Screening of Brazilian soybean genotypes with high potential for somatic embryogenesis and plant regeneration

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Abstract – The aim of this work was to identify Brazilian soybean (Glycine max) genotypes with potential to respond to in vitro culture stimuli for primary somatic embryo induction, secondary embryo proliferation and plant regeneration. Differences among eight tested cultivars were observed at each stage. Two cultivars, IAS-5 and BRSMG 68 Vencedora, were selected for the evaluation of the capacity for embryo differentiation and plant regeneration. These cultivars had high embryo induction frequencies, repetitive embryogenic proliferation, and low precocious embryo germination in the initial experiment. The effect of abscisic acid (ABA) and charcoal addition on plant regeneration was investigated. The addition of ABA to proliferation medium and of ABA and activated charcoal to maturation medium increased embryo differentiation rates, which resulted in a higher number of regenerated plants. The BRSMG 68 Vencedora cultivar was found to have a high potential for embryo induction, embryo proliferation and plant regeneration. The potential of this cultivar for somatic embryogenesis was similar to that observed for cultivar IAS-5, which is currently used for soybean transformation in Brazil. BRSMG 68 Vencedora may be a good alternative genotype for soybean genetic engineering via somatic embryogenesis protocols.

Index terms: Glycine max, Brazilian cultivars, embryogenic potential, genetic engineering, soybean transformation.

Introduction

The absence of highly efficient regeneration procedures is one of the main limiting factors in gene transfer technology in soybean (Tomlin et al., 2002). Proliferating somatic embryos are one of the most suitable targets for soybean genetic manipulation (Sato et al., 1993; Droste et al., 2002; Homrich et al., 2008). Several studies have shown differences among soybean genotypes in their capacity to respond to the different steps of somatic embryogenesis (Bailey et al., 1993; Santos et al., 1997; Simmonds & Donaldson, 2000; Droste et al., 2001; Tomlin et al., 2002; Hiraga et al., 2007; Yang...
et al., 2009). However, the development of genetically engineered soybean has been limited to a small number of cultivars that respond well to in vitro culture (Tomlin et al., 2002).

The efficiency of regeneration and transformation of soybean is genotype-dependent and remains effective mainly for the variety ‘Jack’ and a few other genotypes in the USA (Walker & Parrott, 2001; Schmidt et al., 2005). In Brazil, the IAS-5 and Bragg cultivars have been used in transformation studies (Droste et al., 2002; Homrich et al., 2008).

The genotype-dependence justifies the screening of soybean cultivars that are more responsive to the induction of somatic embryogenesis and regeneration, including the new genotypes constantly produced by genetic engineering.

The objective of this study was to evaluate Brazilian soybean genotypes for their capacity to respond to embryogenesis induction, embryo proliferation and plant regeneration.

**Materials and Methods**

Bragg and IAS-5 are North American adapted cultivars, commonly used in genetic improvement programs, previously evaluated for their embryogenic response (Santos et al., 1997; Droste et al., 2001). They were used in the first experiment together with cultivars BRSMG 68 Vencedora, BRS Torena, BRS 137, BRS 154, Embrapa 48 and MG/BR 46 Conquista, developed and released by Brazilian breeding programs, and recommended for commercial growth in different states of Brazil (Brasil, 2009).

The culture procedure is illustrated in Figure 1. Young pods containing immature seeds of 3–4 mm in length were harvested from field-grown plants. Pod sterilization, cotyledon excision and placement on D40 induction medium (Bailey et al., 1993) were performed according to Droste et al. (2002). Six Petri dishes, containing ten pairs of cotyledons each, were prepared for each cultivar. Cultures were incubated at

**Figure 1.** Soybean somatic embryo induction, proliferation and plant regeneration. A, primary somatic embryos induced from immature zygotic cotyledons; B, cluster of globular embryos on proliferation medium; C, cluster of histodifferentiated somatic embryos, after 30 days on maturation medium; D, somatic embryos germinating on conversion medium; E, regenerated plant.
26±1°C and 22.5 μE m⁻² s⁻¹ light on a 16/8 hour light/dark cycle for somatic embryogenesis induction.

