Antioxidants enzyme activity in Brassica oleracea var. acephala under Cadmium stress

Kanita Šabanović¹, Ahmet Yıldırım², Jasmin Šutković³*
¹,²,³Genetic and Bioengineering, International University of Sarajevo (IUS)

*Corresponding author: jsutkovic@ius.edu.ba

Abstract

When a plant is under heavy metals stress, it has different mechanism of coping with it. Brassica oleracea var. acephala (kale) is a plant that has an ability of heavy metal accumulation and removal of heavy metals from the ground. The plants were exposed to 50, 100, 200, and 500 μM of CdCl₂ for 5 days, in controlled in vitro conditions. Root length was measured to confirm the Cd effect on plant growth. There are five key antioxidants enzymes responsible for the regulation of heavy metals stress: superoxide dismutase (SOD), catalase (CAT), ascorbate peroxidase (APX), Peroxidase (POD) and Polyphenol oxidase (PPO). All enzymes showed significant activity, especially triggered by 500 μM CdCl₂ in both varieties. The domestic sorts seem more resistant if compared to hybrid variety, showing significant lower expression of antioxidants enzymes at higher concentrations. In general, significant percentage of enzymes is more expressed in the hybrid Italian sort, Nero di Toscana, indicating the ability of domestic sorts to be more resistant to heavy metal stress.

Keywords: heavy metals, Brassica oleracea, Antioxidant enzymes, Superoxide Dismutase

1. Introduction

The periodic system of elements contains 90 natural elements. Of those 90, 53 are classified as heavy metals [1]. Some metals are micronutrients – meaning they are essential to human in small doses, but some, such as Pb, As, Hg and Cd are nonessential and toxic to human. However, there is no widely accepted definition of heavy metals, but they are considered as elements that has a density more than 5g/m³ [2].

Out of all heavy metals – Cadmium is one of the most toxic one. It is a non-essential element that can appear it the nature both naturally and by anthropogenic factor [3,4]. Human can be exposed to cadmium through food, water, as well as tobacco, drugs, cosmetics or dietary supplements [5–7]. Main foods through which cadmium is ingested into the body are the vegetables and cereals but can also be through animals – kidney and liver since they accumulate cadmium. Through creating oxidative stress, cadmium is inducing tissue injury, epigenetic changes, inhibiting or upregulating different transport pathways, inhibition of heme synthesis [5–8].

Cadmium and his compounds are, according to International Agency for Research on Cancer (IARC) classified as Group 1 - meaning “carcinogenic to humans”. According to IARC there is enough evidence that cadmium contributes to development of lung cancer, and limited evidence for development of prostate and kidney cancer [9].

The major site for cadmium accumulation in the body are the kidneys [10], and it is associated with kidney cancer [11]. It also affects the gut microbiota, the abundance, and its relative population. In the intestinal walls it induces cell damage and inflammatory response [12]. Cadmium can also accumulate in the endometrium.
Thus, it is a potential causative of estrogen-diseases, such as endometrial or breast cancer, spontaneous abortions, and endometriosis. Higher intake of cadmium is related to higher endometrial cancer risk. In the endometrial cells they play an important role in placentation, embryogenesis and implantation [13]. Cadmium exposure can affect the sperm quality, penetration into oocytes and the embryo development of one derived from natural fertilization [14]. It also affects female reproduction, balance of reproductive hormones and menstrual cycles. Furthermore, during pregnancy, cadmium exposure affects female offspring more than male [15]. The main source of cadmium in foods is coming from plants [4]. It is causing phytotoxicity and by this becoming a threat for both human and animal trough food chains. It is highly mobile in water-soil-plant system since it is more soluble in water than other metals [16].

According to the World Health Organization, The Joint Food and Agriculture Organization of the United Nations (FAO)/WHO Expert Committee on Food Additives (JECFA) the – provisional tolerable monthly intake (PTMI) is 25 µg/kg body weight. Cadmium has a long biological half-life and is a toxican that is cumulative [6]. It has a half-life of 25 years, representing depositions in the organs, not removal from the body [5]. It takes decades for cadmium to be removed from the tissues and organs from the body [17], and is mainly removed with fecal and urinary excretion [8].

