Exogenous Application of Nitrogen and Cytokinin on Growth, Carbohydrate, and Antioxidant Metabolism of Creeping Bentgrass after De-submergence

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Abstract. Recovery from submergence stress is vital for plant regrowth. The objective of this study was to characterize plant growth, carbohydrate, and antioxidant metabolism of creeping bentgrass (Agrostis stolonifera) following application of nitrogen and cytokinin (CK) after de-submergence. Creeping bentgrass (cv. Penncross and O207) were submerged under the water for 14 days and then foliar-sprayed at 1, 2, 3, 7, and 14 days after de-submergence with six types of chemical treatments, respectively: 1) water (W); 2) 10 mM urea (N10); 3) 20 mM urea (N20); 4) 10 μM CK; 5) N10 with CK (N10CK); and 6) N20 with CK (N20CK). Leaves were harvested at 20 days after chemical applications for various measurements. Compared with the nonstressed plants, plant height (HT), chlorophyll index (Chl), leaf dry weight (DW), water-soluble carbohydrate content (WSC), activities of superoxide dismutase (SOD), and ascorbate peroxidase (APX) decreased, but catalase (CAT) and peroxidase (POD) activities, malondialdehyde (MDA), and total soluble protein (TSP) content increased in both cultivars exposed to 14 days of submergence. After de-submergence, plants treated with N alone (N10, N20) or combined with CK (N10CK, N20CK) generally had higher HT, DW, Chl, TSP, and a lower amount of MDA, compared with treatments of W or CK alone, whereas treatment using CK resulted in higher WSC for both cultivars. Foliar applications of N and CK had some effect on SOD, CAT, POD, and APX activities after de-submergence, but the effects were not consistent across chemicals and cultivars. The results indicated that foliar application of N or combined with CK promoted plant growth and reduced lipid peroxidation after de-submergence. The results also suggested a more positive role of foliar N application in comparison with a complex regulation of CK on creeping bentgrass regrowth after de-submergence.

Heavy and frequent rainfall often creates a flooding environment for plants grown in a poorly drained site. As a result, plants can be completely covered by water, which negatively affects plant growth and physiology due to lack of oxygen. Plant survival in a submerged condition depends on severity of stress, species, and genotypes, as well as survival strategies (Bailey-Serres and Voeseke, 2008; Liu and Jiang, 2015; Yu et al., 2012). The mechanisms of escape and quiescence are strategies for plant tolerance to submergence stress (Bailey-Serres and Voeseke, 2008). Although these mechanisms can be crucial for a particular plant growing under submergence stress, recoverability after de-submergence could also play an important role in plant survival (Alpuerto et al., 2016; Gautam et al., 2016; Panda et al., 2008).

On de-submergence, variations of plant growth, physiological, and molecular alterations have been observed in different species or cultivars exposed to various submerged conditions (Alpuerto et al., 2016; Luo et al., 2011; Yu et al., 2012). In alligator weed (Alternanthera philoxeroides), resumed leaf growth was concomitant with restoration of maximal photosynthetic capacity, whereas rapid leaf regrowth in limpgrass (Hemarthria altissima) was found earlier than full recovery of photosynthesis (Luo et al., 2011). In perennial ryegrass (Lolium perenne), recovery of HT and growth rate varied with plant growth habits (fast vs. slow growth), but the carbohydrates fully recovered in all types of plants after 7 d of de-submergence (Yu et al., 2012). The submergence-tolerant M202 (Sub1) and sensitive M202 rice (Oryza sativa) lines had similar levels of decreased photosystem II photochemistry and stimulated the breakdown of protein under submergence; however, the tolerant M202 (Sub1) showed a more rapid return to homeostasis of these parameters than in the sensitive M202 (Alpuerto et al., 2016). After 3 d of de-submergence, the intolerant rice cultivar exhibited a low survival rate, a slow recovery of total ascorbic acid, and increased MDA formation, compared with the tolerant cultivar (Kawano et al., 2002). Collectively, the results suggest that recovery of physiological activities such as carbohydrate and antioxidant metabolism could play a role in whole plant regrowth after submergence stress.

Rapid recovery from submergence stress is a desirable trait that allows plants to more quickly resume normal function. Exogenous application of nutrients or plant growth regulators may promote plant recovery after de-submergence. Nitrogen (N) is a major component of chlorophyll molecules, therefore, exogenous N supply can enhance chlorophyll biosynthesis, which then reduces submergence stress-induced chlorosis. Foliar application of N after de-submergence substantially increased chlorophyll and soluble sugar contents, leaf area, dry matter weight, specific leaf weight, and tiller regeneration ability of rice (Gautam et al., 2014a, 2014b). Exogenous application of N also positively affected the growth and physiology of the plants under waterlogging conditions. Habibzadeh et al. (2013) reported that foliar application of N compounds before and after stress improved growth and antioxidant activities of canola (Brassica napus). In sugarcane (Saccharum officinarum), the addition of N compounds increased leaf area, shoot weight and nitrate reductase activity, and decreased MDA content, compared with untreated plants exposed to waterlogging stress (Radha et al., 2015). These research findings indicated that the exogenous N application could alleviate the adverse effects of flooding stress.

