Living cover crops have immediate impacts on soil microbial community structure and function

D.M. Finney, J.S. Buyer, and J.P. Kaye

Abstract: Cover cropping is a widely promoted strategy to enhance soil health in agricultural systems. Despite a substantial body of literature demonstrating links between cover crops and soil biology, an important component of soil health, research evaluating how specific cover crop species influence soil microbial communities remains limited. This study examined the effects of eight fall-sown cover crop species grown singly and in multispecies mixtures on microbial community structure and soil biological activity using phospholipid fatty acid (PLFA) profiles and daily respiration rates, respectively. Fourteen cover crop treatments and a no cover crop control were established in August of 2011 and 2012 on adjacent fields in central Pennsylvania following spring oats (Avena sativa L.). Soil communities were sampled from bulk soil collected to a depth of 20 cm (7.9 in) in fall and spring, approximately two and nine months after cover crop planting and prior to cover crop termination. In both fall and spring, cover crops led to an increase in total PLFA concentration relative to the arable weed community present in control plots (increases of 5.37 nmol g⁻¹ and 10.20 nmol g⁻¹, respectively). While there was a positive correlation between aboveground plant biomass (whether from arable weeds or cover crops) and total PLFA concentration, we also found that individual cover crop species favored particular microbial functional groups. Arbuscular mycorrhizal (AM) fungi were more abundant beneath oat and cereal rye (Secale cereale L.) cover crops. Non-AM fungi were positively associated with hairy vetch (Vicia villosa L.). These cover crop–microbial group associations were present not only in monocultures, but also multispecies cover crop mixtures. Arable weed communities were associated with higher proportions of actinomycetes and Gram-positive bacteria. Soil biological activity varied by treatment and was positively correlated with both the size and composition (fungal:bacterial ratio) of the microbial community. This research establishes a clear link between cover crops, microbial communities, and soil health. We have shown that while cover crops generally promote microbial biomass and activity, there are species-specific cover crop effects on soil microbial community composition that ultimately influence soil biological activity. This discovery paves the way for intentional management of the soil microbiome to enhance soil health through cover crop selection.

Key words: cover crops—phospholipid fatty acid analysis—soil biological activity—soil health—soil microbial communities

Building healthy, resilient soils is a central focus of US soil conservation programs, and cover cropping is a core strategy promoted to enhance physical, chemical, and biological properties of managed soils (Lehman et al. 2015a; USDA NRCS 2012). While considerable research has shown that including cover crops in a crop rotation influences soil biology, understanding of cover crop species-specific effects on soil microbial community size, composition, and function and the time frame in which these effects are manifest remains limited. There is a critical need to identify how cover crop selection shapes soil microbial communities in order to develop efficient and effective management strategies that will lead to the restoration of healthy microbial communities in agricultural systems (Lehman et al. 2015b). This study examined the effects of eight fall-sown cover crop species grown singly and combined in multispecies mixtures on the size, composition, and function of soil microbial communities in living cover crop stands.

Soil health is defined as “the continued capacity of soil to function as a vital living system, within ecosystem and land-use boundaries, to sustain biological productivity, maintain the quality of air and water environments, and promote plant, animal, and human health” (Doran et al. 1996). This concept recognizes the central role that soil biota play in providing ecosystem services that are critical to agricultural systems such as nutrient retention and provision, disease suppression, enhancement of soil structure and stability, and weed management (Lehman et al. 2015b). A number of soil organisms contribute functions that ultimately provide these services, including a diverse array of soil microorganisms. Though linking structural features of soil microbial communities (i.e., biomass and composition) to specific soil functions remains an open field of investigation (Lehman et al. 2015a), soil health management recommendations frequently target the following two goals for the soil microbial community: increasing microbial biomass and enhancing microbial diversity (USDA NRCS 2012).

