EFFECT OF SALT AND W ATER STRESS ON ROOT INFECTI ON BY Macrophomina phaseolina AND ION COMPOSITION IN SHOOT IN SORGHUM *

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Abstract

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On of salt and water stress to infect the roots by *Macrophomina phase*

position and growth of sorghum (*Sorghum bicolor*) was studied it

19-35[°]C). Treatments consisted of 4 The interaction of salt and water stress to infect the roots by *Macrophomina phaseolina*, and affect the ion composition and growth of sorghum (*Sorghum bicolor)* was studied in a greenhouse experiment (19-35°C). Treatments consisted of 4 levels of salinity (0, 1400, 2100 and 2800 mg of NaCl kg⁻¹ soil) and three water stress levels (3, 7 and 10 irrigation intervals). Infested soil containing 100 viable microsclerotia g^{-1} of a melon isolate of *M. phaseolina* and non-infested soil were used for all treatments. The experiment was arranged in a completely randomized design with four replications. Six-week-old sorghum seedlings after their transferring to infested and noninfested soil were exposed to salt stress, after which, water stress was started. Shoot dry weights were reduced by increasing salinity levels. This reduction was m ore pronounced in infested soil than in non-infested. Increasing irrigation intervals reduced salt injuries. Shoot and root colonization by *M. phaseolina* significantly increased by increasing salinity levels up to 1400 m g of NaCl kg⁻¹ soil. Moreover, salinity and *M. phaseolina* interaction increased the concentrations of Na⁺ and Cl⁻ compared to salt stress *per se*, but negatively correlated with increasing irrigation intervals. Concentration of K^+ was in contrast with Na⁺ and Cl. Also, disease symptoms appeared only in the highest irrigation intervals (7 and 10 days). Consequently, m ore infected crown and root were observed by increasing irrigation intervals and NaCl levels up to 1400 mg kg⁻¹ soil.

Keywords: Matric potential, Osm otic potential, Charcoal rot, *Sorghum bicolor,* Drought.

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Introduction

Macrophomina phaseolina (Tassi) Goid is a soil-borne, m icrosclerotia producing fungus that caus e s root and stem rot on a large number of host plants including sorghum, sunflower, corn, m elon and beans in tropical as well as subtropical regions (Mihail 1992). This fungus usually infects plants that are subjected to severe stresses induced by drought and high tem peratures (Nischwits *et al*. 2002; Olaya *et al*. 1996). Environmental stres ses can predispose p lants to disease from relatively weak parasites, due to their negative effects on host physiology. Drought and salinity are re regard e as the major predisposing factors in plant d iseases (Ma *et al*. 2001, Triky-Dotan *et al*. 2005).

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regarded as the Drought stress is a li mitation to crop productivity. Under this condition, removal of water from the membrane disrupts the normal bilayer structure and results in the m embrane becom exceptionally porous when desiccated. Stress within the lipid bilayer m ay also result in displacem ent of m embrane proteins and this contributes to loss of m embrane integrity, selectivity, disruption of cellular compartmentalization and a loss of activity of enzymes, which are primarily m embrane based. In addition to m embrane dam age, cytosolic and organelle protein m ay exhibit reduced activity or m ay even undergo com plete denaturation when dehydrated. The high concentration of cellular e lectr olytes due to the dehydration of protoplasm m ay also cause disruption of cellular m etabolism (Mahajan and Tuteja 2005).

Soil salinity problem s are widespread often in arid and sem i-arid regions of the world. In the field, salt stress is usually accompanied by water stres s. Irrigation of crop plants with poor quality water often leads to s alt build u p and water s tress (Richardson and McCree 1985). The components of drought and salt stress cross talk with each other as both these stresses ultimately result in dehydration of the cell and osm otic imbalance. Virtually every aspect of

plants physiology as well cellular m etabolism is affected by salt and drought stress (Mahajan and Tuteja, 2005). In saline soils, although water is present it is unavailable to plants because it is retained by the ions in the soil, such as Na + and Cl − (Perez-Le pza *et al*. 2009). High salt concentrations decrease the osm otic potential of soil solution creating a water stress in plants. I n addition to osm otic stress, salinity imposes on plants other stresses such as ion toxicity, as a result of ion entry in excess of appropriate com part mentation, and nutrient imbalances, as commonly seen in the displacement of potassium by sodium. The main damage to plants, however, could result from osmotic stress imposed externally due to high ion concentrations in the soil or internally when excess salt uptake resulted in high salt accum ulation in the intercellular spaces (Zhang *et a l*., 2006).