The presence of cadmium induced the inhibition of growth, oxidative damage, and genotoxicity in the seedlings, increasing the activities of antioxidant enzymes and proline accumulation as well as limiting the uptake of potassium (K) and calcium (Ca) [18]. Furthermore, it causes oxidative stress, inhibition of enzymes and inactivation of biomolecules [19]. Plant can resist to heavy metal stress by avoiding it, meaning being protected externally, and by tolerating it, meaning coping with the stress internally [20]. There are different mechanisms to prevent the entry in the roots, such as thick cuticle, cell walls, trichomes. If the heavy metals, despite the biophysical barriers, enter the plant, the plant will start to synthesize various biomolecules. Those include biomolecules such as chelators or metallochaperones and cellular exudates or metallothioneins. If these, above mentioned mechanisms, are not able to resist the heavy metal contamination, there will be an increase of reactive oxygen species (ROS) [21].

Free radicals are molecules that can exist independently and contains one or more unpaired electrons alone in an electron orbital. The oxidative metabolism is producing free radicals that are oxygen-centered, as well as other reactive oxygen species (ROS) [22]. In plants, ROS can cause DNA damage, unspecific oxidation of membrane lipids and proteins, enzyme inhibition by activation of apoptosis [19].

However, since ROS are playing an important role of the plant defense system, it cannot be totally omitted from the plant. ROS will begin to be harmful when the concentration exceeds the threshold for defense mechanisms [19] and cause oxidative stress [23]. In normal conditions, plants have a mechanism to cope with ROS. But under stress, the production on ROS is increased as the cell homeostasis is disrupted. For a plant to cope with this kind of disorder, the antioxidative mechanisms are activated [24]. An antioxidant is a stable molecule that is donating an electron to the free radical and thus, neutralizing it. By neutralizing the free radicals, the antioxidant is reducing its ability to cause a damage. There are different enzymatic and non-enzymatic antioxidants, which can be found in the diet, or produced by the body metabolism [25]. Non-enzymatic are ones such as glutathione, vitamin C, vitamin E; and enzymatic antioxidant are ones such as peroxidase (POD), catalase (CAT), ascorbate [25], superoxide dismutase (SOD) [22], polyphenol oxidase (POD) [26], ascorbate peroxidases (APX) [27] etc. Levels of ROS-cleaning enzymes or antioxidants are in part determining the ability of a plant to tolerate different environmental stresses [28].

Superoxide dismutase (SOD) is one of the most important enzymes that is enabling organism to cope with the presence of molecular oxygen [23]. It is the first enzyme to detoxicate the ROS generated. [29]. Catalyze (CAT) is an enzyme which is found in almost all living organisms. It is reducing hydrogen peroxide, a harmful byproduct of normal metabolic processes, to hydrogen and water [25]. By removing the hydrogen peroxide, CAT is preserving cell walls from membrane damage and lipid production. [29]. Polyphenol Oxidase (PPO) can be found in fungi, plants, humans, as well as in some higher animals. It is also present in higher plants where it protects the plants against microorganisms and insects [30]. The generation of o-quinone which is mediated by PPO is also connected with generation of ROS (as a secondary reaction product) [31]. Peroxidases are present in plants, animals, and microorganisms. In plants, they are responsible for various functions, such as general stress response, rigidification, wound healing (which appears after stress which will induce POX formation), lignification, suberization, construction, strengthening the cell walls, defense against pathogens or protecting the tissues against physical damage. [32,33].
Ascorbate Peroxidase (APX) is a key enzyme responsible for the removal of excess hydrogen peroxide under both stress and normal conditions. It is responsible for reduction of hydrogen peroxide to water and monodehydroascorbate, using ascorbate as a substrate.

The genus Brassica is including a vast variety of more than 30 species, with different varieties and hybrids. Brassica vegetables contain glucosinolate, polyphenol, carotenoid, provitamin A, Vitamin C, K, B9 and B2, Calcium, as well as some antioxidant enzymes, CAT, POX and SOD.

Brassica oleracea varieties include a variety of vegetables that human is using for consumption, such as broccoli, cauliflower, cabbages, Brussel sprouts and others. They are used as a food, oil production, animal fodder and others. Among the Brassica vegetables, Brassica oleracea var. acephala (kale) has more calcium, vitamin C, K, A, B2 B9 content that the others.

