Physiological Responses of Turfgrass Species to Drought Stress under High Desert Conditions

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Abstract. Broad concerns over water shortages and drought where irrigated urban landscapes are common in high desert regions have focused attention on drought tolerance of turfgrass species. We investigated the physiological responses of kentucky bluegrass (KBG) and tall fescue (TF) under a prolonged drought under high desert conditions. The experimental design was a split plot with three replicates. Two irrigation treatments as a whole plot—well-watered and no water—were applied to subplots of ‘Midnight’ KBG and ‘Gazelle’ TF. Stomatal conductance (_gs_), canopy temperature, and predawn leaf water potential were measured over two seasons. KBG _gs_ and leaf water potential decreased faster and to a greater extent than TF in response to soil drying and KBG was in complete dormancy and brown within 5 weeks after cessation of irrigation. By contrast, TF maintained a green canopy throughout the drought periods. In the no-water plots, TF appeared to consume water from the deepest measured soil profiles (80- to 100-cm depth), whereas KBG used most of the water in the 50- to 60-cm depths. When watered for recovery in late summer, KBG plots were mostly green within 3 weeks after rewetting. The surface temperature of the well-watered plots was 6–13 °C cooler than the no-water plots and TF showed _5–7 °C lower temperature than KBG in no-water plots. TF is suitable for deep soil, exploiting a larger volume of water to avoid drought, whereas KBG’s rapid drought avoidance would likely perform better in shallow landscape soils under drought.

Turfgrass species used in the IMW avoid drought but differ in mechanisms. TF avoids drought because it maintains normal physiological function in water-limiting conditions by developing an extensive, deep root system to extract more water from a deeper and greater volume of soil (Huang and Gao, 2001). This postpones tissue dehydration (Scheffer et al., 1987). TF appears to also reduce water loss from transpiring leaves by rolling its leaves as soil water content declines (Qian and Fry, 1997). By contrast, KBG avoids drought by entering summer dormancy (Ervin and Koski, 1998), sometimes referred to as quiescence. However, once adequate moisture is again available, plants will resume active growth (Laude, 1953).

Osmotic adjustment is another drought tolerance mechanism that grasses use to maintain cellular turgor and allow them to take up water at lower soil water potentials (Perdomo et al., 1996; White et al., 1992). Osmotic adjustment under stress conditions has been reported to occur in both TF (Qian and Fry, 1997; West et al., 1990; White et al., 1992) and KBG (Jiang and Huang, 2001; Perdomo et al., 1996).

Grass responses during prolonged summer drought have long been studied. Most research has been conducted in greenhouses under controlled conditions (Aronson et al., 1987; Brown et al., 2004; Qian and Fry, 1997), whereas some were field investigations (Carrow, 1996; Laude, 1953; Richardson et al., 2008). Traits used to measure drought response have more commonly included morphological responses such as growth reduction, turfgrass quality rating, and root density (Ervin and Koski, 1998; Qian and Fry, 1997; Scheffer et al., 1987). Less often but more recently, physiological responses such as water relations, _gs_, photosynthesis, and hormone (abscisic acid) concentration have been measured (Jiang and Huang, 2000; Perdomo et al., 1996; Volare et al., 2009; West et al., 1990).

Because the mechanisms that KBG and TF use to cope with drought are quite different, a comparison under common field conditions, with detailed measurements, will help us understand the distinct drought tolerance or avoidance mechanisms used by these grasses. The objective of this work was to compare the physiological responses of KBG and TF, which differ in drought-coping mechanisms, that might contribute to persistence of field-grown grasses during a prolonged drought in the IMW.

