Buffalograss is native to the Great Plains of North America (Huff and Wu, 1987). This species is widely used as a low-maintenance turfgrass in parks, cemeteries, and rights of way due to its excellent cold, heat, and drought tolerance (Beard, 1985). Buffalograss is increasingly used on golf course fairways and other sports fields in arid and semiarid regions where water shortage is becoming an important concern (Frank et al., 2000; Pozarnsky, 1983). However, one of the weaknesses of buffalograss for use in golf fairways is poor ball-supporting ability due to low shoot density. Improving the traits related to shoot density has been proposed as one of the goals in buffalo-grass breeding programs (Johnson et al., 2000; Taliaferro and McMaugh, 1993). Buffalograss is considered a “guerilla”-type clonal species (Lovett Doust, 1981), meaning it can spread extensively. The opposite are “phalanx”-type plants [short nodes and compact growth (Lovett Doust, 1981)], which have higher potential for shoot density because of the intensive growth habit. Limited information is available on shoot density affected by the growth type, “guerilla” type versus “phalanx” type, although research has been conducted on the effects of nitrogen and mowing height (Frank et al., 2004). No research has yet been conducted to understand the effect of nutrient and carbohydrate translocation on the shoot density of buffalograss.

Reproduction and population density are often discussed in relation to population biology and ecology of clonal plants. Semi-autonomous young ramets may import nutrients, water, hormones, etc., from the parent until they are established, and thereafter they may be independent of parental support (Pitelka and Ashmun, 1985). In ecological research, members of a clone are considered ramets before they become independent from the parent plant for study of photosynthate partition (Alpert, 1991). Another important characteristic of clonal plants is physiological integration, which is defined as a process of redistribution of assimilated resources among the interconnected ramets according to source-sink relationships (Forde, 1966; Kaitaniemi and Honkanen, 1996; Marshall, 1990). Physiological integration is an important means by which clonal plants adapt to heterogeneous environmental conditions (Kroon et al., 1996; Marshall, 1990). Because buffalograss has a “guerilla”-type growth habit, it may differentially exploit the environment, selecting favorable and avoiding unfavorable sites (Jackson, 1979). As a stoloniferous clonal species, buffalograss clones are often connected for an extended period of time after establishment.

Resource sharing in clonal plants is reflected in the ability of perennial grasses to change phenotypes in response to fluctuating environments and stresses (Bradshaw, 1965). For instance, the specialization of certain ramets in photosynthesis and nutrient-uptaking functions may be enhanced when essential nutrients and light are heterogeneously distributed (Stuefer...
et al., 1994). The consideration of phenotypic changes in turfgrass breeding was discussed by Casler and Duncan (2003) and Bradshaw (1965). An internal gradient was established when connected ramets were exposed to different nutrient availability (Marshall, 1990), and such a gradient may be the driving force for the nutrient redistribution (Marshall and Price, 1999). Water was transported from parent ramets to offspring ramets along the water potential gradient in *Fragaria chiloensis* (Alpert and Mooney, 1986). Photosynthetic also was translocated among connected ramets in many grass species (Nyahoza et al., 1974; St. Pierre and Wright, 1972). However, the mechanisms of physiological integration in grasses are not well understood (Hellström et al., 2006).

Huang (1999) reported that buffalograss performed better than zoysiagrass (*Zoysia japonica*) under localized soil drying and attributed the difference to the more extensive root system in buffalograss. Qian et al. (2009) reported that buffalograss shoot number displayed water integration when connected ramets were grown in media with different soil water contents. The same authors also reported inter-ramet translocation of lipid peroxidation, antioxidants, and proline (Qian et al., 2009). High uniformity is a major quality component of turf. One of the purposes of topdressing and applying wetting agents, among many other turfgrass management practices, is to correct the heterogeneous conditions in the root zone media and to create uniform turf (Karnok and Tucker, 2001; Minner et al., 1997). This integration appears to deal with soil variability to provide more uniform growth in buffalograss—a highly desirable trait. Understanding this process may also help to make precise cultural practices and improve turf uniformity.

