Growth and Physiological Traits Associated with Drought Survival and Post-drought Recovery in Perennial Turfgrass Species

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ABSTRACT. The objective of this study was to determine physiological traits for drought survival and post-drought recovery upon re-watering in two C3 perennial grass species, kentucky bluegrass [KBG (Poa pratensis)] and perennial ryegrass [PRG (Lolium perenne)]. Plants were maintained well watered or exposed to drought stress by withholding irrigation and were then re-watered in a growth chamber. KBG had significantly higher grass quality and leaf photochemical efficiency, and lower electrolyte leakage than PRG during 20 days of drought. After 7 days of re-watering, drought-damaged leaves were rehydrated to the control level in KBG, but could not fully recover in PRG. KBG produced a greater number of new roots, while PRG had more rapid elongation of new roots after 16 days of re-watering. Superior drought tolerance in KBG was associated with osmotic adjustment, higher cell wall elasticity, and lower relative water content at zero turgor. Osmotic adjustment, cell wall elasticity, and cell membrane stability could play important roles in leaf desiccation tolerance and drought survival in perennial grass species. In addition, post-drought recovery of leaf hydration level and physiological activity could be associated with the accumulation of carbohydrates in leaves and rhizomes during drought stress and new root production after re-watering.

Water availability for agricultural or urban irrigation is becoming increasingly limited, which is a common threat for plant growth in many areas. The mechanisms of drought tolerance in terms of plant productivity under moderate drought have been investigated extensively in many crop species, but mechanisms contributing to desiccation tolerance or drought survival in perennial grasses have received limited attention (Volaire and Lelievre, 2001). Improving the persistence and survivability of perennial species is critically important in areas with prolonged periods of drought. In addition, rapid recovery of damaged plant tissue and re-growth of new tissue following drought stress when water becomes available are also important in perennial grass management to ensure rapid stand re-establishment. Post-drought recovery is largely dependent on the recovery of existing leaf tissue and the regeneration of new tissue from crowns, stolons, and rhizomes of grass plants. Plant response to drought stress involves changes in various morphological and physiological factors (Nilsen and Orcutt, 1996; Shinozaki and Yamaguchi-Shinozaki, 1997). Leaf dehydration tolerance has been attributed to at least two mechanisms: osmotic adjustment (involving inorganic ions, carbohydrates, and organic acids), and changes in cellular/tissue elasticity [i.e., bulk elastic modulus (E)] (Touchette, 2006). Osmotic adjustment and high cell wall elasticity can modify the relationship between turgor pressure and cell volume, facilitating the maintenance of cell turgor and cell growth under drought stress (Blake et al., 1991; Saito and Terashima, 2004). Continuing leaf expansion and maintaining turgid leaves under drought stress are highly desirable traits for improving crop production during drought stress and rapid recovery from drought damages upon rewatering (Siopongco et al., 2006).

Drought tolerance and recuperative potential vary with grass species (Carrow, 1996; DaCosta and Huang, 2006; Su et al., 2008; Volaire and Gandoin, 1996). Kentucky bluegrass (KBG) and perennial ryegrass (PRG) are C3 perennial species widely used as turfgrass in cool climatic regions (Beard, 1973; Sun, 2008). It is generally believed that KBG has better drought tolerance than PRG, and some researchers have attributed the

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Materials and Methods

**Plant materials and growth conditions.** 'Midnight' KBG and 'Paragon GLR' PRG were examined. These cultivars of KBG and PRG were selected because both were reported to have good grass quality and drought resistance within the species (Hardebeck and Bigelow, 2008; Wang and Huang, 2004). Mature sods of KBG and PRG were collected from Rutgers University research farm, North Brunswick, NJ, and were planted in plastic containers (60 cm deep and 10 cm in diameter) filled with a mixture (1:1 v/v) of fine sand and silt loam soil (fine, montmorillonitic, mesic, Aquic, Arquidoll) that was sterilized before planting. Plants were maintained in a greenhouse for 40 d under 12 to 14 h natural light conditions with average level of photosynthetically active radiation (PAR) at 760 μmol·m⁻²·s⁻¹ and temperatures of ~23/17 °C (day/night), and were then moved to a walk-in growth chamber where treatments were imposed. The growth chamber (3 x 2.5 m) was set at 23/18 °C (day/night), 60% relative humidity, and a 12-h photoperiod with PAR of 550 μmol·m⁻²·s⁻¹ at the canopy level. Before drought stress treatment, plants in each container were watered three times per week to maintain soil moisture at field capacity and were fertilized biweekly with 100 mL of 20N–8.8P–16.6K soluble fertilizer (Peter’s General Purpose 20–20–20; Grace-Sierra, Milpitas, CA), including micronutrients at a concentration of 5 g·L⁻¹ in 100 mL of the soluble fertilizer (0.05% Mg, 0.0068% B, 0.0036% Cu, 0.05% Fe, 0.025% Mn, 0.0009% Mn, and 0.0025% Zn). The grass was hand-clipped weekly to maintain a height ~6 to 8 cm.

