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Influence of pesticides on *Azospirillum* sp. population and its nitrogen fixation in groundnut (*Arachis hypogaea* L.) soils

ABSTRACT

Aim: To study the impact of selected pesticides on *Azospirillum* sp. population and its nitrification in groundnut (*Arachis hypogaea* L.) soils.

Study design: Black clay and red sandy loam soils with known pesticide history were collected from groundnut (*Arachis hypogaea* L.) cultivated fields and were investigated to elucidate the impact of pesticides on *Azospirillum* sp. population and its nitrification in both the soils.

Place and Duration of Study: The soil samples were collected from groundnut cultivated fields of Anantapur District, Andhra Pradesh (A.P) and the study was carried out for 3 months.

Methodology: Ten gram portions of each soil sample were placed in (25 × 150 mm) test tubes and treated with different concentrations of pesticides, (10, 25, 50, 75 and 100 µg g⁻¹ soil) which were equivalent - 1.0, 2.5, 5.0, 7.5 and 10 kg ha⁻¹. Soil samples without pesticides served as controls. The soils with and without pesticides were incubated at room temperature (28 ± 4°C) in the laboratory and moisture content was maintained at 60% water holding capacity (WHC) throughout the experimental period. After 7 and 14 days of incubation, triplicate soil samples were used to estimate the population size of *Azospirillum* sp. using the MPN method. Five ml aliquots of semi – solid malate medium were added to five MPN tubes and inoculated with 0.5 ml of a soil suspension from 10⁻¹ to 10⁻⁵ soil dilutions, and incubated at 37° C.

Results: The population of *Azospirillum* sp. in both soils increased when pesticides were applied @ 2.5 - 5.0 kg ha⁻¹ and incongruity, when the pesticides concentration increased from 7.5 - 10.0 kg ha⁻¹, the *Azospirillum* sp. population gradually decreased in both soils.

Conclusion: The present study aimed at determining the influence of selected pesticides such as oxydemeton methyl, emamectin benzoate, dithane Z-78 and benomyl on the population of *Azospirillum* sp. and nitrogen fixation in black clay soil and red sandy loam soils in groundnut cultivated fields of Anantapur District, Andhra Pradesh, India. Insecticides and fungicides applied up to 5.0 kg ha⁻¹, enhanced the population of *Azospirillum* sp. and its nitrogen fixation also increased significantly after 7 and 14 days of incubation in both soils. However, the population of *Azospirillum* sp., decreased with increasing period of soil incubation in both treated and untreated soils.

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Keywords: Pesticides, Groundnut (*Arachis hypogaea* L.) soils, *Azospirillum* sp. population, nitrogen fixation activity.

1. INTRODUCTION

Soil is an important system for the biological interactions of various microorganisms, hence the applications of pesticides in the agriculture leads to pessimistic side effects on soil micro flora leading to soil pollution and soil contamination [1]. Pesticides may perturb

21 microorganisms by reducing their numbers, biochemical activity, diversity and change the
22 structure of microbial populations. [2,3,4,5,6]. According to [7], pesticides application starts
23 from pre sowing and post sowing stages of seeds, such as treatment of pesticides includes
24 soil treatment, seed treatment and spraying treatment. About 20% of crop farming
25 production and 60% of fruit production are based on the utilization of pesticides [8].
26 According to the FAO data, discontinuation of pesticide practice, would wither agricultural
27 crop yield by 30-50 % with the damage of about 75 billion dollars [9]. According to the type of
28 pest which shows effectual action, pesticides are grouped into insecticides, herbicides and
29 fungicides [10]. In pure culture and in mixed populations the impact of pesticides on the
30 microbial activities of *Azospirillum* has been studied [11,12]. *Azospirillum* sp. are very
31 important rhizosphere bacteria and many species has been isolated from the roots and
32 rhizosphere of numerous host plants and successfully isolated from bulk soil [13], from the
33 beginning of agricultural research on these species [14].

34
35 *Azospirilla* are free-living rhizobacteria that are able to promote plant growth and increase
36 yields in many crops of agronomic importance. It is assumed that the bacteria affect plant
37 growth mainly by the production of plant growth promoting substances, which leads to an
38 improvement in root development and an increase in the rate of water and mineral uptake
39 [15].

