Changes of Antioxidant Defense System and Fatty Acid Composition in Bermudagrass under Chilling Stress

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ADDITIONAL INDEX WORDS. low temperature, genotype, glutathione, gene expression, reactive oxygen species

ABSTRACT. Bermudagrass (Cynodon dactylon) is a typical and widely used warm-season turfgrass. Low temperature is one of the key environmental stress limiting its utility. However, little information is available about the differences of cold response between bermudagrass genotypes. Here, we analyzed antioxidant defense system and fatty acid composition in cold-resistant genotype WBD128 and cold-sensitive genotype WBDg17 exposed to chilling stress. Low temperature (4 °C) significantly decreased the relative water content, whereas increased the H2O2 and O2•− contents, more profoundly for WBDg17. Under chilling condition, WBD128 had higher anti O2•− activity than WBDg17. Besides, the contents of total glutathione, reduced glutathione (GSH) and its oxidized form (GSSG) were markedly increased by low temperature in both genotypes, whereas WBD128 had significantly higher values of GSH, total glutathione, and GSH/GSSG ratio than WBDg17. Moreover, chilling stress increased saturated fatty acids (SFAs) percentage (palmitic acid and stearic acid) in WBDg17. After chilling treatment, the proportion of linoleic acid decreased in both genotypes, particularly in WBDg17. As for unsaturated fatty acids (UFAs), the percentage of linolenic acid was increased in WBD128. In addition, chilling treatment decreased the values of double bond index (DBI), UFA/SFA ratio as well as degree of unsaturation in WBDg17. Finally, chilling stress altered the expression patterns of the genes, which encode one kind of late embryogenesis abundant proteins (LEA), superoxide dismutase (Cu/Zn SOD), C-repeat-binding factor/DRE-binding factor (CBF1), and peroxidase (POD-2). Collectively, our results revealed that natural variation of chilling tolerance in bermudagrass genotypes may be largely associated with the alterations of antioxidant defense system and fatty acid composition.

Low temperature is one of the most detrimental abiotic stresses, which limits plant growth and productivity as well as distribution (Burke et al., 1976). The mechanisms of cold-induced injury are complex, and vary across different species and hardness degree (Baek and Skinner, 2003; Burke et al., 1976; Ishwari and Palta, 1989). The status of the plasma membrane is crucial to cellular response when a plant is exposed to cold stress (Thomashow, 1999). Numerous mechanisms have been proposed to be involved in membrane damage under cold stress.

This includes structural transitions and membrane phase transitions (Pearce and Willison, 1985), injury to membrane-bound ATPase (Iswari and Palta, 1989), and loss of bound water (Weiser, 1970).

Oxidative stress also has been suggested to be one of the causes of cold-induced damage (Halliwell and Gutteridge, 2015; McKersie and Bowley, 1997), which occurs when there are excessive free cellular radicals. Under normal condition, reactive oxygen species (ROS) are generated on a regular basis and at a low level (Arora et al., 2002). When plants are subjected to stresses, excessive ROS are produced, which can induce plant injury, including peroxidation of cell membrane components, enzymes denaturation, and DNA strands distortion (Halliwell and Gutteridge, 2015). To alleviate the detrimental effects of oxidative stress, plants have developed...
Various and effective antioxidant systems necessary for scavenging ROS (Mittler, 2002), including enzymes and nonenzymatic antioxidants; for example, L-γ-glutamyl-L-cysteinyl-glycine. Antioxidant enzymes and nonenzymatic antioxidants were generally induced by stresses, and the elevated activities enhanced resistance to stress (Allen, 1995). Glutathione and glutathione reductase has been reported to participate in the redox regulation, as components of the ascorbate–glutathione cycle (Foyer et al., 1997). Under chilling conditions, the ratio of GSH to glutathione disulfide decreases. It is because that GSH is used for the reduction of excess H₂O₂ in the ascorbate–glutathione cycle (Foyer et al., 1997). Besides, the increase of H₂O₂ and the succeeding changes of the GSH/GSSG ratio may serve as signals to activate specific transcription factors and antioxidant enzymes (Kocsy et al., 2001). In addition, GSH is involved in numerous physiological processes in plants, which include control of cell division and gene expression (Noctor et al., 1998), transport and storage of sulfur (Rennenberg and Brunold, 1994), and removal of toxic metabolite by the formation of glutathione S (GS) conjugates (Mauch and Dudler, 1993). Changes in the GSH content and GSH/GSSG ratio may modulate these physiological processes in plants’ response to the unfavorable environmental conditions.

