Media Selection and Seed Coating Influence Germination of Turfgrasses under Salinity

Matteo Serena
Department of Plant and Environmental Sciences, New Mexico State University, MSC 3Q, Las Cruces, NM 88003

Bernd Leinauer1
Department of Extension Plant Sciences, New Mexico State University, MSC 3AE, P.O. Box 30003, Las Cruces, NM 88003

Rossana Sallenave
Extension Animal Sciences and Natural Resources Department, New Mexico State University, MSC 3AE, Las Cruces, NM 88003

Marco Schiavon
Department of Plant and Environmental Sciences, New Mexico State University, MSC 3Q, Las Cruces, NM 88003

Bernd Maier
Department of Extension Plant Sciences, New Mexico State University, MSC 3AE, Las Cruces, NM 88003

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Abstract. Germination of five turfgrass species ['Barrister' kentucky bluegrass (Poa pratensis L.), 'Barvado' tall fescue (Festuca arundinacea Schreb.), 'Premier II' perennial ryegrass (Lolium perenne L.), 'Bargusto' bermudagrass (Cynodon dactylon L. Pers.), and ‘Sea Spray’ seashore paspalum (Paspalum vaginatum O. Swartz)] from coated (ZEBA®-cornstarch coating; Absorbent Technologies Inc., Beaverton, OR) and uncoated seeds was evaluated on both filter paper and agar. Final germination percentage (FGP) and germination rate (GR) were determined at salinity levels of 0.6 (tap water, control), 2.2 (saline groundwater from a local shallow aquifer), and 7.0, 12.5, and 22.5 dS m–1 [sodium chloride and calcium chloride (1:1; w:w) dissolved in tap water]. Final germination percentage for kentucky bluegrass, perennial ryegrass, and tall fescue was greater in agar at all salinity levels but was unaffected by the medium at any of the salinities except for 7 dS m–1 for bermudagrass and seashore paspalum. Coated seashore paspalum and coated perennial ryegrass seed exhibited greater germination than uncoated seed at four of the five salinity levels. Seed coating had no effect on FGP of bermudagrass at any salinity level and coated kentucky bluegrass seed showed reduced germination at 0.6 and 7.0 dS m–1. Final germination percentage for seashore paspalum improved from 22% to 54% at 12.5 dS m–1 and from 8% to 20% at 22.5 dS m–1 when coated seed was used instead of uncoated seed. Germination rates were unaffected by salinity levels ranging from 0.6 to 12.5 dS m–1 and were higher on agar (10%/day) than on paper (8%/day). Our study suggests that the choice of medium can influence the outcome of germination tests and that results can also vary depending on the salinity level tested and whether the seed are coated.

