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1	Arbuscular mycorrhizal fungi reduce nitrous oxide emissions from N2O hotspots.					
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# 20 Summary

21	٠	Nitrous oxide (N2O) is a potent, globally important, greenhouse gas, predominantly
22		released from agricultural soils during nitrogen (N) cycling. Arbuscular mycorrhizal
23		fungi (AMF) form a mutualistic symbiosis with two-thirds of land plants, providing
24		phosphorus and/or N in exchange for carbon. Since AMF acquire N, it was
25		hypothesised that AMF hyphae may reduce N <sub>2</sub> O production.
26	•	AMF hyphae were either allowed (AMF) or prevented (non-AMF) access to a
27		compartment containing an organic matter and soil patch in two independent
28		microcosm experiments. Compartment and patch $N_2O$ production was measured both
29		before and after addition of ammonium and nitrate.
30	•	In both experiments, N <sub>2</sub> O production decreased when AMF hyphae were present prior
31		to inorganic N addition. In the presence of AMF hyphae, N <sub>2</sub> O production remained
32		low following ammonium application, but increased in the non-AMF controls. In
33		contrast, negligible N <sub>2</sub> O was produced following nitrate application to either AMF
34		treatment.
35	•	Thus, the main $N_2O$ source in this system appeared to be via nitrification and the
36		production of $N_2O$ was reduced in the presence of AMF hyphae. It is hypothesised
37		that AMF hyphae may be out competing slow growing nitrifiers for ammonium. This
38		has significant global implications for our understanding of soil N cycling pathways
39		and N <sub>2</sub> O production.
40		
41	Keyw	ords: agriculture, arbuscular mycorrhizal fungi (AMF), greenhouse gas, hyphosphere, N

42 cycle, nitrification, nitrogen (N), nitrous oxide (N<sub>2</sub>O)

### 43 Introduction

44 Agricultural soils are a major source of the globally important greenhouse gas, nitrous oxide

- 45 (N<sub>2</sub>O), a gaseous product of the nitrogen (N) cycle (Singh *et al.*, 2010; Hartmann, 2013). In
- 46 fact, the marked global N<sub>2</sub>O atmospheric concentration increases between 1940 and 2005
- 47 were predominantly a result of increased use of N-based fertilisers in agricultural systems (S.
- 48 Park *et al.*, 2012). N<sub>2</sub>O also has a long pertubation lifetime of 121 years (Hartmann, 2013),
- 49 thus it is essential that we understand the soil derived fluxes of  $N_2O$  as, unlike shorter lived
- 50 greenhouse gases (e.g. CH<sub>4</sub>; Hartmann, 2013), any changes in the atmospheric concentration
- of  $N_2O$  will have long term effects. Consequently,  $N_2O$  is viewed as an immediate target to
- 52 achieve greenhouse gas reductions (Wuebbles & Hayhoe, 2002; Reay *et al.*, 2012). However,
- 53 in order to achieve such reductions, an enhanced understanding of the major sources and sinks
- 54 of  $N_2O$  is urgently required.

55

- 56 In recent years, our understanding of N<sub>2</sub>O production in soil systems has significantly
- 57 improved, mostly as a result of the development of isotopic methods for tracing the sources of
- 58 N<sub>2</sub>O (Baggs, 2008; Kool *et al.*, 2011a; Ostrom & Ostrom, 2011). The rate of N<sub>2</sub>O production
- is predominantly controlled by the availability of the inorganic N source (Hino *et al.*, 2010),
- 60 O<sub>2</sub> (Bollman & Conrad, 1998), and other factors that influence microbial activity (e.g.
- 61 temperature, carbon (C) availability and pH (Bollman & Conrad, 1998; Prosser, 2007;
- 62 Thomson *et al.*, 2012)). In addition, recent evidence has revealed that  $N_2O$  reduction is not
- only confined to denitrifers. Other commonly occurring soil bacteria and archaea may also
- 164 utilise exogenous N<sub>2</sub>O, including under aerobic conditions, even though they lack the
- 65 preceding steps in the denitrification pathway (C.M. Jones *et al.*, 2014; Sanford *et al.*, 2012).
- Therefore, it follows that the net  $N_2O$  emitted from soils will be influenced by the presence of microorganisms.

- 69 Arbuscular mycorrhizal fungi (AMF) are a key group of soil microorganisms that form
- symbiotic associations with the majority of land plants (Smith & Read, 2008). Moreover, it is
- now widely acknowledged that these fungi play a previously unrecognised role in nitrogen
- 72 (N) cycling, and can both aquire N for their host plant (Barrett *et al.*, 2011; Herman *et al.*,
- 73 2012) as well as having a substantial N requirement themselves (Hodge & Fitter 2010). There
- is also evidence for reduced nitrate  $(NO_3)$  leaching in the presence AMF (Asghari &

Cavagnaro, 2012; Cavagnaro et al., 2015; Köhl & van der Heijden 2016). Alongside NO<sub>3</sub>, a 75 major output of the N cycle is the potent greenhouse gas, N<sub>2</sub>O. Therefore, it might be 76 expected that these fungi may influence the availability of N substrates (ammonium (NH<sub>4</sub><sup>+</sup>) 77 and  $NO_3^{-}$ ) for N<sub>2</sub>O production. AMF have been shown to be able to acquire both  $NH_4^+$  and 78  $NO_3^-$ , although it appears they may prefer the more energetically attractive  $NH_4^+$ 79 (Govindarajulu et al., 2005; Hodge & Storer, 2015). If these fungi compete effectively with 80 other microorganisms for these inorganic N forms then this could reduce the availability of N 81 substrates for N<sub>2</sub>O producers, leading to a reduction in N<sub>2</sub>O emissions. There is some 82 83 circumstantial evidence to suggest this may be the case. For example, Bender et al. (2014) found a reduction in N<sub>2</sub>O fluxes from soils influenced by AMF colonized roots when 84 compared to soils influenced by roots alone. N2O fluxes are also reduced when rice plants in 85 draining paddies are arbuscular mycorrhizal (Zhang *et al.*, 2015). Collectively, these studies 86 87 suggest that AMF may alter N<sub>2</sub>O emissions in conventional agricultural soils but, thus far, it has not been determined if this is mediated through physiological changes in the AMF 88 89 colonised roots, or as a direct result of the AMF themselves. If AMF hyphae can directly reduce N<sub>2</sub>O production, this could have significant implications for global N<sub>2</sub>O production 90 91 and our understanding of soil N cycling.

