Relationships of Cold Acclimation and Antioxidative Enzymes with Chilling Tolerance in Cucumber (Cucumis sativus L.)

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ABSTRACT. To determine whether chilling tolerance is related to cold acclimation, changes in physiological responses and activity of antioxidative enzymes were investigated in leaves of cucumber (Cucumis sativus L.) grown in controlled environments. Plants were exposed to 15°C (cold-acclimated) or 25°C (nonacclimated) for 3 days, under 50 µmol·m⁻²·s⁻¹ photosynthetic photon flux and 70% relative humidity. Plants were then exposed to 8°C chilling temperature for 3 days, and allowed to recover in a growth chamber at 25°C for 3 days. Measurements of leaf water content, cellular leakage, lipid peroxidation, chlorophyll a fluorescence, isozyme profile, and lipid peroxidation

ADDITIONAL INDEX WORDS. chilling injury, chlorophyll a fluorescence, isozyme profile, lipid peroxidation

Crops of tropical and subtropical origins are now cultivated in areas where temperatures may be below the optimum required for their normal growth and development. Like other thermophilic crops, cucumber (Cucumis sativus) is susceptible to chilling temperatures (Wang, 1990). One method used to improve cold tolerance in plants is cold acclimation (Salveit and Morris, 1990). This technique entails exposure of chilling-sensitive tissue to temperatures slightly above chilling for a certain period of time. Cold acclimation can reduce injury caused by chilling temperature (Gilmour et al., 1988). Acclimation of chilling-sensitive plants, such as maize (Zea mays L.) and tomato (Lycopersicon esculentum [L.] Mill.), to low temperature reduces chilling injury (Anderson et al., 1995; Venema et al., 2000). Although mechanisms of chilling injury and tolerance have been studied in cucumber plants (Erez et al., 2002; Reyes and Jennings, 1994; Terashima et al., 1994), little is known about the role of antioxidative enzymes in chilling tolerance.

Various mechanisms have been suggested to account for chilling injury or tolerance in plants (Basra, 2001; Lee and Lee, 2000; Wise and Naylor, 1987). Active oxygen species (AOS), such as superoxide (O₂⁻), hydrogen peroxide (H₂O₂), hydroxyl radical (OH⁻), and singlet oxygen (¹O₂), may be produced under low temperatures due to chilling-induced photoinhibition (Havaux and Davauv, 1994; Terashima et al., 1994). Plants have also evolved mechanisms to protect cellular membranes and organelles from the damaging effects of AOS (Foyer et al., 1991). Antioxidative enzymes and antioxidant molecules can neutralize AOS (Foyer et al., 1991; Lee and Lee, 2000; Oidaira et al., 2000; Scandalios, 1993; Wise and Naylor, 1987).

Cold acclimation increases tolerance to AOS in cereals and correlates with an increase in antioxidative enzymes (Anderson et al., 1995; Scebba et al., 1999). In chilling-sensitive plants, the ability to defend against oxidative damage is directly correlated with the level of antioxidants such as ascorbate, glutathione, and α-tocopherol (Wise and Naylor, 1987), and the activities of antioxidative catalase (CAT; Upadhyaya et al., 1989), superoxide dismutase (SOD) (Michalski and Kamiu, 1982), and glutathione reductase (GR) (Foyer et al., 1991). Thus, it is important to determine the activities of various antioxidative enzymes during acclimation and chilling to assess their contribution to chilling tolerance.

The objectives of this study were to compare the physiological responses of cold-acclimated and nonacclimated cucumber plants to chilling and the ability of cucumber plants to recover from chilling injury. Leaf wilting, membrane damage, and photosynthetic efficiency were measured to estimate chilling injury. Furthermore, activities of antioxidative enzymes and their isozymes were compared in cold-acclimated and nonacclimated cucumber plants to examine if the enzymes are associated with chilling tolerance.