Loss of productive lands as a result of soil salinization is a growing worldwide problem that affects arid zone regions most acutely. This, coupled with a growing scarcity of fresh water and increased use of effluent or low-quality water for turfgrass irrigation, has prompted turfgrass researchers to develop and test salt-tolerant turfgrass varieties. Establishing turfgrass from seeds under saline conditions is becoming increasingly important, particularly in arid regions of the world. The first step in the process of screening for salinity tolerance in any plant species is to conduct a germination test. According to protocols outlined by the Association of Official Seed Analysts (AOSA, 2009), media considered acceptable for use in standard germination tests include blotter paper, creped cellulose paper, filter paper, sponge rock, vermiculite, sand, and paper toweling. However, it is not clear whether all of these substrata are adequate for screening purposes when the germination test is conducted under saline conditions or when coated seeds are used. A wide variety of substrata have been used to investigate the effect of salinity on germination of turfgrasses. Dewey (1962) used salinized soil to test wheatgrass (Agropyron desertorum Fisch.) germination at 6,000, 12,000 and 18,000 ppm NaCl and CaCl2. Harivandi et al. (1982) studied germination of several cool-season grasses on washed sand and germination pads (blotter paper). The sand substrata can be easily used; however, it has little water-holding capacity, which may cause salts to concentrate over the course of the experiments (Dudeck et al., 1986). Others have used hydroponic culture systems to study the effect of salinity on germination and growth of a number of cool-season grasses (Brilman and Sardar, 2010; Horst and Beadle, 1984; Horst and Dunning, 1989; Horst and Taylor, 1983; Richardson and McCalla, 2008; Wu, 1981; Zhang et al., 2011). Although these studies reported differences in salt tolerance among the turfgrasses tested, the saline aqueous environment in which the seeds were exposed may not accurately simulate saline conditions occurring in soil. Moreover, hydroponic culture experiments are very labor-intensive (Dudeck et al., 1986). Agar gel was first used as a germination medium by Torello and Symington (1984) to test the effect of salinity (10,000 ppm) on germination of several kentucky bluegrass, alkali grass [Puccinella distans (L.) Parl.], and red fescue varieties. Since then, other researchers have used agar to study the effects of various salinity levels on germination of a number of warm- and cool-season grasses (Dai et al., 2009; Dudeck and Peacock, 1985; McCarty and Dudeck, 1993; Peacock and Dudeck, 1989; Wang and Zhang, 2010; Zhang et al., 2011). However, with the exception of Wang and Zhang (2010) and Zhang et al. (2011), only low salinity levels were tested in the aforementioned studies. Blotter or filter paper has also been widely used as a germination medium to screen for salinity tolerance in turfgrasses (Camberato and Martin, 2004; Johnson et al., 2007; Lunt et al., 1961; Marcar, 1987; Qian and Suplick, 2001; Serena et al., 2010; Shahba et al., 2008; Zhang et al., 2011). As reported by Dudeck et al. (1986), blotter paper tends to lose more water than agar media and salinity levels surrounding the seed may be higher in blotter paper than in agar.

Seed coatings were first introduced in the late 1930s and early 1940s (Kaufman, 1991; Ross and Moore, 1975) to improve seeding and establishment of vegetables. More recently coatings have been used to improve the microenvironment of turfgrass seeds (Scott, 1989). Grass seeds have been coated with a number of materials, including limestone (Hathcock...
et al., 1984a), fertilizers (Bruneau et al., 1989; Hathcock et al., 1984b), diatomaceous earth and cytokinins (Greipson, 1999), fungicides (Hummel, 1991; Newell, 1997; Newell et al., 1999), and water adsorvents (Berdhal and Barker, 1980), resulting in increased germination and/or establishment of turf. Since its introduction in 2006, the corn starch coating by the name of ZEBA™ has been promoted for use on warm- and cool-season turfgrass seeds (Ardens, 2007) for its water retention capabilities. The coating consists of a polymer [starch-g-poly (2-propanamide-co-propanoic acid) potassium salt], which, according to the manufacturer, absorbs water up to 400 times its weight (Absorbent Technologies, 2006; Ardens, 2007). Leinauer et al. (2010) documented that ZEBA® coating resulted in increased field emergence of ‘Bengal’ creeping bentgrass and in improved establishment of several cool-season blends and mixes. The authors also showed that increased moisture retention from seed coating could compensate for reduced seeding rates, reduced irrigation during establishment, and for a combination of both. However, it is not known if seed coating will have similar beneficial effects on germination under saline conditions.

To identify the optimal germination medium for salt tolerance screening of turfgrasses, more studies are needed that compare media, species, and seed treatments. Dudeck et al. (1986) compared germination of perennial ryegrass (Lolium perenne L.) on blotter paper vs. agar at salinity levels of up to 5000 ppm. Although the authors found no differences between the two media, only one species was tested, and moderate salinity levels were used. Zhang et al. (2011) compared germination of several varieties of four cool-season species using a maximum salt concentration of 20 g·L⁻¹ NaCl and three different media. To date, no studies have compared germination of both warm- and cool-season grasses under highly saline conditions on more than one medium. In addition, studies are lacking on how seed coatings purported to increase moisture retention influence germination under a wide range of salinities. In this study, we compared germination of coated and uncoated cool- and warm-season grasses on two of the most widely used media under low [potable or electrical conductivity (EC) = 0.6 dS·m⁻¹] to highly saline conditions (EC = 22.5 dS·m⁻¹). The substrata chosen for the study were 1% agar gel and filter paper, and ZEBA® cornstarch material was used on the coated seeds. The objective of this study was to compare the performance of two commonly used media for germinating coated and uncoated turfgrass seed under saline conditions.

