Seed metrics and influence of temperatures and pre-germination treatments on germination of *Libidibia ferrea* seeds

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ABSTRACT: The objective of this study was to evaluate seed metrics, optimum temperature for germination and efficiency of five pre-germination treatments for overcoming dormancy of *L*. *ferrea* seeds. Seeds were subjected to constant temperatures of 15, 20, 25, 30 and 35°C and to the following pre-germination treatments: nicking with pincers; immersion in water for 24 hours at room temperature; scarification with sandpaper; immersion in caustic soda for 60 minutes; and control (untreated seeds). Germination rate was assessed by germination percentage and germination speed index. The biometric characteristics of the evaluated seeds were: longitudinal length, width and thickness, using a digital caliper with a precision of 0.05 mm. A completely randomized design was used with four replicates and means were compared by Tukey test at 5% probability. The best germination performance was obtained in the 15-30°C temperature range and by using chemical scarification with immersion in caustic soda, and mechanical scarification by nicking with pincers and by rubbing on sandpaper.

Keywords: dormancy; scarification; seed dimensions.

Caracterização biométrica de sementes e influência de temperaturas e tratamentos pré-germinativos em pau-ferro

RESUMO: O objetivo do presente trabalho foi avaliar a biometria de sementes, a temperatura ótima e a eficiência de cinco tratamentos pré-germinativos para a superação de dormência das sementes de *L*. *ferrea*, sugeridos na literatura. As sementes foram submetidas a temperaturas constantes de 15, 20, 25, 30 e 35 ºC e aos tratamentos pré-germinativos: corte com alicate no lado oposto ao hilo, imersão em água por 24 horas em temperatura ambiente, escarificação com lixa d’água nº 4, imersão em soda cáustica por 60 minutos e a testemunha (sem tratamento). A germinabilidade foi avaliada pela porcentagem de germinação e o índice de velocidade de germinação. As características biométricas das sementes avaliadas foram: comprimento longitudinal, largura e espessura, utilizando-se paquímetro digital com precisão de 0,05 mm. O delineamento experimental foi inteiramente casualizado, e as médias foram comparadas pelo teste Tukey a 5% de probabilidade. O melhor desempenho germinativo das sementes foi obtido nas temperaturas entre 15 e 30 ºC e nos tratamentos pré-germinativos de escarificação química com imersão em soda cáustica e escarificação mecânica por meio do desponte com alicate e o atrito em lixa.

Palavras-chave: dormência, escarificação, dimensão da semente.

1. INTRODUÇÃO

*Libidibia ferrea* Mart. ex Tul., commonly known in Brazil as pau-ferro or jucá, belongs to the family Caesalpiniaceae and is a typical tree species found in the Caatinga. This tree has high ornamental and medicinal potential (Silva et al., 2015), and its high-density wood is used in the lumber industry. *Libidibia ferrea* can be found from Piauí to Rio de Janeiro states due to its high seed dispersal.

Seed metrics is regarded as an important tool for detecting genetic variability within populations of a given species, thereby relating this variability to environmental factors, which provides important information for defining ecological characteristics such as seed dispersion, dispersion agents, and seedling setting (FERNANDES et al., 2012).

Temperature affects both germination percentage and speed at which germination occurs (Ranzani et al., 2016) owing to its direct impact on water uptake by seeds and on biochemical reactions regulating metabolic processes during germination. According to Carvalho; Nakagawa (2012), there is an optimum temperature range for germination in which the efficiency of the process is maximized, i.e., maximum germination in the least amount of time.

There is an increasing demand for native tree species due to environmental awareness programs (Nascimento et al., 2012); therefore, it is important to know the ideal condition for seed germination, especially because of different responses each tree species can have as a function of
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2. MATERIAL E MÉTODOS

2.1. Seed collection and experimental area

Ripe fruits of *L. ferrea* were manually collected in April 2016, in different trees.

The seeds were manually removed from pods with the aid of a hammer. After processing, seeds were selected and placed in glass containers (amber), sealed up, and stored in a refrigerator for 120 days.

