

Yield Response of African rice genotypes to mycorrhizal fungi and rhizobium inoculation

ABSTRACT

Aims: A short term field study was conducted to investigate the yield performance of selected African rice genotypes inoculated with biofertilizers.

Study design: A randomized complete block design laid out in a split-plot arrangement was used to evaluate response of yield components and grain yield of some selected African rice genotypes will be to mycorrhizal fungi and rhizobium inoculation.

Place and Duration of Study: The study was conducted at the Teaching and Research farm of the Federal university of technology, Akure Ondo state, Nigeria during the 2013 planting season.

Methodology: The study was laid out in a split plot arrangement in Randomized Complete Block Design (RCBD), with mycorrhizal fungi, rhizobium inoculation and control in the main plot, while genotypes (N-U-1, N-U-8, WAB 56-104, OFADA GR and MOROBEREKAN) were in the sub-plot and treatments were replicated thrice. There were three main blocks, each block consist of 15 sub-plots with a size measurement of 2m x 1m and inter sub-plot spacing of 0.5m in between plots. A total of 50 plants were raised per sub plot. Transplanted seedlings were planted with the soil slurry containing rhizobium and mycorrhizal fungi inoculum into planting holes in the field at two seedlings per stand, according to their respective plot at a spacing of 25cm x 25cm. Yield component data collected include; number of days to 90% maturity, number of days to 50% flowering; plant height at maturity, number of primary tillers per plot, number of grains per panicle, number of panicles, number of filled and unfilled spikelet, weight of 1000 filled grains and grain yield per plot.

Results: Result showed significant ($P < 0.05$) single and interactive effect of rhizobium and mycorrhizal fungi inoculation on rice yield and yield components. 61.4% increase in grain yield was observed in rhizobium inoculated genotypes when compared to 37.4% increase in mycorrhized genotypes and the uninoculated control. WAB56–104 and N-U-8 had the best interactive response amongst genotypes inoculated with rhizobium while genotypes WAB56-104 and MOROBEREKAN responded better amongst mycorrhized genotypes in relation to yield components.

Conclusion: The results from this study indicate that African rice genotypes differ in grain yield response and host specificity when inoculated with mycorrhizal fungi and rhizobium inoculums. However, inoculating specific African rice genotypes with mycorrhizal fungi and rhizobium can positively influence their grain yield and yield component development and this could play an important role in improving African rice productivity.

Keywords: Mycorrhizal fungi; rhizobium; biofertilizer; African rice; grain yield

1. INTRODUCTION

Rice (*Oryza sativa*) is a major staple food for millions of people in West Africa and the most in-demand staple amongst cereal crops in Nigeria's food basket [1]; [2]. Rice cultivation and production in Nigeria has increased in recent times due to series of government initiatives, change in policies and increased efforts towards self-sufficiency. However, there has been a considerable lag between production and demand level with imports making up the shortfall. The low productivity of the rice production system in

20 Nigeria is due to a lot of factors such as socio-economic constraint, crop management system and lack or
21 no access to external inputs [3]. Rain fed upland rice production system is the dominant cultivation
22 system across several agro-ecological zones in Nigeria [2]. Several abiotic and biotic factors are
23 responsible for the low productivity of upland rice production system [4]. Studies have shown that nitrogen
24 deficiency, phosphorus fixation, weed build-up, rice blast and drought are the leading constraints to
25 upland rice production in Nigeria [5]; [2]. Small holder farmers, **who are the predominant** rice growers in
26 the country are unable to realize the potentials of recently released improved high yielding African rice
27 genotypes such as NERICA which mine soil nutrients rapidly and have higher nutrient use efficiency than
28 traditional genotypes. Furthermore, smallholder farmers lack the **financial resources** to purchase chemical
29 fertilizer to replenish mined nutrient from the soil. Exploitation of microbial sources such as mycorrhizal
30 fungi and rhizobium as biofertilizers for rice growth promotion and increased yield have been previously
31 tested due to their excellent endophytic plant-microbe interactions [6];[7]. Mycorrhizal fungi are excellent
32 colonizer of plant roots. They help colonized plant in accessing water especially during dry spells and also
33 help in the uptake and solubilization of immobile soil nutrients while the plants supplies carbon as food
34 and energy source to the fungi in a symbiotic relationship [8]; [9]. Phosphorus (P) deficiency and fixation
35 can severely limits rice production; colonization of plant root with mycorrhizal fungi may have an
36 influencing effect on P **solubilization, uptake, and plant growth** [10]. Rhizobium are largely recognized for
37 their role in nodule formation in leguminous crops through biological nitrogen fixation but studies have
38 also shown that they can be inoculated into non-leguminous crops such as rice for plant growth promotion
39 and increase yield [11]; [6]; [7]. They are widely regarded as the most efficient biofertilizer in relation to
40 the quantity of nitrogen fixed. Rhizobium is said to promote plant growth through mobilization and fixation
41 of nutrient, improving plant resistance to abiotic stress, solubilization of nutrients in nutrient fixed soils,
42 release of plant growth-hormones [7]; [10]; [12 13]; [13 14]. Therefore, coating African rice genotype seed
43 or soaking seedlings in soil slurry with mycorrhizal fungi and rhizobium inoculum before planting or
44 transplanting could help in improving nitrogen and phosphorus bioavailability and uptake in deficient soils
45 which would help improve rice yields, increase economic return to farmers and mitigate environmental
46 pollution. **There has been scanty experimental evidence in Nigeria on the ability of Rhizobium which are**
47 **normally associated with leguminous crops and the ability of arbuscular mycorrhizal fungi (AMF) to**
48 **colonize roots of certain cereals e.g. rice, in nutrient deficient soils and promote their growth and yield.**
49 **Increased interest in nitrogen fixing bacteria associated with cereals such as rice, wheat and maize has**
50 **been shown in recent times to reduce the use of expensive mineral fertilizers in cereal production. One of**
51 **the reasons for the success recorded with nitrogen fixation independent of nodule formation in rice**
52 **studies is the observation that nitrogen status of rainfed and irrigated lowland soils under rice cultivation**
53 **has increased due to the activities of nitrogen fixing bacteria which survived under such condition evident**
54 **with increased growth and population of beneficial microbes [14]; [15]. Inoculation and improvement of**
55 **cereals through nitrogen fixing bacteria have been observed in various field studies [16]; [3]; [4]; [17].**
56 However, studies conducted by [18], suggested that response of rice genotypes to inoculation with

57 beneficial organisms may differ due to specificity of plant-bacterial and fungi associations, **gas exchange**
58 **and differences in root exudation**. Therefore, rice genotypes with the best response from inoculations with
59 introduced or native beneficial organisms should be selected for recommendation. With this in hindsight,
60 the study was set up to evaluate the performance of African rice genotypes inoculated with mycorrhizal
61 fungi and rhizobium under field conditions with a view to identify promising and best performing
62 inoculated genotypes in terms of yield for recommendation.

63

64 **2. MATERIALS AND METHODS**

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66 **2.1. Description of Location and Experimental Site**

67 The study was conducted at the Teaching and Research farm of the Federal university of technology,
68 Akure Ondo state, Nigeria during the **planting season** of 2013. **The vegetation is a tropical rain forest with**
69 **an average relative humidity of between 56 and 59% during the dry season and between 80% - 85%**
70 **during the wet season**. **The study site** is located between Latitude 5°08 10.5"E and 7°17' 59.2"N, and at
71 elevation of 140 m above the mean sea level. **The site has an average annual rainfall of about 1613mm**
72 **per annum and an annual mean temperature of 27°C**. However, during the course of the experiment,
73 average annual rainfall fell to 1233mm and temperature level increased to a mean average of 31°C due
74 to fluctuations in weather conditions. The Soil at the experimental site was a Sandy clay loam classified
75 under the soil order alfisol according to [19] soil classification. The experimental site was ploughed,
76 cleared and pegged before transplanting **and** experimental blocks were laid out and sectioned
77 accordingly. Seedlings were transplanted with the soil slurry carefully to prevent root damage and also **to**
78 ensure optimal root colonization by **inoculated inoculums**. Seedlings of each variety were planted in each
79 designated block. There was no pre or post application of herbicides/pesticides and no basal or
80 recommended fertilizer application was added throughout the duration of the experiment. Weeding was
81 done manually by hoe and hand.

