Dormancy cycling and persistence of seeds in soil of a cold desert halophyte shrub

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INTRODUCTION

Survival and regeneration of a species depend on adaptation to its habitat throughout the life cycle. Persistence of soil seed banks is an important adaptive characteristic of plants for avoiding local extinction (Stöcklin and Fischer, 1999; Thompson, 2000), and dormancy is considered to be an effective mechanism for seeds to persist in the soil (Grime, 1981; Pons, 1991; Dalling et al., 2011). Seeds of many temperate-zone herbaceous species exhibit dormancy cycles when buried in soil (Schütz, 1997a, b; Baskin and Baskin, 1998; Carter and Unger, 2003). When coming out of dormancy, the seeds gain the ability to germinate over the full range of conditions possible for the population, at which time they are non-dormant (Baskin and Baskin, 1998). There is a series of transitional states between dormancy and non-dormancy, called conditional dormancy, during which seeds germinate over a narrower range of conditions than do non-dormant seeds (Baskin and Baskin, 1998). Non-dormant seeds of some species may re-enter dormancy via conditional dormancy and annually cycle between non-dormancy and dormancy; thus, seeds of some species may exhibit an annual cycle between non-dormancy and conditional dormancy.

Dormancy cycling is an endogenous mechanism that helps regulate germination timing and thus plays a role in timing of seedling establishment of the species under seasonally varying environments (Baskin et al., 1993; Baskin and Baskin, 1996). Dormancy cycling coordinates the time of seedling emergence with favourable seasons, which increases the chance of seedling survival in seasonally varying environments (Baskin and Baskin, 1980, 1985).

In addition to seed dormancy, the soil seed bank is crucial in the adaptation of plants to deserts, since environmental conditions, especially the amount of precipitation, are highly variable (Archibold, 1995). Many herbaceous plants in arid regions have soil seed banks that allow them to survive in harsh and unpredictable environments (Facelli et al., 2005; Dreber and Esler, 2011). For seedlings to become established, it is important for seeds of desert plants to germinate when the very scarce rainfall events occur (Guterman, 1993, 2000).

It is sometimes assumed that seeds of desert species are ready to germinate as soon as temperature and soil moisture become favourable for them to do so (Went, 1949; Tevis, 1958). However, information is accumulating on the importance of dormancy as a mechanism for delay of germination in seeds of desert species (Clauss and Venable, 2000; Adondakis and Venable, 2004; Huang et al., 2004). One aspect of dormancy in seeds of desert plants that has received very little attention is dormancy cycling. We are aware of only two studies on dormancy cycling in seeds of desert annuals (Baskin et al., 1993; Cao et al., 2012), and to our knowledge there are no reports on
dormancy cycling in herbaceous perennials, shrubs or trees in deserts. In arid environments, it is well established that formation of a seed bank is important for the persistence of annuals, whereas it has been assumed that long-lived shrubs do not need a seed bank to persist at the site (Pake and Venable, 1996). If such a species has a persistent seed bank, does dormancy cycling occur, or do the seeds remain non-dormant (Baskin and Baskin, 1998; Thompson et al., 2003; Fenner and Thompson, 2005)? Some trees (Marks, 1974; Marquis, 1975) and shrubs (Hassan and West, 1986) form a local persistent seed bank, but little is known about dormancy cycling in woody plants. However, Alnus glutinosa, a small tree of relatively mesic northern temperate regions, especially Europe, is an exception. Seeds of this species buried in the field at a depth of 7 cm and exhumed and tested for germination in light and darkness at 15 °C at various intervals for 34 months exhibited reduced germination in light in September and October (autumn) and peaks of germination from January to August; few seeds germinated in darkness (Schütz, 1998). These results indicate the presence of a dormancy cycle, but since only one test temperature was used the type of cycle cannot be determined.

Since halophytic shrubs grow in very harsh environments and since persistent seed banks and dormancy cycling are known to occur in some halophytic herbaceous species (Carter and Ungar, 2003; Cao et al., 2012), we hypothesized that seeds of the cold desert halophytic shrub Kalidium gracile (Amaranthaceae) would form a persistent seed bank and exhibit dormancy cycling.

