



## Induction of arsenic stress tolerance in *Capsicum annum* through *Trichoderma harzianum* sand mix method

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### Abstract

The research was performed at Nusrat Jahan College Rabwah Pakistan to overcome arsenic stress in two varieties "Sanam and Ghotki" of *Capsicum annum* through *Trichoderma harzianum* sand mix method. Seeds of mentioned chili varieties and *Trichoderma* fungus were taken from NARC Pakistan. Cups of sand were mixed with *Trichoderma harzianum* at the rate of  $2 \times 10^7$  CFU. Chili seeds after surface sterilization through mercuric chloride were sown in cups of fungal treated sand. Arsenic oxide (AsO 1: 1mg/L and AsO 2: 2mg/L) stress was applied after one week of sowing. Seedlings were harvested after 30 days of sowing and were preserved in 50mM potassium phosphate buffer. Roots, shoots and leaves were separately preserved. The preserved samples were subjected to different biochemical tests. This study has revealed *Trichoderma* sand mix method very effective method to eliminate Arsenic oxide stress by generating ROS damaging proteins.

**Key words:** Green Chillies, Arsenic, *Trichoderma harzianum*, Seedlings, Sand Mix.

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### 1. Introduction

Arsenic is one of the most toxic heavy metal. It mostly exists in conjugation with various minerals also it is present in pure crystalline forms. Arsenic pollution is result of many different types of anthropogenic activities as well as natural phenomena. Increasing arsenic pollution is constantly contaminating water supplies and soils biochemistry. As a result, crops like rice, wheat, beans, turmeric, chillies, onions, carrots, reddish and many more have been reported to reserve arsenic contents in their edible regions [1]. Large concentrations of arsenic in any plant can hinder normal growth mechanism, seed germination also ceases, and heights of plants also get retarded. Rate of flowering and fruiting as well as number of grains also reduce [2]. Arsenic present in plants disturb various metabolic activities in cells such as associations with sulphohydryl groups and additions, deletions of phosphate groups from ATP. Arsenic develops free radicals and ROS species that cause oxidative damages [3]. In order to overcome such adverse effects of arsenic stress healthy preventive or treating measures are being working out on

many scientific platforms but among all measures, biological means of controlling arsenic pollution are worth appraising as in biological treatments there are no threats of chemical contaminations or side effects. *Trichoderma harzianum* is soil inhabitant ascomycete fungi, that forms dirty green colored colonies. Many researches have shown that this fungi act as growth promoting agent in different plants [4]. Many species of these fungi are known for eliminating different biotic as well as abiotic diseases of plants. Capsicums are one of the major components of humans' routine diet. These provide flavor to food also many vitamins and phenolic compounds that work as antioxidants. Concentrations of antioxidants are dependent on multiple factors such as age, growing scenarios and variations in genotypes and varieties [5].

Our research was planned, considering edible requirements of chillies and these being influenced by arsenic contamination to overcome this contamination using *Trichoderma harzianum* sand mix method.

## 2. Materials and Methods

Botany department, Nusrat Jahan College Rabwah, Pakistan was site of experiment. The experimental design was to induce stress tolerance in chillies of two varieties “SANAM” and “GHOTKI” taken from NARC PAKISTAN against arsenic oxide (1mg/L, 2mg/L) through *Trichoderma harzianum* sand mix method. This fungus was also taken from NARC. Cups of sand were mixed with *Trichoderma harzianum* at the rate of  $2 \times 10^7$  CFU. Chili seeds of both varieties after surface sterilization through mercuric chloride were sown in cups of fungal treated sand. Arsenic oxide (AsO 1: 1mg/L and AsO 2: 2mg/L) stress was applied after one week of sowing. Seedlings were harvested after 30 days of sowing and were preserved in 50mM potassium phosphate buffer. Roots, shoots and leaves were separately grinded and then centrifuged at 14000rpm for 15 minutes. After centrifugation supernatants were subjected to different biochemical tests.

### 2.1 Total Soluble Proteins

Concentration of total soluble proteins was estimated using the method of (Bradford, 1976) [6] with few amendments. The 1ml supernatant was reacted with 2ml Bradford Reagent and incubated for 15-20 min then reading was measured at 595 nm.

### 2.2 Ascorbate Peroxidase Activity (APX)

The APX working was measured using the method of Asada and Takahashi (1987) [7]. The reaction solution (1600 $\mu$ l) was comprised of 50 mM potassium phosphate buffer (pH 7.0), 0.5 mM ascorbic acid, 0.1 mM H<sub>2</sub>O<sub>2</sub> and 400  $\mu$ l of enzyme extract. The absorbance was taken at 290 nm against the blank and the enzyme activity was represented in U $\text{mg}^{-1}$  protein (U=change in 0.1 absorbance min<sup>-1</sup> mg<sup>-1</sup> protein).

