Recurrent Selection for Improved Germination under Water Stress in Russian Dandelion

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ABSTRACT. Russian dandelion [Taraxacum kok-saghyz (TKS)] is a promising candidate for introducing natural rubber production into North America. Seeds normally germinate in a humid microenvironment, such as the thatch layer of a lawn or under a canopy of grass; however, 5% to 15% establishment is often observed on bare soil, presumably due to water stress. Phenotypic selection and half-sib family recurrent selection were conducted for three cycles to improve germination in vitro, under low osmotic potential ($\Psi_{w}$), using a polyethylene glycol (PEG) solution. Populations were then tested for establishment on bare soil in the greenhouse and field. Germination under water stress in vitro increased from 5.8% for the cycle 0 (C0) population to 40.8% and 47.8% for the C3-phenotypic and C3-half-sib family populations, respectively. Soil establishment in the greenhouse and field was improved up to two- and 4-fold, respectively, compared with the C0, in two of four greenhouse experiments and three of eight field experiments. Overall, recurrent selection for germination under water stress in vitro has potential to improve establishment in the field and can be incorporated into current breeding programs to support the overall goal of creating cultivars with high-rubber yield.

Russian dandelion is a cross-pollinated, self-incompatible diploid species (Warmke, 1943) that may be grown as a source of natural rubber in North America. TKS is adapted to southern Canada and the northern United States; however, it requires domestication to be cultivated as a new crop.

Dandelion (Taraxacum sp.) seeds typically germinate and establish in a humid microenvironment protected from direct sunlight, usually in the thatch layer of lawns and pastures (Martinkova et al., 2014). The sowing of TKS seeds on bare soil, which is prone to drying, commonly results in 5% to 15% establishment (Moussavi et al., 2016). Although more than 80% establishment is achieved in greenhouses when initiating transplants, this form of propagation would not be economical for commercial production. Consequently, breeding to improve germination under water stress could result in increased germination and establishment on bare soils in the field and facilitate the domestication of this species.

Studies of germination under water stress in soil can be problematic because consistent and uniform water potentials are difficult to maintain. The use of an osmoticum in solution can simulate drought stress and provide repeatable results in controlled environments (Springer and Goldman, 2016). Of the numerous osmotica available (e.g., PEG, mannitol, sucrose, and sodium chloride), PEG has been identified as most effective because it is nontoxic and has large molecules excluded from the seedcoat pores (Hohl and Schopfer, 1991).

The use of PEG for simulating water stress is very common (Blum et al., 1980), maize [Zea mays (Khodarahmpour, 2012)], and three dandelion species (Luo and Cardina, 2012). Percent germination of Taraxacum officinale, Taraxacum laevigatum, and Taraxacum brevicorniculatum seeds exposed to decreasing water potentials from 0 to −0.8 MPa diminished incrementally from 94% to 2%.

Germination patterns with PEG-induced drought stress are often similar to those in soil. Percent germination and germination rate in PEG were correlated with field emergence in sunflower [Helianthus annuus (Somers et al., 1982)] and wheat (Thill et al., 1979). However, the rate was much slower in soil compared with that in PEG solutions; seeds exposed to −0.4 MPa water potential germinated 4.5% per hour in PEG and 1% per hour in soil. The increased sensitivity to water stress in soil compared with PEG solutions is likely due to low hydraulic conductivity and wetted seed contact area in the soil (Hadas and Russo, 1974).

The potential for in vitro screening methods to improve germination under water stress has been demonstrated. Two cycles of phenotypic recurrent selection at −0.8 MPa increased germination of sand bluestem (Andropogon hallii) nearly 2-fold in vitro (Springer, 2011; Springer et al., 2014). In the field, establishment increased 16.4% compared with the base population (Springer et al., 2012, 2014).

