Water uptake and pre-germination treatments in macaw palm 
(Acrocomia aculeata - Arecaceae) seeds

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ABSTRACT – The aims of this study were to evaluate the influence of embryo adjacent tissues in water uptake and treatments to release dormancy in macaw palm seeds. To assess water absorption with or without the opercular tegument (OT), seeds were immersed in water for 0, 12, 24, 48, 72, 96, 120, 168, 240, 360, 480 and 720 hours and the water content of isolated embryos and seeds was determined. In seeds with or without OT, gibberellic acid (GA₃) solutions at concentrations of 0, 50, 100, 500, 1000 and 2000 mg.L⁻¹ were applied on the OT region using a culture medium. In another experiment, the OT was removed or left intact and the seeds immersed in hydrogen cyanamide solutions (0, 0.5, 1, 1.5, 2 and 2.5%). The percentage germination, germination speed index (GSI) and percentage dead seeds were determined. The OT does not prevent embryo water uptake but reduces water absorption speed. The use of GA₃ in a culture medium applied to the OT region did not effectively release dormancy. The use of cyanamide at these concentrations increased seed mortality, resulting in a low germination. OT removal increased the GSI.

Index terms: imbibition, opercular tegument, GA₃, hydrogen cyanamide.

Introduction

The macaw palm, Acrocomia aculeata (Jacq.) Lodd. ex. Mart. (Arecaceae), is a tropical palm widely distributed throughout Brazil (Loreni et al., 2004) and its occurrence in Minas Gerais state is associated with eutrophic soils (Motta et al., 2002). This species is a very productive oil-producing plant and has a high potential for producing biofuels, especially in dry tropical regions (Moura et al., 2010; Dias, 2011).

The germination of macaw palm seeds is low due to dormancy (Lorenzi et al., 2004; Ribeiro et al., 2011) and, therefore, breaking dormancy is important for producing seedlings on a commercial scale (Loreni et al., 2004).

Water absorption is the first stage in germination and can occur in different ways in seed tissues (McDonald Jr. et al., 1988). According to Bewley and Black (1994), the
embryo absorbs water continuously at a greater speed due to cell elongation and division, in contrast to other seed tissues where there is no tissue expansion. Seed imbition by water immersion can harm embryos due to the rapid absorption and lower diffusion of oxygen (Marcos-Filho, 2005).

Dormancy has been characterized as blocking the conclusion of germination (Bewley, 1997) and can be classified as physiological, morphological, morphophysiological, physical and a combination between physical and physiological dormancy (Baskin and Baskin, 2004; Finnch-Savage and Leubner-Metzger, 2006). Cases of morphological dormancy are common in palm seeds (Orozco-Segovia et al., 2003), but studies with isolated embryos of macaw palm demonstrate that dormancy due to embryo immaturity does not occur in this species (Ribeiro et al., 2012), and it can be classified as nondeep physiological dormancy (Baskin and Baskin, 2004; Finnch-Savage and Leubner-Metzger, 2006).

Various pre-germination treatments are used to overcome dormancy in palm seeds, such as heat treatments (Hussey, 1958; Rees, 1961; 1962; Addae-Kagyah et al., 1988; Meerow, 1991; Robinson, 2009), pre-immersion in water (Meerow, 1991), removal of the operculum (Hussey, 1958; Carpenter et al., 1993; Al-Wasel and Warrag, 1998; Pérez et al., 2008), and of the endocarp (Merrow, 1991; Pérez et al., 2008), and chemical treatments (Nagao et al., 1980; Pérez et al., 2008; Roberto and Habermann, 2010).

The positive effect of gibberellic acid (GA$_3$) and the removal of the opercular tegument were observed in the germination of macaw palm seeds (Ribeiro et al. 2011) and also in the Pritchardia remota palm (Pérez et al., 2008). The use of this plant growth regulator has also proved efficient in breaking seed dormancy in other palm species, such as Euterpe edulis (Roberto and Habermann, 2010), Archontophoenix alexandrae and Ptychosperma macarthurii (Nagao et al., 1980). However, the high cost of GA$_3$ and the difficulty in diffusion of solutions through the seed tissues (Nagao et al. 1980, Ribeiro et al., 2011) has encouraged studies on new ways of applying this growth regulator.

