Alleviation of Zn Toxicity in Germinated Wheat Grains (Triticum aestivum L.) by Seed Priming with Defensine Like Protein

Eman E. Selem¹ and Deyala M.Naguib

Botany Department, Faculty of Science, Zagazig University, Zagazig, Egypt.

The aim of this study is to examine the effect of priming the wheat grains with defensine like protein in a trial to enhance their tolerance of Zn toxicity. The present technique was performed on two groups of wheat grains that were germinated under different concentrations of Zn. The first group included the non-primed grains and the second group included the primed grains with defensine like proteins which was extracted and purified from fenugreek seeds. Growth and some physiological parameters of wheat seedlings were determined. Data revealed that increasing Zn concentrations in the non-primed grains reduced the growth parameters represented by germination ratio and length of both radicle and coleoptile. Also, some hydrolysis processes represented by amylase, acid and alkaline phosphatase were significantly decreased, such decrease was accompanied with a decrease in soluble carbohydrates and phosphorus content. Additionally, there was a significant inhibition in the activity of the antioxidant enzymes such as superoxide dismutase, polyphenol oxidase and peroxidase. In contrast, there was an enhancement in the growth process in the defensine primed grains treated with high Zn concentrations. This was associated with a reduction in the oxidative markers such as malondialdehyde (MDA) and H₂O₂ content. Compared with non-primed grains, an improvement in the activity of the antioxidant enzymes and controlling the excess of the ROS were recorded but no significant effect on the hydrolyzing enzymes was found.

Keywords: Antioxidant enzymes, Defensine, Germination, Hydrolysis processes, Oxidative stress, Zn toxicity.

Introduction

With the expansion of the population, the environmental pollution and toxicity by chemicals become of great concern worldwide. Rapid industrialization and urbanization processes have led to the incorporation of pollutants such as pesticides, petroleum products, acids and heavy metals in the natural resources like soil, water and air thus degrading not only the quality of the environment, but also affecting both plants and animals (Sethy & Ghosh, 2013). Some heavy metals can be highly toxic when their concentrations exceed threshold value, while others are essential micronutrients for plants in low concentrations, but in higher concentrations they may cause metabolic disorders and growth inhibition for most of the plant species (Kumari et al., 2016). The up-regulation of reactive oxygen species (ROS) induced by Zn stress may contribute to inhibitory effects on the activities of antioxidative enzymes including superoxide dismutase (SOD, EC 1.1.5.1.1), catalase (CAT, EC 1.11.1.6), ascorbate peroxidase (APX, EC 1.11.1.11), glutathione reductase (GR, EC 1.6.4.2) (Li et al., 2013). Another mechanism of heavy metal toxicity is their ability to bind strongly with oxygen, nitrogen and sulfur atoms. This binding affinity is related to free enthalpy of the formation of the product of metal and ligand. Thus, due to these activities, heavy metals can inactivate the enzymes by binding to cysteine residues (Wani et al., 2012).

Zinc phytotoxicity showed a considerable attention because it is a part of the long term utilization of fertilizers, also industry increased its content in the surface soil (Tsonev & Lidon, 2012). It is known that Zn can replace Mg²⁺ ions and it has an important role as functional, structural and regulating cofactor (Molnárová & Fargašová, 2016).

Priming is a seed treatment before sowing to improve their germination, it includes soaking seeds in a solution where the first two phases of germination (imbibition and activation phases)
take place without emergence and growth of the radicle. This technique has proven its effectiveness for good crop establishment under stress (Elouaer et al., 2012).

Plant defensins are small, cysteine-rich proteins (45–54 amino acids) that have been isolated from many plant species and tissues (Lay & Anderson, 2005). A variety of functions have been attributed to plant defensins. While many have antifungal activity, plant defensins have also been described with functions in antibacterial activity, zinc tolerance and blocking of ion channels (Carvalho & Gomes, 2009 and van der Weerden & Anderson, 2013).

A defensin like protein from wheat was also found to be induced during cold acclimatization (Koike et al., 2002). Induction of gene expression of plant defensins by drought or salt stresses has also been reported (Yamada et al., 1997 and Maitra & Cushman, 1998).

