

Taproot Elongation in Woody Plant Seedlings: A Factor in Species Encroachment Potential

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Abstract

Encroachment of woody plants into arid and semi-arid grasslands is well-documented. Effects on nutrient cycles and community composition are profound but it remains unclear why some native shrubs have spread dramatically while others have not. We suggest that the capacity for rapid taproot elongation by seedlings under a wide range of moisture conditions is associated with encroachment potential. To test this hypothesis we compared seedling emergence and growth of two species with different encroachment histories in southern Arizona: *Acacia greggii* (not an encroacher) and *Prosopis velutina* (a strong encroacher). Seedlings were grown in a controlled environment for 2.5 weeks, varying the number of watering days to trigger germination ("trigger duration"; 4 levels) and the follow-up watering frequency (2 levels). We hypothesized that taproot elongation would be faster in *P. velutina* (H_1) and that *P. velutina* would be more responsive to water inputs both during (H_2) and after (H_3) germination. Evidence supporting H_1 was weak and there was no support for H_2 . However, there was strong support for H_3 ; taproot elongation increased linearly with trigger duration with a slope 58% higher in *P. velutina* than *A. greggii* ($P = 0.0076$). This suggests that newly emerged *P. velutina* seedlings are highly responsive to the magnitude of the rainfall event which triggers germination, thus reducing the time until seedlings can escape (a) topsoil desiccation and (b) strong below-ground competition from shallow-rooted competitors, such as grasses. By contrast, *A. greggii* seems less able to take advantage of rainfall early in the growing season.

Methods

Site Description and environmental basis for treatments

The experiment was conducted in a greenhouse at the University of Arizona Campus Agricultural Center in Tucson, Arizona. The Sonoran desert climate is semi-arid with most rainfall occurring in the summer monsoon (SRER 2007).

Watering treatments were based on summer precipitation from the Santa Rita Experimental Range (SRER), 40 km south of Tucson (tables 1 and 2). Data sets were provided by the Santa Rita Experimental Range Digital Database (SRER 2007).

Experimental Design

Prosopis velutina, and *Acacia greggii* seeds were chemically scarified to break dormancy: *Prosopis* in 20% H_2SO_4 for 10 minutes and *Acacia* in 90% H_2SO_4 for 20 minutes. They were then soaked 24 hours in distilled water in the dark.

For each species, four seeds were planted into each of 72 pots (Zipset™) filled with a sandy loam soil (Fig. 1). Emerging seedlings were thinned to one per pot. Four pots of each species were randomly assigned within each block (tray of pots). Irrigation was by a drip line system. Each tray was allocated to one of two feeds, operated by manual valves with a flow meter and calibrated to determine the volume of water required for each feed to supply 5mm per pot.

Triggering events started on consecutive days to allow post-trigger watering to begin the same day for all treatments. 5mm was supplied in the morning and 5mm in the evening each day of a triggering event. Thereafter, one feed delivered 5mm water to each of its pots on alternate mornings; the other feed delivered 5mm every morning.

Harvests were staggered, 16 days after 3- and 5-day triggering events and 17 days after 2- and 4-day triggering events. No water was applied the day before harvest, and 5mm water was applied as scheduled two days beforehand. Taproot length was measured. Entire seedling root systems and shoots were oven dried at 105 °C for 48 hours and then weighed.

Mean green-weight seed masses were determined after removing seed coats ($n = 20$).

Analysis

Measurements were converted to units per day of experiment to account for differences in experiment duration. Values for each species were averaged for each block (tray), as watering treatments were applied at the tray level.

Multiple linear regressions were applied to taproot elongation rate, $\ln(\text{root biomass})$, and rate of total biomass increase (root + shoot - mean seed biomass). Block location (distance from cooler pads) was a random effects variable; trigger duration and subsequent watering frequency were fixed effects. An additional regression on taproot elongation rate also included $\ln(\text{root biomass per day})$ as an explanatory variable. JMP 5.0.1a software was used for all analyses (SAS 2002).



Figure 1. Experimental setup.

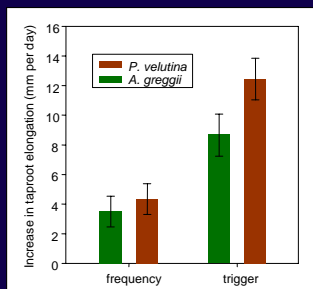


Figure 2. Root elongation responses to increased watering frequency (alternate days to every day) and increased triggering event (2 to 5 days).