In cross-pollinated crops, phenotypic (also known as mass) and family selection are common methods to improve populations (Allard, 1960). For phenotypic selection, plants are selected based on their individual phenotypes. When conducting family selection, a group of related plants is selected based on average performance. Phenotypic selection is considered a simple and easy method and allows for large populations to be screened. However, family selection is best for quantitative traits that are determined by many genes and highly affected by the environment; replication allows experimental error to be controlled and increases heritability. Furthermore, genetic gains may be increased by selecting individuals within superior families.

Seed germination under nonstress as well as stress conditions has been shown to be a quantitative trait, controlled by many loci, in several species (Czyczyl-Mysza et al., 2014;
Foolad et al., 2007; Hayashi et al., 2008; Wang et al., 2010). Consequently, family selection may be superior to phenotypic selection for population improvement. The objectives of this research were to compare phenotypic and half-sib family recurrent selection for enhancing TKS germination under water stress in vitro, and determine if establishment could also be improved on bare soil in the field and greenhouse.

**Materials and Methods**

**Genetic materials and population formation**

TKS accessions (W6 35156, W6 35159, W6 35160, W6 35162, W6 35164, W6 35165, W6 35166, W6 35168, W6 35169, W6 35170, W6 35172, W6 35173, W6 35176, W6 35177, W6 35178, W6 35179, W6 35180, W6 35181, W6 35182, W6 35183) were obtained from the Washington State University Regional Plant Introduction Station of the Agricultural Research Service (Pullman, WA), a division of the U.S. Department of Agriculture. All accessions were planted in the field and open-pollinated, and seed was: 1) bulked from 100 plants for phenotypic selection (C0-phenotypic population) and 2) collected individually from the remaining plants to form 100 half-sib families for family selection (C0-family population). Seed was rubbed against a sieve with a pore size of 425 μm to remove the papus, and then shaken over an air vent to collect the heavy viable seeds.

**Artificial water stress**

Selection for germination under water stress was conducted in vitro. For the two methods of selection described below and experiments to estimate gain from selection, seeds were incubated in 92 × 16-mm petri dishes on Whatman No. 4 filter paper with water for root development. Five selected seedlings from each of 10 plates in each of two replicate chambers resulted in 100 selected plants (selection intensity = 10%). After 14 d, seedlings were planted into 50-cell plug trays filled with a peat-based soil-less medium (Sunshine Mix No. 1; Sun Gro Horticulture, Vancouver, BC, Canada), then grown for 30 d, and transplanted subsequently into 12.7 × 12.1-cm (diameter × depth) round pots. All plants were grown in a greenhouse set to 21/18 °C (day/night) with a 16-h photoperiod, produced by high pressure sodium lamps with a photosynthetic photon flux density of 50 to 70 μmol·m⁻²·s⁻¹ to supplement natural light, and fertigated on alternate weeks with 20N–3.5P–16.6K at a concentration of 1.5 g·L⁻¹. Once flowering initiated, plants were placed inside polyethylene fabric isolation cages (Econet T; Gintec Shade Technologies, Vanessa, ON, Canada) and open-pollinated with bumblebees [Bombus impatiens (Biobest Canada, Leamington, ON, Canada)]. An equal amount of seed from each plant was bulked to create the C1-phenotypic population for the next cycle of selection. The process was repeated to generate C2- and C3-phenotypic generations.

**Among and within half-sib family recurrent selection.** For family selection, 50 seeds from each of 100 half-sib families of the C0-family population were incubated in each of four replicate petri dishes and two petri dishes were placed in each of two growth chambers under conditions described above. Seedlings were selected when ≈10 families had 5% germination in each of the four replicate petri dishes. The first 10 families (among family selection intensity = 10%) that achieved 5% germination, or if any families showed 5% germination simultaneously, those that had the highest percent germination, were selected. Within the 10 selected families, seedlings for the first two to three seeds from each petri dish replicate to germinate were selected for a total of 10 seedlings (within family selection intensity = 5%) from each family (100 total seedlings). These plants were transferred to petri dishes with water, then transplanted, and open-pollinated, as described above. Seed was collected and saved separately from each plant to produce the half-sib families constituting the C1-family population for the next cycle of selection. The process was repeated to generate C2- and C3-family generations.