92

93 AMF hyphae have previously been demonstrated to proliferate in organic matter patches (e.g.; Hodge et al., 2001; Hodge, 2014; Barrett et al., 2014) and have been shown to take up and 94 transfer N in the inorganic form from these patches to their host plant (Leigh et al., 2009; 95 96 Hodge & Fitter, 2010). The two studies described here followed a similar experimental design to Hodge & Fitter (2010) using dried, milled Zea mays L. leaves mixed with an agricultural 97 98 soil (which had a high N<sub>2</sub>O production rate; Storer, 2013), to create organic matter 'patches'. These organic matter patches represent 'N<sub>2</sub>O hotspots' which commonly occur in natural 99 100 systems (Cowan et al., 2015). Both experiments tested the hypothesis that AMF hyphae 101 would reduce N<sub>2</sub>O production from the organic matter patches, while the second experiment 102 further examined the hypothesis that a reduction in N<sub>2</sub>O production was a consequence of reduced nitrification rates in the presence of AMF hyphae. 103

104

### 105 Materials and Methods

106 Microcosm design and growth media

107 To test the hypothesis that N<sub>2</sub>O production was reduced in the presence of AMF hyphae, two 108 experiments were established under glasshouse conditions using compartmented microcosm 109 units. Experiment 1 was designed to determine the impact of AMF hyphae on N<sub>2</sub>O production, 110 whereas Experiment 2 was designed to determine whether AMF hyphae affected N<sub>2</sub>O produced 111 by nitrification and/or denitrification. Organic matter patches were used to create 'hotspots' of 112 N<sub>2</sub>O production; a commonly observed phenomenon under natural conditions.

113

### 114 Experiment 1

Microcosm units (Fig. 1a) were constructed by joining two 1 L (each 145 x 145 x 70 mm<sup>3</sup>) 115 plastic containers via a double-mesh membrane of either 20 µm (John Stanier & Co., 116 Whitefield, Manchester, UK) or 0.45 µm (Osmonics Inc., Minnetonka, USA) pore size. These 117 size membranes either allowed (AMF) or denied (non-AMF) AMF hyphal access between the 118 two compartments. In all cases roots were prevented from passing between the compartments. 119 120 There were three 6 mm drainage holes in the base of each compartment. In one compartment (the 'planted' compartment) a single Z. mays seedling (Incredible F1, Mr Fothergills, 121 122 Newmarket, UK) inoculated with *Rhizophagus irregularis* (PlantWorks Ltd., Kent, UK) was placed, whereas the other compartment contained no plant (the 'unplanted' compartment). 123

124

### 125 *Experiment 2*

Three compartment microcosm units were used (Fig. 1b). Each microcosm consisted of a 126 127 central 'planted' compartment (volume: 2 L, dimensions: 150 x 150 x 150 mm; Thumbs Up Ltd., Bury, UK), containing a single Z. mays plant inoculated with R. irregularis, and on either 128 129 side of the central planted compartment, two unplanted compartments separated from the central compartment by a nylon mesh membrane as in Experiment 1 (volume: 2.6 L, 130 131 dimensions: 140 x 140 x 160 mm; Lock & Lock, Australia PTY Ltd., Blacktown, NSW, Australia). The mesh window either allowed AMF hyphal access (AMF; 20 µm mesh) or 132 133 prevented AMF hyphal access (non-AMF; 0.45 µm mesh) from the central planted to the outer unplanted compartments. A supporting stainless steel mesh (0.25 mm aperture; Mesh Direct, 134 135 Hanscan Ltd., Burslem, UK) was placed inside the plant compartment over the nylon meshes 136  $(0.45 \,\mu\text{m} \text{ and } 20 \,\mu\text{m})$  as a precautionary measure to protect the finer meshes from possible root damage. Thus, each unit had one AMF outer compartment and one non-AMF outer 137

compartment, creating a paired design. The unplanted compartments were covered with a foillayer when the lids were not attached to prevent them from drying out.

140

141 Experiments 1 & 2: Growth media

In both Experiments, the planted and unplanted compartments contained a mix (1:1 v/v) of sand 142 and Agsorb® (Agsorb®; Oil-Dri, USA; a calcined attapulgite clay soil conditioner) that had 143 been rinsed thoroughly in deionised water to remove any excess soluble N and/or P. The planted 144 compartments also had 50 g (Experiment 1) or 90 g (Experiment 2) of a fresh *R. irregularis* 145 inoculum (Plantworks Ltd., Kent, UK) and 0.25 g L<sup>-1</sup> bonemeal (a complex N and P source to 146 encourage mycorrhizal development; 3.5% N, 8.7% P; Vitax, Leicestershire, UK). Three pre-147 germinated Z. mays seeds were added to each planted compartment for both Experiments on 148 25<sup>th</sup> June 2012 and thinned to one per pot after 11 d (Experiment 1) or 14 d (Experiment 2). A 149 sterile centrifuge tube (Experiment 1: 15 cm<sup>3</sup>; Experiment 2: 50 cm<sup>3</sup>) was added to each of the 150 unplanted compartments to create a hole into which the organic matter patches and gas probes 151 could be added at a later date (see 'Organic matter patches and gas probes' section). 152

153

#### 154 Growth conditions

Microcosm units were placed in a randomised block design in a heated, lit glasshouse. The 155 Experiments ran for 78 days between 25<sup>th</sup> June – 10<sup>th</sup> Sept (Experiment 1) and 103 days between 156 25<sup>th</sup> June – 5<sup>th</sup> Oct 2012 (Experiment 2). Photosynthetically active radiation (PAR) was 157 measured weekly for both Experiments at plant level in the centre of each block and averaged 158 141 (Standard Error of the Mean; SEM +/- 15) µmol m<sup>-2</sup> s<sup>-1</sup> (Experiment 1) and 251 (SEM +/-159 45)  $\mu$  umol m<sup>-2</sup> s<sup>-1</sup> (Experiment 2). Overhead lights were used to extend the photoperiod to 16 h 160 per day and the mean daily temperature over the experimental period was 21.9°C (SEM +/-161 0.02) (Experiment 1) and 21.5°C (SEM +/- 0.3) (Experiment 2). The planted and unplanted 162 compartments for all microcosm units were watered daily as required. After two weeks of plant 163 growth, the planted compartments received 50 cm<sup>3</sup> of a reduced N and P nutrient solution as 164 described by Leigh et al. (2009) once a week (Experiment 1 and 2). This was increased to twice 165 weekly at 49 d after planting in Experiment 2 and to full N at 55 d after planting as the plants 166 were starting to show symptoms of N deficiency. In Experiment 2, at 76 d the plants began to 167 show P deficiency symptoms, so a 3/10th P, full N solution was used once a week in addition 168

