Non-mycorrhizal root-associated fungi increase soil C stocks and

2 stability via diverse mechanisms

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- 8 Abstract. While various root-associated fungi could facilitate soil carbon (C) storage and therefore aid climate change
- 9 mitigation, so far research in this area has largely focused on mycorrhizal fungi, and potential impacts and mechanisms for
- 10 other fungi are largely unknown. Here, with the aim to identify novel organisms that could be introduced to crop plants to
- 11 sequesterpromote C sequestration, we assessed the soil C storage potential of 12 root-associated, non-mycorrhizal fungal
- 12 isolates (spanning nine genera and selected from a wide pool based on traits potentially linked to soil C accrual) and
- 13 investigated fungal, plant and microbial mediators. We grew wheat plants inoculated with individual isolates in chambers
- 14 allowing continuous ¹³C labelling. After harvest, we quantified C persistencecontent and stabilitystorage potential, and by
- 15 measuring pools of different origin (plant vs soil) and of different stability with long-term soil incubations and size/density
- 16 fractionation. We assessed plant and microbial community responses, as well as fungal physiological and morphological traits
- 17 in a parallel in vitro study. While inoculation with three of the 12 isolates resulted in significant total soil C increases, soil C
- 18 stability improved under inoculation with most isolates as a result of increases in resistant C pools and decreases in labile
- 19 pools and respired C. Further, these increases in soil C stability were positively associated with various fungal traits and plant
- 20 growth responses, including greater fungal hyphal density and plant biomass, indicating multiple direct and indirect
- 21 mechanisms for fungal impacts on soil C storage. We found more evidence for metabolic inhibition of microbial decomposition
- 22 than for physical limitation under the fungal treatments. Our study provides the first direct experimental evidence in plant-soil
- 23 systems that inoculation with specific non-mycorrhizal fungal strains can improve soil C storage, primarily by stabilising
- 24 existing C. By identifying specific fungi and traits that hold promise for enhancing soil C storage, our study highlights the
- 25 potential of non-mycorrhizal fungi in C sequestration and the need to study the mechanisms underpinning it.

26 1 Introduction

- 27 Despite soils having the capacity to sequester large amounts of atmospheric CO2 and mitigate catastrophic climate change, the
- 28 full potential of soil carbon (C) sequestration is yet to be realised (Field and Raupach, 2004; Scharlemann et al., 2014;
- 29 Schlesinger, 1990). Moreover, rather than being protected, soils are becoming increasingly degraded globally due to intensive

agricultural practices - a situation that may worsen as C loss potentially accelerates with future climate scenarios (Hannula and 31 Morriën, 2022; Lal, 2018). While soil C sequestration is becoming more broadly recognised as an important climate mitigation 32 strategy, and as an approach to recover the multiple ecosystem services provided by soil C (Kopittke et al., 2022), its successful 33 implementation first requires understanding of processes underpinning the storage of C in soil C storage (Dynarski et al., 2020; Smith and Wan, 2019; Von Unger and Emmer, 2018), Knowledge of soil C storage has improved substantially in recent years, 34 35 with it now understood to result from the balance of multiple, dynamic processes (that are further complicated by pedoclimatic 36 context) determining C inputs to soil and their stabilisation (i.e. resistance to decay; -(Cotrufo and Lavallee, 2022; Derrien et 37 al., 2023; Dignac et al., 2005; Dynarski et al., 2020; Jackson et al., 2017; Schmidt et al., 2011). Soil microbes act as key 38 participants of these processes, as : they regulate the stability of soil C is regulated primarily via their abilities to mineralise 39 soil organic matter. , which Thus, soil microbes determine how long C of plant or microbial origin persists in soil, and can also influence how much C is available for stabilisation from their necromass and from plant inputs. However, the soil microbial 40 41 community is complex, and largely unknown; hence, referred to as a "black box" (Mishra et al., 2023; Tiedje et al., 1999). Within this black box, fungi, both free-living and plant-associated, are considered particularly important for soil C storage; 42 43 however, their impacts on soil C storage are both multifaceted and diverse.

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44 The complexity in fungal impacts on soil C storage firstly arises from their abilities to influence both soil C inputs and their 45 stabilitysation-via multiple direct and indirect mechanisms occurring simultaneously (Hannula and Morriën, 2022; Kallenbach et al., 2016; Liang et al., 2019; Starke et al., 2021). In general, fungi that are present in soil (1) all produce hyphae and with 46 them hyphal C inputs, (2) can alter plant health, growth, and C chemistry and allocation to soil, and (3) can influence the rest 47 48 of soil microbial community structure and composition, thus impacting fungal-, plant-, and microbial-derived C, respectively (Clocchiatti et al., 2020; Hannula and Morriën, 2022; Rai and Agarkar, 2016; Stuart et al., 2022). All of these inputs, but 49 50 particularly fungal and plant C, are potentially available for soil C storage but they require stabilisation in order to persist in soil long term. The broad and efficient enzymatic capabilities and extensive mycelial structure of fungi, as compared to the 51 52 rest of the microbial community, allow them to competitively obtain soil C and transform it so that it can be readily sorbed 53 and stabilised onto mineral surfaces (Boer et al., 2005; Hannula and Morriën, 2022). In addition, fungal necromass is considered to have a particularly strong affinity for mineral surfaces and is therefore an important source of stabilisable C 54 55 (Sokol et al., 2019). The impact of fungi on soil structure and spatial heterogeneity, including promoting aggregate formation 56 by enmeshing soil particles with their hyphae and producing various extracellular biopolymers, further protects C by physically constraining microbial decomposition, leading to greater persistence (Berg and Mcclaugherty, 2014; Dynarski et al., 2020; 57 Kleber et al., 2011; Lehmann et al., 2017; Lützow et al., 2006; Schmidt et al., 2011). 58

These various impacts of fungi on soil C storage are further complicated by fungal diversity, which occurs at the inter-genus, 59

inter-species, and even down to the sub-species level (Andrade et al., 2016; Hiscox et al., 2015; Johnson et al., 2012; Juan-60

Ovejero et al., 2020; Plett et al., 2021). In plant-soil ecosystems, fungi exist either as free-living saprotrophs or as plant-

associated fungi, including mycorrhizal, endophytic, and parasitic fungi (Rai and Agarkar, 2016). Saprotrophic fungi are often 63 assumed to promote soil C output, as they decompose soil organic matter due to being outcompeted by mycorrhizal fungi for plant C exudates, but as decomposition can increase the availability of C to be sorbed onto mineral surfaces, thereby fostering 64 65 soil C stability, their net impacts on soil C storage may need further exploration (Frac et al., 2018; Hannula and Morriën, 2022; Lehmann and Rillig, 2015). Meanwhile, much of the research on the impacts of plant-associated fungi on soil C has focused 66 67 on mycorrhizal fungi, particularly arbuscular mycorrhizal fungi and ectomycorrhizal fungi due to their dominance in their 68 respective habitats (Jackson et al., 2017; Smith and Read, 2008). These fungi have additional impacts, to the general fungal impacts outlined above, on the inputs and stabilisation of C. As they transform and funnel plant C belowground, mycorrhizal 69 70 fungi can increase and modify the quality of C inputs, for example by synthesising melanin for cell walls, which is considered 71 to be highly stable and has been associated with decreased hyphal decomposability and increased soil C content (Fernandez and Kennedy, 2018; Fernandez and Koide, 2013; Zak et al., 2019; Zhu and Michael Miller, 2003). Due to their nutrient 72 73 requirements and abilities to mine soil resources, they are thought to be strong competitors against saprotrophs for not only 74 plant C but also soil nutrients, thereby suppressing microbial respiration, and resulting in greater C stability (Gadgil and Gadgil, 75 1971; Averill and Hawkes, 2016). Some mycorrhizal fungi have limited abilities to directly and partially decay organic matter, 76 and they can also prime saprotrophic microbes to decompose pre-existing soil C, thus having the potential to decrease C 77 stability persistence—though their net impact on soil C storage is not well understood (Frey, 2019). Despite the large diversity 78 amongst fungi in plant-soil ecosystems, influences of non-mycorrhizal fungi, particularly other plant-associated fungi, on soil 79 C storage have not been studied in as greatlesser detail compared to mycorrhizal fungi but do hold promise. For example, endophytic fungi could potentially be important for soil C storage due to their abilities to produce melanin and promote plant 80 81 growth (Berthelot et al., 2017; He et al., 2019; Mandyam and Jumpponen, 2005; Rai and Agarkar, 2016). However, similar to 82 mycorrhizal fungi, there are conflicting reports regarding their lifestyles, benefits or harms imposed on host plants, enzymatic 83 and nutrient acquisition ability, or even whether they produce extraradical mycelium, suggesting there may be wide functional 84 variation or plasticity within this fungal group (Addy et al., 2005; Mukasa Mugerwa and Mcgee, 2017; Rai and Agarkar, 2016). 85 To better understand the diversity of fungal impacts on soil C storage, particularly soil C stability, more focus is also needed 86 on fungal types other than mycorrhizal fungi. 87 There is growing interest in searching and screening for organisms that, in addition to supporting plant productivity, may improve soil C storage in agricultural systems (Kaminsky et al., 2019; Islam et al., 2021; Salomon et al., 2022). Thus far, 88 mycorrhizal fungi have received much attention in this area due to their well established better known impacts on plant health 89

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and soil C. However, as discussed above, other fungal types may also offer advantages to soil C storage and plant productivity

but have been largely unexplored. With this objective in mind, in the current study we aimed to determine the net potential

impacts of inoculation with diverse non-mycorrhizal fungi to impacton soil C-stocks, formation (by impacting the origin of C), and stabilitypersistence (by impacting C pools, dynamics, and fractions), and to investigate the mechanisms

94 underpinning these impacts, both direct and indirect. We assessed 12 separate fungal species (spanning nine genera in the 95 orders Chaetosphaeriales, Helotiales, and Pleosporales), isolated from roots collected from multiple soil environments across 96 Australia and screened for traits that may support plant growth and soil C storage, such as capabilities to capture and 97 solubilise nutrients from the soil. These fungi were selected with the specific aim to identify novel organisms that could 98 potentially be introduced to crop plants to improve soil C accrual. In a pot study, we inoculated spring wheat (Triticum 99 aestivum), an important cereal crop, with one of the 12 fungi and grew the plants for a full life cycle in ¹³C-depleted CO₂ 100 growth chambers to homogeneously label the plants during the full growth cycle, in order to distinguish soil C from plant-101 derived soil C. Following harvest, we assessed total C and its isotopic composition, and assessed C distribution among pools 102 of different stability and persistence (labile, intermediate, and resistant) via four-month soil incubations, and evaluated the 103 contribution of soil and plant C to these pools using isotopic analysis. These incubation-based assessments were accompanied 104 by size and density fractionation analyses to quantify mineral-associated organic matter (MAOM), aggregate carbon (AggC), 105 and particulate organic matter (POM). We then measured traits of the fungi and of the plants and microbial community to explore the potential direct and indirect mechanisms behind these impacts, respectively. We hypothesised that if a fungal 106 107 species increased total soil C storage, this would be due primarily to increasing plant C inputs by supporting plant growth and 108 also to stabilising existing soil C - so that fungi-driven increases in total soil C would be associated with more persistence and 109 stable pools and fractions of C. We expected that these changes to soil C would be associated with fungal traits, alluding to 110 direct mechanisms, as well as to increases in plant growth and shifts in microbial community composition, alluding to indirect 111 mechanisms.

2 Materials and methods

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- 114 The overall study design consisted of a wheat growth pot experiment, in which changes to soil, plant, and soil microbial
- 115 communities in response to fungal inoculation were assessed, and a separate in vitro fungal growth assay, to measure fungal
- traits that could potentially be linked to observations made in the main experiment (Fig. A1).