**Treatment and experimental design.** After a 14-d acclimation to growth chamber conditions, plants were exposed to three treatments: 1) in the well-watered control treatment, plants were watered three times per week until water drained from the bottom of the container at each irrigation; 2) in the drought stress treatment, irrigation was withheld until permanent leaf wilting occurred; 3) in the re-watering treatment, plants were initially exposed to drought stress by withholding irrigation until complete desiccation of grass canopy, but then were irrigated every 2 d days to allow for recovery from drought stress.

Treatments and species were arranged as a completely randomized design with two factors (watering treatments and species). Each treatment for each species was repeated in four replicates (four pots).

**Soil water status and physiological responses to drought and re-watering.** Soil volumetric water content (SWC) in a 0- to 20-cm deep soil layer of each container was measured as an indication of the level of soil water deficit using time domain reflectometry (Soil Moisture Equipment, Santa Barbara, CA) by inserting the 20-cm-long wave guide vertically in the top 20 cm soil profile.

Grass quality (GQ), leaf relative water content (RWC), leaf chlorophyll content (Chl), and leaf photochemical efficiency (Fv/Fm) were measured at 0, 5, 10, 15, and 20 d of drought stress and 3 and 7 d of re-watering. During 3 and 7 d of re-watering, these measurements were taken only on existing leaves that were previously exposed to drought stress to examine the recovery potential of stressed tissues.

Grass quality was rated based on a leaf color and turgidity level, which is a nondestructive parameter and the most widely used one in the evaluation of the overall performance and growth of a grass stand and survivability of perennial grass plants under stressful conditions (Beard, 1973; Turgeon, 1996). It was visually rated on a scale from 0 to 9, where the grass quality was rated as zero if leaves were completely brown and desiccated and rated as 9 when leaves were green and fully hydrated.

Leaf water status was determined by measuring RWC calculated as follows: RWC (%) = 100 × (FW – DW)/(TW – DW), where FW is the leaf fresh weight, DW is leaf dry weight for tissues dried at 85 °C for 3 d, and TW is turgid weight of leaves after being soaked in water for 4 h at ~20 °C (Barrs and Weatherley, 1962).

Cell membrane stability was estimated by calculating the percentage EL from leaf cells. About 10 first and second fully expanded leaves per container were excised and cut into 2-cm-long pieces. They were rinsed three times in distilled water and placed into vials containing 20 mL of distilled water. After shaking for 6 h, initial conductivity (Ci) of the bathing solution containing fresh leaves was measured with a conductivity meter (YSI, Yellow Spring, OH). Leaves were then killed in an autoclave at 140 °C for 30 min and were placed on a shaker for 24 h before maximum conductivity (Cmax) of the bathing solution was measured. The EL was calculated as 100 × Ci/Cmax (DaCosta et al., 2004).

Leaf photochemical efficiency expressed as the ratio of variable to maximum chlorophyll fluorescence (Fv/Fm ratio) was determined on five individual leaves in each container with a fluorescence induction monitor (Fim 1500; ADC Bioscientific, Hodderston, UK) after dark adaptation for 30 min. Leaf Chl content was extracted by soaking 0.05 to 0.1 g leaves in 10 mL of dimethyl sulfoxide in the dark for 72 h (Hiscox and
Absorbance of extracted Chl was measured at 663 and 645 nm using a spectrophotometer (Genesys 2; Spectronic Instruments, Rochester, NY) (Jiang and Huang, 2001). The content of total nonstructural carbohydrate (TNC) in leaves of both species, crowns in PRG and rhizomes in KBG, was analyzed following the method described in Nelson (1944) and Somogyi (1945), with some modifications. After plants were washed free of soil, leaves, crowns, or rhizomes were immediately frozen in liquid nitrogen, and then stored at −80 °C. Before analysis, samples were dried in a freeze-drier (Labconco, KS City, MO). Samples were grounded with a mortar and pestle and passed through a 0.35-mm screen. A 40-mg ground sample was hydrolyzed with 2.5 mL of amylase (Labconco, KS City, MO). Samples were grounded with a mortar and pestle and passed through a 0.35-mm screen. A 40-mg ground sample was hydrolyzed with 2.5 mL of amylase for 24 h at 37 °C. The solution was then quickly cooled in water or in an ice bath. The pH was adjusted between 5 and 7 with 0.31 mL of 10 N NaOH, added for tissue digestion for another 18 h. After the solution was neutralized with 3.0 mL of 2 N H2SO4, the solution was brought to 50 mL with distilled water. A 1.0-mL aliquot of the solution was transferred into a flask and the volume was brought to 50 mL with distilled water. The absorbance of the solution was measured at 515 nm using a spectrophotometer (Genesys 2) and the amount of sugar was determined by reference to a standard curve.