82

83 **2.2. Nursery practice**

84 The nursery stage was conducted at the screen house of the Department of Crop Soil and Pest
85 Management of the Federal University of Technology, Akure. Micro polythene pots of about 5cm in
86 diameter and 10cm in length were filled with topsoil and mixed with mycorrhizae (*Glomus intaradices*) and
87 rhizobium strains (RACA 3/5/12) separately to form slurry at a weight of 50g per pot; this was done to
88 ensure maximum colonization of rice roots before transplanting. Rice seeds were sown at 5 seeds per pot
89 and was later thinned to 2 seedlings per pot. The pots were made moist and maintained for about 14
90 days, **and thereafter** germinated seedlings were transplanted to the field.

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95 2.3. Pre-Planting Soil Analysis

96 Soil samples were collected at a depth of 0-15cm and bulked together prior to the determination of
97 physico-chemical properties before planting. Soil pH was determined in 1:2.5 (soil: water) ratio using
98 glass electrode pH meter. Soil organic matter was determined according to [20] method. Total nitrogen in
99 the soil was determined using Kjeldahl method [21]. Available phosphorus was extracted using Bray-1 P
100 followed by molybdenum blue colorimetry. Exchangeable cation (K, Ca, Mg) were extracted with 1 N
101 Ammonium Acetate K and the extract was determined by flame photometry, Ca and Mg were determined
102 by Atomic Absorption Spectrophotometer (AAS) (Table 1).

103
104

Table 1. Physico-chemical properties of experimental soil before planting

Soil properties	Values
Sand (%)	60.4
Clay (%)	26
Silt (%)	13.6
Textural class	Sandy clay loam
Nitrogen (g/kg)	0.28
Organic Carbon (%)	1.90
Organic Matter (%)	3.26
Calcium (cmol/kg)	2.51
Magnesium (cmol/kg)	2.13
Potassium (cmol/kg)	2.04
Phosphorus (mg/kg)	26.58
pH	5.03
CEC	6.68

105 *mean values are presented in the table (n = 4)
106

107 The pre-plant soil physico-chemical properties of the experimental site as shown above indicate that the
108 soil contains 60.4%, 26%, and 13.6% sand, silt and clay respectively and falls into the textural class of
109 sandy clay loam. The soil organic carbon and total nitrogen values were 1.90% and 0.28g/kg respectively
110 which are below critical limit. The soil was acidic with a pH of 5.03 and has a cation exchange capacity
111 (CEC) of 6.68; potassium level of 2.04cmolk⁻¹; phosphorus level of 26.58mg/kg⁻¹; magnesium level of
112 2.13 cmolk⁻¹ and calcium level of 2.51cmolk⁻¹. Analysis indicate the need for nitrogen and phosphorus
113 fertilization as they are deficient in the soil and in an unavailable form for plant use, which justified the
114 need for inoculation with biofertilizers which will help to increase nutrient availability and uptake for
115 enhanced rice yield.

116 2.4. Experimental Design

117 The study was a split plot arrangement in Randomized Complete Block Design (RCBD), with mycorrhizal
118 fungi, rhizobium inoculation and control in the main plot, while genotypes were in the sub-plot (N-U-1, N-
119 U-8, WAB 56-104, OFADA GR and MOROBEREKAN) and the treatments were replicated thrice. There
120 were three main plots, each plot consist of 15 sub-plots with a size measurement of 2m x 1m and inter
121 sub-plot spacing of 0.5m in between plots. A total of 50 plants were planted per sub plot. Transplanted

122 seedlings were planted with the soil slurry into planting holes in the field at two seedlings per stand,
123 according to their respective plot at a spacing of 25cm x 25cm³.

124

125 **2.5. Source and Application of Planting Materials**

126 The different rice seeds (genotypes) were acquired from Africa Rice Centre, International Institute of
127 Tropical Agriculture, Ibadan (IITA). The genotypes collected are the recently released improved high
128 yielding genotypes for rice farmers in the study area. Cultured Arbuscular mycorrhizae fungi (*Glomus*
129 *intaradices*) inoculum with soil as carrier and cultured Rhizobium strains (RACA 3/5/12) were obtained
130 from the International Institute of Tropical Agriculture (IITA) Ibadan, Nigeria.

131

132 **2.6. Culture Media and Growth Condition**

133 The pure cultures of rhizobium strain RACA 3/5/12 was obtained from the (microbial technology culture
134 department), International Institute of Tropical Agriculture, Ibadan Nigeria (IITA). Rhizobium sp, RACA
135 was isolated from root nodules of cowpea, the strain were characterized by biochemical methods.
136 Rhizobium sp, RACA was maintained on yeast extract manitol agar medium. The culture were maintained
137 by periodic transfer and stored in the refrigerator. Photomicrography was used to examine roots of rice
138 genotypes inoculated with mycorrhizal fungi and rhizobium to view and ascertain root colonization by the
139 microbes.

140 **2.7. Mycorrhizal Infection Determination**

141 Portions of rice roots were collected, using a clean knife from all the replicates, in plastic bottles for the
142 mycorrhizal infection determination before the grinding and storage in 50% ethanol. Mycorrhizal fungi
143 staining in the roots was achieved by heating the root samples in 10% KOH, rinsing with distilled water
144 and soaking in 1% HCl for 10 minutes. Trypan blue solution was used to stain the roots. The roots were
145 soaked in the Trypan blue solution for 2 hours, and the stained roots were destained with 50% glycerol.
146 The grid-intersect method of [22] was used to evaluate the percentage of root infection. The data in
147 (Table 2) reveals maximum root colonization in rice genotypes treatment inoculated with introduced
148 mycorrhizal fungi (*Glomus intaradices*) (86%). It was also observed that rhizobium inoculated treatments
149 also recorded a (41%) root colonization by native mycorrhizae fungi while the un-inoculated treatments
150 recorded the lowest root colonization (10%) by native mycorrhizal fungi.

151 **Table 2.** Mycorrhizal fungi infection in roots of rice plants

152

Treatments	% Colonization
Mycorrhizal fungi	86
Rhizobium	41
Control (native mycorrhizal fungi colonization)	10

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156 2.8. Data Collection and Statistical analysis

157 Data collected were number of days to 90% maturity; Number of days to 50% flowering, Plant height at
158 Maturity; Number of primary tillers per plot; Number of grains per panicle; Number of panicles; Number of
159 filled and unfilled spikelet; Weight of 1,000 filled grains (g); Grain yield per plot (kg); this was taken by
160 converting the grain yield per plot into hectare using the formula ($[weight\ in\ grams/m^2] * 10$) [23]. The data
161 collected were statistically analyzed, all data were checked prior to statistical analysis for probable
162 violation of ANOVA assumption, and means were separated using Duncan multiple range test. SPSS 20th
163 edition statistical package was used for the analysis.

