<b>Original Research Article</b>
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# Priming Induction in Neighbouring Plants of Gossypium hirsutum under Salt Stress

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# 5 ABSTRACT

6 Plants are subjected to various types of environmental stresses throughout their lifecycle. It has been 7 found that plants are able to communicate with the neighbouring plants under stress conditions 8 through volatile organic compounds. These volatiles act as signals for the neighbouring plants thus 9 preparing them for the upcoming stress, a phenomenon known as priming. So, the present study 10 explores the effects of salt stress on cotton plants and the resultant induction of priming in the nearby 11 plants. For this purpose, salt tolerant cotton (Gossypium hirsutum) variety was used. Two 12 concentration levels, 100 mM, and 150 mM (what) were used to study the impacts of the stress. The 13 experiment was divided into two steps for each treatment. In the first step, a set of plants (emitters) 14 was given salt stress. A second set of plants (receivers) was placed adjacent to the stressed plants 15 (emitters), while the third set of plants was placed separately as a control for both the treatments. 16 Various physiological and morphological parameters were measured at the beginning and the end of 17 the first step. In the second step, the receiver plants now termed as "primed" were given same levels 18 of stress while a new set of non-primed plants was placed near the primed plants. These non-primed 19 plants were now treated with 100mM and 150mM of NaCl respectively and the results were 20 compared. The results show that plants were able to get signals from neighbouring stressed plants. 21 Plants responded by altering morphology and physiology to prepare themselves for future stress 22 conditions.

#### 23 1. INTRODUCTION

Communication among plants has been an area of interest for quite a long time now. This communication is done by releasing and sensing volatile organic compounds. These compounds are released by the plants under normal circumstances [1] and are found in almost all plant tissues including roots, stem, leaves and flower tissues.

However, under conditions of stress, the level and composition of these volatile organic compounds has been reported to fluctuate [2]. Several studies suggest that plants are also able to receive signals from neighbouring plants in the form of these VOCs. Plants can thus detect any change in the concentration of such signals, from a neighbouring plant, caused by biotic or abiotic stress [3, 5]. Interaction among plants through roots via common mycorrhizal networks (CMNs) has already been reported in several studies

The present study focuses on plant to plant communication as a result of VOC emission through the leaves. VOCs released by leaf tissues are called "Green Leaf Volatiles" or GLVs [6, 7]. Alterations in the levels of GLVs can induce defensive responses in the form of physiological and morphological changes in the unstressed neighbours [8]. Green leaf volatiles mainly consist of alcohols, aldehydes, ketones, monoterpenes and esters [9]. Several studies show the effects of biotic stresses on GLVs concentration such as viral infection [10, 12].

However, there is not much work done over the effects of abiotic stress on the levels of VOCs and the resultant induction of resistance (priming) in the receiver plants. Salinity is an abiotic stress and is associated with several changes in the treated plants at both morphological as well as physiological level. Changes in the levels of isoprene and other volatile organic compounds have been associated with salinity stress. So the following experiment was conducted in order to study effects of salinity stress on both treated and untreated cotton plants as well as the induction of priming in the untreated neighbouring plants.

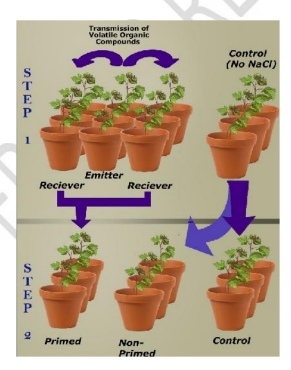
#### 47 2. MATERIAL AND METHODS

The experiment was carried out at Cotton Research Station, Ayub Agricultural Research Institute Faisalabad. Fuzzy seeds (age) of cotton (*Gossypium hirsutum*) variety FH-142 were soaked under water overnight. These seeds were then sown in polythene bags containing a mixture of compost and alluvial soil (mixture of sand silt and clay) for good germination. 210 bags were filled and 4-5 seeds (uniform treatment) were sown in each bag on 02 March, 2018 and placed it in green house.