40
41 Among the oil yielding crops, Groundnut (*Arachis hypogaeae* L.) is one of the important,
42 major, profitable crops grown throughout the year in India and India is a World leader in
43 groundnut farming, with 8 million hectare of cultivated area in the year 2002-03 [16]. It is the
44 single largest source of edible oils in india and constitutes roughly about 50% of the total oil
45 seed production [17]. Groundnut (*Arachis hypogaea* L.) is one of the major cash crops grown
46 in dry land of India [18]. Within Andhra Pradesh state, Anantapur district, a semi-arid region
47 occupies a predominant place in groundnut cultivation [19].

48
49 The current day agriculture involves huge cultivation of the groundnut crop because of its
50 imperative role in edible oil seeds production [20]. The escalating increase of pest problem
51 and demand for agricultural food production entailed the utilization of agrochemicals that
52 ensure high quality and to crop yield [21]. The application of pesticides into the soil
53 environment inflates concern as to their effect on ecological balance in terms of soil fertility
54 [22,21]. The amount of applied pesticides reaching the target organism is about 0.1% while
55 the remaining bulk contaminates the soil environment [23,24]. Globally, about 3×10^9 kg of
56 pesticides is applied annually with a purchase price of nearly \$40 billions each year [25].
57 According to [26], pesticide residues generally persist in the top 15 cm layer of the soil which
58 is the area of greatest activity of soil microflora that is conducive for - interaction of pesticide
59 residues with the flora of the soil ecosystem [27]. The interaction of pesticides with soil
60 microorganisms and their metabolic activities may change the physiological and biochemical
61 behavior of microorganisms in soil [28]. According to [29], the observed changes in the soil
62 activity depend on the intensity and spectrum of activity as well as tenacity of the parent
63 chemicals or its metabolites.

64
65 Microorganisms play a significant role in many soil biological processes, including nitrogen
66 transformations, organic matter decomposition, nutrient release and their availability, as well
67 as stabilize the soil structure and disturb its fertility, investigated by [30,31,32]. Soil
68 microflora is the first biota that undergoes direct and indirect impacts of toxic substances
69 introduced to soil. The predominant feature of soil quality is considered to be the microbial
70 biomass [33]. Microorganisms forms an essential part of soil food web and hence, microbial
71 biomass is considered to be a measure of potential microbiological and ecosystem
72 functioning. [34].

74 Bacteria that belong to the *Azospirillum* genus are known to associate symbiotically with
75 grass forming specialized structures in the roots in which there is conversion of N₂ to NH₃
76 [35]. *Azospirillum* is a free living micro-aerophilic, heterotrophic diazotrophic bacterium that is
77 involved in heterotrophic nitrogen fixation in several grass bacterial associations [36].

78
79 Agrochemicals especially pesticides and herbicides had adverse effect on *Azospirillum*
80 growth [37]. The impact of several pesticides on the growth and nitrogen fixation of
81 *Azospirillum* sp. has been scrutinized in pure culture systems by few workers [38,39,40,41].
82 Bacteria play an important role in maintaining the health status of soil ecosystem by
83 performing many biological processes. Changes on soil microbial activity may be triggered
84 by different management approaches and the study of the effects of such changes on
85 xenobiotics, of non-target populations, may represent a valuable strategy to evaluate their
86 environmental risk potential. Based on these considerations, the objective of the present
87 study was to evaluate the effect of insecticides and fungicides on *Azospirillum* sp. population
88 and its nitrogen fixation in black clay soil and red sandy loam soils of groundnut (*Arachis*
89 *hypogaeae* L.) cultivated fields of Anantapur District.

91 **2. MATERIALS AND METHODS**

92 **2.1 Soils**

93 Soil samples used in this investigation were collected from groundnut (*Arachis hypogaeae* L.)
94 cultivated fields of Anantapur district of Andhra Pradesh, India, to a depth of 12 cm, air dried
95 and sieved through a 2 - mm sieve before use.

