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Manuscript Draft

Manuscript Number: SBB7919R2

Title: Enhancing ecosystem services in sustainable agriculture: biofertilization and biofortification of chickpea (*Cicer arietinum* L.) by arbuscular mycorrhizal fungi

Article Type: Research Paper (FLA)

Keywords: arbuscular mycorrhizal (AM) fungal field inoculation, biofertilization, biofortification, functional diversity, *Funneliformis mosseae*, *Rhizophagus irregularis*, crop yield and quality improvement, local AM fungi (AMF), foreign AMF, sowing time.

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Manuscript Region of Origin: ITALY

September 25, 2013

Manuscript ID: SBB7919R1

Enhancing ecosystem services in sustainable agriculture: biofertilization and biofortification of chickpea (*Cicer arietinum* L.) by arbuscular mycorrhizal fungi.

Authors: Elisa Pellegrino, Stefano Bedini

Regular manuscript

Dear Editor,

We revised the manuscript following the reviewer suggestions and comments. We tested the PERMANOVA as multivariate method of data analysis alternative to RDA. Our data allowed both analyses and both the analyses gave the same picture. Therefore, we preferred to maintain the RDA analysis because it allows also to discriminate among the levels of the treatments. We hope that now our manuscript can be finally accepted for publication.

Yours sincerely,

Elisa Pellegrino

RESPONSE TO REVIEWER

REVIEWER: 1

1) We appreciate the comment of the reviewer about the choice of the multivariate data analysis method. We checked our dataset on the basis of his two main questions:

1. “A limitation of RDA is that it should be used when multiple response variables are normally distributed and if they are intercorrelated, they are correlated in a linear fashion. Is this true?”

- YES, our response variables were all log- or arcsen transformed to fulfil the assumption of normality. In addition, we did a Detrended Correspondence Analysis (DCA) for checking linearity or unimodality behaviour of the response variables (MS-rev: lines 240-244). This method is based on determining the length of gradients describing response variable variation and the results indicated a short variational gradients (< 4) and, hence that the linear relations are a good assumption for our response data (Leps & Smilauer 2003, p. 28).

2. “You state a concern that PERMANOVA may not be reliable if there are differences in multivariate dispersion (the multivariate equivalent of unequal variance). Do your data suffer from this problem?”

- NO, actually, our data do not suffer from differences in multivariate dispersion. We checked this by drawing a triplot (samples, response and environmental variables).

Taking into consideration that on the basis of the above answers (YES/NO) both analyses could have been done, we also performed a PERMANOVA and a PCO ordination (see Table and Figure below). Actually, both the analyses gave the same picture (see below Table and Figure). Therefore, we preferred to maintain the RDA analysis because it allows also the discrimination among the levels of the treatments.

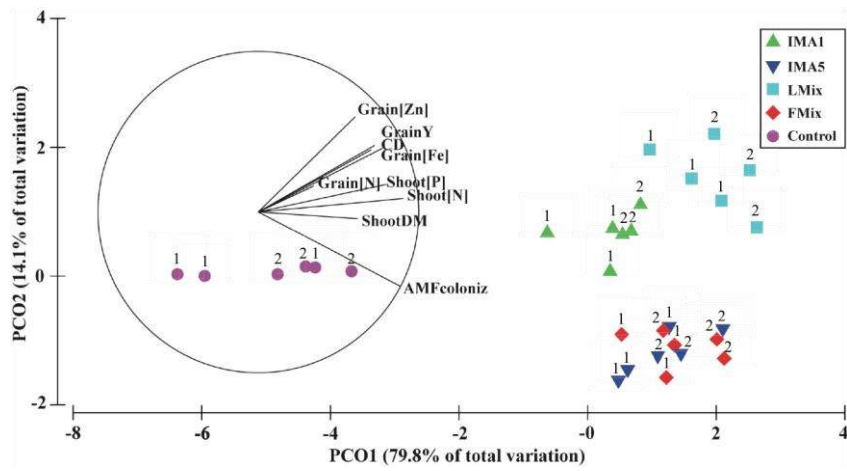
Results of PERMANOVA analyses on the effects, at harvest, of inoculation with arbuscular mycorrhizal (AM) fungi (AMF inoc), sowing time (S time) and their interactions on chickpea (*Cicer arietinum* L.) AM fungal colonisation, plant growth, yield, yield components, plant nutrient uptakes and grain biofortification.

Source of variation	df ^a	SS	MS	Pseudo- <i>F</i>	<i>P</i> (perm)
AMF inoc ^b	4	234.830	58.706	65.110	0.001
S time	1	8.326	8.326	9.235	0.002
AMF inoc x S time	4	2.063	0.516	0.572	0.748
Residual	20	18.033	0.902		
Total	29	263.250			

permutation).

^b Two-way PERMANOVA: AMF inoc, fixed factor; S time, fixed factor. AMF inoc: two single, foreign AMF (*Funneliformis mosseae*, IMA1; *Rhizophagus irregularis*, IMA5); a dual strain inoculum, foreign mixture (FMix): IMA1 + IMA5; a trap-culture-enriched locally-sourced AM fungal community (local mixture = LMix) and a mock inoculum as control; S time: autumn (October) and spring (March).

^d In bold statistically significant values ($P \leq 0.05$). Replicates field plots were three per treatment.



1= Autumn sowing; 2 = Spring sowing.

Minor comments

We changed through the text all instances "at autumn or spring sowing" to "in the autumn or spring sowing". All suggested minor changes were done.

1 **HIGHLIGHTS**

2

3 ▪ AMF field inoculation increased chickpea root colonization, yield & nutrient uptake

4 ▪ AMF improved chickpea nutritional value by protein, Fe & Zn grain biofortification

5 ▪ Local AMF were more effective than foreign for yield & grain N content

6 ▪ Local AMF inoculum was the most efficient in Fe and Zn grain biofortification

1 **Enhancing ecosystem services in sustainable agriculture: biofertilization and**
2 **biofortification of chickpea (*Cicer arietinum* L.) by arbuscular mycorrhizal fungi**

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16 **ABSTRACT**

17 Arbuscular mycorrhizal fungi (AMF) establishing beneficial symbiosis with most crop plants
18 have gained a growing interest as agro-ecosystem service providers able to sustain crop
19 productivity and quality. In this study we tested the agronomic relevance of field-inoculated
20 locally sourced and foreign inocula on chickpea (*Cicer arietinum* L.), one of the most
21 important worldwide grain legumes. The foreign AMF *Funneliformis mosseae* and
22 *Rhizophagus irregularis* were used as single and dual species inocula. Crop growth and
23 productivity, plant nutrient uptakes and protein, Fe and Zn grain biofortification were
24 assessed under a rainfed low-input cropping system after autumn and spring sowings. Uni-
25 and multivariate analyses of data showed that AM fungal field inoculation increased chickpea
26 AM fungal root colonization as well as plant biomass and yield. In addition, AMF were also
27 effective in improving the nutritional value of grain by protein, Fe and Zn biofortification.
28 The locally sourced AM fungal inoculum was more efficient than the foreign ones in Fe and
29 Zn grain biofortification and, in the spring sowing treatment, also in improving yield and
30 grain protein content. These findings enhance our understanding of the field potential role of
31 AMF showing that a mycorrhiza-friendly approach in agriculture may have great potential in
32 biofertilization of crops and biofortification of foods.

36 Key words:

37 arbuscular mycorrhizal (AM) fungal field inoculation, biofertilization, biofortification,
38 functional diversity, *Funneliformis mosseae*, *Rhizophagus irregularis*, crop yield and quality
39 improvement, local AM fungi (AMF), foreign AMF, sowing time.

41 **1. Introduction**

42
43 Over the last 20 years, low-input and organic agriculture has increased worldwide to
44 preserve agro-ecosystem functionality (Altieri, 1999; Lotter, 2003; Crowder et al., 2010;
45 Postma-Blaauw et al., 2010). The central pillar of such an agriculture is a systemic ‘holistic’
46 approach to cropping system management, which is based on the use of the ecosystem
47 services, in order to achieve sustainable yield and crop quality together with high energy
48 efficiency and low environmental impact (Pimentel et al., 2005; Moonen and Bàrberi, 2008).
49 In this view, soil microorganisms, such as arbuscular mycorrhizal (AM) fungi (AMF, phylum
50 *Glomeromycota*), representing a key interface between plant hosts and soil mineral nutrients,
51 have gained a growing interest as *ecosystem engineers* and *biofertilizers* (Gianinazzi et al.,
52 1990; Gianinazzi and Vosátka, 2004; Fitter et al., 2011).