117 2.1 Experiment set up and maintenance

- 118 Twelve fungal isolates were originally obtained isolated from surface-sterilised plant roots of multiple species of grasses and
- 119 shrubs from across diverse natural environments in southeast Australia and screened for traits that may support plant growth
- 120 and soil C storage by Loam Bio Pty Ltd (Orange, New South Wales, Australia). Briefly, the screening process included
- 121 assessing successful colonisation of crop plants (including wheat), testing for responses of soil properties to inoculation, and
- 122 assessing interactions of the fungi with other bacteria and fungi. The fungal isolates, including endophytic fungi and potentially
- 123 saprotrophic or other fungi, comprised: a-Thozetella-species, a-Paraconiothyrium-species, three Darksidea-species, a
- 124 Leptodontidium-species, a Clohesyomyces species, two Phialocephala species, an Acrocalymma species, a Periconia species,
- 124 Leptouoniutum-species, a-Cionesyomyces-species, two I nutocephata-species, arracrocalynima-species, arracrocalynima-species,
- 125 and an Ophiosphaerella species.
- 126 Pure cultures of these isolates were maintained on 1/10 strength potato dextrose agar (PDA). Surface-sterilised (2% NaOCl)
- 127 and moistened seeds of Australian wheat cultivar Condo (Triticum aestivum) were incubated at room temperature for 48 h.
- 128 Clay loam Soil was obtained from an agricultural field where the past 10 years of land use history included wheat, barley,
- 129 canola, and sorghum (4.3% C, 0.39% N, pH 5.85; Table B1). The soil was -sieved through 2 mm, and was a clay loam (4.3%
- 130 C, 0.39% N, pH 5.85; Table B1). not sterilised before use in this experiment.
- 131 The experimental setup consisted of 12 fungal treatments (seven planted replicates per treatment) inoculated with one of the
- 132 12 fungal isolates, and six replicates of an uninoculated planted potstreatment (six replicates) applied to "planted" pots, which
- 133 were distributed among six CO₂-controlled growth chambers (Climatron-1260; Thermoline, Wetherill Park, New South Wales,
- 134 Australia). Each chamber contained one replicate per treatment for replicates 1 to 6, and replicate 7 was distributed among the
- 135 chambers that had been modified to achieve continuous 12 C labeling of plant tissues. The CO2-controlled growth chambers
- were modified using the approach by Cheng and Dijkstra (2007) to achieve continuous ¹³C-labeling of plant tissues. Briefly,
- 137 the chambers were adapted to take an influx of naturally ¹³C-depleted CO₂ (δ¹³C = -31.7 o/oo ± 1.2) during the photoperiod,
- 138 combined with a continuous supply of external CO₂-free air, and set to 450 ppm CO₂ concentration. Chambers were adjusted
- to a 16 h/8 h photoperiod, 22°C/17°C, 60% relative humidity, and 500 µmol m⁻² s⁻¹ light intensity. For "planted" replicates,
- 140 three 7 mm agar squares from actively growing 1/10 PDA fungal culture plates were placed near three sterile seeds in 2 L
- plastic pots (at a depth of 2-3 cm) containing 1800 g of the non-sterile soil. Uninoculated planted pots ("absent/control")
- 142 received three agar squares from uninoculated plates. Each agar square contained approximately 1.3 mg C. Smaller pots

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- 143 (containing 500 g of soil) for "unplanted" control pots (see belowfour replicates per treatment) were set up three days later
- 144 using two agar squares (as they contained less soil than the planted pots), as controls for impacts of fungi in the absence of
- 145 plants, adding to 142 pots in total. After 10 days of growth, seedlings were thinned to one per pot.
- 146 Pots were regularly and uniformly watered with tap water. Pots within each tub and tubs within each chamber were randomly
- 147 relocated repositioned four times throughout the experiment. The chamber atmosphere was sampled weekly to confirm that
- 148 the atmospheric CO₂ was sufficiently depleted in ¹³C via a pump system into a Tedlar® SCV Gas Sampling Bag and δ¹³C
- 149 analysis in a PICARRO G2201i isotopic CO₂/CH₄ analyser (Picarro Inc., Santa Clara, CA, USA).

150 2.2 Harvest and plant biomass measurement

- 151 Once the plants had senesced and the grain had ripened, at 18th_weeks of growth, wheat spikes and shoots were cut off, dried
- 152 at 70°C and weighed. The intact root-containing soil was preserved in the pots by freezing at -20°C immediately after shoots
- 153 were cut to stop all decomposer activity to retain the C status generated by the treatment until ready for subsampling and
- 154 processing. After two days of thawing at 4°C, soil was removed from the pots and a subsample for fractionation analysis was
- 155 collected from near the root crown and oven-dried at 40° C. The main root system was gently shaken of soil and 1/3 of the
- 156 roots were cut, washed, patted dry, frozen at -20°C prior to root morphology measurement. The rest of the soil was
- 157 homogenised before subsamples collection. A subsample for phospholipid fatty acid (PLFA) analysis was frozen at -20°C. A
- 158 subsample for soil moisture content was weighed and dried at 105°C. A subsample for soil incubations was oven-dried at
- 159 40°C and sieved at 2 mm, and of this, a further subsample for isotope analysis was dried at 105°C. To obtain total root mass,
- 160 first the root/soil ratio outside the main root system was estimated by collecting the root mass of the remaining soil (after all
- 161 subsampling) via wet sieving (500 µm) and oven-drying at 40°C. The root mass of the soil subsamples was calculated using
- 162 this ratio and the amount of soil in all subsamples.

163 2.3 Root morphology

- 164 To evaluate root morphology, a potential indirect mechanism for fungal impacts on soil C storage, washed, dried, and frozen
- 165 root subsamples were arranged with minimal overlap for digital scanning (Epson Expression 11000XL scanner, Epson,
- 166 Macquarie Park, Australia). Images were analysed with WinRhizo Pro software 2015 (Regent Instruments Inc., Quebec City,
- 167 Canada) to obtain root average diameter (mm), specific length as the ratio of length to dry mass (cm mg⁻¹), tissue density as
- mass per unit volume (g cm⁻³), specific surface area as the ratio of area to dry mass (cm² g⁻¹), and branching as the number of
- 169 forks per unit of mass (number mg⁻¹). Following root morphology assessment, the root subsample was oven-dried at 40°C
- 170 for determination of total root mass.

2.4 Plant and soil isotope and chemical analysis

- 172 To determine the contribution of soil- versus plant-derived C to total C in soils under wheat, isotopic compositions and C/N
- 173 content of ground shoots and soil were assessed using an elemental analyser interfaced to a continuous flow isotope ratio mass
- 174 spectrometer (UC Davis Stable Isotope Facility, Davis, California, USA). The proportion of original soil C present in the soil
- 175 of each pot after plant growth was calculated via isotopic partitioning following Eq. (1):
- 176 Soil proportion. Soil = $\frac{(\delta^{13}C_{Soil} \delta^{13}C_{UP} Soil)}{\delta^{13}C_{PP} \delta^{13}C_{UP} Soil},$
- 177 where $\delta^{13}C_{Soil}$ is the ^{13}C isotopic composition of soil measured in each planted pot, $\delta^{13}C_{UP-Soil}$ is the mean ^{13}C isotopic
- 178 composition of soil in unplanted controls, and $\delta^{13}C_P$ is the ^{13}C isotopic composition of the plant shoots in each planted pot.
- 179 The plant C proportion (including C from other biological sources) was defined as 1 minus the soil C proportion. These
- 180 proportions were then applied to the measured C concentrations in each pot to calculate plant- and soil-derived C amounts.

181 2.5 Soil incubations

- 182 To evaluate the fungal impacts of fungal isolates on overall C persistence and on C distribution across pools of different
- 183 stability (labile, intermediate, and resistant), we assessed microbial CO₂ production during 135-day laboratory incubations of
- 184 soil harvested at the time of wheat harvest. Headspace samples from incubation jars containing 30 g soil, incubated -under
- 185 standard temperature and moisture conditions (25°C and 42% gravimetric moisture, respectively), were collected on 16
- 186 occasions over the course of 135 days. Following incubation, we and fitted a decay model (i.e. exponential decay equations)
- to estimate decay kinetic parameters. Kinetic parameters derived from mid- to long-term soil incubation are sensitive functional
- 188 measures of changes in the distribution and stability of C pools resulting from previous exposure to experimental treatments
- 189 (Carney et al., 2007; Carrillo et al., 2011; Jian et al., 2020; Langley et al., 2009; Taneva and Gonzalez-Meler, 2008). Measured
- 190 CO₂ production rates over time were fitted to a two-pool exponential decay model to estimate the size of the labile and
- 191 intermediate C pools and their mean residence time (MRT; Cheng and Dijkstra, 2007; Wedin and Pastor, 1993). The size of
- the resistant pool was calculated as the difference between the total measured organic C and the sum of the estimated labile
- and intermediate pools. This same procedure was also applied to the portion of CO₂ that was released from the originally
- present soil C (soil-derived C, i.e. not plant-derived C), which was determined via isotopic partitioning of plant vs. soil-derived
- 195 CO₂. Based on these, we calculated total CO₂ released from plant- and soil-derived C during the full length of the incubation.
- 196 See Supplementary Methods for full details on incubations, isotopic partitioning, and decay curve fittingmodelling.

2.6 Soil fractionation analysis

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- 198 Soil fractionation analysis was performed as an alternative method to soil incubations for understanding fungal impacts on C
- 199 stability. Hereafter we refer to the pools measured via fractionation analysis as "fractions", as opposed to "pools" measured
- 200 via soil incubations. The analysis was performed according to a method developed by (Poeplau et al., 2017; Poeplau et al.,
- 201 2018) and adapted by Buss et al. (2023, in review) involving high throughput physical fractionation into conceptually
- 202 designed soil C fractions mineral-associated organic matter (MAOM), aggregate carbon (AggC), and particulate organic
- 203 matter (POM). See Supplementary Methods for further details.

204 2.7 Soil PLFA analysis

- 205 Total microbial community size and composition are also potential indirect drivers of fungal impacts on soil C storage.
- 206 Microbial PLFAs in soils were extracted from 2 g of freeze-dried soil harvested from the wheat growth experiment, following
- 207 the high throughput method developed and described by Buyer and Sasser (2012; see Supplementary Methods).

208 2.8 In vitro fungal assessment

- 209 To assess morphological and chemical properties of the fungal isolates (used in the wheat growth experiment) as potential
- 210 drivers of fungal impacts on soil C storage, a separate in vitro plate assay was performed using 1/2 PDA plates incubated in
- $211 \quad \text{the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by measuring colony areas every two-points of the dark at } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate was calculated by } 23\text{-}25^{\circ}\text{C (see Supplementary Methods)}. \ \text{Radial growth rate } 23\text{-}25$
- 212 to-three days using ImageJ (National Institutes of Health, Bethesda, Maryland, US; Schneider et al., 2012). Growth rate was
- 213 calculated by subtracting the colony area from an earlier sampling point from that of the following sampling point. Hyphal
- 214 density was calculated as the final fungal biomass per final colony area. C and N content were measured by Dumas combustion
- 215 using a El Vario cube analyser (Elementar, Langenselbold, Germany).

216 2.9 Data and statistical analysis

- 217 ANOVA of soil C properties and experimental variables was performed in R (v. 4.1.2; R Core Team, 2021), followed by
- 218 Dunnett's post-hoc test to determine which treatment groups were significantly different to the uninoculated control group or
- Tukey's post-hoc test to determine significant differences between inoculated groups. Principal component analysis (PCA) of
- 220 soil C property data was performed to identify soil C properties associated with fungi-driven increases in soil C. Redundancy
- 221 analyses (RDA) of soil C property data as response variables and either plant and microbial community data or using *in vitro*
- 222 fungal assessment data as explanatory variables were performed to identify explanatory variables for fungi-driven increases in
- 223 soil C and its stability. Both analyses and redundancy analysis (RDA) were performed using the vegan package in R (Oksanen

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et al., 2020). Missing values (17 values across 46 total variables) in the PCA and RDA datasets were replaced with the variable treatment mean.