97 **2.1.1 Chemicals**

98
99 For incubation studies and for estimating microbial populations such as *Azospirillum* sp.
100 Commercial formulations of oxydemeton methyl (25 % EC), emamectin benzoate (5 % SG),
101 dithane Z-78 and benomyl dissolved in distilled water were used. The details of the
102 pesticides can be found in Table 2.

105 **2.1.1.1 Soil incubation**

106
107 The soil ecosystem stimulating non-flooded conditions consisting of ten gram portions of soil
108 samples were added in test tubes (25 x 150 mm) and moistened to a water potential of
109 0.090 MPa, in order to maintain at 60% water holding capacity [42].

111 **2.1.1.1.1 Population of *Azospirillum* sp.**

112
113 To determine the influence of selected insecticides oxydemeton methyl, emamectin
114 benzoate and fungicides such as dithane Z-78 and benomyl with concentrations of 10, 25,
115 50, 75 and 100 µg g⁻¹ soil on population of *Azospirillum* sp. Ten gram portions of each soil
116 sample were placed in (25 × 150 mm) test tubes and were treated with different
117 concentrations of pesticides, (10, 25, 50, 75 and 100 µg g⁻¹ soil) which were equivalent to
118 1.0, 2.5, 5.0, 7.5 and 10 kg ha⁻¹ [43,44]. Soil samples without pesticides served as controls.
119 The soils with and without pesticides were incubated at room temperature (28 ± 4°C) in the
120 laboratory and moisture content was maintained at 60% water holding capacity (WHC)
121 throughout the experimental period. After 7 and 14 days of incubation, triplicate soil samples
122 were used to estimate the population size of *Azospirillum* sp. using the MPN method
123 described by [45], with MPN values calculated using probability tables [45]. The growth
124 medium (sterile, nitrogen-free, semi-solid malate medium, pH=6.8 [46] contained (per L):
125 Malic acid, 5 g; KOH, 4g; K₂HPO₄, 0.5 g; MgSO₄, 0.2 g; NaCl, 0.1 g; CaCl₂, 0.02 g; FeSO₄

126 , 0.5 g; Na₂MoO₄, 0.02 g; MnSO₄, 0.01 g; 5 % Alcoholic solution of bromothymol blue, 2 ml;
127 agar, 1.75 g). Five ml aliquots of medium were added to five MPN tubes and inoculated with
128 0.5 ml of a soil suspension from 10⁻¹ to 10⁻⁵ soil dilutions, and incubated at 37° C. MPN
129 tubes in which a typical white pellicle developed a few mm below the surface of the medium
130 after incubation for 36 h were scored positive for *Azospirillum* sp.. Microscopic examination
131 of the cultures revealed the characteristic rods adhered to the flat droplets of oil.
132

133 **2.1.1.1.1 Nitrogen fixation by *Azospirillum* sp.**

134
135 Stock solutions of technical grade pesticides, prepared in acetone, were placed in sterilized
136 test tubes (25 × 200 mm) to provide a final concentration of 50µg ml⁻¹ malate medium. After
137 evaporation of carrier solvent, 20 ml portions of the steam-sterilized malate medium were
138 introduced into each test tube under aseptic conditions. The residues were equilibrated for
139 24 hrs to obtain aqueous solutions of the pesticides [47,48]. Medium, in test tubes without
140 the pesticide served as controls. Soil suspensions (1:10 soil to water ratio) from untreated
141 and pesticide-treated (5 kg ha⁻¹ level with commercial formulations) samples, incubated for 7
142 days, were prepared in sterilized distilled water. These suspensions (0.1 ml) were used to
143 inoculate 20 ml portions of malate medium with and without the pesticide. After 3 days (72 h)
144 incubation at 37°C, these test tubes for each treatment were digested with H₂SO₄ to estimate
145 in total nitrogen (N) by the Micro - Kjeldahl method as described earlier [49,50]. The amount
146 of N present in 0.1 ml soil suspensions, used for inoculation, together with that of the
147 medium was deducted from experimental values.
148