Fatty acids can be divided into SFAs and UFAs. UFAs have double bonds (one or more), and more common cis-double bonds help maintain membrane fluidity, which is pivotal for plant survival during cold stress (Cyril et al., 2002; Lehninger, 1977; Vigh et al., 1998). Causally, under cold stress, cellular membranes undergo a phase transition from liquid crystalline (highly fluid) to gel phase (more rigid). In the gel phase, lipids are highly ordered, which hampers regular physiological functions, and makes the cellular membrane more permeable (Cyril et al., 2002). Multiple biochemical and biophysical alterations occur to prevent the phase transition. In addition, the degree of unsaturation dramatically impacts the temperature range in which cell membrane undergo pernicious phase change (Vigh et al., 1998). Thus, increasing the percentage of UFA in membrane fatty acid composition contributes to the maintenance of plasma membrane stability, integrity, as well as function (Steponkus, 1984).

Bermudagrass is a typical warm-season turfgrass species, which grows in warm climatic regions and exhibits remarkably great stress resistance. This turfgrass has been widely applied on account of its superior merits of fast reproduction and traffic tolerance, particularly in parks, sport fields, and lawns (Fan et al., 2014; Hu et al., 2016a; Shi et al., 2014; Zhang et al., 2008). As a warm-season turfgrass, the optimum temperature for growth is 26 to 35 °C, and low temperature is considered as a key environmental factor limiting bermudagrass growth and utility (Fan et al., 2014). However, limited research has been done on chilling resistance in bermudagrass. Therefore, detailed information about chilling stress response in bermudagrass is urgent and crucial for grass engineering (Fagerness et al., 2002; Shi et al., 2014; Zhang et al., 2008, 2011a, 2011b).

This study was designed to investigate the differences of chilling stress response between the cold-sensitive bermudagrass genotype WBDg17 and the cold-resistant genotype WBD128, and explore the possible mechanisms involved in chilling stress response in bermudagrass. Several physiological indicators, combined with fatty acid composition and gene expression patterns were investigated.

**Materials and Methods**

**PLANT MATERIAL AND GROWTH CONDITIONS.** Two wild types of bermudagrass, WBDg17 and WBD128, used in the study had been selected by some colleagues in our laboratory and have been reported by Chen et al. (2010). Initially, 128 bermudagrass accessions were used in the screening test. The plants were treated with 4 °C for 21 d, and the control was maintained at (30 °C day/25 °C night). Several parameters were determined every week including transpiration rate and growth rate. After the first round of screening, 10 relatively cold-tolerant and cold-sensitive accessions were selected. The selected accessions were then treated with –5 °C for 4 h to further screen for the relatively most cold-tolerant and cold-sensitive genotypes. Finally, the most promising cold-tolerant and cold-sensitive bermudagrass genotypes were selected. The cold-sensitive (S) genotype WBDg17 was collected from Xiaojiang City, Zhejiang Province, China (lat. 27°34.258’N, long. 120°27.383’E), whereas the cold-resistant (R) genotype WBD128 was collected from Baise City, Guangxi Province, China (lat. 24°51.397’N, long. 106°33.288’E). Uniform stolons of the two genotypes were planted in plastic pots (7.5 cm diameter and 9.0 cm deep) filled with matrix (brown coal soil) on 5 Apr. 2016. These pots were kept in the greenhouse for 2 months to establish, with growth conditions of (light/dark) 12/12 h, 30/25 °C, 240 μmol·m⁻²·s⁻¹, and 60% relative humidity (5 Apr. to 5 June 2016). During this period, plants were watered every 2 d and fertilized weekly with half-strength Hoagland’s solution (Hoagland and Arnon, 1950), and the plants were not mowed.

