Physiological and Cellular Changes of Stored Cryptocarya aschersoniana Mez. Seeds

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Abstract

Some species of the Lauraceae family produce seeds that are generally sensitive to desiccation, which makes them difficult to store. The objective of this study was to characterize changes in seed quality of C. aschersoniana from two lots, as well in the physiological and cellular aspects during 12 months of storage. Seeds were stored with their original moisture content (MC) or after pre-drying to 35% MC in a cold chamber (5 °C) at a relative humidity of 40%. Seeds were sampled and tested at time 0, 3, 6 and 12 months of storage regarding to moisture content, germination and ultrastructural features. The seeds were dispersed with dormancy that was overcome by the cold storage condition and the reserves in undried seeds were partly consumed during storage. Both undried and pre-dried seeds remained viable for at least 12 months.

Keywords: Lauraceae, seed conservation, storage, ultrastructural analyses.

1. INTRODUCTION AND OBJECTIVES

In order to have an efficient storage, the seed moisture content must be low. However, seeds that are sensitive to desiccation, including seeds of many tropical species, need to be kept sufficiently moist for metabolism not to be interrupted (Walters et al., 2001) and for gas exchange to occur. Because of their active metabolism, desiccation-sensitive seeds can germinate during storage and, even if there is no sufficient water for germination to complete, their reserves continue to be consumed, leading to a loss of viability (Marcos-Filho, 2005). Storage alternatives that have been proposed for these seeds include the removal of free water, a procedure that helps to maintain seed viability (Walters, 2000; Bonjovani & Barbedo, 2008), by lowering the metabolic rates, as well as prevent infestations by microorganisms (Lima et al., 2018). In some cases, this partial drying may also improve the seed quality by provoking a moderate stress condition that generates a certain positive response in the seed (Walters, 2000).

The Lauraceae family is one of the most diverse in Brazilian forestry formations, with species frequently being found in the remaining riparian forests in Southeast Brazil. Some species of this family belong to the ecological group climax (Lorenzi, 1998) and produce seeds irregularly that are generally sensitive to desiccation (Carvalho et al., 2008; Vicente et al., 2016) and, in some cases, with dormancy (Jaganathan et al., 2019). In particular, seeds of Cryptocarya aschersoniana have been reported to be sensitive to desiccation (Carvalho, 2000; Hirano, 2004; Muxfeldt et al., 2012). These seeds are dispersed during the rainy season (Lorenzi, 1998) and often need to be stored, even for short periods, to be sown later in order to synchronize the end of the seedling production cycle with the beginning of the next rainy season. Therefore, the objective of this study was to characterize changes in seed quality of C. aschersoniana, as well in the physiological and cellular aspects during 12 months of storage.

2. MATERIAL AND METHODS

2.1. Characterization of the material

Ripe (yellow) fruits of C. aschersoniana were collected in Lavras, Minas Gerais state, Brazil, in February, in two different years. The fruits were processed under running water
and through a sieve until the pulp was completely removed. The dispersal structures were considered to be the "seeds", formed by the seed itself within a woody endocarp. The seeds were spread in a single layer in the open air until the surface water had evaporated. These seeds were considered as "recently harvested".

Some of the seeds were maintained at their initial moisture content. Other seeds were pre-dried until a target moisture content of 35% was reached. This pre-drying step was performed in drying boxes with silica gel at the bottom, in an air-conditioned room (20 °C). The silica gel was changed whenever the color indicator for humidity (blue) turned pale. The internal condition in the boxes was maintained at 20 °C with a relative humidity (RH) between 13.5% and 40%. The 35% target moisture content was chosen because pilot tests performed in the same laboratory indicated that this level does not cause a significant drop in germination rates, however, below this moisture content, the seeds had an increased risk of loss of viability (data not shown).

Samples of the obtained biological material were taken for the initial physiological characterization tests (moisture content and germination) and image analyses (electronic microscopy), as described below.

### 2.2. Moisture content test

The oven moisture test was performed at 105 °C for 24 h (Brasil, 2009), in 4 replications of 5 cut seeds placed in aluminum foil packets. The moisture content was calculated on a wet basis as the average of the 4 repetitions.