Plant communities are a primary source of the carbon (C) supporting microbial growth and activity; therefore, crop management can be used to promote increases in the size of the soil microbial community in agricultural systems. For instance, a recent meta-analysis demonstrated that crop rotations increase soil microbial biomass by an average of 21% relative to monocultures (McDaniel et al. 2014b). This same analysis also demonstrated a strong influence of cover crops on this rotational effect, as cover crops led to an 8.5% increase in total soil C (McDaniel et al. 2014b). Cover crops are typically unharvested crops planted between cash crops that augment C provisioning to the soil system not only via
unharvested residues, but also during active cover crop growth. During the latter period, C inputs derived from root exudation, fine root turnover, and litter decomposition may be sufficient to promote increases in soil microbial biomass (Wardle et al. 2004; Buyer et al. 2010; Maul and Drinkwater 2010; Kong and Six 2012). Despite this universally recognized link between plant and microbial communities, very few studies have actually examined microbial communities in living cover crop stands (Schutter et al. 2001; Lehman et al. 2012). Much of our knowledge of relationships between cover crops and soil communities is based on studies of legacy effects of cover crop residues, the results of which may be convoluted by complex interactions between cover crops and other management factors that influence microbial communities such as tillage (Treonis et al. 2010; Wortman et al. 2013; Lienhard et al. 2014) and fertility amendments (Grayston et al. 2004; Saison et al. 2006; Carrera et al. 2007). Studies designed to isolate cover crop-specific effects on microbial communities from other influential management practices are essential to building our understanding of interactions between cover crops and microbes.

A diverse soil microbial community is viewed as capable of providing a broad range of functions and demonstrating greater resilience of those functions to physical and chemical disturbance (Lehman et al. 2015a). While there is no “optimal composition” for a healthy microbial community, promotion of certain microbial groups has been shown to enhance soil health. For example, arbuscular mycorrhizal fungi (AM fungi) enhance crop production by protecting host plants from pathogens, improving nutrient uptake, and increasing host plant tolerance to environmental stresses such as drought (Rillig 2004). Previous studies offer evidence that cover crops, and particularly grass species such as oat (Avena sativa L. [Kabir and Koide 2002; Lehman et al. 2012]), cereal rye (Secale cereale L. [White and Weil 2010]), and winter wheat (Triticum aestivum L. [Kabir and Koide 2000]), can, in fact, increase AM fungi in agricultural soils. Promoting the establishment of fungal communities in agricultural soils may also promote soil health more generally. A key indicator of soil health is soil organic matter (SOM), increases in which lead to enhanced nutrient cycling, aggregate stability, and water-holding capacity (Lehman et al. 2015b). Both the quantity and quality of SOM are improved in fungal-dominated microbial communities (Six et al. 2006). Crop rotation has been shown to influence the relative abundances of fungi and bacteria in agricultural systems (Tiemann et al. 2015); however, the significance of cover crops to this outcome remains unknown. Research in unfertilized grasslands has demonstrated that fungal communities respond positively to plant-derived C inputs, suggesting that inclusion of cover crops in a rotation may promote fungal community development (Denef et al. 2009), and previous studies have shown increases in fungal population density in some cover cropping systems (Schutter et al. 2001; Buyer et al. 2010). The fact that certain cover crop species have been associated with increases in fungal groups suggests that cover crop selection could play an important role in managing soils for greater fungal dominance.

Phospholipid fatty acid (PLFA) analysis has been widely used to assess the composition of microbial communities in agricultural systems and community response to cover crops (Bossio et al. 1998; Schutter et al. 2001; Buyer et al. 2010; Treonis et al. 2010; Lehman et al. 2012; Wortman et al. 2013). Phospholipids are a component of all cellular membranes, and signature phospholipids are found in the membranes of specific microbial groups. Quantifying these “biomarkers” provides a means of determining the abundance of different bacterial and fungal groups, as well as protozoa in a soil community (Zelles 1999). Though other molecular techniques may provide a more detailed description of microbial diversity, PLFA offers the advantages of quantifying the total microbial biomass, providing a fungal:bacterial ratio, and lower cost. The recent development of high throughput PLFA analysis is likely to further increase the cost-effectiveness and utility of this tool (Buyer and Sasser 2012).