**Construction of pressure-volume curve for the estimate of cell wall elasticity, ψS at full turgor, and relative water content at zero turgor.** A pressure-volume technique is widely used to study water relations of plants, providing estimation of ψS at full turgor and critical tissue water content during plasmolysis, as well as cell wall elasticity (Barker et al., 1993; Kirkham, 2005; Schulte and Hinckley, 1985; White et al., 1992; Wilson et al., 1979). A total of 30 fully extended leaves were detached from plants in each container and used for the development of the pressure-volume curve for plants exposed to well-watered conditions and 15 d of drought stress when leaf lamina of all plants were completely wilted. The pressure-volume curve was developed for each species by plotting the reciprocal of leaf water potential (1/ψ) against RWC following the methods described by White et al. (1992) and Barker et al. (1993). Leaves were detached from the plant and were rehydrated in distilled water in the dark at 4 °C for 16 h until fully turgidity or a constant weight was reached. Individual leaves were then subjected to the following procedure: 1) they were blotted dry for the determination of turgid weight (TW); 2) they were kept in petri dishes in a growth chamber with a constant temperature of 23 °C, a PAR of 500 μmol·m−2·s−1, and a relative humidity of 70%; 3) they were weighed for leaf FW every 5 min; 4) immediately after FW measurement, leaf ψ was measured using a pressure chamber (Soil Moisture Equipment) following the method described by Kirkham (2005); and 5) after leaf ψ measurement, the leaf was dried in an oven at 80 °C for 48 h and DW was determined.

Leaf ψS at full turgor (ψS100) and relative water content at zero turgor (RWC0) or incipient plasmolysis were derived from each pressure-volume curve (Wilson et al., 1979). Bulk modulus of elasticity of the leaf tissue (ε) was determined from the relationship between pressure potential (ψS) and RWC by assuming an approximate linear relation (Kirkham, 2005) by

\[ ε = \frac{[ψS/RWC] \times RWC₀}{ψS} \]

Turgid weight to dry weight ratio was determined from the turgid weight after full hydration and the dry weight of the leaf. TD/DW ratio indicates the extent of leaf cells to absorb water or cell wall hydration level (Malinowski and Belesky, 2000).

**Production and growth of leaves, tillers, and roots in response to re-watering.** At 3 and 7 d of re-watering, the number of wilted, brown leaves and turgid, green leaves was counted to calculate the percentage of live leaves. Green leaves were separated from brown leaves and leaf area for each group was measured using the digimizer program (version 3.1.2.0; MedCalc Software, Mariakerke, Belgium). The percentage of green leaves to total leaf area was calculated.

At 7 and 16 d of re-watering, roots were washed free of soil and newly regenerated roots were separated from old, existing roots that were exposed to drought stress. Images of the new roots were scanned and analyzed for root number, length, and surface areas using WinRHIZO (Regent Instruments, Loretteville, Canada). Roots were then dried in an oven at 80 °C for 72 h to determine dry weight.

At the end of drought stress (20 d), all shoots were removed at the shoot base in the re-watering treatment (Treatment 3). Plants were re-watered to examine the regeneration of new tillers in response to re-watering. At 16 d after rewatering, the number of new tillers was counted. The survival rate of plants was calculated as percentage of the number of new tillers to the total number of tillers.

