HEAT STRESS-INDUCED ALTERATIONS IN ANTIOXIDATIVE ENZYMES 
OF SOME PLANTS OF CUCURBITACEA FAMILY

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Abstract
The effects of high temperatures on melon cvs. Miranda and Poli, watermelon cv. Crimson Tide and zucchini cv. Asma leaves. The leaves obtained from plants were subjected to 35, 40, 45, 50, 55 and 60°C temperatures with gradual increments every 30-minutes. Samples, obtained at each treatment, were analyzed for ascorbic acid content, NADP(H) oxidase, catalase, gluthatione reductase, peroxidases activities and isoperoxidase patterns. The ascorbic acid content slightly increased parallel to temperature increment in zucchini but did not change in watermelon and in both melon cultivars. Melon cv. Poli exhibited comparatively less oxidative damage than cv. Miranda with a lower NAD(P)H oxidase activity. Heat stress induced NAD(P)H activity in watermelon and zucchini comparing to control plants. Results revealed that antioxidative enzyme activities were increased generally up to 50°C then decreased gradually in melon cultivars. Besides cv. Poli generally had higher enzyme activities than cv. Miranda. The activity of catalaes become prominent in watermelon while the activity of ascorbate peroxidase become prominent in zucchini. Acidic isoperoxidase bands with different relative mobility values were found in all species. Besides, basic isoperoxidase band could not be determined in both melon cultivars and watermelon while a basic isoperoxidase band was found in zucchini.

Keywords: Ascorbic acid, cucurbits, enzymatic antioxidants, high temperature stress, isoenzyme

1. INTRODUCTION
Plants face various stress factors throughout their lifespans. Heat stress is one of the major threat to plant survival especially in the arid regions (Jin et al., 2011). Besides, considerably affecting plant growth and yield (Ahmad and Prasad 2012), high temperature is one of the major concern for crop production due to global warming (Hu et al., 2020). Plant responses to abiotic stresses are multigenic therefore it is difficult to identify, control and manipulate (Ben-Ari and Lavi, 2012). Hence, understanding the complexities of heat tolerance mechanisms is a key to manipulate tolerance in plants (Milner, 2020). The plant responses to heat stress vary depending on the plant and even the tissue or organ within the same plant affected (Peer et al., 2020). The devastating effects of heat stress on plant metabolism includes, disrupting cellular homeostasis and uncoupling major physiological and biochemical processes.
A direct consequence of stress-induced cellular changes is the production of excessive reactive oxygen species (ROS), which are produced in plants in such a way that they are confined to a small area and are also produced in a non-specific pattern in biological responses (Awasthi et al., 2015). The ROS (superoxide; O2−, hydroxyl radicals; •OH, and non-radicals such as hydrogen peroxide;
H₂O₂ and singlet oxygen \(^1\text{O}_2\) are highly reactive and toxic and damage proteins, lipids, carbohydrates, resulting in oxidative stress (Rehman et al., 2020). The rate of ROS production and scavenging is critical in determining the oxidative load of plant tissue. Hydrogen peroxide accumulation itself can trigger higher ROS production by disrupting photosynthesis and activities of nicotinamide adenine dinucleotide phosphate [NAD(P)H]-dependent oxidase (Ara et al., 2013; Du et al., 2013). Catalysing the production of \(\text{O}_2^-\) NAD(P)H oxidase (EC 1.6.3.1) takes place in diverse essential processes in plants (Qu et al., 2017).

Plants can cope with ROS by their antioxidant defence system which consists of non-enzymatic and enzymatic antioxidants (Gill and Tuteja, 2010). Ascorbic acid (AsA), reduced glutathione, (GSH) and proline are some non-enzymatic antioxidants. Besides, superoxide dismutase (SOD, EC 1.15.1.1), catalase (CAT, EC 1.11.1.6), glutathione reductase (GR, EC 1.6.4.2), ascorbate peroxidase (APOX, EC 1.11.1.11) are some enzymatic antioxidants (Hasanuzzaman et al., 2020).