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165

166 3. RESULTS

167

168 3.1. Effects of mycorrhizal fungi and rhizobium inoculation on yield components of rice genotypes

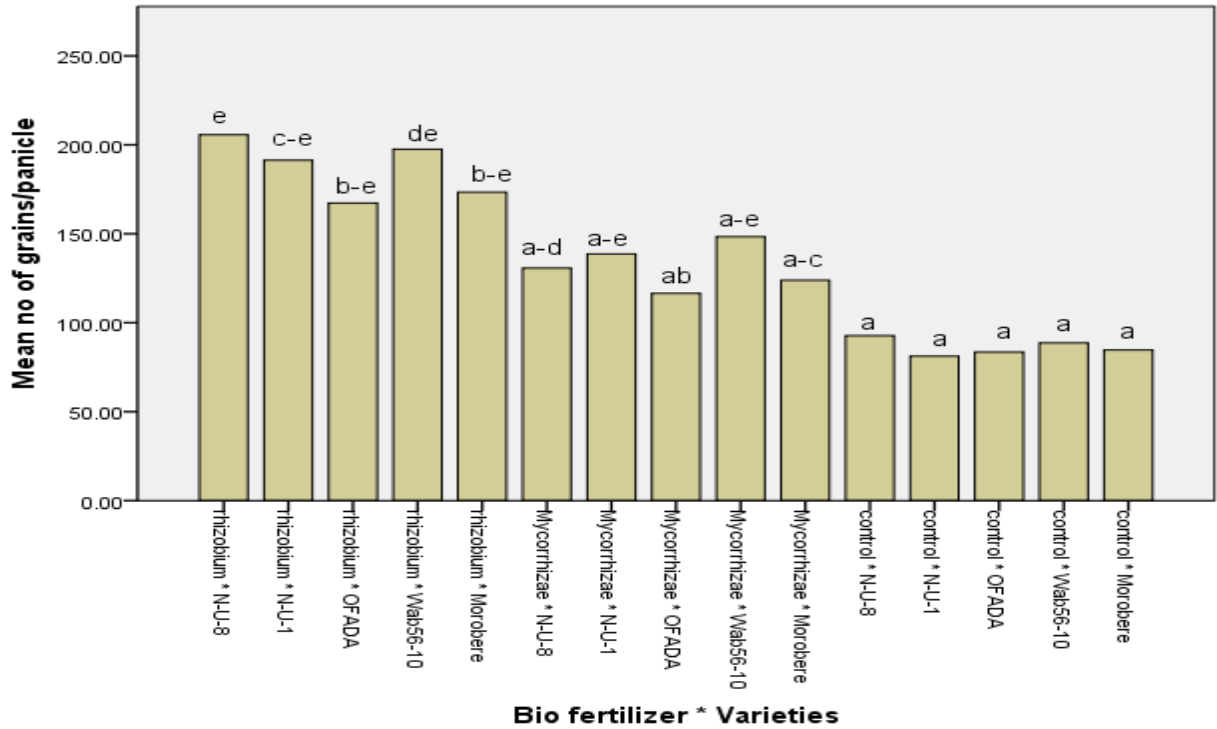
169 3.1.1. Plant Height at Maturity

170 The result presented in Table 3, indicate significant ($P < 0.05$) single effect of rhizobium and mycorrhizal
171 fungi inoculation on plant height at maturity. Rice genotypes inoculated with rhizobium recorded higher
172 plant heights (92.42cm) over the un-inoculated control (88.10cm). With respect to interactions between
173 rice genotypes and biofertilizer treatments, no significant ($P < 0.05$) interaction was observed in plant
174 height at maturity for both mycorrhized and rhizobium inoculated genotypes. However, rice genotype N-
175 U-8 recorded the lowest plant height while Moroberekan a local rice genotype was the tallest and had the
176 best response amongst all genotypes studied.

177

178 3.1.2. Number of grains per panicle

179 Significant interaction ($P < 0.05$) was observed between biofertilizers and rice genotypes (Fig.1),
180 biofertilizer inoculated genotypes had better performance when compared with the un-inoculated control
181 genotypes. Rice genotype (N-U-8) produced the highest number of grains per panicle (210) and genotype
182 (OFADA GR) the lowest number of grains per panicle (160) amongst rhizobium inoculated genotypes.
183 Rice genotype (WAB 56-104) produced the highest number of grains per panicle (150) amongst
184 mycorrhized genotypes. Single effect of mycorrhizal fungi and rhizobium inoculation on rice plants are
185 presented in (Table 3). Significant ($P < 0.05$) differences were observed with respect to number of grains
186 per panicle (Table 3). Rhizobium inoculated genotypes recorded the highest number of grains per panicle
187 187.05 and were significantly different from both mycorrhized genotypes (131.59) and the un-inoculated
188 control (86.15). Mycorrhized genotypes were significantly different with higher number of grains per
189 panicle recorded when compared with the un-inoculated control (Table 3).



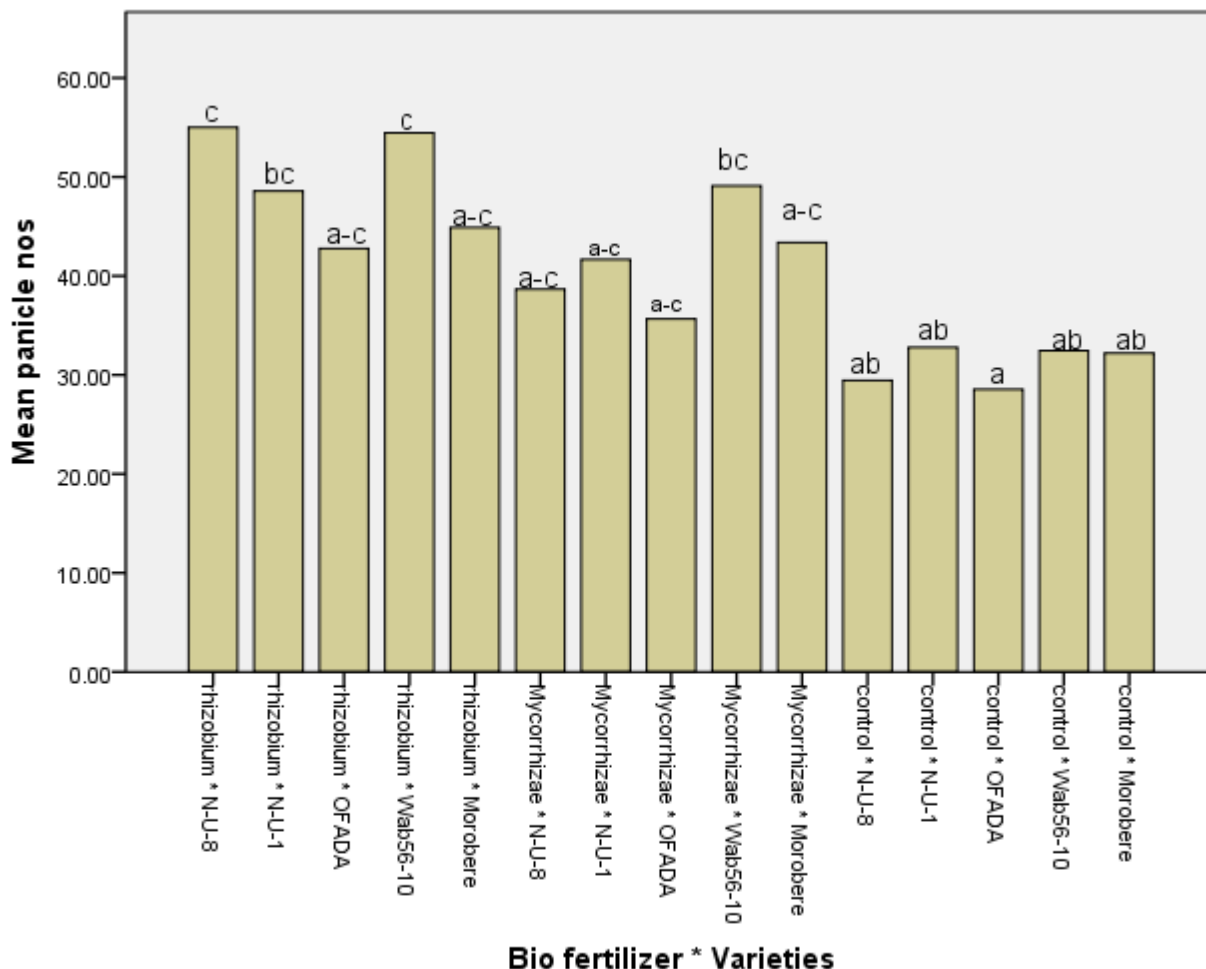
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191 **Figure 1:** Interactive effect of mycorrhizae and rhizobium inoculation on rice genotypes and number of
 192 grains per panicle. Standard error (P=0.05)