**Statistical analysis.** The experiment was a completely randomized plot design with two factors (watering treatments and grass species). Treatment effects, species differences, and treatment × species interactions were determined by analysis of variance according to the general linear model procedure of SAS (version 9.0; SAS Institute, Cary, NC). Means were tested with least significant difference test at a probability level of 0.05. Correlation analysis between physiological parameters was performed using a correlation matrices procedure with STATISTICA Software (StatSoft, Tulsa, OK).

**Results and Discussion**

**Soil water status during drought and re-watering.** Soil water content was measured as an indication of soil water status and it also provides a good estimate of the water depletion rate of a plant (Kirkham, 2005). SWC in a 20-cm soil depth in KBG and PRG declined rapidly as irrigation was withheld, to 4% by 20 d of drought stress in both species (Fig. 1). SWC increased to the pre-stress control level upon re-watering. No significant difference in SWC was found between the two species during the 20-d drying period or re-watering. This indicates that both species were exposed to the same level of water deficit or availability under drought stress and during re-watering, and the water depletion rate was not different between the two species. Therefore, the variation in growth performance and physiological responses to drought stress between KBG and PRG may not be associated with differences in soil water use rate.

**Growth and physiological changes and cultivar variation in drought tolerance and recovery of drought damages leaves.** Grass quality of KBG and PRG declined with...
drought stress and increased upon re-watering, but the extent of change varied between the two species (Fig. 2). Grass quality for KBG decreased to 40.3% of the well-watered control at 20 d of drought stress and then increased to 53.7% and 64.8% of the well-watered control at 3 and 7 d of re-watering, respectively. Grass quality for PRG decreased to 31.9% of the well-watered control at 20 d of drought stress and then increased to 35.2% and 46.3% of the well-watered control at 3 and 7 d of re-watering, respectively. Grass quality of KBG was 26.1%, 52.6%, and 40.0% higher than that of PRG at 20 d of drought stress, 3 and 7 d of re-watering, respectively. The significantly higher grass quality and slower decline under drought and significantly more rapid and greater extent of recovery upon rewatering for KBG suggest that KBG had superior drought tolerance and better recuperative potential from drought damage compared with PRG. Ma and Wang (2001) also reported that after 20 d of drought stress, plants of three PRG genotypes were completely desiccated and died while five KBG genotypes still had an average survival rate of 58.4%. Kanapeckas et al. (2008) also reported faster recovery of grass quality in KBG than PRG after 10 d re-watering. The interspecific difference in overall grass performance under drought stress and recuperative potential could be related to some of the physiological traits examined in this study, as discussed below.

Cell membrane stability (expressed as EL) has been widely used as an indicator of leaf desiccation tolerance (Martin et al., 1987). The level of cell damage increases with the plasma membrane permeability (Li et al., 1994, 2004; Lu and Gao, 1996). Leaf EL for KBG increased significantly above the well-watered control level at 15 d of drought, which was 7.0 times the well-watered control by 20 d of drought stress (Fig. 3). For PRG, significant increases in EL were found at 5 d of drought stress; by 20 d of drought, it increased to 8.0 times the well-watered control. EL of both species decreased at 3 and 7 d of re-watering compared with the level at 20 d of drought, but were still significantly higher than the respective well-watered controls. KBG leaves had significantly lower EL than PRG leaves during drought and re-watering. The EL data suggest that leaves of KBG and PRG experienced permanent cellular membrane damages when leaf RWC declined to ≈25% and could not recover, even after re-watering. Similar results have been observed in two KBG cultivars in a previous study (Wang and Huang, 2004). However, KBG leaves were able to maintain better cell membrane stability than PRG under such severe drought stress. This result is in agreement with the results reported in Ma and Wang (2001).