**MATERIALS AND METHODS**

*Species and seed source*

Kalidium gracile is a small shrub about 20–50 cm in height that grows on alkaline plains and salt-lake shores in northwest China and Mongolia (Zhu et al., 2003). It flowers and fruits from July to September (Zhu et al., 2003). Freshly matured seeds of *K. gracile* were collected on 20 November 2009 and 15 November 2010 from several hundred plants growing in a saline desert on the Ordos Plateau (38° 14′ N, 107° 29′ E, 1311 m a.s.l.) in Inner Mongolia, northern China. This area has a typical continental semi-arid climate with a mean annual precipitation of 250–490 mm, about 60% of which occurs from June to August. Mean annual temperature is about 6.0–9.0°C. Seeds collected in 2009 were air-dried and stored at room temperature (10–18°C, 17–32% relative humidity) until initiation of germination tests on fresh seeds and burial of seeds in soil. After air-drying at room temperature and hand separation, seeds collected in 2010 were stored dry at −20°C until used.

*Seed morphology and mass*

The lengths and widths of ten seeds were measured using a Nikon 80i (Tokyo, Japan) microscope. The 1000-seed mass was determined by weighing five replicates of 1000 seeds to the nearest 0.1 mg.

*Germination tests of fresh seeds*

Freshly collected seeds of *K. gracile* were tested for germination in Petri dishes (diameter 5 cm) on two layers of Whatman No. 1 filter paper moistened with 3 ml of distilled water at 15/5, 20/5, 25/15 and 30/15°C (12/12 h) in light (12 h photoperiod, ~100 μmol m−2 s−1, cool white fluorescent light) and in total darkness. All Petri dishes were wrapped in plastic film to reduce evaporation, and seeds incubated in darkness were placed inside light-tight black bags. The higher temperature coincided with a 12 h light period and the lower temperature with a 12 h dark period. The alternating temperature regimes represented the approximate mean daily maximum and minimum air temperatures for each month during the growing season on the Ordos Plateau: April and October, 15/5°C; May and September, 20/5°C; June and August 25/15°C; and July, 30/15°C (Cao et al., 2012). Four replicates of 25 seeds each were used for each test condition. Germination was monitored daily for seeds incubated in light, and additional water was added as necessary. Final germination percentages were determined after 20 days. Emergence of the radicle (or cotyledons, which sometimes emerge from the seed coat before the radicle) was the criterion for germination in this and all other germination experiments.

Following all germination tests, non-germinated seeds were examined under a dissecting microscope to determine whether the embryo was firm and white, indicating that it was viable, or soft and grey, indicating that it was non-viable. Tetrazolium tests confirmed that the firm, white embryos were viable and the soft, grey ones non-viable. Only viable seeds were used in calculating germination percentages. More than 90% of the seeds were viable.

*Dormancy breaking*

Fresh seeds of *K. gracile* germinated to very low percentages at 15/5°C, but buried seeds germinated to >80% under this temperature regime in spring (see Results), indicating that fresh *K. gracile* seeds have primary dormancy. Thus, we tested the effects of cold stratification, gibberellic acid (GA3) and scarification of the seed coat on germination of fresh seeds collected in November 2010 to determine the kind of dormancy in seeds of this species and the factor(s) that break it.

Cold stratification. Freshly collected seeds were arranged uniformly on two layers of Whatman No. 1 filter paper over moist sand (10–13% water content) in a metal box (20 cm length × 10 cm width × 10 cm depth). The metal box was covered with a metal lid and placed in a refrigerator at 5°C. After 0 (control), 5, 10, and 20 days of cold stratification, four dishes (replicates) each of 25 seeds were arbitrarily chosen to test germination in light at 15/5, 20/5, 25/15 and 30/15°C (12 h/12 h) as described above. Final germination percentages were determined after 20 days. GA3 treatments. Four replicates of 25 freshly collected seeds each were tested for germination in a 0.1 mmol L−1 GA3 solution in light at 15/5°C.

Seed coat scarification. Seed coats of four replicates of 25 freshly collected seeds were scarified with a scalpel, being careful not to injure the embryos, and tested for germination in light at 15/5°C.

*Dormancy cycling*

On 30 November 2009, about 1000 seeds of *K. gracile* were placed in each of 20 nylon bags. Each bag was buried at a depth of 7 cm and exhumed and tested for germination in light at 15/5°C. In arid environments, it is well established that formation of a seed bank is important for the persistence of annuals, whereas it has been assumed that long-lived shrubs do not need a seed bank to persist at the site (Pake and Venable, 1996). If such a species has a persistent seed bank, does dormancy cycling occur, or do the seeds remain non-dormant (Baskin and Baskin, 1998; Thompson et al., 2003; Fenner and Thompson, 2005)? Some trees (Marks, 1974; Marquis, 1975) and shrubs (Hassan and West, 1986) form a local persistent seed bank, but little is known about dormancy cycling in woody plants. However, Alnus glutinosa, a small tree of relatively mesic northern temperate regions, especially Europe, is an exception. Seeds of this species buried in the field at a depth of 7 cm and exhumed and tested for germination in light and darkness at 15 °C at various intervals for 34 months exhibited reduced germination in light in September and October (autumn) and peaks of germination from January to August; few seeds germinated in darkness (Schütz, 1998). These results indicate the presence of a dormancy cycle, but since only one test temperature was used the type of cycle cannot be determined.

