THE EFFECT OF TEMPERATURE AND SCARIFICATION METHOD ON GORSE (ULEX EUROPAEUS L.) SEED GERMINATION

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ABSTRACT

As part of a study of gorse (Ulex europaeus L.) biocontrol, the scarification requirements to maximise germination of freshly harvested seed were investigated. Both time of immersion in concentrated sulphuric acid (36N) and incubation temperature were critical. Optimum incubation temperature was 15°C (mean germination 65%). The sulphuric acid treatments that gave the highest percentage of germinated seeds were 180 and 210 min immersion with incubation at 15°C, giving a mean germination of 81%. The mean control germination, over all incubation temperatures, was 10%. Hot water immersion did not significantly increase germination when compared with the unscarified control treatment. Mechanical scarification gave a mean germination over all temperatures of 35%; the highest germination was again at 15°C (48%).

Keywords: gorse, germination temperature, acid scarification, mechanical scarification.

INTRODUCTION

Gorse (Ulex europaeus L.) is a prolific seeder and seed viability is at least 40 years (Zabkiewicz 1976; Zabkiewicz & Gaskin 1978). There is anecdotal evidence that hard-coated seed may lie ungerminated in the soil for 75 years (W.J. Davis, pers. comm.). The ability of gorse seed to remain viable for many years, just requiring to be brought nearer the surface for germination to occur, means that the weed seedbank can rapidly re-establish gorse infestations (Ivens 1982).

Scarification with concentrated sulphuric acid (H₂SO₄) (36N) may increase gorse seed germination. In Russell lupin (Lupinus polyphyllus x L. arboreus) acid scarification for 30 or 45 min gave more than 75% germination, while unscarified control seed had 20% germination (Tesfaye 1989). With broom (Cytisus scoparius) highest germination was obtained from seed that had been immersed in H₂SO₄ for 5 h (Wan Mohamed 1981). In one experiment with concentrated H₂SO₄ and gorse seed, the highest germination was obtained after 60 min of immersion followed by a 21-day incubation (Butler 1976). In a previous study of gorse seed, the best constant temperature for germination was 15°C (Moss 1959). However, the optimum temperature for scarified seed germinated in light was 16°C. The germination rate declined at temperatures above optimum (Zabkiewicz & Gaskin 1978). Another study showed the optimum constant temperature range for germination of gorse seed was 15-19°C (Ivens 1983). Butler (1976) found more seed germinated under alternating temperatures between 15 and 20°C for 19 weeks than untreated seeds kept at 20°C for 17 weeks.

This study is part of a program that is determining the proportion of gorse seeds that are hard or dead to establish why some gorse seeds are not germinating. These experiments measured the amount of gorse seed that germinated under laboratory conditions with differing temperature, concentrated sulphuric acid, hot water and mechanical scarification.
treatments. The best scarification method and temperature for gorse seed germination will be used in the laboratory to test gorse viability throughout the South Island, New Zealand.

MATERIALS AND METHODS

Gorse seed was collected from plants growing at Onekaka, Golden Bay, New Zealand, site reference 40° 46.31 S, 172° 43.49 E, altitude 68 m. The seed for Experiment 1 was collected on 20 September 2002 and for Experiment 2 on 17 November 2002. Only mature brown pods were collected. Until required the seed was stored in the pods in a refrigerator at 4°C. When required for testing, the pods were dried at 60°C for 24-48 h then threshed in a Kurt Peltz thresher. After threshing, the seed was washed and any that floated or appeared shrivelled were discarded. The seed was carefully examined to exclude any seed that had been attacked by *Cydia succedana* or *Exapion ulicis*, which are both insects that feed on gorse seed.

All equipment was sterilised in an autoclave and only sterilised water was used. To further minimise the risk of infestation by fungi and bacteria, seed samples were washed in 2.5% sodium hypochlorite (NaOCl) for 5 min. The seed was then rinsed and air-dried overnight at room temperature.

Incubation temperature

The incubator temperatures used in both experiments were a constant 5, 10, 15, 20 or 25°C. Seed was germinated for 10 days. For each treatment, 25 seeds were placed on a moist germination paper in a Petri dish. There were four replicates at each temperature of each scarification treatment. Dishes were inspected daily and sterilised water added as required. The incubators were operated with 16 h light and 8 h dark cycle.

Experiment 1: Acid scarification

Two volumes of acid to one volume of seed were used (Hartmann et al. 2002). Seed samples were placed in concentrated sulphuric acid (36N) for 0 (control), 30, 60, 90, 120, 150, 180, 210, 240, 270 or 300 min. Following acid treatment seed was washed in running water until the pH was neutral, and then rinsed in sterile water. After rinsing, the seeds were placed in Petri dishes.

Experiment 2: Hot water and mechanical scarification

Surface-sterilised seed was immersed in hot water at 70, 75, 80, 85, 90, 95 or 100°C for 1, 2 and 5 min. Following hot water immersion, the seed was rapidly cooled by placing it under running cold water. Control seed was only surface sterilised. Another set of seeds was mechanically scarified by placing them in a sealed tobacco tin lined with fine sandpaper through which compressed air was pumped for 10 seconds (R.J. Lucas, pers. comm.). After scarification treatments, the seeds were placed in the Petri dishes.

Data collection and statistical analysis

The number of seeds germinated was counted 10 days after scarification. A seed was counted as germinated when it showed a radicle that was at least the length of the seed. Seedlings were not classified as normal and abnormal. Data were analysed using analysis of variance with the GENSTAT for Windows (release 6) package.