**Response evaluation**

Seven populations (C0, C1–C3 phenotypic, and C1–C3 family) were evaluated simultaneously, as described below, to estimate selection gain precisely. Seeds of all populations were regenerated to eliminate seed age as a factor affecting germination. Each population was produced by intercrossing 200 random plants in polyethylene fabric isolation cages. Seed was produced for all populations simultaneously in Jan. 2015 and germinated on petri dishes with PEG 8000 solution with an Ψₛ of –1.0 MPa. For each population, four replicate petri dishes, each containing 50 seeds, were incubated within each of
four replicate growth chambers under the conditions described above. One control dish, with seeds germinated in 7-mL double distilled H₂O (ddH₂O), was incubated per population per replicate growth chamber. All petri dishes were randomized daily within each growth chamber. Percent germination was estimated daily for each treatment replicate. The experiment was repeated independently.

**Field experiment.** Seeds were planted at the Simco Research Station (SRS), Simcoe, Ontario (lat. 42°51′N, long. 80°16′W, elevation 240.5 m) in a split-split-plot design. Whole plots were soil-type, sand or loam, each replicated twice, where a replicate was a different site at SRS. Soil classifications at Sand Sites 1 and 2 and Loam Sites 1 and 2 were Scotland sand and Berrien sand, and Bookton sandy clay loam and Wattford sandy loam, respectively. Sub-plots were planting dates, 4/5 June 2015 and 25/26 Aug. 2015. Sub-subplots were 14 treatment combinations of seven breeding populations (C₀, C₁, C₁₁−C₁₃ phenotypic, and C₁₁−C₁₃ family) grown with or without irrigation. These were replicated four times in a randomized complete block design within each site x planting date treatment combination. About 15 mm of water was added to each irrigated plot by passing a watering can over the soil once on the day of planting and throughout the experiment when the soil surface was visibly dry. Each sub-subplot row was 50 cm and planted with 100 seeds at a depth of ≈5 mm; rows were spaced 76.2 cm. Lettuce seed was co-olated with the breeding populations to mark the rows and facilitate data collection. Lettuce plants and weeds were removed manually. Establishment was counted 14 d after planting (DAP). The second planting in August had low establishment after 14 d and was also assessed at 21 DAP. Seedlings that died before the counts were not included in establishment estimates. As a control, seeds of each population were incubated in four replicate petri dishes with ddH₂O, as described above.

**Greenhouse experiment.** Fresh seed of the C₀, C₃-phenotypic, and C₃-family populations were generated to conduct the experiment. The C₀ and C₃-family populations were produced in Jan. 2016 and a replicate C₀ population was produced with the C₃-phenotypic population in June 2016. The experiment was arranged in a split-plot design with four replicates in each of two greenhouses under the conditions described above. Whole plots were irrigation treatments (water-saturated or water-deficit) and sub-plots were the four populations (C₁, C₁₁−C₁₃ family and C₀-phenotypic and C₀-phenotypic control). The experiment was repeated independently; the first and second experiments were planted on 30 June and 12 July 2016, respectively. Each treatment combination replicate consisted of 50 seeds planted in a 12.7 × 12.1-cm round pot filled with a dry peat-based medium (Sunshine Mix No. 5, Sun Gro Horticulture) at a depth of 3 mm. Irrigation treatments were accomplished by passing a hose with a water breaker attachment (Ultra-soft Redhead; Dramm, Fenwick, ON, Canada) over whole plots where one pass was equivalent to ≈5 mm of raw water. Water-saturated treatments were irrigated daily with 30 mm of water and water-deficit treatments were given 30 mm of water on the day of planting and 10 mm every 2 d, which ensured that the soil surface dried between waterings. The number of established seedlings was counted daily. For each population, a control petri dish, as described above, was placed in each replicate greenhouse.

