A Preliminary Study on the Germination of *Eurycoma longifolia* Jack (Tongkat Ali) Seeds

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**Keywords:** In vitro germination, jiffy, endocarp, MS culture medium, seedlings

**ABSTRACT**


**INTRODUCTION**

Eurycoma longifolia fruits are borne in large bunches consisting of 200-300 fruits in each bunch. Even though trees produce abundant fruits during the peak fructifying season in September, yet the number of seedlings found growing in the forest floor is very low. Our results indicated that the seeds with hard endocarp sown in a 1:1 soil and sand mixture only start to germinate 43 days after sowing and continue to germinate over a period of 99 days. The soil and sand mixture (1:1) is equivalent to that of the forest sandy soil combination that was optimum for germination of *Eurycoma longifolia* seeds. The seeds with endocarp intact that are sown in jiffy germinated within 35-85 days. However, when the endocarp of seeds were removed, the seeds germinated within two weeks via in vitro culture using basic MS medium. The ripe seeds germinated better when sown in the 1:1 soil and sand mixture than in jiffy pellets. But the unripe seeds with the endocarp removed seemed to germinate faster (CVg = 0.2053) when cultured in vitro in basic MS medium. All the seedlings were found to have the same growth pattern in terms of seedling height, number of leaves produced, and the stem diameter irrespective of germination methods over a period of 120 days.
stigma and always with large but sterile stamens. The male flowers produce five stamens with a sterile pistil. The ovoid shape fruits are borne in a large dangling axillary bunch. Its peak flowering season is from June to July and with peak fruiting in September (Corner 1988).

Even though trees produce abundant fruits and seeds during each fruiting season, the number of seedlings found growing around the adult trees is low. Until now, the germination behavior of the seeds has not been studied. Therefore we wished to determine the general morphology of *E. longifolia* fruits and seeds, and how its structures influenced seed germination behavior. Their capacity to germinate under laboratory conditions and the growth pattern of its seedlings were studied. The possibility of using *in vitro* seed germination as an alternative method for enhancing the seed germination of *E. longifolia* Jack was also investigated.

**MATERIALS AND METHODS**

**Fruit and Seed Morphology**

*E. longifolia* fruits were collected from a secondary forest in Penang, Malaysia at three different sites namely Bayan Lepas, Teluk Bahang and Teluk Kumbar. A study was done on the external morphology and cross-section of the fruit and seed.

**Germination Test**

a. *Effects of Germination Methods and Seed Maturity on Seed Germination*

The fruits were removed from each bunch which consisted of approximately 200 to 300 fruits and grouped as young, unripe, green seeds and matured, ripe, red or dark-red seeds. Twenty seeds were taken randomly from each bunch and from each grouping to study the effect of each of the germination methods on seed germination.

The three germination methods were:

1. The seeds were sown approximately one cm deep in a 1:1 soil and sand mixture.
2. The seeds were sown in jiffy pellets (Jiffy Products Ltd., Norway). These jiffy pellets were made up of peat soil and each seed was placed in each pellet.
3. The seeds were germinated via the *in vitro* technique. For this technique, the epicarp and mesocarp of the fruits were removed. The seeds were washed with detergent, then rinsed in running tap water for 30 minutes. The seeds were then immersed in a 250 ml conical flask containing 20% (v/v) Clorox® solution which contained 5.25% sodium hypochlorite and three drops of tween-20 for 20 minutes, with continuous agitation. This was followed by rinsing three times with sterile distilled water. Surface sterilization of these seeds was repeated with 15% Clorox® solution for 15 minutes and again rinsed three times with sterile distilled water. The sterilized seeds were then placed on the surface of 15 ml Murashige and Skoog basic medium (MS) (Murashige & Skoog 1962) contained in 25x150mm culture tubes capped with autoclavable plastic caps (Jenaerglas, Rasotherm, Germany).