- to two 1/10th N and P additions. In total the plants received either 1.74 kg N ha<sup>-1</sup> or 11.97 kg
- 170 N ha<sup>-1</sup> in Experiments 1 and 2 respectively over the duration of the Experiments (11 weeks and
- 171 14 weeks respectively).
- 172

### 173 Organic matter patches and gas probes

### 174 Organic matter patches

Organic matter patch material comprised of 13 g dry weight (DW) equivalent agricultural soil (sandy loam; 53°92'N, -1°00'E, pH 6.6 in 0.01M CaCl<sub>2</sub> (following Allen, 1974)) mixed with 2 g DW milled *Z. mays* leaves, all enclosed in a 20  $\mu$ m mesh bag (70 mm x 60 mm). The mean C and N content of the mixed organic patches were 1435 mg (SEM +/- 182) and 116 mg (SEM +/- 15) (Experiment 1) or 1200 (SEM +/- 79) and 99 mg (SEM +/- 15) (Experiment 2) respectively, with a C:N ratio of 12:1 in both Experiments. Each patch contained a gas probe (described below) in the centre (Fig. 1c).

182

### 183 *Gas probes*

A stainless steel tube (9 cm long, outer diameter 1 cm, wall thickness 1 mm; Coopers Needle 184 Works Ltd., Birmingham, UK) was welded at the base to form an airtight seal (Fig. 1c). Two 185 186 diametrically opposed holes, 6 mm, were drilled through each tube 13 mm from the base. These holes were covered in a PVDF membrane (0.2 µm, Bio-Rad Laboratories Inc., Hercules, 187 California, USA) that was air permeable but impermeable to water. This fine PVDF membrane 188 was then housed in a supporting silicone tube (wall thickness 0.8 mm, outer diameter 8 mm; 189 Silex Ltd., Hampshire, UK) with access holes exposing the membrane covering the holes. The 190 stainless steel tube was then sealed at the top with a white rubber Suba-Seal® (No. 13, Sigma-191 Aldrich, Poole, UK) to form a gas sampling port. The total internal volume of the gas probe 192 193 was ca.  $4.5 \text{ cm}^3$ .

194

A single organic matter patch and gas probe was placed into the pre-formed holes in the unplanted compartments 2 cm from the mesh window, 7 cm from the surface and covered with sand and Agsorb® media at 29 or 28 d (Experiment 1 or 2 respectively) post-planting.

### 199 Inorganic nitrogen addition

In Experiment 1, half of the organic matter patches were injected with 7 cm<sup>3</sup> of 30 mM NH<sub>4</sub>NO<sub>3</sub> 200 and the other half with 7 cm<sup>3</sup> of deionised water (n = 6 in each case) at 44 d after patch addition. 201 Consequently, the treatments were: AMF + NH<sub>4</sub>NO<sub>3</sub>, AMF + water, non-AMF + NH<sub>4</sub>NO<sub>3</sub> and 202 non-AMF + water. In Experiment 2, at 62 d after organic patch addition (90 d after planting) 203 each patch was injected with one of 7 cm<sup>3</sup> of: 15 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> (NH<sub>4</sub> treatment), 30 mM 204 KNO<sub>3</sub> (NO<sub>3</sub> treatment), 15 mM K<sub>2</sub>SO<sub>4</sub> (K<sub>2</sub>SO<sub>4</sub> treatment) or deionised water (water treatment), 205 where the N treatments were equivalent to 0.196 mg N g<sup>-1</sup> DW patch (n = 10 in each case). In 206 both experiments, two 3.5 cm<sup>3</sup> aliquots of solution were injected into each organic patch with 207 208 an hour gap between each addition to reduce spread into the surrounding sand/Agsorb®.

209

### 210 Gas sampling and analysis

### 211 Experiment 1

The air in the gas probes was sampled before N addition at 44 d post-patch addition. The 212 NH<sub>4</sub>NO<sub>3</sub> and water addition treatments were then added and the gas probes were sampled again 213 at 24, 48 and 96 h post-NH<sub>4</sub>NO<sub>3</sub> addition. Before sample removal, 1 cm<sup>3</sup> of N<sub>2</sub> was added to 214 the probe via the Suba-Seal®, taking care not to disturb the surrounding media. This was left 215 for 10 s before a 1 cm<sup>3</sup> sample was slowly removed from the gas probe, waiting for a further 5 216 s to allow the sample to mix inside the syringes before removing the syringe. Each gas sample 217 was then stored in a pre-filled 3 cm<sup>3</sup> Exetainer (with 6 cm<sup>3</sup> N<sub>2</sub>), over-pressuring the sample to 218 7 cm<sup>3</sup> in total. All gas samples were analysed using a gas chromatograph (GC) which quantified 219 the concentration of N<sub>2</sub>O. The concentration (ppm) values for each sample were calculated by 220 comparing to certified standards that were diluted in parallel in a 1 cm<sup>3</sup> standard: 6 cm<sup>3</sup> N<sub>2</sub> ratio 221 and correcting for this dilution. The concentration values were also corrected for dilution from 222 addition of N<sub>2</sub> to the gas probe just before gas sample removal. 223

224

# 225 Experiment 2

Gas sampling was carried out using both gas probes (as described above for Experiment 1) and
continuous flow loop sampling with an attached Los Gatos Isotopic N<sub>2</sub>O analyser (LGR N<sub>2</sub>O;
Los Gatos Research, Inc., California, USA) which provided an N<sub>2</sub>O concentration once per

second. A gas-tight lid (Fig. 1b) was attached to each of the 80 unplanted compartments in
block sequence for a minimum of 5 minutes, with a minimum of 2 min flushing the system with
air between each compartment measurement. Gas sampling using both methods was carried out
pre-N addition (58-59 and 61 d post-patch addition), and at 48, 96 and 192 h post-N addition
(64, 66 and 70 d post-patch addition respectively).

234

When using the LGR N<sub>2</sub>O analyser, the headspace in the microcosm unit (0.6 L), volume of connecting tubing (0.274 L) and internal volume of the N<sub>2</sub>O analyser (0.850 L) along with the surface area of the soil sampled (0.024 m<sup>-2</sup>) were used in the regression calculation of the N<sub>2</sub>O flux rate in mg m<sup>-2</sup> h<sup>-1</sup>. These fluxes were calculated using values measured between 200 to 280 s after the cover-box lid was attached. All regressions were calculated using SAS (v9.3 SAS institute Inc., North Carolina, USA).