53 Transplantation was done after three weeks from polythene bags to plastic pots having one feet 54 height. These pots were filled with soil, alluvial soil and FYM mixture in 4:4:1 proportion. Selected 150 55 pots and place them under the tunnel and watered it frequently.

56 The experiment was arranged into three plots for treatment 1, 2 and control respectively and these plots were partitioned by polythene paper. Total 36 plants were placed in 1<sup>st</sup> and 2<sup>nd</sup> plot each, while 57 third plot contained 72 plants. In 1<sup>st</sup> and 2<sup>nd</sup> plot, arrange the plants in three rows and each row 58 59 contain 12 plants. The mid row called emitter while other two rows called receiver, we named it 60 because according to hypothesis, emitter release volatile organic compounds when we gave any 61 stress while receiver which is placed at the periphery of emitter receive those compounds (Fig. 1). After two weeks of transplantation a salt stress was given to emitters only. Two levels of NaCl 62 concentration were used, 100mM and 150mM respectively. The emitters placed in the 1<sup>st</sup> plot was 63 given 100 mM solution while other emitters placed in 2<sup>nd</sup> plot was given 150 mM solution while rest of 64 65 the plants were on tap water. 100 ml salt solution to emitters and 100 ml tap water to rest of the plants 66 were given.

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Fig.1 shows the mechanism of experiment

70 Since the whole experiment was carried out in two steps. In step one a group of plants called 71 receivers placed in the proximity of salt stressed plants called emitter for twenty days. Another group 72 of plant called control placed apart from emitter and receiver. In second step, receiver plants were 73 taken from step one (Fig. 1). Now it is called primed plants, gave a salt stress to primed and non-74 primed plants (taken from the control of step 1) and compare it with control. This whole experiment 75 was repeated under different dose of salt concentration.

The parameters that studied are Leaf area, Total dry weight, Total Fresh weight, Relative growth rate,
 Proline content, No. of leaves per plant, Shoot length, Root length, Relative growth rate, H<sub>2</sub>O<sub>2</sub>
 species, Dry shoot weight, Dry root weight, Moisture content, Root to shoot ratio, Leaf Area.

79 The plant material with sample size of six were taken at the beginning of experiment (day 0), at the 80 end of step 1 just after 20 days of experiment started and at the end of step 2, after 20 days of step 1 81 finished. Plants were uprooted carefully, gently washing away the soil and take fresh weight of each 82 sample by using weighing balance. Different morphological trait that are mentioned above was then 83 analyse. Leaf area was measure using leaf area meter. Plant material was then oven dried at 70 °C 84 for 48 hours. Relative Growth rate was measured by using total dry weight according to Pérez [13] 85 method using the formula RGR =  $(\ln W_2 - \ln W_1)/(t_2 - t_1)$ , where W is a total dry weight and t is a time 86 between two reference period which is 20 days.

Proline content was determined by Bates [14] method with little modification. Take 250 g sample of ground leaf and add 10ml of 3% solution of salphosalicyclic acid, then centrifuged it under 3000rpm for 10 min, transfer 2ml of supernatant to new test tube after centrifugation. Add 2ml of 6M ortho phosphoric acid, 2ml acid ninhydrin and 2ml glacial acetic acid. Keep it in water bath for one hour at 100°C. After it add 4ml of toluene, shake it well and wait for few minutes. Separate the upper layer of solution and observe it in spectrophotometer under 520nm. The reading was compared with standards. H<sub>2</sub>O<sub>2</sub> analysis was determined by Orozco-Cardenas and Ryan [15] method.

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# 97 **4. RESULTS**

# 98 4.1 PROLINE CONTENT

99 The experiment was organized in to three treatments with two doses of NaCl concentration and one is 100 control. As we see in Table 1, the emitter and receiver of treatment 1 is non-significant while both are 101 significant with control. It means that the Emitter which is under salt stress produce more proline than 102 control and it induces a receiver plant to produce proline. When we move this receiver to step 2 as 103 shown in fig. 3, it become primed and shows a significant result with control as well as Non-primed 104 but it seems that when plant grow the rate of proline content in primed plant reduces and it produce 105 less amount of proline than control while non primed plant also reduces its rate as compared with 106 control and receiver.