Ascorbic acid is synthesized in mitochondria and transported to almost all parts of the plant (Foyer, 2015). Simultaneous effect of AA, α-tocopherol and GSH, results in detoxification of ROS, which limits oxidative stress in plants (Hameed et al., 2012). Ascorbic acid has different effects under abiotic stress conditions and used by APOX as a substrate. (Tripathy and Oelmüller, 2012). The first defence of plants against ROS is that the SOD enzyme converts \(\text{O}_2^-\) radicals to \(\text{H}_2\text{O}_2\). Then \(\text{H}_2\text{O}_2\) is reduced to water and molecular oxygen by CAT, APOX and peroxides, thus preventing further damage to the cell membrane (Hasanuzzaman et al., 2019). Peroxidases (POX, EC 1.11.1.7) is a family of isozymes found in all plants, which uses \(\text{H}_2\text{O}_2\) or \(\text{O}_2\) for the oxidation of various molecules, has a hem-containing monomeric glycoprotein structure, and is classified as acidic, neutral or basic according to their isoelectric points (Rajput et al., 2021). They are involved in many physiological events such as lignin biosynthesis, response to biotic and abiotic stress (Gaspar et al., 1982; Vicuna, 2005).

The antioxidant system is genotype dependent and is tissue specific. Tolerant cultivars have higher levels of antioxidant enzymes under stress conditions (Chaitanya et al., 2002). In addition, leaf is prone to higher ROS production since it is the plant organ where photosynthetic and transpiration processes take place (Jin et al., 2011). For this reason, leaves undergo significant physiological and biochemical alterations that serve as good biomarkers showing the tolerance of plants during stress exposure (Ara et al., 2015). Characterization of the antioxidative defence system in plant cultivars and organs, as increased production of ROS in response to stress is a common phenomenon, can be used as a diagnostic indicator in breeding programs regarding stress tolerance (Naji and Devaraj, 2011; Queirós et al., 2011; Ara et al., 2015).

Being a rich sources of nutrients, vegetables are consumed as daily staple food in all parts of the World and are very sensitive to abiotic stresses (Ara et al., 2013). Considering their growing periods, one of the most important environmental factors affecting the growth and development of cucurbits is high temperature. This study was conducted to investigate the effects of high temperatures on melon cvs. Miranda and Poli, watermelon cv. Crimson Tide and zucchini cv. Asma leaves.

### 2. MATERIALS AND METHODS

The samples were collected from the plants grown in a field under favourable conditions in Eskisehir, Turkey (longitude: 39°45'38"N, latitude: 30°28'47"E). Leaves were collected from cucurbit plants (Cucumis melo L. cvs. Miranda and Poli, Citrullus lanatus L. cv. Crimson Tide and Cucurbita pepo L. cv. Asma in vegetative stage. The average temperatures were measured as 22-24°
°C when the plants were grown. The controlled heat tests were applied to the leaf samples based on the method of Arora et al. (1998). For this purpose, leaves obtained from plants were collected into pyrex tubes with caps closed and placed into water bath. After a 30-minute habituation at 30°C, the water temperature was escalated to 35, 40, 45, 50, 55 and 60 °C temperatures with gradual increments every half an hour. Samples, obtained at each treatment temperature, were immediately fixed in liquid nitrogen and stored at -80 °C until further analysis.

NAD(P)H oxidase was assayed according to Cakmak and Marschner (1988); leaf samples (0.5 g) were homogenized in 5 mL of extraction buffer by using chilled mortar and pestle. The extraction buffers used were 100 mM potassium phosphate (K-phosphate) buffer (pH 7.0). The homogenate was centrifuged at 15 000 g for 20 min at 4°C. The activity of enzyme was estimated by the oxidation of NAD(P)H or NADH using a spectrophotometer (Perkin Elmer Lambda 25 UV/VIS, Santa Clara, CA, USA) at 340 nm.

