Behavior of lateral buds of *Hancornia speciosa* after cryopreservation by encapsulation-vitrification

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**ABSTRACT.** *Hancornia speciosa* is a fruitful species from Cerrado biome with high economic potential. However, the intense and disordered extractivism have caused a reduction of its population in its endemic area. In addition, seed recalcitrance negatively affects the conventional conservation of the species. Aiming to find alternatives that enable the long-term conservation of this species, the study's objective was to assess the behavior of lateral bud's regrowth after cryopreservation procedures by encapsulation-vitrification technique. Sodium alginate capsules containing lateral buds were pre-cultured in liquid WPM supplemented with 1.0 M glycerol, and subsequently exposed to different concentrations of sucrose (0.3; 0.75 and 1.0 M) for 24 or 48 hours. The capsules were subjected to dehydration in silica gel or airflow hood for 0, 1, 2 and 3 hours before different incubation times in PVS2 (0, 15, 30, 60 and 120 minutes) at 0°C. A high regeneration percentage of lateral buds was observed after cryopreservation of capsules treated with 0.75 M sucrose plus 1.0 M glycerol (24 hours), associated with dehydration in an airflow hood (1 hour) and immersion in PVS2 (15 minutes). Encapsulation-vitrification allowed the long-term conservation, and provided high plant material survival rates after cryopreservation of *Hancornia speciosa* sensitive explants.

**Keywords:** Mangabeira, recalcitrant species, long-term conservation, dehydration.

**Introduction**

Cerrado is a complex Brazilian biome with great diversity of habitats. This rich biodiversity occupies an area over 2 million km², and represents about 23% of the Brazilian national territory (Klink & Machado, 2005). Several plant species of economic importance are found in this biome and many of them, if exploited consciously, have potential for the sustainable production of food and medicine (Machado, Ramos, Caldas, & Vivaldi, 2004). The species *Hancornia speciosa* Gomes, popularly known as ‘Mangabeira’, is a native fruitful tree of the Brazilian Cerrado, belonging to Apocynaceae, and is widely distributed throughout the country, from Amapá to the state of São Paulo (Silva Junior, Xavier, Silva Lédo, Santos Musser, & Silva Lédo, 2007). Among the attractive characteristics of the species are its fruits that, due to its pleasant flavor, stimulate fresh consumption and can also be used in the production of syrups, juices, candy, ice cream...
and jams (Soares et al., 2011). Since it is an iron-rich fruit and a vitamin C source, the mangaba fruit occupies a prominent position among functional foods (Silva Júnior et al., 2007). For all these characteristics, this species has been the target of predatory extraction, which has led to a decline in *H. speciosa* population in its native areas (Soares et al., 2011). In addition, the recalcitrance of its seeds is another detrimental factor for the availability of the species, since it prevents conventional storage and its later use (Barros et al., 2006).

Aiming to overcome the obstacles imposed by the natural characteristics of each species, germplasm conservation strategies have been used in order to prevent the extinction of these species and allow multiple genotypes to remain available for order to prevent the extinction of these species and germplasm conservation strategies have been used in the natural characteristics of each species, later use (Barros et al., 2006).

Given the above, the aim of this study was to establish an efficient long-term conservation method through cryopreservation protocol using the encapsulation-vitrification technique for lateral buds of *Hancornia speciosa*.

**Material and methods**

**Plant material**

*Hancornia speciosa* shoots with 60 days of *in vitro* culture were used as a source of explants. These explants were grown in culture medium composed by *Wood Plant Medium* (WPM) basal salts (Lloyd & Mccown, 1980) supplemented with 8.87 μM 6-benzylaminopurine (BA), 30 g L⁻¹ sucrose and gelled with 7 g L⁻¹ agar, as established by Soares et al. (2011).

**Encapsulation and dehydration of lateral buds**

Lateral buds, with approximately 1.0 mm², were excised from shoots and immersed in sodium alginate matrix (2.5% w/v). Subsequently, with the aid of an automatic pipette, the individual encapsulated units were retrieved and dripped into calcium chloride solution (CaCl₂.2H₂O) (100 mM), remaining for 20 minutes for complexation procedure. The encapsulated units, individually formed by a lateral bud wrapped in sodium alginate matrix, were subjected to three washes with autoclaved distilled water to remove the excess of CaCl₂. Sequentially, the encapsulated units were immersed in potassium nitrate solution (KNO₃) (100 mM) for 15 minutes for decomplexation, and then washed again with autoclaved distilled water. Capsules containing the lateral buds were immersed in pre-culture liquid WPM medium added with sucrose at different concentrations (0.3; 0.75 and 1.0 M) and supplemented with 1.0 M glycerol for different incubation times (24 or 48 hours) under constant stirring of 150 rpm. Subsequently, the capsules were inoculated in regeneration medium (WPM added with 0.2 μM BA) and maintained in a growth chamber with a 16 hours photoperiod, irradiance of 36 μmol m⁻² s⁻¹ and temperature of 25°C ± 2°C. Each treatment consisted of ten Petri dishes containing five encapsulated units per plate, randomly. After 30 days in culture, the regeneration percentage of encapsulated buds was evaluated. After the determination of sucrose optimal concentration...
in the pre-culture medium and the best incubation time, the capsules were subjected to two dehydration procedures: the first in an airflow hood, and the second on silica gel, both for 0, 1, 2 and 3 hours. After dehydration, the capsules were inoculated in regeneration medium (WPM supplemented with 0.2 μM BA) and maintained in a growth room chamber with a 16 hours photoperiod, irradiance of 36 μmol m⁻² s⁻¹ and temperature of 25°C ± 2°C. Each treatment consisted of ten Petri dishes containing five encapsulated units per plate, randomly. After 30 days in culture, the regeneration percentage of encapsulated buds was evaluated.