In treatment 2 (under 150 mM NaCl), emitter shows non-significant with control. The treatment 2 has no effect on proline while it induces proline in receiver. It is assume that emitter plant produce another compound and reduces the amount of proline and that another compound induces a proline in receiver. When we go to step 2, a receiver gives better result than treatment one, its proline content increase up to 104% and becomes equal to control. Non primed plants went to more stress and it proline content rises up to 276%.

# 113 **4.2 H<sub>2</sub>O<sub>2</sub> SPECIES**

At the end of step one, the emitter produce more  $H_2O_2$  than control and have no effect on receiver under treatment 1. When this receiver moves to step 2, it shows a negative response compared with control. The non-primed plant coming from control of step 1 also shows a negative response. Overall treatment 1 do not induces a priming effect on neighbouring plants in case of  $H_2O_2$  species.

118 In treatment 2 emitter produce large amount of  $H_2O_2$  and induces in neighbouring plant. Receiver 119 shows a priming effect in step 2 and increases its production up to 132.7% and tried to near control. 120 Non-primed plants increase H2O2 up to 269.9%.

# 121 4.3 MOISTURE CONTENT

At the end of step 1 fig. 2, salt greatly effect on emitter and this emitter induce a priming effect on receiver. Moisture content decreases compared with control. In step 2 receiver and control undergoes no changes while non-primed plant coming from control of step 1 shows a negative response. Nonprimed plant reduces its moisture content in treatment 1 and 2 up to -17.42% and -44.84 respectively.

# 126 4.4 RELATIVE GROWTH RATE

Relative growth rate means, how much increase in total dry weight per day. The emitter and receiver show non-significant results with each other while both shows a significant result with control. This result shows that the emitter and receiver reduce its growth rate in both treatment but if we compare treatments with each other, it shows that the treatment 2 reduces more growth rate with treatment 1.

In step 2 receiver of both treatments show a non-significant result with control. This shows that receiver which is now primed speed up its growth rate and becomes equal to control. Non-primed plant in treatment 1 shows non-significant with control; this is due to stagnant growth of plants in step 2. Non-primed in treatment 2 greatly reduces its growth rate due to high level of NaCl concentration.

## 135 4.5 PLANT WEIGHT

Plant weight is a major character to study salt stress in plants. Total fresh and dry weight of emitter and receiver in both treatments of step 1 shows a significant reduction in weight. The fresh weight decrease in emitter as well as receiver might be due to decrease in moisture content and dry root weight. In step 2 total fresh weight of receiver shows a non-significant result due to no change in moisture content.

141 Under treatment 2, total fresh and dry weight of emitter and receiver both show non-significant result 142 with each other, which shows that the emitter induces a priming effect on receiver. In step 2 the 143 increase in fresh weight was non-significant because receiver after induction start to increase total dry 144 weight of shoot, while moisture content remains stagnant.

#### 145 4.6 PLANT HEIGHT

At the end of step 1, both root and shoot length increases up to definite length but emitter and receiver of both treatment shows significant reduction in length of both character compared with control. Root length of primed in step 2 of treatment 1 and 2 increases up to 74.03% and 42.13% respectively. While non-primed plant shows no increase in root length compared with control. Shoot length in step 2 show non-significant due to stagnant growth.

#### 151 4.7 NUMBER OF FLOWERS AND BOLLS

152 Number of flowers per six plants was counted at the end of step 1. Controls showed the maximum 153 number of flowers per six plants. The number of flowers was significantly reduced in the emitters and 154 receivers in both the treatments.

The no. of flowers per six plants remained the same for control plants. However, the no. of flowers in treatment 1 primed plants increased from 5 flowers to 8 flowers. No. of flowers in the non-primed plants of treatment 1 decreased from 13 to 11. In treatment 2, no. of flowers in the primed plants increased from 4 to 5 while their no. reduced from 13 to 7.

At the end of step 2, an average of two bolls appeared on the control plants. 6 bolls appeared on the primed while 3 appeared on the non-primed plants of treatment 1. Similarly, 4 bolls were formed on the primed while 1 boll was appeared on average in the non-primed plants of treatment 2.