2.2. Seed characterization

The seeds metrics was measured, with 100 seeds with 100 seeds in each treatment, were longitudinal length, measured from base to tip; width and thickness, both of which measured in the middle of the seed, using a 0.05 mm accuracy digital caliper.

Eight replicates consisting of 100 seeds each were used to determine the number of seeds in a kilogram. Seeds were randomly picked and then weighed on an analytical scale. Seed moisture content was measured by the oven drying method, at 105 ± 3 ºC, during 24 hours, as standardized by the Rules for Seed Analysis (Brasil, 2009). We used four replicates for each treatment, with 50 seeds. Data are expressed as percentage based on fresh weight of the sample.

The experiment was carried out in a completely randomized design (CRD) 5 x 5 factorial experiment (five temperatures x five pre-germination treatments), with four replicates and each replicate consisted of 20 seeds.

2.3. Experimental conditions and treatments

Temperatures used were: 15, 20, 25, 30, and 35°C, in a 12-hour photoperiod using fluorescent bulbs (Phillips 15W) in B.O.D. incubators (model 347 FANEN).

To overcome the coat-imposed seed dormancy, four pre-germination methods were used: nicking with pincers on the opposite side to the hilum; immersion in water for 24 hours at room temperature; scarification with wet/dry sandpaper; and immersion in caustic soda for 60 minutes. Before setting up germination tests, seeds were immersed in a 5% sodium hypochlorite for 5 minutes and then rinsed in distilled water.

2.4. Seeds immersed in water at room temperature

Seeds were immersed in 150 mL of distilled water for 24 hours at room temperature.

2.5. Chemical scarification

A basic solution was prepared using water and caustic soda at 20% (65 g of caustic soda to 325 mL of water) following recommendations of Garcia & Azevedo (1999). First, seeds were placed in a beaker; then, 65 g of caustic soda was added to the beaker containing the seeds; and, finally, 325 mL of distilled water was added to it, stirring the solution with a glass rod and letting it rest for 60 minutes. After removing seeds from the solution, caustic soda residues were rinsed off them.

2.6. Mechanical scarification by nicking

With the aid of pincers, seeds were nicked on the opposite side to the hilum to promote the rupture of the coat without damaging the embryo.

2.7. Mechanical scarification using sandpaper

Seeds were rubbed against a 40-grit wet/dry sandpaper on the opposite side to the micropyle to rupture the coat without damaging the embryo.

Seeds were sown in transparent gearbox-type boxes measuring 10 x 10 x 4 cm (length, width, and depth, respectively) containing vermiculite previously sterilized in oven for 24 hours at 200°C and wetted with distilled water. Moisture content in the substrate was maintained by watering it every two days in accordance with the field capacity of it.

2.8. Germination characteristics

Germination percentage (% GERM), germination speed index (GSI), mean germination time (MGT), the day on which the first germination event occurs (NIG), and the day on which the last germination occurs (NFG) were evaluated.

Germinated seeds were counted daily from the 54rd day on when there were no germination events for three consecutive days. Maguire (1962), developed the following formula for the GSI (Equation 1):

\[
\text{GSI} = \frac{G_1}{N_1} + \frac{G_2}{N_2} + \cdots + \frac{G_n}{N_n} \quad (eq. 01)
\]

where: \(G_1, G_2, G_n\) = number of normal seedlings at first, second and last counting; \(N_1, N_2\) and \(N_n\) = number of days to first, second and last counting.

Labouriau; Valadars (1976) cited the formula used to calculate the germination time as follows (Equation 2):

\[
GT = i = kNi \times Tii = 1kNi \quad (eq. 02)
\]

where \(N_i = \) number of seeds germinated at time \(T_i\) (not the cumulative number, rather the number of the \(i\)-th observation); \(T_i = \) time between sowing and the \(i\)-th observation; \(k = \) last time at which seeds germinated.
Length, width and thickness were subjected to exploratory analysis (descriptive statistics) of estimates. Results were assigned to classes and plotted as frequency distribution graphs ([BEZERRA et al., 2012]). The remaining characteristics were tested by analysis of variance using the F test ($p \leq 0.05$). To compare means of pre-germination treatments, Tukey test at 5% probability was used. Regression models were adjusted to temperatures using SISVAR software (version 5.4) provided that means were significantly different ([FERREIRA, 2014]).

