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# Phenol Production In Maize (Zea mays L.) in Response To Infection Caused By Fusariun verticillioides (Niren.).

# 4 ABSTRACT

5 This study investigated the phenol production in five maize varieties in response to infection caused 6 by Fusarium verticillioides. Pure culture of the pathogen was obtained from Plant Pathology 7 Laboratory of the International Institute of Tropical Agriculture (IITA), Ibadan, Nigeria. The screen 8 house experiment was laid out in a completely randomized design. Dual inoculation was done where 9 soil was infected separately before planting and seedlings were infected separately two weeks after planting with two volumes (10ml and 20ml) containing  $1.4 \times 10^7$  spores/ml suspension of F. 10 *verticillioides*. The maize plants were harvested at 4<sup>th</sup>, 5<sup>th</sup> and 6<sup>th</sup> weeks after inoculation and the 11 12 maize plants were dried at room temperature before determination of phenol content in each of the varieties. Data gathered on the agronomic parameters and phenol contents were subjected to analysis 13 14 of variance (ANOVA) using SAS 9.1 statistical package.

All the maize varieties recorded more than 70% stalk rot incidence while the severity ranged from 15 16 19.01% in variety ART-98-SW1 to 25.21% in ART-98-SW6. ART-98-SW6 showed the most significant (p<0.05) phenol content (21.28mg/g). Soil inoculation method produced highest 17 percentage stalk rot severity while seedling inoculation method showed a significantly higher phenol 18 19 contents across the period of study. Similarly, severity increased with increasing inoculum levels 20 with highest significance attained at 20ml (23.99%) while higher phenol content was obtained at 21 inoculum level 10 ml (18.61mg/g) compare to results at 20ml (15.63mg/g) and control (7.88mg/g). The maize variety; ART-98-SW6 with highest stalk rot severity also produced the highest phenol 22 23 content. Therefore, rate of phenol production in maize corresponds with the extent of severity F. 24 verticillioides infections.

Key words: Maize varieties, Phenol, *Fusarium verticillioides*, Stalk rot incidence and severity.

# 27 INTRODUCTION

Maize (*Zea mays* L.) is one of the oldest and most productive cereals cultivated across the world (Olawuyi *et al.*, 2010; Okoro-Robinson *et al.*, 2014). The crop plays an important role in the diet of millions of African people due to its high yields per hectare, ease of cultivation and its adaptability to different ecological zones coupled with its versatile food uses and storage characteristics (Olakojo and Iken, 2001; Olawuyi et al. 2015). Maize has been well established in the farming system in Nigeria where it is grown as sole crop or as an intercrop with other crops such as root crops, grain legumes, cereals (sorghum, millet, rice) and even vegetables (Gwinner *et al.*, 1996). Despite the importance and wide cultivation of this cereal, its growth and productions are usually impaired by diseases caused by fungal, bacterial, viral or nematode which results in considerable yield loss and decrease grain quality (Akande and Lamidi, 2006).

38 Fungi were ranked as the second most important cause of maize diseases and the major genera 39 commonly encountered on maize in tropical regions are Fusarium, Aspergillus and Penicillium 40 (Ominski et al., 1994; Orsi et al., 2000). Fusarium is considered as the most devastating fungal 41 menace of maize while Fusarium verticillioides is the prevalent specie causing root rot, stalk rot and 42 ear rots in maize (Masuka et al. 2003; Alankova et al., 2008). Meanwhile, the severe economic 43 losses in corn and other cereal crops worldwide, in addition to the potential occurrence of fumonisins and other mycotoxins in consequence of F. verticillioides infection are a matter of 44 45 concern in current mycotoxicology (Ono et al. 2010).

46 In the recent times, the use of fungicide which is a chemical control measures commonly employed 47 in the management of diseases caused by fungi have been discouraged due to its environmental and 48 food contamination (Akanmu et al. 2013). Hence, there is need to embrace a safer and novel 49 biological approach to plant disease management. In order to improve plants' resistance to diseases 50 effort has been directed at the search for new anti-microbial materials from natural sources, which are mostly low-molecular weight secondary metabolites essential for plant disease resistance (Singh 51 52 et al., 2010; Maddox et al. 2010). While accumulation of certain plant secondary metabolites such as 53 phytoalexins is induced upon pathogen attack, the increase accumulation of phenolic phytoalexins in 54 plants can promote host defense against pathogens (Boudet, 2006; Maddox et al. 2010). More so, 55 phenolic compounds with less complex structures, such as catechol and coumarin, have exhibited bactericidal and fungicidal activities (Cowan, 1999). Hence, the ability of plants to release phenol as 56 57 a defense mechanism against infection by pathogens is of importance in determining its resistance 58 (Samapundo, 2007). This study therefore investigates the phenol production by five maize varieties 59 in response to infection caused by Fusarium verticillioides

#### MATERIALS AND METHODS

Experimental Site: The experiment was conducted in the screen house of the Department of BotanyUniversity of Ibadan, Ibadan, Nigeria.