Materials and Methods

Field plot. ‘Midnight’ KBG (Poa pratensis L.) and ‘Gazelle’ tall fescue (Schedonorus phoenix (Scop.) Holub) were planted at the Greenville Research farm in North Logan, UT (lat. 41°45′ N, long. 111°48′ W) in 2003. Temperature and precipitation data at the experimental site are summarized in Figure 1. The soil at the experimental site was a silt loam, Millville series of uniform depth with a pH of 7.5–8.2 (Abdu et al., 2007). The experimental design was a split

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plot with six main plots of 3 × 6 m each divided into six subplots, each 1.5 × 2 m. Subplots were randomly assigned within each main plot and planted with a different turfgrass species.

Two irrigation treatments were applied to the whole plot area: 1) well-watered, irrigated three times a week with 1.2 cm of water; and 2) no-water plots that did not receive irrigation after 12 June in 2007 and 25 June in 2008. Well-watered plots were irrigated by hand to ensure uniformity and grasses were mowed at 7.62 cm and fertilized with ammonium sulfate at a rate of 9.8 kg·ha⁻¹, which was applied per year to the plots—half in the spring and half in the fall.

Plots were 4 years old and well established when measurements were begun. In 2007, measurements were made twice each week from 13 June to 22 Aug. and then irrigation was resumed for 3 weeks to recover from prolonged drought. In 2008, measurements were made twice each week from 26 June to 22 Aug., and then irrigated to allow recovery for 4 weeks.

Measurements. Predawn leaf water potential measurements were made twice a week in 2007 and once a week in 2008 using a pressure chamber (Model 3005HGPL; Soil, Moisture Equipment Corp, Santa Barbara, CA). At predawn, five stems of each species in each subplot were collected by pulling the entire plant including the root, then immediately wrapping them in plastic wrap and storing in a bag filled with ice for transport to the laboratory. Stems were cut slightly above the root and placed in the pressure chamber. Nitrogen gas was slowly applied to increase the chamber’s atmospheric pressure until water appeared at the cut end of the stem. The pressure reading was then taken and used as leaf water potential.

Stomatal conductance measurements were made twice a week in 2007 and once a week in 2008 using a leaf porometer (Model SC-1; Decagon Devices, Inc., Pullman, WA). Measurements were taken between 1100 HR and 1400 HR on a clear day. Four to five blades of grass in each subplot were excised and arranged before clamping side by side with the adaxial side of the leaves facing the porometer chamber. Time used to prepare leaves for measuring after excising the leaves was less than 5 s to prevent the effect of water discontinuity on stomata. Stomata closure in lucerne occurred within 2 h after cutting for hay and when the relative water content in tall fescue leaves remained between 80% and 90%, the gs would not be affected by cutting (Harris and Tullberg, 1980). By using the instrument’s automatic mode, the gs was measured in 30 s. Eight separate measurements were made from each subplot and averaged for a final value.

Surface temperature of each subplot was measured using a digital thermometer (Model 52-II Dual Input Digital Thermometer; Fluke Corporation, Everett, WA) connected with infrared (IR) temperature sensors (Model SI-111; Apogee Instruments, Inc., Logan, UT) after measuring gs at 1400 HR MDT. The IR temperature sensor was held 1 m above the canopy perpendicular to the ground allowing a field of view of 2 m in diameter.

Ambient air temperature data were continuously collected by a weather station in Greenville Research farm with a combination temperature and humidity sensor (Model CR500; Campbell Scientific, Logan, UT). The sensor was scanned every 10 s and averages were recorded every 30 min with a data logger (Model CR1000; Campbell Scientific). VPD and LAVPD were calculated using ambient air temperature, dewpoint temperature, and leaf temperature as described by Murray (1967).

VPD is the difference between saturation vapor pressure and actual vapor pressure of ambient air (e_s – e), whereas LAVPD was calculated from the difference between saturation vapor pressure of the leaf using leaf temperature and actual vapor pressure of the ambient air (e_s – e).

Volumetric soil water content (VWC) was measured in 2008 using a frequency domain reflectometry (FDR) sensor (Diviner 2000; Sentek Sensor Technologies, Adelaide, Australia). One-meter long polyvinyl chloride ride tubes were installed in the center of each subplot. Every day at 1600 HR, the Diviner 2000 probe was inserted into each access tube to measure water content to a depth of 100 cm (the deepest measurement of this study) with measurements made at 10-cm intervals.