### 2.3. Germination test

The germination test was performed in 4 replications of 25 seeds that had been previously washed in 1.0% sodium hypochlorite for 10 min. Seeds were placed into plastic trays (27 × 40 cm) containing autoclaved sand, which were incubated in a Mangelsdorf germinator at 25 °C under continuous light. Seeds were considered to have germinated when they produced normal seedlings within a 120-day period. At the end of the test, seeds that had not germinated were cut and incubated in a solution of tetrazolium (Vetec, Rio de Janeiro, Brazil) at 0.5%, 25 °C, for 24 h. They were considered viable if they stained pink/red in color, indicating respiratory activity in their tissues.

### 2.4. Ultrastructural analysis

The ultrastructure of the samples was analyzed by scanning electron microscopy (SEM). Samples (5 seeds) were prepared according to the following protocol: seeds were cut lengthwise in the area of the embryonic axis. Samples were fixed in a modified Karnovsky solution (2.5% glutaraldehyde, 0.05 mol/L CaCl₂, 0.001 mol/L sodium cacodylate buffer, pH 7.2) for at least 24 h. Samples were transferred to a cryo-protectant solution (glycerol 30% v/v in water) for 30 min and sectioned longitudinally in liquid nitrogen using a scalpel blade. The specimens were dehydrated in a gradient acetone series (1 × 25%, 50%, 75% and 90%, 3 × 100%; 10 min each), and then transferred to a critical-point dryer (Bal-Tec) to evaporate the acetone without loss of tissue conformation. The samples were placed in stubs lined with aluminum foil, over carbon tape, and submitted to gold-sputtering in a Sputter Coater SCB 050 gold evaporator. Finally, the samples were examined by scanning electron microscopy (SEM) (LEO EVO 40 XVP). Observations focused on the transition area from the embryonic axis to the cotyledons.

### 2.5. Storage

After the initial tests were completed, samples with the two moisture contents were stored in a cold chamber (5 °C, 40% RH) in semipermeable packages (sealed plastic bags with holes punched in them). The above evaluations were repeated after 3, 6 and 12 months of storage and compared to the results before storage.

### 2.6. Data analysis

All the experiments were set using a completely randomized design. Data had the normality analyzed using the Shapiro Wilk Test, if no normal distribution was observed, those were transformed into arcsin of √(x/100). Normal data or those normalized by the transformation were analyzed by ANOVA and Tukey's test at 5% probability. Those data whose distribution was not normal, even after transformation, were analyzed by Generalized Linear Models (GLM) and if detected significant differences among the variables, the data were compared by Duncan's test at 5% probability. All analyses were carried using the software R for Windows (R Core Team, 2020).

### 3. RESULTS AND DISCUSSION

From the initial results (Tables 1, 2 and 3), the average moisture content of seeds collected in year one was 40.6%, and the average percentage of germination was 15%. Seeds harvested in year two showed an average moisture content of 47.0%, and percentage of germination of 6%. After pre-drying, the average moisture content and germination
were 35.5% and 9%, respectively, in the first year, and 37.8% and 9%, respectively, in the second year. The moisture content of the seeds remained stable over 12 months of storage, without significant variation (p < 0.05).

After storage, the percentage of germination increased irrespective of the collection year, with the highest increases being observed after 6 months of storage, remaining stable for up to 12 months for both lots (Figure 1). Slight reduction of moisture by pre-drying did not influence the final percentage of germination (p < 0.05) at time zero (Tables 2 and 3), as already reported for the study species (Hirano, 2004; Muxfeldt et al., 2012) and for recalcitrant seeds of *Inga vera* (Faria et al., 2004). After 12 months of storage, although there was no effect of pre-drying on final percentage of germination for both lots, seeds from the second year subjected to pre-drying germinated slower than those that were stored wet (Table 3).

### Table 1. Variation in moisture content of *Cryptocarya aschersoniana* seeds during storage. SD = Standard Deviation, CV = coefficient of variation.

| Collection year | Storage condition | Storage time (months) | Mean ± SD   | CV (%) |
|-----------------|-------------------|-----------------------|-------------|--------|
|                 |                   | 0                     | 3           | 6      | 12     |
| Year 1          | Wet               | 40.6                  | 40.4        | 40.6   | 39.5   | 40.3 ± 0.53 | 4.12  |
|                 | Pre-drying        | 35.5                  | 34.9        | 35.4   | 35.8   | 35.4 ± 0.37 | 2.87  |
| Year 2          | Wet               | 47.0                  | 45.8        | 47.6   | 48.3   | 47.2 ± 1.06 | 3.93  |
|                 | Pre-drying        | 37.8                  | 36.4        | 35.7   | 38.4   | 37.1 ± 1.24 | 6.94  |

