Growth and Physiological Adaptations of Tomato Plants 
(*Lycopersicon esculentum* Mill) in Response to Water Scarcity in Soil

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Abstract. This study aim to investigate fundamentally the growth and physiological responses of tomato plants in responses to two different levels of water deficit, a weak drought stress (−25 kPa) and a severe drought stress (−100 kPa) in soil. The two levels of water deficit were maintained using a micro-irrigation system consisted of soil sensors for the real-time monitoring of soil water content and irrigation modules in a greenhouse experiment. Soil water contents were fluctuated throughout the 30 days treatment period but differed between the two treatments with the average −47 kPa in −25 kPa set treatment and the −119 kPa in −100 kPa set treatment. There were significant differences in plant height between the two different soil water statuses in plant height without differences of the number of nodes. The plants grown in the severe water-deficit treatment had greater accumulation of biomass than the plants in the weak water-deficit treatment. The severe water-deficit treatment (−119 kPa) also induced greater leaf area and leaf dry weight of the plants than the weak water-deficit treatment did, even though there was no difference in leaf area per unit dry weight. These results of growth parameters tested in this study indicate that the severe drought could cause an adaptation of tomato plants to the drought stress with the enhancement of biomass and leaf expansion without changes of leaf thickness. Greater relative water content of leaves and lower osmotic potential of sap expressed from turgid leaves were recorded in the severe water deficit treatment than in the weak water deficit treatment. This finding also postulated physiological adaptation to be better water status under drought stress. The drought imposition affected significantly on photosynthesis, water use efficiency and stomatal conductance of tomato plants. The severe water-deficit treatment increased PSII activities and water use efficiency, but decreased stomatal conductance than the weak water-deficit treatment. However, there were no differences between the two treatments in total photosynthetic capacity. Finally, there were no differences in the number and biomass of fruits. These results suggested that tomato plants have an ability to make adaptation to water deficit conditions through changes in leaf morphology, osmotic potentials, and water use efficiency as well as PSII activity. These adaptation responses should be considered in the screening of drought tolerance of tomato plants.

Key words: drought adaptation, osmotic potential, tomato, water deficit

Introduction

Tomato, *Lycopersicon esculentum* Mill., is one of the most important vegetable crops in the world in terms of consumption per capita. Fruits of tomato are not only eaten directly as a fresh vegetable but also additive ingredient to other food items. They contribute significantly to the dietary intake of vitamins A and C, essential minerals and other nutrients. Furthermore, tomatoes are the richest source of lycopene, a photochemical that protects cells from oxidants causing cancer (Giovannucci, 1999).

It is widely distributed as an annual vegetable crop adapted to diverse climates. However, in spite of its broad adaptation, the production is concentrated in a few warm and rather dry areas particularly in tropical and sub-tropical climates (Cuartero and Fernandez-Munoz, 1999). In the tropical and subtropical regions, environmental extremities such as drought and salinity are major factors contributing and limiting the plant growth and productivity due to disturbing the intercellular water balance (Chaves et al., 2002).
Drought, a substantial water deficit in soil, is an increasingly major constraint to crop productivity and yield stability worldwide. Modern agriculture is affected by several abiotic factors such as drought, extreme temperature, light, and salt stress. Among abiotic factors, drought is a leading stress causing significant lost in crop productivity. Drought affects the quality, quantity, yield and growth related characters of the plant. However, it is presumed that yield-related characters are associated with almost every aspect of the physiology and biochemistry of the plant. Furthermore, plants responses to water deficit are associated with various physiological changes (Bray, 1997).

Usually, plants synthesize and accumulate osmolytes as a physiological response to counter balance the abiotic stresses. In Arabidopsis, the drought signal is mediated through abscisic acid (ABA)-dependent and -independent pathways which is often regulated by the expression of gene under drought. Similarly, during vegetative growth, roots of many angiosperms synthesize ABA and transport to the shoots under prevalent drought conditions which have been reported in various crop plants such as rice, wheat, alfalfa etc. (Leung and Giraudat, 1998; Shinozaki and Yamaguchi-Shinozaki, 1997, 2000). However, physiological responses of drought tolerant plants have not been critically considered so far.

Besides, there is a range of mechanisms of adaptation to drought stress in higher plant mentioned by some workers. For example, drought-induced osmotic adjustment is important to maintain low water potential and to maintain cellular turgor pressure. Likewise, drought-induced reduction of stomatal conductance and transpiration are critical to save internal water under the water deficit condition (Costa Franca et al., 2000). However, studies related to mechanism of adaptation to drought stress in tomato plant are rather scarce.