The role of plant hormones in response to drought stress has been discussed for endogenous (Abreu and Munne-Bosch, 2008) and exogenous (Liu and Huang, 2002; Zhang and Schmidt, 1999) sources. Abscisic acid was found to play a role in water stress-induced antioxidants (Jiang and Zhang, 2002) and morphological responses (Zhang and Davies, 1989). Increased contents of zeatin riboside (ZR) in shoot and root alleviated the heat stress in creeping bentgrass (*Agrostis palustris*) (Liu and Huang, 2002). Gibberellic acid treatment delayed senescence in *Cynodon dactylon* caused by chilling (DiPaola et al., 1981). Research also indicated that hormone allocation was modified in the process of resource sharing in clonal plants of *F. chiloensis* (Alpert et al., 2002).

The primary objective of this study was to assess the effects of differential water stress on water transport and photosynthetic translocation and distribution. A secondary objective was to examine if endogenous hormone translocation and distribution display coordinated changes under differential water stress in buffalograss.

**Materials and Methods**

Plant materials used in this study were clones from a single parent plant of ‘Texoka’ buffalograss vegetatively propagated in a greenhouse (Chinese Forestry Institute, Beijing, China) with temperatures at 25 ± 2 °C and relative humidity at 50% ± 10%, and supplemental light provided by metal halide lamps on a 16-h photoperiod. The connected ramets were produced by generating roots and shoots at the nodes of buffalograss stolons. To summarize, each node of a stolon with four to five nodes was covered with a mixture of sand and peat in equal volumes on 15 June 2007. The rooting medium was fertilized at the beginning of the experiment with a 12N–5.3P–10K water-soluble fertilizer at 50 kg·ha⁻¹ N and was kept moist by frequent watering under a mist spray system. On 5 July 2007, 3 weeks after the initiation of tiller induction, sections containing three connected ramets were harvested by cutting them off the stolons and uniform sections were selected for the experiments. The sections did not include the unrooted newest nodes at the tip of stolons, which were cut off at the harvest. In this study, a ramet is defined as a tiller with developed shoot and roots at one node. The youngest, second, and oldest ramets were denoted as R₁, R₂, and R₃, respectively. The roots of connected ramets were rinsed free of growth media with distilled water and each ramet was cultured in a separate glass flask with 200 mL of half-strength Hoagland solution (Hothem et al., 2003) (\(\psi_u \approx -0.05\) MPa, \(\mathrm{pH} = 5.8\)) for 6 d before the initiation of treatments in different experiments (Fig. 1). Osmotic potential of Hoagland solution was measured using a dew point potential meter (WP4; Decagon Device, Pullman, WA). The solution was changed daily and air was pumped into the hydroponic solution through pressure-regulating valves and PVC tubes with a porous stone cap at the end of each tube in the containers.

Three experiments were included in this study and all followed a similar general setup as follows. The first group of ramets had R₃ cultured in half strength Hoagland solution with 30% of PEG-8000 (previously named PEG-6000; Sigma-Aldrich, St. Louis) (\(\psi_o = -1.2\) MPa), while R₁ and R₂ were kept in half strength Hoagland solution only, and was denoted as \(\text{PR}_3\text{PR}_2\text{PR}_1\). The second group of ramets had R₂ cultured in half strength Hoagland solution with 30% of PEG-8000, while R₁ and R₃ were kept in half strength Hoagland solution only, and was denoted as \(\text{PR}_1\text{PR}_2\text{PR}_3\). The third group of ramets had R₁ cultured in half strength Hoagland solution with 30% of PEG-8000, while R₂ and R₃ were kept in half strength Hoagland solution only, and was denoted as \(\text{PR}_1\text{PR}_3\text{PR}_2\). The fourth group of ramets had all three ramets cultured in half strength Hoagland solution and served as a control, \(\text{PR}_1\text{PR}_2\text{PR}_3\). Polyethylene glycol has been used in a number of studies to decrease the \(\psi_o\) of solutions (Jia et al., 2001; Jiang and Zhang, 2002; Nayyar, 2003). These ramet groups were maintained in a growth chamber with

![Fig. 1. Diagram showing the interconnected ramets of 'Texoka' buffalograss grown in separate containers, where R₃, R₂, and R₁ represent the oldest, second, and youngest ramet, respectively.](image-url)
a 10-h photoperiod, light intensity of 500–600 μmol·m⁻²·s⁻¹, temperature at 24 to 28 °C, and relative humidity at 45% to 60%.