**Seed mass.** To determine the effect of selection on seed mass, 100-seed weight was measured for the C₀ and C₃ populations. Seed was produced independently in Jan. 2015 and 2016, and June 2016 for some or all of the populations as described above. Three replicate seed increases were obtained for the C₀, and two for the C₃-family and C₃-phenotypic populations.

**Statistical analyses.** Data were fit to a generalized linear mixed model using PROC GLIMMIX in SAS (version 9.3; SAS Institute, Cary, NC). The analysis of variance (ANOVA) for the in vitro experiment considered the effect of population as fixed and the effects of experiment and block nested within experiment as random. For the field experiment, the fixed effects were irrigation, population, planting date, and soil-type and the random effects were site nested within soil-type and block nested within site. For the greenhouse experiment, the effects of population and irrigation were considered fixed and the effects of experiment, greenhouse nested within experiment and block nested within greenhouse were random. For the seed mass experiment, population was considered a fixed effect and replication was random. The assumptions of the ANOVA were tested using the Shapiro–Wilk statistic for normality of error, the Levene’s test for homogeneity of error, and by visual analysis of residual plots for random distribution of error. Data with heterogeneous error were fit to a heterogeneous error model. Non-Gaussian data that could not be corrected with this model were analyzed using the Gauss–Hermite Quadrature method specified with a beta distribution and complimentary log-link (Bowley, 2015). A likelihood ratio test was used to determine if random effects differed from zero. Least square means were separated using Tukey’s honestly significant difference. Regression coefficients for the response of the two selection methods in vitro were generated using PROC GLIMMIX and compared using a Student’s t test. Significance for all analyses was determined at P ≤ 0.05. For each experiment, controls incubated in petri dishes with water had ≈90% germination; therefore, no adjustments were made to germination or establishment data to account for variation in seed lots.

![Germination Graph](https://example.com/germination_graph.png)

Fig. 1. Percent germination in Russian dandelion populations, cycle 0 (C₀) to C₃, 14 d after imbibition at −1.0 MPA, in a polyethylene glycol 8000 solution. Populations were subjected to three cycles of phenotypic or half-sib family recurrent selection for germination under low water potential in vitro. Means (n = 32), pooled over experiments, are presented ±SE; letters indicate differences among all populations according to Tukey’s honestly significant difference (P ≤ 0.05).
Results

**In vitro experiment.** The ANOVA indicated the effect of experiment and its interaction with population were not significant for percent germination 14 d after imbibition (DAI) at –1.0 MPa, therefore data were pooled over the two experiments. The fixed effect of population was significant.

Both phenotypic and half-sib family recurrent selection increased percent germination under water stress over three cycles (Fig. 1). Percent germination of C2 and C3 populations were greater than that of the C0, and values for C3 were greater than those of C2 14 DAI. The C2-family population had higher germination than C2-phenotypic; however, the two C3 populations did not differ.

Average gain per cycle 14 DAI in vitro, determined as the slope from regression analysis, was greater for family, 14.9%, than phenotypic selection, 11.6%. The former may be best to improve germination under water stress; however, significant gains are possible with the latter.

Patterns of seed germination differed among populations when incubated in water, or PEG 8000 solution with an Ψw of –1.0 MPa (Fig. 2). At 2 DAI, percent germination for seeds plated in water generally increased with advancing cycles of selection; however, differences decreased over time and all populations had greater than 95% germination 4 DAI (Fig. 2A). Seeds incubated in PEG 8000, at –1.0 MPa, germinated more slowly and had lower percent germination than those in water (Fig. 2A and B). Days to initiate germination at –1.0 MPa generally decreased with increasing selection cycles (Fig. 2B). The C3-family population had the earliest and most rapid germination, starting at 3 DAI, while that for the C3-phenotypic population began at 5 DAI. The C0, C1-phenotypic, and C1-family populations were the latest to initiate germination, beginning at 7 DAI.

