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Gibberellic Acid Promotes Seed Germination in *Penstemon digitalis* cv. Husker Red

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Additional index words. dormancy, seed soak, GA3

**Abstract**. Penstemon seed often shows an inconsistent or a low germination percentage. Although most select cultivars are propagated by cuttings, for export to other countries, seed is preferred. Three experiments were conducted to determine if soaking seed in gibberellic acid (GA3) would increase seed germination of *Penstemon digitalis* cv. Husker Red. GA3 concentrations used were 0, 10, 50, 100, 200, and 500 mg L⁻¹ (first experiment); 0, 500, 1000, and 1500 mg L⁻¹ (second experiment); and 0, 500, and 1000 mg L⁻¹ (third experiment). The first and second experiments were conducted in a growth chamber, whereas the third experiment was conducted in both a growth chamber and greenhouse with seeds either covered or not covered by the mix. In all experiments, GA3 increased the percentage and rate of seed germination. The 1000 mg L⁻¹ GA3 was the best treatment. In the third experiment, percentage and rate of seed germination were the highest for seeds grown inside of the growth chamber, probably as a result of the consistency of temperatures and darkness. In the greenhouse, the percentage of seeds that germinated and the rate of germination were similar whether or not the seeds were covered with mix and whether they received either the 500 or 1000 mg L⁻¹ GA3 treatment.

Over the past few years, various penstemons (from ≈280 species) have become increasingly popular as a garden flower. These plants grow and tolerate a wide range of temperatures and have a variety of flower types and colors. Of particular interest to us is *Penstemon digitalis* cv. Husker Red, because its tall showy flower spikes have potential as a specialty cut flower. Although cuttings are the preferred method of propagating ‘Husker Red’ penstemon, for locations and regions where cuttings are not readily available, seedlings could be screened and then selected for stock plant production. However, seeds of many *Penstemon* species possess a dormancy that limits seed germination. As a result, individual germination requirements vary widely both among and within species (Kitchen and Meyer 1991; Meyer et al., 1995). In perennial production classes at the University of Nebraska–Lincoln, fresh penstemon seed sown without any soaking will take 4 to 5 weeks to germinate with only a maximum of 50% success rate (unpublished data).

Gibberellic acid (GA3, GA4, and GA5) has been shown to break dormancy and increase germination in seeds of several genera (Bewley and Black, 1982, 1985), including certain penstemon (Atwater, 1980; Meyer et al., 1995). Using Intermountain region penstemon species, Kitchen and Meyer (1991) working with five concentrations of GA3 (0 to 500 ppm) in combination with a cold treatment (2 °C) found that seeds that were treated with GA3 and stratified (to simulate natural conditions that a seed must endure before germination) had higher germination percentages. Raeber and Lee (1991) soaked seeds of *Penstemon parryi* in six concentrations of GA3 (0 to 500 ppm) for 24 h and found 87% to 97% seed germination. Neither low temperature stratification nor NaCl soaks were as effective as 500 ppm GA3 for seeds of *P. parryi*. Mexican species such as *P. harwegii* and *P. gentianoides* as well as *P. barbatus* ‘Coccineus’ germinated without stratification at 21 °C (Nau, 1996; Way and James, 1998).

Given the desire to grow ‘Husker Red’ penstemon as a specialty cut flower in a country where cuttings/plugs are not available and difficult to impossible to send from the United States and because seeds of ‘Husker Red’ are usually sown while fresh (when used for breeding and selection purposes) and often result in uneven seed germination (Lindgren, personal communication) or are reported to take up to 8 weeks for germination (Swayne, 2000), three experiments were conducted to determine if GA3 soaks would promote seed germination to facilitate stock plant establishment.