Hydrogen cyanamide (CH$_2$N$_2$) is an organic compound used to induce bud sprouting and release seed dormancy (Shulman et al., 1983; Herrera et al., 1998; Jiménez et al., 2008). The efficiency of this compound in release seed dormancy was demonstrated for the Elaeis guineensis palm, where it stimulated germination (Herrera et al., 1998; Jiménez et al., 2008).

The aim of this study was to evaluate the influence of seed structures on water absorption by the embryo, the effect of GA$_3$ dosages applied by culture medium and of hydrogen cyanamide dosages, in breaking dormancy in macaw palm seeds.

Material and Methods

**Collection and preliminary procedures:** fruits were collected after abscission from 20 individuals of *A. aculeata* from natural populations in Montes Claros county (16°42'34"S; 43°52'48"W), in the north of Minas Gerais state. The fruits were stored in the shade until the experiments were carried out and, remained in this condition for a maximum of eight months.

Seeds were removed from the fruits before each experiment, with the aid of a benchtop vice, were selected and damaged seeds were discarded. Seed water content was determined using the oven method with five replications of ten seeds (Brasil, 2009). The seeds were disinfected in 6% chlorine solution for 10 minutes, followed by three washes in running water. The opercular tegument was removed with a razor blade, using a stereomicroscope to avoid embryo damage (Ribeiro et al., 2011).

**Water absorption:** the fruits for this experiment were collected in 2008, with the seed being extracted from the fruits and the opercular tegument removed from half of them. Seeds, with and without the opercular tegument, were immersed separately in water in 2 L beakers, changing the water daily. Imbibition occurred at ambient temperature. Before immersion and after 12, 24, 48, 72, 96, 120, 168, 240, 360, 480 and 720 hours of imbition, five replications of 10 seeds from each treatment were removed and the embryo was isolated using scalpels in order to determine the water content of the embryo and the rest of the seed (endosperm + tegument), using the oven method (Brasil, 2009). The experimental design was completely random.

**Application of GA$_3$ by culture medium:** the seeds used in this experiment were collected in 2009, extracted from the fruit and kept in polystyrene trays containing vermiculite. Seeds were submitted to pre-imbition in vapor in a humid growth chamber with 95 ± 5 % of RH at 30 °C for one week. After the pre-imbition, the seeds were disinfected with 2% chlorine solution for 10 minutes.

The opercular tegument was removed from half the seeds and 1 mL of culture medium was applied on the micropylar region with a 10 mL hypodermic syringe. The MS medium was used (Murashige and Skoog, 1962), supplemented with 10 g.L$^{-1}$ of agar and autoclaved at 121 °C for 20 minutes, with different dosages of GA$_3$ added (0, 50, 100, 500, 1000 and 2000 mg.L$^{-1}$), filtered in a Millipore® membrane with 0.45 µm mesh and added after the medium had reached 50 °C.

After application of the culture medium, the seeds were returned to the humid growth chamber. The vermiculite was moistened daily with a squeeze bottle. The experimental
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The design was randomized blocks in a 6 x 2 factorial (GA3 concentrations x seeds with or without the opercular tegument), with five replications of 20 seeds per treatment. The seeds were evaluated weekly for four months and the germinated seeds (considering germination as the protrusion of the cotyledonary petiole) and dead seeds were removed after counting. The percentage germination, germination speed index (GSI) (Maguire, 1962) and percentage dead seeds was calculated.

Immersion in hydrogen cyanamide solution: another seed lot collected in 2009 was used for this experiment. Seeds were extracted from the fruits, disinfected and submitted to pre-imbibition as described in the previous experiment. The opercular tegument was removed from half the seeds and these were immersed in hydrogen cyanamide solutions (Dormex®), for 24 hours with different concentrations of the commercial formulation (0, 0.5, 1, 1.5, 2 and 2.5%) and returned to the humid growth chamber. After six weeks, the seeds were removed from the vermiculite and the hydrogen cyanamide treatment was repeated.

The experimental design were randomized blocks in a 6 x 2 factorial (cyanamide dosages x seeds with or without the opercular tegument), with five replications of 20 seeds per treatment. The seeds were evaluated weekly during 13 weeks using the same criteria described for the previous experiment. Those seed embryos which remained hard until the end of the experiment were submitted to the tetrazolium test, using four replications of 10 embryos, which were immersed in a 0.5% solution of 2,3,5-triphenyl tetrazolium chloride at 35 °C for four hours, according to the methodology described by Ribeiro et al. (2010). The tetrazolium test was used in this experiment to evaluate the toxicity of the hydrogen cyanamide on the embryos.