Specially, the physiological response of tomato to drought stress has not been extensively studied particularly throughout vegetative and reproductive periods. Presumably, each stage may be considered as separate traits and may require different evaluation methods. Further it is known that no reliable physiological evaluation procedure can be employed successfully and effectively to examine the drought tolerance in each phenophasis. So the present study describes the growth and physiological responses of tomato plants against drought stress in two phenophasis i.e. vegetative and reproductive phases. Physiological analysis of tomato plants under drought stress throughout the vegetative and reproductive phase may be useful to assess the drought tolerance of other crops using micro-irrigation system. This study aims to fundamentally investigate the growth and physiological responses of tomato plants in responses to two levels water deficit, a weak drought stress and a severe drought stress, in soil.

Materials and Methods

1. Plant growth conditions and treatments

Tomato seeds (Lycopersicon esculentum Mill.) ‘Picco’, a popular cherry tomato, were used in this study which had carried out from 31 March to 12 July 2006. Seeds were sterilized with sodium benlate solution (×1/200) for 30 min. After that, the sterilized seeds were washed with distilled water several times before sowing in wet paper. ‘Picco’ raised in horticultural institute lab in 40 days. Seeds were sown and germinated seedling then transplanted into sub-irrigation system after 45 days. Germinated seeds were transferred into pots having size about 21 × 20 cm. Eighteen pots were prepared with two sets of nine pots, weak water-deficit treatment (−25 kPa) and severe water-deficit treatment (−100 kPa). Pots were arranged randomly and filled with perlite. Plants were grown in the greenhouse especially, located National Institute of Agricultural Biotechnology, Rural Development Administration, Suwon, Republic of Korea (altitude 37.2669 N°, latitude 126.9669 E°). Each treatment was replicated nine times with two plants in each pot. Plants were irrigated with half-strength Hoagland solution while drought treatments were maintained as −47 kPa and −119 kPa by micro-irrigation system, respectively. The treatments were started from 12 June to 12 July, 2006.

2. Drought treatment system using micro-irrigation monitoring system

Precise imposition of drought stress can be achieved thought a controlled micro-irrigation system with water sensors and drip irrigation. Micro-irrigation monitoring system set up the two different water statues, weak water-deficit treatment (−25 kPa) as control and severe water-
deficit treatment (−100 kPa) for 30 days. This system revealed drought imposition significantly affected the water status of the soil. As shown as Fig. 1, the drought treatment (−100 kPa) increased the soil water content compared to weak drought (−25 kPa) as control. Throughout the 30 days treatment period, soil water contents were fluctuated but differed among the two treatments with average −47 kPa in −25 kPa-set treatments and −119 kPa in −100 kPa-set treatments (Fig. 1). It indicated that relative humidity caused significant change in drought condition. However, the average temperature and relative humidity were maintained as 24.3°C and 64.3%, respectively.

3. Sampling and physiological analysis

Plants were harvested on 12 July after 30 days treatment, and height, nodes, fruit rate, leaf area, dry weights, photosynthesis, relative water contents (RWC) and osmotic potential were measured. Total 18 plants were used per each treatment. Whole sample, leaf and fruit were dried at 65°C for 72 hours.

Leaf area was determined using a LI-3100 area meters (Delta Area Meter MK2, Delta-T Devices Ltd, England). Relative water content (RWC) was calculated from gravimetric measurements using the following formula (Catsky, 1960):

\[
\text{RWC} (%) = \frac{\text{Fresh Weight} - \text{Dry Weight}}{\text{Turgid Weight} - \text{Dry Weight}} \times 100.
\]

Randomly selected leaves per plant were weighed immediately after cutting off (fresh weight), rehydrated in distilled water at 95% humidity (turgid weight) for 4 hours and then oven dried at 65°C for 72 hr (dry weight). For the determination of osmotic potential, leaf samples were frozen in liquid nitrogen and kept at −20°C. Systemically, the expressed sap solution was prepared with the homogenates at 12000 g for 20 min at 4°C. Osmotic potential of a 10 μL of the expressed sap was determined using a vapor pressure osmometer (model 5520; Wescor, Inc, USA) and then calculated based on the van’t Hoff relation reported by Nobel (1983). For turgid osmotic potential, the leaf samples were kept in distilled water for 4 hours before frozen and homogenate. Photosynthesis was measured in terms of net assimilation.
rate of CO₂ and quantum yield in both severe water-deficit treatment (−100 kPa) and optimal irrigation treatment (−25 kPa) plants using Y-portable photosynthesis system (Li-6400, Li-Cor Co., Inc., Lincoln, NE, USA). LI-6400 computes stomatal conductance and transpiration rate by measuring air flow rate and chamber H₂O concentration, leaf area, and mole fraction of water in chamber and in stomata. Water use efficiency was measured as net CO₂ assimilation rate, transpiration rate, and photosynthetic parameters measured after light adaptation for 15 min at 600 µmol m⁻² s⁻¹ PPF on 13 July, 2006 on the next day after harvesting.