Sorghum is often grown in areas with relatively low rainfall, high temperatures and saline soils (Netondo *et al.* 2004). Tolerance to salinity is variable among crops (Sanogo 2004). Plant responses to salinity stress depend upon various factors, such as the duration and degree of the stress and growth stage (Trik y-Dotan *et al*. 2005). Most agronomical crops do not function well at a salinity level of $5 \text{ dS} \text{ m}^{-1}$ or higher (Mass 1986). Sorghum's ability to be productive in com parison with oth e r cere als in saline and drought-prone environments has been attributed to several different m orphological and physiological traits (Sm irnoff, 1998). Osmotic adjustment defined as a net increase in tissue solute concentration in response to salt and water s tress has been p roposed as a beneficial salinity and drought tolerance mechanism for several crop species including sorghum (Girm and Krieg 1992). Accum ulation of organic solutes m ainly proline, free a mino acids, and carbohydrates are among the nonspecific mechanism s, which increase under a range of stressful conditions (Khan 2007). This mechanism has

a positive effect on the daily carbon balance of a stressed plant, sinc e it allows the plant to photosynthesize down to lower leaf water potentials than would otherwise have been possible (McCree 1986).

Charcoal rot is a m ajor disease p roblem under drought and saline conditions (Diourt *et al*. 1995). T he effects of salt on plant disease m ay result from its effect on one or m ore of biotic com ponents involved in the disease nam ely the pathogen, the host, m icrobial activity in soils, or abiotic components of soil (Triky-Dotan *et al*. 2005). W ater potentials from -1.2 to -1.5 MPa have been shown to increase predisposition to root rot pathogens. Edmunds (1964) found that charcoal rot was severe in sorghums plants when inoculated near maturity at temperatures of 35-40°C and 25% availab le soil m oisture. W ater stress can predispos e the plant to *M. phaseolina* whenever the defense mechanism of the plant is impaired (Waller 1986).

The objective of the present study was to determine the effects o f several N aCl levels and irrigation intervals on the growth and mineral composition of sorghum and its susceptibility to charcoal rot pathogen.

Materials and methods Soil

A dry virgin, sandy clay soil (pH 7.9, organic matter 2.2%, field capacity 16.52% (w/w), permanent wilting point 3.21% (w/w) and electrical conductivity (EC_e) 0.83 dS m^{-1} was collected from the Agricultural Experim ent Station, Shiraz University in Badjgah, 16 k m north of S hiraz, Iran. To determinate the m axi m um accessib le m oisture for each pot, the soil m oisture curve was obtained at a preliminary experim ent using cell pressure apparatus (Fig. 1) (Cram er *et a l*. 1985). Field capacity (FC) and permanent wilting point (PW P) were found to be 0.0305% and 17.2%, respective ly.

The m aximum access i b le m oisture for each 5000 g pot was 709m l according to equation 1:

water volu me (ml) = water den sity / soil weight (g) \times (FC-PWP) $(0.1723 - 0.0305) \times 5000g = 709ml$

Inoculum production

Archive of the minimizing point (*Archive of the minimizing point* of the amistiking point of the determinate the minimization of the minimization of the minimization of state and minimization of simple and minimization A m elon isolate of *M. phaseolina* (MP) was collected in summer from the Borazjan area, in Boushehr province in southern Iran. Microsclerotia of *M. phaseolina* were obtained free of culture m edium by aseptically p lacing a small agar b lock from an actively growing culture in a flask containing sterile potato dextrose broth (PDB) (Short and Wyllie, 1978). The flask was incubated at room tem perature for 3 months until a thick mat composed predominantly of m icrosclerotia forms on the surface of the broth. The m at was removed, was hed three times using sterile distilled water, and dried at 35°C. The dried microsclerotial mat then were ground with a m ortar and pestle giving a black powder of microsclerotia. The germination of m icrosclerotia was calculated using water agar m edium (WA) prior to the experim ent. The m icrosclerotia were m ixed with 1000 g sterile air-borne sand and stored at 4°C. Before inoculation, the m ixture was added to 104 kg of soil required for infesting soil with the pathogen (to reach 100 viable microsclerotia g^{-1} soil).