3. RESULTS

Weight of 1,000 seeds was 183.1 g, which allows us to infer that a kilogram can contain 5,461 seeds of *L. ferra*. Moisture content was 7.22%, confirming that the seeds are orthodox. Mean seed size as to length, width and thickness were 9.9, 6.9 and 4.0 mm, respectively (Table 1).

Coefficients of variation are below 20%, which means that data are reliable. As for maximum and minimum values of measured variables, we observed differences of 3.4, 4.4 and 1.7 mm, with the least variation for thickness. Regarding the standard deviation, seed thickness varied the least. This lower variability may be a result of genetic conditions and local environmental variations.

Figure 1 shows the frequency distribution of *L. ferrea* seed metrics as to length, width and thickness.

![Figure 1](image1.png)

Figure 1. Length (A), width (B) and thickness (C) of *L. ferrea*, average values of 100 seeds.

Curves yielded from slicing significant interactions between treatments for overcoming seed dormancy and temperatures are shown in Figures 2 and 3. Results indicated that interactions between these two factors had an influence on germination percentage, germination speed index, mean germination time, and number of days to start and finish germination of *L. ferrea* seeds tested at a significance level of 5%. The data are best described by cubic and quadratic equations.

The highest estimated germination percentage (110.0%) was recorded in seeds scarified with caustic soda at 24 °C, more than 100% higher than that recorded in the control group (6.3%); though, from 24 °C forth, germination decreased. As for mechanical scarification by nicking and rubbing against sandpaper, there is an increasing trend of germination up to 102.4% and 91.5% (estimated maximum point) at 24 and 22 °C, respectively.

![Figure 2](image2.png)

Figure 2. Final germination percentage (GERM) and germination speed index (GSI) of *L. ferrea* seeds subjected to different temperatures and pre-germination treatments. Letters different between treatments at each temperature differ from each other by Tukey test ($p \leq 0.05$).

![Figure 3](image3.png)

Figure 3. Percentage of germination at different temperatures for *L. ferrea* seeds under different treatments.

Table 1. Mean length, width and thickness of 100 *L. ferrea* seeds.

<table>
<thead>
<tr>
<th>Variables</th>
<th>Mean</th>
<th>Standard deviation</th>
<th>CV (%)</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length (mm)</td>
<td>9.9</td>
<td>0.6</td>
<td>5.7</td>
<td>8.1</td>
<td>11.1</td>
</tr>
<tr>
<td>Width (mm)</td>
<td>6.9</td>
<td>0.7</td>
<td>9.7</td>
<td>4.2</td>
<td>8.6</td>
</tr>
<tr>
<td>Thickness (mm)</td>
<td>4.0</td>
<td>0.3</td>
<td>8.2</td>
<td>3.3</td>
<td>5.0</td>
</tr>
</tbody>
</table>

Table 1. Valores médios de comprimento, largura e espessura de 100 sementes de *L. ferrea*.
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Seeds treated by nicking and rubbing against sandpaper had higher germination percentages in comparison with the remaining treatments at every temperature tested but 25 °C at which nicking and caustic soda treatments were more efficient. At 20 °C and 30 °C, germination percentage of seeds immersed in caustic soda was not significantly different from nicked seeds; however, untreated seeds and seeds immersed in water had the lowest germination percentages regardless of the temperature. These results demonstrate that *L. ferrea* seeds do have dormancy; therefore, efficient pre-germination treatments are necessary for germination events to occur.