4. Data analysis

Statistical analysis was performed with SAS software (SAS, Statistical Institute, Cary, NC, USA). Significant differences from two treatment means were determined \( P \leq 0.05 \). Least Significant Difference (L.S.D.) was used to identify dissimilar treatment means.

Results and Discussion

1. Effects of drought stress on agronomic parameters

We observed several growth parameters of tomato growing under two levels of water status, −25 kPa and −100 kPa, each. Drought imposition (−100 kPa) showed significant reduction in height but did not show significant difference of nodes between two treatments, −25 kPa and −100 kPa (Fig. 3). That means severe drought stress (−100 kPa) could be affected on inter-nodal length. Interestingly, the plants in the −100 kPa treatment showed higher dry weight than those of −25 kPa treatment with significant reduction of total dry weight and leaf dry weight (Fig. 4B and 5C). In addition, non-significant difference was recorded in shoot/root ratio of *L. esculentum* (data not shown). Fruit sets in both −25 kPa and −100 kPa condition having similar morphology (Fig. 2), however, few fruits were reduced in size at drought stress. There is no significant difference of fruit dry weight between −100 kPa and −25 kPa (Fig. 5). Although the deficit irrigation resulted in substantial yield loss compared to the standard practice, marketable tomato fruits was increased by 27% at −30 kPa treatment compared to the −15 kPa treatment (Nam et al., 2006).

The experiment season was June to July when it is usually very hot and humid. The high humidity during this experiment might affect stomata opening which cause water absorption, leading to large leaf area and dry weight. However, we observed that leaf thickness and leaf dry weight were not affected during the experimental periods (Fig. 4). Non-significant difference was recorded in leaf area per unit dry weights, rate of fruit set, and fruit
dry weight (Fig. 4 and 5). That means tomato ‘Picco’ has ability to keep its growth without any significant yield loss under drought stress.

We conclude that drought imposition did not inhibition of growth inducing tomato fruit setting throughout −25 to −100 kPa during this experiment period using micro-irrigation system. In tomato, most commercial cultivars are sensitive to drought stress throughout the ontogeny of the plant, yet genotypic variation for drought tolerance exists within the cultivar (Wudiri and Henderson, 1985). The cultivar experiments indicate the potential for factors such as maturity, yield potential, and adaptation to generate spurious correlations. It has been also observed that within a set of cultivar having more similar yield potential and adaptation, grain yield in a dry season environment was not closely associated with leaf RWC (Lafitte, 2002). For instance, yield variation that is not related to plant water status may reflect variation in mechanisms such as the tolerance to root or leaf-source signals that reduce grain set under stress, or difference in the response of grain formation to varying levels of carbohydrate mobilization near flowering to fruiting. Moreover, high leaf water status levels allow the plant to grow well, high water status under stressed conditions is not necessarily advantageous for biomass production at least at the young stage (Daniele et al., 2005).

Thus, it is assumed that these responses associated with plant growth stages and cultivars might be the adaptation mechanism of ‘Picco’ tomato plant. There is a range of mechanisms for adaptation to drought stress in higher plants which are maintenance of turgor, low osmotic potential, better osmotic adjustment value, high relative water contents, better water use efficiency, stomatal regulation, photosynthetic performance, etc. (Anyia and Herzog, 2004; Bohnert et al., 1995).

2. Effects of drought stress on physiological parameters

Based on the results of this study, it is presumed that tomato shows its adaptability under −100 kPa drought treatments. This study also goes on to demonstrate appropriate physiological responses such as RWC, osmotic potential, photosynthesis, CO₂ assimilation rate under drought stress. We investigated whether drought imposition affects those physiological parameters inducing the growth difference. The result showed drought imposition significantly affects on the relative water content of L. esculentum. The maximum relative water content was recorded in −100 kPa treated sample compared to −25 kPa water deficit. Similarly, osmotic potential was almost the same in both −25 and −100 kPa treated samples (Table 1). In addition, −100 kPa treatment showed a little higher in turgid osmotic potential than those of −25 kPa. Water use efficiency was significantly increased due to drought compared to −25 kPa unstressed condition.

Maintenance of leaf relative water content in drought resistant plant could be an important strategy under drought stress. Some studies have demonstrated that
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Desert Caragana species have secondary xylem, which is typical in xerophytes and young stems and leaves have a xeromorphic structure (Chang and Zhang, 1997) showing lower transpiration rate and higher water use efficiency than other species. Undoubtedly, in plants that have high water potential, reduction of stomatal conductance and transpiration rate are critical to save water (Costa Franca *et al.*, 2000). Its photosynthesis system is adaptable to drought, high temperature and strong solar radiation (Ma *et al.*, 2004). Furthermore, drought-tolerant plants have a stronger ability to adjust stomatal conductance, enabling them to maintain water balance in an arid environment (Vacher, 1998).

