# Germination pretreatments to break hard-seed dormancy in *Astragalus (Fabaceae)* (#10547)

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# Germination pretreatments to break hard-seed dormancy in Astragalus (Fabaceae) spell name of your study species in full - you didn't look at Astragalus in general

Joseph M Statwick

Conservationists often propagate rare species to improve their long-term population viability. However, seed dormancy can make propagation efforts challenging by substantially lowering seed germination. Here I statistically compare several pretreatment options for seeds of Astraglus cicer: unscarified controls and scarification via physical damage, hot water, fire, acid, and hydrogen peroxide. Although only 30% of unscarified seeds germinated, just physical scarification significantly improved germination, whereas two treatments, hot water and fire, resulted in no germination at all. I recommend that rare species of Astragalus, as well as other hard-seeded legumes, be pretreated using physical scarification. Other methods have the potential to be effective, but may require considerable optimization, wasting precious time and seeds.



## Germination pretreatments to break hard-seed dormancy in Astragalus (Fabaceae)

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## ABSTRACT

1	Conservationists often propagate rare species to improve their long-term population viability.
2	However, seed dormancy can make propagation efforts challenging by substantially lowering
3	seed germination. Here I statistically compare several pretreatment options for seeds of Astraglus
4	cicer: unscarified controls and scarification via physical damage, hot water, fire, acid, and
5	hydrogen peroxide. Although only 30% of unscarified seeds germinated, just physical
6	scarification significantly improved germination, whereas two treatments, hot water and fire,
7	resulted in no germination at all. I recommend that rare species of Astragalus, as well as other
8	hard-seeded legumes, be pretreated using physical scarification. Other methods have the
9	potential to be effective, but may require considerable optimization, wasting precious time and
10	seeds.
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12	Key words: Astragalus; Dormancy; Germination; Milkvetch; Propagation; Scarification



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13	Introduction

Propagating wild species in greenhouses and common gardens for their restoration or reintroduction in native habitats can be an effective method of improving the size and viability of rare or threatened populations (Maunder, 1992; Menges, 2008). Such in situ and ex situ propagation techniques are beneficial, so long as these techniques are successful in establishing additional reproductive adults in novel, degraded, or extirpated sites (Maunder, 1992; Menges, 2008). If, however, reintroduction is unsuccessful (which it usually is (Godefroid et al., 2011)), it accomplishes nothing more than wasting resources and even further threatening the species by removing seeds that would have become the future seed bank. At ~3270 species, Astragalus (Fabaceae) is the largest genus of flowering plants in the world (Watrous and Kane, 2011). Though a few Astragalus are weedy, wide-ranging generalists, specialization on uncommon and infertile soils seems to be a hallmark of the genus (Barneby, 1964). Unfortunately, this specialization appears to restrict many species to small geographic ranges, making them more vulnerable to extinction. In the United States alone, the US Fish and Wildlife service (2014) has listed 3 Astragalus species as under review, 5 as candidate, 5 as threatened, and 16 as endangered. Although the IUCN database (2014) contains less than one half of one percent of known Astragalus species, nearly 40 percent of those with sufficient data are considered "vulnerable" or worse (9 vulnerable, 12 endangered, 18 critically endangered, and 1 extinct). NatureServe (2014), meanwhile, lists 100 vulnerable, 58 imperiled, and 31 critically imperiled species, which combine to nearly a third of the 616 Astragalus species in its database. Astragalus species, like most temperate legumes, as well as species of as many as 15 different plant families, have hard seed coats and physical dormancy, which often require scarification or stratification to break (Baskin et al., 2008; Long et al., 2012). In particular, low



36	germination rate is a known "weak point" in the life cycle of several rare species of Astragalus,
37	including A. nitidiflorus (Vicente et al., 2011), A. bibullatus (Albrecht & Penzagos, 2012), and A.
38	arpilobus (Long et al., 2012). Although prolonged dormancy of the seed bank may contribute to
39	the maintenance of genetic diversity in rare <i>Astragalus</i> such as <i>A. albens</i> (Neel, 2007) in the Any research on what evolutionary advantage of dormancy would be?
40	wild, this dormancy is counterproductive for propagation efforts.
41	Many scarification treatments have been explored in the literature, including dry heat
42	(Albrecht & Penzagos, 2012; Chou et al., 2012; Long et al., 2012), wet heat (Long et al., 2012),
43	stratification (Acharya et al., 2006; Albrecht & Penzagos 2012; Long et al., 2012), physical
44	scarification (Acharya et al., 2006; Albrecht & Penzagos, 2012), acid (Acharya et al., 2006;
45	Long et al., 2012) smoke water (Chou et al., 2012), etc., but it is rare that the results of more
46	than one or two treatments have been compared in the same study. Because different species and
47	even collections within species vary in germination rate, (Acharya et al., 2006; Albrecht &
48	Penzagos, 2012), the results of these studies are not directly comparable to one another in order
49	to determine the most effective scarification treatment. I therefore explored six different pre-
50	planting seed treatments (e.g. chemical and physical scarification) to determine which would best
51	promote germination in the generalist forage crop, Astragalus cicer "Oxley".
52	
53	Methods
54	Astragalus cicer (L.) (cicer milkvetch) is an old-world native that was introduced to
55	North America as a hardy, palatable forage crop (Acharya et al., 2006). "Oxley" is an ecotype
56	that was first collected in the former USSR and introduced to the United States in 1971 (Acharya
57	et al., 2006). Although A. cicer is not rare, it is a suitable model for rare species because it is
58	readily commercially available without threatening wild populations, and because it, like its rare