Inoculation and salt treatment

Ten germinated sorghum seeds were sown in each pot contain ing 1000 g dry soil (15 cm in diameter and 15 c m high). After 6 weeks, all 10 plants were transferred to larger pots (25 cm in diameter and 25cm high) containing 5000g infested (100 viable microsclerotia $g⁻¹$ soil) and non-infested soil. Salt treatm ents consisted of 0, 1400, 2100 and 2800 of m g NaCl kg^{-1} soil. Sodium chloride was gradually added to the soil in aqueous solution for 7 d to reach the desired NaCl levels. For unifor m water distribution, the pots were irrigated through a nylon tubing placed diago nally in the soil with a closed end and several, 1-2 mm diameter, holes on the

Fig. 1. Soil moisture cur ve for the pot soils.

tube (Banihashem i and deZeeuw 1975). The whole salt s olutions wer e retained in the pots and m aintained in field capacity by periodical weighing.

Irrigation treatments

Fig. 1. Soil moisture curve for the pot soils.

Antenia and deZeeuw 1975). The interactions among salt, was infection by *M. phaseolina* we

ed in field capacity by periodical measuring the percentage of segments coloniz Three irrigation treatments using distilled water at 3, 7 and 10 d intervals were imposed after salt treatm ent and pots wer e irrigated with distilled water to a field capacity by periodical weighing throughout the experim ent (Table 6). The number of total irrigation used during the experiment in 3, 7 and 10 d treatm ents were 16, 7 and 5 times, respectively. The experim ent was conducted in a greenhouse with supplemental light provided by fluorescent and incandescent illum ination s. The air temperature d uring the experim ent fluctuated between 19 ˚C and 35 ˚C, however it often was above 30 ˚C during the day.

Data Collection

At the end of the experim ent (9 wk after inoculation), 15-day-old sorghum seedlings were cut a t the soil line and roots were washed free of soil. The shoots were dried at 70 ˚C for 48 h, weighed and ground, dry-ashed and $Na⁺$ and $K⁺$ concentrations were determined by flame photometry. The Cl concentration was determined following the method of Chapman and Pratt (1961). The

interactions among salt, water stress, and infection by *M. phaseolina* were evaluated by measuring the percentage of crown and root segments colonized. Crowns and roots were thoroughly washed free of soil and surface sterilized for 2 m in in 0.5% sodium hypochlorite. Twenty five pieces (0.5 c m) of crown or root were randomly selected from all plants s eparately in each pot and plated on potato dextrose agar (PDA). All plates were incubated for 10 d at 35 ˚C and the percentages of crown and root segments colonized by *M. phaseolina* were recorded.

Statistical Analysis

The experimental layout was a $4 \times 3 \times 2$ factorial arrangem ent in a com pletely randomized design with three replications. Analysis o f variance for the effects of irr igation interva ls, s alt levels, disease infection and their interactions was done using MSTATC software.

Results

The analysis of variance of the effects of irr igation intervals, salt, charcoa l rot and their interactions on the growth and ion concentrations is shown in Table 1.

Results rep orted here indicate that increasing NaCl level led to a significant ($P \leq$ 0.001) rise in Na⁺ and Cl⁻ concentrations of shoot (Tables 2 and 3). Whereas, there was a

Table 1. Analysis of variance for growth responses and ion concentrations in shoot (s) of sorghum **under four s alt levels, three irrigation treatments and with or without fungus.**

(*) Significant at $P \le 0.05$, (**) Significant at $P \le 0.01$, and (***) Significant at $P \le 0.001$ Ns (not significant at $P \leq 0.05$).

Table 2. Effect of salt, irrigation intervals and charcoal rot on the Na⁺ concentration (%) in shoots of **sorghum.**

* Means followed by the same letters are not significantly different at $P \le 0.05$ according to Duncan's Multiple Range Test.

Table 3. Effect of irrigation intervals and charcoal rot on the Cl - concentration (%) in shoots of sorghum.

Irrigation intervals									
$\rm (day)$	Inoculum			NaCl levels $(mg kg^{-1} soil)$					
	$-Mp$	$+Mp$		1400	2100	2800			
	$5.07a*$	4.2ab	3.11ef	2.76 efg	4.24cd	8.43a			
	3.36ab	3.39 _b	1.99gh	2.59fg	3.61de	5.85b			
10	3.13 _b	2.89b	.45h	2.58fg	3.36ef	4.64c			

* Means followed by the same letters are not significantly different at $P \le 0.05$ according to Duncan's Multiple Range Test.

significant ($P \le 0.001$) decrease in shoot K⁺ concentration with applied salt (Table 4). At any salinity level, infected sorghums accumulated more Na^+ and Cl and less K^+ than non-infected ones.