Ascorbic acid concentrations were determined according to Schoner and Krause (1990). Briefly, 0.5 g fresh weight of leaf samples were homogenised in 5 ml ice-cold 4% (v/v) metaphosphoric acid, and centrifuged at 4 000 g for 10 min. One ml of the supernatant was sampled, mixed with 1 mL 50 mM Na citrate buffer (pH 2.6) and 1 mL 10 mM dichlorophenolindophenol (DCPIP) and incubated at 25°C for 1 min. The optical intensity of the solution was determined at 524 nm and compared with those of standard AsA solutions.

For extraction of antioxidative enzymes, leaf samples (0.5 g) were homogenized in 5 mL of extraction buffer by using chilled mortar and pestle. The extraction buffers used were; 100 mM K-phosphate buffer (pH7.0) for CAT, 50 mM K-phosphate (pH 7.6) for GR, and 50 mM K-phosphate buffer (pH 7.8) for APOX activities. The activity of CAT was estimated according to Rao et al. (1996) by recording the decrease of absorption at 240 nm in as H2O2 (ε= 39.4 mM/cm) was consumed. The GR activity was measured spectrophotometrically at 340 nm (ε=6.22 mM/cm) by the method of Cakmak and Marschner (1992), corresponding the oxidation of β-nicotinamide adenine dinucleotide phosphate (NADPH) and the method of Nakano and Asada (1980) was employed for the assay of APOX by monitoring the decrease in oxidized AsA (ε= 2.8 mM/cm) at 290 nm.

The activities of soluble peroxidase (S-POX) and cell wall bound-peroxidase (CWB-POX) in the leaf tissues were determined spectrophotometrically according to Andrews et al. (2000). Shortly, 1.0 g of leaf tissue was extracted using a mortar and pestle and 5 mL 100 mM K-phosphate buffer (pH 7.0). The extract was centrifuged at 15 000 g for 20 min at 4°C. The supernatant was used to assay for S-POX activity. The CWB pellet was washed four-times with the same buffer, and incubated on ice in 5 mL 1 M NaCl for 5 h. The CWB suspension was then dialysed against distilled water at 4°C for 48 h and centrifuged at 10 000 g for 30 min at 4°C. The supernatant was used for enzyme assays. The POX activity was also measured according to Andrews et al. (2000) and was expressed by reference to a standard curve using horseradish peroxidase (Sigma-Aldrich).

The soluble protein content of the crude enzyme extracts was determined by the Bradford assay method using bovine serum albumin (BSA, Sigma) as standard (Bradford, 1976).

A discontinuous polyacrylamide gel electrophoresis (PAGE) under nondenaturing, nonreducing conditions was performed for the detection of isoperoxidases. The method of Gulen and Eris (2003) was used for the POX extraction from leaf samples. In brief, 0.1 g of ground leaf samples were homogenized with 0.6 mL 0.1M K-phosphate buffer (pH 7.5). The homogenate was centrifuged at 21 000 g for 20 min. at 4 °C and the supernatant was used for POX electrophoresis. Samples were subjected to PAGE using a Mini-PROTEAN tetra cell electrophoresis system (Bio-Rad, Hercules,
CA) as described by Davis (1964) and Reisfeld et al. (1962) for acidic and basic POX, respectively. The method of Wendel and Weeden (1989) was used for staining of the gels. The relative mobility (Rf) value of the bands on the gel was calculated by using Rf= 1.0, as the distance to the finishing point of the running and Rf=0.0, as the starting point of the running (Manganaris and Alston, 1992). The experiment was set up as a randomized block design. All of the assays were repeated three times. The mean values of the data were evaluated using Duncan test at p<0.05 using the SPSS software (version 20., Chicago, IL, USA).