59	congenerics, is well known for its slow stand establishment, largely due to low germination rates
60	and prolonged seed dormancy (Acharya et al., 2006).
61	I exposed 50 A. cicer seeds (Granite Seed, Denver, CO) to each of six different
62	scarification treatments, starting March 15, 2013 at Denver Botanic Gardens (DBG) in Denver,
63	Some justificatio, explanation for your chosen methods would be helpful Colorado. The scarification treatments were physical damage, hot water, hydrogen peroxide,
64	acid, fire, and a control. Control seeds were planted in 1 cm <sup>2</sup> germination pots, without
65	scarification, on the surface of a seed starter mix, and covered with approximately 3 mm of
66	vermiculite. Treated seeds, except fire, were planted in the same manner, but after a scarification
67	treatment. I physically scarified seeds by cracking the seed coat opposite the radicle with a pair
68	of infant nail clippers, being careful to not damage the endosperm or embryo. For the hot water  Why this temperature? Note degree symbol missing
69	treatment, seeds were placed in a thermos and covered with boiling (~95 C) water. I closed the
70	Why so long? Is it not the case that many seeds experience mortality at prolonger temperatures above ca, 70C thermos and allowed the seeds to soak for 20 hours before planting. Peroxide seeds were soaked
71	in pure ZeroTol (27% hydrogen peroxide) for one hour before planting. Acid treated seeds were
72	Would dry heat, smoke and smoke + heat not have been better? soaked in lab grade sulfuric acid (98%) for five minutes. Fire treated seeds were scattered on the
73	soil surface of two 10 cm clay pots, and then covered with ~2 cm of dry pine needles and grass.
74	The dry material was lit with a butane torch and allowed to burn until naturally extinguished. How standardised was this in terms of soil surface temperature and burn duration?
75	Approximately 2 mm of ash remained, and the seeds were left undisturbed to germinate in the ls it reasonable to expect seeds on the soil surface to germinate following a fire?
76	clay pots. The total number of seeds germinated in each treatment was recorded approximately
77	Any reason why you didn't try soaking in cold water? twice per week for one month.
78	All seedlings were reared in a propagation greenhouse at DBG. The potting soil was
79	checked daily and kept evenly moist by DBG horticulture staff. Plants were exposed only to
80	natural sunlight, which, given the date and latitude, ranged between approximately 12 hours at
81	the beginning of the trial and 13 hours and a half hours at the end of the trial.



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#### Reference needed for this analysis and the software

Germination data were analyzed with a proportional hazards analysis using JMP v10. This analysis type is well suited to germination data in that it is intended for time series datasets composed of binary data in which each observation is a replicate (i.e. each seed has germinated or not germinated), and compares observed and expected frequencies with a  $\chi$  distribution. Repeated measures ANOVA was not used because calculating the variance of proportions based on grouped binary data is inappropriate in that the proportions are both ordinal and bounded between 0 and 1. RESULTS Seed treatment was an exceptionally strong predictor of seed germination success ( $\chi^2$ =101.4, P<0.0001, df=5, n=300). Physically scarified seeds germinated most quickly, and were more than twice as successful as any other treatment (Table 1), with a final germination You need to describe how your post-hoc pair-wise comparisons were done rate of 74% over 33 days (Figure 1). Statistically similar percentages of unscarified, acid scarified, and peroxide scarified seeds germinated (30%, 34%, and 26%, respectively) (Table 1). No seeds from either hot water or fire scarification treatments germinated. Across all treatments, the bulk of germination occurred within the first 2 weeks, with virtually no germination after that point (Figure 1). DISCUSSION Although many scarification treatments have been attempted for *Astragalus* species, my data show that not all treatments are equal in efficacy. In fact, only one treatment, physical scarification, was significantly better than the control, and both the fire and hot water treatments

were significantly worse than the control, effectively sterilizing all of the seeds.





