Microbial communities in top- and subsoil of repacked soil columns respond differently to amendments but their diversity is negatively correlated with plant productivity

Corinne Celestina1,2, Jennifer L. Wood1,2, James B. Manson3, Xiaojuan Wang3, Peter W. G. Sale3, Caixian Tang3 & Ashley E. Franks1,2

Organic and inorganic amendments with equivalent nutrient content may have comparable fertilizer effects on crop yield, but their effects on the soil microbial community and subsequent plant-soil-microbe interactions in this context are unknown. This experiment aimed to understand the relationship between soil microbial communities, soil physicochemical characteristics and crop performance after addition of amendments to soil. Poultry litter and synthetic fertilizer with balanced total nitrogen (N) content equivalent to 1,200 kg ha\(^{-1}\) were added to the topsoil (0–10 cm) or subsoil layer (20–30 cm) of repacked soil columns. Wheat plants were grown until maturity. Soil samples were taken at Zadoks 87–91 (76 days after sowing) for analysis of bacterial and fungal communities using 16S and ITS amplicon sequencing. The interaction between amendment type and placement depth had significant effects on bacterial and fungal community structure and diversity in the two soil layers. Addition of poultry litter and fertilizer stimulated or suppressed different taxa in the topsoil and subsoil leading to divergence of these layers from the untreated control. Both amendments reduced microbial community richness, diversity and evenness in the topsoil and subsoil compared to the nil-amendment control, with these reductions in diversity being consistently negatively correlated with plant biomass (root and shoot weight, root length, grain weight) and soil fertility (soil NH\(_4^+\), shoot N). These results indicate that in this experimental system, the soil microbial diversity was correlated negatively with plant productivity.

Surface1 and subsoil2–4 placement of nutrient-rich organic amendments has been used as a technique to improve plant productivity on agricultural soils characterized by low organic matter, poor physical structure or low chemical fertility. Crop yield responses to the amendments have been attributed to a combination of improved soil structure, water use, nutrient supply and biological activity2–6. Because of the interplay between the plant, soil and soil biota7, it is expected that the microbial community plays a key role in nutrient transformations and soil aggregation and hence the crop response to organic amendments. Although many authors have reported changes to soil properties and microbial communities8–11 after addition of organic amendments to soil, our understanding of the subsequent plant-soil-microbe interactions and their effect on plant productivity is unclear.

Additionally, the role of nutrients in the crop response to organic amendments may be more significant than previously thought, with a number of recent meta-analyses concluding that organic amendments do not have substantial additional effects on crop yields beyond their fertilizer effects12–15. That is, organic and inorganic amendments with matched macronutrients can have the same effect on crop yields, regardless of whether they are applied to the soil surface or the subsoil16. This suggests that the key contribution of the microbial community to the crop response to amendments may lie in its role in nutrient cycling and decomposition. And, whilst plant
In the present study we aimed to understand the relationship between soil microbial communities, soil physical-chemical characteristics and crop performance after addition of organic and inorganic amendments to repacked soil columns. Poultry litter (‘PL’) and synthetic fertilizer (‘FERT’) with balanced total nitrogen (N) content were added to topsoil (‘TOP’) or subsoil (‘SUB’) layers of soil columns with wheat plants grown until physiological maturity. We hypothesized that the structure and function of the microbial community would differ between different amendments and different soil layers, with the addition of amendments leading to convergence of the bacterial and fungal communities of the topsoil and subsoil layers. Furthermore, we theorized that changes in microbial community structure and diversity compared to the nil-amendment control (‘NIL’) would be positively associated with soil fertility, aggregation and plant growth.

Results

A total of 1,953,805 and 1,079,219 high quality 16S and ITS sequences were obtained from the 18 samples, with sequences per sample ranging from 34,714 to 193,000 for bacteria and 26,744 to 117,578 for fungi. These sequences were clustered into 6,689 bacterial and 570 fungal OTUs at the 97% similarity level with an average of 1,810 bacterial and 150 fungal OTUs per sample. After trimming to remove OTUs with mean relative abundance less than 0.005% there were 538 bacterial and 110 fungal OTUs remaining. The majority of OTUs were rare members of the community, with only 2% of bacterial and 10% of fungal OTUs having mean relative abundance greater than 1%.