The Brassica vegetables have beneficial properties such as anticarcinogenic properties, protecting against cardiovascular diseases, protecting against ageing processes, etc. Some Brassica species has been found out to be good heavy metal accumulators, and thus, used in phytoremediation process. They have the ability to accumulate heavy metals in the parts above the ground, tolerance to the high concentration of heavy metals in the soils, rapidly grow and highly accumulating biomass and they are easy to harvest and to grow as an agricultural crop.

The aim of this study is to examine different sorts of Brassica oleracea var. acephala for their enzymatic activity in kales of domestic and hybrid origins.

2. Materials and methods

2.1 Seed germination and root analysis

The seeds that were used was a hybrid variety kale NT (Nero di Toscana, an Italian sort) and a domestic variety sort from Stolac (village Ravine) from the region of Herzegovina. The seeds were germinated using the tap and paper method. Two layers of paper tissues were placed in each petri dishes (9 cm diameter). Each petri dish was moistened with different concentration of cadmium-chloride (Table 1). The stock solutions were prepared from cadmium-chloride (CdCl₂) from Sigma-Aldrich and distilled water. Thirty seeds were placed in each petri dish. Each petri dish contained seeds with different concentrations of CdCl₂.

| Petri-dish | 1 | 2 | 3 | 4 | 5 |
|------------|---|---|---|---|---|
| Concentration | 0 µM (Control) | 50 µM | 100 µM | 200 µM | 500 µM |

The petri-dishes were then placed in a Growth Chamber for 5 days, at the temperature of 27°C with 16 hours of light per day. After 5 days the seeds were collected, roots were measured for each plant using a ruler. The plants from each petri dish where placed in separate tubes (i.e., concentration) were stored at -80°C fridge, until it is needed for further experiments.

2.2 Plant tissue extraction for enzyme activity determination

Extraction buffer of 100 mM phosphate (pH = 7) is made. The root tissues from plants were placed in a mortar and ground under the liquid nitrogen using a pestle. Weight of 0.3 – 1.0 g of obtained plant powder is transferred to the 1.5 mL Eppendorf tube, and triple amount of phosphate buffer of the plant powder of is added. Stir. The tubes are transferred to the centrifuge and centrifuged at 13,000 x g for 20 minutes and at 4°C. After the centrifugation, the supernatant is carefully taken with the pipette and transferred to new Eppendorf tubes.
2.3 Enzymes activity determination assays

2.3.1 Peroxidase (POX) activity determination assay

A 3 mL of 0.1 M phosphate buffer (pH = 6.5) was added to the tube. Then, 100 μL of 20 mM guaiacol (prepared under the hub) and 100 μL of enzyme was added. To initiate the reaction, 30 μL of 12 mM hydrogen peroxide was added. The process is repeated for each obtained enzyme (i.e., each petri dish). Control group is made, with the hereby reaction mixture, but without the enzyme. A 200 μL from each tube and place it in each well of the 96-well plate. Measure the absorbance at 436 nm at 0 min and 2 min [43]. Peroxidase is catalyzing the hydrogen removal from large number of molecules, as well as decomposition of hydrogen peroxide. POX will bind hydrogen peroxide and the complex obtained, [POD-H2O2], can oxidize different hydrogen donors. By formation of an oxidized compound, among other methods, the activity of POX can be determined. Guaiacol is a non-cancerogenic substrate, which acts as a substrate for POX, where one mole of guaiacol is oxidized by one mole of hydrogen peroxide. The end product of this reaction is tetraguaiacol which is measured spectrophotometrically at 436 nm [43–45]. In the experiment performed, guaiacol and enzyme source are firstly added, where the guaiacol acts as a substrate. For reaction to start, hydrogen peroxide was added, and as a result tetraguaiacol was formed. Thus, it is expected that the absorption is increasing, with the increasing concentration of cadmium (more cadmium, more POX, more tetraguaiacol).

2.3.2 Superoxide dismutase (SOD) activity determination assay

A 1.3 mL of 50 mM of sodium carbonate buffer is added to the test tube. Then, 500 μL of 96 mM Nitroblue tetrazolium (NBT) (pH = 10.0) and 100 μL of 0.6% Triton X-100 was added. To initiate the reaction, 100 μL of 20 mM hydroxylamine hydrochloride (NH₂OH.HCl) (pH = 6.0) was added. A 70 μL of enzyme is added. The process is repeated for each obtained enzyme (i.e., each petri dish). Control group is made, with the hereby reaction mixture, but without the enzyme. A 200 μL from each tube and place it in each well of the 96-well plate. Measure the absorbance at 540 nm at 0 min and 2 min [46]. Hydroxylamine will autoxidize and generate superoxide radicals. NBT, which is colorless, will be reduced by superoxide radicals, and turn NBT to blue formazan, which can be measured spectrophotometrically. The SOD, which is scavenging the superoxide radicals is going to inhibit the blue color formation. Thus, as more SOD is present, less color will appear [46, 47].