3. RESULTS AND DISCUSSIONS
We found that exposure to high temperatures caused differential responses of NAD(P)H oxidase enzyme activity in cucurbit plants (Figure 1). Heat stress caused a decreased the activity of NAD(P)H oxidase enzyme in both melon cultivars comparing to control treatment. In general, melon cv. Poli exhibited comparatively less oxidative damage than cv. Miranda with a lower NAD(P)H oxidase enzyme activity (Figure 1A, p<0.05). Difference with regard to NAD(P)H oxidase between cultivars was more prominent at 50 °C treatment with 17.74 nmol/mg protein/min and 8.13 nmol/mg protein/min in cv. Miranda and cv. Poli, respectively. The NAD(P)H oxidase enzyme activity of watermelon increased comparing to control plants especially after 50 °C significantly (Figure 1B, p<0.05). The greatest NAD(P)H oxidase enzyme activity at 50 °C, 55 °C, and 60 °C treatment (7.22 nmol/mg protein/min, 7.02 nmol/mg protein/min and 6.78 nmol/mg protein/min, respectively) was observed in watermelon cv Crimson Tide. Statistical analysis revealed a significant effect of high temperature treatments on NAD(P)H oxidase enzyme activity of watermelon. The increment of NAD(P)H oxidase enzyme activity with elevating temperature was remarkable at 60 °C in zucchini plants with 9.75 nmol/mg protein/min enzyme activity (Figure 1C, p<0.05).

There are many studies showing that NAD(P)H oxidases, localized in the plasma membrane are major producers of ROS in plants under both normal and stress conditions (Foreman et al., 2003; Sagi and Fluhr, 2006; Wang et al., 2016). Heat stress decreased the activity of NAD(P)H oxidase in *Gossypium hirsutum* plants (Snider et al., 2009). In the present study, cucurbit plants respond

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differentially by the means of NAD(P)H oxidase activity. Since ROS are both substrate and product in NAD(P)H oxidase enzyme reactions, the change in NAD(P)H oxidase activity thought to be related with the ROS scavenging capacity, thus tolerance of plants to heat stress.

The changes in AsA content of the leaf tissues are shown in Figure 2. According to average values, the AsA content did not change in both melon cultivars (Figure 2A) and watermelon (Figure 2B). Statistical analysis revealed no significant effect of temperature, and cultivar on AsA content of melon cultivars. Although the interaction of treatment*cultivar was statistically significant (p<0.05). The AsA content slightly increased parallel to temperature increment (p<0.005) in zucchini (Figure 2C) and was the highest at 55 °C (0.66 mg/gFW).

The AsA scavenge the most dangerous ROS forms, namely •OH, O$_2^-$ and H$_2$O$_2$, through the reaction catalysed by APOX while GSH involves in the maintenance of reduced state of the cellular pool of AsA via the Halliwell-Asada pathway (Asthir, 2015). The absence of change in AsA content of melon and watermelon cultivars in response to heat treatments indicated that AsA has no effect on heat-tolerance of this species. Besides the increment of AsA content in zucchini plants may related to ROS-scavenging capacity. In lily plants AsA concentration was largely unaffected compared to the control at 37 °C and 42 °C, except for a significant increase after 2 h of 42 °C. The AsA concentration showed slight alternation within first six hours and then significantly decreased compared to the control at 47°C (Yin et al., 2008). The AsA content of strawberry plants did not change depending on heat stress (gradual temperature increment or heat shock) and were not effective in gaining tolerance (Ergin et al., 2016).

A gradual increase of temperature resulted in an activity peak at 50 °C, 45°C, 40 °C respectively for CAT, GR, and APOX enzymes that rapidly declined to nearly control levels in both cultivars (Table 1). The highest CAT enzyme activities were 48.55 nmol/mg protein/min and 48.13 nmol/mg protein/min in cv. Miranda and cv. Poli, respectively (p<0.005). Statistical analysis revealed a significant effect of heat treatment, cultivar, and the interaction of heat treatment and cultivar on CAT activity. The peak values of GR enzyme activity were 5.56 units/mg protein/min and 5.54 units/mg protein/min in cv. Miranda and cv. Poli, respectively (p<0.005). Statistical analysis revealed a significant effect of heat treatment, cultivar, and the interaction of heat treatments and cultivar on GR activity. The highest APOX enzyme activities were 0.44 μmol/mg protein/min and
Statistical analysis revealed a significant effect of heat treatment, cultivar, and the interaction of heat treatment and cultivar on APOX activity. Heat stress caused an activity peak at 40 °C and 35 °C for CAT (14.37 nmol/mg protein/min) and GR (40.60 units/mg protein/min) enzyme activities in watermelon, respectively then rapidly declined (Table 1). Although showing fluctuations in the APOX activity, a sharp decrease, from 0.15 µmol/ mg protein/min to 0.01 µmol/ mg protein/min, was present at 35 °C and never reached to control levels. Statistically significant differences were found between heat treatments on CAT, GR and APOX activities of watermelon plants (p<0.05).