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Comparison of microbial community structure. The estimated number of OTUs in each sample (Chao1 richness), Shannon diversity and Simpson’s evenness are shown in Table 1. The type of amendment and placement depth had consistent, significant ($P < 0.05$) effects on the bacterial and fungal communities, but the amendment × depth interactive effect was not as strong. Additionally, placement depth tended to have a stronger effect size than amendment type or amendment × depth interaction. In general, addition of fertilizer and, to a lesser extent, poultry litter to the soil reduced the richness, diversity and evenness of bacterial and fungal communities in both soil layers. As a main effect, averaged across placement depth, the nil-amendment control tended to have the lowest values for all indices. These reductions in Chao1 richness and Shannon diversity are due to reductions in the number of OTUs and the distribution of individuals among OTUs, whereas reductions in evenness correspond to a less evenly distributed community dominated by a few OTUs. In terms of placement depth, microbial communities in the subsoil consistently had lower richness, diversity and evenness compared to those in the topsoil. The exceptions to the trend were Simpson’s evenness in fungal communities: evenness was not affected by amendment type but did differ between topsoil and subsoil layers, with the subsoil significantly more even ($P = 0.022$) because of more equal abundances of all OTUs.

A similar phylum-level composition was observed across all six treatments (Fig. 1). The bacterial community in the topsoil and subsoil was dominated by Proteobacteria (~45%), Acidobacteria (~19%) and Actinobacteria (~16%), whilst Ascomycota (~85%) was the dominant fungal phylum in both soil layers. Fungal community
composition was variable between placement depth and amendment type, whereas the bacterial community was relatively stable in the topsoil but more variable in the subsoil across the three amendment treatments. In the nil-amendment control, the topsoil and the subsoil had similar phylum-level abundances, but differences could be seen in community composition between the two soil layers in the fertilizer and poultry litter treatments.

Principal coordinates analysis of weighted UniFrac distances was used to visualize the effects of amendment type and placement depth on soil microbial communities (Fig. 2). The first two principal coordinates explained a high percentage of the variance for the bacterial (70%) and fungal communities (68%), with variation in microbial community structure associated with both placement depth and amendment type. Within each depth, communities in the nil-amendment control, fertilizer and poultry litter treatments separated. There was also clear separation between the microbial communities of the topsoil and subsoil, although one replicate of the nil-amendment subsoil treatment clustered with the nil-amendment topsoil. Overall, the topsoil communities clustered more closely together and were more similar in terms of species abundance and phylogenetic distance than the subsoil communities.

Permutational multivariate analysis of variance (PERMANOVA) of weighted UniFrac distances confirmed a significant (P < 0.05) global effect of amendment type, placement depth and amendment type × placement depth on microbial communities (Table 2). As a main effect, placement depth (Pseudo-F = 13.79–14.31) had a stronger effect than amendment type (Pseudo-F = 2.66–6.20) and effect sizes were generally higher in bacteria than in
Table 2. Results of permutational analysis of variance (PERMANOVA) of weighted UniFrac distances testing the effect of amendment type and depth on soil microbial communities. P(MC) = P-value based on Monte Carlo random draws.

| Amendment type | Global tests | | | Bacteria | Fungi |
|----------------|--------------|--------|--------|----------|--------|
|                | Pseudo-F     | P-value| Pseudo-F| P-value  |
| Amendment type | 6.20         | 0.001  | 2.66   | 0.003    |
| Placement depth| 14.31        | 0.002  | 13.79  | 0.001    |
| Amendment type × Placement depth | 3.65        | 0.002  | 2.46   | 0.006    |

Pairwise tests

| Amendment type | Topsoil | | | Subsoil | | | | Placement depth within Amendment type |
|----------------|---------|--------|--------|---------|--------|--------|--------|---------|
|                | t-statistic | P-value| t-statistic | P-value| t-statistic | P-value| |
| Nil vs. Fertilizer | 3.42 | 0.002 | 2.33 | 0.006 |
| Nil vs. Poultry Litter | 1.47 | 0.048 | 1.14 | 0.278 |
| Poultry Litter vs. Fertilizer | 2.61 | 0.003 | 1.69 | 0.013 |
| Amendment type within Placement depth | | | | Nil | | | |
| Topsoil | | | | Subsoil | | | |
| Nil vs. Fertilizer | 2.76 | 0.012 | 1.78 | 0.078 |
| Nil vs. Poultry Litter | 1.33 | 0.199 | 1.12 | 0.323 |
| Poultry Litter vs. Fertilizer | 4.51 | 0.002 | 1.30 | 0.253 |
| Subsoil | | | | Placement depth within Amendment type | | | |
| Nil | | | | Topsoil vs. Subsoil | 1.49 | 0.173 | 1.69 | 0.141 |
| Poultry litter | | | | Topsoil vs. Subsoil | 2.25 | 0.032 | 2.00 | 0.036 |
| Fertilizer | | | | Topsoil vs. Subsoil | 4.50 | 0.001 | 8.48 | 0.004 |

There were significant differences in microbial communities between nil-amendment and fertilizer treatments, and poultry litter and fertilizer treatments. Bacterial communities also differed between nil-amendment and poultry litter treatments.