Sources of materials used: Seeds of maize varieties; SWAM1-SR, BR-9928-DMR-SR, ART-98SW1, ART-98-SW6, BR-9943-DR-SR were obtained from maize germplasm of the Institute of
Agricultural and Research Training (IAR&T), Apata, Ibadan. Pure culture of the characterised *Fusarium verticillioides* isolate was obtained from the Plant Pathology Laboratory, International
Institute of Tropical Agriculture (IITA), Ibadan, Nigeria.

68 **Multiplication of the inoculum:** *Fusarium verticillioides* isolate was sub-cultured, maintained on 69 solidified potato dextrose agar (PDA) and incubated at room temperature  $(28\pm2)$  °C for 7 days to 70 produce multiple cultures of the organism. Each of the cultured plates was flooded with 2ml sterile 71 distilled water. A sterile scalpel was used to harvest the spores and mycelia growths by scrapping 72 them into a sterilized flask. The solution was adjusted with sterile distilled water, stirred and filtered 73 out the spore suspension with the use of muslin cloth. The inoculums load of  $1.4 \times 10^7$  spores/ml was 74 counted using haemocytometer.

Soil preparation and planting: Prior to the conduct of the experiment, the screenhouse was adequately sanitized. Agricultural soil of between 0 to 15 cm depth was collected from the experimental site of Botany Department, University of Ibadan, Ibadan. The soil was sterilized using electric soil sterilizer at 120°C for 1 hour, and was filled in polythene bag at 5kg per bag after cooling. Maize seeds were surface sterilized in a beaker containing sodium hypochlorite (3.5% w/v) for 3minutes after which the seeds were dried in between layers of whatmann filter paper for another 5minutes. The seeds were planted at three seed per hole across the varieties.

Experimental Design: The experiment was laid out in a completely randomized design (CRD) in
ten replications. Treatments across the five maize varieties consisted of two inoculation methods;
soil and seedling inoculations. These were carried out at varying volume of the standardized
(1.4×10<sup>7</sup> spores/ml) inoculums at 0 ml, 10 ml and 20 ml of the spore suspensions. The treatment

- with soil inoculation was conducted at the period of planting while seedling inoculation was carried
  out on the respective plots at 2 weeks after planting. The control experiments were treated with
  respective volume of sterile distilled water. Can include number of replication
- 89 Determination of disease incidence and severity: The percentage of disease incidence and severity
  90 were determined using the formulas:

Disease incidence (%) = <u>number of infected maize plants</u>  $\times 100$ 

92	number of maize plants
93 94	Disease severity (%) = <u>area of plant tissue affected</u> $\times 100$
95	total area
96	
97	Extraction of plant samples: Two replicates from each treatment across the varieties were
98	carefully uprooted at 4 <sup>th</sup> , 5 <sup>th</sup> and 6 <sup>th</sup> week after planting (WAP). Soil adhering to the root of maize
99	plant was removed by agitation in water, the plants were then air dried at room temperature. Two
100	gramme (2g) of leaf sample per treatments was treated with 10ml of 80% methanol in a tightly
101	clogged flask. The treated leaf was allowed to stand for 3 minutes after which the liquid fraction was
102	filtered using muslin cloth. The step was repeated three times with 80% ethanol and the supernatants
103	were collected into volumetric flasks. Final volume of the extracts was made to 50 ml with 80%
104	ethanol and all the extracts from each sample was combined and centrifuged at 2000 revolutions per
105	minutes for 20 minutes in a centrifuge.

Determination of phenol content: This was done using the Folin-Ciocalteu method as described by
Singh *et al.* (2012) in which gallic acid served as a standard and total phenol is expressed as mg/g
gallic acid equivalents (GAE). The amount of total soluble phenol present in the sample was
calculated according to Singh et al. (2012).

110 Data collection and statistical analysis: Data were collected on plant height (cm), leaf number 111 (cm), disease incidence (%) and disease severity (%) weekly for 9 weeks. All the data on growth, 112 disease and phenol contents collected were subjected to ANOVA using SAS 9.1 statistical analysis 113 software and Means were separated by Duncan Multiple Range System at 95% confidence interval.

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#### RESULTS

The fitted model for the interactive effect of maize varieties, inoculation methods, inoculums' quantity and duration of experiment produced a significant (p<0.01) result on the growth and disease parameters measured. All the factors evaluated were significant with plant height, number of leaves and disease severity while only inoculum volume and duration of the experiment (WAP) was significant with disease incidence caused by *F. verticillioides* (Table 1).