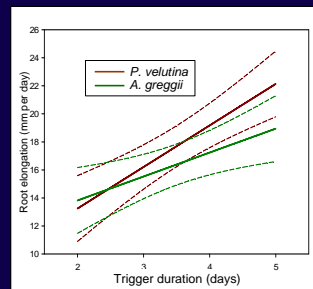


Figure 3. Plot of mean root elongation rate against triggering event, with daily subsequent watering.

Santa Rita Experimental Range precipitation: single monsoon storms			Equivalent experimental germination trigger	
Number of consecutive days' rain in July - Sept	Event frequency (yr ⁻¹)	Average cumulative rainfall during event (mm)	Number of days	Total water delivered (mm)
5	0.32	55.6	5	50
4	1.13	36.3	4	40
3	3.10	25.0	3	30
2	7.26	15.5	2	20

Table 1. Storm duration frequencies and average rainfall, SRER, July through September, 1922-2005; compared with experimental treatments (trigger duration).

Santa Rita Experimental Range precipitation: means of peak rainfall months (mm/day)			Equivalent experimental post-trigger watering:	
	July	August	frequency of 5mm waterings	
lower 95% limit	0.22	0.54	—	
mean	2.60	2.61	Alternate days	
upper 95% limit	4.99	4.67	Every day	

Table 2. Estimated mean daily summer rainfall at SRER, July and August, 1922-2005; compared with experimental treatments (follow-up watering frequency).

Results

Taproot elongation increased linearly with both triggering event and watering frequency ($P < 0.0001$, $P = 0.0001$ respectively). Across the range of treatments, the increase due to triggering event was significantly greater than that due to subsequent watering frequency (figure 1) ($P < 0.0001$). Slope of elongation rate against trigger duration was 43% higher in *P. velutina* than *A. greggii* (95% CI 10% to 89%, $P = 0.0093$; figure 2) but slope of elongation rate against watering frequency was not significantly different between the two species ($P = 0.4110$).

Taproot elongation was positively correlated with $\ln(\text{root biomass})$ ($P = 0.0464$). After taking account of $\ln(\text{root biomass})$, there remained a positive correlation with trigger duration and watering frequency ($P < 0.0001$ and $P = 0.0012$ respectively) and some evidence that *P. velutina* responded more to increased triggering event than *A. greggii* ($P = 0.0547$).

Total biomass was greater in *A. greggii* than in *P. velutina* ($P = 0.0027$, for $\ln(\text{total biomass})$) but biomass accumulation rate was five times greater in *P. velutina* ($P < 0.0001$). *A. greggii* showing only slight increase after accounting for average seed mass (*P. velutina*: 7.96 ± 1.47 mg/day; *A. greggii*: 1.49 ± 1.47 mg/day). There was no clear evidence for treatment effects on total biomass, or of differences in root/shoot biomass allocation according to species or treatment ($P > 0.05$).

Discussion

This study investigated initial seedling establishment – the most vulnerable and tractable phase of the shrub life cycle. In arid and semi-arid environments, rapid access to deeper soil moisture can be expected to be a key factor in woody plant seedling survival and the spread of shrub and tree species into grasslands. Within typical environmental limits, taproot elongation in young *A. greggii* and, especially, *P. velutina* seedlings was more responsive to triggering event duration than subsequent watering, those responses being greater than expected due only to allometric increases with biomass.

Therefore, root depth can be expected to be more dependent on the magnitudes of the larger storms (triggering events) than average wet season rainfall. Deeper soil moisture is less available to grasses and forbs and lasts longer into the dry season, so greater rooting depth can be expected to translate into increased survival through the first dry season. Young *P. velutina* seedlings appear more able than *A. greggii* to capitalize on the water provided by larger storms. This may help explain why environmental changes favoring shrub proliferation in grasslands have benefited some species more than others and, in particular, why *P. velutina* has become a ubiquitous grassland invader in the desert southwest.

It remains to be shown whether the trends shown in this study continue through the seedlings' first year, and whether the root elongation differences between invasive and non-invasive species apply more generally than to just these two species – the subjects of continuing experiments.

References

SAS, 2002. JMP® Software, Version 5. SAS Institute Inc., Cary, NC, USA.
 Santa Rita Experimental Range (SRER), 2007. <http://ag.arizona.edu/SRER/precip.xls>, maintained by SRER, viewed 05/09/2007. Funding for the digitization of these data was provided by USDA Forest Service Rocky Mountain Research Station and the University of Arizona.

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