53 AMF, which could represent 10% or more of the soil microbial biomass, establish a mutual
54 symbiosis with the majority (approx. 80%) of land plant species and agricultural crops (Smith
55 and Read, 2008). They supply mineral nutrients to the plants, mainly phosphate, in exchange
56 for photosynthetically fixed carbon (Bago et al., 2000; Hodge et al., 2010). As an effect of the
57 symbiosis, AMF improve agriculture productivity by enhancing plant growth (Koide, 1991),
58 seed production (Shumway and Koide, 1994) and protecting plants from root pathogenic
59 fungi (Newsham et al., 1995; Linderman, 2000) and drought (Augé, 2001). Moreover, AMF
60 have a direct effect on the ecosystem, driving the structure of plant communities (van der
61 Heijden et al., 1998a,b) and ameliorating the quality of soil by improving its aggregation and
62 organic carbon content (Miller and Jastrow, 1990; Rillig and Mummey, 2006; Bedini et al.,
63 2009).

64 Although the presence of AMF is widespread in agricultural soils, field experiments showed
65 that a further addition of AMF by inoculation can positively affect plant the root colonization

66 and increase the crop productivity (McGonigle, 1988; Lekberg and Koide, 2005; Lehmann et
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67 al., 2012).

68 Beside yield, AMF, due to their role in plant nutrition, could also enhance crop quality not
69 only by enrichment in macronutrients (i.e., N and P) (Karandashov and Bucher, 2005;
70 Veresoglou et al., 2012), but also in micronutrients (White and Broadley, 2009; He and Nara,
71 2007; Antunes et al., 2012). Micronutrient deficiency is a major issue affecting health of
72 billions of people all over the world. Actually, because of the intensity of crop production,
73 agricultural soils have become more and more depleted in micronutrients and, as a
74 consequence, yield mineral element contents are decreasing, compromising the nutritional
75 value of food. Indeed, biofortification, the increase of the concentrations and/or
76 bioavailability of mineral elements in produce, is considered a promising strategy for tackling
77 micronutrient malnutrition, especially in developing countries (White and Broadley, 2009).

78 AM fungal field inoculation studies have been mostly based on the use of selected, foreign
79 AM fungal isolates (Clarke and Mosse, 1981; Edathil et al., 1996; Meyer et al., 2005).
80 However, due to different affinities between host plants and AMF (van der Heijden et al.,
81 1998a; Klironomos, 2003; Munkvold et al., 2004; Avio et al., 2006), the use of a single AM
82 fungal strain is likely not optimal for all crops (Koomen et al., 1987; Jansa et al., 2008, 2009;
83 Smith et al., 2000; Koide, 2000; Maherali and Klironomos, 2007). Moreover, because
84 interactions among different AMF are not always synergistic (Koide, 2000; Jansa et al., 2008,
85 2009), AM fungal inocula have to be evaluated also in the field, where a local community is
86 present. As an alternative, the use of an inoculum based on locally sourced AMF may be a
87 suitable choice because of a better adaptation to the prevailing conditions (Lambert et al.,
88 1980) and also because they could avoid the ecological risks of the introduction of foreign
89 species (Schwartz et al., 2006). Actually, some studies showed higher or similar plant growth
90 and nutritional performances of locally sourced AMF compared to foreign selected ones

91 (Caravaca et al., 2003, 2005; Requena et al., 2001; Tchabi et al., 2010; Pellegrino et al.,
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2 92 2011a).

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4 93 In the present study we evaluated the effectiveness of the inoculation of locally sourced and
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6
7 94 foreign AMF on chickpea (*Cicer arietinum* L.), cultivated under a rainfed low-input system.
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10 95 This crop is one of the most ancient pulse crops domesticated in the Middle East around 7500
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12 96 years ago (Maiti and Wesche-Ebeling, 2001) and one among the five most important
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14 97 worldwide grain legumes with an annual production of about 9.4 million tons (FAOSTAT,
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17 98 2010). This crop is also largely cultivated in the Mediterranean area, where to benefit from
18
19 99 the rainfall season, autumn sowing has been proposed as an alternative to the traditional
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22 100 spring one (Frenkel et al., 2010).

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24 101 Despite several studies reported the beneficial effects of AMF on shoot concentrations of
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26 102 mineral elements, such as iron (Fe) and zinc (Zn) (Tarafdar and Rao, 1997; Al-Karaki, 2000;
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29 103 Ryan and Angus, 2003; Cavagnaro, 2008; Ortas, 2012), the effectiveness of AM fungal
30
31 104 inoculation on agricultural produce is still not resolved (Antunes et al., 2012). Starting from
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34 105 the evidence that chickpea positively responds to single foreign AM fungal inocula (Singh
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36 106 and Tilak, 1989; Weber et al., 1993; Clark and Zeto, 2000; Tufenkci et al., 2005), in the
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38
39 107 present study the field effectiveness of the inoculation of locally sourced AMF on chickpea
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41 108 (*Cicer arietinum* L.), cultivated under a rainfed low-input system was evaluated. The response
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44 109 of autumn and spring-sown chickpea in terms of growth, yield and nutritional profile has been
45
46 110 assessed and compared with the effects of two foreign strains of the AM fungal species
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48 111 *Funneliformis mosseae* and *Rhizophagus irregularis*, used as single or dual species inocula.
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52 53 113 **2. Material and methods**

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57 58 115 *2.1. Fungal and plant material*

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117 The AMF used were: *Funnelliformis mosseae* (T.H. Nicolson & Gerd.) C. Walker & A.
118 Schüßler, isolate IMA1 = BEG12 from UK (collector B. Mosse) and *Rhizophagus irregularis*
119 (N.C. Schenck & G.S. Sm.) C. Walker & A. Schüßler, isolate IMA5 from Italy (collector M.
120 Giovannetti) (new classification by Schüßler and Walker, 2010; the species were formerly
121 known as *Glomus mosseae* and *Glomus intraradices*), used as single or dual species inocula
122 (foreign mixture (FMix) = IMA1 + IMA5) and a locally sourced inoculum (LMix) consisting
123 of AMF originating from the field site. The trap-culture-enriched locally sourced AM fungal
124 inoculum was composed by: *Acaulospora cavernata* (syn. *Acaulospora scrobiculata*), *A.*
125 *spinosa*, *Acaulospora* spp. (syn. *A. rugosa*), *Diversispora spurca*, *Funnelliformis coronatum*
126 (syn. *Glomus coronatum*), *Claroideoglomus etunicatum* (syn. *Glomus etunicatum*),
127 *Funnelliformis geosporum* (syn. *Glomus geosporum*), *F. mosseae*, *Glomus* spp., *Rhizophagus*
128 *clarus* (syn. *Glomus clarus*), *R. irregularis*, *Scutellospora aurigloba* and *S. calospora* and
129 *Septoglomus viscosum* (syn. *Glomus viscosum*) (Pellegrino, 2007; Redecker et al., 2013). The
130 plant species used was chickpea (*Cicer arietinum* L.) cv. Sultano.

132 2.2. Experimental field site

134 The experiment was settled at the Rottaia Experimental Centre of the University of Pisa,
135 Italy (43°30'86"N - 10°19'00"E). The soil is a sandy loam (66.1% sand, 24.4% silt and 9.5%
136 clay) with 8.1 g kg⁻¹ soil organic carbon (Walkley-Black), pH(H₂O) of 8.4 and the following
137 total nutrient concentrations: 0.7 g kg⁻¹ N (Kjeldahl), 36.0 mg kg⁻¹ P and 14.6 mg kg⁻¹
138 available P (Olsen). Climatic conditions are typically Mediterranean with rainfall mainly
139 concentrated in autumn and spring (mean 948 mm year⁻¹) and mean monthly temperature
140 ranging from 11 °C in February to 30 °C in August (mean of 14.5 °C year⁻¹). Before
141 experimental setup, the field site has been uncultivated for about three years.

