Sugarcane (NCo310) Transient Transformation Using uidA Reporter Gene

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ABSTRACT

Background: Sugarcane is a monocotyledonous crop that is cultivated in the tropical and subtropical regions of the world. One of the most important criteria, influencing the efficiency of the sugarcane transformation is known to be related to physical and biological factors during the transformation procedure.

Objectives: The objective of this research was to study the response of callus induction and embryogenic callus production and to identify the major parameters controlling DNA delivery by particle bombardment into sugarcane (Saccharum officinarum L.) cv. NCo310.

Materials and Methods: For callus induction and embryogenic callus production, leaf base segments were subjected to in vitro culture medium supplemented with two plant growth regulators (2,4-D and Dicamba). Results showed that 1 mg L-1 2,4-D was significantly influential in callus induction and embryogenic callus production. Considering both physical and biological factors, the efficiency of DNA (uidA gene) delivery was assessed by scoring transient GUS (gene (β-glucuronidase) expression in bombarded tissues.

Results: The highest transient GUS expression was obtained when callus was bombarded with the construct harboring rice Act1 promoter, and having 9 cm target distance, 25 inHg vacuum pressure, 1 µm gold particles, 12.5 µg DNA per bombardment and one day pre-culture prior to the bombardment.

Conclusions: A bombardment procedure suitable for elite sugarcane varieties was developed, which allowed high-efficiency DNA delivery combined with reduced damage to target tissues.

Keywords: Sugarcane; Callus Induction; Embryogenic; DNA Delivery; Particle Bombardment; Transient GUS Expression

1. Background

Sugarcane belongs to the grass family (Poaceae), which is cultivated primarily for its ability to store high concentrations of sugar in the internodes of its stem (1). Modern sugarcane varieties are complex interspecific hybrids primarily involving crosses between the species S. officinarum L. and S. spontaneum L. that have been produced through intensive selective breeding (2). Varieties of sugarcane are highly heterogeneous and are generally
multiplied vegetatively by stem cutting (3). An advantage of tissue culture amongst many other approaches is the production of high quality and uniform planting material that can be multiplied under disease-free conditions (4). Tissue culture techniques have been widely used in Saccharum spp. for various purposes. Meristem tip or shoot tip cultures (using apical meristems) have been used as a tool to produce virus-free plants (5). Genotype is another important factor influencing the efficiency of callus production and plant regeneration capacity (6). Burner (7) and Gandonou et al. (8) reported that genotype highly affects callusing responses in sugarcane. Somatic embryogenesis offers an efficient regeneration system for the production of a large number of uniform plants within a short period. Somatic embryogenesis in sugarcane has been reported using immature inflorescences (9), apical meristems (10) and young leaves (11). Biolistic mediated transformation is preferred over Agrobacterium-mediated transformation as it does not require construction of a specialized vector. Also, the advantage of biolistic technology for direct gene transfer into plant cell/tissues has facilitated plant transformation on monocots such as maize (12), wheat (13), rice (14), barley (15), palm date (16) and oil palm (17). The efficiency of biolistic transformation depends on certain physical and biological parameters (18).

2. Objectives

In the present study, sugarcane cv. NCo310, which is considered as a widely used cultivar was evaluated for callus induction, embryogenesis and transformation via particle bombardment. Successful transformation was complemented with the desired genes and putative transgenic plants could be selected precisely.

3. Materials and Methods

All chemicals were purchased from Merck (Germany), unless otherwise stated.

3.1. Plant Materials and Explants Preparation

The sugarcane (S. officinarum L.) cv. NCo310 plants were obtained from Iranian Sugarcane Research and Training Institute, and used as the mother plant. Different treatments were used for sterilization of the explants. The 5 mm of the stem apex were surface disinfected with 70% ethanol followed by a subsequent hot water (50°C) treatment for 2 h and submerged in varying concentrations (0.2, 0.5, and 0.7%) of mercuric chloride (HgCl2) for 10 min followed by three rinses with sterile dH2O (10 min each).