The interaction between the evaluated treatments indicated that the highest germination speed index (0.26) was obtained by using mechanical chiseling by topping, at a temperature of 25 °C, followed by immersion treatment, with an index of 0.22, at temperature maximum of 23 °C. In the water immersion treatment, there was a higher germination speed index (0.15), at a temperature of 19 °C, after which there was a decrease up to 30 °C, with an index of 0.03. It was also observed that without the use of efficient pre-scarification treatments, low rates of germination speed of pau-ferro seeds occur.

The water immersion treatment for 24 hours promoted a shorter germination time (0 days) at a temperature of 17 °C, after which there was an increase up to a temperature of 30 °C, with an average of 23 days. While, in the control, the average maximum germination time was 18 days, at 24 °C. The scarified seeds by topping, soda and sandpaper germinated faster, reaching the maximum average time of 5, 6 and 7 days, respectively, after sowing (Figure 3).

Seeds scarified by caustic soda, nicking or sandpaper germinated faster, reaching the maximum mean time of 10 days after sowing. After initiating tests, we recorded germination events up to the 20th and 29th day in control and immersion in water, respectively.

As for the number of days to germination as a function of temperature, regressions were significant only for the treatments control and immersion in water. Germination events of these treatments were delayed compared to the remaining treatments (Figure 4). Maximum mean time to start germination of untreated seeds was 13 days at 27 °C whereas for immersion in water at room temperature, the higher the temperature, the longer seeds take to germinate.

![Figure 3. Mean germination time (GMT) of *L. ferrea* seeds subjected to different temperatures and pre-germination treatments.](image3)

**Figure 3.** Mean germination time (GMT) of *L. ferrea* seeds subjected to different temperatures and pre-germination treatments. Different letters across treatments for each temperature differ from each other by Tukey’s test (p ≤0.05).

![Figure 4. Number of days to germination (NIG) and number of days to finish germination (NFG) of *L. ferrea* seeds subjected to different temperatures and pre-germination treatments.](image4)

**Figure 4.** Number of days to germination (NIG) and number of days to finish germination (NFG) of *L. ferrea* seeds subjected to different temperatures and pre-germination treatments. Different letters across treatments at each temperature differ from each other by Tukey test (p ≤0.05).

The number of days to finish germination as a function of temperatures yielded a significant regression for immersion in water, caustic soda and control (Figure 4). Caustic soda reduced the number of days to finish germination up to 26 °C, and from there on, number of days rose up again, while control treatment had an inverse behavior. NFG increases in the control group up to 27 °C, and from there on, it decreases. Immersing seeds in water at room temperature shortened number of days to finish germination between 15 and 20 °C. From the latter temperature on, subsequent increments varied, then, past 30 °C germination decreased again.

4. DISCUSSION

Moisture content is consistent with that recorded by Lima et al. (2006) and Alves et al. (2009), which reported moisture contents of 7.46% and 6.9% in their studies with seeds of the same species evaluated in our study. This low seed moisture content lengthens viability of seeds (MATOS, 2015).

Seed metrics is fundamental for characterization of a seed lot by providing information about different sizes and storage
content of seeds. According to Silva et al. (2012), these differences might be linked to strategies of using nutrients and available water resources, to local environmental variations, and to the own genotypic diversity of populations that might result in different phenotype characteristics for the species.

Pereira et al. (2011) demonstrated that *Hymenaea stigonocarpa* var. *stigonocarpa* has major variation in fruit size and seed weight. Silva et al. (2012) differentiated *Hymenaea intermedia* Ducke from *Hymenaea maritana* Hayne by comparing seed metrics and emphasized the importance of assessing seed metrics when differentiating species.

Untreated seeds and seeds immersed in water at different temperatures had low germination percentage, which indicates that in the absence of a scarification pretreatment, germination events are slow, uneven and, therefore, unsuitable for seedling production, due to the presence of a coat-imposed dormancy (COELHO et al., 2013). Conversely, the use of efficient pre-germination treatments, such as chemical scarification with caustic soda, and mechanical scarification by either nicking or rubbing against sandpaper, can result in high germination rates.