142

143 2.3. *Mycorrhizal potential of the experimental soil*

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145 Number and infectivity of AM fungal soil propagules of the experimental site were
146 evaluated by mycorrhizal index, AM fungal spore density and mycorrhizal infection potential
147 (MIP). Mycorrhizal index (calculated as percentage of AM fungal colonized root length) was
148 determined by examining the roots of 15 naturally occurring mycotrophic plants (*Bellis*
149 *perennis* L., $n = 3$; *Calendula arvensis* L., $n = 5$; *Daucus carota* L., $n = 5$; *Papaver rhoeas* L.
150 $n = 1$ and *Plantago lanceolata*, $n = 1$). The percentage of AM fungal colonization was
151 assessed, after root clearing and staining, using lactic acid instead of phenol (Phillips and
152 Hayman, 1970) by the gridline intersect method (Giovannetti and Mosse, 1980). The AM
153 fungal spores density was estimated in 50 g of soil from ten core samples by wet-sieving and
154 decanting, followed by sucrose centrifugation (Sieverding, 1991) and spore number was
155 assessed under a Wild dissecting microscope (Leica, Milano, Italy). The MIP was evaluated
156 as follows: *Lactuca sativa* L. seeds were sown in 50 mL sterile plastic tubes filled with 40 mL
157 of 15 soil samples taken up to a depth of 30 cm using a soil corer and air-dried. Six replicate
158 plastic tubes were used for each soil sample. After emergence, *L. sativa* plants were thinned to
159 three. After two week's growth, plants were removed from tubes and root systems were
160 cleared and stained as above, then mounted on microscope slides and examined under a
161 Reichert-Jung (Vienna, Austria) Polyvar microscope. Root length and AM fungal colonized
162 root length were measured using a grid eyepiece. Number of infection units (hyphal entry
163 points at the roots) and number of entry points were assessed at magnifications of x125-500
164 and verified at a magnification of x1250.

165

166 2.4. *AM fungal inoculum production*

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168 The AM fungal inocula required for the field inoculation were produced in 18 L pots filled
169 with soil and Terragreen (calcinated clay, OILDRI, Chicago, IL, USA) (1:1, by volume), as
170 potting substrate. The soil used for the potting substrate was collected at the Rottaia
171 Experimental Centre and was a sandy loam (54.5% sand, 30.1% silt and 15.3% clay) with
172 12.8 g kg⁻¹ soil organic carbon (Walkley-Black), pH(H₂O) of 8.0 and the following total
173 nutrient concentrations: 1.3 g kg⁻¹ N (Kjeldahl), 469.5 mg kg⁻¹ P, 14.6 mg kg⁻¹ available P
174 (Olsen) and 149.6 mg kg⁻¹ extractable K. The potting substrate was steam-sterilized (121° C
175 for 25 min, on two consecutive days) to kill naturally occurring AMF. The single species
176 inocula (IMA1 and IMA5) were produced by inoculating each pot with 500 g of crude
177 inoculum. The dual species foreign inoculum (FMix) was obtained by mixing equal quantities
178 of the two single species inocula (250 g IMA1 and 250 g IMA5). The crude inocula were
179 obtained from pot cultures maintained in the collection of the Soil Microbiology Laboratory
180 of the Department of Agriculture, Food and Environment, University of Pisa, Italy. The
181 locally sourced inoculum (LMix) was produced by inoculating the pots, containing the steam-
182 sterilized potting substrate of soil and Terragreen, with 500 g of soil from the Rottaia field
183 site. In addition a mock inoculum, free of AMF, was prepared to treat the control plots
184 (Control). The mock inoculum was produced by adding to the steam-sterilized potting
185 substrate 500 g of a sterilized mixture of equal quantities (about 170 g each) of each of the
186 two single species crude inocula (IMA1 and IMA5) and of the Rottaia field soil. The inocula
187 were produced in greenhouse using maize (*Zea mays* L.) as host plant (10 plants per each
188 pot). Finally, to ensure a common microflora, all pots received 1.5 L of soil filtrate, obtained
189 by filtering, through Whatman no. 1 filter paper, a mixture of each single species inoculum
190 and of the Rottaia field soil. Pots were supplied with deionized water (irrigation cycle: 4 days)
191 replaced, after two month's growth, by half-strength Hoagland's solution (Hoagland and
192 Snyder, 1933). After four month's growth, maize plants were harvested and the roots
193 removed from the pots. The roots and the potting substrate were air-dried. The roots were,

194 then, cut and mixed with the potting substrate and stored in polyethylene bags at 4° C until
195 field use. The MIP of the AM fungal inocula showed values of infection units ranging from
196 1.10 ± 0.34 to 2.89 ± 0.54 cm⁻¹ root length (mean \pm S.E.) in IMA5 and IMA1, respectively
197 (Pellegrino et al., 2011a).

2.5. Experimental set-up

201 A two-factor design was applied with the inoculum treatment (IMA1, IMA5, FMix, LMix
202 and the control) and the sowing time (autumn and spring sowing) as factors and three
203 replicate plots. Plots (2.5 m x 1.5 m) were dug (10 cm depth) and harrowed (5 cm depth) and
204 then inoculated with 5.3 kg plot⁻¹ of crude inoculum (mock inoculum for the control) along
205 the rows. Plots were sown on October 2004 (autumn sowing) ($n = 15$) and on March 2005
206 (spring sowing) ($n = 15$) with 40 x 10 cm row spacing and 10 cm border spacing (8 rows plot⁻¹)
207 in order to obtain a plant density of 25 plants m⁻². Three seeds were placed in each planting
208 position and, after the emergence, the seedlings were thinned to one. Chickpea plants were
209 harvested at the drying of seed pods on June 2005. During the crop cycle, plots were
210 manually kept weed-free.

2.6. Measurements

214 One month after emergence, the percentage of AM fungal root colonization was assessed,
215 after root clearing and staining, using lactic acid instead of phenol (Phillips and Hayman,
216 1970) by the gridline intersect method (Giovannetti and Mosse, 1980) on a random sample (n
217 = 6) of chickpea plants from each plot. At harvest, 10 plants from each replicate were cut at
218 ground level and air-dried. AM fungal root colonization, shoot dry matter, collar diameter and
219 grain yield were measured. Shoot and seed N and shoot P concentrations were assessed using

220 the Kjeldahl method and the sulphuric/perchloric acid digestion using the photometric method
221 (Jones et al., 1991), respectively. Fe and Zn seed concentrations were detected by inductively
222 coupled plasma-mass spectrometric (ICP-MS) analysis. The acid mineralization was done on
223 0.5 g dry samples. Samples were digested for 10 min into a microwave-digesting machine
224 with a solution composed by 6 ml HNO₃ 65% and 1 ml H₂O₂ 33%. The resulting solutions
225 were filtered and diluted with ultra-pure water to a 25 ml volume and then analyzed with ICP-
226 MS analysis.

227

228 *2.7. Statistics and data analyses*

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230 Data were analyzed by two-way (inoculum treatment and sowing time as fixed factors) or
231 one-way ANOVA (inoculum treatment as factor). Data were ln- and arcsine-transformed
232 when needed to fulfill the assumptions of the ANOVA, which was carried out according to
233 the completely randomized design. Multiple comparisons within the one-way ANOVAs were
234 done with orthogonal contrasts. All the analyses were performed by the SPSS 17.0 software
235 (SPSS Inc., Chicago, IL, USA). Means and standard errors (S.E.) given in tables are for
236 untransformed data.