**TREATMENTS.** All established bermudagrass were transferred to growth chambers (HP300GS-C; Wuhan Ruihua Instrument and Equipment Co., Wuhan, China) for 3 d of adaptation, with conditions of (light/dark) 12/12 h, 30/25 °C, 240 μmol·m⁻²·s⁻¹, and 60% relative humidity. The lights they used were cold light illuminators. On 8 June 2016, plants were subjected to two temperature treatments: the control and chilling treatment, which were performed by two growth chambers. For control chamber, the conditions were (light/dark) 12/12 h, 30/25 °C, 240 μmol·m⁻²·s⁻¹, and 60% relative humidity. And the lights they used were cold light illuminators. Such period, fully extended bermudagrass leaves were collected at 0, 3, 6, 12, 24, and 48 h after chilling treatment were subjected to two temperature treatments: the control and chilling treatment, which were performed by two growth chambers. For control chamber, the conditions were (light/dark) 12/12 h, 30/25 °C, 240 μmol·m⁻²·s⁻¹, and 60% relative humidity. Low temperature (4 °C) was selected according to many previous studies (Fan et al., 2014; Hu et al., 2016a, 2016b; Shi et al., 2014) and the initial screening test of bermudagrass (unpublished data). Actually, the day temperature and the night temperature were 4.9 to 5.6 °C and 4.3 to 4.6 °C, respectively, in the growth chambers for the cold treatment. Plants of each species were subjected to the two treatments for a week (8 to 15 June 2016). During the treatment period, fully extended bermudagrass leaves were collected at 0, 3, 6, 12, 24, and 48 h for RNA extraction, and at 7 d for measurements of RWC, enzyme activity, ROS, and fatty acids.

**MEASUREMENTS.** RWC was determined on 16 June 2016 according to the method described by Barrs and Weatherley (1962) with slight modifications. In brief, leaves were clipped and weighed [fresh weight (FW)], then transferred into small petri dishes filled with deionized water. The leaves were soaked in water for 24 h at 25 °C, thereafter the leaves were weighed after removal of excessive moisture by blotting paper [turgid weight (TW)]. Subsequent drying of leaves was done at 70 °C for 24 h to measure dry weight (DW). Leaf RWC was calculated following the formula: (FW – DW)/(TW – DW) × 100.
Crude enzyme extraction was performed on 17 June 2016 according to Fan et al. (2014). Fully expanded leaves (0.2 g) were ground into powder in liquid nitrogen, and then the powder was transferred into 15-mL tubes filled with 4 mL sodium phosphate buffer (150 mM, pH 7.0). After 5 h extraction at 4 °C, the extraction solution was centrifuged for 25 min at 8000 g at 4 °C. The supernatant was decanted into 2-mL microcentrifuge tubes for the subsequent physiological assays.

The ROS contents were measured on 18 June 2016 by using different kits. The measurements of H$_2$O$_2$ and O$_2$ concentrations were, respectively, performed by using H$_2$O$_2$ assay kits (A064; Nanjing Jiancheng Bioengineering Institute, Nanjing, China), and Plant O$_2$ ELISA Kit (10-40-488; Beijing Dingguo, Beijing, China) according to the manufacturer’s instructions as introduced by Shi et al. (2013). The anti-O$_2^-$ activity was measured by using the assay kit (A052; Nanjing Jiancheng Bioengineering Institute). For the anti O$_2^-$ activity determination, the electron transport material and chromogenic agent were added to react with superoxide anion free radical, which produced by the system of xanthine and xanthine oxidase. After chemical reaction, the red mixture was quantified at 550 nm of absorbance. Moreover, the concentrations of GSH, GSSG, and total glutathione were determined by using the GSH and GSSG Assay Kit (S0053; Beyotime, Shanghai, China) in accordance with the manufacturer’s instructions as described by Shi et al. (2013, 2014). The GSH/GSSG ratio was calculated based on GSH and GSSG concentration.

Total lipid was extracted from bermudagrass leaves on 20 June 2016 by using the method described by Mishra et al. (2015) with some modifications. About 0.3 g bermudagrass leaves were ground into fine powder and then extracted in 10 mL chloroform–methanol–phosphate buffer (1:2:0.9 v/v/v, pH 7.5). Then 50 mL of 2 mg·mL$^{-1}$ heptadecanoic acid (C17:0 as internal standard) was added. The homologous fatty acid methyl esters (FAMEs) of these fatty acids were obtained by transmethylation (Kumari et al., 2013). Transmethylation was conducted by adding 1 mL of NaOH (1% v/v in methanol) in a 15-mL tube that contained extracted lipid, the mixture was thereafter heated at 55 °C for 15 min. Thereafter, 2 mL of methanolic HCl (5% v/v) was added followed by heating at 55 °C for 15 min, then the mixture solution was added with 3 mL of ultrapure water–hexane mixture (1:2 v/v). These derivative FAMEs were extracted in hexane three times and dried with application of a bench-top centrifugal concentrator (Centrivap; Labogene, Lynge, Denmark). Finally, the dried extraction was dissolved in 200 μL of hexane.