149 *Azospirillum* sp. were isolated from untreated and pesticide - treated (4 times at 10 day
150 intervals) soil samples to determine whether the increased nitrogen fixing capacity of
151 *Azospirillum* sp. isolated from soil samples treated with pesticides would continue further, the
152 isolates were subcultured in the semi - solid malate medium 3 times at an interval of 7 days,
153 and their rates of nitrogen fixation were compared with those of fresh cultures obtained
154 immediately after isolation from untreated and pesticide treated soil samples.
155

156 **3. Statistical analysis**

157
158 All data were expressed on an air dry soil basis and were averages of three replicates. Data
159 were analyzed by significant difference ($P < 0.05$) between pesticide - treated and untreated
160 soils using Duncan multiple range (DMR) test [51,52]. If $A + B < AB$, the response can be
161 considered as synergistic interaction. If $A + B > AB$, the response can be considered as
162 antagonistic interaction; if $A + B = AB$, the response can be considered as additive
163 interaction (where, A = the percent stimulation in population of *Azospirillum* sp. caused by
164 pesticide X alone over the control; B = the percent stimulation in population *Azospirillum* sp.
165 caused by pesticide Y alone over the control; and AB = the percent stimulation in population
166 of *Azospirillum* sp. caused by the combination of X + Y over the control). The percent
167 stimulation values were calculated relative to population of *Azospirillum* sp.
168

169 **4. Results**

170 **4.1 Effect of pesticides on population of *Azospirillum* sp. in soils**

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172 The initial size of the population of *Azospirillum* sp. was low in both soils (Table. 3 and 4).
173 The population of *Azospirillum* sp. was significantly higher in soils treated with oxydemeton
174 methyl, emamectin benzoate, dithane Z-78 and benomyl respectively, than in untreated
175 control soils during the course of experiment (table 1). The population of *Azospirillum* sp. in
176 soils increased when pesticides were applied at 2.5 - 5.0 kg ha⁻¹; by contrast, as the
177 concentration of pesticides increased to 7.5 - 10.0 kg ha⁻¹, the population of *Azospirillum* sp.
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179 gradually decreased in both soils. Application of pesticides, singly and in repeated up to 5.0
 180 kg ha⁻¹, profoundly enhanced the population of *Azospirillum* sp. in vertisol soil (Table 3 and
 181 4). For the laterite soil, pesticide concentrations up to 2.5 kg ha⁻¹ increased the population of
 182 *Azospirillum* sp. after 7 and 14 days of incubation (Table 3 and 4). The increase in
 183 population of *Azospirillum* sp. in vertisol soil amended with oxydemeton methyl, emamectin
 184 benzoate, dithane Z-78 and benomyl (i.e. at 1.0, 2.5 and 5.0 kg ha⁻¹) was 100 - 300, 85 -238,
 185 82 - 192 and 115 - 284 %, respectively, over the control treatment after incubation for 7 days
 186 (Table 3). The population of *Azospirillum* sp. in vertisol soil with or without pesticides
 187 decreased gradually after 14 days (Table 3 and 4) compared to that after 7 days. The
 188 corresponding increases in population of *Azospirillum* sp. in laterite soil amended with four
 189 pesticides at 1.0 and 2.5 kg ha⁻¹ were 46 - 203, 64 - 239, 80 - 239 and 84 - 221 %, respectively,
 190 over the control treatment by the end of 7 day interval (Table 3 and 4). The
 191 population of *Azospirillum* sp. also decreased gradually under similar conditions after a 14
 192 day incubation in laterite soil (Table 4). The influence of oxydemeton methyl, emamectin
 193 benzoate, dithane Z-78 and benomyl alone, at different levels on the population of
 194 *Azospirillum* sp. in the two soils was assessed to examine interaction between pesticides.
 195 Interaction responses are generally distinguished on the basis of percent stimulation values
 196 (over control) regarding any parameter in soil treated with single pesticide or in repeated
 197 application at a specified dose in soil. In this study oxydemeton methyl, emamectin
 198 benzoate, dithane Z-78 and benomyl singly (i.e., at 1.0, 2.5, 5.0, 7.5 and 10.0 kg ha⁻¹)
 199 interacted synergistically, additively and antagonistically, respectively (Table 3.4 and 5). It is
 200 clear from these results that the occurrence of interactions between insecticides and
 201 fungicides was dose-dependent, and these interactions were prevailed in soil even after
 202 incubation for 14 days.

