ABSTRACT – Seed dormancy is a frequent phenomenon in tropical species, causing slow and non-uniform germination. To overcome this, treatments such as scarification on abrasive surface and hot water are efficient. The objective of this study was to quantify seed germination with no treatment (Experiment 1) and identify an efficient method of breaking dormancy in Schizolobium amazonicum Huber ex Ducke seeds (Experiment 2). The effects of manual scarification on electric emery, water at 80°C and 100°C and manual scarification on wood sandpaper were studied. Seeds were sown either immediately after scarification or after immersion in water for 24h in a sand and sawdust mixture. Germination and hard seed percentages and germination speed were recorded and analyzed in a completely randomized design. Analysis of germination was carried out at six, nine, 12, 15, 18, 21 and 24 days after sowing as a 4x2 factorial design and through regression analysis. Treatment means of the remaining variables were compared by the Tukey test. Seed germination with no treatment started on the 7th day after sowing and reached 90% on the 23rd day (Experiment 1). Significant interaction between treatments to overcome dormancy and time of immersion in water was observed (Experiment 2). In general, immersion in water increased the germination in most evaluations. The regression analyses were significant for all treatments with exception of the control treatment and immersion in water at 80°C. Germination speed was higher when seeds were scarified on an abrasive surface (emery and sandpaper) and, in these treatments, the germination ranged from 87% to 96%, with no hard seeds. S. amazonicum seeds coats are impermeable to water, which hinders quick and uniform germination. Scarification on electric emery followed by immediate sowing, scarification on sandpaper followed by immediate sowing and sowing after 24h were the most efficient treatments for overcoming dormancy in S. amazonicum seeds.

Index terms: tropical tree, germination, germination speed, hard seeds, dead seeds.
comparadas através do teste de Tukey. A germinação iniciou no sétimo dia e alcançou 90% aos 2310 dias após a semeadura (Experimento 1). Foi observada interação significativa entre tratamentos para superação da dormência e tempo de imersão em água após a escarificação (Experimento 2). Em geral, a imersão em água aumentou a germinação das sementes na maioria das avaliações. A análise de regressão foi significativa para todos os tratamentos, exceto na testemunha e imersão em água a 80ºC. A velocidade de germinação foi maior nas sementes escarificadas em superfície abrasiva (lixa e esmeril) e, nesses tratamentos, a germinação variou de 87 a 96%, sem a presença de sementes duras. Sementes de *S. amazonicum* apresentam tegumento impermeável à água que impede germinação rápida e uniforme. Escarificação em esmeril elétrico seguido de semeadura imediata, escarificação em lixa com semeadura imediata e semeadura após 24 horas de imersão em água, foram os tratamentos mais eficientes na superação da dormência em sementes de *S. amazonicum*.

Termos para indexação: árvore tropical, germinação, velocidade de germinação, sementes duras, sementes mortas.

**INTRODUCTION**

Seed dormancy is a common phenomenon in tropical species (Knowles and Parrotta, 1995; Bruno et al., 2001), causing slow and non-uniform germination (Cruz et al., 2001). On the other hand, seed dormancy is recognized as a survival strategy, through which plants species avoid germination in unfavorable conditions (Fenner, 1993; Schmidt, 2000).

The impermeability of seed coat to water, or hard-seededness, is a common mechanism of dormancy in Leguminosae (Rolston, 1978), but it is also found in several other families including Anacardiaceae, Bixaceae, Cannaceae, Convolvulaceae, Ebenaceae, Geraniaceae, Liliaceae, Malvaceae, Myrtaceae, Rhamaceae, Sapindaceae, Solanaceae and Zingiberaceae (Ballard, 1973; Atwater, 1980).

Several pretreatments have been proven efficient to overcome dormancy of Leguminosae seeds (Baskin and Baskin, 1998; Schmidt, 2000). The mechanical scarification on an abrasive surface has been used to overcome dormancy in many Leguminosae species. For example, in *Hymenaea intermedia* Ducke, 96% of scarified seeds germinated in 26 days, while non-scarified seeds reached similar value only after 418 days (Cruz et al., 2001). In *Bowdichia virgilioides* Kunth, 79% of germination was obtained when seeds were scarified contrasting with 21% in the control treatment (Smiderle and Souza, 2003). In *Sesbania sesban* (L.) Fawc. & Rendle, germination was 82% when scarified compared to 68% in non-scarified seeds (Veasey and Freitas, 2002). Likewise germination in *Senna occidentalis* (L.) Link was 83% and 40%, respectively (Delachiave and Pinto, 2003).