In non-infested soil, water stress decreased the shoot $Na⁺$ and Cl^{\dagger} concentrations (Tables 2 and 3). Conversely, the K + concentration tended to be increased with increasing irrigation intervals (Table 4).

At any irrigation intervals, a higher $Na⁺$ and CI and a lower K^+ concentrations were observed in infected than in non-infected sorghums (Tables 2-4).

In non-salinized treatments, a reduction in the sho ot dry weight with increasing irrigation intervals was observed, however the differences were not statistically significant (Table 5).

The results also suggest that shoot dry weight was significantly (*P* ≤ 0.001) decreased with increasing NaCl levels at each irrigation interval (Table 5). Increasing irr igation interva ls reduced salt injuries. The highest amount of NaCl (2800 mg NaCl kg⁻¹ soil) with 3 d irrigation interval caused a m axi m um reduction in shoot dry w eight. In all the cases, the reduction of shoot dry weight was m ore in infected plants than in non-infected ones.

At the end of the exp erim ent, cro wn and root of plants were evaluated for visible charcoal rot sym ptom s. No visib le disease symptom s were observ ed in infected plants with 3 d irrigation treatment, while with 7 and 10 d irrigation intervals the typical charcoal rot sym ptom s were observed (data not shown). The results of the present study showed that there was less disease development in 7 d treatm ent, as irregular lesions being extended across the s tem and root, whereas the symptoms in 10 d irrigation intervals w as tissue disintegration in lower portion of stems.

The mean percentage of plants with infected cro wn and root (percentage of crown and root in which *M. phaseolina* was isolated) are shown in Figs 2 and 3. In non-salinized treatments, coloniz ation by *M. phaseolina* was negatively correlated with soil water content. In other words, the percentage of coloniz ation significantly (*P* \leq 0.001) increased by increasing irrigation intervals. However, crown had m ore infection than root. In our study, m ore incidence of infected crown and root was observed with increasing NaCl up to 1400 mg kg^{-1} soil but any further

increase significantly ($P \le 0.001$) decreased colonization (Figs. 2 and 3). Our results indicate that infection by *M. phaseolina* caused plants to accumulate higher Na⁺ and Cl and lower K^+ with increasing salinity level, while increasing irrigation intervals resulted in an opposite trend. In addition, incre asing irrigation intervals increas ed crown and root colonization.

nterval (Table 5). Increasing when soil moisture content is revals reduced salt injuries. The of soil moisture depletion for the TMC (2800 mg NaCl te⁻¹ dirrigation in shoot dry weight. In irrigation treatment (Table 8) d Soil m oisture deficit (SMD) at the various irrigation intervals and NaCl levels is shown in Table 6. SMD is approached to zero when soil moistur e content is m ore. The m ean of soil m oisture d epletion for each irrigation interval was calculated using the mean of applied water (Table 8) during the 7 wk irrigation treatm ent (Table 7). To determine the SMD during the growth period, the m ean soil m oisture dep letion at each irrigation interval w as divided by the m axi m um accessible moisture for 5000 g soil /po t (709ml) (Equation 1). F or example, in the 3 d irrigation interval and 1400 mg kg^{-1} soil, the mean of applied water was 6000 cm³. Thus, the mean of applied water and SMD at each irrigation interval during the irrigation treatments were 375 cm^3 and 52.8% $\arccor \text{dip}$ to $6000/(48 \div 3)$ ÷3) and

 $(375/709) \times 100$, respectively.

According to Table 7, at each irrigation interval the loss of soil m oisture in noninfected treatments was more than that of infected ones. It indicates that in non-infected plants, roots were intact and therefore water uptake was m ore. Also, in each irrigatio n interval, the loss of soil moisture was reduced with increasing salinity showing detrimental effect of salinity on the roots as well as water uptake limitation. In addition, the h igher ion concentrations caused more reduction in soil water osm otic potential.