CK regulates a wide array of processes crucial to plant growth and development and abiotic stress tolerance (O’Brien and Benkova, 2013; Zwack and Rashotte, 2015). Ren et al. (2016) demonstrated that application of CK after waterlogging delayed leaf senescence, increased chlorophyll content, and improved photosynthetic performance of summer maize (Zea mays). At the molecular level, expression of the ipt gene controlling CK biosynthesis did not result in CK accumulation in submerged plants until the stress was removed (Huynh et al., 2005). Transgenic wheat (Triticum aestivum) plants carrying the ipt gene were more tolerant of flooding than wild type plants (Tereshonok et al., 2011). CK is beneficial in improving plant growth and development, but the effects of CK application on plant growth and physiology after de-submergence are not clear, particularly in comparison with the effectiveness of N application.

Creeping bentgrass is a widely used cool-season turfgrass on golf greens and fairways in temperate regions. These grass plants can be subjected to submerged conditions due to high and frequent precipitation combined with poor soil drainage. Recoverability from submergence stress is an important trait that influences grass quality and performance, but physiological mechanisms of the plants to de-submergence are not known. In practice, turfgrass managers often have few strategies at hand to recover grass from prolonged submergence stress. Whether foliar applications of N or CK alone, or in combination, promotes plant growth and metabolic activities after de-submergence is not well understood in plant species, including turfgrass. Therefore, the objective of the experiment was to investigate plant growth, carbohydrate, and antioxidant metabolism of creeping bentgrass to exogenous N and CK application after...
de-submergence. The results from the study would reveal physiological mechanisms of perennial plants to recovery after submergence.

**Materials and Methods**

**Plant materials and growing conditions.** Creeping bentgrass cultivars Penncross and 007 were chosen for the experiment. Penncross is a commonly used cultivar but is sensitive to waterlogging stress (Jiang and Wang, 2006). 007 is a newly developed cultivar that shows good overall turf quality (NTEP, 2016). On 2 July 2015, the seeds were sown in plastic pots (10 cm diameter, 9 cm deep) containing top soil in a greenhouse at Purdue University, West Lafayette, IN. Plants were watered four times a week and fertilized once a week with a soluble fertilizer (N–P–K, 24–8–16) (Scotts Inc., Marysville, OH) to supply 290 kg N/ha during the whole experiment. Plants were mowed three times a week at 3 cm. During the growing period, the average air temperatures and photosynthetic photon flux density (PPFD) in the greenhouse were 20 ± 1.5 °C and 700 μmol·m⁻²·s⁻¹, respectively. The grass was cut to 3 cm before submergence stress.

**Submergence stress.** Submergence stress began on 19 Nov. and ended on 3 Dec. 2015. The submerged pots were placed into plastic containers (60 cm length × 43 cm width × 31 cm height) and tap water (pH of 6.5, electrical conductivity of 1.2 dS·m⁻¹) was added to 10 cm above the grass canopy at the beginning of the treatment. During the periods of stress, the average air temperatures and PPFD were the same as the conditions indicated above. The average water temperature was 19.0 ± 1.0 °C. The nonsubmerged control pots were placed in the same size containers, but without the excess water, and irrigated every 2 d. No nutrients were supplied to the plants during the period of submergence stress. Algae were removed if they accumulated. On 3 Dec. 2015 (after 14 d of stress), all submerged plants were removed from the containers, and leaf samplings were made from randomly selected pots for various measurements for both the control and submerged plants, and the remaining pots were allowed for recovery.

**Application of nitrogen and CK after de-submergence.** The control and submerged plants were foliar-sprayed at 1, 2, 3, 7, and 14 d after de-submergence with the following treatments, respectively: 1) water (W); 2) 10 mM urea (N10); 3) 20 mM urea (N20); 4) 15 μM cytokinin (transzeatin riboside, ZR) (CK); 5) 10 mM urea + 15 μM cytokinin (N10CK); and 6) 20 mM urea + 15 μM cytokinin (N20CK). Pots were sprayed with 10 mL of solutions at each date for N and CK applications as described above in this paragraph. At 20 d after spraying, leaves were collected for various measurements.

