Effects of desiccation on Euterpe edulis Martius seeds

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ABSTRACT: Information on desiccation sensitivity of Euterpe edulis seeds under two drying rates is presented. The sensitivity was studied during the course of germination and normal germination. The water content was evaluated for both seeds and embryos. Results showed the following: (a) For both drying treatments and for both germination and normal germination, desiccation sensitivity values were higher for measurements based on the water content of the embryo than for those of the seed. (b) For both drying treatments, desiccation sensitivity were higher for normal germination than for germination based on both the embryo and seed water contents. (c) Under the slow drying treatment and for measurements based on the seed water content, critical water content was visible for normal germination but not for germination; (d) Critical water contents for germination and normal germination were more clearly established in the fast drying treatment than they were in the slow drying method based on both the embryo and seed water contents. Critical water contents were not associated with changes in electrolyte leakage, which suggests that conductivity is not a good indicator of physiological seed quality. From the beginning of both drying treatments, changes in nuclei and vacuoles were observed, but, when seed water content was reduced to below critical values, the cells became severely plasmolyzed, the vacuoles highly distorted, and the nuclei formed an almost homogeneous mass with the chromatin and the nucleoplasm, which suggests irreversible DNA damages.

Introduction

Euterpe edulis exhibits recalcitrant seed-storage behaviour; in other words, seed viability begins to decrease at approximately 39% seed water content and is definitively lost at 21% water content (Andrade, 2001; Martins et al., 2000; Reis et al., 1999). In addition, Panza et al. (2004) report subcellular recalcitrant features of the E. edulis embryo tissues, i.e., cells highly vacuolated (at high water content), almost lacking storage reserves, with abundant endomembranes, and abundant ergastic substances. The differential water content between the embryo axis (and embryo) and the storage tissue presents a difficulty for interpreting the experimental results on recalcitrant seeds (Berjak et al., 1989). This difficulty is exacerbated when the embryonic axis constitutes a small fraction of the total volume of the seed, and its water content is considerably higher than that of the storage tissues (ISTA, 1989), which is the situation with E. edulis seeds. A preliminary study indicates that the whole embryo constitutes only 0.54% of the seed fresh weight with 85% water content, while that of the endosperm is 48.2% (Panza et al., 2001).
When submitted to dehydration, recalcitrant seeds show variable critical values according to the speed of dehydration (Farrant et al., 1985; Pritchard, 1991). However, there are conflicting opinions on this subject (Bonner, 1996; Finch-Savage, 1992; Pritchard et al., 1995).

Transmission electron microscopy shows that recalcitrant seeds exhibit a progressive degradation of membranes during drying (Pammenter and Berjak, 1999). In the recalcitrant seeds of *Acer saccharinum* L. and *Aquilaria agallocha* Roxb, damages in cell membranes increase solutes leakage and, consequently, conductivity (Beewar et al., 1982; Kundu and Kachari, 2000). Although leakage has been correlated with the loss of seed viability and germination, such correlation has been scarcely investigated in recalcitrant seeds.

In orthodox seeds, during the desiccation-tolerant phase, the chromatin is in a highly condensed state while the seeds are desiccation-tolerant. However, this is clearly a reversible process and rehydration to the state of resumption of DNA replication is associated with chromatin decondensation (Pammenter and Berjak, 1999). Chromatin compaction is also observed in recalcitrant seeds with increasing water loss, although here the compaction is irreversible (Pammenter and Berjak, 1999). This suggests that the desiccation-sensitive embryo tissues of recalcitrant seeds lack the orderly and reversible mechanism that provide stability to chromatin in the desiccation-tolerant embryo tissues of orthodox seeds.

The present study examined the germination course in function of the water content of the whole seed and the embryo at two drying rates. Subcellular analysis by transmission electron microscopy (TEM) and electrolyte leakage by conductivity tests were also used to evaluate the dehydration process. To obtain exact knowledge of the desiccation sensitivity of these seeds, we determined the embryo water content at which germination parameters are affected; this information was previously unavailable.

**Material and Methods**

Mature fruits of *E. edulis* were harvested in the Parque Nacional Iguazú, province of Misiones, Argentina (S25°40’50.4’’ W54°09’24.0’) during the month of August for three consecutive years, from 1999 to 2001, and shipped by expedited post to the Germplasm Bank of INTA, Castelar, Buenos Aires, Argentina, where the experiments were conducted.