3.2. Culture Media and Conditions

The MS medium (19) supplemented with different concentrations of 2, 4 Dichlorophenoxyacetic acid (M1: 1 mg.L⁻¹; M2: 2 mg.L⁻¹; M3: 3 mg.L⁻¹; M4: 4 mg.L⁻¹) and Dicamba (M5: 4 mg.L⁻¹; M6: 6 mg.L⁻¹) and 30 g.L⁻¹ sucrose was used for callus induction. The pH was adjusted to 5.8 with 1 N NaOH and all media were solidified with 8 g.L⁻¹ agar before autoclaving for 20 min at 121 °C. Five explants per Petri dish were cultivated and cultures were kept in dark at 28 °C for one month. Callus induction percentage was determined after two weeks. After one month of culture, the number of embryogenic calli was recorded. Three Petri dishes were bombarded for each treatment. Histochemical GUS assays were conducted 24 h after bombardment.

3.3. Plasmid DNA and Microparticle Preparation

The plasmids used for the transient expression experiments are listed in Table 1. Gold (1 µm in diameter) or tungsten particles (1.1 µm in diameter; BioRad, USA) were coated with plasmid DNA following the protocol accompanied by the biolistic PDS-1000/He with some modifications (BioRad, USA). A volume of 50 µL of particle solution (60 mg.L⁻¹ prepared in 50% glycerol), together with 50 µL CaCl2 (2.5 M) and 20 µL spermidine (0.1 M) were added to 1µg of DNA one by one while vigorously vortexed. The microcarriers were allowed to settle for 1 min and pelleted by spinning for 2 s in a microfuge. After discarding the supernatant, the pellet was washed twice by 140 µL of 70% ethanol and 140 µL of 100% ethanol. After adding 48 µL of 100% ethanol and resuspension of pellet by vortex, 6 µL of the DNA-coated microcarrier suspension was loaded into the center of a macrocarrier, air dried and used for bombardment. Unless otherwise stated, particle bombardment was performed with a single bombardment and 1 µg DNA was used. The range of variations from the standard procedure is listed in Table 2. Bombardment parameters were tested independently, maintaining other conditions as in the standard procedure.

<table>
<thead>
<tr>
<th>Plasmid Type</th>
<th>GUS Gene</th>
<th>Promoter Type</th>
<th>Plasmid Size, kb</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>pBI121</td>
<td>+</td>
<td>CaMV 35S</td>
<td>14.7</td>
<td>Clontech, USA</td>
</tr>
<tr>
<td>pBI221</td>
<td>+</td>
<td>CaMV 35S</td>
<td>5.6</td>
<td>Clontech, USA</td>
</tr>
<tr>
<td>pAct1-D</td>
<td>+</td>
<td>Act</td>
<td>7.5</td>
<td>McElroy et al. (20)</td>
</tr>
<tr>
<td>pUB121</td>
<td>+</td>
<td>Ubi</td>
<td>16.7</td>
<td>In this study</td>
</tr>
</tbody>
</table>

Abbreviations: CaMV 35S: cauliflower mosaic virus 35S; Act: actin; Ubi: ubiquitin
Table 2. Particle Bombardment Parameters Tested Independently for Their Effect on DNA Delivery to Sugarcane Callus

<table>
<thead>
<tr>
<th>Bombardment Parameters</th>
<th>Conditions Tested</th>
</tr>
</thead>
<tbody>
<tr>
<td>Macrocarrier type</td>
<td>gold and tungsten</td>
</tr>
<tr>
<td>Chamber vacuum pressure, inHg</td>
<td>23 and 25</td>
</tr>
<tr>
<td>Distance from macrocarrier to target tissue, cm</td>
<td>6, 9, 12</td>
</tr>
<tr>
<td>Number of bombardments</td>
<td>single and double</td>
</tr>
<tr>
<td>Plasmid type</td>
<td>pBI121, pBI221, pAct1-D, pUBI121</td>
</tr>
<tr>
<td>DNA concentration, µg</td>
<td>0.5, 1, 2.5, 5, 12.5, 25</td>
</tr>
<tr>
<td>Osmoticum type</td>
<td>manitol, sorbitol, manitol+sorbitol</td>
</tr>
</tbody>
</table>