Materials and Methods

Two seed germination experiments were conducted in a dark growth chamber in the Plant Science Building at the University of Nebraska, Lincoln, NE (UNL). The first experiment was conducted from 13 to 27 Apr. 2007 and the second from 12 to 26 Aug. 2007. The third experiment was conducted from 23 Sept. to 8 Oct. 2007 in a growth chamber and the UNL double polyethylene greenhouses. Fresh seeds of *Penstemon digitalis* ‘Husker Red’ were obtained from Dr. Dale Lindgren’s breeding program and were soaked for 24 h in 50 mL of one of the following treatments: 0, 10, 50, 100, 200, and 500 [first experiment; used by Raeber and Lee (1991) on *P. parryi*]; 0, 500, 1000, and 1500 (second experiment); and 0, 500, and 1000 (third experiment) mg L⁻¹ GA3. All seed mixtures were gently swirled at the beginning, middle, and end of the 24-h soak period in each experiment. Floating (empty) seeds were discarded. Seed mixtures were then filtered and rinsed under cold distilled water for 2 min.

In the first and second experiments, 15 seeds were placed in a small petri dish (5 cm) with No. 2 Whatman filter paper, which was saturated with 1 mL of distilled deionized (d/d) water initially and then watered as needed. There were 15 petri dishes per treatment placed randomly across four shelves in a dark growth chamber at 21-3 °C.

In the third experiment, seeds were soaked and then sown into six packs (128 cells per tray; each cell holds 80 mL of mix) such that each treatment had eight cells with one seed per cell. Seeds were sown by either placing them on top of the mix (uncovered) or lightly covered by the mix (1:1 peat vermiculite by volume). There were eight flats total with two blocks per flat. Six flats were placed in the greenhouse (24 to 32 °C day/21 to 24 °C night) and two flats were placed in a dark growth chamber (21-3 °C). All flats were covered with a plastic dome and each cell was watered with 4 mL of d/d water initially and then as needed.

For Expts. 1 and 2, seeds were observed daily for germination and were considered germinated when the cotyledons appeared.

For Expt. 3, seeds were considered germinated when the cotyledons appeared.
The experimental design for the first and second experiments was a completely randomized design. The third experiment was a randomized complete block design with the covered and uncovered planting treatments in the greenhouse or in the growth chamber analyzed as four separate experiments. Thus, there were 12 replications for each—the uncovered and covered in the greenhouse and there were four replications for each—the uncovered and covered in the growth chamber.

All experimental data were subjected to a Shapiro-Wilk test of normality to determine if the data needed to be transformed. Residuals versus predicted values were also plotted to assess the assumption of constant variance. Because the variance was found to be constant and the seed count data were normally distributed, the data were not transformed. Statistical analyses were then conducted using analysis of variance implemented in SAS PROC MIXED (SAS Institute, 2006) identifying significant differences through the use of contrasts. Least square means were expressed as percent germination.

The number of germinating seeds out of 15 for Expt. 1 and out of 10 for Expts. 2 and 3 was also modeled as a function of time using a logistic model implemented in SAS PROC NLMIXED (SAS Institute, 2006) from the fitted models, time until 50% germination (T50) was calculated, and differences between T50s for different levels of GA3 were calculated.

### Results and Discussion

**Expt. 1.** Gibberelic acid increased seed germination and the rate of germination as compared with the control (Fig. 1; Table 1). The increase was directly proportional to the increase in GA3 concentrations within the range used. All concentration of GA3 reduced the number of days to 50% seed germination, but particularly 500 mg L\(^{-1}\) (12 d) as compared with the control (23 d; water soak only) (Table 1). However, total germination percentages were not much higher than 50% of the seed sown.

**Expt. 2.** As in Expt. 1, all GA3 treatments were effective for increasing seed germination when compared with the control (Fig. 2). Moreover, seeds from the control treatment (0 mg L\(^{-1}\)) did not start to germinate until 5 d after the beginning of the experiment, whereas seeds from all other GA3 treatments started to germinate after 48 h. The highest percentage of seed germination occurred when seeds were soaked with 1000 mg L\(^{-1}\) GA3 (almost 100% germination after 8 d). Again, all GA3 treatments were effective for increasing the speed of seed germination by 7 to 13 d (Table 2). Seeds germinated quicker when they were soaked in either 1000 or 1500 mg L\(^{-1}\) GA3 (Days 4 and 5).