Seed germination is one of the most susceptible physiological phenomena in the plant life cycle that is affected by various biotic and abiotic environmental factors (Moosavi et al., 2012), so, any adverse effect on seed germination process would be detrimental for the establishment and healthy growth of seedlings (Donohue et al., 2010).

An excessive Zn metals in soils can have a negative impact on seed germination, biomass production, root growth, root morphology and architecture. It can also interfere with the activities of many key enzymes related to normal metabolic processes (Dhankhar, 2011). Study on Zn toxicity on Phaseolus vulgaris and Mung bean revealed that the high concentrations of Zn caused significant decrease in germination and seedling growth represented in root and shoot length (Hojiboland et al., 2006). In the research of Dhankhar (2011) investigating Vigna mungo (L.), it was observed that the increase in the Zn concentration from 0.25 to 1.50mM caused decreasing plumule and radicle lengths. In the experiment of Samuilov et al. (2014) zinc inhibited the growth of both shoots and roots. Also Zn concentrations of 100 – 400µg g⁻¹ (soil d.m.) cause significant decrease in root and shoot growth parameters at different developmental stages of Artemisia annua plants (Khudsar et al., 2004).

Thus, this work attempts to apply the technique of priming with definsine to improve the germination of wheat grains under the influence of Zn toxicity.

Material and Methods

Extraction and purification of Definsine from Fenugreek seeds

The natural defensin from fenugreek seeds (Trigonella Foenum Graecum), was purified according to Oddepally & Guruprasad (2015) with some modification as follows; fine flour (100g) was prepared from the fenugreek seeds in a mill. A protein extract was prepared from this flour using 500ml of extraction buffer (10mM Na₂HPO₄, 15mM NaH₂PO₄, 100mM KCl, 1.5% EDTA, pH 5.4) for 2h at 4°C with constant agitation. The protein extract was centrifuged at 15,000g for 15min. at 25°C, and the supernatant was fractionated by ammonium sulfate with 70% relative saturation at 4°C for 18h. After centrifugation under the same conditions, the precipitate was redissolved in distilled water and heated at 80°C for 15min in a water bath. The heated protein extract was centrifuged at 10,000 g for 15min. at 25°C and the supernatant was recovered and extensively dialyzed against distilled water for three days and then recovered by freeze drying. For peptides, purification was initially performed on a DEAE-Sepharose column (with 100ml of resin) equilibrated with 20mM Tris–HCl (pH 8.0) at flow rate of 60ml/h. The freeze dried protein extract (50mg) was reconstituted in 5ml of the equilibrium buffer and centrifuged (16,000g for 3min at 4°C) and the supernatant was loaded onto the column. A non-retained fraction (D1) was eluted in the equilibrium buffer and examined by SDS protein gel electrophoresis.

Sodium dodecyl sulfate polyacrylamide (SDS) gel electrophoresis (SDS-PAGE) was performed according to the method of Laemmli (1970) as modified by Studier (1973). One-dimensional SDS-PAGE was run in a mini-gel electrophoresis apparatus (Cleaver Scientific Ltd, OmniPAGE Gel Casting System). Gel was scanned and analyzed with Bio-Rad Video Gel documentation.

Plant material and Zn treatments

Wheat grains were sterilized in 10% Na hypochlorite solution for 10min, and the grains were washed thoroughly 4-5 times with sterilized distilled water. The grains were divided into two groups; the first was soaked overnight in the prepared definsine solution and the second was soaked in distilled water.

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Five concentrations of ZnSO$_4$ were used in this study. The selected concentrations were 0.0, 1.0, 10, 50 and 100mM. For each treatment, the pH was adjusted to 6.5.

**Determination of germination parameters**

Grain germination test on filter paper was carried out in glass Petri dishes (15cm) with three layers of filter paper on the bottom. Each dish contained 10ml of ZnSO$_4$ solution, and 20 grains, covered by lid. Petri dishes containing the grains with 10ml of distilled water (free of Zn solution) served as control. Effluents were applied every alternate day. Petri dishes were incubated at 25°C ± 1 with 16h light and 8h dark cycles in a growth chamber for 3 days with five replicates and all parameters (germination ratio, root length and coleoptile length) were recorded at 24, 48 and 72h.