Percent water use in each depth (10-cm interval) was calculated by the following equation:

\[
\% \text{ water use each depth (10 cm)} = \frac{VWC_{\text{initial}} - VWC_{\text{end}}}{\sum_{i=1}^{n} VWC_{\text{initial}} - VWC_{\text{end}}} \times 100\%
\]

where VWC_{initial} is the volumetric soil water content at the beginning of the experiment and VWC_{end} is the volumetric soil water content at the end of the experiment.

Statistical analysis. The experiment was a split plot design with six whole plots with two treatments applied (well-watered and no-water treatments) and three replicates of each. Treatment effects, species differences, and treatments × species interactions were determined by analysis of variance according to the mixed procedure of SAS (Version 9.0; SAS Institute, Cary, NC). Thus, fixed parameters were species × treatments, whereas random parameters were replicates, replicates × treatments, and replicates × species × treatment. Mean differences were tested with least significant difference test at a probability level of 0.05. Slope comparison was tested using the GLM procedure of SAS.
Results

The drought responses of the grasses varied slightly as a result of the difference in weather conditions between 2007 and 2008 (Fig. 1). Average air temperature in 2007 was ≈1 to 2 °C higher than in 2008; however, average air temperature of both years was 2 to 3 °C higher than the 30-year average. Moreover, total rainfall during both years was significantly lower than the 30-year average (Fig. 1). It should be noted that during the study period, there were several rainfall events with minimal effect. Recently, there was a study in the same field, which indicated that it took greater than 13 mm of precipitation to restore active growth in turfgrass (unpublished data).

Because irrigation was withheld in the no-water plots, KBG entered summer dormancy in 5 weeks and all aboveground tissues were brown, whereas TF remained green with some browning for the whole period of each experiment in both years. Each year, gs of KBG in well-watered and no-water plots was equal at the beginning of the experiment but in the no-water plots, conductance decreased rapidly after irrigation stopped (Fig. 2). Stomatal conductance differed from well-watered plots by Week 1 in both years. Unlike KBG, gs of TF in no-water plots was significantly lower than in well-watered plots by Week 2 in both years. Irrigation was resumed on the no-water plots in August to end dormancy and restore active growth. The spike in Figure 2E at Day 48 was caused by 88 mm of rainfall (Fig. 1B) stimulating a rapid increase in gs in no-water TF but not in no-water KBG.

After resumption of irrigation, gs of no-water KBG equalized with that of well-watered plots after 3 weeks in 2007 and 4 weeks in 2008. TF recovered to the same degree of gs after 2 weeks in 2007 and after 4 weeks in 2008. Overall, gs of KBG dropped lower and faster than that of TF in both years. However, it took about the same length of time for KBG and TF to recover from prolonged drought.

Predawn leaf water potential (ψleaf) followed a similar trend to gs. As soil became drier, ψleaf of KBG was more negative and significantly lower than well-watered ψleaf by Week 2 in both years. KBG entered complete summer dormancy when ψleaf dropped to -2.0 MPa. In contrast, the ψleaf of TF in no-water plots was significantly different from those in well-watered plots by Week 3 in 2007 and by the first week in 2008. The sharp decrease of ψleaf in Day 43 in 2008 (Figs. 2D and 2H) was the result of the difficulty of measurements the week before (Day 36) resulting in no data on that day, the initial week for KBG entering dormancy in 2007 and a rapid decline in internal water potential (Fig. 2B). During the recovery period, ψleaf of both species increased (less negative) to well-watered plot levels after 2 weeks of recovery in 2007 and 3 weeks in 2008.