Twenty seeds were used for each germination method and the study was repeated three times. Percentage of germination for each method was recorded over a 120-day period. Germination was determined by the emergence of the radical and epicotyl on the germination medium surface. The effects of germination methods and maturity of seeds and their interactions on percentage of germination were analyzed using analysis of variance (ANOVA).

b. *Influence of Endocarp (testa) on Seed Germination*

The endocarp was removed after the seeds were surface sterilized twice as mentioned above. The seeds with the endocarp removed were again surface sterilized with 5% Clorox® solution for 10 minutes, rinsed three times with sterile distilled water and placed in 25 x 150 mm culture tubes containing 15 ml MS basic medium. Twenty seeds were used for each trial and the experiment was repeated three times. Percentage of germination was recorded over a period of 120 days.

c. *Determination of the Coefficient of Velocity of Germination*

The coefficient of velocity of germination (CVG) was computed based on Hartman and Kaster (1968):

\[
CVG = \frac{\text{Total number of germination}}{A_1T_1 + A_2T_2 + \ldots + A_nT_n}
\]

where \( A = \) number of fresh germination recorded at each day interval

\( T = \) number of days from sowing.
The effects of germination methods and the maturity of seeds, and their interactions on the coefficient of velocity of germination were computed using ANOVA.

The Growth Pattern of Seedlings

Two weeks after germination, the seedlings were transferred to 15 x 23 cm polybags containing a 1:1:2 mixture of organic manure: top soil: sand. These seedlings in polybags were placed in a plant house at a temperature of between 28-30°C. The height of the seedlings was recorded every week starting from the emergence of the epicotyl, while the stem diameter was recorded every month. Plant height was taken as the distance from the tip of the shoot apex to the first node on the plant. Stem diameter was measured with a pair of calipers (Kern, Germany) at the fifth node of the stem. The number of leaves produced over a fortnight period was estimated by counting the last tagged leaf of the previous recording to the most recently produced leaf.

RESULTS AND DISCUSSION

The fruits of *E. longifolia* were borne in a large dangling axillary bunch. The bunches of fruits that were collected consisted of 200-300 seeds per bunch. The fruits were produced in groups of 1-5 on the bunches (Fig. 1).

The fruits were yellow to light green when young, and became red to blackish-red when ripe. The ripe and unripe fruits were distributed randomly in the same bunch. The variation in fruit maturity within the bunch serves to minimize the competition for substrate for successful seed germination at the forest floor which is often overcrowded with secondary growth. As stated by Villier (1972), some seeds appear to be involved in controlling germination by restricting it to periods and conditions most favorable for seedling growth.

The fleshy drupe fruits were ovoid in shape and about 10-20 mm long and 5-12 mm broad. It consisted of a thin shining epicarp, fleshy mesocarp, hard and stony endocarp. The seed consisted of two large expanded cotyledons and a chlorophyllous capitulate embryo. With the endocarp removed, the seed could be seen to be covered with a thin papery covering, which could be easily removed from the inner surface of the endocarp (Fig. 2).

Seeds with the endocarp intact sown in the 1:1 soil and sand mixture started to germinate from the 43 days and continued to germinate until 99 days after sowing (Table 1). The inhibition and delay in germination could be due to a high degree of impermeability of the endocarp to water or oxygen or to both. This phenomenon was similar to that of winged bean seeds (*Psophocarpus tetragonolobus* L.) which showed very low percentage of germination due to impermeability of the seed coat to water (Rudrapal et al. 1992). Rolston (1978) also reported that impermeability of hard seed coats was typical of legume seeds.

Seeds sown in jiffy pellets germinated earlier and within a shorter period of time (35-85 days) as compared to those sown in 1:1 soil and sand mixture (43-99 days). This was because jiffy pellets consisted mainly of peat soil and were able to retain higher moisture content compared to the 1:1 soil and sand mixture, hence allowing more water absorption by the seeds. None of the seeds with endocarp intact germinated when cultured in vitro using the MS culture medium (Table 1). Blackening occurred on the non-germinated seeds with the endocarp intact.

<table>
<thead>
<tr>
<th>Germination methods for <em>E. longifolia</em> seeds</th>
<th>Duration of germination (days)</th>
<th>Ripe seeds</th>
<th>Unripe seeds</th>
</tr>
</thead>
<tbody>
<tr>
<td>sown in soil and sand mixture</td>
<td>45-99</td>
<td>43-93</td>
<td></td>
</tr>
<tr>
<td>sown in jiffy pellets</td>
<td>37-85</td>
<td>35-70</td>
<td></td>
</tr>
<tr>
<td>In vitro culture</td>
<td>No germination</td>
<td>No germination</td>
<td></td>
</tr>
</tbody>
</table>

* seeds used for germination are with endocarp intact
Fig. 1-2. E. longifolia fruits and seeds
1. E. longifolia fruits and groups of 1-5 in part of the bunches. 2. The seed with two large cotyledons and a chlophyllous capitulate embryo covered with a thin papery covering which was easily removed from the stony endocarp.

intact and also in the MS culture medium (Fig. 3). However, the seeds without endocarp did not release any black exudates (Fig. 4). Marbach and Mayer (1974) reported that black exudates released were mainly phenolic compounds and could contribute to the impermeability of seed coats to water, hence preventing germination of seeds.