Leaf chlorophyll and associated photochemical reactions are important factors in determining photosynthetic capacity. Decline in leaf Chl content is associated with leaf senescence induced by drought stress in various plant species (Wise and Naylor, 1987). Increases in EL and loss of chlorophyll usually are attributed to membrane damage and the processes occurring during drought-induced leaf senescence (Liu and Huang, 2000; Simon, 1974). In this study, no significant difference was detected in Chl content after 20 d of drought stress for KBG compared with the well-watered control (Fig. 4). During the re-watering treatment, Chl for KBG decreased, which was 72.2% and 80.5% of the well-watered control at 3 and 7 d, respectively. For PRG, Chl content decreased to 46.5% of the
well-watered control at 20 d of drought, and was only 39.2% and 36.1% of the well-watered control after 3 and 7 d of re-watering, respectively. The decline in Chl content or partial recovery in Chl content after re-watering in both grass species suggests that severe drought may have caused irreversible damage in leaf tissue, negating a recovery of Chl biosynthesis. However, leaf Chl content in KBG was significantly higher than that in PRG during drought and re-watering, which was 71.5%, 59.0%, and 68.0% higher at 20 d of drought stress, 3 and 7 d of re-watering, respectively. Leaf Fv/Fm of KBG decreased to 22.9% of the well-watered control by 20 d of drought stress, and then increased to 80.3% and 96.3% of the well-watered control at 3 and 7 d of re-watering, respectively (Fig. 5). For PRG, Fv/Fm decreased to only 8.9% of the control level after 20 d of drought stress, and increased to 39.6% and 83.5% of the well-watered control at 3 and 7 d of re-watering, respectively. Leaf Fv/Fm in PRG was 61.5%, 51.5%, and 13.1% lower than that in KBG at 20 d of drought stress, 3 and 7 d of re-watering, respectively. The changes in Chl and Fv/Fm ratio indicated that drought stress for 20 d induced severe leaf senescence, but KBG leaves were able to maintain greener and more active photochemical reactions than PRG leaves.

The level of TNC of a plant has been considered an indirect indicator of physiological activity and is related to the recuperative potential of plants from stress damages (Busso et al., 1990; Sheffer et al., 1979). TNC content in leaves of PRG did not exhibit significant changes during drought and re-watering compared with the control (Fig. 6A), while crown TNC content decreased by 30.8% at 20 d of drought stress and 18.3% at 7 d of re-watering (Fig. 6B). These results suggest that carbohydrate accumulation in leaves was unaffected by drought stress, but storage in crowns was inhibited during drought, which may affect the recuperative potential of plants from drought damage. Previous studies have also shown a reduction in concentrations of fructans and other water soluble carbohydrates in the basal stem and crown tissues in PRG and tall fescue (Festuca arundinacea) in response to drought stress (Norris and Thomas, 1982; Spollen and Nelson, 1994; Suzuki and Chatterton, 1993; Volaire and Gandoir, 1996). In contrast, TNC content in leaves of KBG increased by 35.1% at 20 d drought stress and 4.8% at 7 d of re-water treatment compared with the well-watered control (Fig. 6A). In KBG rhizomes, TNC content increased by 3.8% at 20 d of drought stress and decreased by 45.8% at 7 d of re-watering compared with the control (Fig. 6B). The accumulation of TNC in leaves and rhizomes in KBG during drought stress could serve as a carbohydrate reserve and/or used in osmotic adjustment, while the decrease in TNC content in rhizomes during re-watering may reflect the use or remobilization of carbohydrate from rhizomes for tissue re-growth.

Changes in leaf water relations and cell wall modulus during drought and re-watering. Leaf RWC for KBG decreased to 26.1% of the well-watered control by 20 d of drought (Fig. 7). By 7 d of re-watering, RWC for KBG increased to 45.0% of the well-watered control. Leaf RWC for PRG decreased to 25.7% of the well-watered control at 20 d of drought stress, and then increased to 40.8% of the well-watered control at 7 d of re-watering. No significant difference in RWC was found between the two species during drought stress or re-watering. These results indicated that leaves of both species suffered the same level of internal water deficit. Leaves of both species experienced severe water deficit (with RWC, as low as 24%), which could not fully recover even after re-watering. A previous study with KBG found that RWC could not resume fully after 6 d of rewatering when RWC declined below 25% during drought stress; a RWC of 25% was the critical level for whole-plant survival of drought stress for KBG (Wang and Huang, 2004). Prolonged, severe soil drying can ultimately cause cellular disruption to the point that leaves may no longer be fully rehydrated (Volaire et al., 1998a, 1998b). The critical RWC below which tissue physiological injuries and death occurs is ≈50%, but can vary among species and tissue types (Taiz and Zeiger, 1998).

Osmotic adjustment is an important mechanism of dehydration tolerance, helping to maintain cell turgor and volume during drought stress (Clifford et al., 1998). Differences in ψ_S at ψ_50 are associated with species variation in drought tolerance and duration of water deficit (Barker et al., 1993). Osmotic potential at ψ_50 decreased by 17.3% in KBG after 15 d of drought stress compared with the well-watered control, while it did not change significantly in PRG (Table 1).
Decreases in $\psi_{k100}$ in PRG leaves could be due to the inhibition of osmotic solute accumulation or the lack of osmotic adjustment in this species under drought stress.