Surface temperature at midday of no-water vs. the well-watered plots was ≈6 to 13 °C higher in both species, typical of drought-stressed turf, and manifested in a higher leaf and air temperature (Tleaf - Tair) difference in no-water than in well-watered plots (P = 0.0008) (Fig. 3A). Tleaf - Tair in well-watered plots declined similarly in both species as VPD increased as a result of evaporative cooling. This well-established inverse baseline relationship between Tleaf - Tair and VPD for dense, uniform crops (Idso, 1982) has been conceptually refined (Blonsquist et al., 2009) and applied to cool season turfgrass (Martin et al., 2005). In the no-water plots, drought-induced stomatal closure (Fig. 2A–B) reduced evaporative cooling in both species. However, KBG Tleaf - Tair was higher than that of TF across all VPD levels as KBG entered dormancy and lost stomatal function.

In the well-watered plots, gs decreased as the LAVPD increased with no difference in slope (P = 0.204) of both species (Fig. 3B). As drought stress became more severe, LAVPD was greater as a result of stomatal closure, leading to an increase in leaf temperature, in turn causing greater differences in LAVPD (Fig. 3C). However, in no-water plots, the depth of LAVPD increased, indicated by the slope in Figure 3C, which was also not significantly different (P = 0.313) in both species at LAVPD less than 5 kPa. At LAVPD above 5 kPa, TF stabilized gs at ≈150 mmol m⁻² s⁻¹, whereas gs in KBG dropped to lower levels and continued to decrease over a progressively higher range of LAVPD levels than TF until complete gs closure.

The rapid and progressive stomatal closure in KBG in response to soil drying is delineated more sharply when related to water potential (Fig. 4). The ratio of well-watered to no-water of leaf water potential vs. gs of both species showed a similar trend, decreasing as drought-induced gs declined. This figure is similar to that of Kjelgren et al. (2009) but with the well-watered vs. no-water ratio inverted. Stomatal conductance of TF in no-water plants declined along with declining leaf water potential but no-water gs did not fall below 30% of well-watered plants (ratio of well-watered to no water was not more than 3) while maintaining ψleaf above 50% of well-watered plants. This trend was initially apparent in KBG as well, up to a certain depth, then it declined more rapidly and absolute leaf water potential decreased more rapidly in KBG than TF.

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The greatest water use of plants in no-water plots was indicated by the greatest depletion to the lowest percent volumetric soil water content over the course of the study period (Figs. 5 and 6). KBG used more water from the soil at depths between 0 and 90 cm, whereas TF used water down to 100 cm (greatest depth measured in this study) (Fig. 6); the large depletion at 100 cm suggested that TF likely extracted water below the measured root zone. Both species extracted more water at the surface (0 to 30 cm) than at the deeper soil profile, but TF depleted more water at the deepest depths than KBG. In addition, the total amount of water used by TF was 9% higher than by KBG. To better evaluate which depths contributed the most to variation in gs, stepwise regression was used to relate soil depth as independent variables (Xs) and gs as the dependent variable (Y). In well-watered plots, the shallow depths (0 to 20 cm) contributed most to gs in TF (R² = 0.033 and R² = 0.411) and at 60- to 80-cm depths in KBG (R² = 0.014 and R² = 0.509). However, the relationship of depth and gs in well-watered plants may not be meaningful as a result of low R². However, in the no-water plots, the analysis indicated that the deepest depths (80 to 100 cm) are most important for TF regarding variation in gs (R² < 0.0001 and R² = 0.993), whereas the 40- to 60-cm depth for KBG (R² = 0.0003 and R² = 0.991) are most indicating (Table 1). The stepwise regression supported evidence in Figures 5 and 6 indicating that TF extracted more water from the deeper soil and became water stressed when that layer was depleted, whereas KBG gs was more sensitive to soil water depletion in the top soil layer.