2.3.3 Polyphenol oxidase (PPO) activity determination assay

A 1.5 mL of 50 mM phosphate buffer (pH = 6.5) was added to the tube. Then, 200 μL of the enzyme is added. To initiate the reaction, 200 μL of 100 mM catechol was added. The process is repeated for each obtained enzyme (i.e., each petri dish). Control group is made, with the hereby reaction mixture, but without the enzyme. A 200 μL from each tube and place it in each well of the 96-well plate. Measure the absorbance at 495 nm at 0 min and 2 min [48]. For Polyphenol Oxidase, catechol is used as a substrate, which will, in the present of oxygen form quinone. The amount of quinone is measured spectrophotometrically. Thus, as more PPO is present, more quinone is generated [49].

2.3.4 Catalase (CAT) activity determination assay

A 1.9 ml of 50 mM PBH (pH = 7.0) was added to the tube. Then, 0.1 ml of the enzyme was added. To initiate the reaction, 1.0 mL of 0.075% mM hydrogen peroxide was added. The process is repeated for each obtained enzyme (i.e., each petri dish). Control group is made, with the hereby reaction mixture, but without the enzyme. A 200 μL from each tube and place it in each well of the 96-well plate. Measure the absorbance at 240 nm at 0 min and 2 min [50]. Catalase is decomposing hydrogen peroxide to water and oxygen. The decomposition of hydrogen peroxide is measured at 240 nm [50]. Thus, as more CAT is present, less hydrogen peroxide will be present, as the catalase will decompose the hydrogen peroxide.
2.3.5 Ascorbate peroxidase (APX) activity determination assay

A 1.5 ml of 100 mM phosphate buffer (pH = 7.0) is added to the tube. Then, 300 μL of 5mM ascorbate, and 600 μL of enzyme is added to each tube. To initiate the reaction, 0.5 mM of 600 μL of hydrogen peroxide is added. The process is repeated for each obtained enzyme (i.e., each petri dish). Control group is made, with the hereby reaction mixture, but without the enzyme. A 200 μL from each tube and place it in each well of the 96-well plate. Measure the absorbance at 290 nm at 0 min and 2 min [51]. Ascorbate Peroxidase is an enzyme that scavenge hydrogen peroxide, using ascorbate as an electron donor, which will oxidase. The amount of ascorbate is measured spectrophotometrically [52]. As more APX is present, less ascorbate will be available.

3. Results

3.1 Root measurements

![Avarage Root Length](image)

Figure 1. Average root length in domestic and hybrid sorts; with standard deviation

From the Fig. 1 it can be observed that the hybrid sorts have larger roots for all CdCl$_2$ concentrations, except for 500 µM. The root length decreases with the increase of CdCl$_2$ concentration, except for control group in both domestic and hybrid sorts. The root length was largest for the concentration of 50 µM, while the root length was smallest for the concentration of 500 µM, for both domestic and hybrid sorts. Also, root length is decreasing for both domestic and hybrid sorts, from the concentration of 50 µM.

In table 2, descriptive analysis of domestic and hybrid root lengths. The p-value (T<=t) for both one-tail and two tail are greater than 0,05, which represent insignificant difference.

| Table 2. T-Test: two-sample assuming equal variances; Domestic VS. Hybrids root length |
|-----------------|-----------------|-----------------|-----------------|
|                | Average of 2 min – domestic | Average of 2 min - hybrid |
| Mean           | 1,67362258      | 2,25177         |
| Variance       | 0,245413288     | 1,045952791     |
| t Stat         | -1,137048672    |                 |
| P (T<=t) one-tail | 0,144213248    |                 |
| t Critical one-tail | 1,859548038   |                 |
| P (T<=t) two-tail | 0,288426495    |                 |
| t Critical two-tail | 2,306004135   |                 |
3.2 Enzyme activity