Curve fitting of CO₂ rate dynamics was done using the non-linear modelling platform in JMP 16.1.0 and the biexponential four-parameter decay model using all replicates of a treatment. We used nonlinear least square curve fitting to test if the models were significantly different between a fungal treatment and uninoculated control, using the nls function in R.

230 **3 Results**

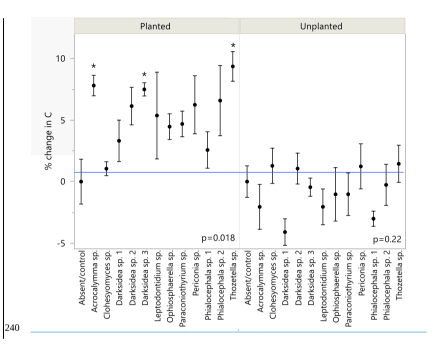
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3.1 Several non-mycorrhizal fungal species increased soil C under wheat plants

months of plant growth, there was a positive but varied effect of fungal inoculation on soil C content compared to the uninoculated control group (p < 0.05; Fig. 1, Table B2). This effect was not observed in soils that received the same fungi but were unplanted (p = 0.22; Fig. 1). We found significant isolate-specific increases in soil C content of the planted treatments under inoculation with *Thozetella* sp., *Darksidea* sp. 3, and *Acrocalymma* sp., relative to the uninoculated control, of 9.4% (percentage of change), 7.5, and 7.8, respectively. Nitrogen levels were generally higher in the soils of the inoculated and planted treatments compared to the uninoculated control and were generally higher in the treatments where C was also higher (Table B2).

We inoculated wheat plants (Triticum aestivum) with one of 12 fungi (non-mycorrhizal) isolated from plant roots. After four



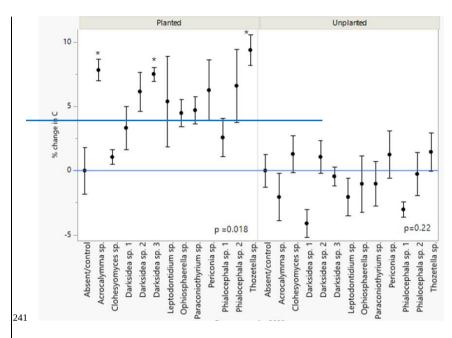
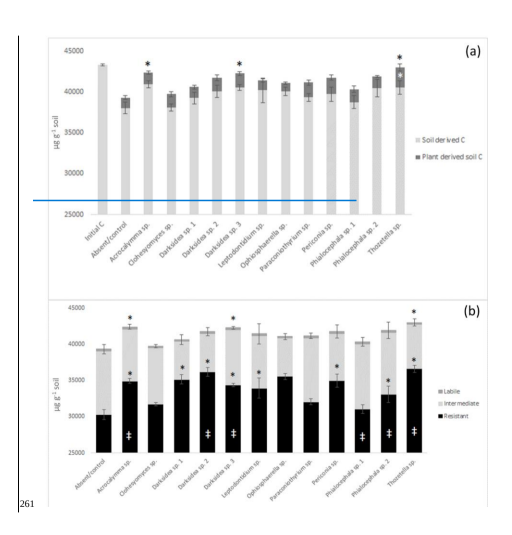


Figure 1. Changes in total soil C under inoculation with different fungal isolates compared to an uninoculated control. Values indicate percentage of change relative to mean of uninoculated control (blue line). Error bars indicate standard error, n=7 for inoculated treatments, n=6 for control. ANOVA results for planted and unplanted are presented. Asterisks indicate significant differences with control (Dunnett test, p < 0.05). C concentrations are presented in Table B2.

3.2 Fungi-dependent increases in soil C are associated with changes in soil C pools, origin; and stabilitypersistence

To understand the underlying mechanisms of the fungal isolate-dependent increases in soil C content and potential shifts in sources and stability of the resulting soil C, we performed C isotope analysis, soil incubations, and soil C fractionation analysis. Isotopic partitioning of C into plant- and soil-derived C revealed how changes in these pools contributed to changes in total soil C (Fig. 2a, Table B2). Planting reduced total soil C, compared to initial C prior to planting (t = 4.13, p < 0.001), as expected due to C inputs stimulating decomposition (rhizosphere priming). This reduction was due to decreases in soil-derived C, which were generally not counteracted by newly added plant-derived soil C - which on average represented 3.8% (± 0.2) of total soil C (Fig. A2a). Soil C increases under fungal inoculation had different origins depending on the fungal treatment. Some increases

in total soil C compared to the planted uninoculated controls could be explained by plant—and soil derived C. Namely, oOne
of the fungal treatments whereby total soil C significantly increased (*Thozetella* sp.) tended to contain higher levels of plant—derived C (p = 0.06)exhibited higher amounts of plant—derived C—at a level that was marginal in its non-significance. However,
overall, the higher total soil C content relative to controls corresponded correlated more closely with higher soil-derived C
(Pearson's R = 0.93, p < 0.01), than with plant-derived C (Pearson's R = 0.02, p = 0.83). All three fungal treatments resulting
in significant increases in total soil C showed increases in soil-derived C but these were not statistically significant.



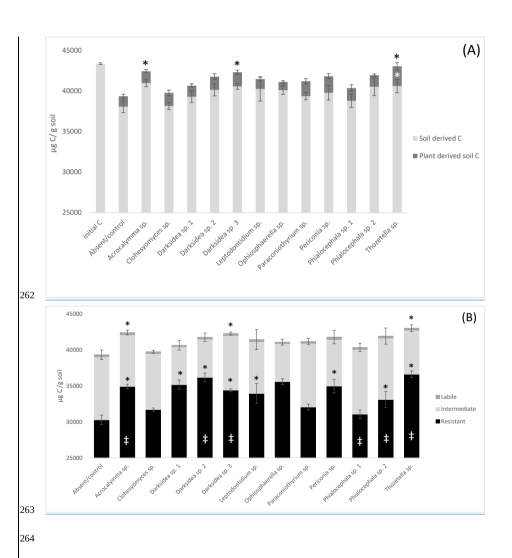


Figure 2. Distribution of total soil C in plant- and soil-derived pools (A) and among labile, intermediate, and resistant pools (B) in soil under inoculation with different fungal isolates or under no inoculation (Absent/control). (A): Plant-

267 and soil-derived C from C isotope partitioning (see Materials and methods). Black asterisks indicate significant 268 differences in total C with control and white asterisks differences in plant-derived soil C with control (Dunnett test, p 269 < 0.1); (B): Pools estimated from decay models derived from soil incubation (see Materials and methods). Crosses 270 indicate significant differences in the dynamics of total C decomposition (decay curves models, Table B3) compared to the uninoculated control. Asterisks indicate significant differences in total C or resistant C against control (Dunnett 271 272 test, p < 0.05). Error bars indicate standard error of total C, n=7 for inoculated treatments, n=6 for uninoculated 273 control. Note y axis scale.

274 Incubation of soils after plant harvest demonstrated impacts of several fungal species on the dynamics of C decomposition and 275 the distribution of C among soil pools of different stability. The dynamics of total C decomposition (decay curves models

276 derived from incubations) were significantly different to the control under half of the isolates (Table B3, Fig. A23). These

included the three isolates that produced higher total C pools: Thozetella sp., Darksidea sp. 3, and Acrocalymma sp. Soil-

derived C decomposition curves (from isotopic partitioning of respiration) were also significantly different to the controls

under the same fungal treatments as well as Leptodontidium sp. Estimated pools from these decay curves showed significantly

280 higher total resistant C (up to 86% of C), compared to controls (76% of C), under eight of the 12 isolates, including the three 281 treatments where total C increased the most (Fig. 2b, Fig. A2b, Table B3). In terms of other pools, MRT of the total labile C

was significantly lower under inoculation with Darksidea sp. 1 compared to the control, whereas MRT of the soil-derived 282 283 labile C was significantly higher under inoculation with Periconia sp. (Table B3). In terms of intermediate pool MRTs, controls

and fungal treatments were not significantly different. 284

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285 Soil incubations and partitioning of respiration revealed fungal effects on the degree of stability persistence of total C, soilderived C, and plant-derived C over time, which we assessed as the proportion of what was present at harvest that was respired 286 287 over the full incubation. Significantly lower proportions of total and soil-derived C were respired under all fungal treatments

compared to the controls (p < 0.001; Fig. A34), indicating increased stabilitypersistence. In contrast, plant-derived respired C 289 was significantly lower (more <u>stable persistent</u>) than the controls only with *Thozetella* sp. (p < 0.05).

290 From fractionation analysis, %C and %N of the AggC fraction, i.e. the fraction of intermediate stability whereby C is protected 291 in aggregates, were found to have significant fungal effects, with Thozetella sp. and Periconia sp. exhibiting significantly

292 higher levels of both C and N, and Ophiosphaerella sp. and Phialocephala sp. 1 exhibiting significantly higher levels of N

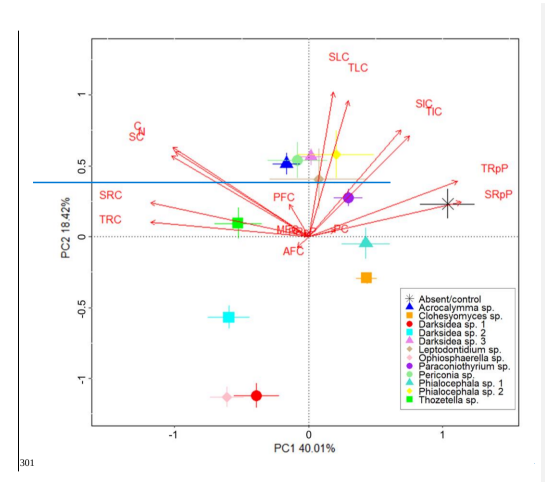
293 compared to controls (Table B4). Significant fungal effects were not observed in the MAOM and POM fractions.

294 We performed PCA to identify soil C properties associated with fungi-driven increases in soil C (Fig. 3). Most of the variance

was explained by PC1 and 2 (58%). Greater total soil C (C) was closely associated with soil-derived C (SC), but not plant-295

derived C (PC), at time of harvest and soil N. Soil C was also related with the resistant C pools (total (TRC) and soil-derived 296

(SRC)). The treatments with lowest total soil C (mainly the control, followed by *Clohesyomyces* sp., and *Phialocephala* sp. 1;
 Fig. 1) were associated with higher proportions of total and soil-derived C respired during incubation indicating that the C remaining at harvest was inherently less persistent stable. %C of the AggC and MAOM fractions, generally considered to be more stable fractions of C, were not clearly associated with soil C or the resistant C pools, nor with any fungal treatments.