Maize variety; BR-9928-DMR-SR followed by ART-98-SW6 recorded the most significant
(p<0.05) growth on plant height and number of leaves across the varieties evaluated. This was</li>
followed by ART-98- SW1 and BR-9943-DR-SR which also recorded similar level of significance
while SWAM 1-SR showed the least growth performances (Table 2).

All the maize varieties recorded more than 70% stalk rot infections with BR-9928-DMR-SR having
highest rate of 73.11%. The stalk rot severity ranged from 19.01% in variety ART-98-SW1 to
25.21% in ART-98-SW6 (Figure 1).

128 Table 3 showed the effect of different inoculation methods and inoculums volume on the growth and 129 disease occurrence in maize plants. While no significant differences was recorded between the inoculation methods with respect to plant height and number of leaves, soil inoculation method 130 131 produced a significant (p<0.05) disease incidence and severity compare to seedling inoculation 132 which showed no significant difference from the control. Whereas, significant (p<0.05) reduction was recorded in the growth rate with increasing inoculums volume. No significant result was 133 134 obtained in the disease incidence with respect to inoculums levels while at 20 ml, there was 135 significant increase in stalk rot severity (Table 3).

There was a significant increase in plant height and leaf number with the increasing weeks afterplanting (Table 4)

Figure 2 showed a linear relationship between the disease incidence caused by *F. verticillioides* in
maize and the period of experiment. A consistent increase in the disease progression was observed

until the 5<sup>th</sup> week when the infection reached its climax and maintained this position till the 9<sup>th</sup> week
of experiment (Figure 2).

The  $r^2$  value of 0.8694 reinstated the reliability of result obtained in the severity of stalk rot with respect to period of experiment. Having received *F. verticillioides* inoculation in the 2<sup>nd</sup> WAP, disease severity increased consistently from 3<sup>rd</sup> week (9.95%) to 9<sup>th</sup> week (36.29%) after planting (Figure 3).

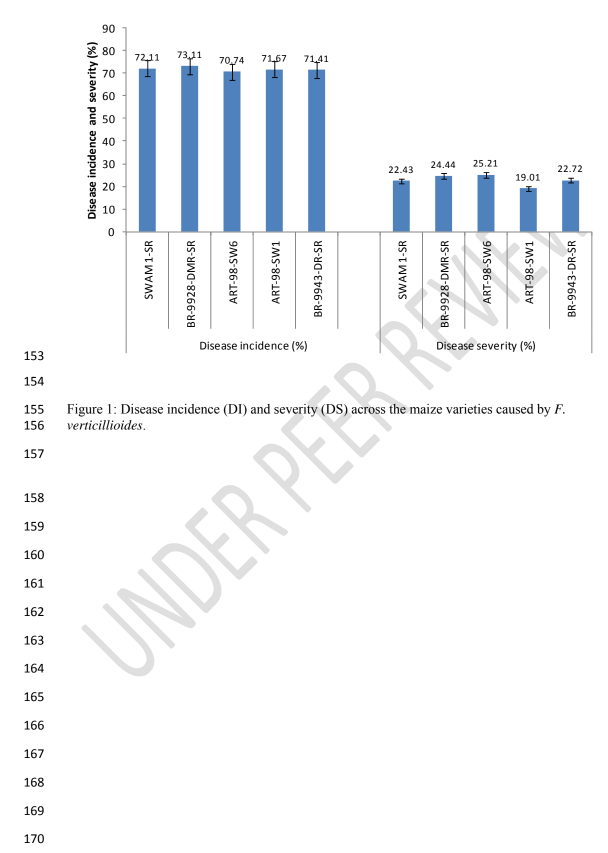
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		Plant height (cm)		Leaf number		Disease incidence		Disease severity	
Sources of variation	df	F value	P value	F value	P value	F value	P value	F value	P value
Model	21	162.46	0.0001**	157.48	0.0001**	210.05	0.0001**	72.74	0.0001**
Maize varieties	4	12.48	0.0001**	14.55	0.0001**	0.30	0.8765	8.82	0.0001**
Inoculums volume	2	43.49	0.0001**	39.33	0.0001**	11.00	0.0011**	34.64	0.0161*
Inoculation method	2	16.16	0.0001**	9.44	0.0001**	0.29	0.5882	5.81	0.0001**
Weeks After Planting	8	405.5	0.0001**	395.34	0.0001**	392.13	0.0001**	172.33	0.0001**
Replicates	9	1.56	0.1693	1.05	0.3864			0.42	0.8372
Error	1327								
Corrected Total	1348				$\frown$				

**Table 1:** ANOVA table of growth and disease occurrence in maize plants after inoculations with *Fusarium verticillioides* 

149 \*Significant \*\* Highly Significant.