Pairwise PERMANOVA revealed a significant interactive effect of amendment type and placement depth. Comparing the two placement depths of each amendment, poultry litter and fertilizer both showed differences (P(MC) < 0.05) in bacterial and fungal community structures between the topsoil and the subsoil. However, the difference between microbial communities in the nil-amendment topsoil and subsoil was not significant (P(MC) > 0.05) due to one replicate from the subsoil layer that clustered with the topsoil samples (Fig. 2). Comparing between amendment types within each placement depth, the bacterial community in the synthetic fertilizer treatment was significantly different to both the nil-amendment and poultry litter treatments in the topsoil and subsoil. This trend was similar for fungal communities in the subsoil, with a significant difference (P(MC) = 0.036) observed between nil and fertilizer treatments and some indication of dissimilarity between poultry litter and fertilizer (pseudo-F = 2.02, P(MC) = 0.071). Amongst fungal communities in the topsoil, there was no significant difference between amendments. However, this was only the case for PERMANOVA of weighted UniFrac distances (based on phylogenetic structure, weighted by OTU abundance). The PERMANOVA of unweighted UniFrac distances (takes into account species presence only), suggests that although similar OTUs were present, the abundance of these OTUs was likely having a strong influence on community structure in this layer (Poultry litter vs. Fertilizer, P(MC) = 0.04; Nil vs. Fertilizer, P(MC) = 0.04) (Supplementary Table S2).

Comparison of microbial communities at OTU level. Univariate tests for each OTU identified 99 bacterial and 6 fungal OTUs (16 and 5% of the total community, respectively) that were observed to differ significantly (P_adj < 0.05) between amendment type and placement depth treatments (Supplementary Table S3). The majority of bacterial OTUs (84 of 99) were responding to the placement depth, whereas only 12 out of 99 OTUs differed in abundance between the three amendments and the remainder responded to the amendment × depth interaction. Most of the bacterial OTUs that responded to amendment type belonged to the phyla Proteobacteria and Candidatus Saccharibacteria (Candidate Division TM7). Those responding to placement depth were mainly from the highly abundant phyla Proteobacteria, Acidobacteria and Actinobacteria (subset shown in Fig. 3a). Among the bacterial OTUs that differed between the amendment types, 8 out of 12 OTUs had a higher abundance in the fertilizer-amended soil than in either the nil or poultry litter treatments, and 7 OTUs were lowest in abundance in the nil treatment. Of the bacterial OTUs responding to placement depth, 55 of 84 were more abundant in the topsoil compared to the subsoil (Fig. 3a, Supplementary Table S3). Of the OTUs that were more...
abundant in the topsoil, the order of abundance (from highest to lowest) was most commonly TOP FERT > TOP PL > TOP NIL > SUB NIL > SUB FERT > SUB PL. Of the OTUs that were more abundant in the subsoil, the order of abundance was most commonly SUB PL > SUB NIL > SUB FERT > TOP NIL > TOP PL > TOP FERT (Supplementary Table S1, S4). Of the 6 fungal OTUs that were observed to differ significantly in abundance, 4 responded to the amendment and 2 to the placement depth (Fig. 3b). Half were from the phylum Ascomycota and another two were classified as Glomeromycota. There were no consistent trends in fungal OTUs responding to amendment type, but the two OTUs that responded to depth were both higher in abundance in the subsoil than the topsoil (Fig. 3b).

**Linking microbial community structure to soil and plant variables.** Canonical correspondence analysis (CCA) was used to relate measured soil and plant variables to bacterial and fungal community structure. After removing highly correlated variables (soil NH$_4^+$, large macroaggregates, small macroaggregates, microaggregates, silt and clay fraction and total root weight), stepwise ordination significance testing on the remaining
measures (soil NO$_3^-$, gravimetric moisture, aggregate mean weight diameter, total shoot weight, root length, root surface area, root diameter, shoot N and total grain weight) identified six environmental variables that were significantly correlated with bacterial and/or fungal community structure (Fig. 4). Soil moisture content ($P = 0.005$), mean-weight diameter of aggregates ($P = 0.005$), root surface area ($P = 0.035$) and root diameter ($P = 0.020$) had a strong influence on bacterial community structure. Soil moisture ($P = 0.005$) and aggregate mean-weight diameter ($P = 0.025$) were also strong drivers of fungal community structure. The constrained ordinations distinctly separated into topsoil and subsoil groups. All six significantly-correlated variables were higher in value in the topsoil than subsoil (Supplementary Table S4) and this corresponds with the direction of the vectors on the ordinations (Fig. 4). The CCA-based variation partitioning analysis indicated that the retained variables (moisture, mean-weight diameter of aggregates, root diameter and root surface area) explained 16, 10, 4 and 7% of the observed variance in bacterial communities, respectively. The variables significantly correlated with fungal community structure (moisture, mean-weight diameter of aggregates) explained 8 and 6% of the community structure. The majority of variation in bacterial and fungal communities was unexplained.