**Biochemical study**

The biochemical parameters were determined at 3 different time intervals (24, 48 and 72h), as follows:

**Effect on antioxidant machinery**

**Determination of hydrogen peroxide (H$_2$O$_2$) content:** The content of H$_2$O$_2$ as a type of reactive oxygen species (ROS) was determined according to the method of Alexievia et al. (2001), in which 1g of germinated wheat seedlings was homogenized in 0.1% trichloroacetic acid (TCA) solution. The homogenate was filtered through Whatman No.1. filter paper, after which 0.5ml of 100mM K-phosphate buffer (pH 6.8) and 2ml reagent (1M KI in fresh bidistilled water H$_2$O) were added to 0.5ml of the leaf extract filtrate. The blank probe consisted of 0.1% TCA in the absence of leaf extract. The reaction was developed for 1h in darkness and the absorbance was measured at 390nm. The amount of H$_2$O$_2$ was calculated using a standard curve prepared with known concentrations of H$_2$O$_2$. The H$_2$O$_2$ content was expressed as µg/g fresh weight.

**Determination of malondialdehyde (MDA) content:** MDA determination (Lipid peroxidation product) was applied according to the method described by Li (2000). 0.2g fresh weight was homogenized in 1.5ml 5% TCA. The homogenates were centrifuged at 13000g for 20min. A reaction mixture of the supernatant (0.5ml) and 1ml 20% TCA and 1ml 0.5% TBA (Thiobarbituric acid) was incubated at 95°C in a water bath for 25min, then cooled immediately before centrifugation. A bsorbance of the supernatants was determined at 450, 532 and 600nm, respectively. Calculation of MDA was based on the following formula:

$$\text{MDA (µm/ml)} = 6.45(A532 - A600) - 0.56 A450$$

**Determination of antioxidant enzymes activity**

**Total cellular enzyme extraction:** Germinated wheat seedlings (5g) were homogenized in 1.5ml extraction buffer (a potassium phosphate buffer 0.1M, pH 7 in ice cold condition). The homogenates were centrifuged at 5000rpm for 10min at 4°C and the supernatant was used as enzymes source (Ma et al., 2012).

**Antioxidant enzymes assay**

**Assay of superoxide dismutase (SOD):** SOD activity was measured by the nitro blue tetrazolium (NBT) reduction method (Beyer & Fridovich, 1987). Test tubes containing reaction solution with 3ml of assay buffer, 60µl of crude enzyme and 30µl of riboflavin were illuminated for 7min in an aluminum foil lined box containing two Fluorescent lamps at 25°C. The absorbance of the blank solution and reaction solution was measured with a spectrophotometer at 560nm. SOD activities were calculated as a following equation:

$$\text{SOD activity (%) } = \left( 1 - \frac{A}{B} \right) \times 100$$

where (A) is the absorbance of sample and (B) is the absorbance of blank.

**Assay of polyphenol oxidase activity (PPO):** It was done according to Beyer & Fridovich (1987). Five ml of assay mixture comprising 125µM of phosphate buffer (pH 6.8), 100µM of pyrogallol and 1ml of crude extract were prepared. After incubation at 25°C for 5min, the reaction was stopped by the addition of 1ml 10% (v/v) H$_2$SO$_4$. The optical density of the produced color was measured spectrophotometrically at 430nm and the enzyme activity was expressed as the change in the optical density/mg protein/min.

**Assay of peroxidase activity (POX):** It was done according to Racusen & Foote (1965). Five ml of the assay mixture comprising 300µM of phosphate buffer (pH 6.8), 50µM catechol, 50µM H$_2$O$_2$, and 1ml of crude enzyme extract were prepared. After incubation at 25°C for 5min, the reaction was stopped by the addition of 1ml 10% (v/v) H$_2$SO$_4$. The optical density of the produced color was measured spectrophotometrically at 430nm and the
enzyme activity was expressed as the change in the optical density/mg protein/min.

**Determination of hydrolysis processes during germination**

*Amylase enzyme assay:* The amount of starch hydrolyzed by the action of amylases was measured according to Johnson (2007). The amylase activity was expressed as the amount of starch hydrolyzed \( \text{min}^{-1} \text{mg protein}^{-1} \).

**Determination of the total soluble carbohydrate content**

Carbohydrates were estimated using phenol sulfuric acid method (Dubois et al., 1956).