Immersion in hot water at temperatures ranging from 60 to 100ºC is an efficient method of overcoming hard-seededness (Bianchetti, 1981; Bianchetti and Ramos, 1981; Bianchetti and Ramos, 1982a; Baskin and Baskin, 1998). *Dimorphandra mollis* Benth. seeds immersed in water at 100ºC for two minutes attained a germination of 64%, while germination of untreated seeds was 12% (Hermansen et al., 2000). Germination of *Dinizia excelsa* Ducke seeds scarified with water at 80ºC for 10 min, was 62%, against 7% in the control treatment (Vastano Júnior et al., 1983). *Mimosa scabrella* Benth. seeds immersed in water at 90ºC showed germination of 79%, while in non-scarified seeds the germination was 17% (Bianchetti, 1981). Immersion time should always be observed to avoid embryo death caused by heat exposure.

*Schizolobium amazonicum* Huber ex Ducke, Leguminosae – Caesalpinioideae, locally known as “paricá” is one of the 350 tropical wood species exploited in the Brazilian Amazon (Martini et al., 1998). It is native to the States of Pará and Amazonas (Ducke, 1949), presenting rapid growth and it is important in sliced veneer production (Falesi and Santos, 1996; Rosa and Pinheiro, 2001). It has been considered a promising species to the reforestation in the Amazon region (Rosa and Pinheiro, 2001). Presently in the Paragominas region (Pará), approximately 30,000ha are planted with *S. amazonicum* and part of this area is being exploited.

According to Pereira et al. (1982) *S. amazonicum* seeds present high germination capacity without the need of any dormancy treatment. However, Maruyama and Ugamoto (1989) observed germination of only 28% at 23 days after sowing. Falesi and Santos (1996) reported germination in this species ranging from 4 to 96% after treatments to overcome dormancy. Maruyama and Ugamoto (1989) demonstrated that seeds of *S. amazonicum* have impermeable coats that delay germination. Bianchetti et al. (1997) evaluated the effect of several treatments to overcome dormancy in *S. amazonicum*
seeds and observed germination ranging from 2 to 67%.

The present study was designed to quantify *S. amazonicum* seed germination, without any treatment to overcome dormancy and to identify an efficient method of breaking seed dormancy in this species. The working hypothesis was that *S. amazonicum* seed germination is improved when seeds are subjected to treatments to overcome dormancy.

**MATERIAL AND METHODS**

Two experiments were carried out to study *Schizolobium amazonicum* Huber ex Ducke seed germination, at Embrapa Amazônia Oriental (1°28’S; 48°27’W), in Belém, Pará, Brazil, in 1995 and 2002. In both, seeds were left to germinate in a laboratory with no control over temperature and relative moisture.

**Experiment 1 – Germination of untreated seeds** – Seed moisture was determined by leaving a random sample of 20 seeds in an oven set at 105±3°C during 24h (Brasil, 1992). Germination type and seedling type were determined as Duke and Polhill (1981). Four replications of 100 seeds were sown at depth of 0.5cm in sand and sawdust (1:1), previously sterilized in hot water (100°C) for two hours and placed in plastic pots (30x22x6cm). These were irrigated every two days. Germination, evaluated on four replications of 100 seeds, was quantified daily during 2310 days and the graphic was carried out considering all germinated seeds during intervals of 154 days. A seed was considered germinated when the first pair of true leaves was visible, with no noticeable seedling abnormalities.

**Experiment 2 – Germination of scarified seeds** – In this experiment the effects of the following treatments were evaluated: manual scarification on electric emery with immediate sowing (T_s) and sowing 24h after immersion in unheated water (T_u); immersion in 80°C water for 2min with immediate sowing (T_b); the same as T_b with sowing 24h after the immersion (T_e); immersion in water at 100°C for 2 min and immediate sowing (T_e); the same as T_e with sowing 24h after immersion in water (T_b); manual scarification on sandpaper and immediate sowing (T_s); the same as T_s and sowing 24h after immersion in water at 29°C (T_o). These treatments were compared with a control treatment (T_c). Seeds from the control treatment were not scarified or immersed in water. Seeds sown 24h after treatment remained in water under ambient conditions of the laboratory. Electric emery scarification was performed at 3450rpm causing soft abrasion at the basal end of the seed for 2s. Scarification with sandpaper was carried out on 80-grit sandpaper. The spot of manual scarification is showed on Figure 1. In the hot water treatments, four parts of water were used to one part of seeds. The 24h immersion treatments used 400mL of water to 210 seeds. Initial seed moisture was determined as in Experiment 1 (Brasil, 1992), but for a sample size of 25 seeds.