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Table 1. Physico-chemical properties of soils used in the present study

Properties	Black clay soil	Red sandy loam soil
Sand (%)	76.50	72.00
Silt (%)	18.00	25.00
Clay (%)	5.50	3.00
pH ^a	8.40	6.30
Water holding capacity (ml g ⁻¹ soil)	0.48	0.34
Electrical conductivity (m.mhos)	266.00	246.00
Organic matter ^b (%)	0.94	0.80
Total nitrogen ^c (%)	0.05	0.03
NH ₄ ⁺ - N(μ g ⁻¹ soil) ^d	8.95	7.80
NO ₂ ⁻ - N (μ g ⁻¹ soil) ^e	0.51	0.35
NO ₃ ⁻ -N(μ g ⁻¹ soil) ^f	1.04	0.19

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^a 1:1.25 (soil:water)

^b Walkley-Black method (Jackson, 1971)

^c Micro-Kjeldhal method (Jackson, 1971)

^d Nesslerization method (Jackson, 1971)

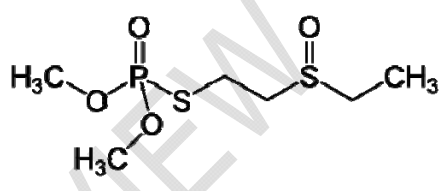
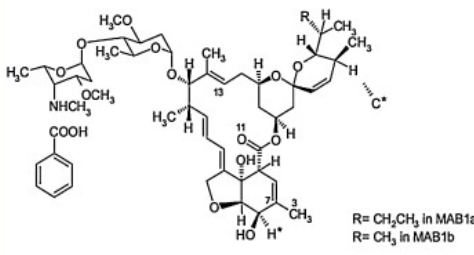
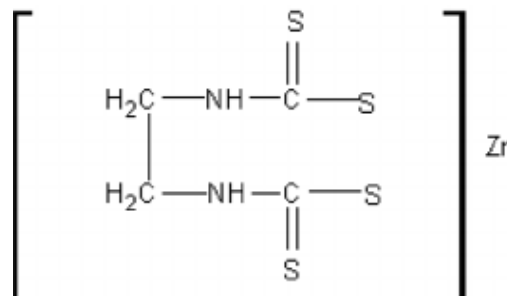
212 ^eDiazotization method (Barnes and Folkard, 1951)

213 ^f Brucine method (Ranney and Bartler, 1972)

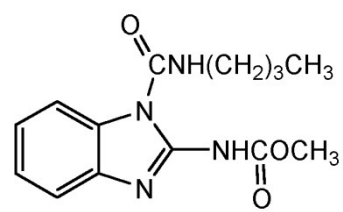
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215 **Table 2. Particulars of the Pesticides used.**

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S.No	PESTICIDE	MOLECULAR FORMULA	STRUCTURE
1.	Oxydemeton Methyl	$C_6H_{15}O_4PS_2$	
2.	Emamectin Benzoate	$C_{51}H_{81}NO_{15}$	
3.	Dithane Z-78	$C_4H_6N_2S_4Zn$	

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4.	Benomyl	$C_{14}H_{18}N_4O_3$	 <chem>CCCCNC(=O)N1C(=NC2=CC=CC=C12)NC(=O)C</chem>
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Table 3. Population (MPN × 10³ g⁻¹ soil) of *Azospirillum* sp. as influenced by the application of pesticides in black soil