Ripe E. longifolia seeds sown in a 1:1 soil and sand mixture or jiffy pellets germinated better than the unripe seeds. Ripe seeds sown in the soil and sand mixture (1:1) reached 58% germination over a 120-day period while only 46% of the unripe seeds sown in the same medium germinated at the same duration. Forty six percent of the ripe seeds sown in jiffy germinated over a 120-day period. Only 29% of the unripe seeds sown in jiffy germinated over the same duration. All the ripe and unripe seeds cultured in vitro did not germinate (Fig. 5). Rudrapal et al. (1992) proposed that the delayed germination of immature seeds was due to lower free
GERMINATION OF EURYCOMA LONGIFOLIA JACK (TONGKAT ALI) SEEDS

Fig. 3-4. Germination of E. longifolia seeds on MS culture medium.
3. Seed with endocarp intact released black exudates into the MS culture medium.
4. Seed without endocarp on MS culture medium clear of black exudates.

gibberellin content in embryo-cotyledon at the immature stage resulting in slower rate of water imbibition. Analysis of variance indicated that the different germination methods significantly affected (p=0.01) the percentage of germination of E. longifolia seeds. However, there were no significant differences in percentage of germination between the ripe and unripe seeds, and there was also no interaction between the ripeness of seeds and the different types of germination methods (Table 2).

The experimental results thus indicated that the stony endocarp did contribute to the germination process of these seeds. This was further supported by our findings on seeds with the endocarp removed starting to germinate 14 days after in-vitro cultured on MS medium. They continued to germinate until 64 days and none of the seeds with endocarp intact germinated on the same MS medium (Table 3). Edwards (1968) reported that most of the inhibition compounds that inhibited seed germination were usually located in the fruit wall or seed coat. Hence, this explained that E. longifolia seeds without endocarp would be able to germinate earlier.

With in vitro germination, the unripe seeds without endocarp germinated faster than the ripe seeds. The unripe seeds without endocarp showed maximum 53% germination while only 30% of the ripe seeds without endocarp germinated in the MS culture medium (Fig 6). This was further supported by the CVG results (Table 4) which indicated that unripe seeds with the endocarp removed, sown via the in-vitro method was the fastest to germinate, (CVG = 0.2053)

Fig. 5. Germination response of the ripe and unripe seeds of E. longifolia to different germination methods
TABLE 2
Analysis of variance for percentage of germination and coefficient of velocity of germination (CVG)

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>df</th>
<th>MS</th>
<th>CVG&lt;sup&gt;x&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment</td>
<td>5</td>
<td>1480.68</td>
<td>0.0159**</td>
</tr>
<tr>
<td>Factor A (method)</td>
<td>2</td>
<td>3489.78</td>
<td>0.0363*</td>
</tr>
<tr>
<td>Factor B (seed type)</td>
<td>1</td>
<td>255.08 ns</td>
<td>0.0015 ns</td>
</tr>
<tr>
<td>A x B</td>
<td>2</td>
<td>84.38 ns</td>
<td>0.0028 ns</td>
</tr>
<tr>
<td>Error</td>
<td>12</td>
<td>151.64</td>
<td>0.0013</td>
</tr>
</tbody>
</table>

<sup>y</sup> analysis based on arc sine value.
<sup>x</sup> analysis considers CVG data for in-vitro method using seeds without endocarp.
** significant at p=0.01.
ns not significant.