One impediment to translating our current knowledge of the relationships between cover crops and microbes into management strategies to effectively promote soil health is the fact that the number of cover crops included in any single study is often limited. This limitation makes it difficult to determine if observed effects are generalizable across cover crop species or actually species-specific. The latter is a reasonable expectation based on evidence that plant influences on belowground communities are often species-specific (Wardle et al. 2004; Berg and Smalla 2009; Eisenhauer et al. 2010; Bezemer et al. 2010; Maul and Drinkwater 2010). Recognizing that management and environmental factors such as soil pH (Fierer and Jackson 2006), soil type (Buyer et al. 2002; Girvan et al. 2003), and climate (Drenovsky et al. 2004; McDaniel et al. 2014a) also mediate soil microbial communities, studies comparing a greater number of cover crops within a uniform management and environmental context are critical to unraveling the links among cover crops, microbes, and soil health.

This study examined the effects of eight fall–summer cover crops grown singly and combined in multispecies mixtures on the size, composition, and function of soil microbial communities. PLFA analysis was used to characterize the structure of microbial communities present in the bulk soil of living cover crop stands approximately two and nine months after planting, prior to cover crop termination. We expected that the presence of cover crops would lead to increased microbial biomass and changes in community composition compared to the arable weed community present in an un Tillage fallow control. We further expected that species-specific plant effects would lead to variation in community composition across treatments. Short-term laboratory incubation was used to determine if changes in microbial community structure resulted in changes in biological activity, an important indicator of soil health. The results of this study provide much-needed information for land managers to guide the selection of cover crop species that will enhance soil health.

Materials and Methods

Field Experiment. We conducted a two-year field study of cover crop diversity at the Russell E. Larson Agricultural Research Center near Rock Springs, Pennsylvania, United States, as described in Finney et al. (2016). A total of 17 (2011 to 2012) and 18 (2012 to 2013) cover crop treatments were planted in adjacent fields (i.e., different fields were used for each year of the study) in late August within an oat–corn (Zea mays L.) cash crop rotation following small grain harvest. We selected a subset of these treatments (15) including the no cover control (an un Tillled weedy fallow), eight species grown in monocultures, and six mixtures composed of the monoculture species for microbial
community analysis (Table 1). Cover crop treatments were established in 9.1 × 6.5 m (30 × 21 ft) plots in a randomized complete block design replicated four times. Each site was moldboard plowed and disked before cover crop planting. We planted all cover crop treatments with a drill fitted with a cone seed distributor and mixed legume seed with dry inoculant before planting. Cover crops were terminated with glyphosate and subsequently incorporated prior to corn planting the following spring. All soil subject to microbial community analysis was collected prior to cover crop termination (see below). We also measured cover crop and arable weed biomass in fall prior to the first killing frost and spring immediately prior to termination by clipping within quadrats. Aboveground biomass data have previously been reported in Finney et al. (2016).

Soil Sampling. Bulk soil samples were collected in fall at the time of peak cover crop growth and spring approximately one week prior to cover crop termination. At each sampling event, we collected 12 cores (0.20 m depth by 0.02 m inside diameter [7.9 in depth by 0.79 in inside diameter]) from each experimental plot (six cores from within the cover crop row and six cores from between cover crop rows) and homogenized cores in a plastic bag. We placed samples in coolers for temporary storage in the field and transport to the lab. In the lab, we stored samples at 4°C (39.2°F) for no more than 24 hours prior to laboratory processing. We sieved homogenized samples to 2 mm (0.07 in). A 50 g (1.8 oz) subsample was transferred to a sterile specimen cup and stored at −20°C (−4°F) for PLFA analysis. A 100 g (3.5 oz) subsample was transferred to a fresh plastic bag and stored at 4°C (39.2°F) for laboratory incubation.