237 To test the null hypotheses about differences between groups on the basis of multiple
238 response variables, along with the univariate methods, we also analyzed the data set with a
239 multivariate constrained approach. Redundancy analyses (RDA) (van den Wollenberg, 1977)
240 were used to investigate, for each sowing time, the influence of the different inoculation
241 treatments (used as explanatory variables) on plant growth, yield, plant nutrient uptake, grain
242 biofortification variables and AM fungal colonization (used as response variables). Since the
243 response variables were in different measurement units and the length of the gradient of the
244 detrended correspondence analysis (DCA) was lower than four, we utilized the RDA linear
245 method (Lepš and Šmilauer, 2003). All data were log- or arcsen-transformed, centered and

246 standardized by the response variables. Monte Carlo permutation tests were performed using
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2 247 499 random permutations (unrestricted permutation) in order to determine the statistical
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4 248 significance of the relations between the whole set of inoculation treatments and the response
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7 249 variables. RDA analyses were done by Canoco for Windows v. 4.5 (ter Braak and Šmilauer,
8
9
10 250 2002). The biplots were drawn by CanoDraw for Windows.

14 252 **3. Results**

19 254 *3.1. Mycorrhizal potential of the experimental soil and chickpea AM fungal root colonization*

24 256 In the experimental soil, AM fungal root colonization of the selected plant species was $7.1 \pm$
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26 257 1.6% , spore density was 3.1 ± 0.4 spores g^{-1} soil, while the MIP test values were: root
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29 258 colonization, $1.3 \pm 0.5\%$; infection units, 0.27 ± 0.07 cm^{-1} root length; entry points, $0.13 \pm$
30
31 259 0.03 cm^{-1} root length.

34 260 One month after emergence, the percentages of colonized root length of control plants in the
35
36 261 autumn and spring sowing treatments were 10.1 ± 1.1 and $12.6 \pm 0.6\%$, respectively, and
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38
39 262 lower than those of mycorrhizal treatments that ranged, in the autumn sowing, from $21.0 \pm$
40
41 263 1.1% to $33.7 \pm 8.4\%$ in IMA5 and FMix, respectively, and, in the spring sowing, from $25.2 \pm$
42
43 264 0.5% to $40.9 \pm 2.3\%$ in IMA5 and IMA1, respectively (data not shown).

46 265 At harvest, mycorrhizal root colonization was significantly affected by AM fungal
47
48 266 inoculation (Table 1). In the autumn sowing treatment, the degree of root colonization ranged
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50
51 267 from 20.4% to 69.7% , while in the spring sowing treatment from 25.1% to 71.7% in control
52
53 268 and FMix, respectively (Fig. 1a). At both sowing times, significant differences were also
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55
56 269 observed between IMA1 and IMA5 and between FMix and LMix (Table 1). In detail,
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58 270 orthogonal contrasts showed that FMix consistently colonized chickpea more than LMix and
59
60 271 single strain inocula (Table 1).

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273 3.3. Plant growth

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7 275 One month after emergence no differences among treatments were observed in root (data
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9 276 not shown) and shoot dry matter. In the autumn sowing treatment, shoot dry matter ranged
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12 277 from 0.29 ± 0.03 to 0.36 ± 0.01 g plant⁻¹ in IMA5 and control, respectively, while in the
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14 278 spring one it ranged from 0.32 ± 0.02 to 0.41 ± 0.07 g plant⁻¹ in IMA5 and in IMA1,
15
16 279 respectively.

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18
19 280 At harvest, shoot dry matter of spring-sown plants was significantly affected by AM fungal
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21 281 inoculation (Table 1). In the spring sowing treatment, inoculated chickpea showed in average
22
23
24 282 25% higher plant biomass than control (Fig. 1b). In both sowing times, larger plant collar
25
26 283 diameters were observed in AM fungal treatments with respect to controls (Table 1). In
27
28
29 284 addition, in the spring sowing treatment, LMix inoculated plants showed larger collar
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31 285 diameter than the FMix ones (Table 1; Fig. 1c).

32

33
34 286 In the spring sowing, shoot dry matter per plant showed a strong correlation with the
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36 287 percentage of AM fungal colonized root length ($R = 0.724$, $F_{1,13} = 14.326$, $P = 0.002$).

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40 289 3.4. Yield

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46 291 As regards yield, in both sowing times, AM fungal inoculation consistently induced better
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48 292 plant responses compared with control (Table 1). Host benefit, calculated, in both sowing
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51 293 times, for each AM fungal inoculation treatment as $[(\text{mycorrhizal treatment} -$
52
53 294 $\text{control})/\text{control}] \times 100$, was 38%, 52%, 69% and 79% in the autumn sowing treatment and
54
55 295 41%, 52%, 86% and 93% in the spring one for IMA5, FMix, IMA1 and LMix, respectively.

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57
58 296 In the spring sowing, LMix chickpea showed higher values of grain yield per plant respect to
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60 297 FMix (Table 1). In detail, LMix inoculated plants showed 27% higher grain yield than the

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298 FMix ones (Fig. 1d). Moreover, in the spring sowing treatment a significant better
299 performance of IMA1 when compared to IMA5 was also observed (32%) (Table 1; Fig. 1d).

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301 *3.5. Nutrient uptake and grain biofortification*

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303 In both sowing times, N and P uptakes, evaluated by shoot nutrient concentrations, were
304 affected by AM fungal inoculation (Table 2). On the basis of N and P shoot concentrations, in
305 the autumn sowing treatment host benefits were on average 106% and 48%, respectively,
306 while in the spring sowing treatment were 110% and 45%, respectively. No significant
307 difference was observed among inoculated treatments, except for shoot N concentration
308 between IMA1 and IMA5 (Table 2). Interestingly, in both sowing times, N and P shoot
309 concentrations were mostly strongly correlated with the percentage of colonized root length
310 (autumn sowing: shoot N and P concentrations $R = 0.781$, $F_{1,13} = 20.310$, $P = 0.001$ and $R =$
311 0.678 , $F_{1,13} = 11.031$, $P = 0.006$, respectively; spring sowing: shoot N and P concentrations R
312 $= 0.793$, $F_{1,13} = 22.042$, $P < 0.001$ and $R = 0.589$, $F_{1,13} = 6.904$, $P = 0.021$, respectively).

313 As regards grain, N concentrations in the spring sowing treatment were affected by AM
314 fungal inoculation (Table 2), with a host benefit of 6 % in average. Interestingly, lower grain
315 N concentrations were observed in FMix in comparison with single AM fungal inoculated
316 plants (Table 2).

317 Fe and Zn seed concentrations were significantly higher in inoculated plants than controls,
318 both in the autumn and spring sowing treatments (Table 3). In detail, Fe and Zn increases
319 were of 5% and 16%, respectively. Interestingly, LMix induced higher concentrations of the
320 analyzed micronutrients in comparison with FMix. These increases, considering both sowing
321 times, were of 4% and 21% for Fe and Zn uptakes, respectively. LMix resulted the most
322 efficient inoculum for biofortification, raising the micronutrient level of about 8% and 36%
323 respect to the control for Fe and Zn, respectively (Table 3).

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2 325 *3.4. Effect of sowing time*
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7 327 The two-way ANOVAs showed no differences in AM fungal root colonization between
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9 328 sowing times (S time) or interaction between AM fungal inoculation (AMF inoc) and S time,
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11 329 both one month after emergence ($P = 0.738$ and $P = 0.483$, respectively) and at harvest (Table
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13 330 4). No differences between S times or interactions were detected in shoot and root dry weights
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15 331 one month after emergence (data not shown).
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19 332 By contrast, at harvest, shoot dry matter and collar diameter were significantly affected by
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21 333 the sowing time (Table 4; Fig. 1b,c). No differences in shoot and grain nutrient uptake
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23 334 between S times and no interaction between AMF inoc and S time were detected (Table 5).
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25 335 Interestingly, grain Fe concentrations were significantly affected by S time, showing higher
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27 336 values in the spring sowing treatment than in the autumn one (Tables 3, 5).
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34 338 *3.5. Main patterns of chickpea traits as affected by AMF*
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39 340 RDA, in line with the univariate tests, showed that AM fungal inoculation explained 62.9%
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41 341 and 69.4% of the whole variance in the autumn and spring sowing treatments, respectively,
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43 342 and that its effect on the response variables was significant ($P = 0.002$). In both sowing
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45 343 treatments, the Monte Carlo permutation test showed that control was significantly different
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47 344 from the other treatments ($P = 0.002$) and LMix from the FMix ($P = 0.002$). In addition, in
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49 345 the spring sowing, we also observed significant differences between IMA1 and IMA5/FMix
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51 346 ($P = 0.008$). The biplots also show that collar diameter, grain yield, shoot P concentration
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53 347 were the most discriminating variables between AM fungal treatments and the control in the
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55 348 autumn sowing treatment (Fig. 2a), whereas grain yield, shoot dry matter and collar diameter
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57 349 were the most discriminating in the spring one (Fig. 2b). The biplots of the autumn and spring
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350 sowing treatment also show that grain Zn and Fe concentrations and AM fungal colonization
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2 351 were highly discriminative between LMix and the other AM fungal treatments (Fig. 2a,b).