**Acid and alkaline phosphatase assays**

The freshly germinated wheat seedlings (5g) were used for enzyme extractions and assays. The activities of all the two enzymes were assayed according to the method of Tominaga & Takeshi (1974). The specific activity was expressed as nkat \( \text{mg}^{-1} \) protein, where one nkat of enzyme activity is defined as one nmol p-nitrophenol liberated \( \text{min}^{-1} \).

**Determination of the total phosphorus content**

The dried powder of germinated wheat seedlings was digested in a mixture of concentrated nitric acid, sulphuric acid and perchloric acid at the ratios 10: 1: 4, respectively. The volume was made up to a constant volume with distilled water according to the method of Chapman & Pratt (1978) with certain modifications. Phosphorus content in the digested samples was determined colorimetrically by ascorbic acid method described by Murphy & Riley (1958). Results were expressed as mg/g dry weight of yielded grains.

**Statistical analysis**

Data was presented as the mean of three replications and standard error (SE) bar. Statistical analysis was carried out using ANOVA test and the means were compared by the least significant difference (LSD) test at a significance level of \( \text{P} \leq 0.05 \).

**Results**

**Definsine separation and characterization**

The purification and separation of definsine-like protein from the fenugreek seeds by SDS-PAGE revealed the appearance of one protein band has molecular weight 10.7kDa as indicated in Fig. 1.

![Fig. 1. Protein profile of the bulk fenugreek seeds extract and the purified part as revealed by SDS-polyacrylamide gel (Lane A= Bulk extract, lane B= The purified extract and M= marker).](image-url)

**Effect on germination**

Germination rate of grains treated with the lowest Zn concentration (1mM) was unaffected when compared to the control. In contrast, germination was severely inhibited at the high Zn concentrations (10, 50 and 100mM) as shown in Table 1. Also, shoot and root lengths of the wheat seedlings were significantly decreased at these high Zn concentrations, compared to the control. The inhibition in the germination rate was approximately 18, 36 and 49%, respectively under these Zn concentrations. Upon application of definsine priming method, the germination inhibition rate was decreased with the three concentrations of Zn to approximately 4.0, 13 and 28%, respectively. This indicates that definsine priming method could alleviate the toxic effect of Zn on the germination rate.

The data also revealed that, root length of the germinated seedling decreased significantly under the high Zn concentrations (10, 50 and 100mM) in a percent of 24.41%, 78.71% and 89.1%, respectively. However, such inhibition was gradually decreased by using definsine priming method recording a percent of 19.5, 21.7, 33.1%, respectively. Similarly, definsine priming method alleviated the toxic effect of Zn concentrations on the shoot germination. Under the high Zn concentrations, the inhibition percentages of shoot lengthswere 52.7, 74.4 and 75.8%, respectively. Definsine priming led to a reduction in such inhibition percentages of shoot length recording inhibition rates of 12.6, 23.1 and 25.1%, respectively.
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Effect on antioxidant machinery

In general, the results of this study showed an increase in antioxidant machinery under all concentrations of Zn in the defensine primed and non-primed germinated grains.

\( \text{H}_2\text{O}_2 \) as a reactive oxygen species increased significantly under the high concentrations of Zn. It increased with the percentages of about 672.5%, 710.8% and 1008.22%, respectively after 24h from germination and this increase was time-dependent. These percentages of increase were found to be 926.35%, 1116.7% and 1428.45%, respectively after 72h from germination, while in the case of defensine priming, the \( \text{H}_2\text{O}_2 \) content increased under the high concentrations of Zn but in a smaller ratio than the non-primed cases. The increase ratios after 24h were 145.5%, 202.16% and 203.89%, respectively, but this increase ratios decreased with time, where they were decreased to 25.5%, 43.096% and 46.44%, respectively after 72h (Fig. 2).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Treatments</th>
<th>Non-primed Grains</th>
<th>Defensine Primed Grains</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
<td>1</td>
<td>10</td>
</tr>
<tr>
<td>Total germination ratio %</td>
<td>97</td>
<td>98</td>
<td>82.*</td>
</tr>
<tr>
<td>Root length</td>
<td>5.12</td>
<td>5.1</td>
<td>3.87*</td>
</tr>
<tr>
<td>coleoptile length</td>
<td>3.98</td>
<td>3.95</td>
<td>1.88*</td>
</tr>
</tbody>
</table>

-Values are means of 5 replicates.
-Means followed by asterisks are significantly different from the control according to paired-samples t test.