Fruits were stored in 30 µm-thick polyethylene bags at 4°C. At this temperature, seeds did not show sensitivity to chilling. During the first days of storage, pericarps were removed. Seeds were carefully selected to assure that the samples were similar in size and germination percentage.

**Desiccation methods**

Two drying methods were used. In both, seeds were maintained in a single layer in plastic mesh bags. Rapid drying was carried out by placing the bags inside silica gel (approximately 2% RH, Vaisala humidity and temperature indicator HMI 31 - Finland) contained in desiccators at 25 ± 1°C. The silica gel/seed ratio was 8/1 (w/w) and silica gel was replenished daily. Seed samples were removed from the desiccators at 24, 48, 72, 96, and 144 h. Slow drying was carried out by placing the bags on a metal mesh in a drying chamber at 21 ± 2% RH and 19 ± 2°C. Samples were removed periodically during 17 days. Seeds from both drying treatments were used for germination testing, conductivity testing, and light and transmission electron microscopy. The water contents (percentage wet weight basis) of the whole seed and the isolated embryos were determined on three replications of five seeds by the ISTA (2005) oven methods (103°C ± 1°C for 17 h).
Germination tests

Germination tests were performed in plastic boxes on moist germination paper and cotton at 25 ± 1°C, 16 h fluorescent light (16 µmol m⁻² s⁻¹). Initially, germination was assessed three times per week. Each test used 3 x 20 seeds. Germination and normal germination were evaluated. Germination corresponds to button protrusion, i.e., emergence of cotyledon tissue forming a wheel around the hole from which the radicle protrudes. Normal germination corresponds to the plantule having its first three leaves (the third leaf being the first one with pinnate structure) and a well-developed radicle (Belin-Depoux and Hering de Queiroz, 1971). Evaluation of normal germination was accessed at weekly intervals during 25 weeks. The critical water contents of the seed and embryo were taken at the point in which the germination significantly decreased.

Transmission electron microscopy

Embryos isolated from seeds of both control and silica gel drying treatment were immediately fixed. In all cases, root and shoot apexes were excised and fixed overnight at 4°C in 2.5% glutaraldehyde in 0.1 M phosphate buffer (pH 7.2). Tissues were post-fixed in 1% OsO₄ in water for 2 h, and then they were dehydrated in a graded ethanol-acetone series and embedded in low viscosity Spurr’s resin. Once germination curves were known, shoot and root apical meristems of those seeds corresponding to: (I) control, (II) initiation of germination decrease, and (III) remarkable germination decrease, were further processed for transmission electron microscopy (five samples for each point). For transmission electron microscopy, ultra-thin sections were mounted on grids, stained with uranyl acetate followed by lead citrate (Maldonado and Lott, 1991), and examined in a Zeiss EM109T microscope.

FIGURE 2. Germination curves of *E. edulis* seeds dried by two different methods. A and B: Seeds dried with silica gel at 25°C and 2% R.H. Germination (G) and normal germination (NG) are expressed in function of A) embryo water content, and B) seed water content. C and D: Seeds dried in drying chamber at 19°C and 21% R.H. Germination (G) and normal germination (NG) are expressed in function of C) embryo water content, and D) seed water content. Vertical bars represent sd of the mean. Different letters indicate significance as calculated by Tukey’s test at ≤0.05. In all curves, an arrow indicates critical water content. In C, an interval between 48.8 and 43% water content is considered as critical for the embryo.
Conductivity tests

Electrolyte leakage from the embryos was determined by placing six embryos in 10 ml of distilled water at 15°C. Conductivity was measured after 3 h of soaking. Results were expressed as μS·mg⁻¹ of dry embryo tissue. Three replications were carried out. Values corresponded to mean ± SD. To avoid damage of imbibitions, embryos were placed for three hours in 100% relative humidity (over water) before immersion in distilled water.

Statistical analysis

To determine the critical water content values, percentage data for germination and normal germination were transformed in arcsin $\sqrt{(x/100)}$ and subjected to analysis of variance and comparison of the means by Tukey’s test at $\alpha = 0.05$.

Results

Shortly after shedding, the germination percentage was 97.5%. The mean water contents were 85.3% for isolated embryos and 48.2% for whole seeds (wet weights).