**Expt. 3.** All levels of GA3 were again effective for increasing seed germination when compared with the control treatment (Table 3). In all treatments, the speed of germination and the number of seeds that germinated was higher for plants grown in the growth chamber compared with plants in the greenhouse (for figures, see Mello, 2009).

All concentrations of GA3, except the control, were effective for increasing the speed of germination when seeds were either not covered or were covered with mix (Table 3). When seeds were not covered with mix, seeds soaked in either the 500 or 1000 mg L\(^{-1}\) GA3 germinated in half the time compared with the control. When seeds were covered with mix, seeds soaked in 1000 mg L\(^{-1}\) GA3 germinated more rapidly.

Seeds flats placed in the greenhouse were slower to germinate than all the seeds that were grown in the dark growth chambers (Table 3). In the greenhouse, it did not matter whether the seeds were covered with mix; either GA3 concentration caused seeds to germinate faster. There was no difference in speed of germination between the two concentrations of GA3.

In these experiments, we have shown that 24-h GA3 soak increases the number of seeds germinated and the speed of seed germination for ‘Husker Red’ penstemon. Similar increases in germination when seeds were soaked at similar concentrations as those used in this research have been shown to effectively increase seed germination for a wide variety of seeds such as *Verbena bonariensis* (Kornegay and Doubrava, 2006), *Sesamum indicum* (Kaya et al., 1995), *Trichocereus terschecckii* (Ortega-Bae and Rojas-Aráchig, 2007), *Sesleria varia* (Casciglioni et al., 2004), *Sesamum indicum* (Kaya et al., 1995), *Ferula gummosa* and *Teucrium polium* (Nadajfi et al., 2005), black mulberry (Koyuncu et al., 2005), *Cyclocarya paliurus* (Fang et al., 2006), *Prunus avium* (Çetinbaş et al., 2006), *Gautheria procumbens* (Ruchala, 2002), and *Galeopsis speciosa* (Karlsso et al., 2006).

So the question arises as to whether *Penstemon digitalis* seeds possess dormancy. According to Nold (1999) in the genus *Penstemon*, dormancy can be removed by subjecting the seeds to temperatures below ≈7 °C (45 °F) for weeks or months depending on the species. Gibberellins (GA3, GA4, and GA7) have been shown to break dormancy in numerous genera of seeds (Bewley and Black, 1982, 1985), including *Penstemon* (Atwater, 1980). Kitchen and Meyer (1991) working with 50, 150, 250, and 500 mg L\(^{-1}\) GA3 recommend treatments with GA3 to remove seed dormancy or shorten the chilling requirement for many, but not all, species of *Penstemon* and recommended the use of gibberellins together with other treatments such as stratification or scarification to increase the effects of GA3 in breaking dormancy. However, according to Atwater (1980), plants in the Scrophulariaceae family to which *Penstemon* belongs have seeds in which the endosperm surrounds the embryo and occupies up to half of the seed. In our data, there was a trend that, over experiments (and thus over time), the percent germination increased for the same treatment, particularly the 0 mg L\(^{-1}\) GA3 treatment. One reason may be that these seeds were undergoing after-ripening. Another possibility could be an increasing permeability of the seedcoat because penstemons have been shown to have thin, fragile seedcoats (Atwater, 1980). However, ‘Husker Red’ seeds, which were acid-scarified for 15 to 60 min at 15-min intervals, did not show improved germination (Mello and Paparozzi, unpublished data). Thus, if there is dormancy, it is probably a physiological one, endodormancy (Baskin and Baskin, 2001).