# 162 4.8 STATISTICAL ANALYSIS

LSD test was used to compare the variation within and between the two treatments for each parameter, while paired t-test was used to compare the step 1 with step 2. These statistical analysis was conducted using the software Statistix 8.1. One-way ANOVA, Mean comparison test and Student's t-test, depending on the dataset, were used to identify significant differences between and within the treatments ( $P \le 0.05$ ).

168 **Table no. 1** 

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Step 1		Control	Treatment 1		Treatment 2	
		Control	Emitter	Receiver	Emitter	Receiver
Physiological	Proline Content (520nm) µmol g⁻¹	0.078 <sup>c</sup>	0.088 <sup>B</sup>	0.091 <sup>B</sup>	0.079 <sup>c</sup>	0.102 <sup>▲</sup>
Data	H <sub>2</sub> O <sub>2</sub> Species (390nm)	0.392 <sup>D</sup>	0.499 <sup>B</sup>	0.363 <sup>D</sup>	0.574 <sup>A</sup>	0.471 <sup>c</sup>
	Moisture Content	15.042 <sup>A</sup>	9.091 <sup>B</sup>	9.629 <sup>B</sup>	6.835 <sup>c</sup>	6.566 <sup>c</sup>
	Relative Growth Rate (g)	0.170 <sup>A</sup>	0.131 <sup>B</sup>	0.137 <sup>в</sup>	0.107 <sup>c</sup>	0.107 <sup>c</sup>
	Total Fresh Weight (g)	19.218 <sup>A</sup>	12.498 <sup>8</sup>	12.418 <sup>B</sup>	9.756 <sup>c</sup>	9.498 <sup>c</sup>
r,	Total Dry Weight (g)	4.177 <b>^</b>	3.407 <sup>B</sup>	3.517 <sup>B</sup>	2.922 <sup>c</sup>	2.932 <sup>c</sup>
l Data	Dry Shoot Weight (g)	3.263 <sup>A</sup>	2.818 <sup>AB</sup>	2.79 <sup>AB</sup>	2.414 <sup>BC</sup>	2.245 <sup>c</sup>
ca	Dry Root Weight (g)	0.913 <sup>A</sup>	0.588 <sup>c</sup>	0.720 <sup>B</sup>	0.507 <sup>D</sup>	0.687 <sup>B</sup>
ogi	Root Length (cm)	14.200 <sup>A</sup>	12.420 <sup>c</sup>	12.440 <sup>c</sup>	12.867 <sup>вс</sup>	13.250 <sup>B</sup>
lo	Shoot Length (cm)	23.125 <sup>A</sup>	21.867 <sup>8</sup>	21.333 <sup>BC</sup>	_20.967 <sup>c</sup>	20.683 <sup>c</sup>
Morphological Data	Root / Shoot in Weight (g)	0.280 <sup>A</sup>	0.209 <sup>B</sup>	0.257 <sup>AB</sup>	0.210 <sup>B</sup>	0.306 <sup>A</sup>
	Root / Shoot in Length (cm)	0.614 <sup>AB</sup>	0.568 <sup>c</sup>	0.583 <sup>BC</sup>	0.614 <sup>AB</sup>	0.641 <sup>A</sup>
	No. of Flowers / 6 plants	13	7	5	6	4
	No. of Leaves	9 <sup>A</sup>	8 <sup>A</sup>	8 <sup>AB</sup>	8 <sup>AB</sup>	7 <sup>B</sup>
	No. of Nodes	7 <sup>A</sup>	6 <sup>AB</sup>	6 <sup>AB</sup>	6 <sup>B</sup>	6 <sup>B</sup>