171Table 2: Effect of *F. verticillioides* on plant heights and leaf number across the maize varieties

	Varieties	Plant height (cm)	Number of leaves
	SWAM 1-SR	43.27 <sup>c</sup>	4.53 <sup>c</sup>
	BR-9928-DMR-SR	51.31 <sup>a</sup>	5.47 <sup>a</sup>
	ART-98-SW6	54.56 <sup>a</sup>	5.68 <sup>a</sup>
	ART-98- SW1	49.52 <sup>b</sup>	5.07 <sup>b</sup>
	BR-9943-DR-SR	48.94 <sup>b</sup>	4.97 <sup>b</sup>
	$R^2$	0.72	0.71
		18,	
Mean v	vith different letters are signi	ficantly different (p≤0.05)	
		$\langle \rangle$	

# 187 Table 3: Effect of inoculation method and inoculum volume on growth and disease occurrence in

188 maize plants

Parameters	Variables	Plant height (cm)	Number of leaves	Disease incidence (%)	Disease severity (%)
	Control	54.11a	5.63a	70.18a	21.51b
Inoculums' volume (ml)	10	48.88b	4.90b	71.78a	22.18ab
(111)	20	41.58c	4.63c	72.65a	23.99a
				I I	
	Control	52.99a	5.23a	69.56b	20.69b
Inoculation method	Seedling inoculation	46.72b	5.06a	70.18b	21.51b
	Soil inoculation	50.58b	5.17a	74.87a	25.59a
	$R^2$	0.72	0.71	0.94	0.53
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190 Means with different letters are significantly different ( $p \le 0.05$ )

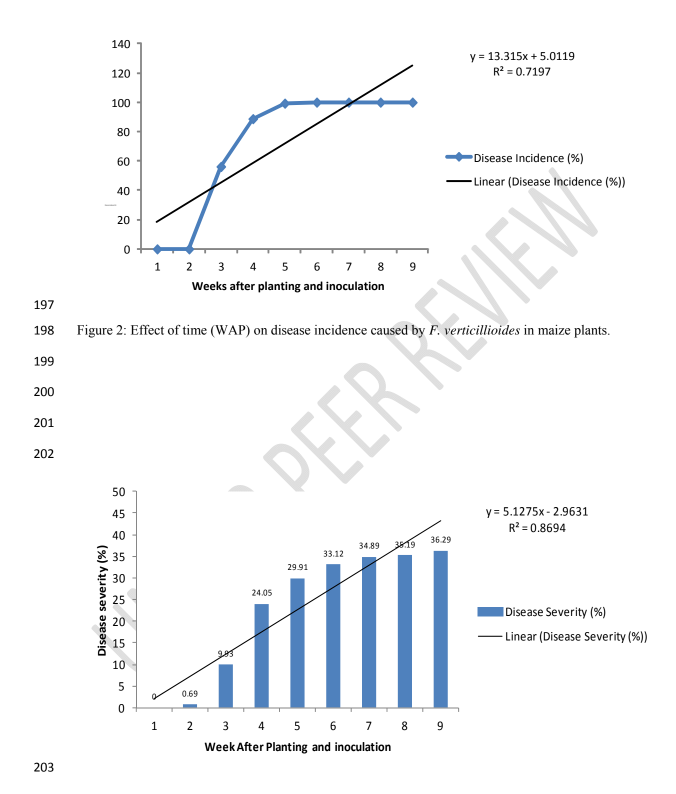
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192 Table 4: Effect of time (WAP explain) on the growth of treated maize plants

Weeks After Planting	Plant height (cm)	Leaf number
	2.08i	0.01i
2	11.78h	1.59h
3	24.61g	3.07g
4	39.63f	4.35f
5	54.69e	5.61e
6	66.92d	6.50d
7	75.94c	7.43c
8	82.57b	8.38b
9	87.65a	9.36a
$R^2$	0.72	0.71

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### 194 Means with different letters in a column are significantly different



**Figure 3:** Effect of time (WAP) on disease severity of *F. verticillioides* in maize plants

The model for the reaction of quantity of phenol produced was significant (p<0.01) through the period of experiment. The effect of maize varieties and inoculums volume also recorded significant result at the 4<sup>th</sup>, 5<sup>th</sup> and 6<sup>th</sup> (WAP). The inoculums' method used showed significance (p<0.01) at 4<sup>th</sup> and 5<sup>th</sup> WAP while it was significant at p<0.05 at 6<sup>th</sup> WAP (Table 5).

209 Maize variety ART-98-SW6 showed the most significant (p<0.05) phenol content across the period

of this study. This result was followed by BR-9928-DMR-SR while ART-98-SW1and BR-9943-

211 DMR-SR showed no significant difference (Table 6).

The seedling inoculation method showed significantly higher phenol contents than soil inoculation, although the two methods produced results which were significantly higher than the control. More so, inoculums volume of 10ml recorded a significantly higher phenol contents than at 20ml (Table 7).