Alpha diversity indices (Chao1 richness estimator, Shannon diversity and Simpson's evenness) for bacteria and fungi were frequently negatively correlated ($P < 0.05$) with environmental variables (Table 3). Reduced OTU richness, diversity and evenness were consistently associated with increasing concentrations of inorganic N in the soil and of total N in shoots, as well as increasing above- and below-ground plant biomass. There were significant ($P < 0.05$), strong negative correlations with soil ammonium, shoot N, root weight, shoot length and grain weight in both the topsoil and subsoil. Conversely, soil moisture was positively correlated ($P < 0.05$) with bacterial and fungal Shannon diversity in both soil layers. In the subsoil only, Shannon diversity and bacterial Simpson's evenness were also found to increase with increasing proportion of silt and clay fractions. In terms of the other measures of soil structure, diversity indices tended to positively correlate with the proportion of soil aggregates and aggregate mean-weight diameter in the topsoil, but negatively correlate with those in the subsoil. Bacterial Chao1 richness and fungal community evenness was weakly correlated with most environmental variables in the subsoil.

**Discussion**

There is evidence that organic and inorganic amendments with equivalent total nutrient content have comparable fertilizer effects on crop yield\cite{12-15}. However, the effects of these amendments on the soil microbial community, and subsequent plant-soil-microbe interactions, are unknown. This experiment aimed to understand the relationship between soil microbial communities, soil physicochemical characteristics and crop performance after addition of amendments with equivalent total N content to topsoil and subsoil.

Amendment type and placement depth affected the diversity of bacterial and fungal communities in the soil, with the depth of placement tending to have a stronger effect than the amendment type. Topsoil communities had a larger number of OTUs and were richer and more evenly distributed than subsoil communities, regardless of whether there was an amendment added to the soil or not. Nevertheless, both organic and inorganic amendments tended to reduce bacterial and fungal community richness, diversity and evenness in both soil layers compared to the nil control because they favored the growth of a few OTUs that dominated the community of the amended soil. The slow-release fertilizer had a larger effect than the poultry litter, reflecting the effect of the nutrient-rich amendments on measured soil and plant variables (e.g. soil inorganic N, shoot weight, root length), which tended to respond in the order of fertilizer > poultry litter > nil-amendment control. Perturbations like the addition of nutrients or labile carbon to a soil have frequently been shown to reduce microbial community diversity and are likely to have a strong selective effect on the community\cite{20-22}. The highly available nutrients in the fertilizer...
Amendment type and placement depth also had significant effects on microbial community structure, and again, the effect of placement (topsoil vs. subsoil) was more pronounced than that of amendment type. Of the OTUs that differed in abundance in response to placement depth, the majority were higher in abundance in the topsoil than the subsoil, indicating a larger stimulatory effect in the topsoil layer. This contradicts other studies reporting a more pronounced response to amendment addition in deeper soil layers under field conditions, leading to convergence of topsoil and subsoil layers. Additionally, all of the OTUs that differed in abundance were inherently more abundant in one soil layer than the other, regardless of whether the soil was amended or not. This suggests that we were primarily observing natural variation in the microbial communities that was arising due to the depth of the soil layer, rather than the type of amendment. Nevertheless, there were obvious trends in each layer caused by the amendments that indicates an interaction between placement depth and amendment type. The poultry litter and inorganic fertilizer treatments formed distinct microbial communities in the topsoil and

| Soil NH₄⁺ | Soil NO₃ | Moisture | Large macroaggregates | Small macroaggregates | Microaggregates | Silt and clay fraction | Aggregate diameter | Root weight | Root surface area | Root diameter | Shoot N | Grain weight |
|----------|---------|----------|----------------------|----------------------|-----------------|----------------------|------------------|------------|-----------------|-------------|----------|-------------|
| −0.51    | −0.45   | 0.52     | 0.17                | 0.43                 | 0.11            | −0.36                | 0.21             | −0.75*     | −0.05           | −0.57       | −0.76*   | −0.46        |
| −0.77*   | −0.40   | 0.92*    | 0.42                | 0.46                 | 0.10            | −0.57                | 0.46             | −0.89*     | −0.59           | −0.52       | −0.91*   | −0.66        |
| −0.62    | −0.17   | 0.93*    | 0.35                | 0.40                 | 0.01            | −0.47                | 0.38             | −0.65      | −0.24           | −0.24       | −0.67*   | −0.48        |
| 0.18     | −0.12   | 0.53     | −0.42               | 0.12                 | 0.09            | 0.27                 | −0.40            | −0.82*     | −0.24           | −0.29       | −0.69*   | −0.67*       |
| −0.66    | −0.28   | 0.71*    | 0.46                | 0.16                 | 0.11            | −0.47                | 0.48             | −0.82*     | −0.34           | −0.34       | −0.83*   | −0.47        |