Figure 1. Behaviour of *Cryptocarya aschersoniana* seeds stored in cold chamber (5 °C / 40% RH) for 0, 3, 6 and 12 months. Regression equations and R² values are as follows: A) Germinated: y = 21.466 + 17.839x - 0.988x² (R² = 0.9309); Dormant: y = 55.9205 - 12.8201x - 0.7039x² (R² = 0.8015); B) Germinated: y = 12.2803 + 16.1061x - 0.8039x² (R² = 0.8812); Dormant: y = 74.3295 - 16.7216x - 0.8933x² (R² = 0.8812); C) Germinated: y = 13.2273 + 17.1288x - 0.9874x² (R² = 0.6889); Dormant: y = 76.645 - 19.741x + 1.169x² (R² = 0.8242); D) Germinated: y = 13.036 + 17.961x - 1.076x² (R² = 0.7247); Dormant: y = 64.227 - 18.038x + 1.068x² (R² = 0.8632).
Table 2. Germination of Cryptocarya aschersoniana seeds (first year) after different storage times. G = Germination; DoS = Dormant seeds; DeS = Dead seeds; MGT = Mean germination time; GSI = Germination speed index.

| Storage time (months) | Storage condition | G  | DoS | DeS | MGT (days) | GSI% |
|-----------------------|-------------------|----|-----|-----|------------|------|
| 0 Wet                 | 15 a 59 a 26 a 96 a | 0.043 a |
| Pre-drying            | 9 a 74 a 17 a 97 a | 0.027 a |
| 3 Wet                 | 70 a 16 a 14 a 52 a | 0.287 a |
| Pre-drying            | 55 b 34 a 11 a 50 a | 0.258 a |
| 6 Wet                 | 90 a 10 a 0 a 33 a | 0.643 a |
| Pre-drying            | 79 b 5 a 16 a 39 a | 0.482 b |
| 12 Wet                | 93 a 3 a 4 a 39 a  | 0.549 a |
| Pre-drying            | 90 a 3 a 7 a 43 a  | 0.509 a |

Mean values followed by the same letter in the column, within each storage time, do not differ from each other. Data were compared by Duncan test at 5% probability.

The total percentage of germination increased by storing seeds for both years of collection, irrespective of their initial condition (wet or pre-dried), simultaneously to the decrease in the percentage of dormant seeds (Figure 1). Pre-drying the seeds did not have any effect on the percentage of dead seeds throughout the storage time (Tables 2 and 3). It can also be observed that the mean germination time (MGT) decreased, coinciding with increasing values of germinating speed index (GSI) along the storage time (Figure 2). In other words, the longer the storage time, the faster the germination. These results suggest that the low initial germination was due to some kind of dormancy that was released during storage (Figures 1 and 3). A similar behaviour was found in seeds of Carthamus tinctorius – Asteraceae (Oba et al., 2019) and of other species of Lauraceae (Carvalho, 2000; Carvalho et al., 2008).

Table 3. Germination of Cryptocarya aschersoniana seeds (second year) after different storage times. G = Germination; DoS = Dormant seeds; DeS = Dead seeds; MGT = Mean germination time; GSI = Germination speed index.

| Storage time (months) | Storage condition | G  | DoS | DeS | MGT* (days) | GSI*% |
|-----------------------|-------------------|----|-----|-----|------------|------|
| 0 Wet                 | 6 a 83 a 11 a 65 a | 0.027 a |
| Pre-drying            | 9 a 69 b 22 a 62 a | 0.049 a |
| 3 Wet                 | 75 a 11 a 14 a 56 a | 0.406 a |
| Pre-drying            | 68 b 7 a 25 a 47 a | 0.484 a |
| 6 Wet                 | 66 a 13 a 21 a 37 a | 0.685 a |
| Pre-drying            | 74 a 4 b 22 a 39 a | 0.655 a |
| 12 Wet                | 79 a 6 a 15 a 23 a | 1.130 a |
| Pre-drying            | 75 a 0 a 25 a 38 a | 0.608 b |

Mean values followed by the same letter in the column, within each storage time, do not differ from each other. Data were compared by Duncan test at 5% probability and *Tukey test at 5% probability.