The inoculation method were positively and significantly (p<0.01) correlated with inoculums volume (r=0.64). Also, inoculation method was significantly associated with phenol production at week 4 (r=0.60), week 5 (r=0.68) and week 6 (r=0.71) after planting. Similarly, inoculums' volume was significantly correlated with phenol content obtained at week 4 (r=0.81), week 5 (r=0.90) and week 6 (r=0.89) of the experiment (Table 8).

Source	df	4th	WAP	AP 5th WAP		6th WAP	
Source	u	F value	P>F	F value	P>F	F value	P>F
Model	7	90.24	0.0001**	227.49	0.0001**	310.19	0.0001**
Maize varieties	4	49.92	0.0001**	67.34	0.0001**	74.98	0.0001**
Inoculation methods	1	9.59	0.0065**	9.79	0.0061**	4.21	0.0560*
Inoculums volume	1	139.13	0.0001**	306.88	0.0001**	267.68	0.0001**
Error	17						
Corrected Total	24						

221 Table 5: ANOVA table for Phenol (mg/g) produced by the maize plants after days of infection

222 **\*\***=Highly significant **\***= Significant

Varieties	4th WAP	5th WAP	6th WAP
SWAM1-BR-SR	7.40 <sup>d</sup>	11.12 <sup>d</sup>	14.04 <sup>d</sup>
BR-9928-DMR-SR	13.80 <sup>b</sup>	16.92 <sup>b</sup>	20.96 <sup>b</sup>
ART-98-SW6	16.16 <sup>a</sup>	19.04 <sup>a</sup>	21.28 <sup>a</sup>
ART-98-SW1	11.20 <sup>c</sup>	15.04 <sup>c</sup>	19.22 <sup>c</sup>
BR-9943-DMR-SR	10.52 <sup>c</sup>	14.54 <sup>c</sup>	18.74 <sup>c</sup>
R <sup>2</sup>	0.97	0.99	0.99

# **Table 6**: Phenol (mg/g) content of the maize plants after days of infection

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Means with different letters are significantly different ( $p \le 0.05$ )

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**Table 7**: Phenol (mg/g) content of the maize plants days after inoculation with different volumes of

# 227 Fusarium verticillioides

Parameter	Variable	4th WAP	5th WAP	6th WAP
	Control	4.72 <sup>c</sup>	5.94 <sup>c</sup>	7.88 <sup>c</sup>
Inoculation method	Seedling	14.32 <sup>a</sup>	16.33 <sup>a</sup>	17.58 <sup>a</sup>
methoa	Soil	12.86 <sup>b</sup>	14.03 <sup>b</sup>	15.48 <sup>b</sup>
	$\mathbf{R}^2$	0.97	0.99	0.99
	Control	4.97 <sup>c</sup>	5.94 <sup>c</sup>	7.88 <sup>c</sup>
Inoculums volume	10ml	13.37 <sup>a</sup>	15.32 <sup>a</sup>	18.61 <sup>a</sup>
volume	20ml	10.81 <sup>b</sup>	13.04 <sup>b</sup>	15.63 <sup>b</sup>
	$\mathbf{R}^2$	0.97	0.99	0.99

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# 229 Means with different letters are significantly different ( $p \le 0.05$ ).

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233 **Table 8:** Extent of association between the maize varieties, pathogen inoculation, and time with

<sup>234</sup> phenol content

Completion	Maize	Inoculation	Inoculums'	Phenol contents			
Correlation	varieties	method	volume	Week 4	Week 5	Week 6	
Maize varieties							
Inoculation method	0.00						
Inoculums' volume	0.00	0.64**					
Week 4	-0.13	0.60**	0.81**				
Week 5	-0.11	0.68**	0.90**	0.98**			
Week 6	-0.72	0.71**	0.89**	0.94**	0.99**		

236 \* Significant, \*\*Highly Significant

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# DISCUSSION

High incidence of stalk rot infection recorded across the Fusarium verticillioides inoculated maize 239 varieties supported the claim that F. verticillioides is the major causal agent of highly variable 240 disease symptoms ranging from asymptomatic plants to severe rotting and wilting in maize (Oren et 241 242 al., 2003; Sobowale et al., 2005; Aliakbari, 2007). Virulence of F. verticillioides that ranged from 25.21% in ART-98-SW6 to 19.01% in ART-98-SW1 demonstrated the variations in the severity of 243 244 systemic infection on the host, while environmental factors also play a major in the activities of this 245 cosmopolitan pathogen (Murillo-Williams and Munkvold, 2008; Thompson and Raizada, 2018). More so, (delete) variation recorded in the growth rate and resistance of maize varieties to F. 246 verticillioides infections is in consonance some earlier findings in which host resistance was 247 248 reported as a major determinant of variations in the activities of the pathogen (Sharma, 2003; Olowe 249 et al. 2017).