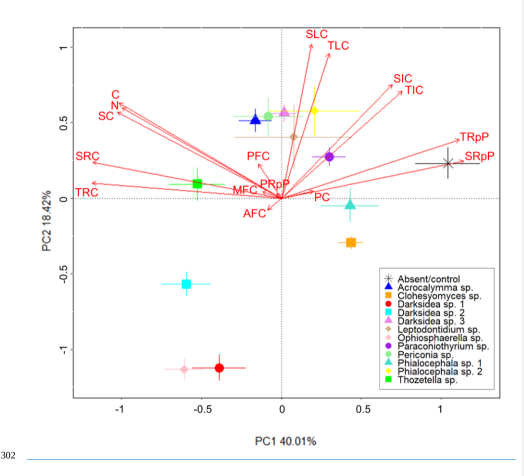


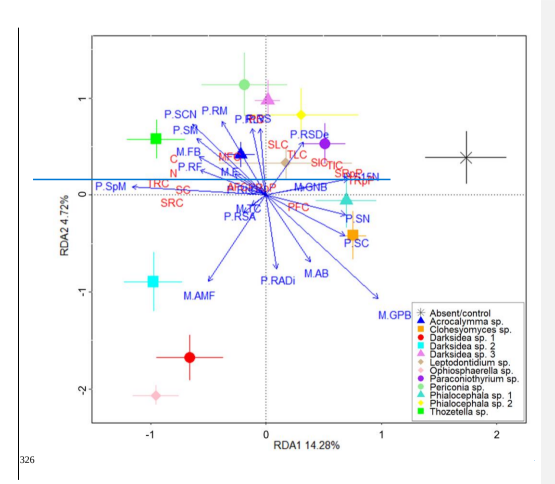
Figure 3. Fungi-dependent increases in soil C largely relate to measures for soil C stability. Principal component analysis showing soil C properties (red text) associated with various fungal isolates (symbols). Soil C properties were measured via isotope analysis, soil incubations, and fractionation analysis of soil from wheat experiment. Soil C property abbreviations: AFC, aggregate C fraction %C; C, %C; MFC, MAOM fraction %C; N, %N; PC, plant-derived C (μ g g⁻¹ soil); PFC, POM fraction – %C; PRpP, plant-derived C respired proportion; SC, soil-derived C (μ g g⁻¹ soil); SIC, soil-derived intermediate C (μ g C g⁻¹ soil); SLC, soil-derived labile C (μ g C g⁻¹ soil); SRC, soil-derived

311 3.3 Fungi-dependent increases in soil C and its stability and persistence are positively associated with plant growth and 312 microbial community composition We assessed plant and microbial community variables, including plant biomass, shoot C/N content, root morphology, and total 313 314 microbial community size and composition derived from PLFA analysis. Overall, while variation among fungal isolates was observed, no significant differences were observed between the inoculated and uninoculated plants for any of the plant or 315 microbial community variables measured, although average spike mass of Thozetella-inoculated plants was significantly 316 317 higher than that of uninoculated plants (Table B5-6). To identify plant and microbial community variables potentially involved in the fungal isolate-dependent changes in soil C 318 319 properties, we performed RDA using plant and microbial community data and the soil C property data used in the PCA (Fig. 320 4). Variance explained by RDA1 and 2 was 14.28 and 4.72%, respectively. The cluster of soil C properties that were found to be closely associated with Thozetella sp. in the PCA (e.g. soil-derived C, resistant C pools; Fig. 3) also trended positively with 321 322 plant biomass and growth (spike and shoot mass, shoot C/N ratio, and root fork number) and with the PLFA-assessed fungal 323 to bacterial ratio. Acrocalymma sp. and Darksidea sp. 3 were more associated with root growth traits, and were also associated 324 with plant-derived C. The low soil C treatments (uninoculated control, Clohesyomyces sp., and Phialocephala sp. 1) and their 325 associated soil C properties (i.e. respired C) were related to shoot C and N.

resistant C (µg C g⁻¹ soil); SRpP, soil-derived C respired proportion; TIC, total intermediate C (µg g⁻¹ soil); TLC, total

labile C (µg g⁻¹ soil); TRC, total resistant C (µg g⁻¹ soil); TRpP, total C respired proportion.

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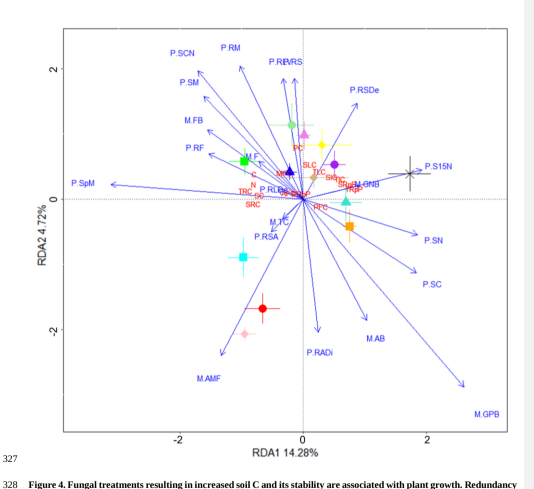
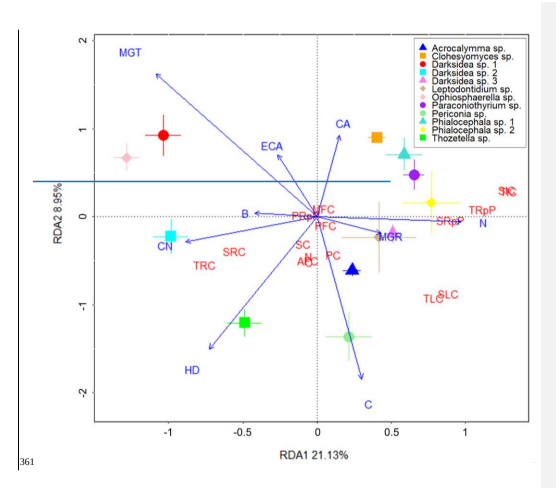


Figure 4. Fungal treatments resulting in increased soil C and its stability are associated with plant growth. Redundancy analysis showing microbial community and plant variables (blue text) driving changes in soil C properties (red text) associated with various fungal isolates (symbols). Soil C properties were measured via isotope analysis, soil incubations, and fractionation analysis of soil from wheat experiment. Microbial community and plant variables were measured using samples harvested from the wheat experiment. Microbial community (M.) and plant (P.) variable abbreviations: M.AB, actinobacteria (% of total community); M.AMF, arbuscular mycorrhizal fungi (% of total community); M.F.

- $334\quad \text{fungi (\% of total community); M.FB, fungal to bacterial biomass \ ratio; M.GNB, \ gram \ negative \ bacteria \ (\% \ of \ total)}$
- community); M.GPB, gram positive bacteria (% of total community); M.TC, total community size (μ g PLFA g^{-1} soil);
- 336 P.RADi. root average diameter (mm): P.RF, root fork number (g⁻¹): P.RLDe, root length density (cm g⁻¹): P.RLV, root
- 337 length per volume (cm m³); P.RM, root mass (g); P.RS, root/shoot ratio; P.RSA, root specific surface area (cm² g²l);
- 338 P.RSDe, root specific density (g cm⁻³); P.S15N, shoot δ15N (‰); P.SC, shoot %C; P.SCN, shoot C/N ratio; P.SM, shoot
- 339 mass (g); P.SN, shoot %N; P.SpM, total spike mass (g). Soil C properties: AFC, aggregate C fraction %C; C, %C;
- 340 MFC, MAOM fraction %C; N, %N; PC, plant-derived C (µg g⁻¹ soil); PFC, POM fraction %C; PRpP, plant-
- 341 derived C respired proportion; SC, soil-derived C (ug g⁻¹ soil); SIC, soil-derived intermediate C (ug C g⁻¹ soil); SLC,
- 342 soil-derived labile C (µg C g⁻¹ soil); SRC, soil-derived resistant C (µg C g⁻¹ soil); SRpP, soil-derived C respired
- 343 proportion; TIC, total intermediate C (µg g⁻¹ soil); TLC, total labile C (µg g⁻¹ soil); TRC, total resistant C (µg g⁻¹ soil);
- 344 TRpP, total C respired proportion.
- 345 3.4 Fungi-dependent increases in soil C and its stability and persistence are associated with denser fungal hyphae and
- 346 <u>higher fungal C/N ratio</u>
- 347 Fungal isolates showed strong differentiation in all of the in vitro-assessed variables relating to growth and C/N content
- 348 (statistically significant effects on all variables, p < 0.001; Table B7). Biomass, colony area, and growth rate tended to be
- 349 positively associated variables, and were higher in several treatments including Acrocalymma sp., Darksidea sp. 3, and
- 350 Phialocephala sp. 1. In contrast, Thozetella sp. and Clohesyomyces sp. tended to have lower values for these variables, but
- 351 Thozetella sp. had significantly higher hyphal density than all other treatments.
- 352 We performed a separate RDA to identify fungal variables potentially involved in increases in fungi-dependent soil %C and
- 353 soil Cits stability increases, using in vitro fungal assessment data and the soil C property data (Fig. 5). Compared to the RDA
- 354 using plant and microbial community data (Fig. 4), greater proportions of variance were explained in this RDA by RDA1 and
- 355 2 (21.1 and 9%, respectively). Fungal colony area and hyphal density were close to opposite in their direction, with the high
- 356 soil C treatment Thozetella sp. closely associated with hyphal density and the low soil C treatment Clohesyomyces sp. more
- 357 associated with colony area. Similarly, fungal colony maximum growth time and rate (denoting slower and faster fungal
- 358 growth, respectively) were in opposing directions. Along this axis, the high soil C treatment Darksidea sp. 3 was closely
- 359 associated with maximum fungal growth rate. Respired C proportions were closely associated with fungal N content and were
- 360 opposite resistant C fractions, which were associated with fungal C/N ratio and hyphal density.



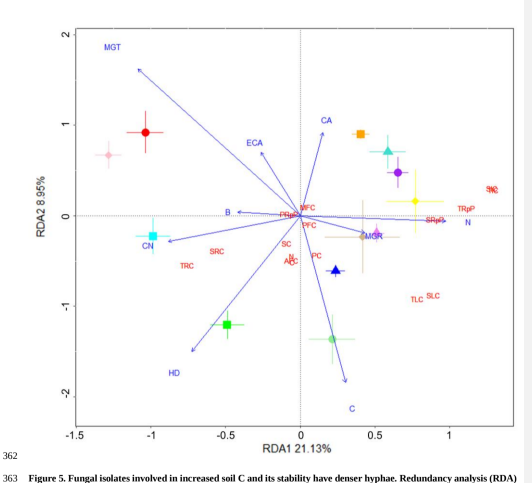


Figure 5. Fungal isolates involved in increased soil C and its stability have denser hyphae. Redundancy analysis (RDA) showing the fungal variables (blue text) driving changes in soil C properties (red text) associated with the various fungal isolates (symbols). Soil C properties were measured via isotope analysis, soil incubations, and fractionation analysis of soil from wheat experiment. Fungal variables were measured in an *in vitro* plate assay and values were averaged for the RDA. Fungal (F.) variable abbreviations: F.B, biomass (g); F.C, %C; F.CA, final colony area (cm²); F.CN, C/N ratio; F.ECA, estimated final colony area (cm²); F.HD, hyphal density (mg cm²); F.MGR, maximum growth rate (cm²)

- ⁶⁹ day); F.MGT, time to maximum growth (days); F.N, %N. Soil C properties: AFC, aggregate C fraction %C; C,
- $370 \quad \text{\%C; MFC, MAOM fraction} \text{\%C; N, \%N; PC, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRpP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRPP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRPP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRPP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRPP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRPP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRPP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRPP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRPP, plant-derived C (μg g$^{-1}$ soil); PFC, POM fraction} \text{\%C; PRPP, plant-derived C (μg g$ g$^{-1}$ soil); PFC, PO$
- $371 \quad \text{derived C respired proportion; SC, soil-derived C (μg\, g^1$\, soil); SIC, soil-derived intermediate C (μg\, C$\, g^1$\, soil); SLC, soil-derived intermediate C (μg\, C$\, g^1$\, soil-derived intermediate C (μg\, C$\, g^1$\, soil-derived intermediate C (μg\, C$\, g^2$\, soil-derived intermediate C (μg\, C$\, g^2$\, soil-derived intermediate C (μg\, C$\, g^2$\, soil-derived C (μg\, C$\, g^2$\,$
- 372 soil-derived labile C (μg C g^{-1} soil); SRC, soil-derived resistant C (μg C g^{-1} soil); SRpP, soil-derived C respired
- 373 proportion; TIC, total intermediate C (μg g⁻¹ soil); TLC, total labile C (μg g⁻¹ soil); TRC, total resistant C (μg g⁻¹ soil);
- 374 TRpP, total C respired proportion.