241

### 242 **Post-harvest analyses**

At harvest, aboveground material was removed at the soil surface and separated into stalk, 243 244 flowers, ear, and leaf material. Roots were extracted from the sand/Agsorb® media, washed and fresh weight (FW) and DW of all plant material were recorded. In Experiment 1, the dried 245 leaves (green leaves only, defined as > 50% green) were milled and analysed for C and N 246 content using an elemental combustion system (Costech Analytical Technologies Inc., 247 California, USA). The gravimetric water content (g g<sup>-1</sup> DW) of soil, sand/Agsorb® and patches 248 for each compartment were measured and the AMF extraradical mycorrhizal hyphae (ERM) 249 were extracted from two 5 g (FW) samples from the organic patches and the surrounding growth 250 medium in the unplanted compartments using a modified membrane filter technique (see 251 Staddon et al., 2003) and acid fuchsin stain. Hyphal lengths were assessed using the gridline 252 intercept method (Miller & Jastrow, 1992) for a minimum of 50 fields of view at x 125 253 magnification (using a 10 x 10 grid of 1 cm side lengths, Graticules Ltd. UK). These hyphal 254 lengths were then converted to ERM length densities (m hyphae g<sup>-1</sup> soil DW). 255

256

# 257 Data analysis

Data were first tested for normality and equality of variance using Kolmogorov-Smirnov and
Levene's equality of variance tests respectively. Statistical analyses were carried out in either

SAS (v9.3 SAS institute Inc., North Carolina, USA) or Genstat (v16 VSN International Ltd., Hemel Hempstead, UK). The pre-N addition fluxes or concentrations were subtracted from the post-N addition fluxes or concentrations respectively to obtain the change in N<sub>2</sub>O flux or concentration following N addition (referred to  $\Delta N_2O$ ).

264

In Experiment 1, where N<sub>2</sub>O concentration and ERM length density data did not fulfil normality 265 or equality of variance assumptions they were  $log_{10}$  transformed. All gas concentration, plant 266 and AMF data were analysed using a two-way ANOVA including block, with Duncan's post 267 hoc tests. However, transformations on changes in N<sub>2</sub>O concentration following N addition 268 failed to normalise the data, and non-parametric equivalent Friedman's two-way ANOVAs, 269 including block, with Wilcoxon post hoc tests were used. Where N<sub>2</sub>O concentrations were 270 measured over time, repeated measures ANOVA including treatment and block was used on 271 log<sub>10</sub> transformed data. Pearson's product moment correlations were used to determine the 272 relationship between variables. Where variables were not normally distributed, Spearman's 273 274 rank order correlations were used. Untransformed data are presented in all figures.

275

In Experiment 2, differences among treatments were analysed using a two-way ANOVA including block with Duncan's *post hoc* tests. ERM length density data were log<sub>10</sub> transformed before analysis. Where the data failed normality or equality of variance assumptions, nonparametric tests were used. A one-sample t-test or a Wilcoxon Signed Ranks test was used to compare absolute values or differences to zero.

281

In Experiment 2, the  $\Delta N_2O$  data were not normally distributed and therefore a Friedman's nonparametric two-way ANOVA, controlling for block with Mann Whitney U (unpaired data) or Wilcoxon signed ranks (paired data) *post hoc* tests and an applied false discovery rate correction was used. Where comparisons in  $\Delta N_2O$  flux or  $\Delta N_2O$  concentration data were made over time, a non-parametric Friedman's repeated measures analysis was used. The relationship between the  $\Delta N_2O$  flux and  $\Delta N_2O$  concentration for each gas sample following N addition (48 h, 96 h and 192 h post-N addition) was determined using a Spearman's rank order correlation.

- 290 There was hyphal breakthrough in one of the non-AMF compartments (treatment: non-AMF,
- $K_2SO_4$ ) therefore this microcosm was excluded from the subsequent data analyses. In
- addition, the N<sub>2</sub>O concentration for one experimental unit in the AMF treatment (treatment:
- 293 (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>) was out of range on the GC at the 48 h post N addition sample and therefore these
- AMF and non-AMF N<sub>2</sub>O concentration values were also omitted.
- 295

### 296 **Results**

- 297 In the AMF treatments, *R. irregularis* colonized the organic matter patches successfully in
- both experiments with ERM length densities of  $1.23 \pm 0.25$  m g<sup>-1</sup> DW in Experiment 1 (non-
- AMF:  $0.31 \pm 0.05$  m g<sup>-1</sup> DW;  $F_{1,12} = 30.77$ , P = 0.0001) and  $0.88 \pm 0.08$  m g<sup>-1</sup> in Experiment
- 300 2 (non-AMF:  $0.35 \pm 0.04$  m g<sup>-1</sup> DW;  $t_{39} = 8.993$ , P < 0.0001).
- 301

# 302 **Pre-N** addition $N_2O$ production

Prior to inorganic N addition there was a greater concentration of N<sub>2</sub>O in the non-AMF 303 patches than in the AMF patches in both experiments (Fig. 2; Experiment 1:  $F_{1,12} = 6.46$ , P =304 0.026; Experiment 2:  $S_{38} = -186$ , P = 0.0076). A similar trend (at the P < 0.1 level) was found 305 for the N<sub>2</sub>O fluxes in Experiment 2, with greater N<sub>2</sub>O fluxes measured from the non-AMF 306 compartments than from the AMF compartments ( $S_{38} = -128$ , P = 0.074). In Experiment 2, 307 N<sub>2</sub>O fluxes measured by continuous flow loop sampling were positively correlated with the 308 patch N<sub>2</sub>O concentrations measured using gas probes ( $r_s = 0.7495$ , P < 0.0001). Since N<sub>2</sub>O 309 310 production is inherently variable, this level of consistency both between and within experiments is striking, particularly because it was observed in the absence of any additional 311 applied inorganic N. In both experiments, there was no significant correlation between the 312 pre-N addition AMF treatment N<sub>2</sub>O concentration or fluxes and the ERM lengths (P > 0.05 in 313

314 each case).