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depth of 2 cm in plastic pots (diameter 20 cm, height 50 cm, with drainage holes) filled with soil collected from the habitat and buried on nearly flat ground at the Ordos Sandland Ecological Research Station of the Chinese Academy of Sciences (OSES) so that the tops were level with the soil surface. Mean daily maximum and minimum monthly air temperatures for each month during the study were obtained from OSES. A bag of buried seeds was exhumed on the first day of each month, beginning on 1 May 2010 and ending on 1 December 2011. Two tests were carried out using the fresh and exhumed seeds.

Test 1: germination of seeds under seasonal temperature regimes. Seeds were placed on filter paper moistened with distilled water in Petri dishes 5 cm in diameter and incubated at 15/5, 20/5, 25/15 and 30/15°C (12/12 h) in light (as described above). Four replicates of 25 seeds each were used in each test condition. Final germination percentages were determined after 20 days.

Test 2: germination of seeds under various salinities. Seeds were tested at salinities of 0, 0.2, 0.4 and 0.6 mol L\(^{-1}\) NaCl. Seeds were placed in Petri dishes on two layers of filter paper moistened with 3 ml of test solution and wrapped with plastic film to reduce evaporation. Four replicates of 25 seeds each were used. The seeds were incubated in light at 30/15°C. Final germination percentages were determined after 20 days.

Germination recovery

Germination recovery was tested for seeds of *K. gracile* that had been dry stored at −20°C for 5 months. Seeds were incubated in 1.0 mol L\(^{-1}\) NaCl and isotonic polyethylene glycol 8000 (PEG-8000) solutions in April 2011. The osmotic potential of 1.0 mol L\(^{-1}\) NaCl solution was determined with the Van’t Hoff equation, \(\mu = cRT\), where \(c\) is osmolality in mol L\(^{-1}\), \(R\) is the gas constant (8.31 J K\(^{-1}\) mol\(^{-1}\)) and \(T\) is temperature (K). The isotonic PEG-8000 solution was prepared according to Michel (1983). For both 1.0 mol L\(^{-1}\) NaCl and isotonic PEG-8000 pretreatments, 25 seeds were placed on two layers of filter paper in each of 12 dishes moistened with 3 ml NaCl or PEG-8000 solution. The dishes were placed in an incubator at a constant temperature of 20°C. This temperature was chosen to represent the average temperature of early summer (before the beginning of the rainy season), when seeds were most likely to experience stress from salinity and drought. A constant temperature rather than alternating temperatures was used to avoid changes in the osmotic potential of PEG-8000 solutions.

After 20, 40 and 60 days of pretreatment in the NaCl and PEG-8000 solutions, four dishes of seeds were arbitrarily chosen and rinsed twice with distilled water and transferred to filter paper moistened with distilled water. The seeds transferred to distilled water were incubated in light at 30/15°C for 20 days and germination percentages determined. Following the recovery germination tests, non-germinated seeds were tested for viability as described above. No seeds germinated in the NaCl or PEG-8000 solution. Recovery germination percentage \((G_p, \%)\) was determined as \(\frac{a(a + b)}{c(a + b)} \times 100\), where \(a\) is the number of seeds germinated during the recovery germination test and \(b\) the number of viable but non-germinated seeds. Viability (V\%) of pretreated seeds was calculated as \(\frac{(a + b)c}{a(a + b)} \times 100\), where \(c\) is the total number of seeds in a dish (i.e. 25). The control consisted of 16 dishes of 25 seeds each without pretreatment with NaCl or PEG-8000 solution stored dry in the 20°C incubator. Four dishes of seeds were incubated in distilled water at 30/15°C after 0, 20, 40 and 60 days of dry storage at 20°C. Final germination percentages and viability of non-germinated seeds were determined after 20 days of incubation.