The fatty acids were determined by using gas chromatography–mass spectrometer [GC-MS (7890A/5975C; Agilent Technologies, Palo Alto, CA)]. In brief, 200 μL of derivative solution was transferred into a DB-5MS capillary (30 m × 0.25 mm × 0.25 mm; Agilent Technologies). The GC-MS was programmed as follows: the initial GC oven temperature was at 70 °C for 2 min, then from 70 to 230 °C with 3 °C·min$^{-1}$ increment and 230 °C maintained for 10 min, and subsequently, from 230 to 270 °C with 10 °C·min$^{-1}$ increase, and then held for 10 min. Injection volume was 1 μL, helium was used as the carrier gas, and the flow rate was set at 1 mg·mL$^{-1}$. The determination was carried out by electron impact ionization at 70 eV in the full scan mode (m/z 30–650). The samples of bermudagrass were quantified against an internal standard (100 μg heptadecanoic acid), and the content of each fatty acid was expressed as a proportion of the total fatty acids present in the sample. The DBI was calculated by using the formula: Index = (16:1) + (18:1) + 2[(16:2) + (18:2)] + 3[(18:3)] (Larkindale and Huang, 2004). Parentheses indicate the proportion of the total fatty acid amount, which was composed of each fatty acid species.

For analysis of real-time quantitative reverse transcriptase polymerase chain reaction (qRT-PCR), total RNA in bermudagrass leaves was isolated and purified with application of Trizol reagent (Invitrogen, Carlsbad, CA) according to the method we described earlier (Hu et al., 2016a). RNA (about 2.5 μg) was reversely transcribed by using M-MLV reverse transcriptase (Promega, Madison, WI) and oligo (dT) primers following the operation manual. Expressions of the selected genes in treated bermudagrass leaves were analyzed by qRT-PCR with application of the fluorescent intercalating dye SYBR Green and StepOnePlus™ Real-Time PCR System (Applied Biosystems, Foster City, CA) for detection. The gene-specific primers of selected genes listed in Table 1 were synthesized based on previously reported sequences, whereas an ACTIN was used as the standard control in the reaction. The RT-PCR procedure was performed according to the method described by Chen et al. (2010).

**Statistical analysis.** Three biological replicates were used in all the experiments, and values were given as mean ± SD. Statistical analyses were performed by one-way analysis of variance combined with independent-samples t test. Means were separated at a significant level of P < 0.05, by using the statistical package SPSS (version 16.0; IBM Corp., Armonk, NY), as well as Excel (2010 for Windows; Microsoft Corp., Redmond, WA).

**Results**

**Physiological response to chilling stress.** As shown in Fig. 1, there was no significant difference in RWC between the two genotypes under control conditions. However, after 7 d of chilling treatment the values were decreased by 11.33% and 2.76% in the cold-sensitive genotype WBDg17 and cold-tolerant genotype WBD128, respectively. This result indicated that leaves of WBDg17 lost water more than WBD128. Consistently, a better growth condition was also observed in the WBD128 (Supplemental Fig. 1).

Table 1. Genes (POD-2, SOD, LEA, and CBF1) involved in response to chilling stress in cold-resistant bermudagrass genotype WBD128 and cold-sensitive genotype WBDg17 under chilling stress (4 °C) and their forward and reverse primer sequences.

| Gene name | Primer sequences (5′-3′) |
|-----------|-------------------------|
| **POD-2** | F AGGCGACGCGGCTGAAAGG R CCCGACGAGTCGTCGAGAA |
| **SOD**   | F TGGGAAACATTGTTGCCAACA R GCCAACAAACACATGGC |
| **LEA**   | F TCGTCCGCTTGGTTTCATCA R GAGGCCGCAACACAGAGA |
| **CBF1**  | F ACCAAATCCGCGGAGCGC R CGATGCGGCAATTTGAGCA |

$^a$POD-2 = gene that encodes one kind of peroxidase; SOD = gene that encodes one kind of superoxide dismutase; LEA = gene that encodes one kind of late embryogenesis abundant proteins; CBF1 = gene that encodes one kind of C-repeat-binding factor/DRE-binding factor.