PLANT GROWTH AND TREATMENT CONDITIONS. ‘Naeso Sam-chuk’ cucumber seeds were obtained from Seminis Korea Co., Ltd. (Seoul, Korea). Seeds were sown in plug trays (50 × 30 × 7 cm) containing commercial soil substrate (Boo-Nong Soil, Seoul, Korea) and allowed to germinate in the greenhouse at 30 ± 3°C/20 ± 3°C day/night temperature. Germinated seedlings

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were transplanted to a 500-mL pot containing commercial soil substrate, and placed in the growth chamber until plants reached three-leaf stage. The growth chamber was maintained at 30 °C/25 °C day/night temperature, 14-h photoperiod, and 70% relative humidity. Light period started at 0600 hr. The photosynthetic photon flux (PPF) on the plant canopy was 250 μmol·m−2·s−1. The seedlings were watered daily and supplied with half-strength Hoagland’s nutrient solution 3 d before treatment.

At the three-leaf stage the plants were acclimated at 15 or 25 °C for 3 d under a 14-h photoperiod with an irradiance of 50 μmol·m−2·s−1. The acclimated plants were exposed to chilling at 8 °C for 3 d. After the chilling treatment, plants were allowed to recover for 3 d under a 14-h photoperiod with 50 μmol·m−2·s−1 PPF at 25 °C. Measurements of physiological variables were taken from the second leaf from the bottom of each plant, 7 h after the onset of light period, from the start of low temperature acclimation to the third day of recovery. Acclimation and stress treatments were also imposed 7 h after onset of light period. Harvesting was done at the same time each day to avoid complications from diurnal fluctuations in biochemical processes. Since chilling injury in cucumber plants usually occurs in the dawn or in the early morning, the plants were subjected to low light condition throughout the experiment. The experiments were at least triplicated each with six plants.

**Evaluation of Chilling Injury.** Chilling injury on leaves was evaluated using several variables. Since typical visual symptom of chilling injury is wilting of leaves (Salveit and Morris, 1990), leaf water content was examined during acclimation, subsequent chilling and then recovery. The leaf water content was simply calculated by the formula [(1 – (leaf dry weight/leaf fresh weight))] × 100.

Chilling injury was also evaluated by measuring cellular leakage. This was accomplished by cutting 4-mm leaf disks (0.2 g fresh weight) with a cork borer, then placing them in a 6-cm-diameter polystyrene petri dish containing 5 mL of 1% sucrose water as substrate, and placed in the growth chamber until plants reached the three-leaf stage. The growth chamber was maintained at 30 °C/25 °C day/night temperature, 14-h photoperiod, and 70% relative humidity. Light period started at 0600 hr. The photosynthetic photon flux (PPF) on the plant canopy was 250 μmol·m−2·s−1. The seedlings were watered daily and supplied with half-strength Hoagland’s nutrient solution 3 d before treatment.

**Lipid Peroxidation.** Lipid peroxidation was estimated by quantifying the amount of malondialdehyde (MDA) production using a slight modification of the thiobarbituric acid (TBA) method as previously described (Buege and Aust, 1978). The second true leaf from each plant (0.5 g) was harvested for each treatment period, and the tissues were homogenized using a mortar and pestle in 5 mL of 0.5% TBA in 20% trichloroacetic acid. The homogenate was centrifuged at 20,000 g for 15 min, the resulting supernatant was collected. The supernatant was heated in a boiling water bath for 15 min and allowed to cool in an ice bath. Following additional centrifugation at 20,000 g, for 15 min, the resulting supernatant was used for spectrophotometric determination of MDA. Absorbance at 532 nm for each sample was recorded and corrected for nonspecific turbidity at 600 nm. MDA concentrations were calculated using a molar extinction coefficient of 156 mM−1·cm−1.

**Chlorophyll (Chl) a Fluorescence and Quantum Yield Measurements.** Chl a fluorescence was measured in vivo at room temperature using a pulse amplitude modulation fluorometer (PAM-2000, Walz, Effeltrich, Germany). Before measuring fluorescence, leaves were adapted in darkness for 5 min to minimize fluorescence quenching associated with thylakoid membrane electron excitation (Krause et al., 1983). Minimal fluorescence yield, F0, was obtained upon excitation of the leaves with a weak measuring beam of 0.12 μmol·m−2·s−1 from a pulse light-emitting diode. Maximal fluorescence yield, Fm, was determined after exposure to a saturating pulse of white light to reduce all reaction centers. The ratio of variable to maximum fluorescence (Fv/Fm) derived from the measurement was used as an estimate of the maximum photochemical efficiency of photosystem (PS) II (Butler, 1978). The quantum yield (Y) of electron transport through PS II (Y = Δ/Fm) was calculated according to Genty et al. (1989).