Fig. 2. Effect of wheat grains priming with defensine on \( \text{H}_2\text{O}_2 \) and malonyl dialdehyde (MDA) at 24, 48 and 72h from germination under different concentrations of Zn. (Values are means of 5 replicates. Pars with asterisks are significantly different from the control according to ANOVA test).
Results presented in Fig. 2 showed that, the MDA increased significantly in the non-primed germinated grains under high concentrations of Zn (10, 50 and 100mM). The ratios recorded increased to 290.38, 453.8 and 496.1%, respectively, while the increase ratios in case of the defensine primed germinated grains were 30.7, 34.6 and 38.46%, respectively.

The defensine priming method enhanced the activity of the antioxidant enzymes including superoxide dismutase (SOD), polyphenol oxidase (PPO) and peroxidase (POX) under all Zn concentrations. The activity of such antioxidant enzymes increased by time in the defensine primed grains, while with the non-primed germinated grains the activity increased with smaller ratio and decreased by time (Fig. 3).

Fig. 3: Effect of wheat grains priming with defensine on antioxidant enzymes (Super oxide Dismutase (SOD), polyphenol oxidase (PPO) and peroxidase (POX)) at 24, 48 and 72h from germination under different concentrations of Zn. (Values are means of 5 replicates. Pars with asterisks are significantly different from the control according to ANOVA test).

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Effect on hydrolysis process

Priming with definsine significantly enhanced the amylase activity under normal conditions as the activity increased about 16% than that recorded in non-primed germinated grains. The amylase activity at the low Zn concentration (1mM) decreased insignificantly either in definsine primed or non-primed germinated grains. However, the high Zn concentrations (10, 50 and 100mM) caused significant decrease in amylase activity in both primed and non-primed germinated grains. The decrease ratios in the non-primed grains were 53.2%, 66.6% and 83.6%, respectively; while, the amylase activity in the defensine primed germinated grains was significantly decreased when compared to that of the non-primed germinated grains where the recorded ratios were 11.5%, 21.5% and 29.0%, respectively (Fig. 4).

![Fig. 4](image_url)

Fig. 4. Effect of wheat grains priming with definsine on amylase activity and carbohydrates content at 24, 48 and 72h from germination under different Zn concentrations. (Values are means of 5 replicates. Pars with asterisks are significantly different from the control according to paired-samples t test).
As shown in Fig. 4, priming with definsine enhanced the soluble carbohydrates content either in normal germination conditions or under the influence of Zn concentrations. In response to Zn concentrations, there was insignificant decrease in the carbohydrate content. However, such decrease was significant in the non-primed germinated grains at high Zn concentrations recording ratios of 56%, 58.2% and 67.3%, respectively.

Data shown in Fig. 5 indicated that phosphatases activity increased significantly in the defensine primed germinated grains. There was a non-significant decrease in the acid and alkaline phosphatases at high Zn concentrations (10, 50 and 100mM). Such decrease was in a small ratio of about 12.05% and 18.4%, respectively in the acid phosphatase and 9.7% and 15.6%, respectively in the alkaline phosphatase. However, the phosphatases activity decreased significantly in the non-primed germinated grains recording a ratio of decrease about 61.9% in acid phosphatase and about 57.8% in the alkaline phosphatase (Fig. 5).

![Graph showing effect of wheat grains priming with definsine on acid and alkaline phosphatase and phosphorus content at 24, 48 and 72h from germination under different Zn concentrations. Values are means of 5 replicates. Bars with asterisks are significantly different from the control according to paired-samples t test.](image-url)
Phosphorus content is related to the activity of the phosphatases. Defensine priming enhanced the soluble phosphate content under normal conditions and Zn concentrations. However, there was a significant decrease in the phosphorous content in the non-primed germinated grains under high Zn concentrations (10, 50 and 100mM) with decrease ratios of about 64.4%, 66.26% and 73.47%, respectively as shown in Fig. 5.