Dynamics of soil seed bank

The soil seed bank of *K. gracile* was monitored monthly from April to November 2011. Ten soil cores (5 cm diameter × 10 cm depth) were collected from arbitrarily chosen places in a monospecific stand of *K. gracile*; sampling plots were about 50 m apart. The soil cores were subdivided into three layers: 0–2, 2–5 and 5–10 cm. The soil samples were washed with water through a 0.5 mm mesh sieve, and seeds were hand-sorted from the residue (roots and other vegetative parts). The number of viable seeds was determined under a dissecting microscope as described above. Only viable seeds were used in calculating seed density in the soil seed bank.

Field emergence of seedlings

Five 4 m × 4 m plots about 50 m apart were arbitrarily established in a monospecific stand of *K. gracile* in April 2011. Newly emerged seedlings of *K. gracile* were marked and recorded monthly in four 20 cm × 20 cm subplots in each of the five plots from April to September 2011. Newly emerged seedlings were monitored twice each month (on the 15th and last day), and the number of emerged seedlings was totalled for each month. Seedlings were labelled and the date was recorded when they were first found. The number of seedlings that survived until the end of the experiment was determined. This experiment was terminated on 30 September 2011, when the plants had turned red, indicating the end of the growing season. On this date, the number of living seedlings that had emerged in each month was recorded.

Soil water content, soil salinity, precipitation and temperature were monitored. Eight soil cores, 5 cm in diameter, 0–5 cm deep and about 50 m apart, were collected arbitrarily within the study population of *K. gracile* on the 15th day of each month from April to September 2011. The moisture content of each soil sample was determined gravimetrically; soil samples were oven-dried at 105°C for 48 h. Salinity was determined by the residue drying quality method (Bao, 2000). Mean monthly air temperature and monthly precipitation were obtained from a national weather station 10 km from the field site.

Data analyses

One-way ANOVA was used to estimate differences in germination percentages and viability of pretreated seeds after different durations of salinity and drought stress pretreatment. Fisher’s LSD test was performed for multiple comparisons when significant differences were found. The independent-samples t-test was used to determine significant differences (or not) between germination percentages and viability of seeds after the same period of exposure to NaCl and PEG-8000 solutions. Data were arcsine square root transformed when necessary to meet assumptions of ANOVA for normality and homogeneity of
results

Seed morphology and mass

Length and width of K. gracile seeds were 1.03 ± 0.02 mm (mean ± s.e.) and 0.80 ± 0.01 mm, respectively, and 1000-seed mass was 155.8 ± 1.5 mg. The seed coat was brown and rough.

Germination tests of fresh seeds

Fresh seeds of K. gracile germinated to <20.0 % at 15/5 °C and to <40 % at 20/5 °C in light but to 95–100 % at 25/15 and 30/15 °C (Fig. 1). Darkness inhibited germination at 15/5 and 20/5 °C (P < 0.05) but not at 25/15 and 30/15 °C (P > 0.05).

Dormancy breaking

Cold stratification significantly (P < 0.05) increased germination percentages at 15/5 and 20/5 °C (Fig. 2). Thus, seeds of K. gracile stratified for 20 days germinated to 84.8 ± 9.0 % at 15/5 °C and 90.4 ± 0.7 % at 20/5 °C (Fig. 2). Darkness inhibited germination at 15/5 and 20/5 °C, and to 14.0–33.3, 9.0–33.2, and 10.0–42.0 % at 15/5 and 20/5 °C in autumn of 2009 (fresh seeds), 2010 and 2011, but to high percentages at these two temperature regimes in the following springs (Fig. 3B). Thus, germination percentages increased at 15/5 and 20/5 °C after exposure to cold stratification in the field in the winters of 2009–2010 and 2010–2011 and decreased at these two temperatures after they were exposed to high temperature in the field in summer (Fig. 3A, B). The pattern of germination in 0.2 mol L⁻¹ NaCl at 30/15 °C in light was almost identical to that in distilled water at 15/5 and 20/10 °C. Thus, germination percentages in 0.2 mol L⁻¹ NaCl were highest in spring and lowest in autumn. Fewer than 20 % of the seeds germinated in 0.4 mol L⁻¹ NaCl throughout the 24-month burial period, and none germinated at 0.6 mol L⁻¹ NaCl (Fig. 3C).

Recovery germination

Non-pretreated seeds of K. gracile had a viability of 96.0 ± 2.3 % and germinated to 95.0 ± 1.9 %. During 0–60 days of immersion in 1.0 mol L⁻¹ NaCl or isotonic PEG-8000 solution, seeds did not change significantly in recovery germination at 30/15 °C or in viability (P > 0.05 ). They had recovery germination percentages of 94.0 ± 1.2 and 97.0 ± 1.0 after 60 days of pretreatment in the solutions of NaCl and PEG, respectively.