Materials and Methods

The study was conducted from Apr. 2011 to July 2011 at the New Mexico Department of Agriculture’s seed testing laboratory. The two germination substrates compared in the study were filter paper and agar. Three cool-season grasses, namely Kentucky bluegrass ‘Barrister’, tall fescue (F. arundinacea Schreb.) ‘Barvado’, and perennial ryegrass (Lolium perenne L.) ‘Premier II’, and two warm-season grasses, Bermuda grass (Cynodon dactylon L. Pers.) ‘Bargusto’ and seashore paspalum (Paspalum vaginatum O. Swartz) ‘Sea Spray’, were included in the study. Uncoated and ZEBA® coated seeds were provided for each species by the seed manufacturer.

The first germination experiment was conducted following the protocol for seed germination on filter paper outlined by the Association of Official Seed Analysts (AOSA, 2009). The five salinity levels tested were: 0.6 dS·m⁻¹ (tap water, control), 2.2 dS·m⁻¹ (saline groundwater from a local shallow aquifer), and three additional saline solutions (7, 12.5, and 22.5 dS·m⁻¹), which were prepared using tap water and sodium chloride (NaCl) and calcium chloride (CaCl₂) in a ratio by weight of 1:1 (Table 1). One hundred seeds were placed on each filter paper, which had been previously saturated with the desired saline water treatment. The filter papers were then transferred to a plastic 10 × 10-cm container (Hoffman Manufacturing Inc., Jefferson, OR), covered with a transparent lid, and the container was placed in the germination chamber.

The second germination experiment was conducted using a 1% Difco Bacto agar substrate (Dudeck and Peacock, 1985). Salinity of the agar medium was adjusted to the aforementioned salinity levels. The agar solution was autoclaved at 120 °C for 30 min before use. The agar was poured into 10 × 1.5-cm Fisherbrand® petri dishes to which 36 seeds were transferred (Dai et al., 2009). Coated and uncoated unsterilized seed from the same seed lots were used in both germination experiments. Petri dishes and plastic boxes were incubated in two different germinators (Stults Scientific Engineering Corp., Springfield, IL), one for warm-season grasses and a second for cool-season grasses. The incubators were programmed to maintain alternating 8 h light at 25 °C with fluorescent light (36 µmol·s⁻¹·m⁻²) and 16 h dark at 15 °C for cool-season grasses and 8 h light at 35 °C with fluorescent light (36 µmol·s⁻¹·m⁻²) and 16 h dark at 20 °C for warm-season grasses (AOSA, 2009). Experiments were conducted using a completely randomized design with four replications for each treatment combination. The positions of the petri dishes and plastic boxes were rotated inside the germinator three times per week to minimize shelf effects on the study. After germinated seeds were counted, water from the corresponding salinity treatment was added to any filter paper that had dried out. Whether blotter paper required rewetting was determined by pressing a finger firmly into the filter paper.

The moisture level in the filter paper was deemed sufficient if a film of water formed almost around the finger (AOSA, 2009). However, if the filter paper was dry and rewetting was necessary, only enough water was added to supply the needed moisture to the seeds but not enough to create puddling on the filter paper or the formation of film around the seeds (AOSA, 2009). Germination data were collected three times per week for 4 weeks. A seed was considered germinated when the root and shoot could be observed with the naked eye (AOSA, 2009; Qian and Suplick, 2001). Germination rate (%/d) was based on seedling counts taken three times per week (Maguire, 1962), and FGP (%) was based on the total number of germinated seeds counted after 28 d. Although FGP provides the total germination after the evaluation period of 4 weeks, GR describes the rate of germination with higher values indicating faster rates. All values were subjected to analysis of variance (ANOVA) using SAS Proc Mixed (Version 9.2, SAS Institute Inc., Cary, NC) followed by multiple comparisons of means using Fisher’s protected least significant difference test at the 0.05 P level.