315

### 316 Post N-addition and harvest

### 317 Experiment 1

- 318 In Experiment 1 the highest patch N<sub>2</sub>O concentrations were observed 24 h after the
- application of inorganic N or water in all treatments except AMF + water, demonstrating the

rapid response of N<sub>2</sub>O producers to treatment application. The patch N<sub>2</sub>O concentrations of 320 321 the non-AMF and AMF + NH<sub>4</sub>NO<sub>3</sub> treatments subsequently decreased over time. In contrast, the AMF + water patch N<sub>2</sub>O concentration remained low. Consequently, there was a 322 significant effect of both time and treatment on patch N<sub>2</sub>O concentration in addition to a 323 significant interaction between these two factors (Fig. 3; time:  $F_{2,30} = 4.37$ , P = 0.023; 324 treatment:  $F_{3,15} = 5.67$ , P = 0.0084; time\*treatment:  $F_{6,30} = 3.23$ , P = 0.015). These results 325 therefore demonstrate how rapidly N<sub>2</sub>O production rates can change over time and emphasise 326 the requirement for repeated measurements following inorganic N application. Two-way 327 ANOVAs at each time point showed that the N<sub>2</sub>O concentration of the AMF + water 328 treatment was lower than all other treatments at 24 h post-treatment application (Fig. 3;  $F_{3,15}$  = 329 4.44, P = 0.020). This effect decreased by the 48 h sample, although the non-AMF + water 330 and non-AMF + NH<sub>4</sub>NO<sub>3</sub> treatments still had a higher  $N_2O$  concentration than that of the 331 AMF + water treatment ( $F_{3,15} = 4.95$ , P = 0.014). At 96 h post-treatment application, the AMF 332 patch N<sub>2</sub>O concentrations were not significantly different from each other but were 333 significantly lower than those of the non-AMF patches ( $F_{3,15} = 7.25$ , P = 0.0031). At 24 h 334 post-treatment application, the  $\Delta N_2O$  concentration was higher in both the AMF + NH<sub>4</sub>NO<sub>3</sub> 335 and non-AMF + NH<sub>4</sub>NO<sub>3</sub> treatments than the AMF + water treatment ( $Q_3 = 8.2, P = 0.042$ ). 336 However, the non-AMF + water treatment was not significantly different from the AMF + 337

338  $NH_4NO_3$  treatment or non-AMF +  $NH_4NO_3$  treatment.

339

There was no relationship between the AMF ERM length densities and N<sub>2</sub>O concentration in 340 341 the AMF patches at any point (P > 0.05 in each case) and the moisture contents of the organic patches did not differ among treatments at harvest ( $Q_3 = 0.05$ , P = 0.827). Additionally, there 342 343 was no significant difference (P > 0.05) in total plant DW or the DW of the various plant tissues (i.e. leaf, total shoot, stalk, total root, root weight ratio, tassel) between the AMF and 344 non-AMF treatments (see Supporting Information Table S1). Neither the addition of NH<sub>4</sub>NO<sub>3</sub> 345 or water had any effect on the leaf C and N content or concentrations or the C:N ratios (P >346 0.05 in each case), therefore, these data were combined for comparison of the AMF versus the 347 non-AMF treatments. Leaf C content did not differ between AMF and non-AMF plants 348 349 (Table 1;  $F_{1,12} = 0.30$ , P = 0.595), although the leaf C concentrations were lower in the AMF compared to the non-AMF treatments (Table 1;  $F_{1,12} = 5.37$ , P = 0.039). Both the N content 350 (Table 1;  $F_{1,12} = 14,18$ , P = 0.0023) and concentration ( $F_{1,12} = 20.06$ , P = 0.0008) of the 351 leaves were higher in the AMF compared to the non-AMF treatments. Consequently, the C:N 352

ratio of the leaves was lower in the AMF compared to the non-AMF treatments (Table 1;  $F_{1,12}$ = 18.51, P = 0.001). However, the organic patch N<sub>2</sub>O concentration was not significantly related to the leaf C or N content or concentration, or to the leaf C:N ratio, either before or after N addition, for both the AMF and non-AMF treatments (P > 0.05 in each case).

357

### 358 *Experiment 2*

There was a significant difference in  $\Delta N_2O$  fluxes among the inorganic N and water 359 application treatments at 48 h post-application (Fig. 4;  $O_7 = 44.85$ , P < 0.0001). In both the 360 AMF and non-AMF patches, more N<sub>2</sub>O was produced following addition of NH<sub>4</sub><sup>+</sup> than any 361 other treatment. Strikingly, however, ca. 2.5 times more N<sub>2</sub>O was produced from the non-362 AMF than the AMF treatment (Fig. 4;  $S_9 = -26.5$ , P = 0.0084). These differences then 363 declined by the 96 h sample and were no longer significant at the 192 h sample, again, 364 illustrating the transient nature of N<sub>2</sub>O release and the importance of following the fluxes over 365 366 discrete timescales (Table 2). There was no significant difference in the % moisture content of either the patch or sand/Agsorb® medium between the AMF and non-AMF treatments at 367 destructive harvest (patch:  $t_{39} = -0.26$ , P = 0.799; sand/Agsorb®:  $S_{39} = -47$ , P = 0.519). 368

369

### 370 **Discussion**

This is the first study to show that N<sub>2</sub>O production is reduced as a direct consequence of the 371 presence of AMF hyphae. Moreover, this reduction was demonstrated in both the presence 372 373 and, notably, the absence of applied inorganic N, indicating that this is a persistent effect. Studies to date have indicated that AMF may influence soil N<sub>2</sub>O production, but this has 374 always been in the presence of plant roots and additional inorganic N (Lazcano et al., 2014; 375 Bender et al., 2015). Critically, the finding that N<sub>2</sub>O production was reduced when AMF 376 377 hyphae, but not plant roots, were present was consistent between the two independent experiments reported here. 378

379

381 mycorrhizosphere (i.e. the soil influenced by AM colonised roots and AMF hyphae), often

only at a single time point after N application, potentially masking cumulative effects (Bender

*et al.*, 2015). AMF hyphae can extend far beyond the plant roots alone, with the ERM being

10 times larger, in biomass terms, than the intraradical mycelium (Olsson et al., 1999). Thus, 384 the influence of AMF hyphae on soils (in the 'hyphosphere') will extend beyond the zone of 385 influence of roots alone, and studies to date have not explored this widespread zone of hyphal 386 influence on N<sub>2</sub>O production in isolation. Furthermore, since the mycorrhizosphere includes 387 both AMF colonised plant roots and AMF hyphae, it is impossible to know whether any effect 388 is a consequence of the AMF hyphae, roots, or both. Rhizodeposition differs between AM and 389 non-AM plants (D.L. Jones et al., 2004), while C exudation from AMF hyphae may also 390 result in quantitative and qualitative changes in the total C flux into the soil (Toljander et al., 391 2007). Moreover, AMF hyphae influence N cycling through the capture of N and subsequent 392 transfer of at least some of this N to their associated host plant (Leigh et al., 2009; Thirkell et 393 al., 2016). C and N are key controls of denitrification and nitrification rates (Bollman & 394 Conrad, 1998; Hino et al., 2010). It is not possible, therefore, to separate AMF and root 395 396 control of N<sub>2</sub>O fluxes in the mycorrhizosphere without first separating the AMF hyphae from the plant roots. 397