At the end of the experim ent, the electrical conductivity of saturated soil extract (EC e) in both non-salinized and salinized treatments was measured at three parts of the pots (Table 9). In salinized treatments, at 3 d irrigation interval, the highest EC_e was

	Irrigation intervals (day)							
NaCl levels (mg)	3				10			
kg^{-1} soil)	$-Mp$ inoculum	$+ Mp$ inoculum	$-Mp$ inoculum	$+ Mp$ inoculum	$-Mp$ inoculum	$+ Mp$ inoculum		
$\boldsymbol{0}$	$0.65bc*$	0.6cd	0.71ab	0.67 _{bc}	0.73a	0.7ab		
1400	0.56def	0.56 defg	0.61cd	0.57 _{def}	0.72ab	0.65 _{bc}		
2100	0.5 efgh	0.4k	0.53 efgh	0.45 ijk	0.5 efgh	0.5 fgh		
2800	0.39k	0.4k	0.44 ijk	0.42 jk	0.49 ghij	0.44 ijk		

Table 4. Effect of salt, irrigation intervals and charcoal rot on the K⁺ concentration (%) in shoots of **sorghum.**

Table 5. Effect of salt, irrigation intervals and charcoal rot on shoot dry weight (g/pot) of sorghum.

2800	0.39k	0.4k	0.44 ijk	0.42 jk	0.49 ghij	0.44 ijk	
Means followed by the same letters are not significantly different at $P \le 0.05$							
Table 5. Effect of salt, irrigation intervals and charcoal rot on shoot dry weight (g/pot) of sorghum.							
				Irrigation intervals (day)			
NaCl levels (mg kg^{-1} soil)		3		7	10		
	$-Mp$ inoculum	$+Mp$ inoculum	$-Mp$ inoculum	$+Mp$ inoculum	$-Mp$ inoculum	$+Mp$ inoculum	
$\overline{0}$	$4.1a*$	3.87ab	3.77ab	3.57abc	3.37abc	3abcd	
1400	2.64abcde	2.07cdefgh	2.95abcd	2.6abcdef	3.07abcd	2.65abcde	
2100	1.8 defgh	1.77 defgh	2.45bcdefg	2.07cdefgh	3abcd	2.52bcdefg	
2800	1.05gh	0.8 _h	1.27 efgh	1.07 fgh	1.62 defgh	1.62defgh	
Means followed by the same letters are not significantly different at $P \le 0.05$ Multiple Range Test.						according to Duncan'	
Table 6. Depletion of soil water holding capacity (%) at the various irrigation intervals and NaC							
levels.							
			Irrigation intervals (day)				

Table 6. Depletion of soil water holding capacity (%) at the various irrigation intervals and NaCl levels.

				Irrigation intervals (day)			
NaCl levels	3				10		
$(mg kg^{-1} soil)$	$-Mp$ inoculum	$+ Mp$ inoculum	$-Mp$ inoculum	$+ Mp$ inoculum	$-Mp$ inoculum	$+ Mp$ inoculum	
Ω	66.9	65.5	83.2	77.5	95.2	91.6	
1400	54.3	52.8	77.5	71.9	90.9	86	
2100	51.4	47.9	76.8	66.9	۸۷,۴	76.1	
2800	45.8	36.6	64.8	59.9	79.5	71.9	

Fig. 2. Effect of salt and irrigation intervals on crown colonization by *Macrophomina phaseolina*. Columns followed by the same letters are not significantly different at $P \leq 0.05$ according to **Duncan's Multiple Range Test.**

Fig. 3. Effect of salt and irrigation intervals on root colonization by *Macrophomina phaseolina*. Columns followed by the same letters are not significantly different at $P \leq 0.05$ according to **Duncan's Multiple Range Test.**

Table 7. The mean of soil water loss (cm³) during the irrigation treatments.

				Irrigation intervals (day)		
NaCl levels $(mg kg-1 soil)$					10	
	$-Mp$ inoculum	$+{\bf Mp}$ inoculum	$-Mp$ inoculum	$+{\bf Mp}$ inoculum	$-Mp$ inoculum	$+{\bf Mp}$ inoculum
	475	465	590	550	675	650
1400	385	375	550	510	645	610
2100	365	340	545	475	620	540
2800	325	260	460	425	565	510

	Irrigation intervals (day)							
	3			7	10			
NaCl levels $(mg kg-1 soil)$	$-Mp$ inoculum	$+ Mp$ inoculum	$-Mp$ inoculum	$+{\bf Mp}$ inoculum	- Mp inoculum	$+{\bf Mp}$ inoculum		
0	7600	7440	4012	3740	3240	3120		
1400	6160	6000	3740	3468	3096	2928		
2100	5840	5440	3706	3230	2976	2592		
2800	5200	4160	3128	2890	2712	2448		