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6 7 353 **4. Discussion**

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12 355 In the present study data showed that: (i) AM fungal field inoculation increased chickpea
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14 356 AM fungal root colonization as well as yield and plant nutrient uptake; (ii) AM fungal
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17 357 inoculation was effective in improving the nutritional value of chickpea grain by protein, Fe
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19 358 and Zn grain biofortification; (iii) locally sourced AM fungal inoculum was the most efficient
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22 359 in Fe and Zn grain biofortification; (v) in the spring sowing treatment, local AM fungal
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24 360 inoculation was more effective than foreign inocula in improving yield and grain N content.

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27 28 29 362 *4.1. Mycorrhizal potential of the experimental soil and chickpea AM fungal root colonization*

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34 364 All AM fungal inocula promoted a rapid increase of the chickpea root colonization that,
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36 365 already one month after emergence, was higher compared to control plots. Since AM fungal
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39 366 inoculation success is expected to be negatively related to the amount of the active AM fungal
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41 367 propagules already present in the soil (Abbott and Robson, 1982, 1991; Gianinazzi and
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44 368 Vosátka, 2004), the higher AM fungal colonization observed in the inoculated plants
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46 369 compared with controls indicated that the mycorrhizal infection potential (MIP) of the soil at
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49 370 the beginning of the experiment was low or sub-optimal. These low MIP values, in line with
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51 371 the those observed in other agricultural soils (Purin et al., 2006; Di Bene et al., 2011;
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53 372 Pellegrino et al., 2011a; 2012; Di Bene et al., 2013; Gosling et al., 2006; Bedini et al., 2013),
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56 373 confirm the detrimental effects of agricultural practices on AM fungal infectivity (Kabir et al.,
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58 374 2005; Plenchette et al., 2005). In addition, the success of the inoculation in terms of AM
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2 375 fungal root colonization indicates that the MIP assay is appropriate as preliminary test in the
3 376 planning of an effective AM fungal inoculation treatment.

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5 377 At harvest, AM fungal root colonization of the inoculated chickpea was up to three-fold
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7 378 respect to the controls. Such colonization increases are in line with previous trials on field
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9 379 AM fungal inoculation of chickpea (Singh and Tilak, 1989; Weber et al., 1993; Saini et al.,
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11 380 2004) and of other crop plants such as lucerne (*Medicago sativa* L.), sorghum (*Sorghum*
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13 381 *bicolor* L.) and white clover (*Trifolium repens* L.) (Hayman and Mosse, 1979; Bagyaray et
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15 382 al., 1979; Rangeley et al., 1982; Hayman, 1984; Pellegrino et al., 2011a; 2012) and strongly
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17 383 support that the colonization of mycorrhizotrophic crops can be effectively improved by
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19 384 field inoculation. Although, all the inocula were effective in raising the level of AM fungal
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21 385 colonization of chickpea, at harvest, we observed differences between the two foreign isolates
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23 386 when individually inoculated. Actually, the higher colonization by IMA5 compared to IMA1
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25 387 is consistent with previous observations both in microcosm (Avio et al., 2006) and field
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27 388 conditions (Pellegrino et al., 2011a) and confirms *R. irregularis* (IMA5) as a good competitor
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29 389 for field inoculation (Alkan et al., 2006; Douds et al., 2011).

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31 390 Interestingly, the linear orthogonal contrast analysis indicated that IMA1 and IMA5 were
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33 391 more efficient in boosting AM fungal colonization when used as dual than as single inocula,
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35 392 suggesting an additive or synergistic effect. However, since the level of root colonization of
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37 393 IMA5 is almost the same of FMix, such an effect is probably due to a dominance of IMA5
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39 394 respect to IMA1. Moreover, the higher root colonization of IMA1/IMA5 dual species respect
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41 395 to the locally sourced inoculum confirms that selected AM fungal species may be more
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43 396 effective in the establishment, survival and spreading of the symbiosis within roots of crops
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45 397 (Sýkorová et al., 2012). However, it should be taken in account that although the colonization
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47 398 effectiveness could be a positive trait for commercial inocula, a high aggressiveness of
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49 399 foreign AMF could determine shifts in composition, structure and functionality of the whole
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400 local soil microbial community (Mummey et al., 2009; Koch et al., 2011; Sýkorova et al.,
401 2012; Veresoglou et al., 2012).

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403 *4.2. Plant growth*

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405 Contrary to our expectation, spring-sown chickpea showed higher plant growth parameters
406 in comparison with the autumn-sown one. In Mediterranean areas, autumn sowing is
407 considered more convenient because the winter rainy season may assure a better crop growth
408 and yield (Singh, 1997). However, especially in the northern part of the Mediterranean areas,
409 it may also result in a higher exposure to the cold and to pathogen diseases such as aschochyta
410 blight that could cause loss of productivity (Nene, 1981). Actually, we observed differences
411 between the two sowing date on the aboveground chickpea biomass that was effectively
412 increased by AM fungal field inoculation only in spring-sown chickpea. AM fungal
413 inoculation resulted much less effective in the autumn sowing treatment respect to spring one
414 (8% vs 25%, respectively) where, inoculation benefits, were consistent with previous
415 chickpea field inoculation trials (Singh and Tilak, 1989; Weber et al., 1993; Zaidi et al.,
416 2003).

417 It is also should be noted that the chickpea AM fungal benefit was lower than what was
418 observed in other pulses such as Egyptian clover and lucerne (mean increases of 93% and
419 83%, respectively) (Pellegrino et al., 2011a, 2012). Taking into account the main role of AMF
420 in plant P nutrition, this lower responsiveness may be due to chickpea minor P requirements
421 (30-80 Kg ha⁻¹ P₂O₅; Foti e Abbate, 2000; Saccardo et al., 2001) in comparison with other
422 fodders (about 120 Kg ha⁻¹ P₂O₅ per year; Masoni et al., 1993). Similarly, the lack of any
423 significant effect on biomass one month after plant emergence may still depend on the large
424 chickpea seed reserves, which could sustain the early growth of the plant (Weber et al., 1993)
425 and thus masking the effect of AM symbiosis.

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426 Despite observing significant differences in root AM fungal colonization, no interspecific
427 differences in plant biomass were detected between IMA1 and IMA5. This may depend on
428 the effects of co-occurring soil AMF as well as on the functional diversity of the two isolates
429 (Klironomos, 2000, 2003; Smith et al., 2004).

430 In line with plant biomass benefits, collar diameter responded positively to AM fungal
431 inoculation. Collar diameter was strongly correlated with N and P uptakes. Since no
432 correlations were observed with shoot dry matter (data not shown), the increase of the collar
433 diameter appears to be associated to a better crop nutritional status of the inoculated plants.
434 Considering that collar diameter is a key trait for plant stress resistance, we can argue that
435 AM fungal inoculation might lead to a stronger resistance of the crop and to lower yield
436 losses due to plant laid down and/or to pathogen attacks.