After 30 days on the induction medium, the cotyledons with primary somatic embryos were transferred to D20 proliferation medium (Wright et al., 1991). After 20 more days, dishes were scored for percentage of embryogenic cotyledon pairs and number of somatic embryos per cotyledon pair. Somatic embryo clusters were then removed from the cotyledons and transferred to fresh proliferation medium, in which they were maintained for five months with subcultures every 14 days.

In order to evaluate the proliferation capacity, a set of four clusters of secondary somatic embryos at the initial globular stage, with 2–3 mm in diameter each, were weighted as a unit and placed in 60x15-mm Petri dishes containing proliferation medium. Five dishes were used for each cultivar. After 28 days, with a subculture on the 14th day, the weight of the four clusters in each Petri dish was again recorded. The response to proliferability was the weight gain (mg), calculated as the difference between the initial and final weights.

To induce histodifferentiation, clusters of proliferating embryogenic tissue, with 2–3 mm in diameter, were placed on modified MS6 maturation medium (Finer & McMullen, 1991), containing MS salts (Murashige & Skoog, 1962), B5 vitamins (Gamborg et al., 1968), 6% sucrose, 0.3% Phytage, at pH 6.4, before autoclaving. Four embryogenic clusters were placed in each Petri dish, and four dishes were used per cultivar. After four weeks, the embryos were separated and transferred to fresh maturation medium for four more weeks. The number of histodifferentiated embryos and the number of precociously germinated embryos was recorded. Precocious germination refers to germination of embryos on maturation medium, i.e., before their desiccation and transfer to conversion medium.

In a second experiment, the IAS-5 and BRSMG 68 Vencedora cultivars were selected for evaluation of germination and conversion capacities. Abscisic acid (ABA) at 50 μmol L⁻¹ was added to D20 proliferation medium in the last month, before the embryo clusters were transferred to maturation medium. ABA concentration was based on a previous study, which showed that globular embryos treated with ABA have higher conversion capability than untreated ones (Weber et al., 2007). Sixteen clusters of secondary somatic embryos at the initial globular stage, of each cultivar, were distributed in four Petri dishes containing proliferation medium with ABA. The same number of embryo clusters was placed on proliferation medium without ABA (control). After 30 days, the embryo clusters treated with ABA were transferred to MSM6 maturation medium with ABA at 50 μmol L⁻¹ and 0.5% activated charcoal (Bailey et al., 1993). The untreated embryo clusters were transferred to MSM6 medium without ABA and charcoal (control). After 30 days in the maturation medium, the treated and the untreated histodifferentiated embryos were individually transferred to fresh MSM6 medium free of ABA and charcoal for four more weeks. A sample of 100 histodifferentiated nonprecociously germinated embryos per cultivar was randomly picked and placed in dry, sterile dishes for a 2-day desiccation treatment. Partially desiccated embryos were plated on MSO conversion medium, containing MS salts, B5 vitamins, 3% sucrose, 0.3% Phytage, at pH 6.4, before autoclaving. After 45 days in the conversion medium, the number of germinated embryos and converted plants were scored. Germination refers to root and shoot emission, while conversion was recorded as the development of the branched root and formation of at least one trifoliolate leaf (Walker & Parrott, 2001).

Data on somatic embryo induction were analyzed using the nonparametric Kruskal-Wallis analysis of variance. Pairwise multiple comparisons of ranked data were performed to compare cultivars, at the 5% significance level. Data obtained for proliferation and histodifferentiation capacity were submitted to ANOVA (Zar, 1999).