### Table 1. The CAT (nmol/mg protein/min.), GR (units/mg protein/min), APOX (µmol/mg protein/min.), S-POX (units/mg protein), and CWB-POX (units/mg protein) activities of cucumber plants at different heat treatments. Values are mean ± SE (n = 3). Different letters indicate significant difference among various temperature treatments of each species/cultivars (p < 0.05)

| Species/Cultivar | Treatment | CAT        | GR         | APOX       | S-POX       | CWB-POX     |
|-----------------|-----------|------------|------------|------------|-------------|-------------|
| **Melon/Miranda** | Control   | 33.07±2.29c | 3.14±0.27d | 0.34±0.03bc | 715.18±23.68ab | 866.29±69.16a |
| 35°C            | 35.87±2.29bc | 3.51±0.22cd | 0.38±0.03ab | 514.55±23.68bc | 735.40±56.47ab |
| 40°C            | 38.97±2.29bc | 4.49±0.22b  | 0.44±0.03a  | 468.49±29.00c  | 627.10±56.47ab |
| 45°C            | 41.93±2.81bc | 5.56±0.27a  | 0.37±0.03ab | 563.70±23.68bc | 649.94±56.47ab |
| 50°C            | 48.55±2.29a | 3.95±0.27bc | 0.44±0.03a  | 655.84±29.00b  | 841.37±69.16a |
| 55°C            | 36.07±2.81bc | 2.44±0.27e  | 0.20±0.03d  | 677.00±23.68ab | 565.93±69.16bc |
| 60°C            | 34.50±2.81c | 3.48±0.27cd | 0.27±0.03dc | 764.29±29.00a  | 350.47±69.16c |
| **Melon/Poli**   | Control   | 23.41±2.29b | 3.27±0.27d | 0.43±0.03bc | 568.99±29.00c  | 459.79±56.47ab |
| 35°C            | 27.47±2.29b | 4.21±0.22bc | 0.48±0.03ab | 668.16±29.00b  | 497.92±69.16a |
| 40°C            | 39.43±2.29a | 5.23±0.27a  | 0.54±0.03a  | 599.03±29.00bc | 463.47±69.16ab |
| 45°C            | 44.10±2.29a | 5.01±0.22ab | 0.47±0.03ab | 487.51±29.00d  | 561.60±69.16a |
| 50°C            | 48.13±2.29a | 5.54±0.27c  | 0.38±0.03c  | 770.81±29.00a  | 488.76±69.16a |
| 55°C            | 40.99±2.29a | 4.88±0.27ab | 0.37±0.03c  | 534.46±23.68cd | 420.26±69.16ab |
| 60°C            | 24.62±2.29b | 4.03±0.27cd | 0.29±0.03d  | 303.66±23.68e  | 330.16±56.47b |
| **Watermelon/Crimson Tide** | Control | 5.71±1.81a | 17.68±8.3d | 0.15±0.01c | 4051.31±41.67a | 0.13±0.19a |
| 35°C            | 6.11±1.47a | 40.60±1.02e | 0.01±0.01a  | 3735.65±41.67a | 0.31±0.24a |
| 40°C            | 14.37±1.81b | 2.61±1.02a  | 0.07±0.01b  | 3071.73±41.67a | 0.24±0.19a |
| 45°C            | 13.96±1.47b | 5.07±0.83ab | 0.02±0.01a  | 3729.31±41.67a | 0.34±0.24a |
| 50°C            | 9.35±1.47ab | 6.38±0.83bc | 0.02±0.01a  | 2937.24±41.67a | 0.11±0.24a |
| 55°C            | 5.79±1.47a | 5.43±0.83abc | 0.06±0.01b | 2683.21±41.67a | 0.30±0.24a |
| 60°C            | 13.2±1.47b | 8.09±0.83c  | 0.03±0.01a  | 6072.63±41.67b | 2.73±0.24b |
| **Zucchini/Asma** | Control | 72.04±8.66b | 13.78±abc  | 0.05±0.02a  | 3186.39±170.62c | 0.08±0.01bcd |
| 35°C            | 127.23±8.66c | 18.80±e   | 0.03±0.02a  | 1473.54±170.62b | 0.09±0.01bcd |
| 40°C            | 57.75±10.61ab | 14.42±abc  | 0.08±0.02a  | 1528.96±170.62b | 0.05±0.01a |
| 45°C            | 78.58±10.61b | 12.41±ab   | 0.31±0.02c  | 1290.83±170.62ab | 0.07±0.01abc |
| 50°C            | 25.55±10.61a | 10.94±a    | 0.14±0.02b  | 1319.86±170.62ab | 0.11±0.01d |
| 55°C            | 87.63±10.61b | 17.35±bc   | 0.14±0.02b  | 911.06±170.62a  | 0.09±0.01cd |
| 60°C            | 29.27±8.66a | 14.24±abc  | 0.16±0.02b  | 2815.35±170.62c | 0.06±0.01ab |