Results and Discussion

The ANOVA revealed that the three-way interactions among grass, germination media, and salinity and among grass, coating, and salinity had a significant effect on FGP (Table 2). The ANOVA further revealed that only the single effects of grass, salinity, and media affected GR. To address the significant three-way interactions, FGP was subsequently pooled over seed coating and is displayed separately for each grass and germination media (Table 3). Data were also pooled over germination media and are shown separately for each grass and seed coating (Table 3).

Final germination percentage. When data were pooled over seed coating and displayed separately for each grass species and germination media, differences in FGP between seeds germinated on filter paper and agar were observed in 17 of 25 treatment combinations (five grasses × five salinity levels). In all 17 combinations, germination was higher in agar than on filter paper (Table 3). Germination was greater in agar at all salinity levels for Kentucky bluegrass, perennial ryegrass, and tall fescue. Germination of bermudagrass and seashore paspalum was unaffected by the type of medium at any of the salinities except for 7 dS·m⁻¹. Our findings of generally higher germination in agar than on filter paper are in part supported by Zhang et al. (2011) who also reported higher germination under saline conditions.

### Table 1. Main chemical constituents of saline waters used in germination tests.

| Main constituents | Electrical conductivity (dS·m⁻¹) |
|-------------------|---------------------------------|
| pH                | 7.6                             |
| Bicarbonate (mg·L⁻¹) | 7.6                             |
| Calcium (mg·L⁻¹)  | 1.5                             |
| Sodium (mg·L⁻¹)   | 1.5                             |
| SAR               | 1.5                             |
Salinity had no effect on final germination percentage for perennial ryegrass when grown on agar and did not vary significantly across salinities levels ranging from 0.6 to 12.5 dS m\(^{-1}\) when germinated on filter paper. Final germination percentages exceeded 75% across all salinity levels and both media (Table 3). These results support the findings of Dudeck and Beadle (1985), Harivandi et al. (1982), and Horst and Dunning (1989) who all found that germination of perennial ryegrass was tolerant to salinity, even to levels found in sea water (Harivandi et al. 1982). Tall fescue exhibited a similar tolerance to salinity with final germination greater than 75% at all salinity levels and media. Our findings differ from those of Horst and Beadle (1984) and Zhang et al. (2011) who reported declining germination with increasing salinity in tall fescue. However, these authors used different methodologies to test salt tolerance: Zhang et al. (2011) used NaCl as the only source of salt and Horst and Beadle (1984) used floating mats in a saline solution. Hydroponic solutions and NaCl have both been shown to affect germination differently than other growth media (Zhang et al., 2011) or salts (MgCl\(_2\) or CaCl\(_2\)) (Lunt et al., 1961; Wu, 1981).

Kentucky bluegrass and bermudagrass maintained high germination in agar across all salinities, but germination dropped below 75% on filter paper at the highest salinity level of 22.5 dS m\(^{-1}\), exhibiting values of 44% and 69%, respectively (Table 3). Final germination percentage of seashore paspalum was low at all salinities tested, ranging from ≈60% at levels lower than 7.0 dS m\(^{-1}\) to 14% at 22.5 dS m\(^{-1}\) (Table 3). Our findings are similar to those of Johnson et al. (2007), who reported a germination percentage of 65% for ‘Sea Spray’ seashore paspalum at salinity levels of 2.0 and 3.0 dS m\(^{-1}\). Shahba et al. (2008) documented generally low germination under highly saline conditions for halophytes, which, according to Dodd and Donovan (1999), can be explained by the high sensitivity of halophytic grasses to osmotic stress during the germination stage. Carpenter (1958) attributed the generally low germination of seashore paspalum to self-incompatibility and the resulting high percentage of sterile seeds.