**Field experiment.** For the two planting dates, 4/5 June and 25/26 Aug., the climate during the experiments represented wet and dry periods, respectively (Fig. 3). For the June planting date, 10 mm of rain fell during each of the three subsequent weeks, respectively (Fig. 3B). Over the duration of the two replicate experiments, air temperature remained constant, averaging 19 °C with a mean maximum and minimum of 24 and 14 °C,

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### Table 1. Fixed and interaction effects included in the PROC GLIMMIX (SAS version 9.3; SAS Institute, Cary, NC) analysis of variance models of percent establishment for Russian dandelion seeds in field (Simcoe Research Station, Simcoe, ON, Canada) and greenhouse experiments.

| Location          | Planting mo. | DAP* | Source of variation | Irrigation | Population | Irrigation × population |
|-------------------|--------------|------|---------------------|------------|------------|------------------------|
| **Field experiments** |              |      |                     |            |            |                        |
| Sand Site 1       | June         | 14   | *                   | *          | *          | NS                     |
| Sand Site 2       | June         | 14   | NS                  | NS         | NS         | NS                     |
|                   | August       | 14   | NS                  | NS         | NS         | NS                     |
| Loam Site 1       | June         | 14   | NS                  | NS         | *          | NS                     |
|                   | August       | 14   | NS                  | NS         | NS         | NS                     |
| Loam Site 2       | June         | 14   | NS                  | NS         | NS         | NS                     |
|                   | August       | 14   | *                   | NS         | NS         | NS                     |
| **Greenhouse experiments** |          |      |                     |            |            |                        |
| 1                 | June         | 14   | *                   | *          | NS         |                        |
| 2                 | June         | 14   | *                   | *          | NS         |                        |
| 3                 | July         | 14   | NS                  | NS         | NS         | NS                     |
| 4                 | July         | 14   | NS                  | NS         | NS         | NS                     |

*Days after planting.

Aug planting date not analyzed because almost all data points were 0.

NS = Nonsignificant or significant at $P \leq 0.05$, respectively.
respectively (data not shown). Relative humidity also did not change, averaging 81%.

The ANOVA of the field experiment, described above with four fixed effects and two random factors, indicated significant interactions between the random effect of replicate site within soil-type and one or more fixed effects (data not shown). Consequently, the eight combinations of site within soil-type × planting date were analyzed as separate experiments, with population, irrigation, and their interaction as fixed effects (Table 1).

For the June planting at Sand Site 1, both selection methods improved percent establishment; the two C₃ populations had more than three times the establishment of the C₀ at 14 DAP (Fig. 4A). Although the C₀ controls had ≈ 5% germination or establishment both in vitro and in the field, respectively, after 14 d, the response of the C₃ in the field was about half that observed in vitro (Figs. 1 and 4A). Only family selection was effective when comparing populations at Loam Site 1 for the June planting date; the C₃ population was improved compared with the C₀ at 14 DAP (Fig. 4B).

Percent establishment of the C₀ at 14 DAP was 3.4% and 33% at Sand Site 1 and Loam Site 1, respectively, for the June planting date, where populations differed (Fig. 4A and B). For the two sites where populations did not differ, Sand Site 2 and Loam Site 2, the C₀ had 40% and 32% establishment, respectively (data not shown). Sand Site 2 was located at the bottom of a slope and remained wet from the significant rainfall during the first planting date experiment. Loam Site 2 also retained moisture well. In contrast, Sand Site 1 was at the top of a slope and predisposed to drying quickly. Therefore, a lack of drought stress at Sand Site 2 and Loam Site 2 may have prevented differences from being observed between the improved and C₀ control populations.

For the August planting date, improved establishment was observed only with family selection at Loam Site 1 (Fig. 4C). The C₃-family population had more than twice the establishment of the C₀. Percent establishment for the C₀ population at Loam Site 1 was 3-fold greater for the June planting date (Fig. 4B), compared with that in August (Fig. 4C), which likely reflects differences in natural rainfall during the experiment (Fig. 3). No establishment was observed at Sand Site 1 for the August planting date while 5% establishment was observed for the C₀ at Sand Site 2 and Loam Site 2, 14 DAP (data not shown).