Drying rates of both embryos and whole seeds were higher in the silica gel (fast drying) treatment than in the drying chamber (slow drying treatment) (Fig. 1). After 6 days of the silica gel treatment, water content decreased to 44.9% for embryos and 29.7% for whole seeds, while the corresponding values for the drying chamber treatment were 54% and 37% (Fig. 1).

Under the silica gel treatment, the germination percentage gradually declined to 80% (Figs. 2A and B) when the water content was decreased to 48.5% (embryo) and 35.9% (whole seed). Under the silica gel treatment, the germination percentage gradually declined to 80% (Figs. 2A and B) when the water content was decreased to 48.5% (embryo) and 35.9% (whole seed). The germination percentage dropped to 22% with slightly more drying: 3.6% in the embryo and 8.2% in the whole seed, in regards to water content. Consequently, the values of 48.5% (embryo) and 35.9% (whole seed) water content were taken to be the critical water contents.

Under the drying chamber treatment, the critical water content for germination was not visible (Figs. 2C and D). However, statistical analysis indicated that the narrow interval between 43 and 48.8% water content could be considered as the critical water content for embryos (Fig. 2C).

Normal germination was always lower than germination for the same water content (Figs. 2A-D). Under the silica gel treatment, normal germination showed a slight increase to 81% that was not statistically significant when water content decreased to 60.6% (embryo) and 40.5% (whole seed). At 55% (embryo) and 38% (whole seed) water content, normal germination markedly decreased, and, therefore, those values were considered as critical water contents (Figs. 2A and B). Under the drying chamber treatment, critical water contents for normal germination were 67% (embryo) and 41.7% (whole seed) (Figs. 2C and D).

For normal germination, critical water contents for both embryos and whole seeds were higher for the drying chamber treatment than for the silica gel treatment, i.e., the susceptibility of normal germination water loss was dependent on the drying rate. Therefore, unlike under slow drying, under fast drying the seed can reach a lower water content without affecting normal germi-
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nation. For germination, comparisons between the critical water contents at the two drying rates could be established only for the embryo water content (Figs. 2A and C). Comparison could not be established for whole seed water content because no critical water content value was observed for drying chamber treatments (Fig. 2D).

Embryo critical water content resulted in similar values for both drying treatments, which suggests that germination is dependent on the water content and not on the drying rate.

When seeds were dried by the silica gel treatment and germination had declined to 80%, cells of the apical meristems showed structural changes indicative of some deterioration (Figs. 3A-D). Compared with freshly harvested seeds, the dried seeds had chromatin that was more compacted, and, consequently, the nucleoli were difficult to distinguish. Some vacuoles became confluent and formed larger compartments (Fig. 3D). Most cells maintained their shape, but some cell walls appeared to be less rigid. The confluence of lipid bodies was characteristic. Also, there was localized withdrawal of the plasmalemma from the cell wall where some vesicles had been extruded.

When seed water content was reduced to below critical values and loss of germination was observed, the general structure of all embryo tissues showed evidence of considerable damage (Figs. 3E and F). The chromatin and the nucleoplasm formed an almost homogeneous mass. Cells were severely plasmolyzed and extra-plasmalemma vesicles, indicative of exocytosis, were visible (Fig. 3F). Vacuoles were reduced in the contracted cytoplasm.

Electrolyte leakage from embryos progressively increased with desiccation, particularly at low water contents, which suggested membrane deterioration (Figs. 4A and B). The critical water contents were not associated with changes in electrolyte leakage.

Discussion

This study compared the susceptibility to desiccation of embryos to that of whole seeds. Berjak et al. (1989) describe the difficulty of comparing water contents when embryo or whole seed are measured. We found that such difficulty was exacerbated in this species because the embryo constitutes only 0.54% of the seed fresh weight. In addition, dehydration curves showed that water loss was faster for the embryo than for the endosperm.