TABLE 3
Effect of seed endocarp on the duration of *E. longifolia* seed germination using in vitro technique

<table>
<thead>
<tr>
<th>Condition of seeds</th>
<th>Duration of germination (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ripe seeds</td>
<td>Unripe seeds</td>
</tr>
<tr>
<td>Seeds with endocarp intact</td>
<td>No germination</td>
</tr>
<tr>
<td>Seeds with endocarp removed</td>
<td>18-50</td>
</tr>
</tbody>
</table>

followed by ripe seeds without endocarp using the same *in vitro* method (CVG = 0.1377). The ripe seeds with endocarp intact sown in the 1:1 soil and sand mixture was the slowest to germinate (CVG = 0.0252).

The ANOVA presented in Table 2 also showed that the different germination methods significantly affected (p=0.01) the coefficient of velocity of germination but there was no significant difference between the maturity of seeds and their interaction with the germination methods.

All the seeds sown in the 1:1 soil and sand mixture and in jiffy pellets showed the same
GERMINATION OF *EURYCOMA LONGIFOLIA* JACK (TONGKAT ALI) SEEDS

### TABLE 4

Mean coefficient of velocity of germination (CVG) for *E. longifolia* seeds germinated with different germination methods

<table>
<thead>
<tr>
<th>Germination methods</th>
<th>Mean CVG w</th>
</tr>
</thead>
<tbody>
<tr>
<td>sown in soil and sand mixture (1:1)</td>
<td></td>
</tr>
<tr>
<td>ripe seeds</td>
<td>0.0252 a</td>
</tr>
<tr>
<td>unripe seeds</td>
<td>0.0362 a</td>
</tr>
<tr>
<td>sown in jiffy pellets</td>
<td>0.0424 a</td>
</tr>
<tr>
<td>ripe seeds</td>
<td>0.0456 a</td>
</tr>
<tr>
<td>unripe seeds</td>
<td>0.2053 c</td>
</tr>
<tr>
<td>In vitro</td>
<td>0.1377 b</td>
</tr>
</tbody>
</table>

* Means separation by Duncan’s multiple range test, p=0.05. Values followed by the same letter are not significantly different.

The number of leaves produced by the seedlings seemed to follow the same growth pattern as that of seedling height irrespective of germination methods. When seedling growth was slower, the number of leaves produced was reduced (Fig. 7 & Fig. 8). Since all the seedlings were planted in polybags and placed in the plant house, the seedlings were exposed to similar environmental condition, hence they have the same growth pattern irrespective of type of germination medium. Chan (1984) also noted that different varieties of *Carica papaya* L. grown in Malaysia had the same growth pattern because the constant environmental conditions encouraged continuous growth and development.

Our study indicated that *E. longifolia* seeds had a low germination rate. This was due to the impermeability of hard stony endocarps of its

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**Fig. 7.** The growth pattern of *E. longifolia* seedlings, derived from seeds germinated in soil and sand mixture and in jiffy pellets, in term of height and number of leaves produced.
seeds to water. The low germination rate of *E. longifolia* seeds could also be the reason why there is poor distribution of its seedlings in the forest floor. Since the seeds germinated faster when cultured in-vitro and there were no differences in the growth pattern of the seedlings, the in-vitro method of germination could be an alternative method for producing faster and more *E. longifolia* seedlings.

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Effect of Methanol and Ethanol Pre-Treatments on Seed Germination and Seedling Development of *Dichrostachys cinerea* (L.) Wight and Arn. (Fabaceae)

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**Keywords:** Seed germination, methanol and ethanol pre-treatments, seedling

**ABSTRACT**

To improve germination as well as achieve high nursery recovery of good quality seedlings, seeds of *Dichrostachys cinerea* were subjected to alcohol scarification for different exposure times. Seeds soaked in methanol for 10 min achieved the highest percentage germination (72%) and germination energy (65%). Similarly, 5 min methanol pre-treatment gave good results. Seedlings resulting from these treatments were mostly of the high vigour class when compared to other alcohol pre-treatments. The 2 min methanol and ethanol pre-treatments gave low germination in percentages and germination energies, and the resultant seedlings were mostly in the low vigour category.

**INTRODUCTION**

*Dichrostachys cinerea* (L) Wight and Arn. Sub-sp *africana* Brenan and Brummitt (Fabaceae), belongs to a small genus of the sub-family Mimosoideae widespread in the tropical savanna of Africa. It is the only known member of the genus in Nigeria. The plant commonly grows as a tree or sometimes as a shrub, often with low branches and dense canopy of branchlets (Keay 1989). This indigenous multipurpose, but under-exploited tree species is important for its fodder and fuel uses, as well as its sand-stabilization ability. Seeds of *D. cinerea* have hard seed coats, which are impermeable to water and gases thereby inhibiting germination.