437

438 4.3. Yield

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440 Overall, chickpea productivity was in line with the mean production reported in previous
441 works (Singh et al., 1989; Weber et al., 1993; Zaidi et al., 2003). In this trial, AM fungal field
442 inoculation was very effective in improving chickpea yield, with increases of about 75% in
443 comparison with the controls. Such increase was much higher of that obtained by Singh and
444 Tilak (1989), who, using *Glomus versiforme* as field inoculum, observed increases of about
445 11% and in contrast with Weber et al. (1993) and Zaidi et al. (2003) who did not register any
446 yield difference. However, similar large benefits were observed for *Trifolium alexandrinum* in
447 the same pedo-climatic conditions by Pellegrino et al. (2011a), who reported seed production
448 increments of about 77% with the same AM fungal isolates and by Saini et al. (2004), who
449 obtained large yield chickpea increases (109%) using a dual inoculum of AMF and rhizobia.
450 This variability on yield benefit may depend on differences in plant-fungal genotype
451 compatibilities (van der Heijden and Sanders, 2002) or on different climatic and soil

452 conditions, which may also include soil mycorrhizal infection potential and rhizobia
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2 453 populations. However, even if the actual impact of AMF on the productivity of agricultural
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4 454 systems seems to be variable, our results indicate that, in the appropriate pedo-climatic
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7 455 conditions and agronomical managements, AM fungal field inoculation can be very effective
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10 456 in improving the yield of grain legumes. This indicates also that, although the modern
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12 457 cultivars of some of the most important crops, such as wheat, seem to be much less
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14 458 responsive to AMF compared to chickpea and other pulses (Ryan and Kirkegaard, 2012), a
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17 459 redesign of agricultural systems in order to enhance the benefits of AM fungal ecosystem
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19 460 services, even considering the additional cost of the inocula, can result in an improved plant
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22 461 growth and yield that could benefit crop economics, in particular in organic and low-input
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24 462 farming systems.

26 463 It is noteworthy that we observed better overall yield performances of chickpea inoculated
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29 464 with the local inoculum compared to the foreign. In this regard, local AMF may be preferable
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31 465 for field inoculation and on-farm inoculum production because of their better adaptation to
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34 466 local prevailing conditions (Dodd et al., 1983; Requena et al., 2001; Caravaca et al., 2003;
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36 467 Douds et al., 2011). Moreover, foreign AM fungal inocula may have negative ecological
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39 468 consequences interfering with the local microbial communities and thus altering agro-
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41 469 ecosystem functions (Schwartz et al., 2006; van der Heijden et al., 2008).

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46 471 *4.4. Nutrient uptake and grain biofortification*

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51 473 Inoculated chickpea plants showed a higher shoot N and P concentration when compared to
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53 474 the controls. A better nutritional status of AM fungal inoculated chickpea plants was already
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56 475 observed by Weber et al. (1992, 1993) and Zaidi et al. (2003) and confirms the key role that
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58 476 AMF can play in plant nutrient uptake even in agricultural conditions.

1 477 Beside the positive effect on shoot N and P, AMF inoculation was effective in raising Fe
2 478 and Zn grain concentrations. To the best of our knowledge, this is the first experiment
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4 479 showing that AMF inoculation enhance the nutritional value of chickpea. Actually, AM
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7 480 fungal inoculation determined increases in Fe and Zn concentration of about 5% and 16%,
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9
10 481 respectively respect with controls. This may be due to the higher AM fungal root colonisation
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12 482 observed in the inoculated chickpea and, consequently, to a larger hyphal network that could
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14 483 have enhanced the transfer and uptake of trace elements (Audet and Charest, 2007). In
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17 484 addition, the larger occurrence of AMF could have determined an acidification of the
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19 485 rhizosphere due to the higher release of organic acids and phenolic compounds with the
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22 486 increases of soil Fe availability (White and Broadley, 2009; Antunes et al., 2012).
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24 487 Consistently, we observed higher grain Fe concentrations in the spring sowing treatment than
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27 488 in the autumn one, in line with the larger AM fungal root colonization of spring-sown
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29 489 chickpea.

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31 490 Beside micronutrient, in the spring sowing treatment, AM fungal inoculation had a
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34 491 significant biofortification effect also on protein (N concentration x 6.25; Antongiovanni,
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36 492 2004), enriching the chickpea grain total protein content about four percent (from 17% to
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39 493 21%). Also for what concern proteins, we can speculate that the higher grain protein contents
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41 494 observed in the inoculated plants could be related to the larger AM fungal hyphal network
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44 495 that has been shown to increase the inorganic and organic soil N mobilization (Harrison et al.,
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46 496 2002; Hodge et al., 2001; Govindarajulu et al., 2005) along with a more efficient N fixation
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49 497 by symbiotic rhizobial bacteria due to the better P nutritional status of the inoculated plants
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51 498 compared with controls.

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53 499 Interestingly, when considering the different inocula, the data suggest that AM fungal
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56 500 biofortification benefits may also depend on other factors than the root colonization rates,
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58 501 such as the functional properties of the different AMF. Actually, in the spring sowing
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61 502 treatment, grain protein, Fe and Zn concentration values of the NMix inoculated plants were
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503 higher than those observed in FMix, despite FMix showed a higher AM fungal colonization
504 indicating that, when considering crop productivity and yield quality, local AMF could be
505 more convenient for field inoculation than foreign strains.

507 **5. Conclusion**

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509 The results obtained in this experiment show that AM fungal field inoculation could be an
510 effective tool to improve the cultivation of chickpea by boosting its productivity and grain
511 nutritional quality. Since chickpea is a basic food for millions of people, AM fungal
512 inoculation, in particular if based on on-farm inoculum production and on local AM fungal
513 strains, could represent a valid biofertilization and biofortification strategy able to benefit not
514 only the economics of the cultivation, but also the healthiness of the agricultural produces.
515 The responses obtained by chickpea to AM fungal field inoculation are a further indication
516 that the potential of a more mycorrhiza-friendly approach in agriculture could be great.
517 However, more research is needed to understand the effectiveness of different AM fungal
518 inoculation strategies in industrial scale agricultural systems and with different crops and soil
519 conditions.

521 **Acknowledgements**

522
523 This work is part of Elisa Pellegrino's PhD thesis project, which was funded by the
524 Sant'Anna School of Advanced Studies. We wish to thanks Dr. Arthur Schüßler for helpful
525 comments and discussion. Special thanks also to Antonio Pellegrino for technical assistance,
526 to the staff of the "Rottaia" experimental station of the University of Pisa for setting up and
527 managing the field experiment and to the staff of "Il Montino" restaurant of Pisa for the moral
528 support and the delicious chickpea pancakes.

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43 822 CAPTIONS

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 46 823 **Figure 1** Arbuscular mycorrhizal (AM) fungal root colonization, shoot dry matter, collar
 47 diameter and grain yield of chickpea (*Cicer arietinum* L.) plants inoculated with the two
 48 824 different single, foreign AM fungi *Funneliformis mosseae* (IMA1) or *Rhizophagus irregularis*
 49 (IMA5), a dual strain inoculum [foreign mixture (FMix): IMA1 + IMA5], a trap-culture-
 50 enriched locally-sourced AM fungal community (local mixture = LMix) and a control (mock
 51 825 inoculum = C). Plants were sampled at harvest (June). Black rectangles: autumn sowing,
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 61 plants sown in October; white rectangles: spring sowing, plants sown in March.
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2 831 **Figure 2** Redundancy Analysis biplots based on plant growth, yield, plant nutrient uptake,
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4 832 grain biofortification and arbuscular mycorrhizal (AM) fungal colonization of chickpea (*Cicer*
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7 833 *arietinum* L.) plants (used as response variable) sown in October [(a) autumn sowing] and in
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9 834 March [(b) spring sowing] and inoculated with the two different single, non-native AM fungi
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11 835 *Funneliformis mosseae* (IMA1) or *Rhizophagus irregularis* (IMA5), the dual strain inoculum
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13 836 [foreign mixture (FMix): IMA1 + IMA5], a trap-culture-enriched locally-sourced AM fungal
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15 837 community (local mixture = LMix) and a control (mock inoculum = Control) (used as
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17 838 explanatory variables). Solid arrows represent plant growth and yield variables: collar
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19 839 diameter, CD; shoot dry matter per plant, ShootDM; grain yield per plant, GrainY. Plant
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21 840 nutrient uptake and grain biofortification are represented by dashed arrows: grain Fe
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23 841 concentration, Grain[Fe]; grain Zn concentration, Grain[Zn]; grain N concentration, Grain[N];
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25 842 shoot N concentration, Shoot[N]; shoot P concentration, Shoot[P]. AM fungal colonization
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27 843 (AMFcoloniz) is represented by dotted arrow. The 1st and 2nd axes accounted for 62.9% and
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29 844 69.4% of the total variance explained by all canonical axes for the autumn (a) and the spring
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31 845 sowing treatment (b), respectively.
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Table 1

P-values of linear orthogonal contrasts for arbuscular mycorrhizal (AM) fungal root colonization, shoot dry matter, collar diameter and grain yield of chickpea (*Cicer arietinum* L.) plants inoculated with the two different single, foreign AM fungi *Funneliformis mosseae* (IMA1) or *Rhizophagus irregularis* (IMA5), a dual strain inoculum [foreign mixture (FMix): IMA1 + IMA5], a trap-culture-enriched locally-sourced AM fungal community (local mixture = LMix) and a control (mock inoculum = C). Autumn sowing, plants sown in October; Spring sowing, plants sown in March.