3.2.1 Peroxidase (POX) Activity

In Fig. 3 we show that the first control group (0 Cd), the values at 0 min and 2 min for domestic sort was 0.036 for domestic sort and 0.051 for hybrid sort. Since no enzyme is present (POX) the reaction is not catalyzed. The activity of POX enzyme is larger in the hybrid sort, whereas in the domestic, it is 2 to 3 times smaller. For both sorts, POX activity is decreasing over time. In the domestic sort, POX activity at both 0 min and 2 min is smallest for the second control group and is greatest at the concentration of 200 at 0 min, and 100 µM at 2 min; after which activity of POX activity is increasing. For the hybrid sort, POX activity is smallest at 500 µM for 0 min and 200 µM for 2 min and is largest at 100 for both 0 min and 2 min.

In Table 3, descriptive analysis of domestic and hybrid for POX activity is shown. The p-value (T<≤t) for both one-tail and two tail are less than 0.05, which represent significant difference.

|                     | Average of 2 min – domestic | Average of 2 min - hybrid |
|---------------------|-----------------------------|---------------------------|
| Mean                | 0.2885                      | 0.63125                   |
| Variance            | 0.000013041666              | 0.000404                  |
| t Stat              | -33.5573458                 |                           |
| P (T<≤t) one-tail   | 0.00000000023               |                           |
| t Critical one-tail | 1.943180281                 |                           |
| P (T<≤t) two-tail   | 0.00000000466               |                           |
| t Critical two-tail | 2.4469118511                |                           |

3.2.2 Superoxide Dismutase (SOD) Activity

In Fig. 4 we show that the first control group (0 Cd), the values at 0 min and 2 min for domestic sort was 0.036 for domestic sort and 0.051 for hybrid sort. Since no enzyme is present (SOD) the reaction is not catalyzed. The activity of SOD enzyme is larger in the hybrid sort, whereas in the domestic, it is 2 to 3 times smaller. For both sorts, SOD activity is decreasing over time. In the domestic sort, SOD activity at both 0 min and 2 min is smallest for the second control group and is greatest at the concentration of 200 at 0 min, and 100 µM at 2 min; after which activity of SOD activity is increasing. For the hybrid sort, SOD activity is smallest at 500 µM for 0 min and 200 µM for 2 min and is largest at 100 for both 0 min and 2 min.
In Fig. 4 we see that the control group the absorption for domestic sort was 0.44 and 0.47 at 0 min and 2 min respectively, while for the hybrid sort the absorption was at both 0.51 for 0 min and 2 min, which is a mistake due to wrong pipetting. The absorption was the smallest for both domestic and hybrid sort. In domestic sort the SOD levels are fluctuating, rising from control to 50 µM, decreasing from 50 µM to 100 µM, again rising to 200 µM, and finally decreasing from 200 µM. In hybrids, SOD levels are increasing until the concentration of 50 µM, after which its levels are decreasing.

In Table 4, descriptive analysis of domestic and hybrid for SOD activity is shown. The p-value (T<\text{\leq}\text{t}) for both one-tail and two tail are less than 0.05, which represent significant difference.

| Table 4. T-Test: Domestic VS. Hybrids - SOD enzyme activity (absorbance) |
| --- |
| **Average of 2 min – domestic** | **Average of 2 min - hybrid** |
| Mean | 0.234 | 0.162 |
| Variance | 0.0000698333333333332 | 0.0002245 |
| t Stat | 8.39349367825105 |
| P (T<\text{\leq}\text{t}) one-tail | 0.0007786530296291 |
| t Critical one-tail | 1.9431802805153 |
| P (T<\text{\leq}\text{t}) two-tail | 0.0015573060592582 |
| t Critical two-tail | 2.44691185114497 |

3.2.3 Polyphenol Oxidase (PPO) Activity

In the first control group the absorption for both domestic and hybrid at 0 min and 2 min were 0.05 (±0.005). The highest values for domestic sorts were at concentration of 100 µM, and at 50 µM for hybrid sorts, after which the absorption decreases, for both 0 min and 2 min. The lowest values, omitting control groups, were at the concentration of 500 µM.