398

Nevertheless, there is some evidence of AMF interacting with soil N<sub>2</sub>O production in the 399 mycorrhizosphere, although results have been inconsistent. Bender et al., (2015) found that 400 the N<sub>2</sub>O flux was lower following the application of  $NO_3^-$  in the AM mycorrhizosphere when 401 402 compared to the rhizosphere of a non-AM control. In contrast, Cavagnaro et al., (2012) found no effect of AM plants on N<sub>2</sub>O production, whereas Lazcano et al., (2014) found a reduction 403 in N<sub>2</sub>O in the mycorrhizosphere of AM plants. Thus, there is support for AMF resulting in 404 405 reduced N<sub>2</sub>O production in the mycorrhizosphere, but the cause of this reduction has so far been poorly understood, likely because of confounding effects of the host plant root system 406 407 also being present. Hypotheses for the decreased N<sub>2</sub>O production in the mycorrhizosphere 408 included a reduction in denitrification (Bender et al., 2015), and increased water use by AM 409 plants (Lazcano et al., 2014).

410

411 In this study, the finding of reduced N<sub>2</sub>O production in the presence of AMF hyphae was

412 evident even prior to inorganic N application. There was also evidence for an increase in both

413 leaf N content and concentration when the AMF had access to the organic matter patches.

414 This suggests that the AMF were supplying their host plant with additional N, presumably

415 from the organic matter patch since all planted compartments received the same quantity of

nutrient solution. Whilst there is a wide range in reported contribution of AMF to plant N
(reviewed by Hodge & Storer, 2015), the findings in this study are in agreement with previous
investigations using <sup>15</sup>N that substantial quantities of N can be transferred from the patch to
the plant via AMF hyphae (Leigh *et al.* 2009; Thirkell *et al.* 2016).

420

The inorganic N applications here were used as a tool to identify the pathway of N<sub>2</sub>O 421 422 production being influenced by the AMF hyphae. The addition of NO<sub>3</sub><sup>-</sup> did not result in increased N<sub>2</sub>O production from any treatment, suggesting that in this study denitrification was 423 424 not a key factor in controlling N<sub>2</sub>O production. There was also no significant difference in gravimetric water content of the organic matter patches, or the surrounding sand/Agsorb® 425 medium at harvest. Thus, these factors were not important controls of N<sub>2</sub>O production in the 426 present study. Instead, we found direct evidence for a reduction in N<sub>2</sub>O produced via 427 nitrification in the presence of AMF hyphae. This is a critical finding and may help to explain 428 variable N<sub>2</sub>O fluxes under field conditions. Since one of the largest sources of N<sub>2</sub>O are 429 agricultural soils, it is highly relevant that the soil used here was agricultural in origin, and the 430 plant material for the organic matter patches was Z. mays, a globally important crop (Leff et 431 al., 2004). 432

433

The soil N<sub>2</sub>O fluxes in this study were predominantly controlled by the availability of NH<sub>4</sub><sup>+</sup>. 434 These fluxes were monitored at intervals up to 192 hours after inorganic N application by 435 which point the N<sub>2</sub>O peak declined back to pre-N application levels, thus ensuring that the full 436 response period was recorded. There was a significantly greater N<sub>2</sub>O flux in response to NH<sub>4</sub><sup>+</sup> 437 addition in the non-AMF than in the AMF treatment, indicating reduced N<sub>2</sub>O production via 438 nitrification in the presence of AMF hyphae. The current understanding of the main pathways 439 440 of N<sub>2</sub>O production in soils (as described in Baggs, 2011, Zhu et al., 2013), are shown in Fig. 5 together with the potential mechanisms by which AMF may interact with N<sub>2</sub>O production. 441 If NH<sub>4</sub><sup>+</sup> elicits N<sub>2</sub>O production but NO<sub>3</sub><sup>-</sup> application does not, by process of elimination the 442 pathway involved in N<sub>2</sub>O production must be a nitrification pathway. 443

444

The links between AMF presence and reduced nitrification rates are in broad agreement with
a series of one field based and three mesocosm based studies by Veresoglou *et al.* (2011). The

- 447 potential nitrification rates (PNR) were lower in the mycorrhizospheres of AM plants
- 448 compared to those from weakly-AM mycorrhizospheres (Veresoglou *et al.*, 2011). The *nirK*
- gene, responsible for N<sub>2</sub>O production has also been shown to be negatively correlated with

450 AMF abundance (Bender *et al.*, 2014). Thus, the presence of AM plants may reduce N<sub>2</sub>O

- 451 production by reducing nitrification rates. Our present study demonstrates, for the first time,
- 452 that AMF hyphae have a direct and limiting influence on soil N<sub>2</sub>O produced via nitrification,
- 453 independent of any plant root influence.

454

- The main 'nitrification' pathways in soil potentially resulting in N<sub>2</sub>O release are nitrifier
- 456 nitrification, and nitrifier denitrification. Nitrifier nitrification is an aerobic process and can be
- 457 carried out by ammonia oxidising bacteria (AOB), archaea (AOA), and organsims capable of
- 458 complete ammonia oxidation (comammox) (Daims *et al.*, 2015; van Kessel *et al.*, 2015).
- 459 AOB and AOA have also been shown to produce N<sub>2</sub>O (Jiang and Bakken, 1999; Jung *et al.*,
- 460 2014). Nitrifier denitrification is also carried out by autotrophic nitrifiers, and can be a
- significant source of N<sub>2</sub>O (Kool *et al.*, 2011b; Wrage *et al.*, 2001). Thus, there are various
- 462 pathways by which the  $N_2O$  in this study may have been produced following the application
- 463 of  $NH_4^+$  and consequently reduced by the presence of AMF hyphae (Fig. 5).