193

194 **3.1.3. Number of panicle**

195 There was no significant ($P < 0.05$) interaction observed between biofertilizer treatments and genotypes.
 196 However, rice genotype (N-U-8) and (WAB 56-104) produced the highest panicle number and genotype
 197 (OFADA GR) produced the lowest panicle number in both treatments respectively (Fig. 2). Significant
 198 ($P < 0.05$) differences were observed in total number of panicles produced by rice genotypes (Table 3).
 199 Rhizobium inoculated genotypes recorded the highest panicle number (49.14) closely followed by
 200 mycorrhized genotypes (41.70) while the un-inoculated genotypes produced the lowest panicle number
 201 (31.08) (Table 3).



202

203 **Figure 2:** Interactive effect of mycorrhizae and rhizobium inoculation on rice genotypes and number of
 204 panicles. Standard error (P=0.05)

205

206

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208 **3.1.4. Number of filled and unfilled spikelet**

209 Table 3, present the effect of rhizobium and mycorrhizae inoculation on number of filled spikelet and un-

210 filled spikelet produced by inoculated rice genotypes. Significant ($P<0.05$) differences were observed

211 amongst treatments, rhizobium inoculated genotypes had the highest filled spikelets (139.17) and un-

212 filled spikelet (47.86), mycorrhized inoculated genotypes also recorded high filled spikelet number

213 (102.10), it however produced lower un-filled spikelet number (29.63) than rhizobium inoculated

214 genotypes. The Un-inoculated genotypes recorded the lowest filled spikelet number (60.81) and un-filled

215 spikelet number (25.07). There was however no significant ($P<0.05$) interaction observed with respect to

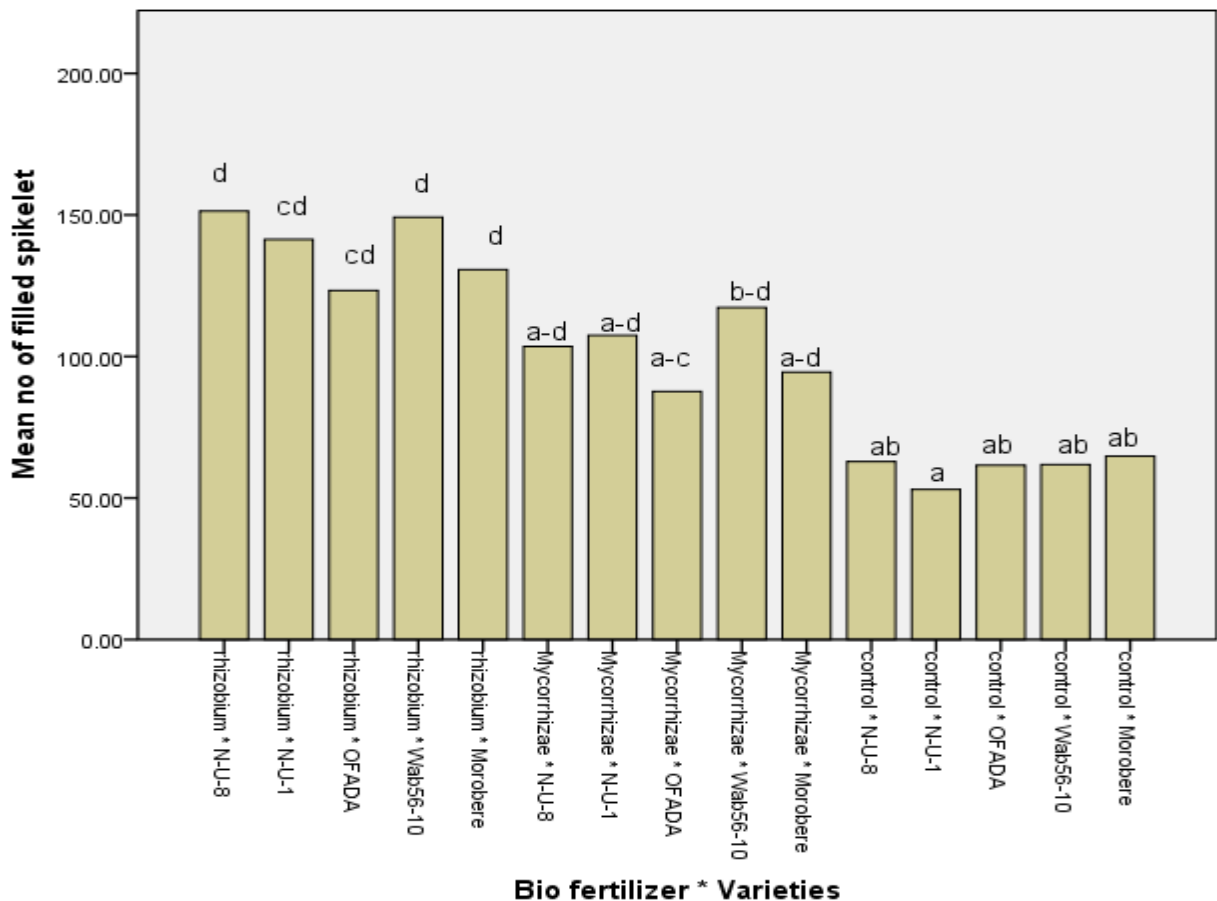
216 number of filled spikelet and un-filled spikelet in both mycorrhized and rhizobium inoculated genotypes

217 (Figs. 3 & 4). Rice genotype (N-U-8) produced the highest filled spikelet and genotype (OFADA GR)

218 produced the highest unfilled spikelet amongst rhizobium inoculated genotypes. Amongst mycorrhized