Discussion

The present study was performed to examine the mechanisms of the beneficial effect of defensine priming on the germinated wheat grains exposed to different Zn concentrations in a trial to alleviate the Zn phytotoxicity. The life cycle of higher plants begins with seed germination, which depends on perceptions of different environmental stimuli. Seed germination could be affected negatively or positively by many factors including dormancy, embryonic inadequacy, germination inhibitors, water, temperature, gases and light as well as organic and inorganic chemicals, such as zinc, which naturally exist in earth’s crust and atmosphere (Srivastava, 2002).

Data of the present study showed that the low concentration of Zn (1mM) showed no significant harmful effect on the germination process of wheat plant. This result corroborates with that of Kösesakal & Ünal (2012) who reported that the low concentrations of ZnCl₂ (1mM) cannot reduce the tomato seeds germination but increases the germination ratio. Such result is in correlation to the fact that Zn at low concentrations being as one of the most essential micronutrients playing a significant role in many vital metabolic processes. For instance, Zn is a cofactor for several enzymes, such as anhydrases, dehydrogenases, oxidases and peroxidases (Aravind & Prasad, 2003). However, in the present results high Zn concentrations (10, 50 and 100mM) markedly decreased the germination process in the non-primed germinated wheat grains, such decrease is ensured as the germination process significantly delayed in the non-primed germinated grains, this decrease can be attributed to the toxic oxidative stress caused by the Zn which significantly appeared in the increase of the oxidative stress markers contents (H₂O₂ and MDA) under Zn treatment and their content increased with time. These results are in accordance with those of Molnárová & Fargašová (2016) who stated that Zn phytotoxicity has come recently into greater focus because it is a part of the long term utilization of fertilizers. Next to this source also industry increased its content in the surface soil. High Zn concentrations create cytotoxic effect on plant growth and metabolism (Arif et al., 2016). Similar results were obtained by Liu et al. (2016) upon germination of Solanum nigrum in the presence of high concentrations of Zn. Also, Munzuroglu & Geckil (2002) reported that with the increase in Zn concentration, the germination ratio, coleoptile and hypocotyl length of wheat grains decreased. Excess Zn causes harmful effects on plants; it inhibits plant growth and root development. Moreover, high Zn levels could cause cellular oxidative damage and membrane lipid peroxidation in plant cells and affect the activity of many antioxidative enzymes and antioxidant contents in plants (Cherif et al., 2011). However, in the present study the defensine priming enhanced the germination process under Zn-stressed conditions. Similar studies indicated that seed priming has been found a double technology to enhance rapid and uniform emergence and to achieve high vigor and better yields in vegetables and floriculture and some field crops (Lemrasky & Hosseini, 2012). In addition to better establishment, farmers reported that primed crops grew more vigorously, flowered earlier and yielded higher (Farooq et al., 2008).

Under non-priming conditions, Zn stress causes excess production of ROS which does not affect the antioxidant enzymes only but also negatively affects the hydrolysis enzymes, as recorded in this study, where the amylase and phosphatases activity decreased as a result of high concentrations of Zn treatment. This decrease was spontaneously accompanied with decrease in soluble carbohydrates and soluble phosphate contents and so, the germination process was delayed. Similar result was reported by Laware & Raskar (2014) who showed that upon germination of onion seeds treated with titanium dioxide, the hydrolysis enzymes activity decreased as a result of excess production of the ROS.

The constructive role of the ROS to the enzymes attributed to their ability to cause damage to biomolecules such as lipids, proteins and DNA. These reactions can alter intrinsic membrane properties like fluidity, ion transport, loss of enzyme activity, protein cross-linking, inhibition of protein synthesis, DNA damage, and ultimately resulting in cell death (Sharma et
al., 2012). However, the present results showed 
that seed priming using defensine enhanced the 
germination process under high concentrations of 
Zn, this could be attributed to the enhancement 
in the activity of antioxidant enzymes which in 
turn regulated the ROS production represented 
in the excess of H$_2$O$_2$ at the beginning of the 
germination which plays a messenger role to 
activate the defense mechanism (Sharma et al., 
2012). Zeng et al. (2011) showed that regulation 
of antioxidative enzymes and control of reactive 
y oxygen species help heavy metal resistance in 
the hyper accumulator plants.