Dynamics of soil seed bank

Seeds of K. gracile were most abundant in the 0–2 cm level of soil, and seed density decreased with increasing soil depth (Fig. 4). Seed density in the 0–2 cm level of soil decreased in the growing season (April to September) and increased in autumn (October and November). More than 70 % of the seeds were depleted from the soil seed bank (0–10 cm) during the growing season, decreasing gradually from 7030 ± 1131 seeds per m² in April to 1987 ± 433 seeds per m² in September.

Field emergence of seedlings

Seedlings emerged in the field in July, August and September, when soil salinity was relatively low and soil water content relatively high (Fig. 5). The rate of emergence was 1300 seedlings m⁻², and 83 % of the seedlings that emerged did so in July, concurrently with the first major precipitation event, lowest soil salinity and highest mean monthly temperature of the growing season (Fig. 5). Only 9.7 % (105/1083) of the seedlings that emerged in July were alive at the end of the growing season. The mean emergence rate was 28 and 190 seedlings m⁻² in August and September, respectively, but none of the seedlings survived until the end of the growing season. Thus, 1301 of the 7030 (18.5 %) seeds m⁻² in the soil seed bank at the beginning of the growing season (April) produced seedlings, of which only 1.4 % (105/7300) survived until the end of the first growing season.

Discussion

Our hypothesis that seeds of K. gracile form a persistent seed bank and have dormancy cycling was supported. Fresh seeds of K. gracile have primary conditional dormancy, and the germination percentage can be increased at low incubation...
temperatures by a short period of cold stratification, GA3 or scarification of seed coat. Thus, *K. gracile* seeds have non-deep physiological dormancy (Baskin and Baskin, 1998). Seeds of many herbaceous annual species with non-deep physiological dormancy exhibit dormancy cycles (Baskin and Baskin, 1998), and it has recently been demonstrated that cycling also occurs at the hormonal–molecular level via differential expression of genes in response to environmental cues (Cadman et al., 2006; Footitt et al., 2011). Although seeds of *K. gracile* maintained high germination at 25/15 and 30/15°C throughout the year, they varied seasonally in germination at 15/5 and 20/5°C. Thus, seeds in the soil had an annual non-dormancy/conditional dormancy (ND/CD) cycle. To our knowledge, this is the first report of seed dormancy cycling in a cold desert shrub.

In addition to being a polycarpic perennial, *K. gracile* can form a seed bank, and the buried seeds can undergo annual dormancy cycling. These three life history traits help ensure the persistence of *K. gracile* in its harsh saline habitat. Rainfall not only increases soil water content, but also decreases soil salinity in saline deserts; however, the timing and intensity of these changes is unpredictable in deserts. The ND/CD dormancy cycle in seeds of *K. gracile* enables them to germinate throughout the growing season whenever there is available rainfall for them to do so. Seeds can only germinate to relatively high percentages at salinities equal to or lower than about 0·2 mol L$^{-1}$ NaCl, a condition that is especially unpredictable due to the unpredictability of the timing of rainfall. Thus, the ability to maintain high germinability at high temperatures ensures field emergence of this species in the growing season. On the other hand, the decrease in the ability of conditionally dormant seeds to germinate at low temperatures contributes to persistence of the soil seed bank for this species by preventing seed germination in the unfavourable autumn season, when there would be no chance of seedling survival. In this study, many fewer seedlings emerged in August and September than in July. Thus, the inability of most seeds in the soil to germinate in autumn contributes to the persistence of seeds in the soil.

**Fig. 2.** Germination percentages of *Kalidium gracile* seeds incubated at various alternating temperature regimes in a 12 h photoperiod after 0, 5, 15 and 20 days of cold stratification at 5°C in continuous darkness. Error bars are means ± s.e.

