for 2 minutes in *Agrobacterium* (pBinAR-*cry2Aa*) and cocultivation for 3 days in MS medium with acetosyringone (100µM) showed transformation efficiency of 12.5%. The presence and integration of *npt II* and *cry2Aa* transgenes into the bhendi genome was confirmed by polymerase chain reaction (PCR). Work is in progress to grow the T₀ seeds for molecular characterization of the inserted transgene using southern and western blot methods.

Keywords: Bhendi, cotyledon, npt II, cry2Aa

Introduction

Bhendi/okra (*Abelmoschus esculentus* (L.) belongs to Malvaceae family. It is a widely grown valued and nutritious vegetable. It is cultivated throughout the world especially in all tropical and subtropical regions of Asia and Africa. In India, okra is a major vegetable crop and 4.8 million tonnes of pods are produced from an area of 0.45 million hectares (Narendran *et al.*, 2013) [10]. It green pod contains dietary fiber, minerals, vitamins and oxidation inhibitors like xanthin and lutein (Kumar *et al.*, 2010; Kaur *et al.*, 2013) [3, 4]. Seeds are the rich source of edible oil (13–22%) and protein (20–24%). Okra mucilage, a high molecular weight monosaccharide polymer, can be used as food additive to modify the food quality in terms of food stability, texture and appearance properties by acting as emulsifier, thickener and gelling agent or texture modifier (Noorlaila *et al.*, 2015). In pharmaceutical industry, it is being considered as potential hydrophilic polymer and binding agent for tablets. In dairy industry, mucilage could be used as ice cream stabilizer to retard ice recrystallization and melting down of ice cream (Giese, 2014).

Among of them Yellow Vein Mosaic Virus (YVMV) transmitted by whitefly (*Bemisia tabaci*) is the most serious disease of okra (Ali *et al.*, 2000). Okra is severely affected by the shoot and fruit borer (*Helicoverpa armigera*), and the larvae bore into shoots or fruits (pods) and consume the internal contents, causing the withering of the plant and reduction in marketable value of the pods (Brar *et al.*, 1994). Frequent use of pesticides by the farmers, without recognizing the vector(s) may create toxic residues in the food chain. Recognizing its economic importance there is a massive scope of utilizing modern biotechnology for further improvement of okra, because conventional plant breeding methods are difficult and time consuming. Therefore, genetic transformation technology could be used to overcome this problem; hence a regeneration protocol is prerequisite (Mitra, 2011).

In-planta transformation is an efficient, quick and tissue culture independent system for crop plants improvement. It is a useful system for those plants that lack tissue culture and regeneration system. Therefore, the direct in-planta methods are commonly used for the transformation of many important genes into several plant species such as *Glycine max*, *Arachis hypogeae*, *Arabidopsis thaliana*, *Raphanus sativus*, *Cicer arietinum*, *Beta vulgaris*, *Gossypium hirsutum*, *Solanum lycopersicum* and *Brassica juncea*. The main objective of this study is to transform okra cv. MDU-1 *via* sonication and vacuum infiltration of sprouted seeds

with *Agrobacterium tumifaciens* harboring a synthetic *cry2Aa* gene for resistance against fruit/pod borer.

Materials and methods

Surface sterilization and explant preparation

The healthy MDU1 seeds were acquired from Vegetable Research Station, Palur. These seeds were surface sterilized using 0.1% mercuric chloride (HgCl₂) for 1 min and in 70% (v/v) ethanol for 1 min and then washed thoroughly for 6 times with sterile double distilled water to eliminate the sterilants completely. The surface sterilized seeds were kept overnight in 500 ml Erlenmeyer flask containing 100 ml liquid half strength MS medium (Murashige and Skoog, 1962) ^[9] in an orbital shaker (120 rpm) under complete darkness (Pre-cultured seeds). The pre-cultured seeds were used as explants for transformation events.

Effect of kanamycin on bhendi seeds of cv. MDU 1

For selection of transformed plants from explants an optimal concentration of kanamycin was determined culturing the seeds on MS medium supplemented with different concentrations of kanamycin (15, 30, 50, 70, 90, 100, 120 mg/l). A total of twenty explants were used with five replications per treatment and the experiment was repeated thrice. For each treatment, data on drying and survival percentage of explants were scored 2 to 3 weeks after initiation of culture.

Preparation of Agrobacterium culture

Single colony of *A. tumefaciens strain* EHA 105 harbouring pBinAR-*cry2Aa* gene (1.863 kb) flanked by CaMV35s promoter and OCS polyA terminator and *nptII* as selectable markers was used for transformation (Figure 1).