In the present study, −100 kPa increased the net CO₂ assimilation rate and photo system II (PSII) values of tomato compared to stomatal conductance where significant reduction was recorded (Table 2). In particular, PS II, the main reaction centre of the photochemical efficiency of photosynthesis, which is located on cell membranes, is very sensitive to water stress (Glynn and Colin, 2002). It is presumed that physiological responses such as RWC, osmotic potential, stomatal conductance, photosynthetic rate, water use efficiency are all associated responses under drought environments especially for the cultivar which are better adopted as shown in the present study (Table 2).

Many experiments have shown that stomata close in response to hydraulic and chemical signals produced when soil moisture decreased (Chaves *et al.*, 2002; Davies *et al.*, 2000). The soil water content and the root water status directly affect stomatal conductance, which diminishes considerably before any observable change in leaf water potential occurs. Stomatal conductance generally decreases as the soil water content falls below an adequate level to sustain normal plant water uptake. In dry soils, stomata opens and closes very rap-

**Table 1.** Effect of drought imposition on the relative water content, osmotic potential and turgid osmotic potential of c.v. ‘Picco’ *L. esculentum*.

<table>
<thead>
<tr>
<th>Drought treatment</th>
<th>Relative water content (%)</th>
<th>Osmotic potential (ψₛ)</th>
<th>Turgid osmotic potential (ψₛ₉₉)</th>
</tr>
</thead>
<tbody>
<tr>
<td>−25 kPa</td>
<td>89.51*</td>
<td>719.03</td>
<td>637.65</td>
</tr>
<tr>
<td>−100 kPa</td>
<td>90.92</td>
<td>716.63</td>
<td>668.44</td>
</tr>
<tr>
<td>LSD* 0.05</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
</tr>
</tbody>
</table>

* Least significant difference at P = 0.05.
* NS* Non significant or significant difference between −25 kPa and −100 kPa treatments (P = 0.05).
* Values are mean of 18 plants, each.

![Fig. 5.](image)

*Fig. 5.** Effect of drought imposition on the fruit set, fruit dry weight, total dry weight of *L. esculentum* on 30 days after micro-irrigation. Vertical lines on the bar graph represents standard error (n = 18).
idly after the beginning of an irrigation event in response to changes in soil water conditions (Zavala, 2004). Since stomata influence the influx of CO$_2$ into leaves, the reduction in stomatal conductance to conserve water inevitably indicates a lower photosynthetic rate since stomata influence the influx of CO$_2$ into leaves. However, -100 kPa treatment showed the reduction in stomatal conductance but did not induce the less photosynthetic rate (Table 2). This is probably due to the high degree of co-regulation of stomatal conductance and photosynthesis in ‘Picco’.

Moreover, all physiological attributes indicate the usefulness of this study on adaptability of tomato to drought. It can be thought that the cultivar optimizes its biomass production before plant growth due to its higher water use efficiency, so as to mobilize stored assimilates when the stomata are closed. Among varieties, a high relative transpiration level, generally associated with a low water potential, is often observed in the most productive plant varieties under drought. In addition, cell membrane tolerance has proven to be an important trait regardless of the type of water stress (Bajji et al., 2002), and could be associated with other favorable attributes for drought adaptation in plant (Daniele et al., 2005).

We explored the growth, yield and their subsequent physiological response of tomato in drought stress. The present study found drought has less significant effects on growth, yield and physiological responses of tomato ‘Picco’ that has various capacity of adoptability. Furthermore, it is expected to contribute for evaluation of drought tolerance in tomato genotype or other crops. Earlier, species naturally adapted to drought, such as bean or groundnut, has improved their agronomic performance (CostaFranca et al., 2000). This study is a first try to establish the adopted mechanism of vegetable crops as tomato using unique micro-irrigation system. However, further molecular studies related to drought tolerance mechanism of ‘Picco’ tomato plant are needed.

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**Literature Cited**


마토 'Picco'가 엽형태의 변형과 삼투압, 수분활용도와 PSII의 활성을 통해 수분결핍상태에서 적응할 수 있게 만드 능력을 보여준다. 본 연구결과에서 나타난 토마토의 수분스트레스 적응 메커니즘은 토마토의 가뭄저항성 스크린에 있어서 고려되어야 할 것으로 보인다.

주제어 : 가뭄적응성, 삼투압, 수분결핍, 토마토