**Measurements.** Various measurements were made for both control and submerged plants as well as for the plots that received post de-submergence chemical applications. HT was recorded by measuring the length of the longest leaf blades. Chl was measured on five randomly selected leaves per pot using the
SPAD 502 Plus Chlorophyll Meter (Spectrum Technologies, Inc., IL). Leaf tissues were collected for various measurements including DW, carbohydrate, and enzyme activity. The total WSC content was determined using the anthrone method (Koehler, 1952) with some modifications (Yu et al., 2012). Briefly, WSC was extracted from 50 mg of ground leaf tissue with 1 mL double-distilled water. The extract was shaken for 10 min and centrifuged at 11,000 g for 10 min, and the supernatant was collected. The extraction was repeated three times for all samples and the supernatants from each extraction were pooled for each sample. A 1 mL aliquot of extract was mixed with 7 mL freshly prepared anthrone [200 mg anthrone + 100 mL 72% (w/w) H2SO4] and placed in a boiling water bath for 8 min. After cooling, absorbance at 625 nm was read. The standard curve was determined using glucose in a range of 5 to 250 μg·mL⁻¹.

To extract the soluble protein, leaf tissues were frozen using liquid nitrogen, and ground into a fine powder. About 50 mg powder was mixed with 1 mL of extraction buffer (50 mM potassium phosphate, 1 mM ethylenediaminetetraacetic acid, 1% polyvinylpyrrolidone, pH 7.8). The mixture was centrifuged at 15,000 g for 30 min at 4 °C, and the supernatant was collected for enzyme assay. The TSP content was measured using the method of Bradford (1976). The activities of SOD, CAT, POD, and APX were assayed by using the methods of Zhang and Kirkham (1996) with minor modifications (Liu and Jiang, 2015).

Lipid peroxidation was measured in terms of MDA content (Dhindsa et al., 1981) with some modifications (Liu and Jiang, 2015). A 0.5 mL aliquot of supernatant was mixed with 2 mL of 20% trichloroacetic acid containing 0.5% thiobarbituric acid. The mixture was heated at 95 °C for 30 min, quickly cooled, and then centrifuged at 10,000 g for 10 min. The absorbance was read at 532 and 600 nm (Heath and Packer, 1968). MDA content was calculated using an extinction coefficient of 155 mm⁻¹·cm⁻¹.

**Results and Discussion**

Submergence and cultivar effects before de-submergence. ANOVA indicated significant submergence effects on all traits and cultivar effects on HT and WSC, but no submergence by cultivar interactions were observed for all traits except for APX (data not shown). Across cultivars, HT, Chl, DW, WSC, SOD, and APX activities significantly decreased by 52.4%, 50.9%, 44.1%, 43.6%, 54.4%, and 67.2%, respectively; whereas TSP, CAT, and POD activities, and MDA content increased by 18.0%, 1.2-fold, 96.0%, and 8.1-fold at 14 d of stress, respectively (Tables 1 and 2).

Other studies have shown that submergence either decreases or enhances plant growth, depending on the type of species and survival strategy (Bailey-Serres and Voesenek, 2008). Creeping bentgrass is tolerant to low cutting, which could allow the plants to conserve energy to increase survival under a short-term submergence stress. However, severe reductions in HT, DW, Chl, and WSC in Penncross and 007 indicated that physiological damages occurred to creeping bentgrass under the water. Decreased WSC of shoots and roots were also observed in ber-mudagrass (Cynodon dactylon) exposed to an increase of depth and duration of submergence (Tan et al., 2010). Moreover, large natural variations of plant growth and leaf color were found in perennial ryegrass accessions under 7 d of submergence, demonstrating diverse survival mechanisms for this species (Yu et al., 2012).

In this study, submergence stress altered the activities of antioxidant enzymes and increased MDA content, potentially leading to lipid peroxidation in creeping bentgrass (Table 2). Antioxidant enzymes play an important role in the removal of toxicity caused by an accumulation of reactive oxygen species under stress conditions. As shown in previous studies, the activities of antioxidant enzymes increased, decreased, and remained unchanged in different plant species, cultivars, and plant
tissues under submergence stress (Liu and Jiang, 2015; Panda and Sarkar, 2013; Tan et al., 2010). In rice, submergence increased MDA content (Lal et al., 2015), but the tolerant cultivar had higher antioxidant enzyme activities and chlorophyll content and lower MDA content than that of the intolerant cultivars during submergence (Panda and Sarkar, 2013).

Submergence, cultivar, and chemical effects after de-submergence. ANOVA indicated significant submergence, cultivar, and chemical application effects were found for most traits except for no cultivar effects on TSP, APX, and MDA, no submergence effects on POD, and no chemical application effects on POD, CAT, and SOD (data not shown). No submergence × cultivar × chemical application interactions were identified for all traits except for Chl, WSC, and MDA (data not shown).