**Protein Extraction.** Frozen leaves (0.5 g) were pulverized in liquid N, using a mortar and pestle, and then resuspended in 3 mL of 100 mM potassium phosphate buffer (pH 7.5) containing 2 mM ethylenediaminetetraacetic acid, 1% polyvinylpyrrolidone–40, and 1 mM phenylmethylsulfonyl fluoride. For ascorbate peroxidase (APX) assay, the extraction buffer also contained 5 mM ascorbate. The suspension was centrifuged at 15,000 g for 20 min at 4 °C and the resulting supernatant was used directly as an enzyme source. For SOD assay, however, the supernatant was eluted through a Sephadex G-25 m minicolumn (PD-10; Pharmacia, Uppsala, Sweden) at 4 °C using 100 mM potassium phosphate buffer (pH 7.5) to remove low molecular weight inhibitors (Anderson et al., 1995). Protein concentration was determined according to the method of Bradford (1976) using bovine serum albumin as a standard.

**Enzyme Assays.** All enzymes were assayed in a 1-mL cuvette at 24 to 26 °C. Activity was linear with respect to time and enzyme concentration. Spectrophotometric determinations were carried out on a Shimadzu dual wavelength spectrophotometer model UV-2401 operated in the split beam mode.

SOD activity was determined based on its capacity to inhibit cytochrome c by superoxide generated by xanthine-xanthine oxidase reaction (Sypchalla and Desborough, 1990). One unit of SOD was defined as the amount of enzyme which inhibited the rate of cytochrome c reduction by 50%. CAT activity was assayed according to the method of Mishra et al. (1993) by monitoring the decline in absorbance at 240 nm (ε = 36 μM−1·cm−1) as a result of H2O2 degradation. APX activity was estimated with the method of Chen and Asada (1989) by monitoring the decline in absorbance at 340 nm as NADPH (ε = 6.2 μM−1·cm−1) was oxidized. GR was measured with the method of Rao et al. (1996) by monitoring the decline in absorbance at 340 nm as NADPH (ε = 6.2 μM−1·cm−1) was oxidized.

**Native Polyacrylamide Gel Electrophoresis and Activity Staining.** Isozymes of CAT, SOD, APX, and GR were separated on nondenaturing polyacrylamide gels according to Laemmli (1970) with slight modifications. Equal amounts of protein extracts, mixed with bromophenol blue and glycerol to provide a marker for each gel, were electrophoresed at 4 °C for 3 h with a constant current of 30 mA. For APX, however, 2 mM ascorbate was added to the electrode buffer (pH 8.3) and the gel was prerun for 30 min before the sample was loaded. MDA concentrations were calculated using a molar extinction coefficient of 156 mM−1·cm−1.

**Chlorophyll (Chl) a Fluorescence and Quantum Yield Measurements.** Chl a fluorescence was measured in vivo at room temperature using a pulse amplitude modulation fluorometer (PAM-2000, Walz, Effeltrich, Germany). Before measuring fluorescence, leaves were adapted in darkness for 5 min to minimize fluorescence quenching associated with thylakoid membrane electron excitation (Krause et al., 1983). Minimal fluorescence yield, F0, was obtained upon excitation of the leaves with a weak measuring beam of 0.12 μmol·m−2·s−1 from a pulse light-emitting diode. Maximal fluorescence yield, Fm, was determined after exposure to a saturating pulse of white light to reduce all reaction centers. The ratio of variable to maximum fluorescence (Fv/Fm) derived from the measurement was used as an estimate of the maximum photochemical efficiency of photosystem (PS) II (Butler, 1978). The quantum yield (Y) of electron transport through PS II (Y = ΔFm) was calculated according to Genty et al. (1989).
CAT activity was detected using the procedure of Anderson et al. (1995). APX and GR isozymes were visualized by the methods described by Rao et al. (1996).

Each staining reaction was stopped with 7.5% glacial acetic acid. The gels were then stored at 4 °C in plastic boxes containing 7.5% acetic acid solution until photographic recording.