Table 8. The mean of applied water (cm³) during the irrigation treatments at the various irrigation **intervals and NaCl levels.**

Table 9. Electrical conductivity (EC_e) (dS m⁻¹) of saturated soil extract in control various salinity **treatments.**

					Irrigation intervals (day)					
NaCl levels $(mg kg-1)$ soil)	3				7			10		
	upper	middle	bottom	upper	middle	bottom	upper	middle	bottom	
$\bf{0}$	0.83	0.94	0.97	0.9	0.95	0.805	0.89	0.98	0.8	
1400	2.73	5.45	9.5	6.41	6.62	4.91	6.76	8.97	3.51	
2100	3.79	8.17	11.72	7.38	7.23	6.61	7.21	13.42	12.15	
2800	4.85	10.88	13.93	8.34	7.84	8.3	7.64	17.87	15.78	
obtained at the bottom third of the pots. It indicates that during irrigation, salts had accumulated in the bottom of the pots, where the roots were active. While in 7 and 10 d irrigation intervals the salt amount in middle part of the pots was more where the end of nylon tubing was placed that shows inadequate of irrigation water to wash the salt.								reports (Ma et al. 2001, Mayek-Prez et al. 2002, Triky-Dotan et al. 2005). Defense mechanisms may be slowed down more in water-stressed than in non-stressed plants (Blodgett et al. 1997). Water stress cause many complex changes in plant metabolism. For instance, stress can affect cell growth, cell division, cell wall synthesis, and plant		

Discussion

As demonstrated in this work, the percentage of plants that developed charcoal rot was negatively correlated with the s oil water content during the experim ent. In other words, the predisposing of plants to the disease was greater at low than high soil water content. These results indicate the inhibitory effect of higher soil m oisture on infection of crown and root and are in agreement with the previously published

reports (Ma *et al*. 2001, Mayek-P rez *et al*. 2002, Triky-Dotan *et al*. 2005). Defense mechanisms may be slowed down more in water-stressed than in non-stressed plants (Blodgett *et al*. 1997). W ater stress cause m any complex changes in plant metabolism . For ins tance, stress can affect cell growth, cell division, cell wall synthesis, and plant hormonal balance, and these m ay affect plant resistance to diseases (Schoeneweiss, 1981). In sorghum, resistance to *M. phaseolina* depends on the m aintenance of physiological vigor caused by stable transpiration rates (Edmunds 1964).

In addition, drought stress increases the negative effects caus e d by *M. phaseolina* in plants (Mayek-Perez *et al*. 2002). More infection under water stress m ay result from the effect of m atric potential on this fungus. Sclerotial germination and m ycelial growth of

M. phaseolina increase with decreasing matric potential up to -1.2 MPa (Goudarzi *et al*. 2008). T here is a rapid loss of viability of *M. phaseolina* sclerotia at high s oil water levels (Shokes *et a l*. 1977). Saturated conditions m ay also affect the sclerotial survival of other plant-pathogenic fungi, including *S clerotinia minor* (Abawi *et al*. 1985), *Verticillium dahliae* (Ioannou *et al*. 1977), and *Rhizoctonia solani* (Ploetz and Mitchell 1985). Reduction of the survival of sclerotia at higher soil water potentials is related to the inability of sclerotia to regulate their water content and the absence of constitutive dormancy (Cook and Al-Hamdani 1986).

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i Similar charcoal rot symptoms that were observed in this work, were found in other susceptible hosts (Mayek-Perez *et al*. 2002). In our view, crown and root charcoal rot that were not expr essed in 3 d irrigation interval sug gesting the possibility o f a latent or quiescent infection. This pathogen apparently infects early, but rem ains laten t until the host is stress ed, at which time the disease progresses rapidly. Subjecting sorghum plants to water stress after inoculation and during reproductive growth may ensure that any latent infections develop scorable regions. Drought stress (high soil tem perature s and low soil m oistures) caused a m arked reduction in total stalk sugars in sorghums which correlated with increased development of charcoal rot. The infected plants show early m aturity, reduced head size, and less number of grains setting. V isibility of the sym ptom s depends upon the severity of infection (Khan 2007).

