Influence of Seed Size, Testa Color, Scarification Method, and Immersion in Cool or Hot Water on Germination of *Baptisia australis* (L.) R. Br. Seeds

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Abstract. A series of experiments was performed to examine the germination responses of *Baptisia australis* (L.) R. Br. seeds. Germination tests were conducted at 23 °C and numbers of germinated seed were counted daily for 21 days. Seeds were separated into two size fractions using standard sieves. Seeds in the large-seeded fraction were heavier than those in the small-seeded fraction, but seed size/weight did not affect the germination percentage at 21 days (G21), the number of days to 50% of final germination (T50), or the number of days between 10% and 90% germination (T90 – T10). Seeds were classified into two groups based on testa color. Light-brown seeds (17% of total) were heavier and had lower G21 and higher T50 and T90 – T10 values than medium- to dark-brown seeds (83% of total). Seeds scarified mechanically germinated nearly 100% and had lower T50 and T90 – T10 values than untreated seeds. Untreated seeds had a higher T90 value than seeds soaked overnight in 20°C water, but the G21 and T90 – T10 values were similar for the two treatments. Mechanical scarification followed by overnight soaking in 20°C water yielded a G21 value of only 12%, and the low germination percentage was attributed to inhibition damage. When seeds were scarified in concentrated H2SO4 for 0, 1, 5, 20, 40, or 80 min, G21 values increased quadratically while T50 and T90 – T10 values decreased quadratically as the immersion time increased. To test the effects of moist heat on germination responses, seeds were immersed for 0, 0.5, 1, 2, 4, or 8 minutes in 85 °C water. G21 values increased linearly as the immersion period increased from 0 to 2 min but remained similar when the immersion time exceeded 2 min. The duration of immersion in hot water did not affect the T50 values whereas T90 – T10 values decreased linearly as the immersion period increased. We conclude that physical dormancy is responsible for temporal variation in germination of *B. australis* seeds. Scarifying seeds in concentrated H2SO4 for 20 to 80 minutes may be the most practical means of treating bulk lots of *B. australis* seeds to obtain rapid and uniform (≥85% germination).

*Baptisia australis* (Fabaceae, subfamily Papilionoideae) is a herbaceous perennial native to rich woods and alluvial thickets from Georgia to Pennsylvania, West Virginia, southern Indiana, and Kentucky (Fernald, 1915). In the northern U.S., plants are 0.6 to 1.2 m tall and produce terminal racemes of indigo-blue flowers in late May and June (Cumming and Lee, 1960; Nau, 1996). It is a widely cultivated garden plant and is the most common *Baptisia* species grown as an ornamental (Armitage, 1989).

*Baptisia australis* is propagated primarily by seed (Nau, 1996). Seeds of *B. australis* sown in germination trays in a greenhouse without temperature controls in Lexington, Ky. continued to germinate over an eight-year period (Baskin and Baskin, 1998; Dirr (1987) reported ≥90% germination when *B. australis* seeds were collected at maturity and immediately subjected to scarification (mechanical or acid), stratification, or soaking in cold or hot water. Bratcher et al. (1993) obtained 80% germination for *B. australis* seeds stratified at 5 °C for 10 weeks, but the germination percentage was only marginally lower (about 70%) for nonstratified seeds. The present study was conducted to determine the effects of seed size, seed color, scarification method (mechanical and H2SO4), and immersion in cold or hot water on the germination responses of *B. australis* seeds.

**Materials and Methods**

*Plant material.* Seeds were harvested 1 Oct. 2004 from a field plot of *B. australis* plants growing at the Univ. of Massachusetts, Amherst, Mass. (lat. 42°22.5′N). All plants in the plot were propagated from seed collected in August 1998 from a single plant growing on the Univ. of Massachusetts campus.

Seeds were cleaned and separated into two size fractions using U.S. standard test sieves. For the large-seeded fraction, 100% of the seeds passed through a No. 6 sieve (3.35 mm nominal opening) and 100% were retained on a No. 8 sieve (2.36 mm nominal opening). For the small-seeded fraction, 100% of the seeds passed through a No. 8 sieve and 100% were retained on a No. 10 sieve (2.00 mm nominal opening). Seeds were stored at 20°C and 30% relative humidity until experiments commenced. Germination experiments were performed between mid-October 2004 and January 2005.