**Fig. 3.** (A) Mean monthly maximum and minimum air temperatures at the seed burial site. (B, C) Germination percentages (mean ± s.e.) of *Kalidium gracile* seeds incubated (B) in distilled water at four temperature regimes in a 12 h photoperiod and (C) in 0, 0·2, 0·4 and 0·6 mol L$^{-1}$ NaCl solutions at 30/15°C in a 12 h photoperiod at time 0 (fresh seeds) and following 5–24 months of burial in soil.
Seeds of *K. gracile* in the soil seed bank vary in sensitivity to salinity (sensitive↔insensitive) during the year, which was well correlated with the ND/CD cycle. Thus, seeds germinated to their highest percentages in 0.2 mol L⁻¹ NaCl when they were non-dormant and to their lowest percentages when they were conditionally dormant (Fig. 3). The annual cycling at 30/15°C in germination response to 0.2 mol L⁻¹ NaCl shows that cyclic changes are occurring in the seeds’ physiological response to stress that are not detectable when seeds are incubated in distilled water (Fig. 3C). In a previous study, an annual cycle of sensitivity to salinity was found in the black seed morph of the seed-dimorphic halophytic summer annual herb *Suaeda corniculata* subsp. *mongolica* (Amaranthaceae), which has an annual D ↔CD ↔ND seed dormancy cycle (Cao et al., 2012) and co-occurs with *K. gracile* in the saline desert on the Ordos Plateau. Seeds of both species exhibited less tolerance for germination to salinity when they were in conditional dormancy than when they were non-dormant. This characteristic of halophyte seeds may play a role in preventing germination in autumn, which is an unfavourable time for seedling establishment.

Soil seed banks are an important component of the regeneration strategies of many species inhabiting unpredictable environments (Leck and Schutz, 2005; Evans et al., 2007). *K. gracile* produces large numbers of seeds, and >7000 seeds per m² were present in the soil seed bank at the beginning of the growing season in 2011. Although 71.7% of seeds of *K. gracile* in the soil were depleted in the growing seasons, those that remained after 1 and 2 years of burial were part of a persistent soil seed bank,*sensa Thompson and Grime (1979). The maintenance of a persistent soil seed bank of this species may be attributed to dormancy cycling (see above) and to the ability to maintain seed viability under salinity stress. The ability to maintain viability under salinity stress is required for a species to persist in its saline habitats (Engels et al., 2011).

Seeds of *K. gracile* maintain viability under high salinity stress and then germinate to high percentages at favourable temperatures and moisture conditions. The ability of seeds to commence germination when salinity decreases following a precipitation event is an overriding factor in seedling emergence of this species, given that the seeds can germinate to high percentages at prevailing habitat temperatures throughout the growing season. In the natural habitat of *K. gracile*, the top layer of soil dries quickly after rainfall due to the high evaporation rate. The ability of seeds of *K. grasicle* to commence germination once rainfall decreases soil salinity increases the possibility of seedling establishment before the concentration of salts in the soil becomes inhibitory for germination. The small radicles of *K. caspicum* are very vulnerable to salinity (Tobe et al., 2000), and this is probably the case with *K. gracile*. This being so, it is necessary for...
Fig. 5. (A) Mean monthly air temperature and precipitation, (B) soil water content and soil salinity and (C) number of *Kalidium gracile* seedlings that emerged in each month from April to September 2011.

Fig. 6. Conceptual model of seed and seedling dynamics in a population of *Kalidium gracile*. The percentages indicate proportions of seeds in a soil seed bank that gave rise to seedlings and of those that entered the persistent soil seed bank. CD, conditional dormancy; D, dormancy; ND, non-dormancy. The dash–dot arrows represent events that may occur.
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the species to complete germination and to establish seedlings before salinity at the surface soil increases. In addition, the ability of the seeds to germinate in 0.2 mol L\(^{-1}\) NaCl may play a role in the persistence of this species at the site, since soil salinity was greater than 0.2 mol L\(^{-1}\) NaCl in the field during most of the growing season (Fig. 5). Thus, most seedlings emerged in July when rainfall decreased soil salinity to about 0.2 mol L\(^{-1}\) NaCl.

The results of our study on seed and seedling dynamics of K. gracile in the field are summarized in a conceptual model (Fig. 6). Seeds have CD when dispersed at maturity in autumn, and they become ND by exposure to low temperatures in winter. Thus, seeds are ND in spring. However, no seedlings emerged until July, when the concentration of salts in the soil was diluted by rain. Seeds re-entered CD in late spring–early summer, but retained the ability to germinate to 100% at prevailing habitat temperatures (Fig. 3). Thus, seedling emergence occurred in summer after major precipitation events in July, August and September.

Although the seeds were ND, they did not germinate during the low-rainfall events in April to June, when soil moisture content was low and soil salinity high. The high seed bank present in April 2011 was depleted by 71.7% during the growing season, but retained the ability to germinate to 100% at prevailing habitat temperatures (Fig. 3). Thus, seedling emergence occurred in summer after major precipitation events in July, August and September.

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LITERATURE CITED