Queiroz et al. (1986) indicate that during germination of fresh E. edulis seeds, the percentage of button protrusion is high and positively correlated with plumule emergence. However, we observed that the development of a seedling with three leaves did not always follow button protrusion, and some death could occur after plumule emergence. A plantule with three leaves seems to be the most appropriate growth stage for germination evaluation; i.e., normal germination was lower than germination at most embryo and whole seed water contents. Consequently, two germination stages were
evaluated: 1) button protrusion (germination) and 2) a subsequent stage represented by the plumule emergence, i.e., seedlings exhibiting both a well-developed radicle and a shoot with the first three leaves, the third leaf being the first one with pinnate structure (normal germination). Using both germination stages we found the following results: (a) For both drying treatments, desiccation sensitivity values were higher for both germination and normal germination based on the embryo water content, i.e., the embryo desiccation sensitivity was masked for measurements based on the whole seed water content. This result demonstrates the convenience of basing evaluations on both the embryo and the whole seed water contents to obtain a better knowledge of the desiccation sensitivity of these seeds. (b) For both drying treatments, desiccation sensitivity was higher for normal germination than for germination based on both the embryo and the whole seed water contents; therefore, a reliable evaluation of drying effects on germination should include the stage with the plantule with three leaves. (c) Under the slow drying treatment and based on the whole seed water content, critical water content was visible for normal germination and not for germination; therefore, germination was not appropriate for detecting critical water content based on the whole seed water content. (d) Critical water contents for germination were most clearly established in the silica gel treatment based on both the embryo and whole seed water contents. Under the drying chamber treatment, critical water contents were only established for normal germination based on both the embryo and the whole seed water contents. (e) For both drying treatments, critical water content was higher for normal germination than for germination. (f) For the drying chamber treatment, when germination was based on the whole seed water content, critical water content was not visible.

Some studies report that critical values for recalcitrant seeds are subject to variation according to the speed of dehydration (Farrant et al., 1985; Pritchard, 1991). However, there are conflicting opinions on this subject (Bonner, 1996; Finch-Savage, 1992; Pritchard et al., 1995). For *E. edulis* seeds, we found that the critical water content for normal germination depended on the drying method, but that for germination did not. Under the drying treatments used here, critical points for the viability of the *E. edulis* embryo could be more precisely determined when the germination parameters were evaluated in function of the water content of the embryo, but not of the whole seed. These results highlight the need to evaluate the water content based on the embryo.

Queiroz et al. (1986) found that, when *E. edulis* seeds with initial water content of 51.3% are slightly dried, their germination percentage significantly increases from 81% to 98%. We used seeds with an initial water content of 48.5% and observed a slight increase in the germination percentage only for normal germination (based on the embryo water content) at initial drying; however, the increase was not statistically significant. We suggest that, for newly collected seeds that differ in water content and physiological state (as those supposedly used by Queiroz et al., 1986), the initial germination percentage could be increased by slight desiccation.

As indicated by the progressive increase in electrolyte leakage and by the transmission electron microscopic images, cell membrane damages occurred especially at high desiccation levels. Critical water contents could not be associated with any abrupt change in the conductivity curve, which suggests that conductivity is not an appropriate method for detecting the critical water content in this species.

TEM images of shoot apical tissues of the *E. edulis* axis show that, in mature seeds, chromatin is present in a natural incipient condensed state, which might be related to the relative quiescence exhibited by the apical meristems (Panza et al., 2004). As in orthodox seeds, this is a reversible state where the rehydration is associated with decondensation of chromatin (Pammenter and Berjak, 1999). During *E. edulis* seed dehydration, gradual intracellular injury of organelles and cytoplasm was accompanied by an increase in chromatin compaction.

Water stress increase is conducive to DNA damage and results in germination decrease, as in *Avicennia marina* is observed by Boubriak et al. (2000). Crévecouer et al. (1976) observe that, during the germination of corn, chromatin compaction in the embryonic tissues accompanies dehydration damage. In *E. edulis* embryo tissues, chromatin damage was also visible during desiccation. In fact, increased chromatin compaction was one feature of the initially reversible deterioration of the *E. edulis* seeds, and this feature preceded the critical water content of germination. Boubriak et al. (1997) propose that the capacity of *Secale cereale* and *Avena fatua* dehydrated embryos to repair DNA when they are re-hydrated is a critical event in the maintenance of a functional genome and is an essential component in the drying tolerance mechanism. Similarly, Boubriak et al. (2000) show the loss of the capacity for DNA repair when 22% of the total water is lost from highly recalcitrant *Avicennia marina* root primordium. After a critical degree of tissue dehydration, the capacity for DNA repair in *E. edulis* embryos may also be lost and, thus, may be a possible cause for the abrupt decrease in germination.
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