Soil sampling and laboratory processing were conducted in an aseptic manner to the extent possible. Specifically, we wore latex gloves during field sampling and rinsed soil probes with 70% reagent alcohol prior to collection from each plot. Similarly, we wore latex gloves for lab activities and changed gloves during field sampling and rinsed soil probes with 70% reagent alcohol prior to collection. Lab equipment (soil sieves and spoons) was washed with soap and water between samples. Lab equipment (soil sieves and spoons) was washed with soap and water between samples. Lab equipment (soil sieves and spoons) was washed with soap and water between samples. We also measured cover crop and arable weed biomass in fall prior to the first killing frost and spring immediately prior to termination by clipping within quadrats. Aboveground biomass data have previously been reported in Finney et al. (2016).

High-throughput PLFA was performed as described in Buyer and Sasser (2012). Briefly, ~2 g (0.1 oz) lyophilized soil was extracted with 4 ml (0.135 fl oz) Bligh-Dyer extractant and the resulting liquid phase separated with 1 ml (0.034 fl oz) each of chloroform and deionized water. The top aqueous phase was aspirated and the bottom lipid-containing phase dried. Lipid classes were separated by solid phase extraction (SPE) chromatography in a 96–well plate with 50 mg (0.002 oz) silica gel per well (Phenomenex, Torrence, California). Lipids were dissolved in chloroform and samples passed through the silica gel prior to washing with 1 mL (0.034 fl oz) chloroform and 1 mL acetone. Phospholipids were then eluted with 0.5 ml (0.017 fl oz) 5:5:1 methanol:chloroform:deionized water into 1.5 ml (0.051 fl oz) vials and dried in vacuo. Following addition of 0.2 ml (0.007 fl oz) transesterification reagent and incubating for 15 minutes at 37°C, 0.4 ml (0.014 fl oz) 0.075 M acetic acid was added, and samples were extracted twice with 0.4 ml chloroform. Chloroform was removed by drying and lipids dissolved in 75 μl hexane prior to transfer to gas chromatography vials with conical inserts. Samples were stored at −20°C (−4°F) as needed during the extraction process and until analysis.

Samples were analyzed on an Agilent (Agilent Technologies, Wilmington, Delaware) 6890 gas chromatograph (GC) outfitted with an autosampler, split-splitless injector, and flame ionization detector. MIS Sherlock software (MIDI, Inc., Newark, Delaware) was used in conjunction with Agilent Chemstation to control the system for the analysis. Fatty acids were separated on an Agilent Ultra 2 column (25 m × 200 μm inside diameter × 0.33 μm film thickness) using hydrogen (H) gas as the carrier (1.2 ml min−1 [0.04 oz min−1] flow rate) and identified using the MIDI PLFAD1 calibration mix and naming table. We also ran random samples on a Clarus 500 GC-MS (Perkin-Elmer, Waltham, Massachusetts, United States) to confirm fatty acid identification following the procedure of Buyer et al. (2010).

Laboratory Incubation. We conducted a seven day incubation of fresh soil to measure daily respiration rate, an indicator of soil biological activity (Weitzman et al. 2014). At initiation, 20 g (0.7 oz) of fresh soil was placed in a 100 mL (3.38 fl oz) Wheaton serum bottle and adjusted to 50% water holding capacity (WHC) using deionized water. Each vial was capped with a butyl stopper, and an aluminum (Al) ring was used to create an air-tight seal. Vial headspace was sampled (1 mL) with a syringe at 24 hour intervals for the first 72 hours. Following each sample collection, the stopper was removed in order to flush accumulated carbon dioxide (CO2) and replaced after approximately one hour. Following the 72 hour collection and subsequent flushing, the stopper was replaced and the bottle sealed until the final headspace sampling. Carbon dioxide concentrations in the headspace gas were quantified...
immediately after collection with an infrared
gas analyzer (LI7000, LiCor, Inc., Lincoln,
Nebraska). Incubation bottles were stored at
room temperature in a dark cabinet through the
course of the incubation. Cumulative CO$_2$-C released during the first seven days of
incubation was calculated by linear inter-
polation between headspace sampling dates.
Results are presented as the daily respiration
rate in grams of dry soil (μg CO$_2$-C g$^{-1}$ soil
d$^{-1}$ ± 1 standard error of the mean).