**Encapsulation-vitrification**

Encapsulated lateral buds were pre-cultured in liquid WPM supplemented with 0.75 M sucrose and 1.0 M glycerol, under constant stirring of 150 rpm for 24 hours, with subsequent dehydration in an airflow hood for 1 hour. The capsules were treated with Loading Solution (LS) (2.0 M glycerol + 0.4 M sucrose) for 20 minutes before immersion in Plant Vitrification Solution number 2 (PVS2), composed by 30% glycerol, 15% ethylene glycol, 15% Dimethyl sulfoxide (DMSO) and 0.4 M sucrose (Sakai, Kobayashi, & Oiyama, 1990) at 0°C for different incubation times (0, 15, 30, 60 and 120 minutes). After the PVS2 treatment, the capsules were inserted into 2.0 mL cryovials and plunged into liquid nitrogen (LN), remaining for 90 minutes. Rewarming was performed in a water bath at 40°C for three minutes. Following, the capsules were then treated in a Petri dish containing Recovery Solution (RS), also known as unloading solution (1.2 M sucrose) at 25°C for 15 minutes. After cryopreservation, the capsules were inoculated in post-culture medium, consisting of WPM basal salts supplemented with 0.3 M sucrose for 24 hours in the absence of light. Subsequently, the capsules were inoculated in bud regeneration medium (WPM supplemented with 0.2 μM BA). The explants were maintained in a growth room chamber with a 16 hours photoperiod, irradiance of 36 μmol m⁻² s⁻¹, at a temperature of 25°C ± 2°C. Each treatment consisted of ten Petri dishes containing five encapsulated units per plate, entirely at random. The variables evaluated after 30 days in culture were the percentage of survival and the percentage of shoot formation.

**Statistical analysis**

The collected data were statistically analyzed using SAS (Statistical Analysis System [SAS], 2001), and the means were compared by Fisher's exact probability test (p ≤ 0.05).

**Results and discussion**

**Encapsulation-dehydration**

After pre-culture of sodium alginate capsules containing the explants in liquid WPM media with different sucrose concentrations, the highest percentage of regeneration was obtained in treatment with 0.75 M sucrose after 24 hours of pre-culture (Figure 1).

The different dehydration conditions of alginate capsules, both in silica gel and airflow hood, showed significant differences among treatments. The highest survival percentage (78%) was observed in the explants exposed to dehydration in an airflow hood for 1h, where the capsules had a water content of 34% (Table 1).

**Table 1.** Percentage of water content and recovery of encapsulated units of *Hancornia speciosa* subjected to drying, either in silica gel or airflow hood, for different times.

<table>
<thead>
<tr>
<th>Time (hour)</th>
<th>Water content (%)</th>
<th>Recovery (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Airflow</td>
<td>Silica</td>
</tr>
<tr>
<td>Control</td>
<td>99.7</td>
<td>99.4</td>
</tr>
<tr>
<td>1</td>
<td>34.0</td>
<td>33.9</td>
</tr>
<tr>
<td>2</td>
<td>23.5</td>
<td>9.5</td>
</tr>
<tr>
<td>3</td>
<td>11.8</td>
<td>8.4</td>
</tr>
</tbody>
</table>

Means followed by the same lowercase letter in the column and uppercase letter in the row do not differ between each other by Fisher’s exact test (p ≤ 0.05).

Among pretreatment techniques described for the plant tissues cryopreservation, encapsulation-dehydration has numerous advantages, such as easy handling of the explants and a large storage capacity for fragile tissues (Sakai & Engelmann, 2007). The importance of dehydration protocols is related to reduce the water content of the plant material (Silva, Paiva, Swennen, Andrè, & Panis, 2014), in order to enable tissue cooling, and prevent injuries caused by the expansion of frozen water and the formation of...
extra- and intracellular ice crystals (Sakai & Engelmann, 2007). Therefore, dehydration acts alleviating these critical steps and is considered one of the key points for the success of cryopreservation. Thus, the optimum water content after dehydration ranges from 20 to 30% of the total water content in the tissues (Vertucci & Farrant, 1995).