Cell wall elasticity is another important factor regulating cellular turgor and cell growth (Blake et al., 1991; Saito and Terashima, 2004). It is known that some plant species with elastic cell walls have a high inherent drought tolerance (Fan et al., 1994; Zimmermann and Steudle, 1978). Cell wall elasticity is negatively correlated to $\varepsilon$, which is often used to estimate cell wall elasticity. Leaf $\varepsilon$ values increased by 81.5% in PRG under drought stress compared with the well-watered control, while no significant difference in $\varepsilon$ was detected in KBG between the control and drought treatment (Table 1). PRG leaves had higher $\varepsilon$ values than KBG leaves under drought stress. The increase in $\varepsilon$ indicated that the cell wall in PRG leaves was less elastic than KBG leaves, thereby restricting cell growth in PRG leaves more than in KBG. The unchanged $\varepsilon$ in KBG leaves under drought stress suggested that these leaves could be able to maintain cell wall elasticity for turgor maintenance under drought stress. In fact, RWC at zero turgor of KBG was significantly lower than that of PRG, particularly under drought stress (Table 1), suggesting that KBG leaves were able to maintain turgidity with lower water content than PRG leaves. Cell turgor maintenance may be due to osmotic adjustment through active accumulation of solutes and/or an increase in solute concentration through reduced cell volume (elastic adjustment) (Grammatikopoulos, 1999). The increases in osmotic adjustment coupled with the maintenance of cell wall elasticity could help maintain growth during drought and facilitate growth during recovery from drought stress upon re-watering (Clifford et al., 1998).

The increases in leaf $\varepsilon$ values in PRG leaves were accompanied by decreases in TW/DW ratio. The TD/DW ratio indicates the water absorbing ability of a cell wall or a cell wall hydration level (Malinowski and Belesky, 2000). The TW/DW ratio in PRG leaves decreased by 18.8% during drought stress compared with the well-watered control. Decreases in TW/DW ratio have been observed in different genotypes of zoysiagrass (Zoysia japonica) exposed to drought stress compared with the pre-stress values (White et al., 2001). Such changes in cell wall properties could be associated with changes in cell wall thickness or cell wall constituents, as reported in tropical forage grass species (Wilson et al., 1980). No significant differences in TW/DW ratio for KBG were detected between the control and drought treatment. The lack of changes in TW/DW suggested that KBG leaves were better able to maintain cell wall elasticity through osmotic adjustment.
hydration level under drought stress compared with PRG leaves (Malinowski and Belesky, 2000).

**Correlation analysis of physiological factors under drought.** Correlation analysis was performed between grass quality as an overall drought performance indicator and EL, which is commonly used as an indicator for cellular damage under drought (DaCosta and Huang, 2006). Correlation between parameters estimated from the pressure-volume curve and EL for leaves of both species exposed to drought stress was also analyzed. For both grass species, EL was negatively correlated with grass quality ($r^2 = 0.92$ for KBG, $r^2 = 0.97$ for PRG). The $\psi_{e100}$ has been shown to be correlated to drought tolerance and suggested as a rapid and economical selection tool for drought tolerance (White et al., 2001). In this study, $\Psi_{e100}$ was negatively correlated with EL ($r^2 = 0.80$) or positively correlated with cell membrane stability under drought stress in KBG, but the correlation was not significant in PRG leaves ($r^2 = 0.37$). These results indicated that osmotic adjustment may be involved in protection of cell walls from drought damage or, in turn, stable cellular membranes may facilitate solute accumulation or prevent electrolyte leakage from leaves exposed to drought stress in KBG. In contrast, RWC$_0$ was negatively correlated to cell membrane stability and positively correlated to EL in both KBG ($r^2 = 0.83$) and PRG ($r^2 = 0.66$), suggesting that leaves with stable cell membranes could lose turgor at a lower RWC or maintain turgid with lower water availability.

Leaf $\varepsilon$ was positively correlated with EL in KBG ($r^2 = 0.74$) and PRG ($r^2 = 0.99$). The negative correlation between $\varepsilon$ and cell membrane stability in both grass species suggested elastic cell walls accompanied by stable membranes may play an important role in leaf dehydration tolerance as the coupled changes may prevent plasma membrane separation from cell walls or plasmolysis, which is critical for the maintenance of cell turgor and cell growth (Nilsen and Orcutt, 1996).