In Table 5, descriptive analysis of domestic and hybrid for PPO activity is shown. The p-value (T<\text{\leq}\text{t}) for both one-tail and two tail are less than 0.05, which represent significant difference.

| Table 5. T-Test: Domestic VS. Hybrids - PPO enzyme activity (absorbance) |
| --- |
| **Average of 2 min – domestic** | **Average of 2 min - hybrid** |
| Mean | 0.0780625 | 0.06775 |
| Variance | 0.000023515625 | 0.0000104166 |
| t Stat | 3.540685574 |
| P (T<\text{\leq}\text{t}) one-tail | 0.006104385 |
| t Critical one-tail | 1.943180281 |
| P (T<\text{\leq}\text{t}) two-tail | 0.002208771 |
| t Critical two-tail | 2.446911851 |
3.2.4 CAT activity

In the first control group the absorption for both domestic and hybrid at 0 min and 2 min were 3.45 (±0.005). The CAT activity did not change through time and remained almost the same with the change of Cd concentration for the domestic sort. For the hybrid sort, the absorption decreased through time, indicating more CAT utilization; but remained relatively stable through time. Furthermore, in hybrid, at 0 min, CAT levels were slightly increasing for 0 min, and slightly decreasing for 2 min.

In table 6, descriptive analysis of domestic and hybrid for CAT activity is shown. The p-value (T<=t) for both one-tail and two-tail are less than 0.05, which represent significant difference.

Table 6. T-Test: Domestic VS. Hybrids - CAT enzyme activity (absorbance)

|                      | Average of 2 min - domestic | Average of 2 min - hybrid |
|----------------------|----------------------------|---------------------------|
| Mean                 | 3.5548125                  | 3.4835                    |
| Variance             | 0.00009714                 | 0.00016                   |
| t Stat               |                            |                           |
| P (T<=t) one-tail    | 0.000057498957             |                           |
| t Critical one-tail  | 1.9431803                  |                           |
| P (T<=t) two-tail    | 0.000115                   |                           |
| t Critical two-tail  | 2.4469119                  |                           |

3.2.5 Ascorbate peroxidase (APX) activity

Figure 7. APX activity in domestic and hybrid sorts
The values for the first control groups were 2.53 and 2.65 at both 0 min and 2 min, for domestic and hybrid sort, respectively. The amount of APX through time is increasing, for both domestic and hybrid sorts. The amount of APX throughout different concentrations is fluctuating. In domestic sort, the highest values of APX were at 100 µM at both 0 min and 2 min, after which the amount of APX is decreasing. In hybrid sort the highest values were at control group and 100 µM at 2 min and 0 min, respectively. After concentration of 100 µM, the amount of APX is also decreased, but is again rising after 200 µM.

In table 7, descriptive analysis of domestic and hybrid for APX activity was done. The p-value (T<=t) for both one-tail and two tail are greater than 0.05, which represent insignificant difference.

| Table 7. t-Test: Domestic VS. Hybrids - APX enzyme activity (absorbance) |
|-----------------------------|-----------------------------|
| Mean                        | Average of 2 min – domestic | Average of 2 min - hybrid |
| Mean                        | 2.24981                     | 2.38538                     |
| Variance                    | 0.03663                     | 0.03093                     |
| T Stat                      | -1.04309                    | 0.16855                     |
| P (T<= One Tail)            | 1.94318                     | 0.3371                      |
| T Critical One-Tail         | 2.44691                     |                             |
| P(T<=T) Two-Tail            |                             |                             |

4. Discussion

The aim of this study was to examine the enzyme activity of domestic and hybrid sorts of Brassica oleracea var. acephala under cadmium stress. The damage can be repaired by antioxidants, which can be non-enzymatic and enzymatic. The presence of heavy metals cause H2O2 production, which will increase the level of oxidative enzymes [25].

Since the root is in direct contact with the heavy metal, they are affected more than other parts [38]. This study showed that the root length was highest for the concentration of 50 µM, followed by 100, µM, 0 µM (control), 200 µM and 500 µM. [38]. In the study conducted by Meng et al.,2009, the root length was higher at small concentrations (10 µM of Cd), positively affecting it [53].

According to them, the Cd was transported to shoots, which can explain this phenomenon. Also, a similar study explained this phenomenon as the disruption of the homeostasis occur, which will lead to overcompensation response [38].

The CAT activity for domestic kale was rising until the concentration of 100 µM, at both 0 min and 2 min, after which it started to decrease. These results are in consistence with the study of Taylor et al., 2013 [54], which showed that the CAT levels were rising until the concentration of 100 µM CdCl2 in Brassica juncea. However, for the hybrid sort, the CAT levels decreased after the concentration of 200 µM and 50 µM at 0 min and 2 min, respectively.