The optimum temperature range to germinate seeds is within what Brancalion et al. (2010) reported because seeds of most tropical and subtropical forest species have maximum potential for germination in the 25 to 30 °C range. Germination events occur more rapidly and efficiently within an optimum temperature range, which, however, depends on the species and the species’ regions of origin.

Temperature influences biochemical reactions of germination, which may affect both capacity and speed of seed germination. Seeds germinate most readily when they are in the characteristic temperature range of the species; however, the time taken to obtain the maximum germination percentage is dependent on the temperature. Increasing temperature makes water more fluid and with more kinetic energy, facilitating its movement from the outer to the inner side of the seed and, in consequence, water uptake is increased, so is the speed of metabolic reactions.

Based on findings of this study, we recommend the use of chemical scarification by immersion in caustic soda, mechanical scarification by nicking with pincers and by rubbing against sandpaper as the most efficient techniques to overcome seed dormancy within the optimum temperature range of 20 to 30 ºC. The optimum temperature for germination is a physiological adaptation of seeds to local environmental conditions where the species occurs or is grown; thus, this temperature may have a direct relationship with the biome in which the seeds were produced. Species with different ecological and geographical distributions produce seeds varying as to temperature requirements for germination (BRANCALION et al., 2010).

Mechanical scarification of the seed coat was efficient when breaking the dormancy of seeds of various species of the family Fabaceae such as, *Cassia fistula* L. (Bezerra et al., 2014; Guedes et al., 2013), *Erythrina velutina* Wildl (Santos et al., 2013), and *Centrosema pluminer* Benth (GAMA et al., 2011).

The decision between chemical or mechanical scarification should be based upon factors other than efficiency. Chemical scarification is advantageous due to its speed and decreased labor requirements. Mechanical scarification is more laborious and takes longer when done manually; though, mechanical methods do not use chemical products, thereby avoiding accidents with these products. Oliveira et al. (2017) pointed out that manual techniques, such as chemical scarification by nicking, either on the opposite side to the hilum or at the tip where the seed attached to the pod, is more desirable since both have equal efficiency to overcome dormancy of *L. ferrea* seeds.

Generally, depending on the site where seeds are collected, due to soil characteristics of each region and to autecology of the sites, seed coat hardness and treatments to break dormancy can also vary (NASCIMENTO, 2018).

The highest germination speed indexes were obtained at 23 °C and 25 °C with immersion in caustic soda and scarification by nicking, respectively, which is consistent with Brancalion et al. (2010) who affirm that Brazilian arboreal species have a maximum potential for germination next to the 25 to 30°C range. Oliveira et al. (2014) evaluated the seed germination of four native Caatinga tree species (*Sidrostylon obtusifolium* (Roem & Schult.), *Mycorrhizon urundeuva* (All.), *Amburana cearensis* (All.) and *Schinopsis brasiliensis* (Engel.), verified that the seeds of different species from the same ecological and climatic conditions present different germinative behavior as to the ideal germination temperature, but the optimum temperature range was on average from 20 to 30 °C. Regarding seeds immersed in water, the behavior was inversed as GSI decreased from 20 °C on, and then, there is an increasing trend past 30 °C. In the water immersion treatments and in the control, there were low levels of germination speed indexes of pau-ferro seeds, not being efficient.

Chemical and mechanical scarification treatments started and ended germination in a smaller number of days, compared to water immersion and control treatments. Scarification ruptures the integument through which larger amounts of water enter the seed in a shorter period of time, leading to turgidity and in consequence, hydrolytic enzymes are activated and the germination process begins.

By comparing scarification methods (nicking by pincers, rubbing against sandpaper and immersion in caustic soda), Zucareli et al. (2010) highlighted nicking as the most economically feasible and safest method; nonetheless, nonetheless, the nicking a large number of seeds might be little feasible.

5. CONCLUSIONS

Metrics of *L. ferrea* seeds is more variable as to length and width, and less variable as to thickness. Chemical scarification with immersion in caustic soda and mechanical scarification through topping with pilers provided the highest percentages and germination speeds, between temperatures 23 and 25 °C, and the shortest time to germination.

7. REFERENCES


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