105	Based on my data, I recommend that propagation efforts involving Astragalus species use
106	physical scarification as the primary method for breaking seed dormancy. Whereas other
107	scarification treatments have been effective in certain circumstances, physical scarification has
108	generally been shown to be the most effective treatment in studies that have compared it to
109	What species were these studies on? alternative methods (Acharya <i>et al.</i> , 2006; Albrecht & Penzagos, 2012). The only downside to
110	physical scarification, the labor-intensive nature of damaging the seed coat with sandpaper, a
111 De	scribe this equipment. Did your nail clipper treatment effectively mimic a process that could be reproduced on a large scale razor blade, or nail clippers, can be overcome with commercial equipment, if necessary, although
112	at the cost of slightly higher seed loss to excessive damage (Acharya et al., 2006).
113	Whereas other studies have demonstrated that methods involving cold, heat, acid, etc.,
114	can improve germination over controls, I recommend against their use in Astragalus, as the
115	studies comparing different durations and intensities (temperature, concentration) of these
116	treatments have found a relatively narrow range of optimal conditions (Albrecht & Penzagos,
117	2012; Chou et al., 2012; Long et al., 2012). Treatments of insufficient duration or intensity
118	appear to be incapable of breaking seed dormancy, whereas treatments of excessive duration or
119	intensity damage not only the seed coat, but the embryo as well, causing a loss of viability
120	(Albrecht & Penzagos, 2012; Chou et al., 2012; Long et al., 2012). This is evidenced in our own
121	study by the apparently insufficient acid and peroxide treatments compared to the apparently
122	excessive fire and hot water treatments.  This conclusion makes it doubly important that you more thoroughly justify your chosen treatments in the Methods



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CONCLUSIONS	
Physical scarification is a simple and reliable way to improve germination rates in	
Astragalus species with hard seed dormancy. I advise that, particularly for rare species for wh	iich
seeds are limited, attempting to optimize other techniques is an unnecessary waste of resource	es
when physical scarification is equally if not more effective.	
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164 FIGURES

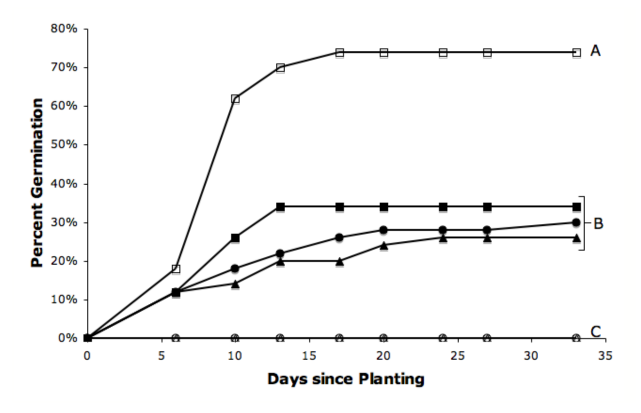


Figure 1: Germination rates over time for different scarification treatments for *Astragalus cicer*. The treatments include an unscarified control (closed circles) and seeds scarified with hot water (open circles), sulfuric acid (closed squares), nail clippers (open squares), hydrogen peroxide (closed triangles), and fire (open triangles). Letters indicate statistically different treatments via proportional hazards analysis.



Table 1: Pairwise risk ratios for treatments, expressed as the ratio of the germination success of
the row relative to the column. n=50 for each treatment. \* represents statistical significance at the
P<0.001 level.

Treatments	Control	Hot	Sulfuric	Nail	Hydrogen	Fire
		Water	Acid	Clippers	Peroxide	
Control	1	>100*	0.85	0.32*	1.17	>100*
Hot Water	<0.01*	1	<0.01*	<0.01*	<0.01*	1
Sulfuric Acid	1.17	>100*	1	0.37*	1.38	>100*
Nail Clippers	3.17*	>100*	2.69*	1	3.72*	>100*
Hydrogen Peroxide	0.85	>100*	0.72	0.27*	1	>100*
Fire	<0.01*	1	<0.01*	<0.01*	<0.01*	1



Table of raw data: Number of germinated seeds (out of 50) for each pretreatment, on a given day

## of the experiment.

Treatment	0	6	10	13	17	20	24	27	33
Control	0	6	9	11	13	14	14	14	15
Hot water	0	0	0	0	0	0	0	0	0
Acid	0	6	13	17	17	17	17	17	17
Physical	0	9	31	35	37	37	37	37	37
Peroxide	0	6	7	10	10	12	13	13	13
Fire	0	0	0	0	0	0	0	0	0