When data were pooled over germination media but analyzed separately for each grass, seed coating affected FGP negatively in only two of 25 treatment combinations (five grasses × five salinity levels). Coated seashore paspalum and perennial ryegrass seeds exhibited increased germination at four of the five salinity levels. Germination of coated tall fescue seeds was higher than for uncoated seeds at 0.6 and 12.5 dS m\(^{-1}\) (Table 3). Seed coating had no effect on FGP of bermudagrass at any salinity levels (Table 3). Kentucky bluegrass was the only grass tested that showed reduced germination from coated seed, but only at two of the five salinity levels (0.6 and 7.0 dS m\(^{-1}\)). Tall fescue and perennial ryegrass maintained FGP greater than 75% across all salinities and seed coatings as did bermudagrass and Kentucky bluegrass at salinities between 0.6

### Table 2. Results of analysis of variance testing the effects of grasses, seed coating, germination media, salinity, and their interactions on final germination and germination rate.

| Coating | Final germination | Germination rate |
|---------|------------------|-----------------|
| *Coating* | *** | NS |
| Grass | *** | *** |
| *Coating* | *** | NS |
| Media | *** | *** |
| Coating*media | NS | NS |
| Grass*media | *** | NS |
| *Coating* | *** | NS |
| Salinity | *** | * |
| *Coating*salinity | ** | NS |
| Grass*salinity | *** | *** |
| *Coating* | ** | NS |
| Media*salinity | *** | NS |
| Grass*media*salinity | *** | NS |
| Coating*grass*media | NS | NS |
| Coating*grass*media*salinity | NS | NS |

*Significant F test at the 0.05 level of probability.
**Significant F test at the 0.01 level of probability.
***Significant F test at the 0.001 level of probability.
NS = Nonsignificant at the 0.05 P level.

### Table 3. Final germination (%) of coated and uncoated bermudagrass, Kentucky bluegrass, perennial ryegrass, seashore paspalum, and tall fescue seed in two media (agar and filter paper) at salinities of 0.6, 2.2, 7.0, 12.5, and 22.5 dS m\(^{-1}\).

| Salinity (dS m\(^{-1}\)) | Germination media | Seed coating |
|--------------------------|-------------------|--------------|
|                          | Agar               | Paper        | Coated | Uncoated |
| Bermudagrass             |                   |              |        |          |
| 0.6                      | 94 a*              | 90 a         | 91 ab  | 93 a     |
| 2.2                      | 94 a               | 89 a         | 92 a   | 91 ab    |
| 7.0                      | 89 abA             | 80 b         | 85 bc  | 84 b     |
| 12.5                     | 83 b               | 84 ab        | 82 c   | 86 b     |
| 22.5                     | 73 c               | 69 c         | 70 d   | 72 c     |
| Kentucky bluegrass       |                   |              |        |          |
| 0.6                      | 93 aA              | 82 abB       | 83 aB  | 92 aA    |
| 2.2                      | 91 aA              | 84 aB        | 87 a   | 89 a     |
| 7.0                      | 92 aA              | 83 aB        | 84 ab  | 91 aA    |
| 12.5                     | 89 abA             | 77 bB        | 84 a   | 83 b     |
| 22.5                     | 86 bA              | 44 cB        | 67 b   | 63 c     |
| Perennial ryegrass       |                   |              |        |          |
| 0.6                      | 96 A               | 89 aB        | 95 A   | 90 B     |
| 2.2                      | 96 A               | 91 aB        | 96 A   | 91 B     |
| 7.0                      | 96 A               | 90 aB        | 94     | 91       |
| 12.5                     | 96 A               | 89 aB        | 96 A   | 90 B     |
| 22.5                     | 96 A               | 84 bB        | 93 A   | 87 B     |
| Seashore paspalum        |                   |              |        |          |
| 0.6                      | 64 a               | 61 a         | 69 aA  | 55 aB    |
| 2.2                      | 59 a               | 61 a         | 66 abA | 54 aB    |
| 7.0                      | 60 aA              | 50 bB        | 59 bc  | 52 a     |
| 12.5                     | 46 b               | 50 b         | 54 cA  | 22 bB    |
| 22.5                     | 14 c               | 14 c         | 20 dA  | 8 cB     |
| Tall fescue              |                   |              |        |          |
| 0.6                      | 88 A               | 82 B         | 88 A   | 83 abB   |
| 2.2                      | 88 A               | 81 B         | 84     | 85 a     |
| 7.0                      | 88 A               | 84 B         | 86     | 86 a     |
| 12.5                     | 88 A               | 80 B         | 86 A   | 81 bB    |
| 22.5                     | 88 A               | 81 B         | 86     | 84 ab    |