$^b$Primer sequences. F and R represent forward and reverse, respectively.
H$_2$O$_2$ and O$_2^-$ are two components of ROS, which if produced in excess is harmful for plant. In the study, there were no differences in H$_2$O$_2$ and O$_2^-$ concentrations between the two genotypes under normal condition (Fig. 2A and B). However, when exposed to chilling treatment for 7 d, H$_2$O$_2$ content was 28.70% and 21.82% higher in WBDg17 and WBD128 than the control, respectively. Accordantly, O$_2^-$ contents were significantly increased in both genotypes, particularly in WBDg17 (by 76.86% for WBDg17 and 49.98% for WBD128). Similar result was also obtained by nitro blue tetrazolium chloride staining (Supplemental Fig. 2). These results showed that the cold-induced ROS concentration was higher in WBDg17. Anti-O$_2^-$ activity was used to evaluate the activity of scavenging superoxide anion in the study. As shown in Fig. 2C, chilling stress increased the activities of anti-O$_2^-$ in both genotypes, and to a greater extent in WBD128.

**CHANGES IN ANTIOXIDANT ACTIVITY UNDER CHILLING STRESS.** As shown in Fig. 3A, chilling treatment dramatically increased GSH content by 49.29% and 210.74% for WBDg17 and WBD128, respectively. Similarly, the contents of GSSG were increased in both genotypes by chilling exposure (by 153.03% for WBDg17 and 142.11% for WBD128), the level in WBD128 was lower than that in WBDg17 (Fig. 3B). Furthermore, chilling stress decreased the GSH/GSSG ratio by 119.18% in WBDg17, but increased the value by 16.64% in WBD128 (Fig. 3D).

**CHANGES IN FATTY ACID COMPOSITION UNDER CHILLING STRESS.** In the present study, four major fatty acids were identified and quantified. There were two SFAs, including palmitic acid (C16:0) stearic acid (C18:0), and two UFAs, containing linoleic acid (C18:2) and linolenic acid (C18:3). As shown in Fig. 4, the palmitic acid content was 16.46% in WBDg17 and 20.10% in WBD128 under control condition. However, after chilling stress exposure the percentages were 21.79% and 19.97% in WBDg17 and WBD128, respectively (Fig. 4A). The stearic acid content was increased from 5.22% to 6.8% in WBDg17 by end of 7 d of chilling treatment (Fig. 4B). In WBD128, a minor increase from 5.32% to 6.06% was observed. In absence of chilling stress, the linoleic acid (C18:2) accounted for 7.38% and 13.06% of total fatty acid content in WBDg17 and WBD128, respectively. After 7 d of chilling treatment, the linoleic acid contents were 2.37% in WBDg17 and 8.54% in WBD128 (Fig. 4C). As for C18:3, the content was 70.50% in WBDg17 and 61.52% in WBD128 under control condition.
However, chilling treatment decreased the percentages to 68.69% in WBDg17, inversely, increased to 64.85% in WBD128 (Fig. 4D).

The degree of unsaturation was 77.88% in WBDg17 and 74.58% in WBD128 under control conditions (Fig. 5A). Under chilling stress, the level was decreased by 8.67% in WBDg17 but almost unaltered in WBD128. To explore further the difference in fatty acid composition between the two genotypes, the UFA/SFA ratio was calculated. In absence of chilling stress, the UFA/SFA ratio was higher for WBDg17 than that of WBD128 (Fig. 5B). Chilling treatment significantly decreased the value by 30.20% for WBDg17, whereas had no notable effects in WBD128. Consequently, WBD128 had a higher level of UFA/SFA ratio than WBDg17. Similarly, chilling stress decreased the DBI value by 6.78% for WBDg17, whereas there was no obvious difference in the value for WBD128 (Fig. 5C). These results suggested that maintenance of higher total UFAs composition, particularly linoleic acid, maybe superior prerequisite for superior chilling tolerance in WBD128.