Data showed that increasing soil salinity results in growth characteristics reduction such as shoot dry weight and a greater red uction occurred in infested soil, while increasing irrigation intervals did not statistically affected plants growth. Sorghum has been considered to be a most droughtresistant (Bhaskaran *et al*. 1985) and a moderately salinity-tolerant crop (Da shti *et al*. 2009). Possibly the extensive waxy bloom on the lower surface of sorghum leaves increases the boundary layer resistance. Although the transpiration rate of sorghum is high when well water ed, it is quic kly reduced when the plants are subjected to water stres s. In sorghum, the upper leaf surface bears only about two-thirds as m any stomata as the lower surface. The stomata on the upper surfaces of well watered plants app ear m ore sensitive to environmental factors than those on the lower surfaces (Bh askaran *et al*. 1985).

As stated earlier, plant responses to salinity and water defi cit are closely related with several overlapping m echanisms. One of the physiological responses to drought and salinity is the accum ulation of intracellular organic solutes. This increase is considerable for proline (Dashti *et al*. 2009). There is a strong correlation between increased cellular proline levels and the capacity to survive both water deficit and the effects of high environmental salinity. It m ay also serve as an organic nitrogen reserve that can b e utilized during recovery (Sairam and Tya gi 2004). Without this mechanism, it is doubtful that the sorghum plants would have continued to add biomass (McCre 1986).

The effect of salinity on plant growth is a com plex syndrome that involves osm otic stress, ion toxicity, an d m ineral deficiencies (Netondo *et al*. 2004). Reduction in dry weight of p lant tissues reflects the increased m etabolic energy cost and reduced carbon gain, which are associated with salt adaptation (Netondo *et al*, 2004; Richardson and McCree 1985). It also reflects salt impact on tissues (Greenway and Munns, 1980), reduction in photosynthetic rates per unit of leaf area (McCree 1986, Netondo *et al*. 2004), and attainm ent of m axim um salt concentration tolerated by the fully expanded leaves (Munns 2002). Leaf growth is generally m ore sensitive than the root growth. Reduced leaf expansion is beneficial to plants under water deficit condition, as less leaf area is exposed resulting in reduced transpiration. In

accordance, m any m atur e plants, for example cotton subjected to drought respond by accelerating senescence and absciss ion of the older leaves. This process is also known as leaf area adjustm ent (Mahajan and Tuteja 2005).

changes, such as water relations,

tolerance during accumulation
 $A \, t \, dl$. 2000). The above changes
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 Archive Domes are to soil sal There was a significant correlation ($P \leq$ 0.001) between colonization of crown and root and increasing salt up to 1400 mg kg^{-1} soil. High salt m ay i mpose an environmental stress and affect plant physiology via morphological, anatomical, m etabolical, and biochem ical changes, such as water relations, ion homeostasis, salt accum ulation, m etabolic pathways, enzym es, and nucleic acids (Hasegawa *et al*. 2000). The above changes might be associated with increased susceptibility to the pathogen (Rasmussen and Stanghellini 1988, Triky-Dotan *et al*. 2005). The response to soil salt found in our experiment, are consistent with the previous work dem onstrating enhanced charcoal rot in melon exposed to salt (Nischwitz *et al.* 2002). Thus, high salt levels m ay be a factor in charcoal rot development leading to higher disease incidences.

Salt alone tended to increase $Na⁺$ and Cl - concentrations (Snapp *et al* 1991). Sodium is the m ajor cation that accum ulated in sorghum roots and stems as NaCl increased $(Netondo$ *et al.* 2004). High $Na⁺$ concentration strongly inhibited uptake and accumulation of K^+ and Ca^{2+} and to a lesser extent of Mg^{2+} by roots. Because K^+ is a macronutrient involved in turgor control, inhibition of K^+ uptake should stunt growth (Renault *et al.* 2001). Also, K^+ frequently is connected with resistance to pathogens (Munns 2002).

High Na⁺ levels in the external medium greatly reduce the physiochemical activity of dissolved calcium and may thus displace Ca^{2+} fro m t h e plasm a m embrane of root cells (Cram er *et al*. 1985). In turn, displacement of Ca^{2+} from root membranes by Na⁺ affects Na/K uptake selectivity in favor of sodium. A low Ca^{2+} concentration under saline

conditions m ay severely affect the functions of m e mbranes as barriers to ion loss from cells (Boursier and Läuchli 1990). Also, the altered ions and water relations have a severe impact on the photosynthetic perform ance of the plant (Netondo *et a l* 2004).