### Results and Discussion

Somatic embryos were induced in all genotypes (Table 1). The embryogenic potential expressed as the number of somatic embryos per cotyledon pair

<table>
<thead>
<tr>
<th>Cultivar</th>
<th>Percentage of embryogenic cotyledon pair</th>
<th>Somatic embryos per cotyledon pair</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bragg</td>
<td>41.7±19.4</td>
<td>2.2±1.5b</td>
</tr>
<tr>
<td>BRSMG 68 Vencedora</td>
<td>66.2±39.5</td>
<td>6.6±5.8a</td>
</tr>
<tr>
<td>BRS Torena</td>
<td>52.2±22.3</td>
<td>1.3±1.2b</td>
</tr>
<tr>
<td>BRS 137</td>
<td>38.7±17.6</td>
<td>0.9±1.0b</td>
</tr>
<tr>
<td>BRS 154</td>
<td>52.2±22.3</td>
<td>2.5±1.5b</td>
</tr>
<tr>
<td>Embrapa 48</td>
<td>40.0±21.0</td>
<td>1.0±1.0b</td>
</tr>
<tr>
<td>IAS5</td>
<td>70.0±22.8</td>
<td>10.6±6.6a</td>
</tr>
<tr>
<td>MG/BR 46 Conquista</td>
<td>64.0±18.2</td>
<td>8.3±2.4a</td>
</tr>
</tbody>
</table>

Mean±SD followed by equal letters in the columns do not differ by multiple comparison tests, at 5% probability.

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*Table 1. Capacity to produce somatic embryos from immature cotyledon pairs of eight Brazilian soybean cultivars*
varied significantly among cultivars, and IAS-5, BRSMG 68 Vencedora, and MG/BR 46 Conquista produced the highest mean numbers (p<0.001). Previous studies also reported the high capacity of IAS-5 to produce somatic embryos (Santos et al., 1997; Di Mauro et al., 2000; Droste et al., 2001). This is the first report on the embryogenic potential of cultivars BRSMG 68 Vencedora, BRS Torena, BRS 137, BRS 154, Embrapa 48 and MG/BR 46 Conquista. In soybean, differences among cultivars in capacity to produce somatic embryos from immature cotyledons have been shown in several reports (Komatsuda & Ohyama, 1988; Bailey et al., 1993; Santos et al., 1997; Di Mauro et al., 2000; Simmonds & Donaldson, 2000; Droste et al., 2001; Meurer et al., 2001; Tomlin et al., 2002; Hofmann et al., 2004; Hiraga et al., 2007; Yang et al., 2009).

A clear influence of the genotype on the capacity for repeated embryogenesis was also detected. Proliferative cultures could not be established for cultivars BRS Torena and BRS 137. However, continuous proliferation was obtained for Bragg, BRSMG 68 Vencedora, BRS 154, Embrapa 48, IAS-5 and MG/BR 46 Conquista, after five months in the proliferation medium. The growth rate of the embryogenic tissues in a 28-day period is presented in Table 2. All analyzed cultivars showed embryogenic proliferation capacity. Although no significant differences were observed among cultivars, MG/BR 46 Conquista had a relatively lower growth rate.

Proliferative embryogenic cultures were established for the Bragg, BRS 154 and Embrapa 48 cultivars, despite their low capacity for somatic embryo induction (Table 1). The absence of correlation between somatic embryo induction and repetitive proliferation of embryogenic cultures has been previously reported (Bailey et al., 1993; Simmonds & Donaldson, 2000; Droste et al., 2001; Yang et al., 2009).

The highest number of histodifferentiated embryos was obtained for the cultivar BRSMG 68 Vencedora, while MG/BR 46 Conquista had the lowest number (Table 3). All cultivars showed precocious germination, and MG/BR 46 Conquista showed the highest germination percentage (34%). Embryos that germinate before sufficient maturation often display poor development and reduced plant regeneration (Merkle et al., 1995). Therefore, certain cultivars, such as MG/BR 46 Conquista, must be discarded as candidates for successful embryogenesis.

In previous studies on soybean, ABA was considered to promote embryo growth, development, maturation, and improved embryo germination, when used in the globular stage (Tian & Brown, 2000), while ABA supplied during advanced maturation did not increment conversion frequencies (Schmidt et al., 2005). However, Weber et al. (2007) reported an increment on plant conversion when ABA was employed in both proliferation and maturation stages.