*Values followed by the same letter are not significantly different from one another (Fisher’s protected least significant difference, α = 0.05). Lower case letters denote differences between salinity levels separately for each grass (in columns).

*Upper case letters denote differences of the mean values between germination media (agar and paper) and between coated and uncoated seed (in rows).
and 12.5 dS m⁻¹. Final germination percentage for seashore paspalum improved greatly at the highest salinity levels when coated seed was used, increasing from 22% to 54% at 12.5 dS m⁻¹ and from 8% to 20% at 22.5 dS m⁻¹ when coated seed was compared with uncoated (Table 3). When potable water was used, coating had a negative effect on germination of Kentucky bluegrass and a positive effect on tall fescue (Table 3). These results differ from those reported by Richardson and Hignight (2010), who found that ZEBA coating had no effect on seedling emergence of tall fescue in three soil types or on Kentucky bluegrass emergence in two of three soil types. Contrary to Richardson and Hignight (2010), Leinauer et al. (2010) found increased field emergence of coated ‘Bengal’ creeping bentgrass and improved establishment of several cool-season blends and mixes when seed was coated. However, no other published reports are currently available on the effect of seed coating on the germination, emergence, or establishment of warm- and cool-season grasses.

Germination rate. All main effects except seed coating affected GR (Table 2). Germination rates for perennial ryegrass and bermudagrass were highest (13% and 12%/d, respectively) followed by tall fescue and Kentucky bluegrass (9% and 7%/d, respectively). Seashore paspalum showed the lowest germination rate, averaging 4%/d (Table 4). Our findings support those of Peacock and Dudeck (1989) who also found GR of 13%/d for bermudagrass at salinity levels as high as 9 dS m⁻¹. When data were pooled overall grasses, media, and seed coatings, and listed separately for each salinity level (Table 5), salinity levels ranging from 0.6 to 12.5 dS m⁻¹ did not affect GRs. Only the highest salinity treatment of 22.5 dS m⁻¹ reduced GR to an average of 7%/d (Table 5). These results support those of Camberato and Martin (2004), who reported decreased GRs with increasing salinity for rough bluegrass. Our findings are also similar to those of Horst and Dunning (1989) who found that perennial ryegrass exhibited a 20% GR reduction at 23 dS m⁻¹ when compared with tap water.

Generally, germination rates were higher on agar than on paper, averaging 10% and 8%/d, respectively (Table 6). This differs from results reported by Zhang et al. (2011) who found no differences in germination rates between media for several cool-season grasses. One possible reason could be that in contrast to our study, the authors used NaCl as the sole source of salinity. Several authors (Camberato and Martin, 2004; Dudeck et al., 1986; Zhang et al., 2011) have reported on the tendency of filter paper to lose water. In our study, we observed dew formation on the lids of the plastic containers; this water evaporation resulted in the filter paper drying out. The repeated rewatering of dry filter papers with saline water is required by the testing protocol (AOSA, 2009) and may have increased salinity levels over time. Agar plates did not require such a procedure. The potential increase in salinity caused by frequent rewatering of filter papers was also mentioned by Dudeck et al. (1986) and Zhang et al. (2011).