**Re-greening and rehydration of drought-damaged leaves and regeneration of new roots in response to re-watering.** Leaf senescence during drought stress may conserve water by decreasing transpirational leaf area, but the attribute of re-greening and rehydration of senescent, desiccated leaves may play a critical role in the resumption of plant growth when water becomes available (Munne-Bosch and Alegre, 2004). At 3 and 7 d of re-watering, rehydrated, green leaf percentages of drought-damaged leaves in terms of number, area, and length of leaves (rehydrated leaves in proportion to all leaves) were all significantly higher in KBG compared with those in PRG (Fig. 8). These data indicated that desiccated, senescent leaves in KBG induced by drought stress were better able to re-green and be rehydrated upon re-watering compared with PRG leaves, which may also reflect less physiological or cellular damages in KBG leaves under drought stress.

At 7 d of re-watering treatment, new roots were found in KBG and PRG plants. The new roots in PRG were 1.33 times greater in total surface area, 1.27 times higher in total length, and 1.35 times more in total dry weight than KBG roots; the number of new roots, however, were significantly lower than in KBG (Fig. 9). At 16 d of re-watering, new roots for PRG were 25.8% higher in total length, 85.5% higher in total surface area, 1.66 times higher in total dry weight, but root number was 15.9% lower compared with those in KBG. These data demonstrate that PRG exhibited more rapid elongation of new roots while KBG was able to regenerate a greater number of new roots in response to re-watering.

The survival rate of plants from drought stress, expressed as percentage of the number of new tillers produced after re-watering to the total number of tillers, was also evaluated at 16 d of re-watering. A significant amount of new tillers were regenerated in both species in response to re-watering even though plants were exposed to prolonged periods (20 d) and severe drought stress (SWC = 4%, RWC = 24%). The survival rate for KBG (90.6%) was significantly higher than that for PRG (82.3%). These results indicated that both perennial grass species had high survivability, but KBG was better than PRG. The differential responses of root growth and tiller production between the two species suggested that new root formation

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**Table 1. Osmotic potential at full turgor ($\psi_{e100}$), relative water content at zero turgor (RWC$_0$), bulk modulus of tissues elasticity ($\varepsilon$), and the ratio of turgid weight to dry weight (TW/DW) determined under well-watered control and drought stress at 15 d of treatment for kentucky bluegrass (KBG) and perennial ryegrass (PRG).**

|                | $\psi_{e100}$ (MPa) | RWC$_0$ (%) | $\varepsilon$ (MPa) | TW/DW (ratio) |
|----------------|---------------------|-------------|---------------------|---------------|
| KBG Control    | –1.44 ab$^*$         | 88.44 a     | –0.63 b             | 4.11 c        |
| Drought        | –1.69 a              | 89.55 b     | –0.64 c             | 4.08 b        |
| PRG Control    | –1.35 b              | 91.03 ab    | –0.54 c             | 5.28 a        |
| Drought        | –1.24 b              | 93.01 a     | –0.98 a             | 4.29 b        |

$^*$Means followed with the same letters within the column were not significantly different at $P \leq 0.05$ level based on LSD test.

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![Fig. 8. Percentage of green part to total leaf area and length, and live leaf percentage of existing leaves of kentucky bluegrass (KBG) and perennial ryegrass (PRG) under re-watering treatment. Means followed by the same letters for each parameter were not significantly different based on LSD test ($P \leq 0.05$).](image-url)
could be more critical, relative to root elongation, for leaf rehydration or regeneration of new tillers upon re-watering.

**Summary**

Kentucky bluegrass exhibited superior drought survival and recuperative potential than PRG, as manifested by the maintenance of higher grass quality, leaf Chl, and Fv/Fm under the same level of water deficit during drought stress and more rapid and greater extent of recovery in all these physiological parameters upon re-watering. Osmotic adjustment, cell wall elasticity, and cell membrane stability were closely correlated to leaf desiccation tolerance or drought survival in perennial grass species. Post-drought recovery in leaf hydration level and physiological activities could be associated with the accumulation of carbohydrates in leaves and rhizomes during drought stress and new root production after re-watering.

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Fig. 9. Area, length, dry weight, and numbers of newly regenerated roots at 7 and 16 d of re-watering in kentucky bluegrass (KBG) and perennial ryegrass (PRG). Means followed by the same letters for each parameter were not significantly different based on LSD test (*P* ≤ 0.05).
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