There are various ways by which plants can keep endogenous levels of ions like Cland Na⁺ low. Reduced influx at the root cell plasm a m embrane, efflux from roots, and retranslocation from the leaves to roots are possible mechanis m s. In addition, salt tolerance during accumulation of $Na⁺$ and Cl⁻ at the cellular level can be achieved through loading in vacuoles. Sequestration of Cl and $Na⁺$ in the leaf sheath of grasses is another mechanism of salt tolerance. Exclusion of Na⁺ from leaf blades protects the delicate photosynthesizing tissues as m uch as possible from the potentially toxic ion. Increased NaCl treatments reduced K⁺ concentration in leaf sheaths but not in leaf blades (Netondo *et al*., 2004). According to Munns (2002), osm otic stress is effective in the beginning of exposure to salt, and ion toxicity becom es i mportant in affecting plant growth after prolonged exposure.

Our observation showed that root infection affected ion uptake. Under salt stress, higher amount of $Na⁺$ and Cl and lower amount of K^+ was accumulated in plant tissue when fungus was present. Young leaves were able to m aintain a low Na/K ratio even under the combination of high salt and greatest d isease d evelopment. The only discern able effect of inoculation was to increase concentration of chloride in young leaves at the end of the season. Tom atoes m ay have developed mechanism s to block off ion uptake through infected tissue or to reduce toxic ion accum ulation by ion-recirculation and root exudation. Sodium uptake and exudation rates were both accelerated in safflower plants with severe Phytophthora root rot (Snapp *et al.* 1991).

Our results showed that the inter action between soil water and s alt s tress affected ion concentration in different ways. Salt increased $Na⁺$ and Cl but decreased $K⁺$ accumulation, whereas increasing irrigation intervals reversed them. This data of $Na⁺$ and Cl concentration support by finding of Saadatm and *et al*. (2008). They presented evidence that effect of salt treatment in predisposing shoot and root colonization to Verticillium wilt was greater at high than at low soil water content.

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shower dev At present, increasing irrigation intervals reduced salt injury, so that trend of growth reduction was decreased in low soil water content. Salinized plants have greater ability to continue leaf expansion and carbon gain under water stress that can be attributed primarily to slower development of water stress, which prolonged the osmotic adjustment. The slower water stress development in the salinized plants is due to a lower water loss rate per plant, which in turn was due to both smaller leaf area and reduced water los s per unit of leaf area. Under wate r stres s, the water loss decreased less rapidly in the salinized plants because of the less rapid decrease in water potential, and the greater volume of water rem aining in the growth m edium. Thus, the combined effect o f salinity and water s tress m ay be less detrimental to plant growth and carbon gain than sum of the separate effects of salinity and water s tress (Richardson and McCree 1985). Such result from interactive effects of salt and water stress was reported by Saadatm and *et al .* (2008) in pistachio. It is concluded that although irrigation of sorghum with saline water inhibits plant growth in comparison with non-saline water, it also inhibits water loss and allows a greater degree of osmotic

adjustm ent, so that the plant are able to continue growing longer and reach lower leaf water potentials between irrigations. As shown in the results, the greatest amount of shoot Na⁺ or Cl⁻ and K⁺ accumulation was obtained in 3 and 10 d irrigation intervals, respectively. The greater $Na⁺$ and Cl ions accum ulation in the sorghum shoots at 3 d as com pared to 7 and/or 10 d irrigation intervals was probably responsible for less salinity induced suppression with a rise in irrigation intervals.

As the results showed, the EC_e of saturated soil extract depends on the irrigation intervals. Regarding the way for applying NaCl and irrigation through nylon tubing, soil salinity was greater at the bottom third of the pots and around the roots, but at the highest irrigation interval the greatest amount of NaCl accum ulation was obtained at the m iddle third. Thus, it is the reason of reduced growth of plants under salt stre s s at the low irr igation interval.

In conclusion, artificially induced water stres s identified as a m ajor predisposing factor of sorghum to infection by *M*. *phaseolina*. Knowledge of the predisposition factors for charcoal rot development is essential for the elaboration of accurate screening methods (Olaya *et al*. 1996). W ater m anage m ent can have a significant effect on root colonization by *M. phaseolina.* Charcoal rot is readily controlled by irrigation despite high tem peratures (Norton and Frank 1953). Given the importance of water stress for development of charcoal rot symptoms it seems essential tha t water stress be evaluated as a contributing variable when testing for resis tance to charcoal rot (Diourt *et al*. 1995).

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