Our study suggests that the choice of medium can influence the outcome of germination tests and that results will also vary depending on the salinity level tested and whether the seed are coated. Filter paper is currently the medium recommended by the Association of Official Seed Analysts, but agar improved both FGP and GR for most turfgrasses in several of the salinity levels investigated. Germination of turfgrasses investigated in our study varied widely under salinities ranging from 0.6 to 22.5 dS m⁻¹. Seashore paspalum, a species described by numerous researchers as a halophyte with superior salt tolerance, exhibited poorest germination under both potable and saline conditions. Despite the significant differences in germination between bermudagrass and seashore paspalum reported in this study, no corresponding differences in establishment have been observed between these two grasses when grown under saline irrigation (Schiaon, unpublished data). This suggests that germination tests alone may not be good predictors of salinity tolerances during establishment. The opposite also appears to be true in that plants such as seashore paspalum, which are well adapted to saline conditions once established, may be very sensitive to salinity at the germination stage. Furthermore, the question regarding which of the two media is more representative of field conditions remains unanswered. Additional studies are needed to compare laboratory germination to field emergence, establishment, and long-term performance under saline conditions. Only studies that include both germination and field trials will help to determine the usefulness of germination tests in providing relevant information on establishing and maintaining turf under saline conditions.

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### Table 4. Germination rates (%/d of five turfgrass species. a

| Grass         | 0.6 | 2.2 | 7.0 | 12.5 | 22.5 |
|---------------|-----|-----|-----|------|------|
| Bermudagrass  | 12 a| 10 a| 9 a | 9 a   | 7 b  |
| Kentucky bluegrass | 7 b | 10 a| 9 a | 9 a   | 7 b  |
| Perennial ryegrass | 13 a| 10 a| 9 a | 9 a   | 7 b  |
| Seashore paspalum | 4 c | 10 a| 9 a | 9 a   | 7 b  |
| Tall fescue   | 9 b | 10 a| 9 a | 9 a   | 7 b  |

Data are pooled over five salinity levels, two seedling media, two seed coats, and four replications, and represent an average of 80 data points.

Values followed by the same letter are not significantly different from one another (Fisher’s protected least significant difference, α = 0.05).

### Table 5. Germination rates (%/d of several turfgrasses at five salinity levels. a

| Salinity (dS m⁻¹) | 0.6 | 2.2 | 7.0 | 12.5 | 22.5 |
|-------------------|-----|-----|-----|------|------|
| Bermudagrass      | 10 a| 10 a| 9 a | 9 a   | 7 b  |
| Kentucky bluegrass | 10 a| 10 a| 9 a | 9 a   | 7 b  |
| Perennial ryegrass | 10 a| 10 a| 9 a | 9 a   | 7 b  |
| Seashore paspalum | 9 a | 9 a | 9 a | 9 a   | 7 b  |
| Tall fescue       | 9 a | 9 a | 9 a | 9 a   | 7 b  |

Data are pooled over five turfgrasses, two media, two seed coatings, and four replications, and represent an average of 80 data points.

Values followed by the same letter are not significantly different from one another (Fisher’s protected least significant difference, α = 0.05).

### Table 6. Germination rates (%/d of several turfgrasses on two germination media. a

| Media       | 0.6 | 2.2 | 7.0 | 12.5 | 22.5 |
|-------------|-----|-----|-----|------|------|
| Agar        | 10 a| 10 a| 9 a | 9 a   | 7 b  |
| Paper       | 8 b | 9 a | 9 a | 9 a   | 7 b  |

Data are pooled over five turfgrasses, five salinity levels, two coatings, and four replications, and represent an average of 200 data points.

Values followed by the same letter are not significantly different from one another (Fisher’s protected least significant difference, α = 0.05).
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