3.4. Histochemical GUS Assay and Assessment of Transient Expression

Bombarded calli remained on the same plates for 24 h before assay. GUS expression analysis was performed by adding 1 ml of X-gluc staining solution to transformed calli, followed by incubation for 24 h at 37 °C, as described by Jefferson et al. (21) with some modifications. The staining solution consisted of phosphate buffer (500 mM), EDTA (10 mM), Triton-X100 (0.1%) and X-gluc (100 mM) dissolved in dimethylformamide (DMF). The pH of the final solution was adjusted to 7.5. Transient GUS activity was recorded as the number of blue spots in each treatment under a Stemi 2000-C binocular microscope (Zeiss, Germany).

3.5. Statistical Analyses

Samples (Petri dishes) in an experiment were completely randomized with 3 replicates. Statistical analysis of data was conducted using one-way ANOVA by the PROC-GLM program of SAS (22). Analysis of variance was performed for each treatment. Means were separated at the 5% probability level with the Tukey test when a significant F ratio occurred (P < 0.05).

4. Results

The explants were treated with different aseptic conditions before transfer to the callus induction medium. The treatment of the explants with hot water (50 °C) for 2 h before incubation in different concentrations of mercuric chloride resulted in an efficient surface disinfection. It was also observed that the lowest percent of subsequent contaminations could be obtained when the leaf sheath explants were treated with 0.7% HgCl2 for 10 min (Figure 1). Each explant showed different responses on different combinations of growth regulators. The optimum conditions for the generation of embryogenic calli in NCo310 cultivar were determined. Effects of three concentrations of 2,4-D (1 mg.L⁻¹; 2 mg.L⁻¹ and 4 mg.L⁻¹) and two concentrations of dicamba (4 mg.L⁻¹ and 6 mg.L⁻¹) on the production of callus and embryogenic calli (yellow and compact) were investigated. Results showed that 1 mg.L⁻¹ 2,4-D had significant effects on the total number of explants, which produced embryogenic callus compared to other concentrations of 2,4-D and dicamba (Figure 2A and 2B). Most of the explants cultured on callus induction medium exhibited swelling after 5 - 7 days followed by callusing within the next two weeks and embryogenesis (the yellow and compact embryogenic calli) after one month (Figure 3). These embryonic calli were selected as target tissues to establish a transformation system for sugarcane cultivar NCo310 by optimizing the physical and biological parameters affecting transient expression of uidA gene, defined by the number of blue spots (Figure 4).

![Figure 1. Effect of % HgCl2 on Explant Sterilization](Image)

4.1. Particle Bombardment Parameters

4.1.1. Effect of Promoter Type

Comparison of transient expression level of the GUS gene under control of different promoters in sugarcane calli was performed. It was interesting to check the effectiveness of two monocot-derived promoters (Act1 and Ubi) and CaMV 35S promoter in sugarcane calli. For this reason, four plasmids with three different promoters (pBI221 and pBI121 harboring CaMV 35S, pAct1-D harboring rice Act1, and pUBI121 harboring rice ubiquitin promoter) carrying the GUS reporter gene were bombarded after precipitating with gold particles. Among the four constructs tested, pAct1-D and pUBI121 vectors showed significant increase in GUS expression with 69 ± 8 and 38 ± 3 blue spots, respec-
tively. Whereas, no significant difference was observed between the two other plasmids, pBl221 and pBl22i with 11 ± 2 and 5 ± 4 blue spots, respectively (Figure 5A).