Table 3. Pearson’s correlation coefficients between microbial diversity indices and environmental variables in the topsoil and subsoil. Asterisks (*) indicate significant correlations between variables at $P < 0.05$ level.
subsoil layers. Our observations here were directly contrary to our hypothesis: the nutrient-rich amendments led to divergence instead of convergence of the microbial communities of the top- and subsoil. Thus, the two soil layers responded differently to the addition of amendments. Where an OTU was higher in abundance in the topsoil, there was a tendency for that OTU to be more abundant in the fertilizer and then poultry litter treatments compared to the nil-amendment control. Similarly, where an OTU was higher in abundance in the subsoil, the poultry litter and then nil treatments tended to be higher in abundance than the synthetic fertilizer. The addition of nutrients or organic matter has been shown to stimulate microbial activity and alter the abundance of select species. It appears that poultry litter had a stimulatory effect on selected microbes in both soil layers, but the fertilizer had a stimulatory effect in the topsoil and an inhibitory effect in the subsoil. This could be due to the chemical composition of the amendments, since differing C chemistry and nutrient content is known to affect microbial communities. The inorganic fertilizer contained highly available macro- and micronutrients whereas the poultry litter contained both nutrients and carbon. Nutrients from the fertilizer might have leached down the soil profile, resulting in an increase of solutes in the soil solution that might have negatively affected the microbial communities there.

Soil and plant properties played a role in shaping microbial community structure in this experiment, with plant roots and soil structure being identified as key drivers. In contrast, soil inorganic N concentration and aboveground plant variables were not strongly correlated with either bacterial or fungal community structure. The structure and function of soil microbial communities are known to be strongly linked to the physical soil environment and to plant roots, with complex interactions occurring between soil biota, roots and aggregates. However, much of the variation in bacterial and fungal community structure in this experiment remained unexplained. It is likely that other edaphic variables – such as pH and soil carbon – may also have been key drivers of microbial community structure after amendment but these were not analyzed in the present study.

Microbial diversity was found to be negatively correlated with soil chemical fertility and plant growth in this experimental system. The richness, diversity and evenness of the bacterial and fungal communities almost always decreased as N concentrations in the soil and shoot, and shoot and root biomass, increased. These results suggest that microbial diversity was not necessarily a critical determinant of increased plant biomass and yield in the present study, since alpha diversity indices were not positively correlated with measures of plant productivity. Many authors have reported strong positive relationships between measures of microbial diversity, soil fertility and plant productivity, indicating that diversity is critical to maintaining ecosystem services. However, others contend that this relationship is complex and should not be generalized and that changes in community composition and activity, rather than diversity, affect ecosystem processes. This experiment demonstrates that a loss of species or reduction in the diversity and evenness of the microbial community does not always have a negative impact on plant growth or yield. Increased root growth and exudation may actually have been a driver of reduced diversity in the soil due to plant selection pressures on soil biota. Additionally, variation in plant growth and microbial communities could be attributed to differences in carbon and nutrient availability of the different amendments, since we suspect that this might have affected microbial community structure and diversity. Because nutrients have a first-order control on both plant productivity and soil microbial communities, manipulating nutrients as we have done here could result in both direct and indirect effects on microbial community structure and diversity, complicating our interpretation of the plant-microbial relationship.

Microbial diversity-plant productivity relationships in degraded ecosystems due to changes in soil nutrient condition and soil inorganic N concentration and plant roots and soil structure being identified as key drivers. In contrast, soil inorganic N concentration and aboveground plant variables were not strongly correlated with either bacterial or fungal community structure. The structure and function of soil microbial communities are known to be strongly linked to the physical soil environment and to plant roots, with complex interactions occurring between soil biota, roots and aggregates. However, much of the variation in bacterial and fungal community structure in this experiment remained unexplained. It is likely that other edaphic variables – such as pH and soil carbon – may also have been key drivers of microbial community structure after amendment but these were not analyzed in the present study.