*Germination methods.* Seeds were sown in 15-cm glass petri dishes on top of a single layer of blue blotter paper (Anchor Paper Co., St. Paul, Minn.). To inhibit fungal growth, seeds were treated one day after sowing with 3α,4,7,7-tetrahydro-2-[(trichloromethyl)thio]-1H-isodine-1,3(2H)-dione (Captan) at 0.24 mg/100 mL solution. Germination tests were performed in an incubator (model I-35LVL; Percival Scientific, Boone, Iowa) kept at 23 ± 0.5 °C with cool-white fluorescent lamps providing 50 ± 8 μmol·m−2·s−1 photosynthetic photon flux for 12 h daily. Blotter paper was moistened as needed with deionized water (dH2O). Mechanical scarification was performed by nicking seeds with a razor blade at the end opposite the radicle. Five experiments were conducted. The large-seeded fraction was used in all experiments except Expt. 1, which compared the germination responses of the two size fractions.

**Data collection.** Four 50-seed replications (plates) were used per treatment. Numbers of germinated seeds (radicle ≥1 mm) were counted daily for 21 d. Germinated seeds were discarded daily. The germination percentage at 21 d (G21), number of days to 50% final germination (T50), and number of days between 10% and 90% germination (T90 – T10) were calculated for each treatment. In Expts. 1, 2, and 5, seeds that had not germinated on day 21 were mechanically scarified on the 21st day and the final germination percentage was determined 7 d later (G28). T50 and T90 – T10 values were calculated using data from the first 21 d.

**Seed size (Expt. 1).** Eight 50-seed replications were collected from the large- and small-seeded fractions. Fresh weights were obtained for each replication and four of the replications were retained for germination studies.

**Seed testa color (Expt. 2).** Seed testa color was characterized qualitatively with the Royal Horticultural Society (RHS) Colour Chart (RHS, 1966). Seeds of the large-seeded fraction were divided into a light brown group and a medium- to dark brown group. The percentage of light brown seeds in the large-seeded fraction was determined for six 100-seed replications. Seed fresh weight was collected on eight 50-seed replications per color group. Seeds were not separated by color in any of the remaining germination experiments.

**Mechanical scarification and soaking in cool water (Expt. 3).** Seeds received one of the following four treatments: (1) mechanical scarifi-
Acid scarification (Expt. 4). Seeds were immersed for 0 (control), 0.5, 1, 2, 4, or 8 min in 18 M (concentrated) sulfuric acid (H₄SO₄). About 250 seeds were placed in 30 mL H₂SO₄ and stirred with a wooden dowel during the treatment period. Afterwards, seeds (including the control) were rinsed in a stream of tap water for 15 min, dried on a lab bench in glass petri dishes, and sown the following day.

Immersion in hot water (Expt. 5). Seeds were immersed for 0 (control), 0.5, 1, 2, 4, or 8 min in 85 ± 0.05 °C tap water. A constant water temperature was maintained using a digital-control device, and temperature bath (model RTE-140; NELLAB Instruments, Inc., Portsmouth, N.H.). Each 200-seed lot that comprised a single treatment was placed in a glass scintillation vial and the top was secured with polypropylene mesh that was secured with fine wire. Vials were filled with water within 2 s after placement in the water bath. Vials were drained immediately after the appropriate exposure period was completed. Seeds were dried in glass petri dishes and sown the following day.

Data analysis. Data were transformed (arc-sin for G21 and G28 and square root for T₅₀ and T₉₀ – T₁₀) before analysis. Data were analyzed using the General Linear Models (GLM) or Analysis of Variance (ANOVA) procedure of SAS (SAS Institute, Cary, N.C.). Pooled t tests were used to compare fresh weight values in Expts. 1 and 2. In Expts. 4 and 5, coefficients for linear and quadratic terms were calculated using the Interacting Matrix Programming (IML) software of SAS.