Statistical analysis: an analysis of variance was done with the data on percentage germination and dead seeds transformed to arc sine √x/100, and the means were compared using Tukey’s test at 5% probability (SAS Institute, 1990). Water absorption and embryo viability data were submitted to a regression analysis and the model adjustment was tested at the 5% significance level and evaluated with the determination coefficient (R²) using the SigmaPlot v.11.0 software.

Results and Discussion

Water absorption: before imbibition, the isolated embryos and the rest of the seeds (endosperm + tegument) had water contents of 7.3 and 6.5% respectively. During imbibition, the highest absorption rate occurred in seeds without the opercular tegument up to approximately 240 hours of imbibition (Figure 1). After this period, the embryos from seeds with and without the opercular tegment contained water contents close to, and higher than the rest of the seeds, reaching percentages higher than 49% after 720 hours of imbibition. The endosperms + tegments, in both conditions, showed similar imbibition behavior, with a slow and gradual increase in water content until a plateau was reached after 360 hours imbibition, with 20.9 and 22% of water content in seeds with and without opercular tegment, respectively, after 720 hours imbibition.

Application of GA3 by culture medium: the seeds had water content of 22.4% before the experiment. The removal of the opercular tegument had a positive effect on germination and the GSI (Table 1), but without effect of the GA3. The removal of the opercular tegument and the GA3 affected the mortality percentage of seeds, with an interaction between these variables (Figure 2). In seeds which had the intact opercular tegument this percentage was unaffected by the use of GA3, whereas in seeds without the opercular tegument there was higher mortality without or with higher dosages of GA3.

Table 1. Germination percentages (G%) and germination speed index (GSI) in seeds of A. aculeata, with or without the opercular tegument (OT).

<table>
<thead>
<tr>
<th>Seed condition</th>
<th>G%</th>
<th>GSI</th>
</tr>
</thead>
<tbody>
<tr>
<td>with OT</td>
<td>8.3 b</td>
<td>0.03 b</td>
</tr>
<tr>
<td>without OT</td>
<td>13.3 a</td>
<td>0.06 a</td>
</tr>
</tbody>
</table>

The same letters indicate an absence of significant differences in the columns according to the Tukey test at 5% probability. The means were obtained from the dosages of GA3 with or without removal of the OT.

Figure 1. Water contents of seeds (endosperm + tegment) and embryos of A. aculeata, with or without the opercular tegument (OT), after different periods of water immersion.
noted that hydration of the embryonic axes is greater than for other structures in soybean seeds. Embryo imbibition is quick due to their chemical composition and is influenced by tissue expansion (McDonald Jr. et al., 1988; Bewley and Black, 1994).

**Immersion in hydrogen cyanamide solution:** the seeds used in this experiment had water content of 18.9%. Removal of the opercular tegument and immersion in the cyanamide affected the germination rate and there was also an interaction between these variables. Germination did not increase with immersion in different dosages and there was no difference between the cyanamide dosages for germination but only an effect with the removal of the opercular tegument in the control (Figure 3). Repeating seed immersion in the cyanamide dosages did not stimulate seed germination.

The GSI was statistically greater in seeds with the opercular tegument removed, presenting a value of 0.05 for these seeds and 0.02 for seeds with the opercular tegument intact, but no effect was observed for immersion in hydrogen cyanamide. The effect on mortality was only seen with cyanamide immersion, there being no effect with the removal of the opercular tegument (Fig. 3). The percentage of dead seeds increased with increasing the dosages of cyanamide. The control showed a lower mortality, which differed significantly for dosages greater than 0.5%.

The tetrazolium test showed a tendency for the percentage of viable embryos to decrease with increasing concentrations of cyanamide (Figure 4), demonstrating the harmful effect of this product on the embryo with increases in concentration.