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The highest CAT enzyme activity (127.38 nmol/mg protein/min) of zucchini was observed at 35 °C (Table 1). Similarly, the highest GR enzyme activity (18.80 units/mg protein/min) was found at 35 °C. The activity of APOX enzyme increased up to 0.31 µmol/mg protein/min at 45°C. Statistical analysis revealed a significant effect of heat treatment on CAT, GR, and APOX activities (p<0.05). Increased ROS production due to abiotic stresses causes changes in their homeostasis (Polle, 2001), but also acts as signals to activate stress responses. To overcome possible oxidative stress, plants produce enzymatic (APX, CAT, GR and SOD) and non-enzymatic, antioxidants (Choudhary and Agrawal, 2014). Under heat stress conditions seedlings of watermelon cv. Crimson Sweet showed much higher CAT activity than Charleston Gray and Fairfax cultivars, although, parallel to increasing temperatures, CAT activity in the cotyledons and embryos of all watermelon cultivars increased (Dantas et al., 2015). Ara et al. (2015) found that, when plants of heat-tolerant Cucurbita moschata, thermolabile Cucurbita maxima and their moderately heat-tolerant interspecific inbred line “Maxchata” genotypes were exposed to moderate (37 °C) and severe (42 °C) heat shocks, the enzyme activities of CAT and APOX in their leaves were increased with heat stress in tolerant genotypes of Cucurbita moschata followed by “Maxchata”. Different responses of leaf antioxidative enzymes were also reported in strawberry plants. While CAT and APX activities increased with high temperatures, GR activity was almost unchanged (Ergin et al., 2016). The activities of CAT, GR, and APOX enzymes in the leaf extracts of control and high temperature (40 °C) were assayed and activities of all enzymes were high in all the mulberry cultivars in response to high temperature (Chaitanya et al., 2002).

The S-POX activities of both melon cultivars indicated significant differences between heat treatments (Table 1). The S-POX activity was significantly greater at 60 °C in cv. Miranda (764.29 units/mg protein) and at 50 °C in cv. Poli (770.81 units/mg protein). Thus, the effects of heat treatments on S-POX enzyme activity were found statistically significant (p<0.05). The highest CWB-POX enzyme activity was detected at control treatment with 866.29 units/mg protein in cv. Miranda and at 45 °C in cv. Poli (561.60 units/mg protein). Statistical analysis revealed a significant effect of heat treatments on CWB-POX enzyme activity of cv. Miranda was statistically significant. Besides, statistical analysis revealed that, the differences between cultivars and heat treatment*cultivar interaction were statistically significant (p<0.05)

The effects of heat treatments on S-POX and CWB-POX enzyme activities of watermelon is shown in Table 1. The highest enzyme activities were at 60 °C with 6072.63 units/mg protein and 2.73 units/mg protein for S-POX and CWB-POX enzymes, respectively. In terms of S-POX and CWB-POX enzyme activities, statistically significant differences were detected between heat treatments (p<0.05).