Figure 2. Germination speed index (GSI) and mean germination time (MGT) of Cryptocarya aschersoniana seeds stored in cold chamber (5°C / 40% RH) for 0, 3, 6 and 12 months. Regression equations and R² values are as follows: A) GSI: y = 0.013119 + 0.143698x - 0.008179x² (R² = 0.8457); MGT: y = 96.306 - 16.770x + 1.004x² (R² = 0.9563); B) GSI: y = 0.01584 + 0.10678x - 0.00545x² (R² = 0.8412); MGT: y = 95.4254 - 15.2399x + 0.9088x² (R² = 0.9381); C) GSI: y = 0.08773 + 0.09079x (R² = 0.7049); MGT: y = 64.262 - 3.589x (R² = 0.6294); D) GSI: y = 0.064171 + 0.148392x - 0.008626x² (R² = 0.7049); MGT: y = 56.137 - 1.765x (R² = 0.3077).
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Figure 3. Germination of Cryptocarya aschersoniana seeds stored in cold chamber (5 °C / 40% RH) for 0, 3, 6 and 12 months.

C. arshersoniana seeds probably have nondeep physiological dormancy, caused by the presence of the endocarp. According to Baskin and Baskin (2014), covering structures may reduce the rate of imbibition, restrict movement of oxygen to the embryo, mechanically restrict embryo growth and contain germination inhibitors.

This behavior may be related to the ecology of the species, and may reflect its adaptation to naturally occurring conditions. C. aschersoniana is typical of Brazilian subtropical forests and, in lower latitudes, often appears in higher altitude forest (Lorenzi, 1998). Seeds are released at the end of the rainy season (Hirano, 2004) and they need to survive through a cold and dry season and remain dormant until the conditions are favorable for germination and seedling development, as reported by Jaganathan et al. (2019), who described the behaviour of seeds of several Lauraceae species that are dispersed with nondeep physiological dormancy that can be released by cold stratification. Similar adaptations are common in recalcitrant seeds of temperate species, as seen in acorns oak (Pasquini et al., 2011, 2012). These seeds generally disperse in the autumn and must survive through the winter without germinating, which involves a process of dormancy and some strategy to reduce water loss (Pammenter & Berjak, 2000; Bewley et al., 2013).

Similar results for storing Lauraceae seeds were found for 4 species after 4 months at 4.0 to 10.0 °C (Hirano, 2004), Nectandra nitidula after 90 days and Nectandra lanceolata after 180 days of storage in a cold chamber (Carvalho, 2006). Persea pyrifolia and Cryptocarya aschersoniana seeds overcome dormancy when stored moist for 3 months in a cold chamber (Carvalho, 2000). For Quercus ilex, a species taken as a model for recalcitrance and dormancy, storing the seeds in wet sand at 3.0 °C encouraged germination after 1 month of storage (Pasquini et al., 2011).

The SEM findings (Figure 4) show that there was no damage to the cell structure after 12 months of storage, however, suggest that seeds stored at their original moisture (Figure 4B) had lower reserves of starch granules compared to seeds that were pre-dried before storage (Figure 4D).

To preserve seeds in the long term, their metabolic rates must be reduced. This reduction can be achieved and the germination process impeded by reducing both the amount of free water (Walters, 2000; Walters et al., 2005) and the temperature (Hong & Ellis, 2002). However, even at reduced temperatures, the metabolism may remain active during storage (Bonjovani & Barbedo, 2019) and is probably more intense in seeds that had not been pre-dried. Wet storage expose the seeds to a stress condition and, according to Oliveira et al. (2020), when the amount of water available is not enough to complete germination, it may occur a disordinated consumption of reserves, leading to viability loss.
4. CONCLUSIONS

Taken together, the results of this study suggest that *C. aschersoniana* seeds are released with dormancy and, one way to overcome this dormancy is storing the wet seeds in plastic bags in cold chamber (5 °C) at 40% RH. Under this storage condition, the viability of the seeds can be kept for at least 12 months, regardless of whether the seeds are stored with their initial moisture content or pre-dried to a moisture content of 35.5%.

Microscopic analyses suggested that *C. aschersoniana* seeds stored in a cold chamber with their initial moisture partly consume their reserves by metabolism, which may lead to a loss of viability during longer storage periods.

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