![Figure 2](image2.png)

**Figure 2.** Effect of Different Concentrations of 2,4-D (M1, 1 mg.L-1; M2, 2 mg.L-1; M3, 3 mg.L-1; M4, 4 mg.L-1) and Dicamba (M5, 4 mg.L-1; M6, 6 mg.L-1) in MS medium on Callus and Embryogenic Callus Induction; Means Within Column Followed by the Same Letter are not Significantly Different (P > 0.05)

![Figure 3](image3.png)

**Figure 3.** Embryogenic Callus Induced From Leaf Explants of Sugarcane cv. NC310

A, Explants adaptation; B, Callus induction; C, Callus propagation; D, Embryogenic callus developed after four weeks of culture in the MS medium

4.1.2. Osmotic Conditioning of Callus

Callus culture on MS medium supplemented with 2 mg.L-1 2,4-D and 0.2 M sorbitol + 0.2 M manitol significantly improved transient GUS expression (56 ± 17) in comparison with calli cultured on 0.4 M sorbitol (23 ± 3) and 0.4 M manitol (22 ± 7) (Figure 5B). Osmotic treatments of embryogenic calli were used four hours before bombardment and one day after bombardment followed by histochemical GUS staining

![Figure 4](image4.png)

**Figure 4.** Transient GUS Expression Pattern Inembryogenic Calli of Sugarcane Bombarded With the Construct Carrying the Ubi Promoter

4.1.3. Quantitation of DNA and Particle Type

To evaluate the effect of plasmid DNA concentration for efficient coating of the microcarrier, we used different concentrations of pAct1-D at 0.5, 2.5, 12.5 and 25 µg per shot. It was found that 12.5 µg DNA per shot gave the highest transient GUS expression (702 ± 43). Lower concentration of DNA (0.5 µg) was found to give lower blue foci (43 ± 3). This observation could be due to the insufficient amount of DNA in proportion to the microparticles for delivery. The average numbers of blue foci for other concentrations, 2.5 and 25 µg were 109 ± 4 and 232 ± 35, respectively (Figure 5C). Gold (1 µm) particles showed significantly higher number of the blue foci per shot (205 ± 23) than tungsten (1.1 µm) particles (48 ± 10) (Figure 6A).

![Figure 5](image5.png)

**Figure 5.** Osmotic Conditioning of Callus A, Explants adaptation; B, Callus induction; C, Callus propagation; D, Embryogenic callus developed after four weeks of culture in the MS medium

4.1.4 Effect of Times of Bombardments and Chamber Vacuum Pressure

There was no significant difference in GUS expression carrying out single (105 ± 24) or multiple (two) bombardments (148.8 ± 9) on sugarcane callus (Figure 6B). Also we found that DNA delivery at 25 Inch of Mercury (inHg) significantly increased the efficiency of transient GUS expression (422 ± 26) in comparison with lower vacuum pressure (23 inHg) resulting in 215 ± 19 blue spots (Figure 6C).
**Figure 5.** Effect of Biological Parameters on Transient expression of the GUS Gene in Bombarded Sugarcane Embryogenic Calli

![Bar chart A](image1)

A, Plasmid type; B, Osmoticum type; and C, DNA quantity per shot. Data represent the Mean ± SD determined from at least three replicates and were analyzed using one-way ANOVA (PROC-GLM program of SAS). Different letters indicate significant differences at the level of P < 0.05 within a parameter.

**Figure 6.** Effect of Physical Parameters on Transient Expression of the GUS Gene in Bombarded Sugarcane Embryogenic Calli

![Bar chart B](image2)

A, particle type; B, bombardment number; C, vacuum pressure and D) distance from stopping screen to target tissue. Data represent the Mean ± SD determined from at least three replicates and were analyzed using one-way ANOVA (PROC-GLM program of SAS). Different letters indicate significant differences at the level of P < 0.05 within a parameter.
4.1.5. Selection of Appropriate Distance From Microcarrier Launches Assembly to the Target Tissue

There are three adjustable distances that influence the delivery of particles using the PDS-1000/He system, namely, gap distance (between rupture disk and macrocarrier), macrocarrier travel distance (between macrocarrier and stopping screen) and the target distance (between stopping screen and target plate). When the distance between calli and macro carrier was 9 cm, a significantly higher number of blue spots (112 ± 29) were detected in comparison with 6 and 12 cm distances which showed 71 ± 5 and 44 ± 7 blue spots, respectively (Figure 6D).