In some cases, irrigation had an effect on percent establishment. At the dry Sand Site 1, irrigation increased values 2.4% for the June planting date (Fig. 4A). An opposite effect was observed at the two loam sites for the August planting date 14 (data not shown) and 21 DAP (Fig. 4C); establishment was reduced an average of 4.1%.

GREENHOUSE EXPERIMENT. For percent establishment 14 DAP, the ANOVA indicated the 3-way interaction between greenhouse nested within experiment, population, and irrigation was significant (data not shown). Therefore, each greenhouse was analyzed as a separate experiment. Subsequent ANOVAs for percent establishment 14 DAP indicated that the effects of population and irrigation were significant in Greenhouse 1 and population, irrigation and their interaction were significant in Greenhouse 2 (Table 1).

Differences between C₀ and C₃ populations were observed for the water-deficit treatment in both greenhouses (Fig. 5). In Greenhouse 1, percent establishment in the C₃-family population was nearly twice that of the C₀, whereas the C₃-phenotypic population did not appear improved (Fig. 5A). Conversely, the C₃-phenotypic population had double the percent establishment...
compared with the C₀ in Greenhouse 2, whereas the C₃-family population did not appear improved (Fig. 5B).

For the water-saturated treatment, percent establishment did not differ between the improved C₃ populations and their respective C₀ controls in both greenhouses (Fig. 5). The C₃-phenotypic and C₀-phenotypic populations had lower establishment than the C₃-family populations under both irrigation treatments, indicating a seed lot effect as the seed increases for each population and control were conducted at different times. All populations, except C₃-family in Greenhouse 1, had lower establishment in water-deficit compared with water-saturated treatments.

Seed mass. One hundred seed weight of the improved and control populations did not differ and ranged from 47 to 49 mg (data not shown). Therefore, selection had no effect on seed mass.

Discussion

Three cycles of phenotypic and half-sib family recurrent selection for germination under low water potential in vitro improved the trait more than 7-fold in TKS. In soil, under greenhouse and field conditions, establishment was improved up to 2- and 4-fold, respectively, for about half of the replicated experiments. Field establishment was optimum on loam rather than sandy soils; however, less than 50% establishment was observed when rainfall was plentiful before and after planting. Consequently, factors other than moisture may be affecting field germination and establishment. Overall, selection for germination in vitro under water stress can be a viable method to improve establishment in the field.

In vitro experiments measured germination while field and greenhouse studies estimated establishment. For the field experiment, individual seeds and seedlings were not monitored daily and germinated seedlings that died subsequently could not be determined. Consequently, field establishment not only accounts for germination but also survival. The positive responses observed with improved breeding populations suggest selection in vitro has a role for improving the germination component of establishment in the field. The effect on field survival remains to be determined.

Where differences were detected among populations in replicate field and greenhouse experiments, advanced populations from family selection differed from the C₀ more often than those developed with phenotypic selection. Interestingly, the C₃-phenotypic and C₃-family populations rarely differed for the trait. However, average gain per cycle in vitro was greater for family than phenotypic selection. This may be, in part, due to increased selection intensity, first 10% among families, then 5% within selected families, compared with selecting 10% of individuals with phenotypic selection. Overall, selection with defined Ψₛ may provide a uniform water stress environment among petri plates to minimize experimental error and maximize genetic gain, giving positive results with both selection methods. With phenotypic selection, the ease and simplicity of conducting experiments may compensate for decreased advances compared with family selection. Increased effort to manage and replicate families, however, can maximize gains per cycle compared with phenotypic selection.