The current study showed a reduction in the 
production of MDA and H$_2$O$_2$ (oxidative stress 
markers) as a result of defensine priming of wheat 
grains, these results are generally in agreement 
with those of Anwaar et al. (2015), who found 
a decrease in the MDA and H$_2$O$_2$ and electrolyte 
leakage in the Zn tolerant cotton plants suggesting 
that protection of cotton plants from Zn toxicity 
by fighting the induced oxidative damage. Also, 
Garg & Kaur (2012) reported that Zn toxicity 
can be countered by alleviating oxidative stress 
through up regulation of antioxidant enzymes. 
Defensine priming of wheat grains in this study 
resulted in a regulation of antioxidant enzymes 
and in controlling the excess production of the 
ROS which protected the hydrolysis enzymes 
destruction. Additionally, the activities of 
amylase and phosphatases were insignificantly 
affected by high concentrations of Zn and also in 
addition to this activity the contents of the soluble 
carbohydrates and phosphate were not affected. 
This regulation in hydrolysis process also has 
a pronounced effect on gibberellin signaling, 
inducing a change in hormonal balance that results 
in germination enhancement (Bahin et al., 2011).

The protective role of the defensine like 
protein against Zn toxicity has also been studied 
by Mirouze et al. (2006) who stated that defensins 
are inducible by Zn. It seems that these defensins 
play a role in the mechanism of metal tolerance, 
specifically Zn mechanism and they suggest that 
defensins confer Zn tolerance in plant, like the 
metal transporters or chelators, defensines found 
to be involved in metal tolerance and also exist 
in non-tolerant plants. Two main hypotheses 
may be proposed to understand the mechanism 
underlying the role of defensins in plant metal 
physiology. The first is a Zn chelation hypothesis, 
like the metallothioneins, defensins contain 
cysteine-rich motifs that are potentially involved 
in metal binding (Cobett & Goldsbrough, 2002). 
The second hypothesis relies on the structure 
similarity between plant defensins and some 
channel-blocker peptides such as scorpion b– 
toxins (del Rio-Portilla et al., 2004 and Zhu et 
al., 2005), which were shown to interfere with 
divalent cation channels (Carlier et al., 2000 and 
Castle et al., 2003). A significant amount of work 
is still necessary to detail the mechanism of action 
of defensins with respect to Zn tolerance.

Conclusions

Results of the present study can ensure that the 
adverse effects caused by Zn toxicity in the 
germinated wheat grains could be alleviated by 
defensine priming method which increased the 
Zn tolerance in wheat germinated grains through 
alleviating the oxidative stress by up regulation of 
the antioxidant enzymes. This could be unforeseen 
role for defensins which opens up new horizons 
for the investigation of defensive mechanisms of 
action.

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التخفيض من سمية الزنك في حبوب القمح النامية بواسطة (Triticum aestivum L.) معاملة البذور بالديفينسين

إيمان السيد سليم و ديالا محمد نجيب
قسم النبات - كلية العلوم - جامعة الزقازيق - الزقازيق - مصر.

تهدف هذه الدراسة لإختبار تأثير طريقة عمر حبوب نبات القمح قبل الزراعة بشبيه البروتين الدفينسين لتحسين مقاومة نبات القمح نسبته عصر الزنك. تم تطبيق التجربة على مجموعتين من حبوب نبات القمح المعرضة لإجهاد تركيزات مختلفة من الزنك، المجموعة الأولى تحتوي على الحبوب الغير مغمورة في شبيه البروتين الدفينسين والثانية المغمورة به. تم دراسة النمو و بعض العوامل الفسيولوجية لحبوب القمح النامية. وكشف النتائج أن زيادة تركيزات الزنك في الحبوب الغير مغمورة بشبيه البروتين الدفينسين أخفضت نسبتهات النمو. كما أن تناقص في الكربوهيدرات القابلة للذوبان وأسلاك النبات. بالإضافة إلى ذلك، كان هناك تأثير كبير في تخفيض الإنتاج من حبوب القمح النامية بالتمييز بالديفينسين. وتم التأكد من هذه النتائج من خلال دراسة الدفعات في الخلايا نحو الحمض. وتم تسجيل نشاط الإنزيمات المضادة للأكسدة مثل سوبر أكسيدديسموتاز. والبيروكسيديز بالنمو في الحبوب المغمورة بالديفينسين. وتم تسجيل تحسن في نوعية الأكسدة في الحبوب المغمورة بالديفينسين.

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