5. Discussion

The commercial sugarcane is not considered as pure S. officinarum but a hybrid species; complex polyploid with a large number of chromosomes. This does not provide a reliable system for genetic manipulation through conventional means. There are a few reports on successful genetic transformation of the sugarcane (23, 24). One of the most important criteria, influencing the efficiency of the transformation is known to be related to physical and biological factors during the transformation procedure. At present, transformation of sugarcane is mostly achieved by microprojectile bombardment (25). In this study, we have tried to determine which combination of physical and biological parameters gives the highest transient expression of the introduced uidA reporter gene in NCO310 sugarcane. The expression level of the introduced gene highly depends on the strength of the promoter that controls transcription of the transgene. For this purpose, transient expression of the uidA reporter gene was evaluated under the control of a number of promoters. Although the CaMV 35S promoter has been shown to be active in most tissues of several plant species (26), in this research it was found to be the weakest of the promoters tested in sugarcane. While differences in transient gene expression level were observed among the monocot promoters, the highest activity was obtained with the Act1 promoter, which is in agreement with previous reports on the Act1-D promoter activity in other monocots such as date palm (16) and rice (27). In conclusion, the histochemical staining results indicated that the rice Act1 promoter is preferred for stable expression of foreign genes in sugarcane callus. The distance from the microprojectile launch site to the biological target can affect the velocity of microprojectiles and consequently transformation rates (16, 28). Our results showed that 9 cm is the most effective distance in particle bombardment of NCO310 by pAct1-D harboring plasmid. To improve the coverage of targeted areas, multiple bombardments were used. Although this can increase the efficiency of transformation when the primary delivery is not efficient, yet at the same time, it may result in an increase in the damage done to the target tissues (29) and thus are undesirable for stable transformation. In cassava (30), banana (31) and Brazilian maize inbred lines (28), double bombardment was shown to be more effective in transient expression than single. But our results showed no significant difference between single and double bombardment which is in agreement with the reported results on oil palm (32), and date palm (16). Non-metabolisable osmotica such as mannitol, sorbitol and PEG or combinations of these are commonly used to minimize cytoplasm leakage from target cells (33, 34). We found that the combination of mannitol and sorbitol significantly increased the average numbers of blue foci. Our results showed a significant increase in GUS expression when a gold particle was used as a microcarrier. Gold microcarrier is biologically inert, non toxic and uniform in size and shape. It generates better performance in particle bombardment than tungsten that could be potentially harmful to some cell types, via acidification and degradation of DNA bonds (18). Vacuum pressure has an important role in the acceleration of the microcarrier from stopping screen to the target tissue (32). A partial vacuum is generally applied in the bombardment chamber because the residual gas in the chamber could cause rubbing forces inducing a deceleration of particles (18). In our research 25 inHg significantly increased the efficiency of transient GUS expression. Lower vacuum pressures did not allow particles to reach the target tissue. This is in agreement with previous results obtained on Catharanthus roseus cells and wheat tissues (25). Increasing the concentration of DNA which was shown to have no significant effect on transgene expression, could instead be considered un economical and cause more particle aggregations resulting in poor efficiency of cell penetration and cell injury (34). Increasing the concentration of DNA from 2.5 µg up to 25 µg in bombarded calli of date palm increased the average number of blue foci but with no significant difference among them (16). This is in agreement with our data showing that 12.5 µg DNA per shot gave the highest transient GUS expression. In conclusion, we provided in this work an optimized protocol for embryogenic callus production and biolistic-mediated transient transformation of sugarcane callus, which allows one to obtain a high number of transformed cells. Such an approach will be very useful to introduce agronomical important genes into tropical sugarcane lines.

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Authors’ Contribution

Mostafa Motallebi and the authors have conducted the
Sugarcane Transient Transformation
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