4 Discussion

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376 Discussions on soil C sequestration as a climate change strategy have largely focused on one side of the soil C storage system 377 - increasing C inputs into soil (promoting soil C formation). However, increased soil C storage can also be achieved through 378 reductions in soil C outputs due to the complex and dynamic nature of soil C, reductions of soil C outputs (or, increases in soil 379 C stability and persistence) must also be attained in order to foster soil C storage. In this study, we drew our attention to fungi 380 that have potential in improving soil C storage but that are often overlooked in this area of research, using a high resolution, 381 multifaceted approach combining isotopic labelling, soil incubations, and soil fractionation analysis, as well as an in vitro 382 study in parallel. Our study supports the notion that inoculation with non-mycorrhizal root-associated fungi can improve soil 383 C storage via multiple direct and indirect mechanisms determining C inputs and stabilisation. Mechanisms that increased the 384 stability of existing C were more common across the diverse fungal treatments than those increasing the input of new C.

Despite our finding that bulk soil C increased significantly under only three fungal treatments, in support of our hypothesis 385 386 our incubations revealed significant increases in directly and functionally assessed soil C stability (i.e. increases in resistant 387 pools and decreases in respired C during incubation) under most of the fungal treatments, with the stabilised C being original 388 soil C, not new inputs of C. Thus, as well as contributing to evidence that fungial inoculation can lead to increased soil C 389 content (e.g. Kallenbach et al., 2016), our study provides direct evidence from plant-fungi soil systems for non-mycorrhizal 390 fungi-driven improvements to soil C storage primarily via enhanced stability of soil C. This is emphasised by our findings that 391 the treatments whereby soil C content was the lowest (control, Clohesyomyces sp., and Phialocephala sp. 1) were associated 392 with higher proportions of total and soil-derived C respired during incubation - indicating that the C remaining at harvest under 393 these treatments was inherently more prone to decomposition (i.e. less persistent stable). Increased stability and persistence of soil C primarily results from inhibition of microbial decomposition (Cotrufo and Lavallee, 2022), which can occur by a variety 394 395 of reasons including reduced saprotrophic activity due to microbes being outcompeted for nutrients (Boer et al., 2005), 396 increased input of fungal, more readily stabilised C (Sokol et al., 2019), and increased soil aggregation (Lehmann et al., 2020). 397 We investigated multiple potential mediators for the observed increases in soil C stability and persistence in our study and 398 found some leads. We found that increased fungal C/N ratio and hyphal density may be important for stability of soil C (while 399 fungal N corresponded with decreased stability). Fungi with denser hyphae can promote soil aggregation, as soil particles get more entangled and stabilised in dense hyphae (Dignac et al., 2017). Our study substantiates previous assertions that fungal 400 401 trait expression is relevant to soil C stability: fungi that exhibited an exploitative growth strategy (denser hyphae) were found 402 to more closely associated with soil C stability and persistence, while fungi that exhibited a more exploratory strategy (faster 403 growth) were positively associated with respired C and less stable C pools (Camenzind et al., 2020; Fernandez et al., 2019; 404 Fernandez and Koide, 2013; Jackson et al., 2017; Lehmann et al., 2020; Schmidt et al., 2011; Zanne et al., 2020). These 405 findings support the notion that an exploitative growth strategy may be more conducive to competition with saprotrophs for 406 nutrients, leading to reduced decomposition (Bödeker et al., 2016).

407 Our PLFA-assessed finding regarding fungal to bacterial ratio points towards a second likely mechanism for the increases in 408 soil C stability - increased input greater proportion of fungal C, which becomes stabilisable necromass. Fungal necromass is a significant source of soil C inputs, and can in some cases make up the majority of SOM (Wang et al., 2021). Substrates with 409 410 high C/N ratios, such as fungal biomass or necromass, are generally associated with reduced decomposition rates, although C/N ratio is not the sole determinant of substrate decomposition and C/N ratios can in fact be altered by, rather than alter the 411 412 activity of, soil microbial communities (Marañón-Jiménez et al., 2021; Smith and Wan, 2019; Schnecker et al., 2019). 413 Compared with other substrates, however, necromass is a particularly stabilisable form of C as it can bind to the surfaces of MAOM or be stabilised on aggregates, where it is physically protected from decomposition (Sokol et al., 2019). For these 414 415 reasons, we expected to see positive associations between soil C stability and aggregate and MAOM soil fractions, which are 416 considered to signify increased and longer-term stability and persistence (Dynarski et al., 2020; Hemingway et al., 2019; Islam 417 et al., 2022; Poeplau et al., 2018; Poeplau et al., 2017). However, in our study these fractions were not strongly associated with 418 soil C content, or its distribution in pools (persistence), nor were they as influential on differences between fungal treatments. 419 While this lends support to the notion that microbial decomposition of soil C was metabolically inhibited (as discussed above). rather than physically limited, our findings may be explained to some extent by methodology. A potential explanation for our 420 421 findings is that although fungal necromass may have been abundant, the experimental conditions may have been unsupportive 422 of MAOM formation (e.g. the high C content of the unplanted soil may have meant that MAOM content was already at 423 saturation level and new MAOM was not able to form). Other potential explanations are that the MAOM fraction could 424 possibly take longer than the experimental timeframe to change substantially, or that the MAOM estimation method may carry 425 greater error, thus making detection of responses more difficult. Nonetheless, our study detected increases in total C, and C 426 resistance stability that were not associated with MAOM, suggesting that soil fractionation analyses do not entirely accurately 427 reflect natural soil C distribution and stability which can be detected functionally via soil incubations. Further studies utilising 428 the combined approach of soil incubations and soil fractionation analysis, such as studies using soil with lower C content or 429 studies over a longer time period, may shed light on how findings from the two methods can be compared. However, our 430 findings call for caution in directly equating operationally defined MAOM pools and their size with C stability and suggest 431 that functionally assessing C dynamics may be more effective in some cases.

In terms of improvements to soil C content, of the three fungal treatments whereby soil C increases were significant, these
only one waswere accompanied by increases in plant-derived C only under inoculation with Thozetella sp.]. While we
expected that there would be some variation in the fungal impacts on soil C storage due to the diversity amongst the fungi
included in this study, this finding is in contrast to our expectation that increases in plant-derived C would be the main
mechanism involved in C increase. As plant growth promotion and changes in nutrient uptake is a well-known characteristic
of some fungi (Hossain et al., 2017), the increase in plant-derived C with Thozetella sp. may have been related to the increases
in quantity or quality of plant inputs related to the shifts in plant variables of Thozetella sp. (spike mass, shoot biomass, and

shoot C/N ratio). Our results from the isotopic partitioning of respiration from soil incubations further indicate that the plantderived C present in soil and that contributed to total soil C increase under inoculation with *Thozetella* sp. was more persistent
stable compared to the control or other treatments. Fungal-derived C could also have contributed to size and persistence
stability of plant-derived C, if the fungi took up plant-derived C. Thus, in addition to increasing plant inputs, *Thozetella* sp.
appears to have been more active in stabilising those inputs via the mechanisms discussed above.

444 Our study addresses key knowledge gaps in the ways fungi affect soil C storage. We have explicitly demonstrated that inoculation with non-mycorrhizal fungi can improve soil C content and, moreover, soil C stability - supporting the general 445 446 agreement in this field that microbial transformations of soil C and microbially driven changes to soil structure are as important, 447 if not more important, than the characteristics of the inputs themselves for soil C storage (Dynarski et al., 2020; Hannula and 448 Morriën, 2022). When it comes to evaluating the potential of fungi to support soil C storage, our findings indicate that it is 449 important to consider not only increases in soil C but also their impact on the stability and persistence of C. Among the diverse fungi studied, these improvements in soil C stability largely resulted from reductions in C outputs by increasing stable C pools 450 451 and resistance of existing soil C to decomposition. We emphasise that these findings from our study are net outcomes of fungal 452 inoculation, which can impact soil C either via direct mechanisms, or indirect mechanisms, including interactions of the fungi 453 with the surrounding soil ecosystem. While potential mechanisms behind these the improvements in soil C stability depended 454 on fungal identity, our study points towards metabolic inhibition (rather than physical limitation) of microbial decomposition 455 for which growth characteristics such as density of fungal hyphae and fungal C/N ratio may be important indicators - thus, fungal trait expression may be a proxy for fungal influences on soil C storage. However, more work is needed to test whether 456 457 or not physical limitation of microbial decomposition leads to enhanced soil C stability by these fungi. More rarely, the 458 improvements to soil C storage involved the effects of fungal inoculation on host plant growth and C inputs (directly as plant or plant-derived fungal C). While total soil C content increased significantly only under a minority of fungal treatments, the 459 460 significant and common fungi-driven increases in stability we observed could potentially lead to even greater increases in soil 461 C content and its persistence over time - however experiments with longer timeframes are needed to test this idea. This study 462 and continued work will advance knowledge of these mechanisms and support the search and potential implementation of 463 root-associated fungi to improve soil C storage, which will aid soil C sequestration strategies.

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Appendices 471

Appendix A

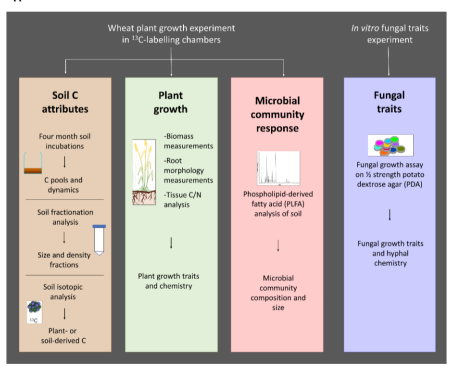
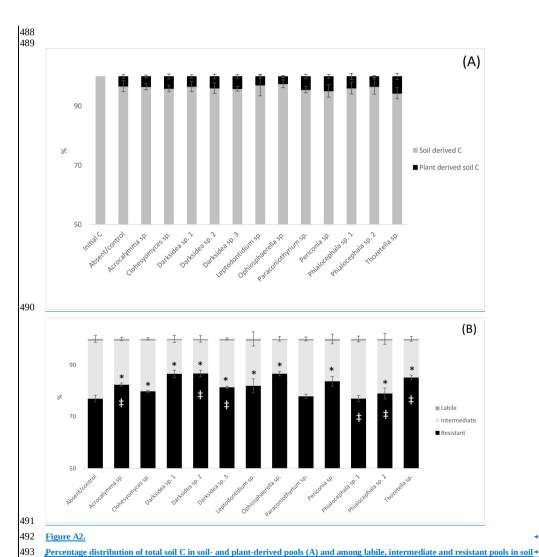


Figure A1. Overview of the study design, measured traits, and methodology used. C, carbon, N, nitrogen.



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under inoculation with different fungal isolates or under no inoculation (absent/control) (B). (A): Percentages of soil- and plant-

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derived C from C isotope partitioning (see Materials and methods). (B): Percentage distributions of pools estimated from decay
models derived from soil incubations (see Materials and methods). Crosses indicate significant differences in the dynamics of total
C decomposition (decay curves models, Table B3) compared to the uninoculated control. Asterisks indicate significant differences
in total C or resistant C against control (Dunnett test, p < 0.05). Error bars indicate standard error of total C, n=7 for inoculated
treatments, n=6 for uninoculated control. Note y axis scale.