The interaction between pre-treatments and the degree of hard seededness varies between seeds of the same or different species, and within the same seedlot (Gill *et al.* 1982). The difference in response to dormancy breaking pre-treatments by the seed depends on environmental conditions, the degree of maturation of the seeds and the duration of storage (Gunn 1990). Various methods have been employed in terminating dormancy in seeds with hard seed coats. Alcohol pre-treatments have been reported to be effective in the breakage of dormancy and improvement of germination in seeds, particularly those of the Fabaceae (Etejere *et al.* 1982; Mayer and Poljakoff-Mayber 1989; Gill *et al.* 1990; Idu 1995). However, not much has been reported on the effect of such pre-treatments on germination energy of the seeds and development of the resultant seedlings.

The present study evaluates the effect of various alcohol pre-treatments on germination and seedling vigour of *D. cinerea*. 

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Bagi memperbaiki percambahan sebaik mungkin untuk mencapai pemerolehan pembenihan anak benih yang berkualiti, biji benih *Dichrostachys cinerea* didedahkan kepada penggemburan tanah dengan alkohol dalam masa pendedahan yang berlainan. Biji benih direndam dalam metanol selama 10 minit mencapai peratusan percambahan (72%) dan tenaga percambahan (65%) yang paling tinggi. Begitu juga, prarawatan metanol 5 minit memberi keputusan-keputusan yang baik. Hasil keputusan anak benih daripada rawatan-rawatan tersebut adalah kebanyakannya kelas tenaga tinggi apabila dibandingkan dengan prarawatan alkohol yang lain. Prarawatan metanol dan etanol 2 minit memberi percambahan yang rendah dalam peratusan dan tenaga percambahanm dan anak-anak benih tersebut kebanyakannya adalah dalam kategori tenaga rendah.

To improve germination as well as achieve high nursery recovery of good quality seedlings, seeds of *Dichrostachys cinerea* were subjected to alcohol scarification for different exposure times. Seeds soaked in methanol for 10 min achieved the highest percentage germination (72%) and germination energy (65%). Similarly, 5 min methanol pre-treatment gave good results. Seedlings resulting from these treatments were mostly of the high vigour class when compared to other alcohol pre-treatments. The 2 min methanol and ethanol pre-treatments gave low germination in percentages and germination energies, and the resultant seedlings were mostly in the low vigour category.
MATERIALS AND METHODS

Seeds for the study were collected from Gieri, Adamawa State, Nigeria (12°, 20'E, 90°, 14'N). Seeds were removed from ripe pods and stored at constant temperature 28 ± 3°C throughout the experimental period.

Six hundred seeds were divided into two sub-samples of 300 seeds for the methanol and ethanol treatments. Each sub-sample was further divided into one hundred seeds for 3-treatment groups (2, 5 and 10 min) exposure period with 5 replicates of 20 seeds each. 100 seeds of 5 replicates served as controls for each pretreatment.

Cleansed seeds were subjected to 70% concentrated methanol and ethanol pre-treatments for 2, 5 and 10 min, respectively and continually stirred. After the designated exposure period, the alcohol was drained and the seeds rinsed thoroughly (five times) in several changes of distilled water before being put up for germination.

Germination technique was as outlined by Dasgupta et al. (1976) and Marunda (1990). Treated and untreated (control) seeds were placed on moist filter paper in Petri-dishes under continuous fluorescent light at 10cm above bench level at room temperature. Three (3) mm radicle emergence served as criterion for germination (Idu & Omonhinmin 1999).

Germination was recorded daily for 30 days. After germination and following a randomized design, 10 seedlings were transplanted into segmented wooded trays (240 x 120 x 30cm) filled with sterile soil (pH=6.90) at a planting depth of 2 cm.

Watering was done daily with Harris culture medium. Seedling height measurements were done at 3-day intervals. Seedlings were grouped into vigour categories based on germination and seedling height data. The vigour index of germination was estimated by calculating the daily germination energy percentage maximum (Seward 1980). Seedling height measurement was stopped after 10 weeks.