RESULTS

Experiment 1: Acid scarification

Gorse seed required both acid scarification and incubation temperatures of up to 15°C to obtain good germination. Germination temperatures above 15°C reduced germination; at 25°C the mean germination was only 8.9% (Table 1).
TABLE 1: The mean germination of gorse seed (%) after 10 days at different constant germination temperatures, after scarification in concentrated sulphuric acid (36N) for between 0 and 300 min.

<table>
<thead>
<tr>
<th>Temperature (ºC)</th>
<th>Mean Germination (%)</th>
<th>SE</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>45.9</td>
<td>0.011</td>
</tr>
<tr>
<td>10</td>
<td>62.6</td>
<td>0.020</td>
</tr>
<tr>
<td>15</td>
<td>64.9</td>
<td>0.022</td>
</tr>
<tr>
<td>20</td>
<td>36.5</td>
<td>0.011</td>
</tr>
<tr>
<td>25</td>
<td>8.9</td>
<td>0.011</td>
</tr>
</tbody>
</table>

Germination increased with increased time in acid (Table 2). The highest germination was after 180 and 210 min in acid with 81% germination at the optimum incubation temperature of 15ºC (Fig. 1).

TABLE 2: Mean germination of gorse seed (%) after different times of immersion in concentrated sulphuric acid averaged over all incubation temperatures.

<table>
<thead>
<tr>
<th>Immersion in acid (min)</th>
<th>Mean germination</th>
<th>SE</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>10.2</td>
<td>0.032</td>
</tr>
<tr>
<td>30</td>
<td>40.4</td>
<td>0.034</td>
</tr>
<tr>
<td>60</td>
<td>46.0</td>
<td>0.028</td>
</tr>
<tr>
<td>90</td>
<td>48.6</td>
<td>0.036</td>
</tr>
<tr>
<td>120</td>
<td>57.0</td>
<td>0.043</td>
</tr>
<tr>
<td>150</td>
<td>49.6</td>
<td>0.044</td>
</tr>
<tr>
<td>180</td>
<td>49.6</td>
<td>0.039</td>
</tr>
<tr>
<td>210</td>
<td>50.6</td>
<td>0.031</td>
</tr>
<tr>
<td>240</td>
<td>47.2</td>
<td>0.023</td>
</tr>
<tr>
<td>270</td>
<td>40.0</td>
<td>0.032</td>
</tr>
<tr>
<td>300</td>
<td>42.8</td>
<td>0.032</td>
</tr>
</tbody>
</table>

FIGURE 1: Germination of gorse seed (%) in response to time of immersion in sulphuric acid when germinated at 15ºC for 10 days. Individual Petri dishes are represented by dots and the line shows the predicted percentage of germinated seed. The regression equation for this line is: Percentage germination = 28.2 + 0.594 (min in acid) – 0.00167 (min in acid)^2, S = 13.9, R^2 = 58.6%, R^2 (adj) = 56.6%.

None of the hot water treatments increased germination significantly over the unscarified control seed. Mechanical scarification gave a germination of 35%, which was significantly different (P<0.001) from unscarified seed, which had a germination of 10%.
DISCUSSION

These experiments were conducted in order to find the method of scarification and temperature that would cause hard seeds to germinate. Previous studies of gorse seed reported that scarification or heat enhances germination (Zabkiewicz & Gaskin 1978; Ivens 1983). Experiment 1 showed that acid scarification significantly increased seed germination. It also indicated that the optimum temperature for acid scarified gorse seed germination was about 15°C, with germination declining above 15°C. Ivens (1983) found that the optimum temperature for germination was 18°C and then the germination decreased up to 26°C; above 26°C germination was inhibited. Experiment 2 showed that, over the range of temperatures and times tested, hot water scarification did not increase gorse seed germination.

Mechanical scarification utilises abrasion, particularly by rough surfaces, and is the most common treatment for impermeable seeds. Scarification can change the percentage of germinated seeds from less than 20% to more than 90% (Rolston 1978). However, in this case, germination after mechanical scarification was only 35%.

Previous experiments have demonstrated that gorse seeds germinate under extreme heat (Zabkiewicz & Gaskin 1978; McAlpine & Timmins 2002). McAlpine & Timmins (2002) found that gorse seed that had been exposed to dry high temperatures (up to 160°C for up to 10 min) still germinated, with the most favourable temperature being 140°C for 5 min where there was 50% germination. However, the current experiment with hot water treatment did not find that the wet high temperatures caused more seed to germinate. This indicates that the gorse seed requires a higher temperature to break the hard shell than that provided by this hot water experiment.

In these experiments, seeds were taken from pods, which were picked from gorse bushes. As a result some seed may not have been fully mature despite the fact that only black pods were selected. This is reflected in the low (10%) germination of unscarified seeds. Since mean germination rates of up to 81% were obtained for some scarification treatments, most of the seed used in these experiments (approximately 70%) was hard. In previous work Ivens (1983) took gorse seed from the soil, collected seed from trays placed under gorse bushes and took seed from ripe pods. It was found that there was a higher proportion of hard seed in seed from pods than in seed collected from seed trays, and very little of the seed from the soil was hard.

The results from this experiment show that the best method of scarifying gorse seed was with concentrated sulphuric acid, giving 81% germination when the seed had been soaking in the acid for 180–210 min. The temperature that gave the best germination was 15°C. This coincides with previous experiments on gorse seed (Ivens 1983) as well as other moderate legume shrubs (Wan Mohamed 1981; Tesfaye 1989). The methods that produced the best results in this experiment will be used in sample tests to calculate the percentage of viable gorse seeds throughout South Island, New Zealand.

ACKNOWLEDGEMENTS

We thank Ross Patching for the source of gorse pods, Mike Bowie for assistance in the laboratory and Richard Sedcole for statistical analysis.

REFERENCES