# 170 Table No. 2

			Treatment 1		Treatment 2	
	Step 2		Primed	Non- Primed	Primed	Non- Primed
Physiolo gical	Proline Content (520nm)	0.205 <sup>B</sup>	0.183 <sup>c</sup>	0.139 <sup>D</sup>	0.209 <sup>в</sup>	0.294 <sup>A</sup>
Data	H <sub>2</sub> O <sub>2</sub> Species (390nm)	1.029 <sup>B</sup>	0.777 <sup>c</sup>	0.569 <sup>D</sup>	1.096 <sup>B</sup>	1.45 <sup>A</sup>
	<b>Moisture Content</b>	13.514 <sup>A</sup>	10.206 <sup>c</sup>	12.421 <sup>B</sup>	6.951 <sup>E</sup>	8.297 <sup>D</sup>
	Relative Growth Rate (g)	0.061 <sup>BC</sup>	0.088 <sup>A</sup>	0.065 <sup>B</sup>	0.045 <sup>c</sup>	0.011 <sup>D</sup>
	Total Fresh Weight (g)	18.916 <sup>A</sup>	14.754 <sup>c</sup>	17.907 <sup>B</sup>	10.782 <sup>E</sup>	12.701 <sup>D</sup>
ata	Total Dry Weight (g)	5.403 <sup>A</sup>	4.548 <sup>B</sup>	5.485 <sup>A</sup>	3.832 <sup>c</sup>	4.404 <sup>B</sup>
	Dry Shoot Weight (g)	4.154 <sup>B</sup>	3.385 <sup>B</sup>	4.235 <sup>A</sup>	2.816 <sup>c</sup>	3.266 <sup>вс</sup>
ca	Dry Root Weight (g)	1.249 <sup>A</sup>	1.163 <sup>АВ</sup>	1.250 <sup>A</sup>	1.015 <sup>₿</sup>	1.138 <sup>AB</sup>
ođ	Root Length (cm)	23.400 <sup>A</sup>	21.650 <sup>₿</sup>	14.300 <sup>D</sup>	18.833 <sup>c</sup>	14.260 <sup>D</sup>
	Shoot Length (cm)	23.683 <sup>B</sup>	22.750 <sup>c</sup>	24.800 <sup>A</sup>	20.967 <sup>D</sup>	23.220 <sup>BC</sup>
Morphological Data	Root / Shoot in Weight (g)	0.301 <sup>A</sup>	0.343 <sup>A</sup>	0.295 <sup>A</sup>	0.360 <sup>A</sup>	0.348 <sup>A</sup>
	Root / Shoot in Length (cm)	0.988 <sup>A</sup>	0.952 <sup>A</sup>	0.577 <sup>c</sup>	0.898 <sup>8</sup>	0.614 <sup>c</sup>
	No. of Flowers / 6	13	8	11	5	7

plants					
No. of Balls / 6 Plants	2	6	3	4	1
No. of Leaves	9 <sup>AB</sup>	8 <sup>AB</sup>	10 <sup>A</sup>	7 <sup>D</sup>	7 <sup>CD</sup>
No. of Nodes	8 <sup>AB</sup>	7 <sup>AB</sup>	9 <b>^</b>	7 <sup>B</sup>	7 <sup>AB</sup>

# 171 Table No. 3

Change in reading in % from step 1 to step 2			Treatment 1		Treatment 2	
		Control	Primed	Non- Primed	Primed	Non- Primed
	Change in Proline Content (520nm)	162.82*	101*	78*	104*	276*
Physiological Data	Change in H2O2 Species (390nm)	162.5*	114*	45.1*	132.7*	269.9*
	Change in Moisture Content	-10.15 N.S	5.99 N.S	-17.42 N.S	5.86 N.S	-44.84 *
_	Change in Total Fresh Weight (g)	-1.59 N.S	18.81 N.S	-7.32 N.S	13.51 N.S	-51.31 *
Data	Change in Total Dry Weight (g)	29.35 *	29.31 *	31.31 N.S	30.69 *	5.43 N.S
Morphological	Change in Dry Shoot Weight (g)	27.3 *	21.32 *	29.78 N.S	25.43 N.S	0.09 N.S
	Change in Dry Root Weight (g)	36.8 *	61.52 *	36.91 N.S	47.74 *	24.64 N.S
Morp	Change in Root Length (cm)	64.79 *	74.03 *	0.7 N.S	42.13 *	0.42 N.S
	Change in Shoot Length (cm)	2.41 N.S	6.64*	7.24 *	1.37 N.S	0.41 N.S

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174 Fig. 2 shows the step 1 of experiment