Discussion

Turfgrass has limited stomatal control over transpiration as a result of low height and a thick boundary layer where the leaf surface is completely decoupled from conditions in the air outside the boundary layer (Javis and McNaughton, 1986). Thus, the evapotranspiration continued even when stomata closed (Harris and Tullberg, 1980), during which water is plausibly lost through the cuticle (Cowan, 1977) resulting in rapid water depletion in soil.

As the soil dried, both TF and KBG initially approached drought stress at the same rate, indicated from gs reduction (Figs. 2 and 4). In addition, both species did not moderate internal water potential through stomatal closure, which means the rate of water potential decline in plants was rapid even when stomatal closure occurred immediately after withholding irrigation. Rapid decline in water potential was likely the result of greater boundary layer control over total transpiration from the canopy than stomatal aperture (Zhang et al., 2007).

Both species showed different responses when the soil dried and conditions became severe. KBG stomata seemed to be very sensitive to soil drying, because they closed more rapidly and absolute leaf water potential fell more rapidly than TF. KBG rapidly went dormant with all aboveground tissues...
turning brown as a result from increasing $T_{\text{leaf}} - T_\text{air}$ (Fig. 3C) when $g_s$ fell below 50% of well-watered levels in potentially a feed-forward process (Fig. 3B). A small reduction in $g_s$ from soil drying will reduce transpiration cooling, increasing $T_{\text{leaf}} - T_\text{air}$ and LAVPD, and diminishing the boundary layer through increased eddy turbulence convection. Increased heating and a decreased boundary layer in turn would push conductance even lower and LAVPD higher in a feed-forward loop resulting in more and faster browning of tissues (Figs. 2 and 3). KBG appeared to reduce $\psi_{\text{leaf}}$ more than TF, suggesting that it can extract soil water at lower contents in the top layers to the point of triggering the feed-forward stomatal closure cascade. At that point, the plants became dormant—a drought tolerance mechanism to avoid greater physiological damage to meristematic tissue and roots from water stress (Fry and Huang, 2004). As irrigation was resumed, the rapid resumption of growth occurred indicating that the rapid KBG dormancy allowed the meristematic growing points to survive under severe stress until the first fall rains coupled with cooler temperatures under natural field conditions (Laude, 1953). The physiological recovery time in no-water plots in our study was 3 to 4 weeks after irrigation was resumed. Deeper rooting and maintenance of green foliage showed that TF did not reach the point over two growing seasons, one

Fig. 2. Stomatal conductance ($g_s$) and predawn leaf water potential in well-watered and no-water plots of ‘Midnight’ kentucky bluegrass (KBG) in (A, C) 2007 and (B, D) 2008 and of ‘Gazelle’ tall fescue (TF) in (E, G) 2007 and (F, H) 2008. Well-watered plots are represented by a short dash line and no-water plots a solid line. The vertical line indicates irrigation resumption on the no-water plots. ns represents non-significantly different ($P < 0.05$), whereas the asterisk represents significantly different ($P < 0.05$) between pairs of well-watered and no-water values at each day.
being exceptionally hot, where meristematic growing points were injured. Thus, when the water resumed, rapid resumption of normal growth occurred almost at the same time as KBG.

TF could be classified as a drought-evader (Fry and Huang, 2004). It kept the above-ground tissues green because the deep root system allowed the plants to extract more water from deeper soil (Figs. 5 and 6). Initially TF responded to soil drying similarly to KBG but reached a steady state between reduced transpiration and higher temperature (convective heat dissipation). Drought-stressed TF thus maintained open stomata at ≈30% of well-watered levels and presumably continued photosynthesis that appeared to be in a steady-state balance with water potential at approximately half of well-watered levels (Fig. 4) through extracting from increasingly deeper soil depths (Figs. 5 and 6).

In several studies, TF has been shown to have significantly more root mass, as measured by root length, at all depths under drought conditions compared with KBG (Ervin and Koski, 1998; Sheffer et al., 1987) and had three to 12 times greater root length in the lower profile (60 to 80 cm) in the field condition (Su et al., 2008). This larger root system and greater ability to obtain water enable TF to maintain consistent rates of transpiration resulting in cooler surface temperatures (Fig. 3C) during drought compared with KBG.