Finally, there were no obvious large-scale shifts in the functional potential of the microbial community arising from the different amendments applied and their interaction with placement depth. It was expected that the bacterial and fungal OTUs responding to the experimental treatments would have functional roles related to nutrient cycling, carbon decomposition, plant growth promotion and so on. Instead, extrapolating from the OTUs that were observed to differ in abundance between treatments, we observed disparate changes in function that did not support this theory. A small minority of bacterial and fungal OTUs was observed to differ in abundance based on amendment type or placement depth. Many of these OTUs belonged to phyla we know little about, such as Candidatus Saccharibacteria (Candidate division TM7) or unclassified Acidobacteria taxa. Others we can make functional predictions for, such as the Ascomycota which are known to influence soil structure via their hyphae or the N-fixing Xanthobacteraceae, but these observations are not consistent and do not point to a large-scale shift in function after addition of amendment. Further investigation of shifts in microbial community function after addition of amendments requires a more direct approach utilizing techniques such as qPCR or omics-based tools to target functional changes more directly. These degraded ecosystems may have some parallels to the heavily-disturbed experimental units used in this experiment, albeit they occur at vastly different temporal and spatial scales.

**Conclusion**

This experiment revealed that the addition of poultry litter and inorganic fertilizer significantly altered microbial community structure and diversity in the topsoil and subsoil of repacked soil columns. The amendments changed the composition of bacterial and fungal communities by stimulating or suppressing taxa, leading to divergence of the amended soils. Substantial inherent heterogeneity in microbial communities in topsoil and subsoil layers was also observed. However, microbial diversity was found to be negatively correlated with plant productivity in this experimental system. We hypothesize that these findings could be due to the chemical composition of the amendments and the differing release rates of nutrients, which affected both the microbes and the plant.
Methods

Treatments and experimental design. This controlled environment experiment with wheat plants grown in repacked soil columns used a factorial design with 3 amendments (nil-amendment control, poultry litter, slow-release synthetic fertilizer) \times 2 depths of placement (topsoil, subsoil) treatments. There were three replicates of each treatment for a total of six treatments: (1) topsoil + nil-amendment control (‘TOP NIL ’), (2) topsoil + poultry litter (‘TOP PL ’), (3) topsoil + fertilizer (‘TOP FERT’), (4) subsoil + nil-amendment control (‘SUB NIL ’), (5) subsoil + poultry litter (‘SUB PL ’) and (6) subsoil + fertilizer (‘SUB FERT’). There were two complete sets of six treatments to allow destructive harvest at two time points. All columns were planted to spring wheat (*Triticum aestivum* cv. Gauntlet). The addition of \pm plant treatments was not possible due to experimental constraints.

The soil used was a Solonetz that was collected from a property in Ballan in south-eastern Australia. Topsoil (0–15 cm) and subsoil (20–40 cm) were collected and then all soil was air-dried, crushed and sieved to 2 mm. The topsoil had pH(1:5 CaCl\(_2\)) 4.7, organic C 38 g kg\(^{-1}\), total N 4.1 g kg\(^{-1}\), Olsen P 34 mg kg\(^{-1}\), electrical conductivity (EC, 1:5 water) 0.39 dS m\(^{-1}\), cation exchange capacity (CEC) 8.3 cmol kg\(^{-1}\) and the texture was 57% sand, 23% silt and 21% clay. The subsoil had pH(CaCl\(_2\)) 4.9, organic C 8.6 g kg\(^{-1}\), total N 1.2 g kg\(^{-1}\), Olsen P 5.8 mg kg\(^{-1}\), EC 0.06 dS m\(^{-1}\), CEC 4.2 cmol kg\(^{-1}\) and the texture was 53% sand, 26% silt and 21% clay.

The poultry litter was sourced from a broiler operation and sieved through a 4-mm mesh screen. Poultry litter was applied at a rate of 53.3 g per column (equivalent to 30 t ha\(^{-1}\)). The slow-release synthetic fertilizer Macracote Orange (Langley Fertilizers; Perth, Australia) was applied at 15 g per column to achieve a similar total N as the poultry litter equivalent to 1,200 kg N ha\(^{-1}\). The poultry litter contained (% w/w): 34 C, 4.5 N, 1.7 P, 2.7 K and 0.68S. The slow-release synthetic fertilizer contained (% w/w) 16 N (7.8 as CH\(_4\)N\(_2\)O, 4.8 as NH\(_4\)+ and 3.4 as NO\(_3^-\)), 3.5 P (3.2 water-soluble P and 0.3 citrate-soluble P), 10 K and 5.2S (as K\(_2\)SO\(_4\)).