Springer (2011) and Springer et al. (2012, 2014) also used selection under low Ψₛ to improve germination in sand bluestem. After two cycles of phenotypic recurrent selection, germination and establishment increased from 21.5% to 38.4% 7 DAI at –0.8 MPa in vitro and from 52.5% to 61.1% in the field, respectively. Improvements in TKS reported here were greater than those observed for sand bluestem; field establishment increased up to 4-fold after three cycles of recurrent selection.

Selection appeared to have no effect on percent establishment for about half of the field and greenhouse experiments. This may be explained by large variation among replicates within certain field sites or greenhouses that reduced the detection of differences among populations. Germination or establishment experiments in soil are difficult to conduct due to inherent variation in soil structure and compaction which may affect water availability to seeds, as well as surface crusting that restricts emergence. In this study, nonuniform compaction of seedbeds produced puddles at sections of some plots after rainfall, resulting in high emergence only in those areas and inflation of experimental error. Varying moisture availability or soil properties at different sites may have also resulted in some locations with decreased water stress and diminished differences between control and improved populations.

If the detection of differences between control and improved populations at certain sites was only the result of random error, percent establishment of the C₀ controls would have been significantly greater than that of improved populations at some of the sites. Observing no differences among the C₀ and improved populations or only improved populations with greater establishment than the control is consistent with genetic advance from selection and large error in some experiments or overall moisture variation among sites. Continued cycles of selection (e.g., C₀) could increase the differences between the C₀ and improved populations relative to the experimental error and ultimately result in consistent, significant enhancement of establishment across multiple experiments. In the future, the methodology of the water-saturated and water-deficit greenhouse treatments could also be improved to increase precision.
For the field experiment planted in June, when rainfall exceeded the 15 mm·week⁻¹ minimum germination requirement recommended for TKS (Whaley and Bowen, 1947), establishment 14 DAP ranged from 3.5% to 40% for the C₀ (Fig. 4A and B, and data not shown), whereas germination with water in petri dishes was over 90%. Consequently, the 80 mm of rain that fell in nine events during the 14 d that followed planting (Fig. 3A) was insufficient to promote germination or establishment at a level similar to that observed in vitro. Soil crusting or intermittent drying of the surface, even on the loam soil, could have prevented optimum results.

Although establishment for the August planting date, during a period of drought, was less than half that of the June planting when rainfall was plentiful, a value of 25% was observed in August for the C₃-family population 21 DAP. Despite the lack of natural rainfall, the observed establishment, although delayed compared with that in June, suggests TKS has potential to develop on loam soils under adverse conditions. Continued cycles of selection would be necessary to increase germination and establishment to acceptable levels.

The high and low establishment in June and August, respectively, demonstrated the importance of rainfall and sufficient soil moisture. Not surprisingly, irrigation had very little effect during the June planting, establishment increased 2.4% at 14 DAP, at the dry Sand Site 1. During the August planting, when conditions were dry, irrigation had an adverse effect, causing a reduction at some sites. This likely resulted from crusting at the soil surface, caused by rapid drying postirrigation, which prevented seedlings from emerging. Similar research conducted on cotton (Gossypium hirsutum L.) also found soil crusting after irrigation inhibited emergence (Nabi et al., 2001).

Conflicting results, both positive and negative correlations between seed weight and germination, have been reported in different crops (Nordon et al., 2008; Rees, 1994; Venable and Brown, 1988). Springer (2011) and Springer et al. (2014) indicated sand bluestem seed weight increased with selection. However, in this study TKS seed weight did not vary among populations generated for improved germination under water stress. Consequently, seed weight would not be a useful indirect selection criterion.

Overall, genetic gains in germination and field establishment in TKS were achieved using phenotypic and half-sib family recurrent selection under PEG-induced water stress in vitro. Complexities of assessing germination and establishment in soil are also apparent and could benefit from further study. Future research should also investigate the relationship between germination or establishment and rubber content to determine the feasibility of multitrait selection. Assuming no negative correlation is observed, in vitro selection under controlled water stress can be applied to high-rubber populations for development of elite cultivars.

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