The highest recovery rate of encapsulated lateral buds after dehydration occurred at a water content around 34% (Table 1), which is considered viable for the capsules hydration levels to be cryopreserved. However, *H. speciosa* lateral buds seems to be more sensitive to dehydration. Successful cryopreservation was observed in shoot apices of *Rabdosia rubescens* encapsulated and precultured in MS culture medium with 0.4 M sucrose, in combination with 2.0 M glycerol and dehydrated for 1 hour in silica gel, reaching a water content of 21% and a recovery rate of 85% (Ai, Lu, & Song, 2012). Encapsulation-vitrification of apical meristems of date palm (*Phoenix dactylifera* L) in PVS2 for 15 minutes and immersion in LN, ensured increased survival and recovery of explants (40%) (Fki et al., 2013). Shoot apices of *Vitis vinifera* encapsulated were pre-cultured in half-strength MS medium supplemented with increasing concentrations of sucrose (0.25-1.0 M) and then dehydrated in an airflow hood for different periods, in order to determine the optimum capsules dehydration time (Wang, Tanne, Arav, & Gafny, 2000). These authors found that only the shoot apices with water content lower than 22.5% survived. Wang, Laamanen, Uosukainen, & Valkonen (2005) observed that sodium alginate capsules containing apical buds of *Rubus idaeus* L., pre-cultured in MS medium containing sucrose, had an initial moisture content of 66%, and the water content decreased to 22.2% in the first 4 hours in an airflow hood, and to 14.3% after 8 hours of dehydration. According to these authors, the highest recovery rates of the apical buds were obtained by dehydration for at least 3 hours in an airflow hood.

Similar to the observed in the present study, the encapsulation-dehydration technology has been successfully applied in the cryopreservation of different species, such as *Beta vulgaris* (Vandenbussche & Proft, 1998), *Apple* (*Malus x domestica* Borkh.) (Paul, Daigny, & Sangwan-Norreel, 2000) and *Citrus* (Wang, Batuman, Li, Bar-Joseph, & Gafny, 2002). One of the advantages of this method is that the recovery of cryopreserved buds occurs directly, quickly and usually without callus formation.

**Encapsulation-vitrification**

The cryopreservation of lateral buds was successfully obtained with the encapsulation-vitrification technique, using PVS2 incubation for 15 minutes before plunge into LN. On average, it was possible to observe 67% of explants survival (Figure 2A) and 89% of shoot formation (Figure 2B). In addition, after 15 days from the beginning of the dehydration and cryopreservation treatment, a disruption of the capsule occurred and shoots start to grow (Figure 3A-B). At the end of 60 days of cultivation, normal plants, which means possessing shoot and roots as well, were formed from encapsulated-vitrified lateral buds (Figure 3 C-E).

In the last few years, the encapsulation-vitrification technique has stood out in the cryopreservation of several species of tropical and temperate plants (Sakai & Engelmann, 2007). However, like other cryopreservation techniques, some limiting factors reduce the efficiency of this process. Osmotic stress and chemical toxicity promoted by cryoprotectants are two major problems associated with the low efficiency of this technique. One of the ways to obtaining better results is optimizing the cryoprotectant concentration before the explants immersion in LN, aiming to increase survival after the application of cryopreservation techniques (Papadopoulos et al., 2002; Castro et al., 2011).
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Among the results obtained by this study, the incubation in PVS2 for 15 minutes was good enough to lead to cryoprotection in the encapsulated explants, not interfering in the capsules structure. After 30 minutes of incubation in PVS2, the capsules had a gelatinous aspect and disrupted spontaneously, which may be the probable cause of the recovery failure (Figure 2).

Recent cryopreservation studies with *Hancornia speciosa* lateral buds by encapsulation-dehydration technique indicated unsuccessfully regrowth rates (Sartor et al., 2012). In this case, the absence of satisfactory results may be due to the lack of adequate procedures for vitrification, combined with the wrongly rewarming of encapsulated explants at room temperature. Both mistakes may have led to intracellular ice formation by means of crystallization during cooling, and recrystallization during reawarming (Fahy et al., 1984). Vitrification procedures must be used, instead only dehydration, for sensitive tissues as *Hancornia speciosa* lateral buds. In this article, it was proved that encapsulation-vitrification is an adequate procedure for cryopreservation of this kind of tissue. Moreover, several studies have shown that the encapsulation-vitrification technique is quite efficient for different explants and species (Wang et al., 2000; Sakai & Engelmann, 2007; Ai et al., 2012; Fki et al., 2012).

**Conclusion**

This protocol describes an efficient method for cryopreservation of *Hancornia speciosa* germplasm by encapsulation-vitrification technique. The optimized parameters here reported enabled to obtain cryopreserved lateral buds and plants regeneration up to 67%. These results also emphasized the importance to investigating the acclimatization phase and material genetic stability after cryopreservation and *ex vitro* development. Besides, this study is a promising approach for cryopreservation using sensitive explants produced...
under in vitro conditions, and may facilitate the conservation of many other recalcitrant species.

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