Results and Discussion

Seed size (Expt. 1). Seeds in the large-seeded fraction were heavier (P < 0.0001) than those in the small-seeded fraction. Values for 50-seed weights were 0.68 ± 0.01 g and 0.84 ± 0.01 g (mean ± SD) for the small- and large-seeded fractions, respectively. The germination responses, however, were similar for both groups (P > 0.05). G21, G28, T₅₀, and T₉₀ – T₁₀ values were 41%, 90%, 10.3 d, and 14.8 d, respectively, when averaged over the two seed-size fractions.

Seed testa color (Expt. 2). Light-brown seeds ranged from RHS Colour Chart value 164A to 165B while the medium- to dark-brown seeds ranged from 165A to 166A/B. Light-brown seeds comprised 17.1 ± 2.8% (mean ± SD) of the light-brown seeds having a slightly greater fresh weight and a lower percentage of imbibed or germinated seeds at 21 d after sowing than dark-brown seeds. Since the light-colored, more dormant seeds were heavier than the darker, less dormant seeds, it seems highly unlikely that a lighter testa color can be attributed to seed immaturity. Further research is needed to determine the mechanism(s) for greater seed dormancy in the lighter-colored seeds.

Fruits (pods) of B. australis are 3 to 4 cm long at maturity and have two rows of anatropous ovules (Fernald, 1987; Wood, 1974). Inspection of pods at seed harvest revealed substantial heterogeneity for testa color within individual pods whereas pods collected at different positions along the raceme exhibited similar patterns of seed color heterogeneity. These observations suggest that heterogeneity for testa color and seed dormancy is established within individual pods and is independent of fruit position within the raceme. Heterogeneity for seed dormancy contributes to temporal variation in germination, which serves a positive role under natural conditions by minimizing risks associated with simultaneous germination for species that occupy habitats subject to droughts, fires, or other natural catastrophes (Baskin and Baskin, 1998). From a horticultural perspective, uniform and rapid germination are desirable attributes and treatments that reduce temporal variation in germination are actively sought.

Mechanical scarification and soaking in cool water (Expt. 3). Mechanically scarified seeds germinated more rapidly than seeds in other treatments and germinated nearly 100% by day 21 (Table 1, Fig. 1). T₅₀ and T₉₀ – T₁₀ values were significantly lower for mechanically scarified seeds compared to controls. Seeds soaked overnight in 20 °C water exhibited a lower T₅₀ value than control seeds, but the G21 and T₉₀ – T₁₀ values for the two treatments were

Table 1. Effect of mechanical scarification and immersion in 20 °C water overnight on the germination responses of Baptisia australis seeds 21 d after sowing (Expt. 3).

| Treatment                      | Germination (%) | T₅₀ (d) | T₉₀ – T₁₀ (d) |
|-------------------------------|-----------------|--------|---------------|
| Control                       | 50 a            | 10.8 a | 16.0 a        |
| Scarified                     | 96 a            | 3.8 c  | 4.3 b         |
| Immersed overnight in H₂O     | 51 b            | 7.0 b  | 15.0 a        |
| Scarified and immersed overnight| 12 c           | 4.3 c  | 4.3 b         |

Means followed by different letters indicate significant differences using Tukey’s studentized range (HSD) test, P ≤ 0.05. The HSD was applied to transformed means (arc-sin for germination percentage and square root for T₅₀ and T₉₀ – T₁₀) and the results presented with the original means.

**Significant at P ≤ 0.001.

![Fig. 1. Influence of mechanical scarification and/or soaking seed overnight in cool (20 °C) H₂O on the germination percentage of Baptisia australis seeds 21 d after sowing (Expt. 3). Data are means of four 100-seed replications.](Image)
similar (Table 1, Fig. 1). Mechanical scarification followed by an overnight soaking in 20 °C water yielded a low germination percentage (Table 1, Fig. 1).

Scarifying the testa before imbibing seeds in water was detrimental to germination, and likely due to imbibition damage. An intact testa acts as a barrier to diffusion and thus restricts water uptake. Removal or injury to the testa permits an increased velocity of water uptake in the outer cells of the cotyledons, resulting in membrane damage to cotyledonary cells and solute leakage from the embryos (Bewley and Black, 1978; Powell, 1998). Imbibition in 85 °C water was detrimental to germination, and 0.0001) effect on the G21, T50, and T90 – T10 values. G21 values increased quadratically whereas T50 and T90 – T10 values decreased quadratically as the duration of exposure to H2SO4 increased from 0 to 8 min (Fig. 2).