Sowing, substrate and environmental conditions were the same as in Experiment 1. Germination was quantified daily during 24 days as well as the percentage of abnormal seedlings and dead seeds (Brasil, 1992), the number of days to germination onset, the mean time of germination (Edmond and Draphala, 1958), and the germination speed (Maguire, 1962).

Data were subjected to the Brown & Forsyth homogeneity of variance test (Statsoft, 1995) and an arcsine $\sqrt{(x + 0.5)/100}$ transformation was performed on mean germination time and percentage of abnormal seedlings. These transformed data were analysed in a completely randomized design with four replicates of 50 seeds each. The statistical analysis of germination percentage was carried out at 6, 9, 12, 15, 18, 21 and 24 days after sowing as a 4x2 factorial design (four types of scarification: electric emery, sandpaper, immersion in water in 80 and 100°C water; and

![FIGURE 1. Schizolobium amazonicum seed. The arrow indicates the spot of manual scarification used in T_c, Belém, 2002.](image-url)
two sowing times: immediate and 24h after the scarification treatments). Treatment means were compared by the Tukey test using Statistica (Statsoft, 1995). Data were back-transformed for presentation purposes.

RESULTS AND DISCUSSION

Experiment 1 – Germination of untreated seeds
– Seed moisture content was 5.3% (±0.2), germination was epigeal and seedlings were phanerocotylar. Germination was non-uniform and slow, beginning 7 days after sowing. The whole germination process took 2310 days, when germination achieved 90% (Figure 2). Germination started with primary root development, achieving 5 to 7cm, when the hypocotyls began their development on the soil surface. Most seeds released their coats before hypocotyl development, though a small proportion of seeds kept their coat attached to the cotyledon, thus restraining seedling development. The germination curve was sigmoid (Tipton, 1984), with an initial phase of slow germination followed by a rapid increase in germination and a final phase presenting no significant germination increase, a pattern similar to the one reported for Micropholis cf. venulosa Mart. & Eichler (Cruz et al., 2003). Long germination periods have been reported in Caesalpinioideae (Cruz et al., 2001).

Experiment 2 – Germination of scarified seeds
– Seed moisture content at the start of the experiment was 11.3% (±2.2). There was interaction between scarification and post-scarification treatments. In general, the germination of seeds immersed in water for 24h was higher than the germination of seeds sown immediately after scarification (Table 1). Post-scarification treatment also affected the onset of germination, which was higher than 40% on manually scarified seeds on the 6th day, and nil on those sown immediately after scarification (Table 1). Twelve days after sowing seeds scarified on emery and sandpaper, but not immersed in water, there was similar or higher germination than in seeds scarified and immersed in water.

The regression analysis was significant for all treatments, for the control (T1) and immersion in water at 80ºC followed by immediate sowing (T4), which showed 1% and 2% germination, respectively. These two treatments presented 97.5 and 96% hard seeds, respectively.

Seeds scarified on emery and sown immediately (T2) showed significant increase in germination until the 15th day (Figure 3A), whereas germination of seeds scarified and immersed in water occurred until the 9th day (Figure 3B).


<table>
<thead>
<tr>
<th>Treatments</th>
<th>IS 6</th>
<th>12</th>
<th>18</th>
<th>21</th>
<th>24</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual scarification on emery</td>
<td>0.0 bA</td>
<td>43.5 aA</td>
<td>30.0 bB</td>
<td>84.0 aA</td>
<td>89.5 aA</td>
</tr>
<tr>
<td>Immersion in water at 80ºC</td>
<td>0.0 bA</td>
<td>1.5 aB</td>
<td>0.5 bC</td>
<td>8.0 aB</td>
<td>1.0 bC</td>
</tr>
<tr>
<td>Immersion in water at 100ºC</td>
<td>0.0 bA</td>
<td>3.5 aB</td>
<td>5.5 aC</td>
<td>15.5 aB</td>
<td>11.5 bB</td>
</tr>
<tr>
<td>Manual scarification on sandpaper</td>
<td>0.0 bA</td>
<td>40.5 aA</td>
<td>61.0 bA</td>
<td>89.0 aA</td>
<td>94.5 aA</td>
</tr>
</tbody>
</table>

TABLE 1 (continued).