464

Regardless of the process, the response to NH<sub>4</sub><sup>+</sup> application in the AMF treatments suggests 465 that there was either a reduction in N<sub>2</sub>O production, through reduced function or number of 466 nitrifiers, or that nitrifier activity was masked by an increase in activity of N<sub>2</sub>O reducers 467 which can cause some soils to become N<sub>2</sub>O sinks (Domeignoz-Horta et al., 2017). It's also 468 feasible that the presence of AMF hyphae modified the microbial community shifting it away 469 from N<sub>2</sub>O producing nitrifiers or nitrifier denitrifiers, perhaps towards organisms capable of 470 471 complete nitrification (van Kessel et al., 2015), or N<sub>2</sub>O reduction (C.M. Jones et al., 2014; Sanford et al., 2012; Domeignoz-Horta et al., 2017). 472

473

Domeignoz-Horta *et al.* (2017) found that N<sub>2</sub>O hotspots were predominantly controlled by
changes in the microbial communities, whereas lower N<sub>2</sub>O producing areas were more likely
to be controlled by variation in soil properties. Using similar organic patches as in the present
study, Nuccio *et al.* (2013) found that while there was no overall change in bacterial diversity,

the presence of AMF hyphae significantly modified the bacterial community. Interestingly, 478 Gemmatimonadetes and Deltaproteobacteria were two of four bacterial phylum that had a 479 higher relative abundance in response to the presence of AMF hyphae in the litter (Nuccio et 480 al., 2013). Both the Gemmatimonadetes and Deltaproteobacteria have subsequently been 481 found to posses nosZ genes, and can, thus, utilise exogenous N<sub>2</sub>O as an electron acceptor 482 (C.M. Jones et al., 2013; D. Park et al., 2017). AMF abundance has also been found to 483 positively correlate with nosZ gene abundance (Bender et al., 2014). This together with the 484 large export of N from the patch by the AMF hyphae and the resulting modifications in the 485 486 physico-chemical environment in the decomposing litter patch may all contribute to a reduction in N<sub>2</sub>O emissions. 487

488

Given the evidence that AMF are known to have a high N demand (Hodge & Fitter, 2010), 489 one hypothesis may be that AMF hyphae were eliciting a longer term control on the nitrifying 490 community as nitrifiers are inhereantly slow growing, taking from 8 h up to a number of days 491 to double in number (Belser & Schmidt, 1980; Woldendorp & Laanbroek, 1989; Prosser, 492 2007; Prosser & Nicol, 2012). AMF hyphae are thought to predominantly take up inorganic N 493 in the form of NH<sub>4</sub><sup>+</sup> (Govindarajulu *et al.*, 2005; Tanaka & Yano, 2005), and AOB are 494 generally thought to be poor competitors for  $NH_4^+$  (Verhagen *et al.*, 1995; Bollmann *et al.*, 495 2002). The AMF hyphae may therefore have reduced the amount of available  $NH_4^+$  in the 496 hyphosphere resulting in a reduction in the population of active AOB. If AOB were the main 497 N<sub>2</sub>O producers, this may explain the reduced N<sub>2</sub>O production prior to inorganic N application 498 499 when the AMF hyphae were present. It may also explain the lack of N<sub>2</sub>O production in the presence of AMF hyphae when NH<sub>4</sub><sup>+</sup> was applied, i.e. the AOB population may have been 500 501 small and too slow-growing to respond to the inorganic NH<sub>4</sub><sup>+</sup> supplied, which may have, 502 instead, been taken up by the N-rich AMF hyphae.

503

504 While AMF may increase or decrease the pH of surrounding media, thought to be a 505 consequence of  $NO_3^-$  or  $NH_4^+$  uptake respectively (Bago *et al.*, 1996; Li *et al.*, 1991), the 506 relative importance of pH effects on N<sub>2</sub>O production if C,  $NH_4^+$  or  $NO_3^-$  are limiting are not 507 clear (reviewed by Šimek & Cooper, 2002) with both increased and decreased nitrification-508 derived N<sub>2</sub>O production reported under low pH conditions (Y. Cheng *et al.*, 2013; Mørkved *et 509 al.*, 2007). The patch pH was not measured in this study, and potential changes in pH cannot 510 be fully discounted. However, the implications of N, and more importantly the form of N,

511 exported by AMF on the local physico-chemical properties including pH, warrants more

512 attention. This may also aid in explaining the differing impacts reported for AMF on

513 decomposition processes, and their importance not only for N, but also C cycling and

stabilization processes (Hodge, 2001; Hodge *et al.*, 2001; L. Cheng *et al.*, 2012).

515

In order to fully understand the mechanism for the reduction in N<sub>2</sub>O production via 516 nitrification observed in the presence of AMF hyphae found in this study, further research 517 should focus upon gene expression and the responses of the microbial community including 518 nitrifier communities, AOA, AOB and potential non-denitrifying N<sub>2</sub>O reducers. Monitoring 519 would also help to establish if nitrifier populations were suppressed by the presence of AMF 520 521 hyphae, as we suggest. Furthermore, field based studies using a wider range of soil types and environmental conditions are an essential next step to determine the global scale and 522 significance of this interaction in both natural and agricultural systems. 523

524

525 In conclusion, using two independent glasshouse based experiments, we have found that the presence of AMF hyphae reduced the production of the globally important greenhouse gas, 526 N<sub>2</sub>O. Cropped agricultural soils cover a significant proportion of land area, representing 527 28.4% of agricultural land, or 10.9% of the total global land area in 2011 (FAO, 2017). The 528 diversity of AMF is reduced in agricultural soils (Helgason et al., 1998), and these soils are 529 one of the largest contributors to N<sub>2</sub>O emissions. This study suggests that a reduction in AMF 530 presence may contribute to further increases in N<sub>2</sub>O production. This could have significant 531 implications for better management of agricultural soils in the future. Given the ubiquity of 532 the AM association, including under agricultural situtations, these findings have global 533 implications not only for our fundamental understanding of the mechanisms of soil N cycling, 534 but also for greenhouse gas management and climate change mitigation. 535

536

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541

### 542 Author Contribution

K.S., P.I. and A.H. designed the research; K.S. performed the research and conducted all data
analysis, A.C. performed practical work for Experiment 1. K.S. and A.H. wrote the
manuscript.