**Statistical Analysis.** We used analysis of
variance (ANOVA) to determine the effect
of cover crop treatment on the size and activity
of the soil microbial community. Total
PLFA (the sum of all detected fatty acids
expressed in nmol g$^{-1}$ soil) was the indicator
of microbial community size. PLFA data
were log transformed for analysis to meet
assumptions of normality and least square
means were back-transformed for presenta-
tion. The daily respiration rate in the seven
day incubation (μg CO$_2$-C g$^{-1}$ soil d$^{-1}$) was
an indicator of biological activity. We used a
mixed model with fixed effects of treatment
and year and their interaction and a random
effect of block nested within year to identify
differences among treatments and between
years within each season (i.e., separate analyses
were conducted on fall and spring data).

To account for an unbalanced design in
PLFA data across years, interaction tests were
performed excluding soybean (Glycine max
[L.] Merr.) and sunn hemp (Crotalaria juncea
L.) monocultures for which data were available
only in Year 1. We also used contrast analysis to
detect differences in size and activity between
the no cover crop control and cover cropped
treatments within each season. Separation
of least square means was performed using Tukey’s honestly significant difference (HSD)
at α = 0.05 (Seale et al. 1980).

Using PLFA data, we assessed the effect
of cover crop treatment on microbial community
composition. For these analyses, PLFA peak areas were combined into bio-
marker groups following Buyer et al. (2010)
as shown in table 2. We used ANOVA to
detect differences in the concentration of
biomarker groups between treatments using
the mixed model approach described above.
We also calculated the ratio of fungal to bac-
terial biomarkers for each community as the
sum of AM and non-AM fungi biomarker
concentrations divided by the sum of Gram-
positive, Gram-negative, and actinomycete
biomarker concentrations. We performed
ANOVA on fungal:bacterial ratios using
the same mixed model previously described
to detect differences among treatments. To
assess the influence of cover crop treatment
on microbial community composition, we
performed distance-based redundancy analysis (db-RDA) on biomarker groups.
Redundancy analysis allowed us to assess
microbial community responses specific to
environmental changes associated with each
cover crop treatment without incorporating
community responses to other factors in
the environment that were not influenced
by treatment (McCune and Grace 2002).

Distance-based RDA was differentiated from
RDA by its use of distance measures other
than Euclidean distance. Biomarker group
concentrations were Hellinger-transformed
prior to analysis. Treatment significance
was tested using permutation tests (Borcard
et al. 2011). We used a second db-RDA to
determine how the quantity and identity of
aboveground biomass impacted microbial community composition. Environmental
variables included in this analysis were cover
crop biomass by species and weed biomass.
All redundancy analyses were performed by
season using Gower distances and condi-
tioned on block nested within year.

Spearman rank correlations were used to
identify relationships between aboveground
biomass production and total PLFA con-
centration, as well as relationships between
microbial community structure (i.e., size as
indicated by total PLFA concentration and
composition as indicated by fungal:bacterial ratio) and daily respiration rate. All statistical
analyses were conducted using R statistical
software (R Development Core Team 2013).

**Results and Discussion**

**Cover Crop Effects on Soil Microbial Biomass.** The presence of cover crops
increased microbial biomass (indicated by
total PLFA concentration) relative to the no
cover crop control. While total PLFA concentra-
tion did not vary among treatments in the
fall, total concentrations were higher in cover
cropped soils compared to the no cover crop
treatment (estimated difference = 5.37 nmol
$g^{-1}$; p = 0.04; table 3) within approximately
two months of planting. Results were similar
in the spring; though total PLFA concentra-
tion did not vary among treatments, cover
crops led to an average increase of 10.22
nmol g$^{-1}$ relative to the control (p < 0.01;
table 4). Total PLFA concentration varied by
year in the spring, indicating greater overall
microbial biomass in Year 1 compared to Year
2 (table 2). Such variability due to year was
not unexpected given differences in climate
and soil conditions between sites (Buyer et al.
2002; Drenovsky et al. 2004).