<table>
<thead>
<tr>
<th>Treatments</th>
<th>15</th>
<th>18</th>
<th>21</th>
<th>24</th>
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</thead>
<tbody>
<tr>
<td>Manual scarification on emery</td>
<td>95.5 aA</td>
<td>86.5 bA</td>
<td>95.5 aA</td>
<td>86.5 bA</td>
</tr>
<tr>
<td>Immersion in water at 80ºC</td>
<td>1.0 bC</td>
<td>27.0 aC</td>
<td>2.0 bC</td>
<td>28.5 aC</td>
</tr>
<tr>
<td>Immersion in water at 100ºC</td>
<td>16.0 bB</td>
<td>70.5 aB</td>
<td>20.0 bB</td>
<td>79.0 aB</td>
</tr>
<tr>
<td>Manual scarification on sandpaper</td>
<td>94.5 aA</td>
<td>91.5 aA</td>
<td>94.5 aA</td>
<td>91.5 aA</td>
</tr>
</tbody>
</table>

Same capital letters in the column and lower case letters in the line, do no differ statistically at 5% (Tukey).
Scarification with sandpaper combined with immersion in water has been shown to increase germination of *S. parahyba* (Vell.) S.F. Blake (Lorenzi, 1992). Scarification on an abrasive surface was also an efficient method to overcome dormancy in *H. intermedia* (Cruz et al., 2001). In *Peltophorum dubium* (Spreng.) Taub. scarification on an abrasive surface was an efficient method of promoting seed germination, but germination was reduced when seeds were scarified for longer than 8s (Bianchetti and Ramos, 1982b). Seeds scarified on sandpaper showed a similar pattern of germination to seeds scarified on emery (Figure 4A and 4B).

Scarification with water at 80ºC and immersed in water for 24h (T5) presented the lowest final germination, reaching only 31.5% (Figure 5). Immersion in water at 100ºC was not so effective at promoting seed germination (Figure 6A), but when followed by a 24-hour water immersion period, germination reached 82.5% (Figure 6B).

The use of hot water to overcome dormancy in seeds with impermeable coats has been suggested for some forest species (Fowler and Bianchetti, 2000), but the efficiency of such a treatment depends on the species, water temperature and immersion time during scarification (Schmidt, 2000). The use of water at 80ºC followed by immersion in water for 18h has been recommended as an efficient treatment to scarify seeds of *Acacia longifolia* (Andrews) Willd. (Medeiros and Zanon, 1999) and *M. bimucronata* (DC.) Kuntze (Fowler and Carpanezzi, 1998). Higher water temperatures (95-100ºC) are an effective method for some species. In *S. parahyba*, seed...
Scarification in 95°C water during 4-10 min promoted germination, which ranged from 84.1 to 88.3% (Bianchetti and Ramos, 1981), while in *Acacia mearnsii* De Wild. 83.3% germination occurred when seeds were scarified in water at 100°C for 6 min (Bianchetti and Ramos, 1982a). However, for *M. bimucronata* the use of temperature above 80°C caused embryo death in some seeds (Fowler and Carpanezzi, 1998).

The Table 2 shows days to germination onset, mean germination time, germination speed index and germination, hard seed, dead seed, and abnormal seedling percentages. There were statistical differences among all treatments except for abnormal seedlings. In general, days to germination onset took 6-8 days after sowing corroborating the description by Pereira et al. (1982). However, when seeds were scarified in water at 80°C and sown immediately (T4), germination started on the 13th day. This treatment appears to delay the beginning of the germination process.

Mean germination time duration ranged from 6.6 days, when scarification was performed by sandpaper plus immersion water, to 14.8 days, when seeds were scarified in water at 80°C followed immediate sowing. Seeds scarified on emery (T3) and sandpaper (T9), followed by immersion in water, required fewer days to reach 50% of germination.