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**Table 1.** The actual percentage of seed germination after 15 d and the estimate of the number of days it would take to 50% seed germination when seeds of *Penstemon digitalis* ‘Husker Red’ were soaked for 24 h in varying concentrations of GA3.

<table>
<thead>
<tr>
<th>GA3 (mg L(^{-1}))</th>
<th>Germination (%)</th>
<th>Estimate (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>8.63</td>
<td>22.83 ± 0.84</td>
</tr>
<tr>
<td>10</td>
<td>21.18</td>
<td>18.11 ± 0.34</td>
</tr>
<tr>
<td>50</td>
<td>18.43</td>
<td>19.6 ± 0.50</td>
</tr>
<tr>
<td>100</td>
<td>26.47</td>
<td>17.00 ± 0.30</td>
</tr>
<tr>
<td>200</td>
<td>33.33</td>
<td>15.82 ± 0.23</td>
</tr>
<tr>
<td>500</td>
<td>54.31</td>
<td>12.10 ± 0.13</td>
</tr>
</tbody>
</table>

*All pairwise comparisons between treatments were significantly different from each other at Pr < 0.01.*

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**Fig. 1.** Percentage of *Penstemon digitalis* ‘Husker Red’ seeds that germinated each day after soaking for 24 h in 0, 10, 50, 100, 200, or 500 mg L\(^{-1}\) gibberelic acid (GA3). Seeds were sown on moistened filter paper in petri dishes and germinated in a dark growth chamber at 21 to 24 °C.


Mello, A.M. 2009. Vernalization and gibberelic acid in the germination and development of Penstemon digitalis cv Husker Red. PhD Diss. Federal University of Santa Maria, Brazil.


Table 2. The actual percentage of seed germination after 15 d and the estimate of the number of days it would take for 50% seed germination when seeds of Penstemon digitalis ‘Husker Red’ were soaked for 24 h in varying concentrations of GA3.

<table>
<thead>
<tr>
<th>GA3 (mg L⁻¹)</th>
<th>Germination (%)</th>
<th>Estimate (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>24.00 ± 1.00</td>
<td>17.94 ± 0.82</td>
</tr>
<tr>
<td>500</td>
<td>71.00 ± 1.00</td>
<td>10.19 ± 0.25</td>
</tr>
<tr>
<td>1000</td>
<td>95.00 ± 1.00</td>
<td>4.17 ± 0.17</td>
</tr>
<tr>
<td>1500</td>
<td>88.00 ± 1.00</td>
<td>5.43 ± 0.21</td>
</tr>
</tbody>
</table>

Table 3. The actual percentage of seed germination after 15 d and the estimate of the number of days it would take for 50% seed germination when seeds of Penstemon digitalis ‘Husker Red’ were soaked for 24 h in varying concentrations of GA3 and then sown in soilless mix.

<table>
<thead>
<tr>
<th>GA3 (mg L⁻¹)</th>
<th>Greenhouse</th>
<th>Growth chamber</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Uncovered</td>
<td>Covered</td>
</tr>
<tr>
<td></td>
<td>Germination (%)</td>
<td>Estimate (days)</td>
</tr>
<tr>
<td>0</td>
<td>50.00 ± 1.00</td>
<td>16.85 ± 1.00</td>
</tr>
<tr>
<td>500</td>
<td>100.00 ± 1.00</td>
<td>46.88 ± 1.00</td>
</tr>
<tr>
<td>1000</td>
<td>84.38 ± 1.00</td>
<td>75.00 ± 1.00</td>
</tr>
</tbody>
</table>

Fig. 2. Percentage of Penstemon digitalis ‘Husker Red’ seeds that germinated each day after soaking for 24 h in 0, 500, 1000, 1500 mg L⁻¹ gibberellic acid (GA3). Seeds were sown on moistened filter paper in petri dishes and germinated in a dark growth chamber at 21 to 24 °C.