GENE EXPRESSION OF CULTIVAR VARIATION IN RESPONSE TO CHILLING STRESS. To explore further the differences in gene expression pattern of the two genotypes response to chilling stress, we analyzed the expression of several genes. In the present study, an increase in transcripts of CBF1 was observed in both genotypes after chilling treatment at 4°C (Fig. 6A). In WBD128, the expression of CBF1 reached peak at 6 h of chilling treatment, whereas at 24 h in WBDg17. The peak value for WBD128 was significantly higher than that for WBDg17. Thereafter, levels of CBF1 expression in both genotypes declined to lower level, which was 3-fold higher than the control level for WBDg17 and 2-fold for WBD128.

A relatively slow upregulation of LEA at 24 h of chilling treatment was observed in WBDg17 (Fig. 4D). Differently, the expression of LEA in WBD128 rapidly declined at 3 h of chilling treatment, then substantially increased and reached
peak at 6 h, thereafter decreased to the control level (Fig. 5B). The level of LEA expression was significantly higher in WBD128 at 0, 3, 6, and 48 h.

In addition, two genes encoding antioxidant enzymes were analyzed. Generally, chilling stress induced both Cu/Zn SOD and POD-2 transcriptions. In WBDg17, expression of Cu/Zn SOD rapidly increased and reached the peak at 3 h of chilling treatment, which was 3-fold higher than the control. Thereafter, the expression declined from 6 to 24 h, and maintained at a relatively lower level at 48 h. However, in WBD128, a slight increase of Cu/Zn SOD expression was observed from 3 to 6 h of chilling treatment, then decreased from 12 to 24 h, and finally increased mildly at 48 h. Compared with the cold-sensitive genotype, Cu/Zn SOD transcription in WBD128 was significantly lower at 3, 6, 12, and 24 h of chilling treatment, whereas higher at 24 and 48 h. Similarly, in both genotypes, chilling dramatically induced POD-2 expression at 3 h, and eventually declined to the least level at 6 h, then up-regulated at 12 h, and finally declined moderately at 24 to 48 h. Comparison of POD-2 expression in the two genotypes revealed higher expression in WBD128 than that in WBDg17 at 3, 24, and 48 h.

Discussion

Low temperature is one of the most detrimental environmental stresses, which can induce cell damage and restrain plant growth (Catalá et al., 2014; Shi et al., 2012; Yang et al., 2006). The mechanisms of chilling stress injury are exceedingly complex, and the causes of injury are not clearly understood. To resist cold stress, multiple changes occur in plant, which include membrane stability, enzymes activity, gene expression, and metabolism (Fan et al., 2014; Hu et al., 2016a; Shi et al., 2014). During evolutionary process, plants have acquired adaptive responses to survive extreme temperatures. Many plants from temperate regions increase in freezing tolerance upon exposure to low, nonfreezing temperatures, which is a phenomenon known as cold acclimation (Thomashow, 1999). The process of cold acclimation is extremely complex, which involves numerous physiological and biochemical changes, such as alterations in fatty acid composition and the accumulation of sugars, amino acid, and other osmolytes (Catalá et al., 2014). Bermudagrass, an important warm-season turfgrass species, may take at least 3 to 4 weeks to complete cold acclimation process, and it may experience a short period of chilling stress in late fall and early spring in some regions.

RWC, a vital indicator of the internal water status of plant, is used to evaluate the hydration state or internal water deficit degree in plant during dehydration (Rachmilevitch et al., 2006). In the present study, low temperature decreased the RWC in both genotypes, particularly in WBDg17. Our results indicated that WBD128 had significantly higher internal water status, which can be beneficial by providing a suitable internal environment for physiological function in plants (Fig. 1).

As one of the major environmental stresses, cold stress causes oxidative stress by rapid and excessive production of ROS. As observed in Fig. 2, chilling treatment dramatically increased H2O2 and O2− contents in both genotypes, and the levels were significantly higher in WBDg17. Meanwhile, the activities of anti O2− were enhanced by low temperature in both genotypes, especially for WBD128. Luo et al. (1997) indicated that the injury is mitigated when activity of scavenging superoxide anion is enhanced in plants exposed to adverse environmental stress. Our result was consistent with Jeon et al. (2010), Hu et al. (2013), and Shi et al. (2014). Plants have evolved complicated antioxidant defense systems to efficiently eliminate excessive ROS, enzymatic antioxidants and non-enzymatic antioxidants are the high-efficiency ways to protect plant from ROS damage (Apel and Hirt, 2004; Mittler, 2002). In our previous study, chilling treatment significantly induced activities of SOD, POD, ascorbate peroxidase (APX), and GS.