	AM fungal root colonization (%) ^a	Shoot dry matter ^{a,b} (g plant ⁻¹)	Collar diameter ^a (mm)	Grain yield (g plant ⁻¹)
<i>Treatment comparisons (P-values of linear orthogonal contrasts)</i>				
Autumn sowing				
C vs M ^c	<0.001 ^d	0.234	0.001	0.004
FMix vs Single ^e	0.008	0.564	0.613	0.992
FMix vs LMix	<0.001	0.946	0.057	0.242
<i>F. mosseae</i> IMA1 vs <i>R. irregularis</i> IMA5	0.001	0.726	0.532	0.144
Spring sowing				
C vs M	<0.001	0.003	<0.001	0.001
FMix vs Single	0.002	0.111	0.448	0.483
FMix vs LMix	<0.001	0.930	0.004	0.049
<i>F. mosseae</i> IMA1 vs <i>R. irregularis</i> IMA5	<0.001	0.389	0.624	0.029

^a At harvest (June).

^b Leaf and stem dry weight.

^c M, AM fungal inocula.

^d In bold statistically significant values according to the linear orthogonal contrasts ($P \leq 0.05$).

^e Single, single AM fungal inocula.

Table 2

Shoot and grain N and shoot P concentrations of chickpea (*Cicer arietinum* L.) plants inoculated with the two different single, foreign arbuscular mycorrhizal (AM) fungi *Funneliformis mosseae* (IMA1) or *Rhizophagus irregularis* (IMA5), a dual strain inoculum [foreign mixture (FMix): IMA1 + IMA5], a trap-culture-enriched locally-sourced AM fungal community (local mixture = LMix) and a control (mock inoculum = C). Plants were sampled at harvest (June). Autumn sowing, plants sown in October; Spring sowing, plants sown in March.

	Shoot N (%)	Shoot P (%)	Grain N (%)	Shoot N (%)	Shoot P (%)	Grain N (%)
	Autumn sowing			Spring sowing		
C	0.98 ± 0.07 ^a	0.25 ± 0.03	2.75 ± 0.03	0.93 ± 0.08	0.25 ± 0.03	2.76 ± 0.02
FMix	1.94 ± 0.06	0.35 ± 0.02	2.77 ± 0.09	1.87 ± 0.06	0.35 ± 0.01	2.81 ± 0.06
LMix	2.09 ± 0.04	0.40 ± 0.01	2.84 ± 0.14	1.94 ± 0.03	0.36 ± 0.02	2.92 ± 0.10
IMA1	2.13 ± 0.05	0.35 ± 0.03	2.89 ± 0.08	2.05 ± 0.07	0.39 ± 0.02	3.01 ± 0.01
IMA5	1.91 ± 0.00	0.38 ± 0.01	2.88 ± 0.08	1.96 ± 0.03	0.35 ± 0.00	2.95 ± 0.04
<i>Treatment comparisons (P-values of linear orthogonal contrasts)</i>						
C vs M ^b	<0.001 ^c	0.001	0.381	<0.001	0.001	0.022
FMix vs Single ^d	0.234	0.506	0.314	0.077	0.552	0.029
FMix vs LMix	0.063	0.134	0.606	0.372	0.730	0.191
<i>F. mosseae</i> IMA1 vs <i>R. irregularis</i> IMA5	0.010	0.260	0.938	0.282	0.154	0.462

^a Values are means ± SE of three replicate plots per treatment.

^b M, AM fungal inocula.

^c In bold statistically significant values according to the linear orthogonal contrasts ($P \leq 0.05$).

^d Single, single AM fungal inocula.

Table 3

Grain Fe and Zn concentrations of chickpea (*Cicer arietinum* L.) plants inoculated with the two different single, foreign AM fungi *Funneliformis mosseae* (IMA1) or *Rhizophagus irregularis* (IMA5), a dual strain inoculum [foreign mixture (FMix): IMA1 + IMA5], a trap-culture-enriched locally-sourced AM fungal community (local mixture = LMix) and a control (mock inoculum = C). Plants were sampled at harvest (June). Autumn sowing, plants sown in October; Spring sowing, plants sown in March.

	Fe	Zn	Fe	Zn
	($\mu\text{g g}^{-1}$)	($\mu\text{g g}^{-1}$)	($\mu\text{g g}^{-1}$)	($\mu\text{g g}^{-1}$)
	Autumn sowing		Spring sowing	
C	472.67 \pm 5.36 ^a	23.57 \pm 0.90	487.33 \pm 2.19	23.33 \pm 0.88
FMix	492.33 \pm 1.45	26.27 \pm 0.09	500.33 \pm 2.60	26.33 \pm 0.18
LMix	512.33 \pm 1.45	32.67 \pm 0.88	523.67 \pm 6.64	31.10 \pm 0.63
IMA1	484.33 \pm 4.70	25.07 \pm 0.26	494.33 \pm 2.03	25.67 \pm 0.09
IMA5	488.67 \pm 5.49	25.40 \pm 0.38	504.00 \pm 2.08	24.60 \pm 0.15
<i>Treatment comparisons (P-values of linear orthogonal contrasts)</i>				
C vs M ^b	0.001 ^c	<0.001	0.001	<0.001
FMix vs Single ^d	0.275	0.354	0.796	0.078
FMix vs LMix	0.006	<0.001	0.001	<0.001
<i>F. mosseae</i> IMA1 vs <i>R. irregularis</i> IMA5	0.475	0.732	0.085	0.182

^a Values are means \pm SE of three replicate plots per treatment.

^b M, AM fungal inocula.

^c In bold statistically significant values according to the linear orthogonal contrasts ($P \leq 0.05$).

^d Single, single AM fungal inocula.

Table 4

Effects of inoculation with arbuscular mycorrhizal (AM) fungi (AMF inoc), sowing time (S time) and their interactions on chickpea (*Cicer arietinum* L.) AM fungal colonization, plant growth, yield and yield components at harvest.

Factors ^a	AM fungal colonization	Shoot dry matter ^b	Collar diameter	Grain yield ^c
AMF inoc ^d	< 0.001 ^e	0.007	< 0.001	< 0.001
S time	0.095	0.013	< 0.001	0.473
AMF inoc x S time	0.801	0.193	0.794	0.941

^a Two single, foreign AM fungi (*Funneliformis mosseae*, IMA1; *Rhizophagus irregularis* IMA5); a dual strain inoculum, foreign mixture (FMix): IMA1 + IMA5; a trap-culture-enriched locally-sourced AM fungal community (local mixture = LMix) and a mock inoculum as control; S time: autumn (October) and spring (March).

^b Leaf and stem dry weight.

^c Measurements per plant.

^d Two-way ANOVAs: AMF inoc, fixed factor; S time, fixed factor.

^e *P*-values of the two-way ANOVAs: in bold statistically significant values ($P \leq 0.05$). Replicates field plots were three per treatment.

Table 5

Effects of inoculation with arbuscular mycorrhizal (AM) fungi (AMF inoc), sowing time (S time) and their interactions on chickpea (*Cicer arietinum* L.) plant nutrient uptakes and grain biofortification.