Under well-watered conditions, both species showed similar responses to the high desert environments during summer. When water became limited, TF showed the ability to extract water from deeper in the soil profile as water content in the top layer depleted (Table 1; Figs. 5 and 6). This allowed TF to maintain green leaves longer than KBG. TF relies on the deep root system to tolerate drought and may become fatally stressed in typically shallow urban landscape soils because the ability to avoid drought with deep rooting is lost. KBG, in contrast, goes dormant rapidly and preserves the growing point, which makes it more suitable for shallow soil. However, its rapid feed-forward descent into dormancy in response to emergence of localized soil drying from nonuniform irrigation application (Kjelgren et al., 2000) may result in overirrigation to avoid the browning of aboveground tissue.

Under high desert conditions of the IMW, TF has the ability to extract water from deep in the soil profile thereby minimizing irrigation as well as labor and other inputs associated with irrigation. Where water conservation and maintenance of green cover is the highest priority, TF may be better adapted than KBG as long as TF is able to root deeply. As a result, aboveground tissues of TF would stay green as long as there is water in the soil profile. However, TF is likely to perform less well if the soil is shallow and its root system is not able to penetrate deeply, causing soil water depletion to occur more rapidly. Therefore, in this case, KBG would be better suited because

Fig. 3. The difference between air and leaf temperature over the range of VPD in well-watered of both species (solid line), in no-water ‘Midnight’ kentucky bluegrass (KBG) (dash-dot-dot line), and in no-water ‘Gazelle’ tall fescue (TF) (short dash line) (A); and the relationship between leaf-to-air vapor pressure difference (LAVPD) and stomatal conductance (gs) in (B) well-watered plots and (C) no-water plots with the equation for LAVPD less than 5 kPa. The gs at LAVPD greater than 5 kPa was shown in two lines, in which the short dash line represented ‘Gazelle’ TF and the dash-dot-dot line represented ‘Midnight’ KBG. VPD = vapor pressure difference.

Fig. 4. The ratio of well-watered:no-water leaf water potential to stomatal conductance (gs). Kentucky bluegrass lines divided into the gs ratio lower than 2 (a short dashed line) and the gs ratio higher than 4 (a dash-dot-dot line) of gs.
it would go dormant rather than suffer damage as TF would be liable to do.

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**Table 1.** Correlation efficiency and *P* value of stomatal conductance (*g*<sub>S</sub>) with volumetric soil water content at 20-cm interval depths.

| Species × treatment | Soil depth (m) | 0–0.2 | 0.2–0.4 | 0.4–0.6 | 0.6–0.8 | 0.8–1.0 |
|---------------------|---------------|-------|---------|---------|---------|---------|
| Well-watered 'Midnight' KBG | 0.367* | 0.546 | 0.673 | **0.713*** | 0.570 |
| | 0.267 | 0.082 | 0.023 | **0.014** | 0.067 |
| Well-watered ‘Gazelle’ TF | **-0.642** | -0.488 | -0.331 | -0.019 | 0.306 |
| | 0.033 | 0.128 | 0.320 | 0.956 | 0.360 |
| No-water ‘Midnight’ KBG | 0.905 | 0.980 | **0.996** | 0.967 | 0.955 |
| | 0.034 | 0.003 | **0.0003** | 0.007 | 0.012 |
| No-water ‘Gazelle’ TF | 0.921 | 0.385 | 0.942 | 0.982 | 0.997 |
| | 0.023 | 0.008 | 0.002 | <0.0001 | <0.0001 |

*Correlation coefficient.

*P* value.

**Bold type indicates the depth with the greatest contribution to changes in *g*<sub>S</sub> using stepwise regression.**

*KBG = Kentucky bluegrass; TF = tall fescue.*
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