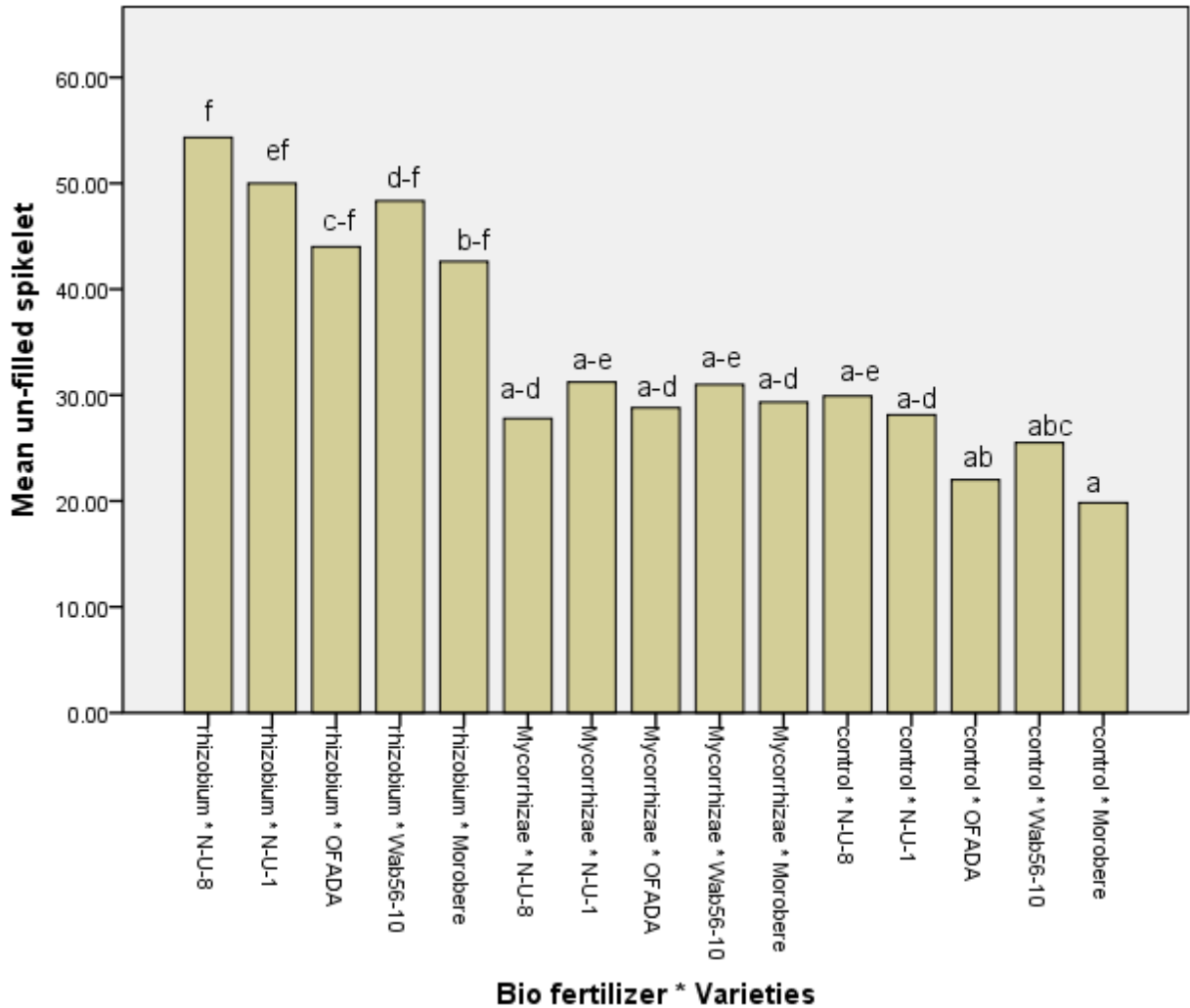
219 genotypes, no significant ($P < 0.05$) interaction was observed between biofertilizers and genotypes.
 220 However, rice genotype (WAB 56-104) recorded the highest number of filled spikelet, while genotype N-
 221 U-8 was recorded to have the highest number of unfilled spikelet.

222



223

224 **Figure 3:** Interactive effect of mycorrhizae and rhizobium inoculation on rice genotypes and number of
 225 filled spikelets. Standard error ($P=0.05$)
 226



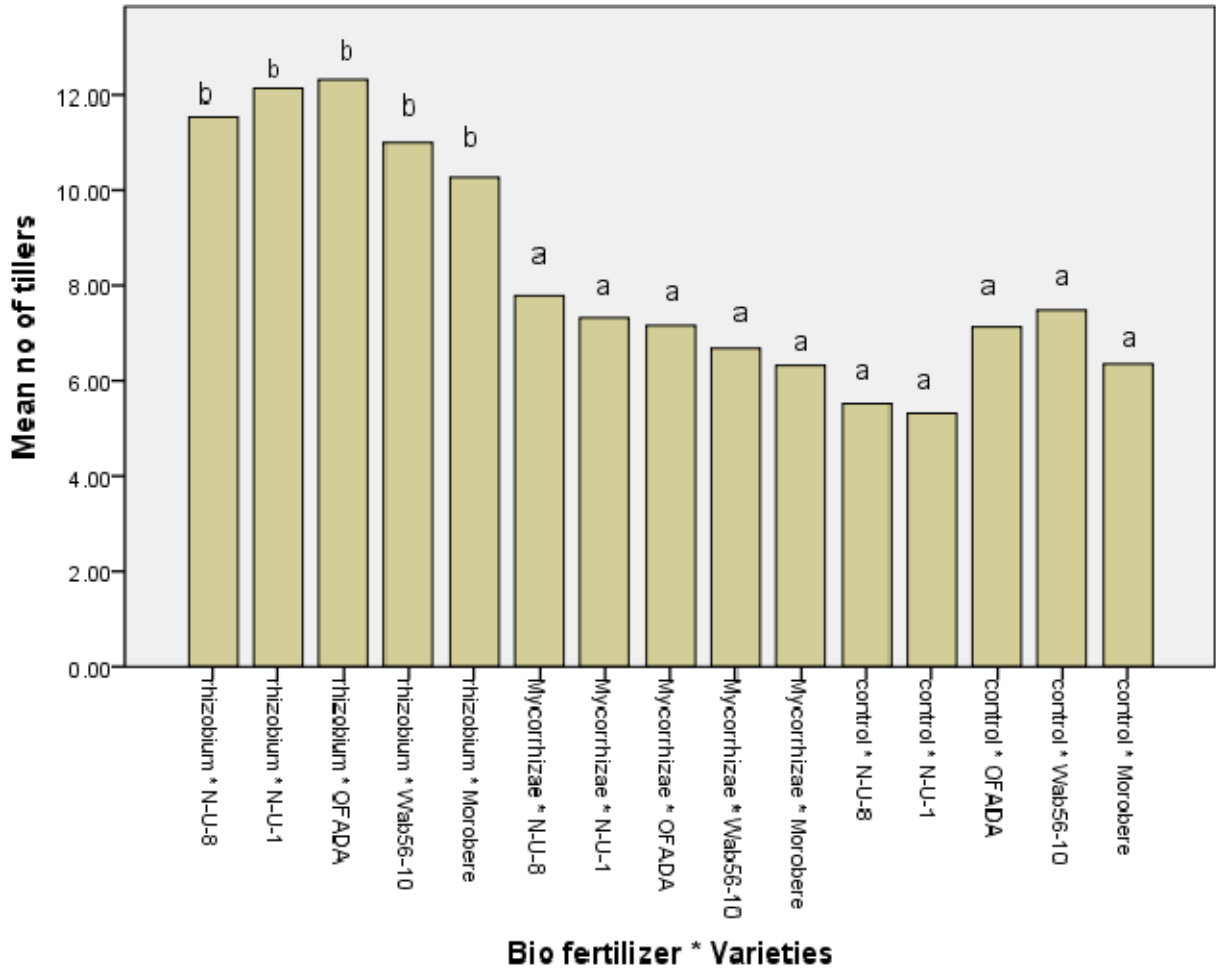
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228 **Figure 4:** Interactive effect of mycorrhizae and rhizobium inoculation on rice genotypes and number of
 229 unfilled spikelet. Standard error (P=0.05)

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231 **3.1.5. Number of primary tillers**