The higher water uptake in the embryos compared to the other tissues were observed by McDonald Jr. et al. (1988), who
The tegument is not a barrier to water uptake, which characterizes the absence of physical dormancy. The tegument of A. aculeata presents cells with thin and non-lignified walls, which do not give any mechanical resistance to the seed (Moura et al., 2010). The opercular tegument was also not a barrier to water absorption, considering that both the seeds and embryos, which did not have the opercular tegument removed, had the same final water content as those with an intact opercular tegument. However, the favorable effect on seed germination of the macaw palm was observed by Ribeiro et al., (2011), caused by removing the opercular tegument, may be related to the higher speed of water absorption by the seed embryo, where the opercular tegument was removed, as observed in the current study, as well as to a reduction in the restrictive mechanical effect (Haigh and Barlow, 1987; Welbaum and Bradford, 1990).

The efficiency of the GA3 in stimulating germination of macaw palm seeds has been described by Ribeiro et al. (2011), who observed an increase in germination with immersion in high concentrations of this growth regulator. This effect was also seen in seeds of Archontophoenix alexandrae and Ptychosperma macarthurii (Nagao et al., 1980) and Euterpe edulis (Roberto and Habermann, 2010), demonstrating that this growth regulator accelerates germination in these palms but its effectiveness is limited by its ability to penetrate in the seed tissues (Nagao et al., 1980).

Embryo viability may be compromised by prolonged immersion of the diaspores in water or solutions, with a risk of damage to cell membranes and solute leakage, together with possible restrictions in the oxygen supply (Ferreira and Borghetti, 2004; Marcos-Filho, 2005). These damages may be avoided by applying the GA3 to seeds using a culture medium (Pinheiro et al., 2001).

The low germination percentage observed in this experiment may be due to the restricted penetration of the GA3 into the seeds, since there was no direct contact of the culture medium supplemented with the GA3 with the embryos, due to the presence of micropylar endosperm, which was not removed.

The opercular tegument may limit the water and gas flows in some species and offer mechanical resistance to embryo elongation (Hussey, 1958; Al-Wasel and Warrag, 1998), which may not be observed in A. aculeata with respect to limitation of water absorption. The high percentage of dead seeds occurring with the removal of the opercular tegument may be due to a greater exposure of the embryo and to use of the MS medium, which may have served as a substrate for microorganisms. Macaw palm seeds with a high water content are more susceptible to microorganism contamination, increasing mortality rate (Rubio Neto et al., 2012), and, therefore, seed water content may have affected the percentage of dead seeds.

The use of hydrogen cyanamide to release dormancy in palm seeds has already been described by Herrera et al. (1998) and Jiménez et al. (2008), who obtained high germination percentages in Elaeis guineensis. In peach-palm (Bactris gasipaes), treatment with hydrogen cyanamide harmed germination even for short exposures times (Villalobos et al., 1992).

Hydrogen cyanamide stimulates plant tissue metabolism, increasing respiration (Goldbach et al., 1988; Or et al., 2000), and its activity in breaking seed dormancy may be due to increased lipid degradation (Amberger, 1984 apud Herrera et al., 1998). The higher mortality observed with increasing dosages of hydrogen cyanamide may be due to the long product exposure and high concentrations harming the seed (Villalobos et al., 1992; Herrera et al., 1998). The retreatment of seeds with cyanamide may have intensified these effects. The rapid metabolic activation and accumulation of peroxide caused by the product (Goldbach et al., 1988; Amberger, 1984 apud Herrera et al., 1998) may have intensified the effects of oxidation stress on the embryo, causing seed death.

Removal of the opercular tegument has also been described in other palms as a germination stimulant (Hussey, 1958; Carpenter et al., 1993; Al-Wasel and Warrag, 1998; Ribeiro et al., 2011). The removal of this layer resulted in an increase in germination speed in Rhipidophyllum hystrix (Carpenter et al., 1993) and a higher germination rate in seeds of Phoenix dactylifera (Al-Wasel and Warrag, 1998) and Elaeis guineensis (Hussey, 1958). The inefficiency of the removal of the opercular tegument, together with immersion in hydrogen cyanamide, demonstrated the harmful effect of this product in A. aculeata embryos.

Conclusions

The tegument and opercular tegument were not a barrier to water uptake by A. aculeata seeds, but the removal of the opercular tegument increases the speed of embryo water absorption.

The application of GA3 by culture medium on the micropylar region did not increase the germination of A. aculeata seeds.

Hydrogen cyanamide did not break the dormancy of A. aculeata seeds at the dosages tested and harmed embryo viability.

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References


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