Experimental unit construction. PVC columns of 15 cm in diameter and 40 cm tall were filled with 8.7 kg soil in four layers to simulate the texture-contrast profile of the Solonetz soil. The layers were constructed as follows: 0–10 cm of topsoil, 10–20 cm of a 1:1 topsoil:subsoil blend, 20–30 cm of subsoil and another 30–40 cm of subsoil (Fig. 5). The column was tapped to compact the soil to the desired bulk density of 1.5 g cm\(^{-3}\) in the deep subsoil. The poultry litter and slow-release synthetic fertilizer were thoroughly mixed into either the topsoil (Fig. 5a) or subsoil (Fig. 5b) layer.

Basal nutrients were mixed into the 0–10 cm layer of each column at the following rates (mg kg\(^{-1}\) soil): 180 KH\(_2\)PO\(_4\); 120 K\(_2\)SO\(_4\); 180 CaCl\(_2\); 2H\(_2\)O; 50 MgSO\(_4\); 7H\(_2\)O; 15 MnSO\(_4\); 7H\(_2\)O; 6 CuSO\(_4\); 5H\(_2\)O; 0.4 Na\(_2\)MoO\(_4\); 2H\(_2\)O; 5.5 FeEDTA. A watering pipe was fitted in the center of each column to allow watering of deeper soil layers and moisture probes were inserted into the top of the 10–20 cm and/or 30–40 cm layers of some of the columns to monitor soil moisture throughout the experiment. The soil surface was covered by 2 cm of high-density polyethylene beads to minimize evaporation and a thin layer of beads was added at each 10 cm layer to distinguish them at harvest.

Growing conditions. The experiment was carried out in a controlled environment room with a 14-hour photoperiod, day-time temperature of 22.5 ºC and night temperature of 18.5 ºC. The irradiance increased from 350 to 650 \(\mu\)mol m\(^{-2}\) s\(^{-1}\) at the canopy level as plant height increased. All columns in the controlled environment room had been grouped into three blocks (three shelves) with re-randomization within each shelf twice a week. The room had an atmospheric CO\(_2\) concentration of 480–500 ppm and a relative humidity of 56%.

Twelve pre-germinated seeds of wheat of similar size were sown in each column on Day 0. These were thinned to five plants per column 13 days after sowing (DAS) and two plants per column 30 DAS. The young shoots were
cut at the soil surface, leaving their roots to decompose in the soil. Urea was applied to all columns at a rate of 30 mg N kg⁻¹ soil 37 DAS to alleviate possible N deficiency in organic amendment treatments.

All columns were watered daily to approximately 80% of field capacity. The topsoil was watered from the surface and watering tubes were used to wet the subsoil layers.

**Sampling and measurement.** The experimental units were destructively harvested. One set of columns was harvested at 76 DAS (hard dough to early ripening, Zadoks 87–91⁴⁷) and the second set at 107 DAS (physiological maturity, Zadoks 99⁴⁷). Shoots were cut at the soil surface and dried at 70°C for 72 hours before recording the mass. For the second harvest, all ears were removed and threshed to separate the grain kernels before the total grain weight was recorded.

Detailed measurements were conducted for the samples of the first harvest at 76 DAS. A subsample of shoot biomass was ground with a ball mill and N concentration was measured with a Series II CHNS/O Analyser (PerkinElmer; Colorado, USA). Soil from each column was sectioned into the four 10-cm layers and a subsample was collected for analysis of the soil microbial community, gravimetric moisture content, inorganic N and water-stable aggregates. The rest of the soil from each layer was sieved and washed to retrieve the roots. Subsamples of roots were scanned and analyzed with WinRHIZO (Regent Instruments Inc.; Canada) for root length, surface area and diameter. Roots were then dried at 70°C for 72 hours before recording the mass.

Soil inorganic N was measured with a QuickChem® 8500 Series II FIA Automated Ion Analyser (Lachat Instruments; USA) after extraction with 2 M KCl (1:5 soil to solution ratio), shaking for 1 hour and filtering with No. 1 Whatman filter paper. Aggregate size distribution was determined using a standard wet-sieving apparatus whereby 25 g of 10 mm-sieved, air-dried soil was placed on top of a nest of pre-weighed sieves and immersed in deionized water for 10 min. The stack of sieves was then subjected to automatic vertical movement for 15 min at 70 rpm. Four aggregate size fractions were collected: large macroaggregates (>2 mm), small macroaggregates (0.25–2 mm), microaggregates (0.053–0.25 mm) and the silt and clay fraction (<0.053 mm). Mean-weight diameter (MWD) of soil aggregates was calculated using the standard formula:

\[
MWD = \sum x_i w_i
\]

where \(x_i\) is the mean aperture of the adjacent sieves and \(w_i\) is the mass fraction remaining on each sieve⁴⁸.