Pesticides	Soil incubation in days, after pesticide application												
	0*	7 Days						14 days					
	0**	1.0	2.5	5.0	7.5	10.0	0**	1.0	2.5	5.0	7.5	10.0	
Oxydemeton methyl	2.2	6.5 a (100)	13.0 b (200)	18.0 b (277)	26.0 c (400)	15.0 d (231)	10.0 c (154)	5.2 a (100)	9.4 b (181)	12.0 c (231)	16.0 d (308)	9.3 e (179)	8.1 f (156)
Emamectin benzoate	2.2	6.5 a (100)	12.0 b (185)	16.0 c (246)	22.0 d (338)	31.0 c (477)	8.6 f (132)	5.2 a (100)	8.5 b (163)	11.0 c (211)	14.0 d (269)	12.0 e (231)	7.3 f (140)
Dithane Z-78	2.2	6.5 a (100)	12.0 b (182)	15.0 c (231)	19.0 d (292)	13.0 e (200)	9.1 f (338)	5.2 a (100)	8.2 a (179)	11.0 b (288)	13.0 c (346)	10.2 d (188)	6.3 f (138)
Benomyl	2.2	6.5 a (100)	14.0 b (215)	18.0 c (215)	25.0 d (384)	15.0 c (231)	9.1 f (338)	5.2 a (100)	9.3 b (179)	15.0 c (288)	18.0 d (346)	9.8 e (188)	7.2 f (138)

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*Initial 0-day population

**Concentration of the pesticide, kg ha⁻¹

Figures, in parenthesis, indicate relative productive percentages.

Means, in each row, obtained for each sampling, followed by the same letter are not significantly different (*P* < 0.05) from each other

according to (Duncan's Multiple Range) DMR test.

Values in the table are means of triplicates.

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Table 4. Population (MPN × 10³ g⁻¹ soil) of *Azospirillum* sp. as influenced by the application of pesticides in red soil

Pesticides	Soil incubation in days, after pesticide application												
	0*	7 Days						14 days					
	0**	1.0	2.5	5.0	7.5	10.0	0**	1.0	2.5	5.0	7.5	10.0	
Oxydemeton methyl	2.2	5.6 a (100)	8.2 b (146)	17.0 c (303)	12.0 d (214)	8.5 e (152)	5.0 f (89)	4.2 a (100)	7.3 b (174)	13.1 c (312)	9.4 d (224)	6.5 e (155)	3.2 f (76)
Emamectin benzoate	2.2	5.6 a (100)	9.2 b (164)	19.0 c (339)	14.0 d (250)	12.0 e (214)	4.2 f (75)	4.2 a (100)	7.3 b (174)	14.0 c (333)	11.0 d (262)	6.8 e (162)	3.6 f (86)
Dithane Z-78	2.2	5.6 a (100)	10.0 b (180)	19.0 c (339)	16.0 d (286)	12.0 e (214)	4.3 f (76)	4.2 a (100)	7.1 b (169)	11.3 c (269)	9.4 d (224)	6.2 e (188)	3.5 f (83)
Benomyl	2.2	5.6 a (100)	10.3 b (184)	18.0 c (321)	15.0 d (268)	12.0 e (214)	4.4 f (78)	4.2 a (100)	6.2 b (147)	12.0 c (286)	7.8 d (186)	7.9 d (188)	3.7 e (88)

232 *Initial 0-day population

233 **Concentration of the pesticide, kg ha⁻¹

234 Figures, in parenthesis, indicate relative productive percentages.

235 Means, in each row, obtained for each sampling, followed by the same letter are not significantly different (*P* < 0.05) from

236 each other according to (Duncan's Multiple Range) DMR test.

237 Values in the table are means of triplicates.

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Table 5 : Influence of selected pesticides on nitrogen fixation (mg N g⁻¹ malate) by *Azospirillum* sp.

Soil Type	Cultures from untreated soil		Culture from pesticide treated soil	
	Untreated	**50 µg ml ⁻¹	Untreated	**50 µg ml ⁻¹
Oxydemeton methyl				
Black Soil	7.80 a	11.89 b	10.98 b	14.24 c
Red Soil	5.32 a	08.78 b	09.24 c	11.82 d
Emamectin benzoate				
Black Soil	6.82 a	10.34 b	11.22 b	13.21 c
Red Soil	4.82 a	07.78 b	09.02 b	11.32 c
Dithane Z-78				
Black Soil	5.78 a	09.78 a	12.01 c	12.86 c
Red Soil	4.92 a	08.71 b	09.02 b	11.32 c
Benomyl				
Black Soil	6.24 a	10.31 b	11.24 c	11.83 c
Red Soil	4.89 a	08.24 b	09.85 b	10.54 c

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*The soil sample was treated with commercial formulation of the four pesticides (5 kg ha⁻¹) and culture was isolated after 7 days.