546

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**Table 1.** Mean leaf N and C total content, concentration and C:N ratio of leaves from AMF and non-AMF

treatments in Experiment 1  $\pm$  standard error of the mean (n = 12). Different letters within rows represent

737 significant di	fferences at $P = 0.05$ (	(in bold)	as determined	using two-way	y ANOVAs.
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		AMF	Non-AMF
Leaf N	Total content (mg)	$13.8\pm0.8^{\rm a}$	$10.2\pm0.9^{\rm b}$
	Concentration (mg g <sup>-1</sup> DW)	$11.3\pm0.6^{\rm f}$	$\pmb{8.8 \pm 0.5^{\rm g}}$
Leaf C	Total content (mg)	$503.2\pm19.9^{j}$	$488.1\pm27.2^{j}$
	Concentration (mg g <sup>-1</sup> DW)	$413.4\pm2.8^{m}$	$422.9\pm3.7^{\rm n}$
	Leaf C:N ratio	$37.6 \pm 2.0^{x}$	$50.0 \pm 3.0^{\text{y}}$

738

- 740 Table 2. Experiment 2 Friedman's test statistics controlling for block comparing the post-N minus pre-N (61 days
- post-patch addition) patch N<sub>2</sub>O concentrations ( $\Delta$ N<sub>2</sub>O concentrations) or compartment N<sub>2</sub>O fluxes ( $\Delta$ N<sub>2</sub>O fluxes)

	Time since N addition								
	48 h		96 h			192 h			
	Q	df	Р	Q	df	Р	Q	df	Р
Patch ΔN <sub>2</sub> O	28.89	7	0.0002***	14.35	7	0.045*	3.79	7	0.804
concentration									
Compartment	44.85	7	<0.0001***	25.63	7	0.0006***	4.80	7	0.684
$\Delta N_2 O$ flux									

among N addition treatments, for each of the gas sampling events.

743 Q = Friedman's test statistic, df = degrees of freedom, n = 10. Significant results are indicated in bold at P = 0.05

744 (\*P < 0.05, \*\*P < 0.01, \*\*\*P < 0.001).

745

#### 747 Figure & Table Legends

Fig. 1 The microcosm units used in Experiments 1 (a) and Experiment 2 (b) and the organic 748 matter patches and gas probes used in both experiments (c). In Experiment 1 the planted 749 compartment was planted with a single Zea mays plant and contained the AMF inoculum whilst 750 the unplanted compartment either allowed or prevented AMF hyphal access. In Experiment 2 751 the central compartment was also planted with a single Z. mays plant and contained the AMF 752 inoculum. From the central, planted compartment, the AMF hyphae could access one outer, 753 unplanted compartment (AMF) but not the other (non-AMF). The gas probe was placed within 754 a mesh bag (the 'organic matter patch') which contained a mix of dried, milled Z. mays leaves 755 and agricultural soil (c). The gas probe and organic matter patch designs were used in both 756 experiments. 757

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**Fig. 2** Mean N<sub>2</sub>O concentration (ppm) in AMF access (AMF) and no AMF access (non-AMF) organic matter patches (a) at 43 d post-patch addition in Experiment 1 and (b) at 58 d postpatch addition in Experiment 2. Error bars represent  $\pm$  standard error of the mean (a: n = 12, b: n = 39). Different letters represent significant differences at P < 0.05 as determined using (a) two-way ANOVAs and (b) by comparing the  $\Delta$ AMF value to zero (Wilcoxon Signed Rank test).

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Fig. 3 Mean patch N<sub>2</sub>O concentration at 24, 48 and 96 h following inorganic N (NH<sub>4</sub>NO<sub>3</sub>: filled symbols) or water (open symbols) addition for AMF access patches (AMF; solid lines) and no AMF access patches (non-AMF; dashed lines) shown over time. Error bars represent  $\pm$  standard error of the mean (n = 6). Asterisks represent a significant difference among treatments within each sample period (\*P < 0.05, \*\*P < 0.01) as determined using a two-way ANOVA. Different letters within each sample timing represent significant differences between treatments for that sample timing (P < 0.05).

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**Fig. 4** Mean difference between 48 h post-N addition (64 d post-patch addition) and pre-N addition (61 days post-patch addition) N<sub>2</sub>O flux ( $\Delta$ N<sub>2</sub>O flux) for AMF access (AMF; solid bars) and no AMF access (non-AMF; open bars) treatments, split by N addition treatment. The N addition treatments were: (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> (labelled as NH<sub>4</sub>), KNO<sub>3</sub> (labelled as NO<sub>3</sub>), K<sub>2</sub>SO<sub>4</sub> or water. Bars with different letters are significant at P = 0.0018 as determined using Mann Whitney U or Wilcoxon signed rank *post hoc* tests with a False Discovery rate correction applied. Asterisks below the bars indicate significant differences to zero (\*P < 0.05, \*\*P < 0.01). Error bars are ± standard error of the mean (n = 10).

782

Fig. 5 Summarized potential interactions between AMF hyphae and soil N<sub>2</sub>O producing 783 784 processes as described in Baggs (2011) and Zhu et al. (2013). The solid and dashed bold lines represent AMF effects that could result in an increase or decrease in N2O production 785 respectively. AMF can affect the availability of nitrogen (N), phosphorus (P), copper (Cu) 786 and iron (Fe) in soils as well as potentially change soil pH. Nitrifier nitrification is generally 787 788 carried out by ammonia oxidising bacteria (AOB) and archaea (AOA). Dissimilatory reduction of nitrate to ammonium (DNRA) may produce N<sub>2</sub>O as a side product. DNRA is 789 790 also known as nitrate ammonification. There are various pathways and organisms capable of 791 carrying out these roles, but for simplicity, they are grouped by factors affecting the rate of  $N_2O$  production (i.e. availability of  $O_2$ , or carbon (C)). 792

793

Supporting Information Table S1. Mean plant biomass parameters from AMF and non-AMF
 treatments in Experiment 1.



organic matter patch

Figure 2



Figure 3.



Time since  $NH_4NO_3$  or water addition (h)

Figure 4.





### New Phytologist Supporting Information

Article title: Arbuscular mycorrhizal fungi reduce nitrous oxide emissions via impacts on nitrification.

Authors: Kate Storer, Aisha Coggan, Phil Ineson, Angela Hodge

The following Supporting Information is available for this article:

**Table S1.** Mean plant biomass parameters from AMF and non-AMF treatments in Experiment 1  $\pm$  standard error of the mean (n = 12). There were no significant differences between treatments for any of the parameters (P > 0.05 in each case) as determined using two-way ANOVAs.

	AMF	Non-AMF
Leaf DW (g)	$1.22^{a}\pm0.05$	$1.16^{a} \pm 0.07$
Stalk DW (g)	$2.08^{\rm b}\pm0.08$	$1.97^{b} \pm 0.11$
Tassel DW (g)	$0.03^{\circ} \pm 0.01$	$0.02^{c} \pm 0.003$
Total shoot DW (g)	$3.89^{\text{d}} \pm 0.32$	$3.82^{d} \pm 0.33$
Root DW (g)	$1.24^{e} \pm 0.29$	$1.68^{e} \pm 0.45$
Total plant DW (g)	$5.13^{\rm f} \pm 0.41$	$5.50^{\rm f} \pm 0.57$
Root weight ratio	$0.23^{\text{g}} \pm 0.03$	$0.27^{g} \pm 0.05$