Experiment 1 was intended to monitor the direction of water movement in the connected ramets when one of them was under water stress from the addition of PEG-8000. Acid fuchsin dye (EMD Chemicals, Gibbstown, NJ) has been used to trace xylem-mediated water transport in many species (Baum et al., 2000; Warren et al., 2008), where water transport followed the route of dye tracer. In this study, acid fuchsin dye was added to the culture solutions, one ramet at a time within each group, at a concentration of 0.25 g·L⁻¹. A superscript D was added to R₃R₂R₄ to denote dye addition to its culture solution. The acid fuchsin tracing treatments were arranged in a randomized complete block design with three replications. Each treatment unit contained three plants. The roots, leaves, and internodes of all ramets were visually observed 4 h after the dye was added and the organs were recorded as positive if red color was observed.

The purpose of Expt. 2 was to assess hormonal distribution in shoots and roots of different ramets affected by water deficit stress in one ramet. In this experiment, the treatments also were arranged in a randomized complete block design with three replications. Each treatment unit contained 10 plants. The leaves, roots, and internodes of R₁, R₂, and R₃ were harvested 4 h after the PEG treatment was applied. The samples were ground with an ice-cold mortar and a pestle in 6 mL of ice-cooled 80% methanol (v/v) containing 1 mmol·L⁻¹ butylated hydroxytoluene (BHT) to prevent oxidation and incubated overnight at 4 °C before centrifugation at 5000 g, for 15 min at 4 °C. After removing the supernatant, the residues were suspended with ice-cooled extraction solution in darkness for 4 h and centrifuged again at 4 °C. The two supernatants were combined and passed through a Chromosep C18 column (Waters, Milford, MA). The efflux was dried in a freeze dryer (Labconco, London) and was then dissolved in 2 mL of phosphate-buffered solution (PBS) containing 0.1% (v/v) polysorbate 20 (Tweeoctr; Fisher Scientific, Pittsburgh) and 0.1% gelatin (Fisher Scientific; pH 7.5). The solution was used for determination of concentrations of indole-3-acetic acid (IAA), gibberellic acid (GAs), tans-ZR, and abscissic acid (ABA) by enzyme-linked immunoabsorbent assay (ELISA) following the methods described by Yang et al. (2001). The mouse monoclonal antibodies and antibodies against ZR, IAA, GAs, and ABA used in ELISA were produced at the Phytohormones Research Institute (China Agricultural University, Beijing, China). IgG-horseradish peroxidase was purchased from Sigma-Aldrich. The absorbance was measured at 490 nm using an ELISA recorder (model DG-3022; Huadong Electron Tube Factory, Shanghai, China). Calculations for the concentration of IAA, GAs, ZR, and ABA were performed following Weiler et al. (1981). The recovery of each hormone was monitored by adding a known amount of standard hormone to a split extract. All recoveries were over 90%. Furthermore, all sample dilution curves were parallel with the standard curves, indicating the absence of nonspecific inhibitors in the extracts.

In Expt. 3, ¹⁴CO₂ was fed to the leaves of R₁, R₂, and R₃ ramets, respectively, at the initiation of each PEG treatment to monitor the translocation of photosynthate. A superscript L was added to R₃R₂R₁ to denote ¹⁴C label. The treatments were arranged in a randomized complete block design with three replications. Each treatment unit contained three plants. The ramet to be labeled was first enclosed in an air-tight transparent polyvinylchloride gas-sampling bag (20 × 5 cm), and 10 mL of ¹⁴CO₂ was injected into each bag through a septum stopper via a syringe and needle. The ¹⁴CO₂ was generated by adding 0.1 M H₂SO₄ into a vial containing NaH¹⁴CO₃ (749 kBq) (PerkinElmer, Waltham, MA). The air in the bags was pumped out and the residual ¹⁴CO₂ was absorbed by soda lime after the ramets were incubated in ¹⁴CO₂ for 30 min. Four hours after ¹⁴CO₂ labeling, the shoots, roots, and internodes were harvested and immediately oven-dried at 95 to 100 °C for 10 min and then at 80 °C for 48 h. The samples were ground with a mortar and pestle and were then measured with a BH1216 low-background α, β-scintillation counter (Beijing Nuclear Instrument, Beijing, China). The sample radioactivity was expressed as counts per gram of dry tissue. The percentage of ¹⁴CO₂-labeled photosynthate in each part of the ramet was adjusted by the percentage of weight relative to the whole plant.