**Semi-solid malate medium was supplemented with technical sample of the pesticides (50 µg ml⁻¹ medium) before incubation with the culture.

Means (n = 3), in each row, are significant (P < 0.05) from each other according to Duncan's Multiple Range (DMR) test.

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Table 6. Impact of subculturing of *Azospirillum* sp. isolated from pesticide-treated soil samples on nitrogen fixation (mg N g⁻¹ malate)

Soil type	Fresh isolate from untreated soil**	Isolate from pesticide-treated soil*	
		Fresh	After third subculturing**
Black Soil			
1.Oxydemeton methyl	8.80 a	18.78 b	17.92 b
2.Emamectin Benzoate	9.65 a	19.24 b	19.05 b
3.Dithane Z-78	7.94 a	18.23 b	17.98 b
4.Benomyl	8.24 a	17.68 b	16.98 b
Red Soil			
1.Oxydemeton methyl	7.76 a	17.34 b	16.88 b
2.Emamectin Benzoate	8.64 a	18.34 b	17.94 b
3.Dithane Z-78	7.68 a	17.42 b	16.82 b
4.Benomyl	7.24 a	17.08 b	16.24 b

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*Soil samples were treated four times with pesticides at 5 kg ha⁻¹ level.

**Semi-solid malate medium was supplemented with technical sample of the pesticides (50µg ml⁻¹ medium) before incubation with the culture.

Means (n = 3), in each row, are significant (P < 0.05) from each other according to Duncan's Multiple Range (DMR) test.

293 **5. Discussion**

294

295 In the present study, four pesticides applied to soil, singly at concentrations ranging from 1.0
296 to 5.0 kg ha⁻¹, had no deleterious effect on *Azospirillum* sp. A similar individual instigate
297 effect of monocrotophos and chlorpyrifos was previously demonstrated on the population of
298 *Azospirillum* sp. [53]. Similarly, observations with other organophosphorus and pyrethroid
299 insecticides and fungicides have also been reported [44,41]. Interactions between different
300 agrochemicals applied in repeated application on microorganisms and their activities in soils
301 have received little attention in comparison to effects of a single agrochemical. There were
302 no differences in degree of diversity in bacterial populations from the application of a
303 combination of five pesticides, including chlorfenviphos and glyphosate, to field plot of 20
304 years[54]. In the present study the application of pesticides to the soils at certain
305 concentrations was not harmful to the population of *Azospirillum* sp. Some reports have
306 been published on interactions between pesticides and their solvents, pesticides and their
307 degradation products, and two different pesticides on growth of organisms in pure culture
308 studies of fungi, algae and cyanobacteria [55,56,57,58,59,60,61]. In all these studies, a
309 variety of interaction effects such as synergistic, additive and antagonistic were observed,
310 depending on concentration of the interacting chemicals. For instance, the combination of
311 permethrin and its degradation product interact to yield antagonistic, additive and synergistic
312 interactions towards the growth of fungi in pure culture [60], because the degradation rate of
313 an individual pesticide may be changed due to the combinations of pesticides, ultimately
314 leading to different types of interactions. In the present study, similar types of interactions
315 occurred by selected pesticides on population of *Azospirillum* sp. in two soils. A increase in
316 the population of *Azospirillum* sp. at high concentrations (100 ppm) of benomyl or 2-
317 aminobenzimidazole (a hydrolysis product of benomyl) were also reported in paddy soil
318 [36,38]. [39], noticed a provoking response in *Azospirillum* sp. population, when treated with
319 benomyl at lower concentration (5 ppm) in alluvial, laterite and saline soils, and carbofuran in
320 alluvial soil only.