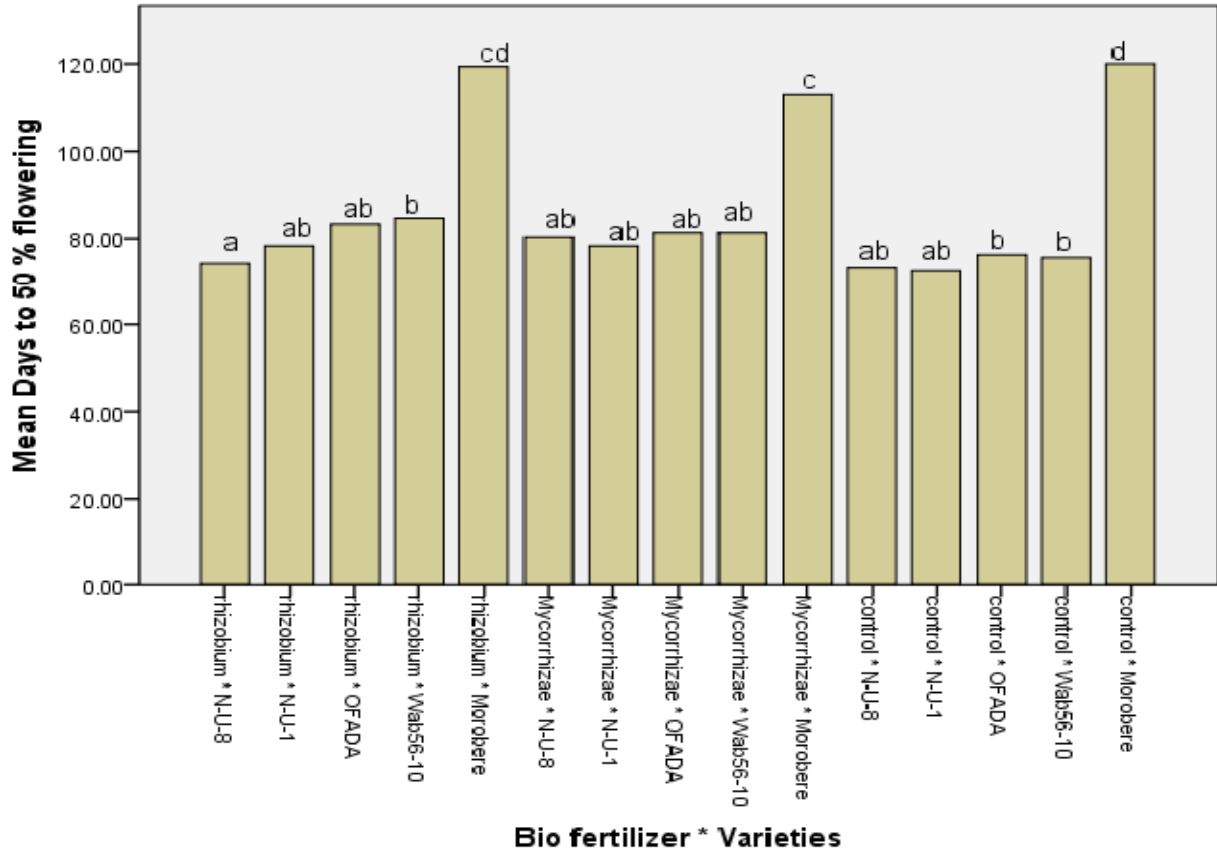
232 Significant ($P < 0.05$) differences were observed in treatments with respect to number of primary tillers
 233 produced by rice genotypes (Table 3). **Single effects of biofertilizer inoculation** indicate that rhizobium
 234 inoculated genotypes produced more tillers (11.46) when compared with un-inoculated genotypes (6.37).
 235 Mycorrhized genotypes were statistically significant and also produced more tillers (7.06) than the un-
 236 inoculated genotypes (Table 3). **Significant interactions were observed between biofertilizer inoculation**
 237 **and genotypes (Fig 5). Genotype OFADA GR inoculated with rhizobium produced more tillers amongst**
 238 **rhizobium inoculated genotypes and across all treatments.** Genotype N-U-8 recorded the highest number
 239 of tillers amongst mycorrhized genotypes while MOROBEREKAN recorded the lowest tiller numbers in
 240 both rhizobium inoculated and mycorrhized genotypes respectively (Fig. 5).



241
 242 **Figure 5:** Interactive effect of mycorrhizae and rhizobium inoculation on rice genotypes and number of
 243 primary tillers. Standard error ($P=0.05$)
 244

245 **3.1.6. Numbers of days to 50% flowering and 90% maturity**

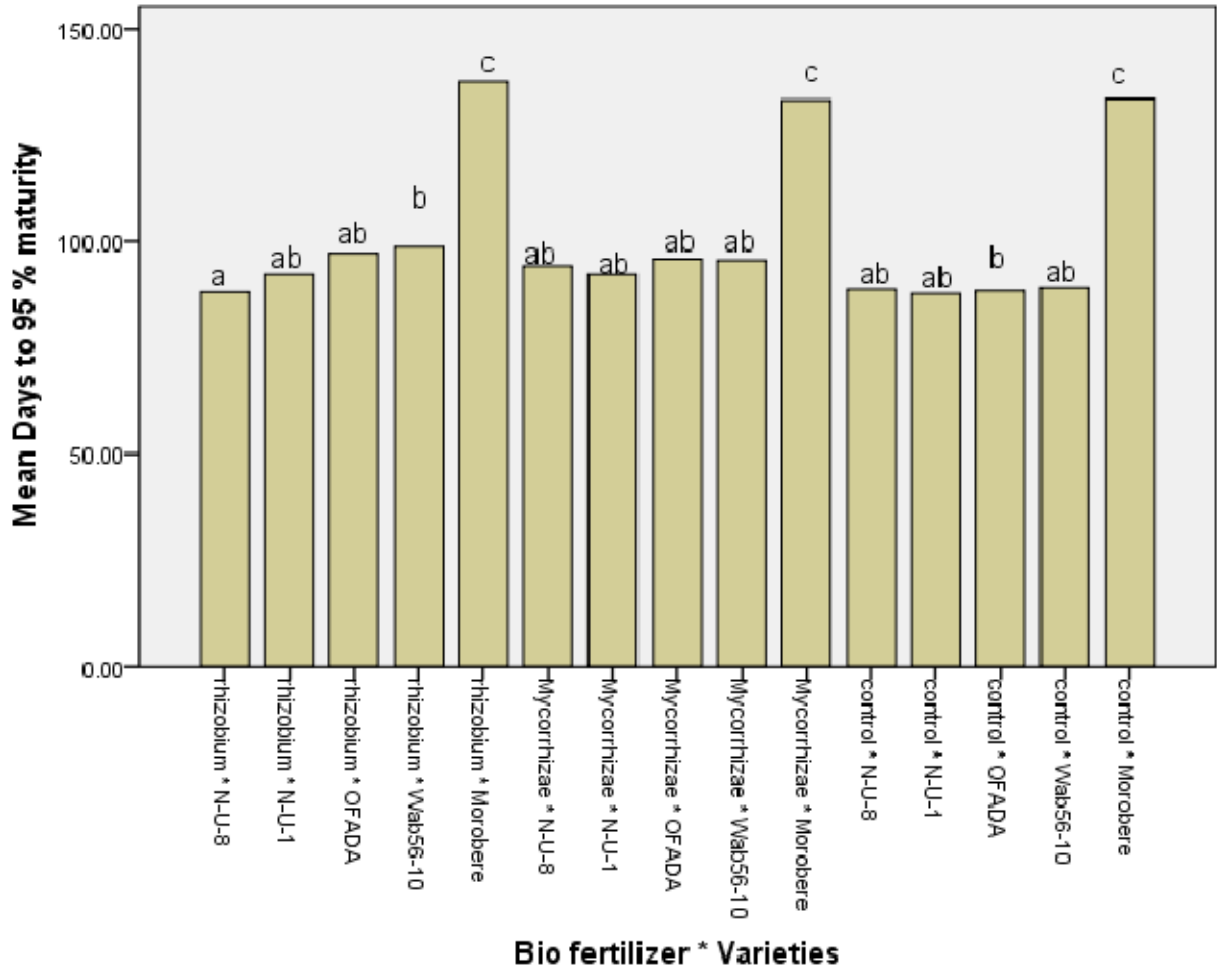
246 There was no significant ($P<0.05$) difference observed with respect to days-to-50% flowering and days-to-
 247 90% maturity amongst treatments (Table 3). However, results indicate that the un-inoculated treatment
 248 flowered and matured earlier (71.40 and 85.20 days respectively) than mycorrhizal treatment (86.87 and
 249 102.33 days) and rhizobium treatments (88 and 102.80 days respectively). Fig. 6 and 7 shows the
 250 significant interaction between biofertilizers and genotypes in reaching days to 50% flowering and days to
 251 95% maturity. Significant ($P<0.05$) interaction was only recorded in days to 50% flowering for both
 252 rhizobium inoculated and mycorrhized genotypes. Rice genotype (N-U-8) flowered the earliest amongst
 253 rhizobium inoculated genotypes, while genotype (MOROBEREKAN) flowered late. With respect to days to
 254 90% maturity, genotype N-U-8 also matured the earliest and MOROBEREKAN matured late. In
 255 mycorrhized genotypes, significant ($P<0.05$) interaction was observed with respect to days to flowering,
 256 N-U-1 flowered earlier while MOROBEREKAN flowered late.