**Results**

**Physiological responses to cold-acclimation and chilling.** The leaf water contents in all plants were equivalent during 3 d of acclimation in cold-acclimated and nonacclimated leaves (Fig. 1A). However, subsequent chilling at 8 °C reduced leaf water contents progressively in nonacclimated leaves with increasing duration of chilling. Furthermore, water contents in these leaves were not restored completely after 3 d of recovery at 25 °C. In contrast, cold-acclimated leaves did not wilt during the chilling process (Fig. 1A). Although in cold-acclimated leaves, leaf water contents decreased 3 d after chilling, the leaves recovered completely 1 d after terminating the chilling treatment.

There was no difference in cellular leakage between these leaves during the acclimation period (Fig. 1B). However, cellular leakage was consistently lower in cold-acclimated leaves than in nonacclimated leaves during the chilling period. After 3 d of chilling, cellular leakage from nonacclimated leaves was about four times higher than that of cold-acclimated leaves. Cellular leakage from chilled leaves declined when the plants were allowed to recover at 25 °C (Fig. 1B). After 3 d in the recovery environment, cellular leakage from cold-acclimated leaves returned to the baseline levels observed during acclimation. Nonacclimated leaves, however, exhibited a higher level of cellular leakage after the recovery period than during acclimation.

**Lipid peroxidation.** The level of MDA production was determined to estimate lipid peroxidation during acclimation, and subsequent chilling and recovery. Levels of MDA indicated that lipid peroxidation did not occur during acclimation in cold-acclimated and nonacclimated leaves (Fig. 1C). In nonacclimated leaves, however, lipid peroxidation occurred during chilling and increased with increasing chilling duration. The level of lipid peroxidation also increased during the first day of recovery and then declined thereafter (Fig. 1C). However, little or no lipid peroxidation occurred in cold-acclimated leaves during the chilling and the subsequent recovery.

**Chl a fluorescence and quantum yield.** During acclimation,
tion, the Chl a fluorescence (Fv/Fm) values in cold-acclimated and nonacclimated leaves were only similar (Fig. 2A). However, subsequent exposure to chilling temperature rapidly reduced Chl a fluorescence in nonacclimated leaves, whereas the Chl a fluorescence in cold-acclimated leaves was slightly affected. The Chl a fluorescence of cold-acclimated and nonacclimated leaves was almost fully restored after 3 d of recovery.

Similar to the changes observed in Chl a fluorescence, quantum yields in nonacclimated leaves decreased rapidly during chilling and was mostly restored after the recovery period (Fig. 2B). In contrast, quantum yields of cold-acclimated leaves were not affected by chilling temperatures.

**Antioxidative Enzyme Activity.** Changes in antioxidative enzyme activities of SOD, CAT, APX, and GR activity in cold-acclimated and nonacclimated leaves during chilling and subsequent recovery are summarized in Fig. 3. The level of SOD activity was initially similar in all plants. However, 3 d after cold-acclimation and 2 d after chilling, SOD activity was higher in cold-acclimated leaves than in nonacclimated leaves (Fig. 3A). During recovery, SOD activity increased in both treatments, but the level of SOD activity in cold-acclimated leaves was higher than in nonacclimated leaves. CAT activity was higher in cold-acclimated leaves than in nonacclimated leaves during the acclimation period (Fig. 3B). CAT activity was higher in cold-acclimated leaves than in nonacclimated leaves during chilling and recovery. The effect of acclimation on CAT activity in cucumber leaves was most prominent during the period of recovery.

Similar to CAT, changes in APX activity were observed between cold-acclimated and nonacclimated leaves during acclimation and subsequent chilling (Fig. 3C). APX activity in cold-acclimated leaves was higher than in nonacclimated leaves toward the later stages of acclimation and this continued during chilling. However, there was no difference in APX activity between cold-acclimated and nonacclimated leaves after recovery. In general, no difference was observed in GR activity between cold-acclimated and nonacclimated leaves after chilling and at the end of the recovery period (Fig. 3D).