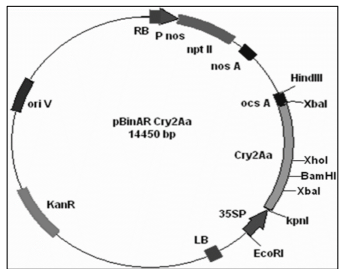


Fig 1: Physical map of pBINAR harboring cry2Aa gene

This construct was obtained from Dr. Rohini Sreevasthsa, senior scientist, NRCPB, New Delhi. Single colony of *A. tumefaciens* strain was inoculated in 3 ml of YEP broth and allowed to grow overnight in a rotary shaker at 37°C for 125 rpm. 1ml of overnight grown culture was inoculated in 30 ml of YEP broth and subculture is done. The cell suspension is maintained at 0°C for 20 minutes by keeping it on ice. Then, it is centrifuged at 5000 rpm for 10 minutes at 4°C. The supernatant was discarded and the pellet was resuspended in freshly prepared infiltration solution for immediate use.

Transformation via Sonication and Vacuum infiltration

The in planta transformation protocols in bhendi using vacuum infiltration sonication and durations, acetosyringone concentrations were followed according to previous protocols with some slight modifications (Manickavasagam *et al.*, 2015) ^[6]. The sterilized explants were transferred into Agrobacterium suspensions containing acetosyringone (100µM). Sonication was carried for 0, 1, 2, 3, 4, 5 and 6-min using a bath sonicator (model 1510 Branson, Branson Ultrasonics, Kanagawa, Japan). Then the seeds were transferred into fresh Agrobacterium suspension and subjected to vacuum infiltration for 0, 1, 2, 3, 4, 5 and 6-min at 750 mm of Hg using a desiccator (Tarsons, Kolkata, India) connected to a vacuum pump (Indian high vacuum pumps, Bangalore, India). The sonicated cum vacuum-infiltrated explants were incubated in Agrobacterium suspension medium supplemented with optimized concentration of acetosyringone 100 µM for 1 h under dark for Agrobacterium infection. Following this, the Agrobacterium infected seeds were blot dried and then co-cultivated in MS medium without hormone containing 100 µM of acetosyringone for 3 days. The infected seeds were washed with sterile distilled water containing 500 mg/l cefotaxime (Alkem laboratories, Mumbai, India) and blot-dried on a sterilized filter paper and inoculated onto solid MS basal medium containing appropriate hygromycin or kanamycin as selectable marker and incubated at 25 ± 2 °C under 16 h photo period. The explants were sub-cultured twice at 5 days interval. After 4 weeks, the well rooted survival seedlings were transferred to earthen pots containing the potting mixture (1:1:1 of red soil, sand and farm yard manure) kept in green house and maintained 85% relative humidity at 25 ± 2°C for acclimatization.

Molecular analysis of putative transgenic plantlets

The antibiotic resistant plants were subjected to molecular confirmation for the transgene integration using PCR with gene specific primers. The primers used for amplification of the *nptII* gene were 5'-GAGGCTATTCGGCTATGACTG-3' and 5' ATCGGGAGGGGCGATACCGTA-3', generating a 750 bp product and the primers for *cry2Aa* gene were 5'-GGG CAC TGT GTC CTC CTT CCT CCTC-3' and 5' GGG GAG ATG GTG AAG CCG GTG TAG-3'resulting in a 1200 bp product.

DNA extraction

Extraction of the DNA from the samples was carried out according to the procedures of Doyle and doyle (1990). The DNA pellet was resuspended by using 100 μ l TE buffer and kept at -20°C for long term use. The concentration of DNA was estimated spectrophotometrically. In spectrophotometric analysis, 5 μ l of DNA was diluted to 3000 μ l of TE buffer. The spectrophotometer readings were recorded at 260 and 280 nm. DNA concentration was calculated using OD values at 260 nm using the following formula concentration of DNA (μ l/ml) = OD at 260 nm X 50.

Results and discussion

A single report on genetic transformation of okra requires lot of time to develop transformed okra plants (Narendran *et al.* 2013) ^[10]. Hence, to establish an efficient, less time consuming in planta transformation method was performed based on our previous reports on sugarcane and brinjal (Mayayan *et al.* 2013; Subramanyam *et al.* 2013) ^[2, 7]. In

addition, year round availability, easiness of explant handling, and efficient selection of transformed plantlets raised from seeds motivated us to select seed as an explant for in planta transformation.

In the present study 50 mg/l kanamycin concentrations in the medium caused a drastic decrease in both the frequency of regeneration and number of shoots per explant, hence this concentration was used for the selection of transformed shoots (data not shown). Using npt II as a selectable marker gene and kanamycin as a selection agent is widely used system for screening transformants in a large variety of plants like mulberry (Bhatnagar and Khurana, 2003) [1], chickpea (Mehrotra *et al.*, 2011) [8]. Here, also we used the same selectable marker and were able to achieve complete suppression of non-transformed plants with optimized dose of kanamycin (50 mg/l). The identification and development of

cry genes in transgenic crops for pest management has turned out to be a major accomplishment. A major limitation has been specificity of the *Bt* toxins to only a certain group of lepidopterans pests.