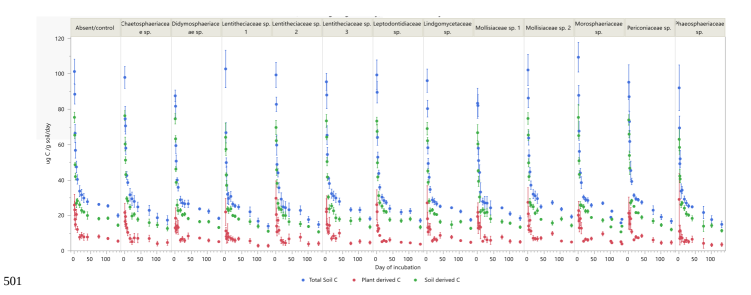


Figure A23. Total soil respiration and its soil- and plant-derived components during laboratory soil incubations of soils collected after plant growth with inoculation of 12 fungal species and a control (Absent/control). Data points are means (n=7 for 33noculated pots; n=6 for controls). Soil and plant components calculated from isotopic partitioning based on planted and unplanted soil δ^{13} C. Error bars are standard error. Family (Genus): Chaetosphaeriaceae sp. (*Thozetella* sp.); Didymosphaeriaceae sp. (*Paraconiothyrium* sp.); Lentitheciaceae sp. 1 (*Darksidea* sp. 1); Lentitheciaceae sp. 2 (*Darksidea* sp. 2); Lentitheciaceae sp. 3 (*Darksidea* sp. 3); Leptodontidiaceae sp. (*Leptodontidium* sp.); Lindgomycetaceae sp. (*Clohesyomyces* sp.); Mollisiaceae sp. 1 (*Phialocephala* sp. 1); Mollisiaceae sp. 2); Morosphaeriaceae sp. (*Acrocalymma* sp.); Periconiaceae sp. (*Periconia* sp.); Phaeosphaeriaceae sp. (*Ophiosphaerella* sp.)

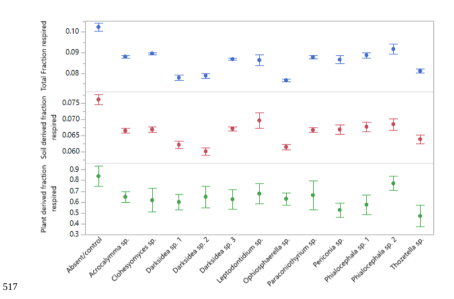


Figure A34. Fraction of soil carbon (C) respired over the course of 135-day incubation of soils under wheat and 12 types of fungal inoculum. Total C is all C respired, and soil- and plant-derived C were obtained from isotopic partitioning of respiration over time (See Materials and methods). Values are means of n=7 for treatments and n=6 for control. Error bars are standard error.

Appendix B529

 ${\bf Table~B1.~Chemical~and~physical~analysis~of~pre-planted~soil~used~in~wheat~experiment.~Analysis~was}$

performed by Environmental Analysis Laboratory (East Lismore, Australia).

Parameter	Units	Value	
Phosphorus	mg kg ⁻¹	151	
pH		5.85	
Electrical conductivity	dS m ⁻¹	0.232	
Estimated organic matter	% OM	7.5	
-	cmol kg ⁻¹	8.9	
Exchangeable calcium	kg ha ⁻¹	4010	
	mg kg ⁻¹	1790	
	cmol kg ⁻¹	2.9	
Exchangeable magnesium	kg ha ⁻¹	795	
	mg kg ⁻¹	355	
	cmol kg ⁻¹	3.1	
Exchangeable potassium	kg ha ⁻¹	2719	
	mg kg ⁻¹	1214	
	cmol kg ⁻¹	0.32	
Exchangeable sodium	kg ha ⁻¹	164	
	mg kg ⁻¹	73	
	cmol kg ⁻¹	0.02	
Exchangeable aluminium	kg ha ⁻¹	3.1	
	mg kg ⁻¹	1.4	
	cmol kg ⁻¹	0.06	
Exchangeable hydrogen	kg ha ⁻¹	1.2	
	mg kg ⁻¹	<1	
Effective cation exchange capacity	cmol kg ⁻¹	15	
Calcium	%	58	
Magnesium	%	19	
Potassium	%	20	
Exchangeable sodium	%	2.1	
Aluminium	%	0.1	
Hydrogen	%	0.36	
Calcium/magnesium ratio		3.1	
Total carbon	%	4.3	
Total nitrogen	%	0.39	
Carbon/nitrogen ratio		11	
Basic texture		Clay loam	
Basic colour		Brownish	
Chloride estimate	(equiv. mg kg ⁻¹)	148	

Table B2. Properties of soil in which inoculated wheat plants were grown for four months. P-values from ANOVA are displayed in the bottom row.

Asterisks/dots in other rows (if present) indicate significant differences to uninoculated controls as determined via Dunnett's post-hoc test (. p < 0.1, * p < 0.05, ** p < 0.01, *** p < 0.001). C, carbon, N, nitrogen.

Treatment	%С	%N	δ ¹³ C (‰)	δ ¹⁵ N (‰)	Plant-derived C (µg/g soil)	Soil-derived C (µg/g soil)
			-25.31 ±			38060.63 ±
Absent/control	3.93 ± 0.07	0.36 ± 0.01	0.03	9.72 ± 0.04	1279.03 ± 247.66	712.28
		0.39 ± 0.003	-25.33 ±			$40966.09 \pm$
Acrocalymma sp.	$4.24 \pm 0.03 *$	**	0.02	9.65 ± 0.01	1448.55 ± 188.76	416.19
, 1			-25.33 ±			
Clohesyomyces sp.	3.98 ± 0.02	0.36 ± 0.003	0.03	9.58 ± 0.03 .	1611.13 ± 319.08	38142.72 ± 394.1
			$-25.32 \pm$			$39281.97 \pm$
Darksidea sp. 1	4.07 ± 0.06	0.37 ± 0.004	0.03	9.61 ± 0.06	1364.06 ± 220.06	668.04
		0.38 ± 0.004	-25.35 ±			$40122.22 \pm$
Darksidea sp. 2	4.18 ± 0.06		0.03	9.62 ± 0.03	1635.09 ± 320.66	683.05
		0.38 ± 0.003	$-25.37 \pm$			$40544.37 \pm$
Darksidea sp. 3	$4.23 \pm 0.02 *$	*	0.02	9.69 ± 0.02	1747.74 ± 243.68	332.86
			$-25.34 \pm$			$40246.15 \pm$
Leptodontidium sp.	4.15 ± 0.13	0.38 ± 0.01	0.04	9.72 ± 0.03	1208.67 ± 207.32	1395.36
			$-25.29 \pm$			$40094.79 \pm$
Ophiosphaerella sp.	4.11 ± 0.04	0.38 ± 0.003	0.04	9.82 ± 0.03	1004.45 ± 142.31	501.62
			$-25.39 \pm$			$39356.27 \pm$
Paraconiothyrium sp.	4.12 ± 0.04	0.38 ± 0.004	0.03	9.72 ± 0.03	1830.47 ± 282.22	415.96
			$-25.44 \pm$			
Periconia sp.	4.18 ± 0.09	0.38 ± 0.01	0.04	9.75 ± 0.05	2038.42 ± 288.09	39760.5 ± 820.79
			$-25.36 \pm$			$38769.63 \pm$
Phialocephala sp. 1	4.04 ± 0.05	0.37 ± 0.01	0.05	9.81 ± 0.03	1582.66 ± 368.69	739.07
		0.38 ± 0.01	-25.35 ±			40511.25 ±
Phialocephala sp. 2	4.19 ± 0.10	*	0.02	9.71 ± 0.03	1422.66 ± 130.89	998.06
, ,	4.30 ± 0.04	0.39 ± 0.01	-25.47 ±			40592.71 ±
Thozetella sp.	**	**	0.04 *	9.69 ± 0.03	2434.52 ± 418.15 .	756.54
	≤0.0 <u>5</u> 2	<u><0.05</u> ‡	≤0.0 <u>5</u> 3			***
p-value (ANOVA)	*	*	*	<0.001 ***	0.06.	0.15

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			Decomposition dynamic					
			p-value				Labile	Labile
			(comparison	Resistant	Intermediate	Intermediate	C	C
		Model	with absent	C	C	C MRT	(μg/g	MRT
	Treatment	R ²	/control group)	(µg/g soil)	(μg/g soil)†	(days)	(μg/g soil)†	(days)
Total C			,	30276 ±	(I-B-B)	247 ±		3.07 ±
	Absent/control	0.89	NA	655	8777.69	74	285.57	0.40
				34923 ±		210 ±		$2.70 \pm$
	Acrocalymma sp.	0.89	< 0.001 ***	304 ***	7195.55	67	295.37	0.33
	, .			$31704 \pm$		246 ±		$2.63 \pm$
	Clohesyomyces sp.	0.91	ns	206	7797.19	67	252.13	0.28
								$1.51 \pm$
				$35164 \pm$		164 ±		0.22
	Darksidea sp. 1	0.84	ns	613 ***	5275.69	51	206.06	**
				$36182 \pm$		160 ±		$2.51 \pm$
	Darksidea sp. 2	0.88	< 0.001 ***	556 ***	5322.69	44	252.16	0.37
				$34398 \pm$		222 ±		$3.01 \pm$
	Darksidea sp. 3	0.87	< 0.01 **	195 **	7620.96	65	272.88	0.42
				33941 ±		227 ±		3.04 ±
	Leptodontidium sp.	0.89	ns	1285 **	7216.05	69	297.45	0.37
		0.50		35583 ±	#04# O 4	161 ±	100.10	2.09 ±
	Ophiosphaerella sp.	0.79	ns	380 ***	5317.96	60	198.12	0.45
	D	0.00		32053 ±	006662	291 ± 97	266.24	3.25 ±
	Paraconiothyrium sp.	0.89	ns	379 34970 ±	8866.63	97 196 ±	266.34	0.41 4.17 ±
	Periconia sp.	0.87	n.c	34970± 859 ***	6485.94	196 ± 77	342.66	4.17 ± 0.81
	r ericonia sp.	0.67	ns	31058 ±	0463.94	309 ±	342.00	3.76 ±
	Phialocephala sp. 1	0.79	< 0.001 ***	540	9011.62	193	282.05	0.77
	1 ниносерпин вр. 1	0.79	0.001	33098 ±	9011.02	249 ±	202.03	2.73 ±
	Phialocephala sp. 2	0.88	< 0.01 **	1041.	8563.14	79	271.87	0.35
	1 maiocephaia sp. 2	0.00	< 0.01	36615 ±	0303.14	182 ±	2/1.0/	3.41 ±
	Thozetella sp.	0.86	< 0.001 ***	439 ***	6127.71	54	284.05	0.53
Soil-				31337 ±		258 ±		2.70 ±
derived	Absent/control	0.95	NA	712	6517.67	55	205.43	0.22
C		0.0	. 0. 001 ***	35086 ±		234 ±		2.90 ±
	Acrocalymma sp.	0.9	< 0.001 ***	416 *	5660.13	77	219.30	0.34
		0.94	na	$32351 \pm$		252 ±		$2.99 \pm$
	Clohesyomyces sp.	0.94	ns	394	5586.36	60	205.31	0.25
		0.85	ns	$34436 \pm$		206 ±		$2.78 \pm$
	Darksidea sp. 1	0.65	115	668.	4669.97	75	175.08	0.43
		0.92	< 0.001 ***	$35757 \pm$		181 ±		$2.86 \pm$
	Darksidea sp. 2	0.92	V.001	683 **	4165.06	45	199.37	0.33

D 1:1 2	0.93	< 0.001 ***	33927 ±	6280.46	277 ± 78	227.75	3.18 ±
Darksidea sp. 3			332 34232 ±	6389.46	78 235 ±	221.15	0.30 3.13 ±
Leptodontidium sp.	0.92	< 0.001 ***	1395	5791.95	58	221.83	0.32
	0.87	ns	$35804 \pm$		169 ±		$3.10 \pm$
Ophiosphaerella sp.	0.07	***	501 **	4113.89	52	175.91	0.56
	0.95	ns	$32887 \pm$		281 ±		$2.64 \pm$
Paraconiothyrium sp.			415	6258.33	64	209.99	0.19
	0.96	ns	$34874 \pm$		187 ±		$3.58 \pm$
Periconia sp.			820 *	4644.09	37	242.11	0.34 *
	0.01	0.001 ****	$32988 \pm$		241 ±		$3.14 \pm$
Phialocephala sp. 1	0.91	< 0.001 ***	739	5584.94	74	196.62	0.38
	0.93	< 0.001 ***	33891 ±		270 ±		$2.94 \pm$
Phialocephala sp. 2	0.93	< 0.001	998	6399.73	72	220.25	0.27
	0.94	< 0.001 ***	$35864 \pm$		184 ±		$3.05 \pm$
Thozetella sp.	0.94	< 0.001 ***	756 **	4509.96	37	217.77	0.29

Table B4. Properties of <u>earbon-C</u> fractions of soil in which inoculated wheat plants were grown for four months. Properties were measured using soil fractionation analysis. P-values from ANOVA are displayed in the bottom row. Asterisks/dots in other rows (if present) indicate significant differences to uninoculated controls as determined via Dunnett's post-hoc test (. p < 0.1, * p < 0.05, ** p < 0.01, *** p < 0.001). C, carbon, N, nitrogen, AggC, aggregate carbon, MAOM, mineral-associated organic matter, POM, particulate organic matter.