Mechanical or acid scarification has been reported to enhance germination of Baptisia species other than B. australis. Thetford (1999) examined the germination responses in B. hirsuta and found that, at 21 d after sowing, mechanical scarification yielded 75% germination while seeds scarified for 15 to 25 min in H2SO4 had only about 10% germination. Mechanical scarification yielded rapid and uniform (~90%) germination in B. leucantha (Voigt, 1977) and B. tinctoria (Voß et al., 1994). Similar results were obtained in the current experiments, with mechanical scarification yielding 96% germination and a low (4.3 d) T90 – T10 value. Acid scarification was considerably more effective with B. australis (Fig. 2) than with B. hirsuta (Thetford, 1999).

Seed testa thickness may vary between species and may also vary according to environmental conditions during seed maturation. Hence, we would expect considerable variation among species (and also among seed lots within species) for the duration of acid treatment that elicits maximum germination.

Immersion in hot water (Expt. 5). G21 values increased linearly as the immersion period in 85 °C water increased from 0 to 2 min but remained similar when the exposure period was extended from 2 to 8 min (Fig. 3A). The period of immersion did not affect the T50 (P = 0.88) or G28 (P = 0.14) values. However, T90 – T10 values decreased linearly as the immersion period was increased (Fig. 3B). The G28 and T50 values were 87.9% and 11.0 d when averaged over all treatments.

Thetford (1999) placed B. hirsuta seeds in boiling water which was allowed to cool to room temperature over the following 24 h; seeds were sown the following day and 19% germinated by day 21. Immersing B. australis seeds in 85 °C water for 2 to 8 min yielded markedly higher germination percentages (Fig. 3) than Thetford (1999) obtained with treating B. hirsuta seeds with boiling water. Differences between B. australis and B. hirsuta in seed testa thickness may account for the differential germination responses of these two species to moist heat. Lack of treatment differences for G28 values in Expt. 5 indicates that immersion in 85 °C water for up to 8 min did not affect seed viability.

With B. australis, moist heat for 2 to 8 min (Fig. 3) was slightly less effective than acid scarification for 20 to 80 min (Fig. 2) and moderately less effective than mechanical scarification (Fig. 1). From a horticultural perspective, acid scarification for 20 min elicited slightly greater germination (84%) than the most effective (8 min) moist heat treatment (G21 = 74%) (Figs. 2 and 3). However, seeds scarified in H2SO4 for 20 min germinated more uniformly than seeds dipped in 85 °C water for 8 min (T90 – T10 = 2.0 vs. 13.0 d, respectively).

These experiments demonstrate that untreated seeds of B. australis germinate over a prolonged interval. Softening the seed coat by soaking in 85 °C water or scarifying the seed coat mechanically or chemically markedly enhanced germination (Figs. 1, 2, and 3). The high G21 and low T90 – T10 values exhibited by mechanically scarified seeds (Table 1) indicate that cold stratification is not a requirement for uniform and rapid

![Fig. 2. Influence of treatment duration of acid scarification on (A) the germination percentage at 21 d after sowing, (B) days to 50% of final germination (T50), and (C) days between 10% and 90% germination (T90 – T10) for Baptisia australis seed (Expt. 4). Data are means of four 100-seed replications.](image-url)
Fig. 3. Effects of immersing Baptisia australis seeds in hot (85 °C) \( \text{H}_2\text{O} \) for various intervals on the (A) germination percentage at 21 d after sowing, and (B) days between 10% and 90% germination (\( T_{90} - T_{10} \)) (Expt. 5). Data are means of four 100-seed replications.

Seed dormancy induced by a water-impermeable testa and/or pericarp is referred to as physical dormancy (Baskin, 1998). Baptisia australis is classified in the Papilionoideae (Fernald, 1987), a subfamily of the Fabaceae that contains numerous species with physical seed dormancy (Baskin and Baskin, 1998; Baskin et al., 2000). Our results demonstrated that scarification or immersion in hot water reduced physical seed dormancy in B. australis. Additional experiments are needed to test B. australis seeds from diverse sources to determine their responses to acid scarification and 2) extend these studies to other Baptisia species with ornamental value.

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