# 175 **5. DISCUSSION**

Priming is a phenomenon whereby plants once exposed to a particular stress, become tolerant
 towards it as a result of physiological and morphological changes. Several studies have been
 conducted to study plant-to-plant communications caused by biotic and abiotic stresses [16]

This experiment was conducted to study whether the neighbouring cotton plants were able to receive air borne signals from the salinity stressed plants. As indicated by the table, although the receivers were not directly exposed to the salt stress, their dry and fresh weights were still reduced compared to

- the controls. Salinity has been associated with lowering of both fresh and dry weights and it is inversely proportional to the salt concentration [17, 19].
- 184 The reduction in relative growth rate of the receivers compared with the control indicates the 185 possibility of some sort of air borne communication between the emitters and the receivers.

As indicated (Table 1), the fresh and dry weights of cotton emitter and receiver plants affected by
150mmol salt concentration were much lower than those affected with 100mmol salt concentration.
Similar effects were observed in the case of dry root weight. Salt stress has been found to be
associated with reduction in total moisture content of the plant [20, 21].

Same was the case with the emitters in both the treatments. Interestingly, the moisture content of the neighbouring plants were also lowered indicating possibility of plant to plant communication. Salinity causes reduction in root and shoot length leading to stunted growth [22, 25].

By the end of step 1, flowering had been induced in the cotton plants. The number of flowers in the emitter and receiver plants of both the treatments was much less than the control plants.

#### 195 **5.1 STEP 2**

- 196 Step 2 was conducted to check whether the receivers in step 1 had developed resistance against the 197 same stress they were exposed to as receivers.
- 198 The data for various morphological and physiological parameters was collected after a 20 days 199 interval.

Therefore, 100mmol treatment had a little effect on the non-primed and primed plants. Whereas, 150mmol treatment caused significant decline in the moisture content of non-primed plants. No significant effect was observed on the primed plants indicating that the plants had developed resistance against the stress.

Similar to the moisture content, no significant change was observed in primed and non-primed plants in treatment 1 while the moisture content of non-primed plants was significantly lower than the primed plants. This can be attributed to the loss in moisture content due to salinity stress.

There was no increase in total dry weight of non-primed plants in both the treatments. While the dry weight increased for control and the primed plants, it remained stagnant in the non-primed plants indicating stagnant growth. Root length is reduced by salt stress. Though the root length increased normally in the control and the primed plants, there was no significant increase in the root length of the non-primed plants in both the treatments. Low levels of salinity have positive effects on shoot length at certain time of development.

When the treatment 1 primed and non-primed plants were treated with 100mmol salinity, there shoot length increased. There was no significant increase in the shoot length of treatment 2 primed and non-primed plants, however, in comparison with the non-primed plants, the shoot length increased more in the primed plants. Flowering is inversely affected by salinity.

As indicated by the graphs, the flowering was greatly reduced in treatment 2 non primed plants in comparison to the controlled. Salinity induces boll formation in the cotton plants. Rapid boll formation in the primed plants in both treatments indicates acclimatization to salinity.

#### 220 **5.2 CHANGES IN PHYSIOLOGY**

In step 1, the proline content of both the emitters and the receivers increased slightly at 100mmol concentration. At 150mmonl concentration, there was no increase in the proline content but the proline content of the receivers was much higher [26].

This is probably due to the formation of some other compound at higher salt concentration [28] Salinity causes increase in the  $H_2O_2$  species [27].

- As indicated, the H2O2 species increased at both treatment levels. Interestingly, the H2O2
- 227 concentration also increased in the receivers indicating communication.

In step 2, the proline content of treatment 2 non primed plants was more than the primed plants

- indicating preparedness. However, the proline content of control also increased which was due to
- 230 growth of the plant [28]. Same was true for  $H_2O_2$  species.

# 231 6. CONCLUSION

From our study it is concluded that cotton plants were able to perceive stress signals emitted from stressed neighbours and responded by changing physiology and morphology, possibly preparing them for a future stress. The effects were more prominent at 150mM concentration. The result showed that the plants might have communicated with each other through airborne signals and priming was induced on reception of those signals. It is assumed that the tolerance of cotton plants against abiotic stresses may be increased by artificially applying volatile organic compounds.

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