The germination speed index was higher for seeds scarified on emery (T3) and sandpaper (T9), with immersion in water, 6.56 and 6.78, respectively, showing the efficiency of these treatments to speed up germination in *S. amazonicum* seeds.

The hard seed percentage was high in all treatments, except for the immersion in water 100°C with sowing after 24 h (T7) with 13% of hard seeds. In general, the dead seed percentage was low, except when seeds were scarified on emery and immersed in water (12.0%).

Veasey and Freitas (2002) observed that treatments not only did not promote germination of *Sesbania sesban* (L.) Fawc. & Rendle, *S. rostrata* Bremek. & Oberm. and *S. virgata* (Cav.) Pers. but damaged a high number of seeds, but in *S. parahyba*, Bianchetti and Ramos (1981) did not detect any significant increase in dead seed percentage in treatments that did not promote seed germination.

Treatments tested in the present study did not increase the percentage of abnormal seedlings due to the small differences among them, typically, ranging from 0.7 to 3%, and non-significant.
TABLE 2. Number of days to germination onset (DGO), mean germination time (MGT), germination speed index (GSI), and germination (G), hard seeds (HD), dead seeds (DS) and abnormal seedlings (AS), in *Schizolobium amazonicum*.

<table>
<thead>
<tr>
<th>Treatments</th>
<th>DGO</th>
<th>MGT</th>
<th>GSI %</th>
<th>G</th>
<th>HD %</th>
<th>DS %</th>
<th>AS %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>8.0 ab</td>
<td>9.5 abc</td>
<td>0.12 e</td>
<td>1.0 f</td>
<td>97.3 c</td>
<td>1.0 a</td>
<td>0.7 a</td>
</tr>
<tr>
<td>Manual scarification on electric emery with immediate sowing</td>
<td>8.0 ab</td>
<td>10.3 bc</td>
<td>4.75 b</td>
<td>96.0 a</td>
<td>*</td>
<td>1.0 a</td>
<td>3.0 a</td>
</tr>
<tr>
<td>Manual scarification on electric emery and sowing 24h after</td>
<td>6.0 a</td>
<td>6.7 a</td>
<td>6.56 a</td>
<td>86.5 bc</td>
<td>*</td>
<td>12.0 d</td>
<td>1.5 a</td>
</tr>
<tr>
<td>Immersion in water at 80°C with immediate sowing</td>
<td>13.3 b</td>
<td>14.8 c</td>
<td>0.10 e</td>
<td>2.0 f</td>
<td>96.0 c</td>
<td>2.0 ab</td>
<td>*</td>
</tr>
<tr>
<td>Immersion in water at 80°C with sowing 24 hours after</td>
<td>6.0 a</td>
<td>12.8 bc</td>
<td>1.39 d</td>
<td>31.5 e</td>
<td>65.5 b</td>
<td>0.5 a</td>
<td>2.5 a</td>
</tr>
<tr>
<td>Immersion in water at 100°C with immediate sowing</td>
<td>8.2 ab</td>
<td>12.7 bc</td>
<td>0.88 de</td>
<td>20.0 e</td>
<td>71.0 b</td>
<td>6.5 bc</td>
<td>2.5 a</td>
</tr>
<tr>
<td>Immersion in water at 100°C with sowing 24 hours after</td>
<td>6.0 a</td>
<td>13.5 c</td>
<td>3.28 c</td>
<td>82.5 c</td>
<td>13.0 a</td>
<td>1.5 ab</td>
<td>2.5 a</td>
</tr>
<tr>
<td>Manual scarification on sandpaper with immediate sowing</td>
<td>7.5 a</td>
<td>9.2 ab</td>
<td>5.20 b</td>
<td>94.5 a</td>
<td>*</td>
<td>2.5 abc</td>
<td>3.0 a</td>
</tr>
<tr>
<td>Manual scarification on sandpaper with sowing 24 hours after</td>
<td>6.0 a</td>
<td>6.6 a</td>
<td>6.78 a</td>
<td>91.5 ab</td>
<td>*</td>
<td>7.5 cd</td>
<td>1.0 a</td>
</tr>
</tbody>
</table>

Treatment means sharing the same letter within columns are not significantly different (P>0.05).
* Values omitted from variance analysis for being zero

CONCLUSION

Scarification either by electric emery or sandpaper, followed by immediate sowing and sowing after 24h, is the most efficient treatment to overcome dormancy in *S. amazonicum* seeds.

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