**Antioxidative Isozyme Profiles.** To relate changes in antioxidative enzyme patterns to chilling injury or tolerance in cold-acclimated and nonacclimated leaves, we analyzed SOD, CAT, APX, and GR isozymes on native polyacrylamide gels (Fig. 4). Two SOD isozymes were detected in cucumber leaves during acclimation and subsequent chilling regardless of acclimation temperature (Fig. 4A). Two additional SOD isozymes were detected in all leaves during recovery. All four isozymes were identified as Mn-SOD type because of their insensitivity to both KCN and H$_2$O$_2$ (Britton et al., 1978). Other SOD isozymes belonging to the Cu/Zn-SOD family...

Fig. 3. Changes in SOD (A), CAT (B), APX (C), and GR (D) activities in cold-acclimated and nonacclimated cucumber leaves during acclimation, chilling, and recovery. The plants were exposed for 3 d at 15°C (cold-acclimated) or 25°C (nonacclimated), chilled for 3 d at 8°C, and allowed to recover at 25°C. Values are the mean ± se of three replications. In some cases, the error bar is obscured by the symbol.

Fig. 4. Changes in SOD (A), CAT (B), APX (C), and GR (D) isozyme profiles cold-acclimated (lower panel) and nonacclimated (upper panel) cucumber leaves during acclimation, chilling, and recovery. The plants were exposed for 3 d at 15°C (cold-acclimated) or 25°C (nonacclimated), chilled for 3 d at 8°C, and allowed to recover at 25°C.
were also present, as indicated by their sensitivity to KCN and H$_2$O$_2$
(data not shown). Fe-SOD was not observed in any treatment. There
was no significant difference in the level of the isozyme activity
or their expression pattern in cold-acclimated and nonacclimated
leaves during acclimation, chilling, and recovery.

Two isozymes of CAT were visualized and distinctive differences
in isozyme expression pattern and activity were observed between
cold-acclimated and nonacclimated leaves (Fig. 4B). The band inten-
sity of CAT isozymes was higher in cold-acclimated leaves than in
nonacclimated leaves during acclimation, subsequent chilling,
and recovery. The CAT isozyme 1 (upper band) appeared one day
after acclimation, 2 d after chilling, and two to 3 d into the recovery
period in cold-acclimated leaves. In contrast, CAT isozyme 1 ap-
peared only once (2 d after chilling) in nonacclimated leaves.

Three isozymes of APX were observed (Fig. 4C). APX isozyme
2 (middle band) was faint, but APX isozymes 1 and 3 were
prominent. Isozymes 1 and 2 appeared only during the recovery
period. The band intensity of APX isozymes was distinctly higher
in cold-acclimated leaves than in nonacclimated leaves two and 3
d after acclimation and during chilling.

Three GR isozymes were observed in cucumber leaves in both
acclimation temperatures (Fig. 4D), but GR isozyme 1 (upper
band) was faint. The band intensity of GR isozyme 2 was higher
in nonacclimated leaves than in cold-acclimated leaves during
acclimation. In contrast, the band intensity of GR isozyme 3 was
higher in cold-acclimated leaves than in nonacclimated leaves
during chilling and 1 to 2 d after recovery.

**Discussion**

To understand the mechanisms of plant species responses to
chilling, previous research had compared metabolic differences
between chilling-susceptible and chilling-tolerant cultivars as model
systems (Jahnke et al., 1991; Walker and McKersie, 1993). However,
this model system was confounded by genetic differences between
sensitive and tolerant species. To properly identify physiological
responses to environmental stress such as chilling, the use of one
cultivar that is chilling-sensitive, but can be acclimated, would be
more appropriate.

Chilling initially damages the photosynthetic apparatus as a
function of reduced CO$_2$, fixation and altered fluorescence patterns
of chlorophyll a (Walker et al., 1991). Other effects, including loss
of water and chlorophyll contents, occur more gradually as leaf
conductivity changes or leaves turn yellow (Koscieniak, 1993).
Visual symptoms of chilling damage include waterlogged appear-
ance, wilting, acceleration of senescence, and inhibition of seedling
growth (Saltveit and Morris, 1990).

Cucumber plants exposed to chilling temperature exhibited all
of these symptoms. However, cold-acclimated plants in this study
were more tolerant to chilling stress than nonacclimated plants.
This conclusion was based upon the variables measured including
leaf water content, cellular leakage, lipid peroxidation, $F_v/F_m$ ratio,
and quantum yield (Figs. 1 and 2). Cold-acclimated plants also
recovered faster than nonacclimated plants. These results suggest
that in cold-acclimated plants oxidative damage is reduced during
chilling and, therefore, the plant recovers from injury. A similar
acclimation phenomenon has been demonstrated in other low
temperature-sensitive species, such as maize and tomato (Leipner
et al., 1997; Venema et al., 2000).