The S-POX enzyme activity of zucchini was the highest at control treatment and decreased with increasing temperatures (55 °C), then increased up to 2815.35 units/mg protein at 60 °C (Table 1). The highest and the lowest CWB-POX enzyme activity were observed at 50 °C (0.11 units/mg protein) and 40 °C (0.05 units/mg protein), respectively. Statistical analysis revealed a significant effect of heat treatments on both S-POX and CWB-POX activities (p<0.05).

Total peroxidase activity of plants is as a biomarker for stress treatments. However, results are not clear, due to the differentially regulated isoenzymes (Mika et al., 2010). Besides, the relationship of CWB-POX activity to heat stress treatments remains unclear (Turhan et al., 2014). In the present research, S-POX had effective means in determining heat tolerance of melon and zucchini plants. Previously, Turhan et al. (2014) showed that the S-POX activity was greater in the first bloom stage than in the yield stage in tomato cultivars. In contrast to S-POX activity, the CWB-POX activity
was greater in the yield stage than in the first bloom stage and generally CWB-POX activity was high in the samples in response to heat stress treatments (Turhan et al., 2014). Gulen and Eris (2004) reported that total and specific POX activities were significantly increased by high temperatures in strawberry plants.

A Native PAGE was performed to observe the acidic and basic isoperoxidase profiles of leaves of cucurbit plants (Figure 3). Six acidic isoperoxidase bands were found in melon cultivar with Rf values of 0.14, 0.42, 0.45, 0.48, 0.52, 0.55. Two bands (Rf=0.45 and Rf=0.48) were common for both melon cultivars. The intensity of the bands was decreased with heat treatments in cv. Miranda. However, in cv. Poli the bands were disappeared at 60 °C (Figure 3A). There were no basic isoperoxidase band was found in both melon cultivars.

There were four acidic isoperoxidase bands (Rf= 0.06, 0.20, 0.76, 0.88) in watermelon plants, though the appearance of the bands were not affected by the heat treatment (except 40 °C) (Figure 3B). There were no basic isoperoxidase band was found in both watermelon cultivars.

The effects of heat treatments on isoperoxidase profiles of zucchini plant are shown in Figure 3C-D. Four acidic (Rf= 0.31, 0.36, 0.40, 0.59) and one basic (Rf=0.73) isoperoxidase bands were found in leaf tissues of zucchini plants. The intensity of the acidic isoperoxidase bands were decreased with heat treatments. Besides, the basic isoperoxidase band was disappeared at 60 °C (Figure 3D).

![Figure 3](image.png)

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(Walter, 1992). In previous studies, one basic isoperoxidase band (Rf =0.22) with different intensity due to heat treatments was detected in strawberry plants (Gulen and Eris, 2004). Similarly, Ergin et al., (2012) reported that there was no acidic isoperoxidase band was observed in strawberry plant under heat stress conditions, and a single sharp basic isoperoxidase band with Rf= 0.55 was observed in all treatments except 60 °C treatment.

4. CONCLUSIONS
In conclusion, NAD(P)H oxidase and antioxidative enzyme activities were increased generally after 50 °C. The results clearly show that the cv. Poli is superior with respect to its antioxidant defense systems and should be more tolerant than cv. Miranda due to higher ROS-scavenging systems. The activity of POXs (APOX and S-POX) become prominent in melon and zucchini while the activity of CAT become prominent in watermelon. Acidic isoperoxidase bands with different relative mobility values were found in all species. Besides, basic isoperoxidase band could not be determined in both melon cultivars and watermelon while a basic isoperoxidase band was found in zucchini. The cucurbit species/cultivars showed different responses to the stress imposed, suggesting different adjustment and tolerance mechanism. Additional researches must be performed regarding other antioxidant responses to each cultivar studied in this work, as well as other pathways involved in stress tolerance. Besides, the results obtained in this research forms a basis for further studies at the molecular level.

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