The levels or SOD in our study were rising, decreasing, rising, and decreasing in domestic, while in hybrid it was rising, and then decreasing. For POX, its levels were rising, and then dropping after concentration of 100 µM for domestic, and for hybrid rising and dropping, after the concentration of 50 µM. In the study conducted by Zhang et al., 2019 which was conducted on Brassica juncea L. (Indian mustard) and Medicago sativa L. (alfalfa), the SOD levels were increasing until the concentration of 150 mg/kg of Cd, after which the levels of that enzyme were dropping until the final concentration of 600 mg/kg [55]. For POX, the levels dropped until the concentration of 75 mg/kg of Cd.

In this study, the CAT activity was also examined, in which its levels were rising until 150 mg/kg, and then dropping. According to them, the rising and dropping levels of SOD and POD enzymes is a usual defense mechanism that plants have against stress caused by Cd. The CAT levels also remained relatively stable for different concentrations in Indian Mustard, with levels decreasing from the concentration of 100 µM. The trend of decreasing of CAT may be because CAT is most sensitive to heavy metal stress, which activity is firstly inhibited, which will result in H2O2 clearance obstruction, so that the superoxide can be only removed by the POD enzyme, and thus POD levels are increased. But with the Cd levels being increased, the
accumulation of ROS results in membrane lipid peroxidation, which is also increased, and thus POD will finally decrease.

The APX level were fluctuating, but the final increase of APX was recorded at the concentration after 100 µM for hybrid, and final decrease after 100 µM for domestic. A study conducted by Mobin & Khan, 2007of Varuna cultivar of Brassica juncea L., the APX levels were also increasing with the Cd increasing [56]. However, there is a threshold, a sub-lethal concentration, which induced maximum level of APX, after which the APX levels are decreasing. The APX, together with SOD is one of the key enzymes in H₂O₂ scavenging. The APX levels were higher in plants that are known as metal accumulators, such as Brassica juncea, Sedum alfredii, Triticum aestivum, as the upregulation of APX increases the ability to overcome Cd stress [57]. Furthermore, Armas et al., 2015 stated that in B. juncea, APX and CAT are not competing, but rather different enzyme classes that scavenge H₂O₂; APX for fine ROS modulation, and APX for excess ROS that accumulated during stress [58].

For PPO enzyme, this study showed that its levels decreased from concentration of 50 µM and 100 µM for hybrid and domestic sorts, respectively. However, in a study conducted by Kapoor et al., 2014 on Brassica juncea, the PPO levels were fluctuating, as they were rising, decreasing and again rising, with the highest levels at the concentration of 600 µM of Cd [59].

Finally, our study showed that out of 5 enzymes, taking in account samples at 0 min and 2 min, PPO and SOD showed more activity in domestic sorts, as well as POX for 0 min. Furthermore, considering both 0 min and 2 min, 50% of enzymes showed decrease at 100 µM, and 30% of enzymes showed decrease from 200 µM for domestic sorts; not considering CAT, which levels were rising. But for hybrid sorts, 66% present showed decreasing levels of enzymes from the concentration of 50 µM, and 16 % decreasing levels from the concentration levels of both 100 µM and 200 µM; not considering CAT and APX, which levels were rising. The decreasing effects of enzymes at higher concentrations can be explained in a way that plants start to die at higher concentrations, as it was explained by Hayat et al., 2007, where the plant starts to die at 200 µM and 250 µM. Lastly, for 4 out of 5 (80%) enzymes there was significant difference between the enzyme activity in domestic and hybrid sorts [60].

5. Conclusion

Heavy metals that are present in the nature can be accumulated by plants and cause various damaging effects on them. Present of heavy metals cause oxidative stress to occur. However, plants have certain mechanism by which they cope with different stresses that affect them, and among them – heavy metal stress. The most effective way of protection is the initiation of antioxidants proteins, both non-enzymatic and enzymatic. Our study compared domestic and hybrid sorts of Brassica oleracea var. acephala to examine which are more resistant to heavy metal stress by examining their enzyme activity. The conclusion that was made from this study is that 80% of enzymes showed significant difference between domestic and hybrid sorts, meaning that domestic sorts are more resistant to heavy metal stress. The domestic sorts are slightly more resistant if compared to the hybrids, showing decreased enzyme activities at higher concentrations.

6. References

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