Fig. 5. Changes of (A) degree of unsaturation, (B) the ratio of unsaturated fatty acid (UFA) to saturated fatty acid (SFA), and (C) double bond index (DBI) in cold-sensitive genotype WBDg17 and cold-resistant bermudagrass genotype WBD128 under chilling stress. CK was the control with (light/dark) 12/12 h and 30/25 °C. Chilling represented (light/dark) 12/12 h and 4/4 °C conditions. All the regimes were treated for 7 d. Mean values and SD were calculated from three biological repeats. Different letters represented statistical difference significant among different regimes by Duncan’s multiple range tests (P < 0.05).
transferase in bermudagrass (Hu et al., 2016a, 2016b). GSH has been reported to react chemically with ROS, which functions as a free radical scavenger and is vital for cellular defense against ROS (Şahin and Gümüşlü, 2004). A high ratio of GSH to GSSG is conducive to improved chilling tolerance (Kocsy et al., 2001). Walker and Mckersie (1993) indicated that the GSH content and the GSH/GSSG ratio were significantly higher for WBDg17 and WBD128 based on independent-samples t test.

SOD catalyzes the process of converting two superoxide radicals (O$_2^-$) to H$_2$O$_2$ and O$_2$, including Cu/Zn SOD, Mn-SOD, as well as Fe-SOD (Alscher et al., 2002). These isoenzymes have different molecular structures but catalyze the same reaction process. Thereafter, the H$_2$O$_2$ was removed by APX, POD, GPX resulting in production of H$_2$O and O$_2$ (Apel and Hirt 2004; Foyer et al., 1997; Mittler, 2002). Increased expression of antioxidant systems has been suggested to render the plant (Thomashow, 1999). Among this pathway, CBF can activate numerous downstream cold-regulated (COR) genes, which can render plant resistance to cold stress. In the study, the level of CBF1 expression induced by chilling stress was generally higher for WBD128 than that for WBDg17 (Fig. 6A). The result revealed that the more rapidly and greater increase of CBF1 expression may activate more COR genes transcription in WBD128, which is beneficial to enhance resistance to chilling stress. LEA has been confirmed to play a crucial role in plant response to cold stress, especially for stabilization of cell membrane (Tunnelciffe and Wise, 2007). In this study, the generally lower level of LEA expression in WBDg17 can lead to less accumulation of LEA protein, which may contribute to enhance the sensitivity to chilling stress (Fig. 6B).

Biological membrane is considered as a primary target site for destruction by cold stress (Cyril et al., 2002), whereas electrolyte leakage (EL) and malondialdehyde (MDA) are the typical indicators to evaluate the integrity of cell membrane. In our previous studies, chilling treatment significantly increased the values EL and MDA in both genotypes, and to a higher extent in WBDg17 (Hu et al., 2016a, 2016b). To further explore the possible mechanism underlying membrane peroxide, the lipid composition was analyzed. It has been determined that there are changes in lipid composition of membranes when plants are exposed to cold stress (Cyril et al., 2002; Vigh et al., 1998). Sufficient evidence indicate that the biophysical state of cellular membrane, mainly composed of two lipid bilayers, is pivotal for maintaining membrane integrity under cold stress (Vigh et al., 1998). It has been postulated that the primary cold injury results from membrane lipid transition from fluid phase to gel phase (Lyons, 1973). Different with SFAs, UFAs are