Sonication and vacuum-infiltrated were performed on the seeds of bhendi as explants. The explants pre-incubated with Agrobacterium suspension + acetosyringone were sonicated for various time durations (0-60 min) using bath sonicator. After standardization of sonication, the sonicated seeds were vacuum-infiltrated along with pre-incubated further suspension-containing Agrobacterium optimized concentration of acetosyringone (100 uM) for different time durations (0–6 min) at a constant pressure (100 mm Hg) using vacuum chamber. Combination of sonication and vacuum infiltration was also tested for transformation efficiency (Table 1).

Table 1: Influences of sonication duration and vacuum infiltration duration on in planta transformation efficiency of bhendi cv. MDU 1

Sonication duration (min)	Vacuum infiltration time (min)	No. of seeds infected	Mean no. of seeds germinated ^a	Mean no. of <i>cry2Aa</i> positive explants	Transformation efficiency (%)
0	0	100	10	-	-
10	-	100	14	-	-
20	-	100	23	-	-
30	-	100	34	2	6
40	-	100	30	-	-
50	-	100	25	-	-
60	-	100	20	-	-
-	1	100	14	-	-
-	2	100	42	-	-
-	3	100	35	-	-
-	4	100	30	2	5.5
-	5	100	24	-	-
-	6	100	20	-	-
30	1	100	18	-	-
30	2	100	48	-	-
30	3	100	31	4	12.5
30	4	100	24	-	-
30	5	100	18	-	-
30	6	100	13	-	-

^a- explants on section medium (MS+ kanamycin 50 mg/l)

In sonication treatments, the transformation efficiency gradually increased with increasing sonication duration up to 30 min, which resulted in the transformation efficiency of 8.5%. It was hypothesized that a longer duration of sonication has an inhibitory effect on seed germination. The obtained results were accordance with the previous report, where 30 min sonication was found to be optimal to achieve maximum transformation efficiency of 43%, and beyond 30 min sonication, the seed germination rate and the transformation efficiency was reduced in bhendi (Manickavasagam *et al.*, 2015) ^[6].

In infiltration treatments, among various infiltration durations tested, 3 min vacuum infiltration at a constant pressure (100 mm Hg) using vacuum chamber recorded higher transformation efficiency of 6%. Vacuum infiltration creates a negative atmospheric pressure which removes the air and creates empty spaces within the seed. An increase in the pressure allows the *Agrobacterium* suspension into the empty spaces of the explant/seed, and the Agrobacterium infects the meristematic tissue effectively (Subramanyam *et al.* 2011;

Mariashibu *et al.* 2013; Jaganath *et al.* 2013) [11, 7, 2]. Beyond 3 min vacuum infiltration, due to the severe *Agrobacterium* infection, the germination rate was reduced, which ultimately led to low transformation efficiency. The obtained results were in concurrence with the previous reports (Subramanyam *et al.* 2011; Mariashibu *et al.* 2013; Jaganath *et al.* 2013) [11, 7, 2]. Combined effects of sonication (30 min) and vacuum infiltration (3 min), enhanced the transformation efficiency up to 12.5% in bhendi.

To confirm the integration of the *cry2Aa* transgenes into the bhendi genome, PCR was performed on the genomic DNA isolated from randomly selected 25-day-old putatively transformed and control plants. For PCR analysis, the primers were used to amplify the 1200-bp coding region of the *cry2Aa* gene. The obtained PCR results revealed a 1200-bp amplified fragment from putatively transformed bhendi plants DNA (Fig. 3a, lanes 3–7). However, no such amplified fragment was observed from the control plant DNA. The obtained results proved that the *cry2Aa* gene was successfully integrated into the bhendi genome (Figure 2).

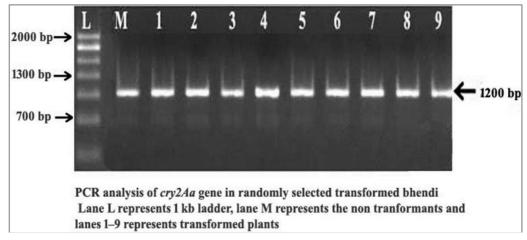


Fig 2: PCR analysis of putative transformed bhendi for amplification of cry2Aa gene

Conclusion

A simple, reproducible, and an improved Agrobacterium mediated in planta transformation system has been established for recalcitrant bhendi cv. MDU 1 by optimizing various factors influencing the transformation efficiency. Optimized parameters, sonication (30 min), vacuum infiltration (3 min), acetosyringone (100 μm), and co-cultivation (3 day) enhanced transformation efficiency of 12.% in bhendi when compared with previously published reports.

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