Similar rate of maize growths observed in both seedling and soil inoculation methods is contrary to the results obtained by Khan et al. (2012) in which a significant increase in the growth parameters of plants that received *Alternaria inocula* through different methods was reported. Also, higher incidence and severity of stalk rot recorded in the soil inoculation method conforms to the earlier claims that attributed varying severities caused by artificial inoculation of *F. verticillioides* with different inoculation methods (Drepper and Renfro, 1990; Sobowale, 2011). Furthermore, the increase in inoculums' quantity which resulted in corresponding decrease in growth rate of maize plants revealed varied efficiency as regards inoculum delivery with different inoculums' level (Sobowale et al., 2007).

The prevalence of stalk rot incidence observed to reach its peak at 5<sup>th</sup> week after planting affirmed that fungus colonizes maize stalks systemically without necessarily causing visible disease symptoms and that *F. verticillioides* strains can be vertically transmitted through seed-to-plant transmission and systemic stalk infection (Munkvold, 1997; Murillo-Williams and Munkvoid, 2008). Meanwhile, the severity rate of 36.29% at 9th week after inoculation justifies the epidemiological claim that *Fusarium* stalk rot reduces output in maize by 10% typically and by 30– 50% in severely affected areas (Li et al. 2010).

266 The result of maize variety ART-98-SW6, followed by BR-9928-DMR-SR, ART-98-SW1, BR-267 9943-DMR-SR and SWAM1-BR-SR as the decreasing order of phenol production after inoculated 268 with F. verticillioides was in agreement with some earlier reports that both constitutive or induced 269 synthesis contribute to abundance and composition of phenolic compounds in cereal grains and this is highly variable depending on the species, variety and environmental conditions (Adom and Liu 270 2002; Lattanzio et al. 2006). The observation of Reddy and Sireesha (2013) that nutritional status 271 272 and concentration of biochemical constituents in plants prior to infection determines the severity of disease possibly explains the reasons maize varieties ART-98-SW6 and BR-9928-DMR-SR with 273 274 higher stalk rot severities also produced higher phenol contents than other varieties tested. This 275 further corroborated the reports that plants respond to pathogen invasion through the activation of 276 complex defense strategies such as the accumulation of flavonoids, phytoalexins and phenolic 277 compounds (Delledonne et al., 2001; Hefny et al., 2012). In this work, total phenol produced by the 278 maize varieties acts as biochemical markers to analyse disease incidence and severity which

conforms to the work done by Singh et al. (2012) on biochemical response and host-pathogen
relationship of stalk rot fungi in early stages of maize (*Zea mays*).

Higher phenol contents produced in seedling inoculation method compare to soil inoculation could 281 282 be associated with plants' reaction to direct inoculation which possibly resulted into a more effective 283 delivery of F. verticillioides inoculums. Thus, the possibility of the pathogen inducing phenol production could be substantiated by the report that biochemical resistance, tolerance or 284 susceptibility in plants against any disease depends mainly on preexisting, preformed or induced 285 286 substances by the pathogen in the host (Reddy and Sireesha, 2013). In this study, the higher phenol 287 production that occurred at lower inoculum volume was consistent with the findings of Perveen et 288 al. (2010) who reported a decrease in total phenol of the leaves of *M. arvensis* with increase in initial 289 inoculum of S. sclerotiorum, a situation attributed to the altered rates of synthetic activity because of 290 infection by pathogen (Howlett, 2006).

The inoculation methods and inoculums volume used in this study effectively delivered infective stalk rot inoculum dosage, the rate of which were found strongly correlated with phenol production in maize plants at 4<sup>th</sup>, 5<sup>th</sup> and 6<sup>th</sup> week after inoculation. Thus, agreed with the claim that majority of phenolic compounds are a part of the preformed general defense system against potential pathogens (Stuper-Szablewska et al. 2017).

In general, phenol production in the maize varieties corresponds with the rate of severity of *F*. *verticillioides* infections. Therefore, close association exists between stalk rot disease and maize plants' resistance to infection through phenol production. Hence, researches that will increase phenol production in crops especially maize, may thus be suggested.

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# REFERENCES-Please check with the journal format, Journal representation uniform Adom KK, Liu RH. (2002). Antioxidant activity of grains. J Agric Food Chem. 50:6182–6187 Akande S.R. and Lamidi G.O. (2006). Performance of quality protein maize varieties and disease reaction in the derived-savanna agro-ecology of South-West Nigeria. *African Journal of Biotechnology*, 5(19), 1744-1748.