257

258 **Figure 6:** Interactive effect of mycorrhizae and rhizobium inoculation on rice genotypes and days to 50%
 259 flowering. Standard error (P=0.05)

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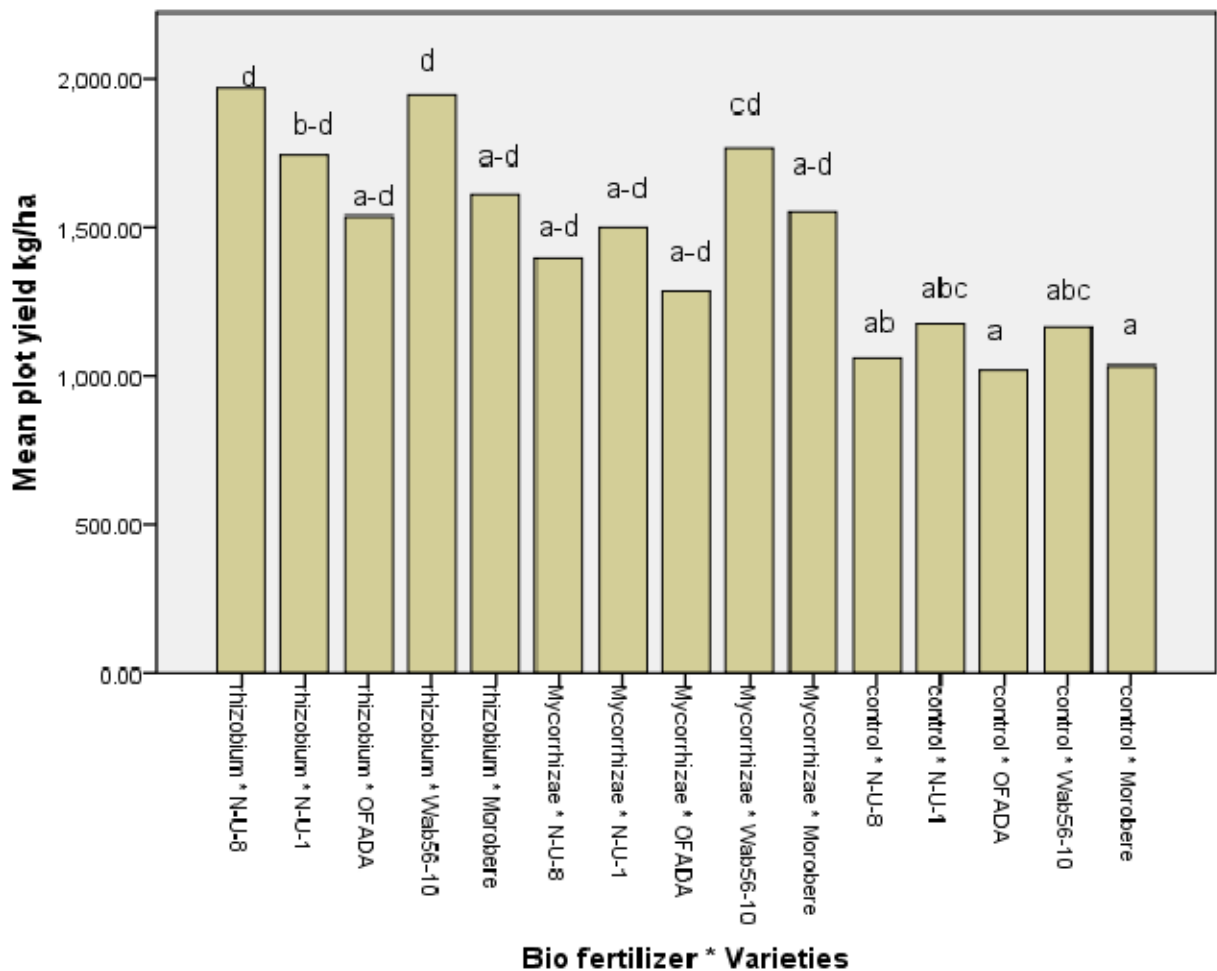


261
 262 **Figure 7:** Interactive effect of mycorrhizae and rhizobium inoculation on rice genotypes and days to 95%
 263 maturity. Standard error ($P=0.05$)
 264

265 **3.1.7. Grain yield and 1000 grain weight**

266 Table 3, shows the significant ($P<0.05$) effect of treatments on grain yield and weight of rice grain
 267 produced. Rhizobium inoculated treatments recorded higher grain yield than mycorrhizal treatment and
 268 the un-inoculated control treatment. Mycorrhizal treatments however had higher 1000 grain weight which
 269 wasn't significantly different from rhizobium treatment and the un-inoculated treatment. There was no
 270 significant ($P<0.05$) interaction observed in rhizobium inoculated genotypes with respect to grain yield
 271 (Fig 8). Rice genotype (N-U-8) produced the highest grain yield while genotype (OFADA GR) produced
 272 the lowest. With respect to 1000 grain weight, no significant ($P<0.05$) interaction was observed between
 273 treatment and genotypes however, genotype N-U-8 and N-U-1 produced the highest grain weight while
 274 genotype (MOROBEREKAN) produced the lowest grain weight. Amongst mycorrhized genotypes, no
 275 significant ($P<0.05$) interaction was observed with respect to grain yield. However, genotype (WAB 56-
 276 104) produced the highest grain yield. Significant ($P<0.05$) interaction was observed between

277 mycorrhizae and genotypes for 1000 grain weight. Rice genotype (N-U-1) recorded the highest 1000
 278 grain weight while genotype (WAB 56-104) weighed the lowest.