![Fig. 6. Changes of (A) relative expression of gene that encodes one kind of C-repeat-binding factor/DRE-binding factor (CBF1), (B) late embryogenesis abundant proteins LEA, (C) superoxide dismutase Cu/Zn SOD, (D) peroxidase POD-2 in cold-sensitive genotype WBDg17 and cold-resistant bermudagrass genotype WBD128 under chilling stress. CK was the control with (light/dark) 12/12 h and 30/25 °C. Chilling represented (light/dark) 12/12 h and 4/4 °C conditions. All the regimes were treated for 7 d. Total RNA were isolated from leaves treated at 4 °C for 0, 3, 6, 12, 24, and 48 h, respectively. Quantitative real time polymerase chain reaction was repeated for three times. Columns marked with star indicated statistical significance between WBDg17 and WBD128 based on independent-samples t test.](image-url)
arranged loosely in the lipid bilayers, which results from the nonlinearity of the fatty acid arrangement introduced by double bonds (Lehninger, 1977). It has been reported that a positive association of UFA content and membrane fluidity is vital for proper cellular metabolism and biological function under environmental stresses (Hoekestra et al., 2001). During chilling exposure, UFAs are vital for maintaining membrane fluidity. This might be attributed to the function of lowering the transition temperature, partly by introducing bends in the linear fatty acyl chains (Cyril et al., 2002). This protects the acyl chains of adjacent lipids from tight packing, which is beneficial for maintaining molecular interactions (Cyril et al., 2002). In the present study, as observed for the two SFAs, chilling stress remarkably increased palmitic acid (16:0) and stearic acid (C18:0) proportions of total lipid in the WBDg17, whereas had no significant effects on the two lipids proportion in WDB128 (Fig. 4A and B). Increase of C16:0 and C18:0 relative contents indicated that the membrane rigidity is enhanced in WBDg17, which results from linearity of lipid arrangement in cell membrane phospholipid phase (Cyril et al., 2002; Lehninger 1977). This result was consistent with the study of Zhong et al. (2011). Compared with WBDg17, the lower percentage and minor change of SFAs might have contributed toward maintenance of cell membrane liquidity in WDB128.

The percentages of linoleic acid (C18:2) was higher in WDB128 than that in WBDg17 under normal condition (Fig. 4C). After 7 d of chilling treatment, the percentages decreased in both genotypes, and WDB128 had a significantly higher level of linoleic acid. For linolenic acid, chilling stress observably induced the rise of percentage in WBD128, whereas there was no obvious change in WBDg17 (Fig. 4D). These results suggested that the biosynthesis of linolenic acid might be regulated in plant response to chilling stress. The process of linolenic acid synthesis is complex and catalyzed by a battery of desaturase enzymes. First, stearoyl-acyl-carrier-protein (ACP) desaturase converts stearic acid to oleic acid (C18:1). Then C18:1 is catalyzed to form linoleic acid (C18:2) by oleoyl-phosphatidylcholine (PC) desaturase. Finally, the formation of linolenic acid (C18:3) is carried out by linoleoyl-PC desaturase (Cyril et al., 2002). On the basis of the observed results, we hypothesized that the activities of desaturases may be regulated by chilling stress. DBI, an indicator for evaluating unsaturation level of fatty acids (Zhong et al., 2011), was applied in the present study. Under chilling condition, the value of DBI declined in WBDg17, whereas there was no significant change in WDB128 (Fig. 5). Similar results were observed in the parameters of UFA/SFA ratio and degree of unsaturation. In addition, values of the two indicators were operably induced the rise of percentage in WBD128, whereas there was no obvious change in WBDg17 (Fig. 4D). These results suggested that the biosynthesis of linolenic acid might be regulated in plant response to chilling stress. The mechanism in bermudagrass response to chilling stress is complex. Superior resistance to low temperature in WBD128 may be mainly related to greater response of antioxidant defense system and relatively higher proportion of UFAs. They are jointly conductive for stability maintenance and integrity of membrane to sustain normal functions, thus to enhance chilling resistance in bermudagrass.

Conclusion

The mechanism in bermudagrass response to chilling stress is complex. Superior resistance to low temperature in WBD128 may be mainly related to greater response of antioxidant defense system and relatively higher proportion of UFAs. They are jointly conductive for stability maintenance and integrity of membrane to sustain normal functions, thus to enhance chilling resistance in bermudagrass.

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Supplemental Fig. 1. Representative leaves of cold-resistant (R) bermudagrass genotype WBD128 and cold-sensitive (S) genotype WBDg17 under different conditions. CK was the control with (light/dark) 12/12 h and 30/25 °C. Chilling represented (light/dark) 12/12 h and 4/4 °C conditions. All the regimes were treated for 7 d.

Supplemental Fig. 2. Nitro blue tetrazolium chloride staining images of representative leaves of cold-resistant (R) bermudagrass genotype WBD128 and cold-sensitive (S) genotype WBDg17 under different conditions. CK was the control with (light/dark) 12/12 h and 30/25 °C. Chilling represented (light/dark) 12/12 h and 4/4 °C conditions. All the regimes were treated for 7 d.