- Akanmu A. O., Abiala M. A., Akanmu A.M., Adedeji A. D, Mudiaga P. M and Odebode A. C.
  (2013) Plant Extracts Abated Pathogenic *Fusarium* Species of Millet Seedlings. *Archives of Phytopathology And Plant Protection*, 46 (10) 1189-1205,
- 310
- Alankoya A.E. Monda E.O. and Ajanga S. (2008). Variation in *in vitro* fumonisin B1 production by
   different *Fusariumverticillioides* isolates in Kenya. *American-Eurasian Journal for Agriculture and*
- 313 Environmental Science. 4: 368-371.
- Aliakbari F., Mirabolfathy M., Emami M., Mazhar S.F. and Karami-Osboo R., (2007). Natural
  Occurrence of *Fusarium* species in Maize Kernels at Gholestan Province in Northern Iran. *Asian Journal of Plant Sciences*, 6: 1276-1281.
- Boudet A (2006) Evolution and current status of research in phenolic compounds. Phytochemistry
  68:2722–2735. (fullstop)
- 319
- Cowan M (1999) Plant products as antimicrobial agents. Clin Microbiol Rev 12:564–582. (fullstop)
   321
- Delledonne, M., J. Zeier, A. Marocco and C. Lamb (2001). Signal interactions between nitric oxide
  and reactive oxygen intermediates in the plant hypersensitive disease-resistance response. Proc.
  Natl. Acad. Sci. USA., 98: 13454-13459.
- 325
- 326 Drepper W.J. and Renfro B.L. (1990). Comparison of Methods for Inoculation of Ears and Stalksof
  327 Maize with *Fusarium moniliforme*. *Plant Dis*. 74:952-956.
- 328
- Gwinner J., Harnisch R., Much O. (1996). Manuel sur la manutention et la conversation des grains
   apres-recolte. GTZ, Eschborn, Germany. P 38.
- Hefny M., Attaa S., Bayoumi T., Ammar S.H. and El- Bramawy M. (2012). Breeding Maize for
  Resistance to Ear Rot Caused by *Fusarium moniliforme*. *Pakistan Journal of Biological Sciences*,
  15: 78-84.
- Howlett B.J. (2006) Secondary metabolite toxins and nutrition of plant pathogenic fungi Cur.
  Opinion Plant Biol., 9, pp. 371-375. (fullstop)
- 336
- Khan M.M., Khan M.R. and Mohiddin F.A. (2012). The Relative Performance of Different
  Inoculation Methods with *Alternaria brassicae* and *A. brassicicola* on Indian Mustard. *Plant Pathology Journal*, 11: 93-98.
- Lattanzio V, Lattanzio V.M., Cardinali A. (2006). Role of phenolics in the resistance mechanisms of
   plants against fungal pathogens and insects. Phytochem: Adv Res. 661:23–67.
- Li W.J., He P and Jin J.Y. (2010) Effect of potassium on ultrastructure of maize stalk pith and young
  root and their relation to stalk rot resistance. Agricultural Sciences in China, 9:1467–1474.
- Maddox C.E., Laur L.M. and Tian L. (2010) Antibactarial activity of phenolic compounds against
   the phytopathogen Xylella fastidiosa. Curr Microbiol. 60(1):53-58.
- 346
- Masuka A.J., Cole D.L., and Mguni C. (2003). List of plant diseases in Zimbabwe. Plant Protection
  Research Institute.
- Munkvold, G. P., McGee, D. C., and Carlton, W. M. (1997). Importance of different pathways for
  maize kernel infection by Fusarium moniliforme. Phytopathology 87:209-217. (fullstop)
- 351 Murillo-Williams A. and Munkvoid G.P. (2008) Systemic Infection by Fusarium verticillioides in
- 352 Maize Plants Grown Under Three Temperature Regimes. Plant Disease 92 No. 12, pages 1695 -
- 353 1700 . (fullstop)