279
 280 **Figure 8:** Interactive effect of mycorrhizae and rhizobium inoculation on rice genotypes and grain yield.
 281 Standard error (P=0.05)
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 284
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286

287

Table 3. Single effect of Mycorrhizal fungi and Rhizobium inoculation on yield and yield components of rice

Treatments	Number of grains/panicle	Panicle Number	Number of filled spikelet	Number of un-filled spikelet	Plant height at maturity (cm)	Number of primary tillers	Number of days to 50% flowering	Number of days to 95% maturity	Grain yield (kg/ha)	1000 grain weight (g)
Control	86.15c	31.08c	60.81c	25.07c	88.10b	6.37c	71.40a	85.20b	1089.60c	27.1b
Rhizobium	187.05a	49.14a	139.17a	47.86a	92.42a	11.46a	88.00a	102.80a	1759.20a	29.6a
Mycorrhizae	131.59b	41.70b	102.10b	29.63b	86.18b	7.06b	86.87a	102.33a	1497.60b	30.1a

288 *Means with the same letter in same column are not significantly different from one another using Duncan Multiple Range Test (DMRT) (P = 0.05)

289 4. DISCUSSION

290
291 **Increase in yield** observed in some African rice genotypes in this study could be attributed to the positive
292 host-plant response to microbial inoculation, biological N fixation and production of plant growth
293 promoting hormones by introduced root colonizing organism. **Findings from our study are consistent with**
294 **field evaluations conducted in Israel and other semi-arid regions [24]; [25] on performance of cereals such**
295 **as wheat inoculated with biofertilizer (*Azospirillum* strain Cd) where significant increase in yield were**
296 **observed.** However, some cereal crop genotypes may portray significant differences in their ability to
297 associate with nitrogen fixing bacteria or mycorrhizal fungi. This assertion was observed in our study as
298 genotype WAB56–104 and N-U-8 had better association with rhizobium inoculant, while WAB56-104 and
299 MOROBEREKAN had better association with mycorrhizal fungi inoculants amongst all five genotypes
300 evaluated. **This finding was further corroborated by [26], who also observed better response, growth**
301 **promotion and an increase in dry plant weight and total N in grain of two select genotypes of sorghum,**
302 **inoculated with three different strains of *Azospirillum*.** Findings from our study indicate that development
303 and yield components of some African rice genotypes were significantly influenced by inoculation with
304 mycorrhizal fungi and rhizobium. These inoculated genotypes recorded higher statistical values over the
305 un-inoculated control, with the rhizobium inoculated rice genotypes recording a 61.4% increase in grain
306 yield over the un-inoculated rice genotypes. Our result established the effectiveness of the introduced
307 rhizobium strain in improving the development and yield of some NERICA lines and two other indigenous
308 genotypes used in the study. The increase in growth, development and yield parameters in response to
309 rhizobium inoculation endorsed the fact that they have one or more growth and yield promoting
310 mechanisms. However, [26] reported that rice plants inoculated with *A. lipoferum*, AI 121 and *A.*
311 *brasilense* did not influence rice growth or grain yield. A field study conducted by [27] in which rice plants
312 were inoculated with nitrogen fixing bacteria indicated that addition of low input of mineral N fertilizer
313 increased rice yield, nitrogen use efficiency and biological nitrogen fixation under flooded lowland
314 conditions. The increase in our studied characters could be ascribed to improvement in soil nutrient
315 availability and nutrient uptake due to the secretion of auxins or hormones and nitrogen fixation by
316 mycorrhizal fungi and bacteria inoculation [28]; [29]. **Findings from our** study are in agreement with [11]
317 who reported 16% increase in number of panicles and grains/panicle per plant of rice and suggested that
318 the improvement was due to increased availability of nutrients and phytohormones like indole acetic acid
319 and ethylene. **Our findings was also corroborated by studies conducted by Peng et al., 2002 [30] who**
320 **observed that inoculation of rice varieties with rhizobium enhanced their stomatal conductance which led**
321 **to an increased photosynthetic rates of about 12% and a 16% increase in grain yield at harvest. Their**
322 **study also observed a positive correlation between increased grain yield and photosynthetic rate without**
323 **nitrogen fertilizer application. Furthermore, [7] also reported that inoculation of rice with different**
324 **rhizobium strains such as *Rhizobium leguminosarum* bv. *trifolii* E11, *Rhizobium* sp. IRBG74 and**
325 ***Bradyrhizobium* sp. IRBG271 increased rice grain and straw yields by 8 to 22 and 4 to 19%, respectively,**
326 **at different nitrogen fertilizer rates and a further 10 - 28% increase in N, P and K and 15 - 64% increase in**

327 Fe uptake partitioned in the tested rice grain and straw. The increase in 1000 grain weight in **our study**
328 observed with inoculations with rhizobium and mycorrhizal fungi could be attributed to reduced spikelet
329 number produced by inoculated genotypes which consequently resulted in increased grain filling due to
330 adequate amount of photosynthetic material assimilated [31]; [32]. **Our result** also agrees with [33] who
331 observed up to 23.63% increase in developments of rice such as number of grains per panicle, filled
332 spikelets, panicle lengths and tillering over un-inoculated control and argued that indole acetic acid and
333 gibberellins production could be the key mechanism for that improvement. Maximum yield in inoculated
334 plants may be attributed to the symbiotic relationship of rhizobium (bacteria) with the roots of the plants,
335 which fixed atmospheric nitrogen into the roots of rice and thus the yield was increased. Early flowering
336 and maturity observed in the un-inoculated control than inoculated genotypes is suggested to be an
337 induced phenotypic response to limiting abiotic stress, such as moisture stress and high temperature.
338 Mycorrhizae inoculated genotypes was observed to have benefitted greatly through increased yield
339 component and also a 37.4% increase in grain yield. This positive influence on inoculated genotypes
340 could be attributed to increased phosphorus, nitrogen uptake, phytohormones such as cytokinins,
341 essential micro-nutrients e.g Fe, Zn, Cu by rice plants which lead to better development response and
342 yield. **Our result was also** in agreement with [5], who reported that inoculation of AMF resulted in
343 comparatively better performance in growth, development and yield of some selected drought tolerant
344 upland rice genotypes investigated in the rainforest transitory zone of Nigeria. However in the un-
345 inoculated control, where the soil was phosphorus and nitrogen deficient and no biofertilizer added the
346 plants grew poorly and yield was low. The potential benefit of exploiting this endophytic plant-bacterium
347 association for cereal production also extends to decreased environmental pollution and health risks
348 originating from excessive use of mineral N fertilizers to achieve high grain yield [6]. Finally the study has
349 demonstrated that the single use of rhizobium and arbuscular mycorrhizae fungi can enhance rice growth
350 and yield through changes induced in growth physiology and root morphology of rice genotypes. Further
351 studies are required to test this study across differing agro-ecologies and use of more genotypes and
352 different strains of rhizobium and mycorrhizae for efficient selection and appropriate recommendation.

353

354 **5. CONCLUSION**

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356 This study reveals that inoculation with biofertilizers resulted in comparatively better performance in
357 relation to yield components of African rice genotypes inoculated than the un-inoculated. The yield of
358 genotypes N-U-8, N-U-1, WAB56-104, OFADA Gr and MOROBEREKAN were statistically similar
359 irrespective of the different biofertilizer treatment applied. However, single rhizobium inoculated
360 genotypes had slightly marginal better performance over mycorrhizal inoculated genotypes. In rhizobium
361 inoculated genotypes, WAB56–104 and N-U-8 had the best response, while in mycorrhizal inoculated
362 genotypes, WAB56-104 and MOROBEREKAN recorded better response with respect to yield. **Results**
363 **from this study indicate that African rice genotypes differ in grain yield response and host specificity when**
364 **inoculated with mycorrhizal fungi and rhizobium inoculums. However, inoculating specific African rice**

365 genotypes with mycorrhizal fungi and rhizobium can positively influence their grain yield and yield
366 component development and this could play an important role in improving African rice productivity.
367 Authors acknowledge that the present study was a short term experiment which requires further field
368 studies for validation before recommendations and scientific inferences can be made. Furthermore, future
369 studies should be also be conducted to ascertain reported synergistic effect and performance of dual
370 inoculation of mycorrhizal fungi and rhizobium on rice growth and yield and their comparison with mineral
371 fertilized rice genotypes.

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