Analysis of variance for a complete randomized design was carried out on the height data for seedlings grown from ethanol and methanol pre-treated seeds and to test for difference in treatment effect. A comparison of treatments' effect on mean height was carried out using the least significance difference (LSD).

### Table 1

<table>
<thead>
<tr>
<th>Pre-treatment</th>
<th>% Germ</th>
<th>Germ En.</th>
<th>HV</th>
<th>LV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methanol 2 min</td>
<td>31</td>
<td>15</td>
<td>22</td>
<td>9</td>
</tr>
<tr>
<td>Methanol 5 min</td>
<td>56</td>
<td>35</td>
<td>32</td>
<td>24</td>
</tr>
<tr>
<td>Methanol 10 min</td>
<td>72</td>
<td>65</td>
<td>48</td>
<td>24</td>
</tr>
<tr>
<td>Ethanol 2 min</td>
<td>22</td>
<td>9</td>
<td>9</td>
<td>13</td>
</tr>
<tr>
<td>Ethanol 5 min</td>
<td>16</td>
<td>6</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Ethanol 10 min</td>
<td>33</td>
<td>15</td>
<td>15</td>
<td>18</td>
</tr>
<tr>
<td>Control</td>
<td>12</td>
<td>5</td>
<td>6</td>
<td>8</td>
</tr>
</tbody>
</table>

(Data are average of five replicates)

**% Germ** - Percentage germination

Germ. En-P - Germination Energy

Experimental mean height = 6.45 cm after 30 days

NG - Non-germinated seeds

HV - High Vigour above mean

LV - Low Vigour below mean

### Table 2

<table>
<thead>
<tr>
<th>Pre-treatment</th>
<th>Ranked mean</th>
<th>LSD(H)+Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>5.80 a</td>
<td>7.00</td>
</tr>
<tr>
<td>Ethanol 2 min</td>
<td>7.40 b</td>
<td>8.60</td>
</tr>
<tr>
<td>Ethanol 5 min</td>
<td>7.60 bc</td>
<td>8.80</td>
</tr>
<tr>
<td>Ethanol 10 min</td>
<td>8.60 c</td>
<td>9.80</td>
</tr>
<tr>
<td>Methanol 5 min</td>
<td>8.87 d</td>
<td></td>
</tr>
<tr>
<td>Methanol 2 min</td>
<td>8.90 d</td>
<td></td>
</tr>
<tr>
<td>Methanol 10 min</td>
<td>9.10 d</td>
<td></td>
</tr>
</tbody>
</table>

** Mean followed by the same letter are not significantly different at 5% (LSD)

RESULTS AND DISCUSSION

Table 1 shows the percentage germination achieved after 30 days, germination energies after 8 days and vigour categories after 30 days. The methanol treatment for 10 min achieved the highest germination percentage of 72% and germination energy of 65%. Five min soaking of seeds in methanol gave 56% germination and germination energy of 35%. The majority of the seeds germinated within the first 9 days. Two min of methanol treatment and 2, 5 and 10 min ethanol pre-treatments gave lower percentage germination and energies. The vigour categories show the methanol pre-treated seeds (2, 5 and 10 min) to be in a higher vigour class than the ethanol pre-treated seeds.
Alcohol stimulates germination in hard coat seeds, particularly those of the Fabaceae, by softening the waxy seed coat, thereby allowing the inflow of water, gaseous exchange and unrestricted expansion of embryonic parts (Mayer and Poljakoff-Mayber 1989). The pre-treatments with high germination energies (5 and 10 min methanol) can be applied in nursery settings to produce uniform planting stock to ensure maximum nursery recovery of high quality seedlings. Poor germination percentage and energies recorded for the ethanol and 2 min methanol pre-treatment may be due to reduced severity of the treatment, which did not render the seed coat soft and permeable to water (Marunda 1990; Idu 1995). Such treatments produce seedlings of variable height in the nursery, resulting in poor recovery of good quality planting stock.

The control treatment produced seedlings with the lowest mean height, which suggests poor germination energy from the start of the experiment. This is a further indication that the seeds of *D. cinerea* require pre-treatments before better seedling performance can be achieved.

In conclusion, it is evident that methanol and ethanol pre-treatments had different effects on germination and vigour of *D. cinerea* seeds. The methanol at 10 minutes treatment gave better results and will be an ideal pre-treatment for effective germination of high quality seedlings of *D. cinerea*.

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