The data were subjected to analysis of variance (ANOVA) using the general linear model in SAS (version 9.1; SAS Institute, Cary, NC). Fisher’s protected least significant difference (LSD) values were calculated. Percentage data were subjected to arcsine square-root transformations before the analysis, and final results were converted to the non-transformed values for clarity.

Results and Discussion

When acid fuchsin was added to the culture solution of a ramet, the shoot and roots of that ramet were always dyed regardless of the water stress from PEG (Table 1). When none of the three connected ramets were under water stress, the acid fuchsin moved from older ramet to younger ramet (Table 1). When a ramet was under water stress from PEG, in addition to apical movement, the dye also moved toward the shoot of the stressed ramet, as shown from the comparisons R₁R₂R₃ versus R₁R₂R₄, R₁R₃R₄ versus R₂R₃R₄, and R₂R₃R₄ versus R₃R₂R₄ (Table 1). When an older ramet was stressed, the apical movement of dye was suppressed, as shown in comparisons of R₁R₃R₄ versus R₂R₃R₄ and R₁R₂R₄ versus R₃R₂R₄ (Table 1). Therefore, our results indicated that interramet water integration happened when one of the connected ramets was under water stress, though it also seems that integration occurs without water stress if ramets differ in age. In other words, integration is one-way in some cases.

The statistical results from ANOVA showed that there were significant differences for the endogenous hormone concentration in the shoots and roots among treatments (Table 2) as well as the shoot/root ratio of endogenous hormones (Table 3).

The content of ZR decreased in PEG-treated roots, but increased in the roots of untreated youngest ramets, as in R₁R₃R₄ and R₂R₃R₄, or untreated oldest ramets, as in R₁R₂R₃ (Table 2). However, because ZR is mainly synthesized in roots and final results were converted to the non-transformed values for clarity.

The levels of ZR in shoots were increased compared with the untreated control except in the shoot of R₃ and R₁ in the treatment R₃R₂R₄. The low content of ZR in the R₃ shoot may be due to the increased transport stream within the vascular bundle from R₃ roots to younger ramets (Table 1). The shoot/root ratios of ZR content were all increased compared with the control except for the youngest ramet in treatment R₃R₂R₄, where the roots of the youngest ramets were treated with PEG (Fig. 2).

The ABA content in shoot was increased in R₃R₄ and R₂R₃R₄ treatments compared with the control (Table 2). When
the roots of the youngest ramet were treated with PEG, the ABA content was not increased in the oldest and youngest shoot, but only increased in the middle shoot, indicating a different pattern of translocation. However, ABA content was increased in the roots of all treatments compared with the control (Table 2). The shoot/root ratio of ABA were decreased in all treatments compared with the control (Fig. 2). This overall increase of ABA content in roots or decrease in shoot/root ratio was probably because ABA is mainly a stress hormone (Guy, 1990; Hasegawa et al., 1987; Tanino et al., 1990; Zhang and Davies, 1989) and can be transported in xylem and phloem and also in parenchyma cells outside the vascular bundles (Salisbury and Ross, 1992).

Water stress in one of the three ramets from PEG caused reduction of IAA content in shoots and roots, especially in the ramet being stressed (Table 2). As a result, the ratio of IAA levels between shoot and root also changed. When the R3 ramet was stressed, the shoot-to-root ratio of IAA in the R2 ramet increased, while the ratios in R1 and R3 decreased compared with non-stressed control (Fig. 2). When the R2 ramet was stressed, the shoot-to-root ratio of IAA in the R3 ramet increased, while the ratios in R1 and R2 decreased compared with the non-stressed control (Fig. 2). When the R1 ramet was stressed, the shoot-to-root ratio of IAA in all ramets decreased compared with the non-stressed control (Fig. 2). Because IAA transportation is usually in the parenchyma cells at a relatively slow speed (Salisbury and Ross, 1992), it is not clear if the changes were due to direct inter-ramet integration of IAA.