### 2.3 Total Phenolic Contents

Total phenolics were evaluated with the help of Folin-Ciocalteu protocol (Wolfe *et al.* 2003) [8] with few amendments. Samples were mixed with 5 ml Folin-Ciocalteu reagent (previously diluted with water 1:10 v/v) and 4 ml (75 g/l) of sodium carbonate. The tubes were shaken for fifteen seconds and were permitted to stand for 30 min at 40°C so that the color develops. Then absorbance was taken at 765 nm on spectrophotometer.

### 2.4 MDA Contents

Malondialdehyde (MDA) was determined in accordance to method proposed by Dhindsa *et al.* (1981) [9]. In the 2 ml TCA, added 2 ml of 0.6% thiobarbituric acid. It was heated at 100 degree centigrade for 20 minutes in water bath. After heating immediately cooled for 20 minutes and then centrifuged at 10000 rpm for 10 minutes. The resulting color was taken at 532 nm on spectrophotometer.

### 2.5 Hydrogen Peroxide Concentration

H<sub>2</sub>O<sub>2</sub> concentration was determined according to the protocol of (Velikova *et al.*, 2000) [10]. The 0.1 ml of supernatant were added to 0.1 ml of 10 Mm potassium

phosphate buffer (PH 7.0) and 1M IKI. The absorbance was taken 390nm.

## 3. Results

### 3.1 Total Soluble Proteins

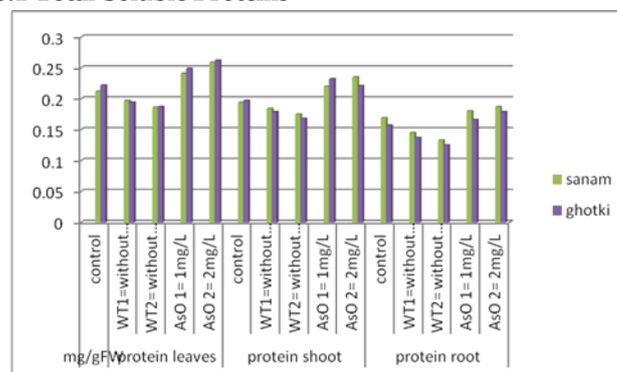


Figure 1: Mean values of protein concentrations of chillies at different treatments

According to Figure 1. Protein concentration enhanced many fold in both varieties of chillies in leaves, shoots and roots at 1mg/L and 2mg/L of arsenic oxide stress. This increase in protein amount is because of the fact that *Trichoderma* application has helped chillies in overcoming both stress levels of arsenic oxide. While (WT= without *Trichoderma*) group which is without *Trichoderma* treatment could not cope up with arsenic stress and as a result protein content decreased in them as compared to control.

### 3.2 Ascorbate Peroxidase Activity (APX)

Results of Figure 2 Show that in all parts of both sanam and ghotki chillies of *Trichoderma* APX activity was increased at both levels of arsenic stress as compared to control group on the other hand WT set showed decline in APX activity because in this group *Trichoderma* was not mixed in sand thus this group was unable to raise APX activity to reduce stress levels.

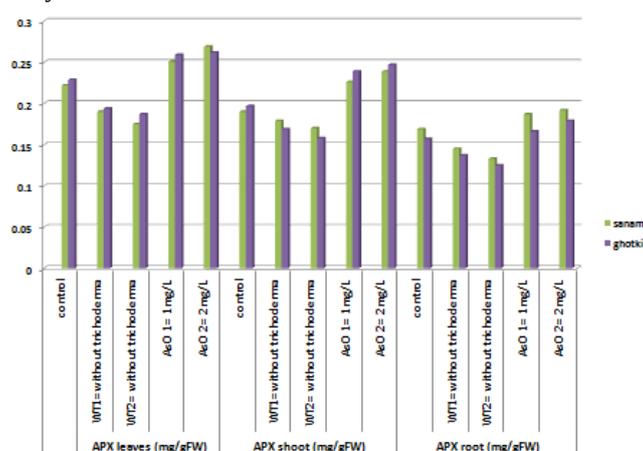


Figure 2: Mean values of APX activity of chillies at different treatments.