Treatment	AggC fraction –	AggC fraction – %N	MAOM fraction – %C	MAOM fraction – %N	POM fraction – %C	POM fraction – %N
Absent/control	1.96 ± 0.05	0.16 ± 0.01	0.57 ± 0.02	0.05 ± 0.002	0.92 ± 0.07	0.06 ± 0.01
Acrocalymma sp.	2.18 ± 0.10	0.18 ± 0.01	0.48 ± 0.02	0.04 ± 0.001	0.98 ± 0.05	0.07 ± 0.004
Clohesyomyces sp.	2.14 ± 0.07	0.18 ± 0.01	0.51 ± 0.02	0.05 ± 0.002	0.94 ± 0.05	0.06 ± 0.003
Darksidea sp. 1	2.09 ± 0.06	0.17 ± 0.01	0.58 ± 0.04	0.05 ± 0.003	0.87 ± 0.04	0.06 ± 0.003
Darksidea sp. 2	2.13 ± 0.03	0.17 ± 0.002	0.54 ± 0.05	0.05 ± 0.004	0.89 ± 0.03	0.06 ± 0.002
Darksidea sp. 3	2.13 ± 0.05	0.17 ± 0.004	0.60 ± 0.02	0.05 ± 0.002	1.00 ± 0.06	0.07 ± 0.004
Leptodontidium sp.	2.12 ± 0.07	0.17 ± 0.01	0.53 ± 0.02	0.05 ± 0.002	0.98 ± 0.04	0.06 ± 0.003
Ophiosphaerella sp.	2.18 ± 0.04	0.19 ± 0.004 *	0.55 ± 0.03	0.05 ± 0.003	0.96 ± 0.04	0.07 ± 0.003
Paraconiothyrium sp.	2.15 ± 0.05	0.18 ± 0.004	0.56 ± 0.03	0.05 ± 0.002	1.00 ± 0.06	0.07 ± 0.01
Periconia sp.	2.25 ± 0.06 *	0.19 ± 0.01 *	0.55 ± 0.05	0.05 ± 0.004	0.89 ± 0.03	0.06 ± 0.002
Phialocephala sp. 1	2.22 ± 0.06	$0.19 \pm 0.01 **$	0.53 ± 0.02	0.05 ± 0.002	0.86 ± 0.09	0.06 ± 0.01
Phialocephala sp. 2	2.09 ± 0.07	0.17 ± 0.01	0.56 ± 0.03	0.05 ± 0.003	0.86 ± 0.03	0.06 ± 0.002
Thozetella sp.	2.37 ± 0.07 ***	0.20 ± 0.01 ***	0.52 ± 0.04	0.05 ± 0.003	0.91 ± 0.10	0.06 ± 0.01
p-value (ANOVA)	0.03 *<0.05 *	0.002 **<<0.01 **	0.63	0.62	0.65	0.41

Table B5. Plant variables potentially influencing soil (in which inoculated wheat plants were grown for four months). P-values from ANOVA are displayed in bottom rows. Asterisks/dots in other rows (if present) indicate significant differences to uninoculated controls as determined via Dunnett's post-hoc test (. p < 0.1, * p < 0.05, ** p < 0.01, *** p < 0.001). C, carbon, N, nitrogen.

	Number	Average	Total		_		-		
	of	spike	spike	Shoot	Root	Root/shoot	Shoot	Shoot	Shoot
Treatment	spikes	mass (g)	mass (g)	mass (g)	mass (g)	ratio	δ ¹³ C (‰)	δ ¹⁵ N (‰)	%C
	$5.50 \pm$	$1.52 \pm$	$7.36 \pm$	$16.38 \pm$	$2.23 \pm$	$0.14 \pm$	$-32.27 \pm$	$9.74 \pm$	$38.30 \pm$
Absent/control	0.91	0.28	1.06	1.97	0.20	0.01	0.92	0.24	0.42
	$4.86 \pm$	$1.82 \pm$	$8.81 \pm$	$16.81 \pm$	$1.83 \pm$	$0.11 \pm$	$-32.47 \pm$	$9.39 \pm$	$37.81 \pm$
Acrocalymma sp.	0.43	0.07	0.81	1.77	0.33	0.01	0.91	0.15	0.40
	$4.14 \pm$	$1.85 \pm$	$6.60 \pm$	$13.28 \pm$	$1.44 \pm$	$0.11 \pm$	$-31.94 \pm$	$9.38 \pm$	$38.21 \pm$
Clohesyomyces sp.	0.65	0.25	0.77	1.26	0.22	0.01	1.02	0.18	0.49
	$3.86 \pm$	$2.13 \pm$	$8.11 \pm$	$15.54 \pm$	$1.75 \pm$	$0.11 \pm$	$-32.27 \pm$	$9.44 \pm$	$38.07 \pm$
Darksidea sp. 1	0.24	0.10	0.38	0.95	0.17	0.01	1.03	0.18	0.28
	$4.43 \pm$	$2.20 \pm$	$9.41 \pm$	$16.88 \pm$	$2.00 \pm$	$0.12 \pm$	$-32.19 \pm$	$9.64 \pm$	$38.08 \pm$
Darksidea sp. 2	0.45	0.14 <u>f</u>	0.68	1.55	0.25	0.01	0.84	0.34	0.49
•	$4.14 \pm$	$1.63 \pm$	$6.37 \pm$	$15.46 \pm$	$1.86 \pm$	$0.14 \pm$	$-32.73 \pm$	$9.89 \pm$	$37.72 \pm$
Darksidea sp. 3	0.84	0.20	1.17	1.62	0.34	0.02	1.13	0.13	0.52
1	5.57 ±	$1.72 \pm$	$8.15 \pm$	$16.42 \pm$	$2.02 \pm$	$0.12 \pm$	-33.53 ±	$9.21 \pm$	$37.73 \pm$
Leptodontidium sp.	0.90	0.25	0.66	0.80	0.44	0.03	0.76	0.48	0.59
	4.43 ±	$1.92 \pm$	$8.32 \pm$	$15.68 \pm$	$1.63 \pm$	$0.10 \pm$	$-32.76 \pm$	$9.37 \pm$	$37.57 \pm$
Ophiosphaerella sp.	0.28	0.11	0.26	1.17	0.40	0.02	1.08	0.24	0.32
1 1 1	$3.86 \pm$	$2.12 \pm$	$7.43 \pm$	$14.01 \pm$	$1.73 \pm$	$0.12 \pm$	$-32.32 \pm$	$9.66 \pm$	$37.21 \pm$
Paraconiothyrium sp.	0.51	0.23	0.40	1.03	0.35	0.02	0.95	0.38	0.36
, 1	$3.86 \pm$	1.93 ±	$7.36 \pm$	$15.96 \pm$	$1.83 \pm$	$0.12 \pm$	$-32.42 \pm$	$10.23 \pm$	$38.17 \pm$
Periconia sp.	0.51	0.20	1.07	1.48	0.23	0.02	0.86	0.26	0.32
1	4.43 ±	$1.98 \pm$	$7.85 \pm$	$15.82 \pm$	1.93 ±	$0.12 \pm$	-32.42 ±	$9.15 \pm$	$38.43 \pm$
Phialocephala sp. 1	0.60	0.25	0.60	1.34	0.36	0.02	0.96	0.16	0.35
r	$4.00 \pm$	$2.26 \pm$	8.56 ±	15.95 ±	$2.19 \pm$	0.14 ±	$-32.68 \pm$	$9.80 \pm$	$37.64 \pm$
Phialocephala sp. 2	0.54	0.20	0.85	1.90	0.28	0.01	0.86	0.19	0.33
r	4.14 ±	$2.48 \pm$	9.82 ±	18.57 ±	2.55 ±	$0.14 \pm$	-32.58 ±	9.31 ±	$37.66 \pm$
Thozetella sp.	0.51	0.15 *	0.66	1.55	0.36	0.02	1.07	0.23	0.41
p-value (ANOVA)	0.66	0.12	0.14	0.75	0.74	0.82	1.00	0.32	0.84
				Root					
			Root	specific	Root	Root	Root		
		Shoot	length	surface	average	length per	specific		
	Shoot	C/N	density	area	diameter	volume	density		
	%N	ratio	(cm/g)	(cm^2/g)	(mm)	(cm/m ³)	(g/cm ³)	Root fork 1	number (/g)
Treatment	P.SN	P.SCN	P.RLDe	P.RSA	P.RADi	P.RLV	P.RSDe	P.RF	
	0.49 ±	83.32 ±	3315.39 ±	490.13	0.48 ±	515.85 ±	0.17 ±	5878.38 ±	
Absent/control	0.05	8.44	307.45	± 30.83	0.02	65.77	0.01	870.62	