The GA content in shoots and roots of all treatments increased compared with the control (Table 2). It is not clear whether this was due to upregulation of GA synthesis in shoots and roots or transport from connected ramets. The shoot/root ratio of GAs decreased in all treatments compared with the control (Fig. 2). In R3PR1 and R3R2R1 treatments, the ratio decreased for the two ramets that were not treated with PEG, and remained at the level of the control for the ramet that was treated with PEG (Fig. 2). The distribution and allocation of endogenous hormones were shown to be affected by differential water stress in the interconnected ramets. Those hormones may be responsible for the regulation of the physiological integration, although the mechanisms were not clear.
The distribution of 14C-labeled photosynthate was not only different between different treatments, but was also affected by the location of labeling (Table 4). When the oldest ramet was under water stress, transport of labeled photosynthate was not detected from the youngest ramet to the roots and shoots of older ramets when the youngest was labeled; however, translocation was found from the shoot of the second ramet to the shoot of the third ramet (Table 5). Yet, in R3PLR2R1, translocation of labeled photosynthate was detected from the shoot of the third ramet to the second ramet. This indicated that the youngest ramet was the sink, but when the oldest ramet was under stress, there might be photosynthate translocation toward the old ramet. The increased translocation of labeled photosynthate toward the shoot of stressed ramet also was shown in treatments R3R2PR1 and R3LR2PR1. The photosynthate retention in and translocation to the stressed roots were increased when the roots of the second ramet were stressed. When the roots of the youngest ramet were under stress, the photosynthate retention and translocation to the shoot of stressed ramet increased compared with the control (Table 5). While the translocation of photosynthate toward the stressed ramets was increased, the partition to the youngest ramet was in shoot and root, but the partition to the oldest ramet was more in the shoot. This pattern of source-sink relationship also was reported in red clover (Trifolium repens) (Ryle et al., 1981).

Unlike in the case of timothy grass (Phleum pretense), where defoliation stress had a minor effect on the distribution of labeled photosynthate to the predetermined sink structure (St. Pierre and Wright, 1972), the transport of photosynthate in buffalo grass under differentiated water stress demonstrated a strong integration toward stressed ramets, although the test was only conducted 4 h after the labeling.

The above results showed that buffalo grass was able to integrate according to the new source-sink relationship under heterogeneous conditions. Therefore, the fate of connected ramets was not independent in this clonal species. Even for established populations as in turf areas, the advantage may be still significant as long as the ramets are connected. Although the agronomic significance of this characteristic needs further evaluation, from a botanical point of view, the “phalanx”-type may create higher shoot density in turf because of the intensive

Table 3. Analysis of variance of the shoot/root ratios of endogenous hormone concentration in the ramets of ‘Texoka’ buffalograss with the roots of one of the three connected ramets treated with 30% polyethylene glycol (PEG-8000; Sigma-Aldrich, St. Louis).

| Source | DF | \( R_1 \) | \( R_2 \) | \( R_3 \) |
|--------|----|-------|-------|-------|
|        | F  | \( P > F \) | F  | \( P > F \) | F  | \( P > F \) |
| Replication | 2 | 4.23 | 0.0713 | 0.42 | 0.6748 | 5.79 | 0.0398 |
| Stress | 3 | 1018.96 | <0.0001 | 914.55 | <0.0001 | 4660.13 | <0.0001 |
| CV (%) | 1.56 | 2.39 | 1.61 | ABA |
| Replication | 2 | 0.14 | 0.8714 | 0.06 | 0.9391 | 0.23 | 0.8024 |
| Stress | 3 | 164.16 | <0.0001 | 319.16 | <0.0001 | 4362.84 | <0.0001 |
| CV (%) | 9.36 | 4.32 | 1.44 | IAA |
| Replication | 2 | 0.07 | 0.934 | 1.69 | 0.2612 | 1.56 | 0.2847 |
| Stress | 3 | 115.46 | <0.0001 | 809.26 | <0.0001 | 794.03 | <0.0001 |
| CV (%) | 3.78 | 2.94 | 2.99 | GAs |
| Replication | 2 | 0.00 | 0.9969 | 0.15 | 0.8619 | 2.12 | 0.2008 |
| Stress | 3 | 25.28 | 0.0008 | 454.12 | <0.0001 | 48.11 | 0.0001 |
| CV (%) | 7.80 | 4.87 | 8.79 |

\( R_3, R_2, \) and \( R_1, \) represent the oldest, second, and youngest ramet, respectively.
\( ZR, ABA, IAA, \) and \( GAs \) represent transzeatin riboside, abscisic acid, indole-3-acetic acid, and gibberellic acid, respectively.
\( CV \) Coefficient of variation.