### 3.3 Total Phenolic Contents

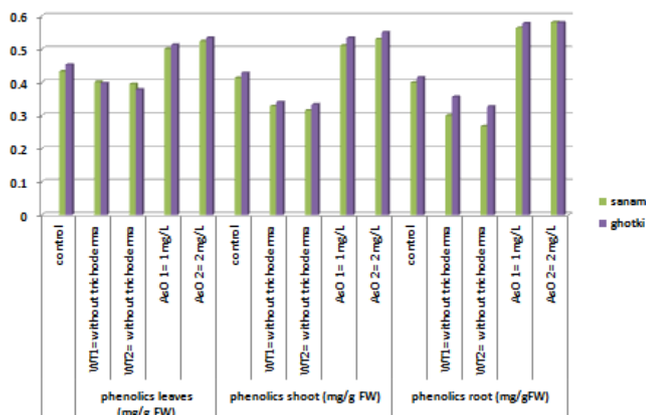


Figure 3: Mean values of total phenolics content of chillies at different treatments

At both stress levels of arsenic *Trichoderma* sand mixed group showed increase in phenolics content as compared to WT group and control group. All parts, roots, leaves and shoots of sand mixed group showed enhancement in phenolics content.

### 3.4 MDA Contents

Table 1: represent mean values of MDA content in roots, shoots and leaves of green chillies

Mean values of MDA in roots, shoots and leaves of green chillies		
Unit:umol/gFW	SANAM	GHOTKI
Control Leaf	0.139	0.143
WT1=without trichoderma	0.145	0.156
WT2= without Trichoderma	0.152	0.162
AsO 1= 1mg/L	0.126	0.132
AsO 2= 2mg/L	0.118	0.126
Control Shoot	0.132	0.127
WT1=without trichoderma	0.142	0.167
WT2= without trichoderma	0.165	0.172
AsO 1= 1mg/L	0.125	0.124
AsO 2= 2mg/L	0.119	0.106
Control Root	0.104	0.121
WT1=without trichoderma	0.139	0.147
WT2= without trichoderma	0.147	0.159
AsO 1= 1mg/L	0.109	0.112
AsO 2= 2mg/L	0.096	0.103

According to Table 1 *Trichoderma* sand mixed group of chillies showed decline in MDA content as compared to non-treated group in all parts of seedlings of both varieties. Proving that this Fungi application prevents chillies from oxidative damage.

### 3.5 Hydrogen Peroxide Concentration

Table 2: Represent mean values of H<sub>2</sub>O<sub>2</sub> concentration in roots, shoots and leaves of green chillies

Mean values of hydrogen peroxide in leaves, shoots and roots of chillies		
Unit:umol/gFW	SANAM	GHOTKI
Control Leaf	0.121	0.104
WT1=without Trichoderma	0.147	0.139
WT2= without Trichoderma	0.159	0.147
AsO 1= 1mg/L	0.112	0.109

AsO 2= 2mg/L	0.103	0.096
Control Shoot	0.127	0.132
WT1=without Trichoderma	0.167	0.142
WT2= without Trichoderma	0.172	0.165
AsO 1= 1mg/L	0.124	0.125
AsO 2= 2mg/L	0.106	0.119
Control Root	0.143	0.139
WT1=without trichoderma	0.156	0.145
WT2= without trichoderma	0.162	0.152
AsO 1= 1mg/L	0.132	0.126
AsO 2= 2mg/L	0.126	0.118

According to Table 2. Hydrogen peroxide concentration was reduced at both levels of stress in *Trichoderma* sand mix group as compared to WT group in all parts of seedlings of both varieties of chillies.

### 4. Discussions

Output of this study reveals that *Trichoderma harzianum* fungi is resistant to arsenic oxide contamination at levels of 1mg/L and 2mg/L. Being itself resistant to this heavy metal contamination, this fungi has effectively combated arsenic oxide stress in both Ghotki and Sanam varieties of green chillies. *Trichoderma harzianum* sand mix method has proven to be very positive and effective biological treatment of preventing green chillies and other plants from harms of arsenic oxide. Increase in protein content of *Trichoderma* sand mix chillies even under stress of arsenic and reduction in protein concentrations of chillies without *Trichoderma* clearly indicate that this fungi has triggered auxins and other hormones which promote healthy growth and biomass of chillies along with developing tolerance against arsenic stress. *Trichoderma harzianum* has activated defense mechanism against arsenic stress by raising levels of phenolic compounds and peroxidases as is evident by results of total phenolics content and ascorbate peroxidase activity. These phenolic compounds and peroxidases act as ROS scavenging substances because they trigger different ROS deteriorating proteins such as glutathione – S-transferase, etc. Thus due to activation of ROS damaging species MDA content and hydrogen peroxide concentration is reduced as a result all *Trichoderma* treated chillies have been prevented from oxidative damages. In conclusion it is portrayed from results of this research activity that for eradicating arsenic harms from plants *Trichoderma harzianum* sand mix method is very effective safe and cheap method.

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