	0.43 ±	90.51 ±	3563.82 ±	530.07	$0.48 \pm$	492.79 ±	$0.16 \pm$	6456.09 ±
Acrocalymma sp.	0.03	7.10	247.20	± 31.47	0.01	95.89	0.01	1283.54
	$0.45 \pm$	$91.07 \pm$	$4044.30 \pm$	561.07	$0.46 \pm$	$499.66 \pm$	$0.17 \pm$	$7056.00 \pm$
Clohesyomyces sp.	0.04	7.69	627.70	± 63.37	0.03	102.50	0.01	1385.96
, , 1	$0.44 \pm$	$90.30 \pm$	$3544.01 \pm$	539.47	$0.49 \pm$	$586.57 \pm$	$0.16 \pm$	6748.77 ±
Darksidea sp. 1	0.04	6.73	390.12	± 52.13	0.02	61.95	0.01	1228.20
•	$0.40 \pm$	$97.22 \pm$	$3872.21 \pm$	557.82	$0.48 \pm$	$620.39 \pm$	$0.16 \pm$	$8050.86 \pm$
Darksidea sp. 2	0.02	6.10	461.38	± 39.54	0.02	123.60	0.01	1549.33
•	$0.58 \pm$	$82.65 \pm$	$3912.67 \pm$	562.39	$0.47 \pm$	$570.09 \pm$	$0.15 \pm$	7540.25 ±
Darksidea sp. 3	0.12	12.54	356.62	± 27.00	0.02	136.56	0.01	1301.61
_	$0.46 \pm$	$85.82 \pm$	$3779.06 \pm$	540.19	$0.47 \pm$	$615.66 \pm$	$0.16 \pm$	6972.52 ±
Leptodontidium sp.	0.04	6.59	475.55	± 41.41	0.03	145.93	0.01	1670.66
	$0.43 \pm$	$89.68 \pm$	$4718.73 \pm$	632.58	$0.45 \pm$	$698.43 \pm$	$0.15 \pm$	$9458.82 \pm$
Ophiosphaerella sp.	0.02	5.32	906.96	± 83.92	0.02	146.81	0.01	2376.20
	$0.44 \pm$	$93.43 \pm$	$3721.05 \pm$	541.97	$0.47 \pm$	$440.31 \pm$	$0.16 \pm$	$6278.34 \pm$
Paraconiothyrium sp.	0.05	10.56	352.69	± 40.66	0.02	85.04	0.01	1226.28
	$0.59 \pm$	$75.07 \pm$	$3629.11 \pm$	520.13	$0.47 \pm$	$465.06 \pm$	$0.17 \pm$	6273.79 ±
Periconia sp.	0.11	8.24	390.34	± 38.44	0.02	89.46	0.01	1414.99
	$0.41 \pm$	$96.97 \pm$	$3170.61 \pm$	469.51	$0.47 \pm$	$382.08 \pm$	$0.19 \pm$	$4430.48 \pm$
Phialocephala sp. 1	0.03	7.95	220.70	± 30.03	0.01	67.80	0.01	488.78
	$0.45 \pm$	$91.12 \pm$	$4648.09 \pm$	631.31	$0.45 \pm$	$748.74 \pm$	$0.15 \pm$	9350.21 ±
Phialocephala sp. 2	0.05	9.15	804.77	± 76.97	0.02	106.18	0.01	1855.27
	$0.39 \pm$	$99.44 \pm$	$3651.81 \pm$	521.36	$0.47 \pm$	$697.98 \pm$	$0.17 \pm$	$6835.67 \pm$
Thozetella sp.	0.03	7.41	353.05	± 30.21	0.02	92.43	0.01	1146.69
p-value (ANOVA)	0.47	0.86	0.75	0.68	0.10	0.98	0.55	0.69

Table B6. Microbial community variables potentially influencing soil (in which inoculated wheat plants were grown for four months). P-values from ANOVA are displayed in the bottom row. Asterisks/dots in other rows (if present) indicate significant differences to uninoculated controls as determined via Dunnett's post-hoc test (. p < 0.1, * p < 0.05, * * p < 0.01, * * p < 0.001).

Treatment	Total community size (µg PLFA /g soil)	Fungal to bacterial biomass ratio	Gram positive bacteria (% of total community)	Gram negative bacteria (% of total community)	Actinobacteria (% of total community)	Fungi (% of total community)	Arbuscular mycorrhizal fungi (% of total community)
Absent/control	8.30 ± 0.33	0.22 ± 0.01	19.50 ± 0.01	26.19 ± 0.55	8.20 ± 0.14	10.19 ± 0.47	2.41 ± 0.09
Acrocalymma sp.	8.59 ± 0.57	0.23 ± 0.01	19.88 ± 0.01	26.10 ± 0.72	7.68 ± 0.74	10.44 ± 0.42	2.45 ± 0.07
Clohesyomyces sp.	8.35 ± 0.28	0.22 ± 0.01	20.38 ± 0.01	26.48 ± 0.48	8.48 ± 0.14	10.11 ± 0.28	2.52 ± 0.07
Darksidea sp. 1	8.54 ± 0.30	0.22 ± 0.01	20.14 ± 0.01	26.06 ± 0.61	8.37 ± 0.11	9.98 ± 0.26	2.63 ± 0.10
Darksidea sp. 2	7.72 ± 0.32	0.21 ± 0.01	20.10 ± 0.01	26.59 ± 0.47	8.23 ± 0.16	9.79 ± 0.32	2.71 ± 0.12
Darksidea sp. 3	7.50 ± 0.71	0.22 ± 0.01	19.03 ± 0.01	25.32 ± 0.40	7.90 ± 0.08	9.54 ± 0.34	2.41 ± 0.08
Leptodontidium sp.	7.89 ± 0.51	0.23 ± 0.01	20.01 ± 0.01	26.02 ± 0.57	8.16 ± 0.20	10.36 ± 0.41	2.62 ± 0.07
Ophiosphaerella sp.	8.61 ± 0.21	0.24 ± 0.01	19.28 ± 0.01	26.27 ± 0.33	8.21 ± 0.17	10.97 ± 0.47	2.72 ± 0.08
Paraconiothyrium sp.	7.98 ± 0.27	0.21 ± 0.01	20.65 ± 0.01	26.64 ± 0.43	8.69 ± 0.15	9.88 ± 0.29	2.65 ± 0.05
Periconia sp.	8.50 ± 0.34	0.21 ± 0.01	20.37 ± 0.01	27.02 ± 0.34	8.25 ± 0.09	9.83 ± 0.34	2.61 ± 0.09
Phialocephala sp. 1	8.69 ± 0.29	0.21 ± 0.01	20.52 ± 0.01	26.34 ± 0.42	8.30 ± 0.09	9.79 ± 0.27	2.75 ± 0.09 .
Phialocephala sp. 2	8.75 ± 0.20	0.23 ± 0.01	19.30 ± 0.01	25.89 ± 0.27	8.25 ± 0.19	10.16 ± 0.43	2.62 ± 0.09
Thozetella sp.	8.27 ± 0.37	0.22 ± 0.01	19.39 ± 0.01	26.23 ± 0.50	8.23 ± 0.11	9.80 ± 0.24	2.53 ± 0.09
p-value (ANOVA)	0.72	0.50	0.45	0.81	0.61	0.50	0.13

Table B7. Fungal variables potentially influencing soil (in which inoculated wheat plants were grown for four months). P-values from ANOVA are displayed in the bottom row (. p < 0.1, * p < 0.05, ** p < 0.01, *** p < 0.001). Different letters indicate significant differences between treatments as determined via Tukey's post-hoc test. † indicates variables calculated using treatment averages. C, carbon, N, nitrogen.

	Estimated final colony area	Maximum growth rate	Time to maximum growth	Biomass	Final colony area	Hyphal density			
Treatment	(cm ²)†	(cm ² /day)†	(days)†	(g)†	(cm ²)†	(mg/cm ²)†	%C†	%N†	C/N ratio†
	53.58 ±	4.61 ±	12.02 ±	0.12 ±	49.17 ±	2.42 ±	51.96 ±	2.67 ±	19.53 ±
Acrocalymma sp.	1.26 c	0.03 de	0.26 bcd	0.01 ab	0.55 abc	0.23 b	0.37 ab	0.06 cd	0.36 bc
,I	38.64 ±	$2.05 \pm$	17.42 ±	$0.04 \pm$	$29.76 \pm$	1.18 ±	49.11 ±	3.81 ±	12.93 ±
Clohesyomyces sp.	1.72 d	0.08 g	0.28 a	0.01 e	1.78 d	0.23 b	0.49 cd	0.09 a	0.41 f
, , ,	59.49 ±	3.39 ±	$18.04 \pm$	$0.08 \pm$	$47.43 \pm$	$1.61 \pm$	45.99 ±	$2.32 \pm$	19.91 ±
Darksidea sp. 1	1.94 bc	0.09 f	0.36 a	0.003 cd	1.14 bc	0.09 b	0.23 e	0.07 de	0.57 bc
•	69.82 ±	$4.89 \pm$	$16.87 \pm$	$0.09 \pm$	$53.58 \pm$	$1.70 \pm$	$46.96 \pm$	$2.55 \pm$	$18.53 \pm$
Darksidea sp. 2	0.84 ab	0.09 cd	0.09 a	0.01 bcd	0.96 ab	0.12 b	0.18 e	0.10 d	0.77 cd
•	58.39 ±	$5.12 \pm$	$12.93 \pm$	$0.07 \pm$	$52.52 \pm$	$1.35 \pm$	$52.81 \pm$	$2.66 \pm$	19.91 ±
Darksidea sp. 3	1.04 bc	0.06 cd	0.10 bc	0.004 cde	0.63 ab	0.08 b	0.30 a	0.04 cd	0.35 bc
•	53.01 ±	$4.00 \pm$	$16.20 \pm$	$0.08 \pm$	$43.02 \pm$	$1.80 \pm$	$52.68 \pm$	$2.06 \pm$	$25.54 \pm$
Leptodontidium sp.	2.42 c	0.21 ef	0.20 a	0.01 cde	2.40 c	0.23 b	0.32 a	0.03 e	0.28 a
	$70.45 \pm$	$6.37 \pm$	$13.63 \pm$	$0.13 \pm$	$54.45 \pm$	$2.44 \pm$	$50.42 \pm$	$2.09 \pm$	$24.16 \pm$
Ophiosphaerella sp.	1.50 ab	0.02 b	0.22 b	0.01 a	0.24 a	0.24 b	0.52 bc	0.03 e	0.03 a
	74.83 ±	$7.54 \pm$	$10.19 \pm$	$0.09 \pm$	$50.25 \pm$	$1.86 \pm$	$47.43 \pm$	$3.02 \pm$	$15.83 \pm$
Paraconiothyrium sp.	3.68 a	0.11 a	0.27 de	0.01 abcd	0.67 ab	0.15 b	0.46 de	0.15 bc	0.66 e
	66.92 ±	$7.28 \pm$	$9.81 \pm$	$0.09 \pm$	$48.01 \pm$	$1.82 \pm$	$52.54 \pm$	$3.24 \pm$	$16.24 \pm$
Periconia sp.	2.66 ab	0.04 a	0.32 e	0.004 bcd	0.41 abc	0.09 b	0.17 a	0.07 b	0.17 de
	$60.76 \pm$	$5.35 \pm$	$13.51 \pm$	$0.10 \pm$	$53.34 \pm$	$1.87 \pm$	$46.51 \pm$	$2.38 \pm$	$19.58 \pm$
Phialocephala sp. 1	2.03 bc	0.17 c	0.15 bc	0.003 abcd	1.43 ab	0.08 b	0.19 e	0.02 de	0.26 bc
	58.61 ±	$5.12 \pm$	$12.32 \pm$	$0.12 \pm$	$53.46 \pm$	$2.15 \pm$	$45.87 \pm$	$2.30 \pm$	$19.98 \pm$
Phialocephala sp. 2	1.74 abc	0.06 cd	0.16 bcde	0.01 abc	1.10 ab	0.13 b	0.44 e	0.02 de	0.14 bc
	28.02 ±	$2.16 \pm$	$11.33 \pm$	$0.06 \pm$	$13.95 \pm$	$4.59 \pm$	$50.97 \pm$	$2.42 \pm$	$21.10 \pm$
Thozetella sp.	4.16 d	0.19 g	1.05 cde	0.01 de	1.17 e	0.54 a	0.35 abc	0.02 de	0.35 b
	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001
p-value (ANOVA)	***	***	***	***	***	***	***	***	***

Code availability Scripts for data and statistical analyses will be made available according to the journal's policies when the manuscript is accepted for publication. **Author contribution** YC, JP and LCG designed the study; ES, LCG and WB performed the research; ES wrote the first draft of the manuscript, and all authors contributed to revisions. Competing interests The research was partially funded by SoilCQuest2031 who provided the fungal cultures and soil. This funding was provided independently of research findings. SoilCQuest2031 did not attempt to influence the interpretations or conclusions of the work. The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest. Acknowledgements This project was supported by Western Sydney University Research Partnership Program and by SoilCQuest2031 (Orange, New South Wales, Australia). We acknowledge assistance from Guy Webb and Suresh Subashchandrabose for providing soils and cultures. We also thank Andrew Gherlenda for assistance with the growth chamber experiment, Russell Thomson for help with nonlinear least square curve fitting, UC Davis Stable Isotope Facility, Environmental Analysis Laboratory, and Pushpinder Matta for running the nutrient analyses, and Sophia Bruna, Hui Zhang, and Asel Weerasekara for help with the experiment harvest.

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