Genomic DNA was extracted from 0.25 g soil samples using the Mobio PowerSoil DNA Isolation Kit as per the manufacturer's instructions. DNA concentration and purity were determined using an Implen P330 NanoPhotometer (Implen GmbH, Munich, Germany) and Qubit 3.0 Fluorometer (ThermoFisher Scientific; Invitrogen, MA, USA). 16S rRNA and ITS diversity profiling was performed by the Australian Genome Research Facility (Melbourne, Australia) on an Illumina MiSeq (Illumina Inc., CA, USA) platform. A 300 bp target was amplified from the V3-V4 region of the 16S rRNA gene using primers 341 F (5′- CCTAYGGGGRBGGASCAG) and 806 R (5′- GGACTACNNGGTATCTAAT)⁴⁹,⁵⁰ and an approximately 230 bp target was amplified from the ITS1-ITS2 region of the internal transcribed spacer (ITS) using primers ITS1f (5′- CTTGTTACATTAGGAAGTAA) and ITS2 (5′- GCTGCGTTCTTCATCGATGC) (White et al. 1990, Gardes and Bruns 1993).

**Bioinformatics.** Raw, demultiplexed fastq files from the Australian Genome Research Facility were re-barcoded, joined and quality filtered using the UPARSE pipeline⁵¹. Joined paired-end reads were quality-filtered by discarding reads with total expected errors > 1 and removing singletons. Operational taxonomic units (OTUs) were clustered with a minimum cluster size > 2 and 97% similarity cut off to enable detection of community-level changes using the UPARSE clustering algorithm. Taxonomic assignments were performed using the USEARCH UTAX algorithm with reference databases created using the RDP 16S (version 16) and MUSCLE⁵². OTUs identified as chloroplasts and mitochondrial DNA were removed from the data set and all OTUs with lower than 80% taxonomic confidence threshold were denoted as ‘Unassigned.’ A phylogenetic tree was constructed using the UPGMA algorithm in MUSCLE⁵².

**Statistical analysis.** All analyses were carried out using R version 3.5.0⁵³ and PRIMER version 7 software with the PERMANOVA + add-on⁵⁴,⁵⁵. Plots were produced in R with the assistance of packages ggplot2⁶⁰ and RColorBrewer⁶¹. Using package phyloseq⁶⁸, spurious reads were removed using a 0.005% relative abundance cut-off⁶⁹ and the bacterial and fungal data sets were rarefied to 17,000 and 3,500 reads, respectively (Supplementary Fig. S1). Alpha diversity analyses (Chao1 richness estimator⁷⁵, Shannon diversity index (H)⁴⁸,⁵⁹) were performed on bacterial and fungal OTU tables using the estimate_richness function in the package phyloseq⁶⁸. Simpson’s evenness (E) was calculated by dividing the inverse Simpson’s index by observed species richness⁵⁹. Weighted and unweighted UniFrac distances between samples⁶⁸ were calculated in phyloseq⁶⁸. Principal coordinates analysis of weighted UniFrac distances was used to visualize the relationships and differences between treatments. Global and pairwise permutational multivariate analysis of variance (PERMANOVA) on the distance matrices was performed in using the PERMANOVA + add-on in PRIMER⁶⁸. All PERMANOVA tests used 9999 permutations from unrestricted permutation of raw data. Where there were fewer than 99 unique permutations for a meaningful test, approximate Monte Carlo P-values (PM(C)) were obtained from an asymptotic permutation distribution⁵⁵. Package mvabund⁶¹ was used to determine which microbial OTUs differed significantly (Padj < 0.05) in abundance between amendment and depth treatments. For this procedure, un rarefied sequence counts were modelled on negative binomial distributions in the generalized linear models. Pearson’s correlation analysis was performed using the rcorr function in package Hmisc⁶² to determine relationships between alpha
diversity indices and environmental variables. Function oridstep in package vegan was used to select significant ($p < 0.05$) drivers of microbial community composition from a standardized matrix of the 15 environmental variables (Supplementary Table S4) using both forward and backward selection. Highly correlated ($R^2 > 0.90$) variables were removed prior to model selection. Canonical correspondence analysis (CCA) was carried out using vegan to correlate sample ordination with the retained variables. Variation partitioning analysis was used to quantify the effects of the significantly correlated environmental variables on the microbial community composition using varpart function in vegan.

Data Availability
The datasets generated during the current study are available from the corresponding author on reasonable request.

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**Author Contributions**

J.B.M., X.W. and C.T. designed and conducted the original planted soil column experiment; C.C designed and conducted the microbial component of the experiment; C.C. wrote the manuscript with leading contribution by J.L.W. and input from J.B.M., X.W., P.W.G.S., C.T. and A.E.F.

**Additional Information**

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