321 These observations are in agreement with the results of the present study. The overall
322 influence of pesticides on microbial activities in soil may be subject to interactions between
323 pesticides (i.e. additive, synergistic and antagonistic) and may differ from the response of the
324 individual pesticide components [62]. In the present study similar types of interactions
325 occurred between selected insecticide and fungicides in two soils. Although the mechanisms
326 of interactions are not known, interaction patterns may have a profound influence on soil
327 microflora and their activities, thereby affecting soil fertility. Pesticides added to soil undergo
328 degradation to metabolites in the course of time. For instance, monocrotophos is hydrolysed
329 to N-methyl acetoacetamide [63]. Pesticides are generally applied simultaneously or serially
330 for crop protection, hence the degradation behavior of a pesticide may be changed after it
331 interacts with other pesticides (or their degradation products) already present in the soil;
332 such changes in pesticide degradation may have different side effects on biological
333 processes, such as nitrification and on microbial populations. The presence of chlorothalonil
334 has been suggested as altering the degradation behavior of chlorpyrifos - degrading
335 microbes [64]. The persistent interaction responses recorded in the present study cannot be
336 attributed exclusively to parent pesticides, since metabolites may also have biological
337 effects. Generally pesticides are recalcitrant (not easily degradable) substances, hence they
338 persist for long periods in the soils. This may be one of the main reasons for persistent
339 interactive effects in soil. The present study further accentuates the need for a systemic
340 study on the interactive effects of pesticides used extensively, as well as their metabolites.
341 The results of the present investigation clearly indicate that the selected pesticides –
342 oxydemeton methyl, emamectin benzoate, dithane Z-78 and benomyl, respectively at levels
343 ranging from 1.0 to 5.0 kg ha⁻¹ significantly increased the population of *Azospirillum* sp.
344 .Furthermore, these pesticides, singly and in repeated application, at levels of 1.0 to 10.0 kg
345 ha⁻¹ exerted synergistic, additive or antagonistic interactions towards population of

346 *Azospirillum* sp. in these soils. *Azospirillum* sp. cultures obtained after 7 days of soil
347 incubation, from unamended soils exhibited appreciable nitrogen fixing activity (Table 5). A
348 significant stimulation of nitrogen fixation was evident in cultures from soils treated with the
349 four pesticides at a level of 5 kg ha⁻¹ when compared with cultures from untreated soils. The
350 extent of nitrogen fixation by the cultures observed in the present study are comparable with
351 those of *Azospirillum* cultures isolated from the same soils amended with monocrotophos
352 and quinolphos for 7 days [40], and those cultures isolated from a rice soil amended with
353 benomyl and incubated for 30 days[36]. The cultures from untreated soil, when inoculated
354 into the medium supplemented with four pesticides (Oxydemeton Methyl, Emamectin
355 Benzoate, Dithane Z-78 and benomyl) at 50 µg ml⁻¹, exhibited greater nitrogen-fixing activity.
356 However, the stimulation in nitrogen fixation was more pronounced in cultures of
357 *Azospirillum* sp. isolated from four pesticides treated (5 kg ha⁻¹) soil and inoculated to the
358 medium containing 50 µg ml⁻¹ of the pesticide (Table 5).
359 An attempt was made to determine whether the observed nitrogenase activity would
360 continue upon subsequent subcultures of the diazotroph. Although, fresh cultures from the
361 pesticide-treated soil exhibited greater nitrogen-fixation when compared with those from
362 untreated soils, subculturing of the isolates 3 times had no effect on nitrogen-fixation in the
363 cultures of *Azospirillum* sp., exposed to the selected pesticides (Table 6).
364 The present study clearly shows that soil application of pesticides (Oxydemeton Methyl,
365 Emamectin Benzoate, Dithane Z -78 and benomyl) increased the population of *Azospirillum*
366 sp., isolated from treated with four pesticides, last for longer periods.
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369 6. CONCLUSION

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371 The results of present investigation clearly indicate that the selected pesticides at levels
372 ranging from 2.5 to 5.0 Kg ha⁻¹ significantly increased the population of *Azospirillum* sp. and
373 nitrification in both the soils. Furthermore, increase in the concentration above 2.5 or 5.0 K g
374 ha⁻¹ exerted synergistic, additive or antagonistic interactions towards population of
375 *Azospirillum* sp. and nitrification in these soils.
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378 COMPETING INTERESTS

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380 Authors have declared that no competing interests exist.

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382 Ethical: NA
383 Consent: NA

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UNDER PEER REVIEW