It is known that chilling-induced AOS triggers a series of deleteri-
ous processes, such as lipid peroxidation and degradation of proteins
and DNA damage in the cell (Halliwell and Gutteridge, 1986).

Lipid peroxidation, as a measure of cellular injury, was 2-fold
higher in nonacclimated plants compared to cold-acclimated plants
during chilling and recovery (Fig. 1C). This confirms that chilling
induced oxidative stress in cucumber plants. In maize seedlings,
for example, there was a 2-fold increase in lipid peroxidation in
nonacclimated plants compared to nonstressed and cold-acclimated
seedlings, during chilling (Prasad, 1996).

High SOD activity has been associated with stress tolerance in
plants where overproduction of superoxide is involved (Bowler et
al., 1992). SOD activity in cold acclimated spinach and wheat plants
was higher compared to nonacclimated plants during exposure to
low temperature (Scebbba et al., 1999; Schöner and Krause, 1990).
Induction of SOD activity by cold stress may not occur in all species.
For instance, SOD activity in wheat subjected to chilling stress was
described to significantly modify by cold acclimation (Scebbba et al., 1998).
In our study, SOD activity was higher in cold-acclimated leaves than in
nonacclimated leaves 3 d after acclimation, 2 d after chilling, and
one and 3 d after recovery (Fig. 3A). Various SOD isozymes are
expressed in different cells, tissues, or organelles and activities of
the isozymes may be affected differently by chilling (Scandalios,
1993). Up to five SOD isozymes were observed during the course
of this experiment on nondenaturing polyacrylamide gels, but none
of them were significantly affected by cold acclimation (Fig. 4A).
Similar to our results, new SOD isozymes were not detected in
cold-acclimated wheat seedlings (Scebbba et al., 1998). However,
new SOD isozymes were induced in spinach by cold acclimation
(Schöner and Krause, 1990).

The higher CAT activity (Fig. 3A) and band intensity (Fig.
4A) in leaves of cold-acclimated plants suggest a more efficient
scavenging of H$_2$O$_2$, which resulted in better protection against
peroxidation. Similarly, CAT activity in cold-acclimated material has
been reported to remain similar to (O’Kane et al., 1996) or higher
than (Anderson et al., 1995) that of nonstressed plants. In many
cases, activity of CAT has been observed to decline in response to
sudden exposure to chilling temperature without prior acclimation
(Feierabend et al., 1992; Mishra et al., 1993). This was also true in
chilled cucumber seedlings (Lee and Lee, 2000).

The important role of APX in relation to increasing oxidative
tolerance has been observed in many plants (Feierabend et al., 1992;
Gupta et al., 1993). In our study, APX activity was significantly
affected by acclimation temperature and chilling treatment (Fig. 3C).
The high band intensity of APX isozymes from cold-accli-
mated plants supported the fact that APX activity was higher in
cold-acclimated plants than in nonacclimated plants (Fig. 4C); i.e.,
the enzyme profile was similar, but expression level was elevated in
cold-acclimated plants.

GR has been implicated as an important protection agent against
oxidative damage in many plants (Foyer et al., 1991). In our study,
however, there was no difference in total GR activity between cold-
acclimated or nonacclimated cucumber plants at any time during
the duration of the experiment, except for 3 d after chilling (Fig.
3D). Exposure to low temperature resulted in altered GR isozyme
profiles (Fig. 4D). GR isozyme activity equalized the total GR activity
in cucumber with or without cold acclimation (Fig. 4D).

Our results indicate that chilling-sensitive cucumber plants can
be made cold-tolerant by cold acclimation which is paralleled by
induced activities of antioxidative enzymes CAT and APX. How-
ever, the mechanism of chilling stress protection is complex and
may act in concert with other physiological mechanisms which
improve cellular membrane integrity, alter the composition of cel-
lular fluids, or maintain the capability for physiological function
under chilling temperature.

Literature Cited