- Okoro-Robinson M.O., Olawuyi O.J., Bello W.O. and Babalola B.J. (2014). Comparative evolution
  of organic manure on growth and yield of maize. *Agricultural and Biological Research*. 30(1): 6073.
- Olakojo S.A. and Iken J.E. (2001). Yield performance and stability of some improved maize (*Zea mays* L.) varieties. *Moor Journal of Agricultural Research*. 2: 21-24.
- Olawuyi O.J., Bello O.B., Ntube C.V. and Akanmu A.O. (2015) Progress from selection of some
  maize cultivars' response to drought in the derived savanna of Nigeria. *Agrivita*, 37 (1) 8-17.
  (fullstop)
- Olawuyi O.J., Odebode A.C., Alfar A., Olakojo S.A. and Adesoye A.I. (2010). Performance of
   maize genotypes and arbuscularmycorrhizal fungi in Samara District of Southwest Region of Doha Qatar. *Nigerian Journal of Mycology*, 3(1): 86-100.
- Olowe, O.M., Odebode, A.C., Olawuyi, O.J. and Sobowale, A.A. (2017) Molecular Variability of
   Fusarium verticillioides(Sacc.) in Maize from Three Agro-Ecological Zones of Southwest Nigeria.
   American Journal of Molecular Biology, 7, 30-40. (fullstop)
- OnoE.Y.S., Fungaro M.H.P., Sofia S.H., Miguel T.D., Sugiura Y., Hirooka E.Y. (2010). *Fusarium verticillioides* strains isolated from corn feed: characterization by fumonisin production and RAPD
   fingerprinting. Braz. arch. biol. technol. vol.53 no.4 Curitiba
- 371
- Oren L., Ezrati S., Cohen D., and Sharon A. (2003) Early Events in the *Fusarium verticillioides* Maize Interaction Characterized by Using a Green Fluorescent Protein-Expressing Transgenic
   Isolate. Appl Environ Microbiol. 2003 Mar; 69(3): 1695–1701.
- 375
- Orsi R.B., Correa B., Possi C.R., Schammass E.A., Nogueira J.R., Dias S., Malozzi M.A.B. (2000).
  Mycoflora and occurrence of fumonisin of freshly harvested and stored hybrid maize. *Journal of storage and Product Research*. 36-87.
- 379
- Perveen K., Haseeb A. and Shukla P.K. (2010) Effect of Sclerotinia sclerotiorum on the disease
  development, growth, oil yield and biochemical changes in plants of *Mentha arvensis*. Saudi Journal
  of Biological Sciences, 17, Issue 4, Pages 291-294
- 383
- Reddy M.N. and Sireesha C.H. (2013). Role of oxidative enzymes and biochemical constituents in
   imparting resistance to groundnut (*Arachis hypogea* L.) against stem rot of diseases caused by
   *Sclerotium rolfsii*. Bioresearch Bulletin. 36-41pp.
- Samapundo S., De Meulenaer B., Osei-Nimoh D., Lamboni Y., Debevere J., Devlieghere F. (2007).
  Can phenolic compounds be used for the protection of corn from fungal invasion and mycotoxin contamination during storage? Food Microbiology 24 (2007) 465–473.
- 390
- Sharma, T.R. (2003) Molecular Diagnosis and Application of DNA Markers in the Management of
   Fungal and Bacterial Plant Diseases. Indian Journal of Biotechnology, 2, 99-109. (fullstop)
- 393
- Singh N., Ambika R., Meena S. and Girish M. 2012. Biochemical response and host-pathogen
  relation of stalk rot fungi in early stages of maize (*Zea mays L.*). *African Journal of Biotechnology*.
  11 (82): 14837-14843.
- Singh P., Shukla R., Prakash B., Kumar A., Singh S. and Mishra P.K. (2010). Chemical profile,
  antifungal, antiaflatoxigenic and antioxidant activity of *Citrus maxima* Burm. and *Citrus sinensis* (L)
  Osbeck essential oils and their cyclic monoterpene, DL-limonene. *Food and chemical toxicology*. 48
  (6), 1734-1740.

- Sobowale A.A. (2011). Determination of infective, non-lethal dosage of *Fusarium verticillioides* in
   maize (*Zea mays*) stem and effective inoculation method in the screenhouse. Journal of Agriculture
   and Biological Sciences Vol. 2(5) pp.118-122. (fullstop)
- 404

Sobowale A.A., Cardwell K.F., Odebode A.C., Bandyopadhyay R., Jonathan S.G. (2007).
Persistence of *Trichoderma* species within maize stem against *Fusarium verticillioides*. Arch. *Phytopathol. Plant Prot.* 40, 3: 215-231.

- 408
  409 Sobowale A.A., Cardwell K.F., Odebode A.C., Bandyopadhyay R., Jonathan S.G. (2005). Growth
  410 inhibition of *Fusarium verticillioides* (Sacc.) Nirenberg by isolates of Trichoderma Pseudokoningii
  411 strains from maize plant parts and its rhizosphere. Journal of Plant Protection Research, 45 (4) Pages
  412 249-265.
- 413 Stuper-Szablewska K, Kurasiak-Popowska D, Nawracała J, Perkowski J. (2017). Response of non-414 enzymatic antioxidative mechanisms to stress caused by infection with *Fusarium* fungi and chemical
- 415 protection in different wheat genotypes. Chem Ecol. 33:949–962.
- 416 Thompson M. and Raizada M. (2018). Fungal Pathogens of Maize Gaining Free Passage Along the
- 417 Silk Road. Pathogens, 7, Number 4, 81. (fullstop)