When added to the medium, activated charcoal presumably adsorbs auxins released from developing tissues and may promote a more normal morphology and increased germination ability (Merkle et al., 1995). Combination of ABA and activated charcoal has been observed to be beneficial for the development of Picea abies somatic embryos (Pullman et al., 2005).

For both cultivars tested, the addition of ABA and activated charcoal to the medium increased the number of histodifferentiated embryos (Table 4). The percentage of converted plants obtained for cultivar BRSMG 68 Vencedora (45%) was slightly higher than that of cultivar IAS-5 (42%), currently used for soybean transformation in Brazil (Droste et al., 2001, 2002; Homrich et al., 2008). In a previous experiment,

### Table 2. Weight gain (mean±SD) of embryogenic tissue after 28-day subculture period.

<table>
<thead>
<tr>
<th>Cultivar</th>
<th>Initial weight (mg)</th>
<th>Final weight (mg)</th>
<th>Growth rate&lt;sup&gt;(1)&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bragg</td>
<td>35.1±6.3</td>
<td>125.3±33.6</td>
<td>3.5±0.4</td>
</tr>
<tr>
<td>BRSMG 68 Vencedora</td>
<td>29.1±2.9</td>
<td>117.3±40.8</td>
<td>4.0±1.3</td>
</tr>
<tr>
<td>BRS 154</td>
<td>40.1±12.7</td>
<td>179.9±36.2</td>
<td>4.9±1.6</td>
</tr>
<tr>
<td>Embrapa 48</td>
<td>34.4±7.2</td>
<td>119.9±47.3</td>
<td>3.5±1.1</td>
</tr>
<tr>
<td>IAS5</td>
<td>24.9±4.5</td>
<td>93.5±22.8</td>
<td>3.9±1.2</td>
</tr>
<tr>
<td>MG/BR46 Conquista</td>
<td>30.8±4.7</td>
<td>76.7±29.9</td>
<td>2.6±1.4</td>
</tr>
</tbody>
</table>

<sup>(1)</sup>Quotient between final and initial weight.

### Table 3. Histodifferentiation and precocious germination after two months in maturation medium.

<table>
<thead>
<tr>
<th>Cultivar</th>
<th>Histodifferentiated embryos</th>
<th>Precociously germinated embryos (%)&lt;sup&gt;(1)&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bragg</td>
<td>709</td>
<td>6.1</td>
</tr>
<tr>
<td>BRSMG 68 Vencedora</td>
<td>819</td>
<td>16.2</td>
</tr>
<tr>
<td>BRS 154</td>
<td>473</td>
<td>19.7</td>
</tr>
<tr>
<td>Embrapa 48</td>
<td>565</td>
<td>11.6</td>
</tr>
<tr>
<td>IAS5</td>
<td>531</td>
<td>5.7</td>
</tr>
<tr>
<td>MG/BR46 Conquista</td>
<td>119</td>
<td>34.5</td>
</tr>
</tbody>
</table>

<sup>(1)</sup>Percentage refers to the number of germinated embryos over the total of histodifferentiated embryos.
in which ABA was added to the proliferation and maturation medium, Weber et al. (2007) obtained the same conversion frequency (42%) for cultivar IAS-5.

Considering the number of histodifferentiated embryos and the conversion percentage obtained in the present work, it is possible to estimate the expected number of plants for each treatment (Table 4). The increment of histodifferentiated embryos obtained from the ABA/charcoal treatment represents a double probability for plant regeneration in transformation experiments.

This work identified BRSMG 68 Vencedora and confirmed IAS-5 as genotypes with high potential for somatic embryogenesis and plant regeneration. Selection of such cultivars should provide a reasonable probability of success at any laboratory initiating soybean embryogenic protocols.

Conclusions

1. Cultivar BRSMG 68 Vencedora and IAS-5 are suitable genotypes for successful embryogenesis and transformation.

2. Addition of ABA to proliferation medium and of ABA plus activated charcoal to maturation medium increases embryo histodifferentiation, resulting in a higher percentage of regenerated plants.

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