Fig. 2. Shoot-to-root ratio of endogenous hormone concentration (\( \mu g \cdot kg^{-1} \) fresh wt) in the ramets of ‘Texoka’ buffalograss. POO, OPO, and OOP are the R3R2R1, R3R2R1, and R3R2R1 treatments, respectively. R3, R2, and R1 represent the oldest, second, and youngest ramet, respectively, and the superscript P represents PEG-8000 (previously named PEG-6000; Sigma-Aldrich, St. Louis) treatment in the roots. Vertical bar represents Fisher’s protected least significant difference at the 0.05 level. ZR, ABA, IAA, and GAs represent transzeatin riboside, abscisic acid, indole-3-acetic acid, and gibberellic acid, respectively.

The distribution of 14C-labeled photosynthate was not only different between different treatments, but was also affected by the location of labeling (Table 4). When the oldest ramet was under water stress, transport of labeled photosynthate was not
Table 4. Analysis of variance of the 14C labeled photosynthates transportation among the ramets of ‘Texoka’ buffalograss with the roots of one of the three connected ramets treated with 30% PEG-8000 (previously named PEG-6000, Sigma-Aldrich, St. Louis, MO).

| Source       | Shoot 1 | Shoot 2 | Shoot 3 | Root 1 | Root 2 | Root 3 |
|--------------|---------|---------|---------|--------|--------|--------|
|              | F       | P > F   | F       | P > F  | F      | P > F  |
| Replication  | 2       | 0.6     | 0.5781  | 1.5    | 0.2535 | 2.2    |
| Stress location (S) | 3       | 37.8    | <0.0001 | 32.6   | <0.0001| 47.0   |
| Label location (L) | 2       | 25,203.2| <0.0001 | 24,695.2| <0.0001| 23,420.0|
| S x L | 6      | 48.7    | <0.0001 | 23.4   | <0.0001| 14.1   |
| CV (%)       | 3.65    | 3.65    | 3.63    | 20.23  | 17.41  | 11.95  |

Table 5. 14C-labeled photosynthate transport among connected ramets of ‘Texoka’ buffalograss when one ramet was under osmotic stress with 30% polyethylene glycol (PEG-8000; Sigma-Aldrich, St. Louis, MO). Data presented are percentage of radioactivity in parts relative to the whole plant.

| Treatment | Shoot 3 | Shoot 2 | Shoot 1 | Root 3 | Root 2 | Root 1 |
|-----------|---------|---------|---------|--------|--------|--------|
|           | F       | P > F   | F       | P > F  | F      | P > F  |
| R3R2R1    | 0.047   | 0.048   | 84.154  | 0.001  | 0.002  | 14.226 |
| R3R2R1    | 0.142   | 0.074   | 97.77   | 0.059  | 4.555  | 0.137  |
| R3R2R1    | 89.256  | 0.000   | 0.000   | 9.777  | 0.000  | 0.003  |
| R3R2R1    | 0.014   | 0.027   | 95.328  | 0.005  | 0.000  | 2.487  |
| R3R2R1    | 0.232   | 0.050   | 93.830  | 0.117  | 0.009  |
| R3R2R1    | 95.477  | 0.245   | 0.004   | 4.352  | 0.037  |
| R3R2R1    | 0.002   | 0.076   | 96.468  | 0.000  | 0.050  | 1.221  |
| R3R2R1    | 0.009   | 0.015   | 97.012  | 0.003  | 0.003  | 0.269  |
| R3R2R1    | 0.004   | 0.014   | 97.658  | 0.006  | 0.000  | 0.214  |
| R3R2R1    | 97.192  | 0.049   | 0.695   | 0.177  | 0.003  | 0.043  |
| LSD0.05   | 0.014   | 0.014   | 0.006   | 0.010  | 0.011  |

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