Factors ^a	Shoot		Grain		
	N ^b	P	N	Fe	Zn
AMF inoc ^c	< 0.001 ^d	< 0.001	0.071	< 0.001	< 0.001
S time	0.098	0.764	0.182	< 0.001	0.158
AMF inoc x S time	0.506	0.326	0.942	0.859	0.569

^aTwo single, foreign AMF (*Funneliformis mosseae*, IMA1; *Rhizophagus irregularis* IMA5); a dual strain inoculum, foreign mixture (FMix): IMA1 + IMA5; a trap-culture-enriched locally-sourced AM fungal community (local mixture = LMix) and a mock inoculum as control; S time: autumn (October) and spring (March).

^bConcentrations.

^cTwo-way ANOVAs: AMF inoc, fixed factor; S time, fixed factor.

^d*P*-values of the two-way ANOVAs: in bold statistically significant values ($P \leq 0.05$). Replicates field plots were three per treatment.

Figure
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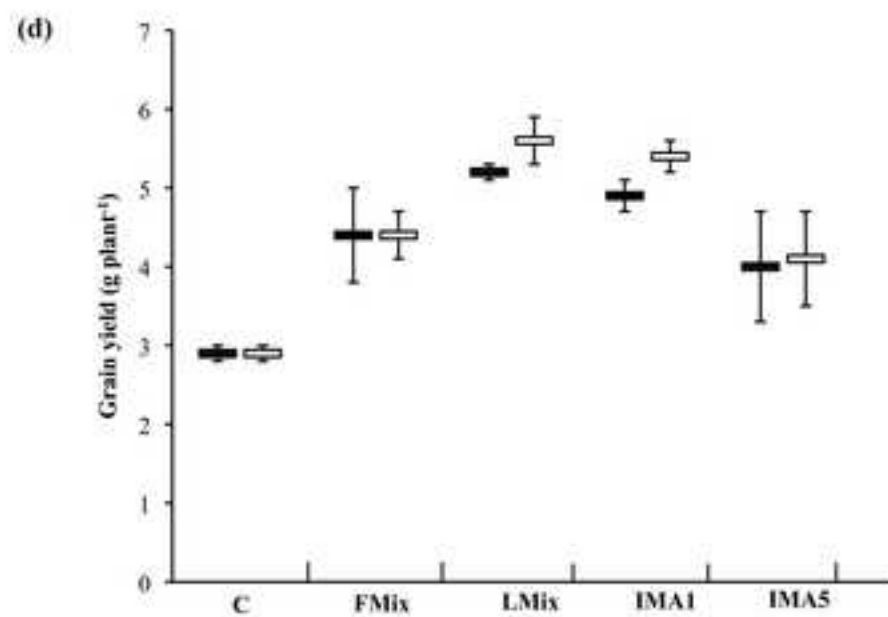
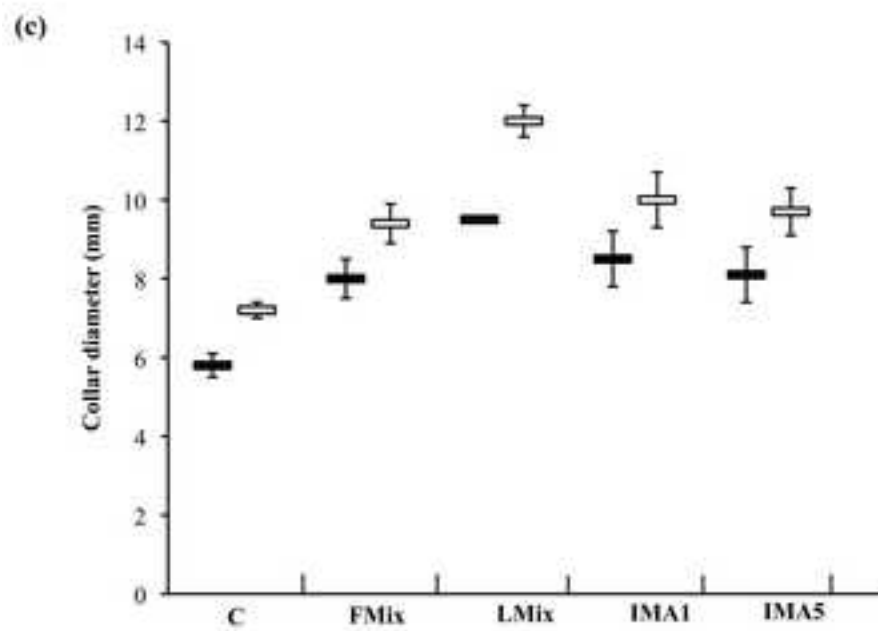
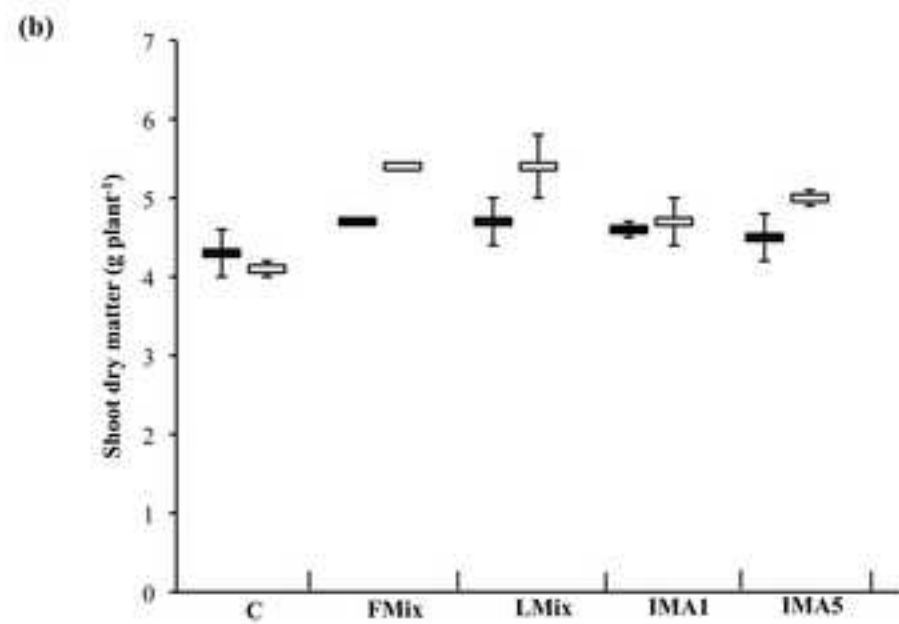
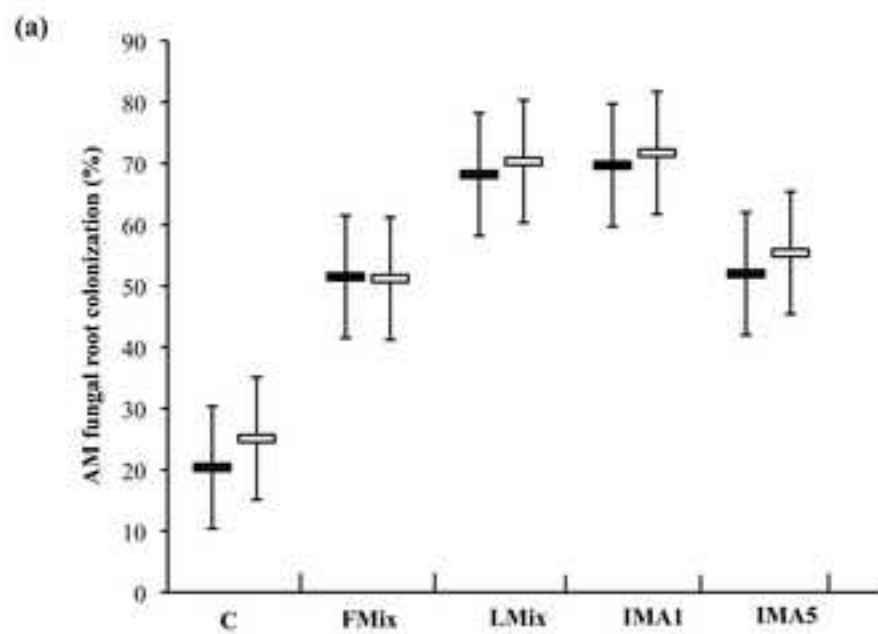


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