Nitrogen and CK effects on growth and Chl after de-submergence. Different chemical applications resulted in significant differences in HT, DW, and Chl for plants previously exposed to nonstress or de-submergence conditions for both cultivars (Fig. 1). After 20 d of de-submergence, higher HT, DW, and Chl were generally found in plants treated with N10, N20, N10CK, or N20CK, whereas the lowest HT and DW were observed in plants treated with S-W or S-CK alone for both cultivars (Fig. 1). For example, compared with plants sprayed with S-W, Chl was ∼20%, 35%, 31%, 31%, 31%, 31%, 31%, 57%, and 61% higher for 007 under S-N10, S-N20, S-N10CK, and S-N20CK, respectively. The results demonstrated that separate W or CK treatments had no effect on the recovery of HT, DW, and Chl, but foliar applications of N alone or in combination with CK promoted plant growth and maintained leaf color. In a previous study, the positive effects of foliar N application were also noted in rice plants exposed to de-submergence (Gautam et al., 2014a, 2014b).

Nitrogen and CK effects on WSC, TSP, and MDA after de-submergence. Plant WSC varied with chemical applications under both nonstress and de-submergence conditions (Fig. 2). On recovery, WSC was not consistent between S-N10 and S-N10CK and between S-N20 and S-N20CK in Penncross, but CK alone led to higher values of WSC along with S-N10CK and S-N20CK. For 007, WSC was significantly higher under S-CK treatment than all other chemical applications at de-submergence. The N application promoted recovery of WSC in rice (Gautam et al., 2014a, 2014b), but our results showed that CK was more effective on WSC than N, at least for 007. Higher Chl found in S-N10CK and S-N20CK than in S-N10 and S-N20, along with positive CK effects on WSC as opposed to growth traits such HT and DW suggested a complex regulation of CK to the recovery of regrowth and physiology of creeping bentgrass. In maize (Z. mays), application of CK after dewaterlogging delayed leaf senescence, increased chlorophyll content, and improved photosynthetic performance (Ren et al., 2016). It seemed that CK effects on plant recovery largely depended on stress intensity and type of measurements after flooding stress. Variations of TSP were found among various chemical applications for plants previously exposed to nonstress and de-submergence conditions, with the exception of 007 at recovery (Fig. 2). The highest value of TSP was found with S-N20CK and the lowest with W; whereas for 007, S-N20, and S-N20CK had higher TSP than other applications. The application of higher N concentrations resulted in higher TSP in both cultivars. However, similar levels of TSP between S-W and S-CK, between S-N10 and S-N10CK, and between S-N20 and S-N20CK suggested that it was N, not CK, that accounted for the TSP recovery after submergence.

Generally, S-W or S-CK exhibited higher MDA content, whereas S-N10, S-N20, S-N10CK, or S-N20CK had lower MDA content in plants exposed to nonstress and de-submergence conditions (Fig. 2). The lowest MDA content found was under N10CK for Penncross and N20CK for 007 after de-submergence. The results indicated that W or CK alone did not limit the level of lipid peroxidation, but the application of N or N plus CK could alleviate oxidative stress injury.
after de-submergence. This was further supported by the higher HT, DW, and Chl levels under N or N plus CK than that of W or CK alone (Fig. 1). According to Gautam et al. (2014a, 2014b), an increased MDA content was also found in rice after de-submergence, but to a greater extent in the susceptible cultivar, suggesting that a potentially higher degree of oxidative injury contributed to poor tolerance of the susceptible cultivar.

Nitrogen and CK effects on antioxidant enzyme activities after de-submergence. All chemical applications did not alter activities of antioxidant enzymes in both cultivars under nonstress conditions, except for higher SOD for Penncross under N10CK (Fig. 3). On de-submergence, activities of the antioxidant enzymes were not consistent in the two cultivars of creeping bentgrass across various chemical applications. Specifically, higher SOD activities were found in Penncross treated with S-W or S-N10CK with lower SOD activities under S-N20 and S-N20CK; whereas plants treated with S-W, S-N20CK, or S-CK had higher SOD activities in 007 after de-submergence. S-N10CK significantly increased CAT activities in Penncross, but CAT activities remained unchanged across applications for 007 after de-submergence. S-N10CK significantly increased CAT activities in Penncross, but CAT activities remained unchanged across applications for 007 after de-submergence. After de-submergence, activities of the antioxidant enzymes were not consistent. The enhanced recovery by foliar application of N, or combined with CK, may be accomplished mainly through promoting growth, Chl, and TSP and reducing lipid peroxidation after de-submergence.

Literature Cited
Ren, B., Y. Zhu, J. Zhang, S. Dong, P. Liu, and B. Zhao. 2016. Effects of spraying exogenous hormone 6-benzyladenine (6-BA) after waterlogging on grain yield and growth of summer maize. Field Crops Res. 185:96–104