There was a positive correlation between
total aboveground plant biomass (the sum
cover crop and weed biomass) and total
PLFA within each season ($r = 0.43; p <
0.01$). This result is consistent with the
broadly recognized link between above-
and belowground productivity (Wardle et
al. 2004) and supports the general consen-
sus that the presence of living plant biomass
provides resources to support microbial popu-
lations during periods that would otherwise
be fallow in an agricultural system (Lehman
et al. 2015b).

Previous studies have found that cover
crops can support increases in soil microbial
biomass (McDaniel et al. 2014b); however,
many cover crop studies have measured
microbial biomass following cover crop
termination (Ingels et al. 2005; Wang et al.
2007; Nair and Ngouajio 2012), or only
observed cover crop effects after incorpotation (Schutter et al. 2001). In this study we
measured soil biological properties in living
cover crop stands in order to isolate cover
crop effects on microbial communities from

### Table 2

<table>
<thead>
<tr>
<th>Biomarker group</th>
<th>Signature fatty acids</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gram-positive bacteria</td>
<td>Iso and anteiso branched fatty acids</td>
<td>Zelles 1999</td>
</tr>
<tr>
<td>Gram-negative bacteria</td>
<td>Monounsaturated fatty acids, cyclopropyl 17:0 and 19:0</td>
<td>Zelles et al. 1997</td>
</tr>
<tr>
<td>Actinomycetes</td>
<td>10-methyl 16:0 and 10-methyl 18:0</td>
<td>Zelles 1999</td>
</tr>
<tr>
<td>Arbuscular mycorrhizal (AM) fungi</td>
<td>16:1 ω5 cis</td>
<td>Olsson 1999</td>
</tr>
<tr>
<td>Fungi (non-AM fungi)</td>
<td>18:2 ω6 cis</td>
<td>Frostegard and Baath 1996</td>
</tr>
<tr>
<td>Protozoa</td>
<td>20:3 ω6 cis and 20:4 ω6 cis</td>
<td>Ringelberg et al. 1997</td>
</tr>
</tbody>
</table>
In the fall, concentrations of non-AM fungi, AM fungi, and protozoa in Year 1, though the difference was not significant for AM fungi in the fall (tables 3 and 4). In the fall, abundances of Gram-positive bacteria and actinomycetes were higher in Year 2 compared to Year 1 (table 3). The opposite pattern was observed in spring; abundances of Gram-positive bacteria and actinomycetes as well as Gram-negative bacteria were higher in Year 1, following the same pattern as total PLFA concentration (table 4).

Distance-based RDA indicated that cover crop treatment was a significant predictor of microbial community composition in both fall and spring (fall: $p < 0.01$; spring: $p < 0.01$). In both seasons, the first axis was significant (fall: $p < 0.01$; spring: $p < 0.01$) and separated treatments with high proportions of Gram-positive bacteria and actinomycetes from those with high proportions of Gram-negative bacteria, non-AM fungi, AM fungi, and protozoa (figures 1a and 1b). A differentiation of communities based on proportions of Gram-positive versus Gram-negative bacteria likely reflects the dominant metabolic strategy used by each of these bio-

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### Table 3

Effect of year and cover crop on least square means and standard errors (standard errors are values in parentheses) of phospholipid fatty acid (PLFA) concentration (nmol g$^{-1}$) in bulk soil in fall approximately two months following cover crop planting in August of 2011 (Year 1) and August of 2012 (Year 2) in central Pennsylvania.

<table>
<thead>
<tr>
<th>Year/crop</th>
<th>Total PLFA</th>
<th>Gram + bacteria</th>
<th>Gram - bacteria</th>
<th>Actinomycetes</th>
<th>Non-AM fungi</th>
<th>AM fungi</th>
<th>Protozoa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year 1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No cover crop</td>
<td>71.66 (0.69)</td>
<td>17.40 (0.18)$^b$</td>
<td>21.90 (0.24)</td>
<td>9.95 (0.10)$^b$</td>
<td>1.21 (0.04)$^a$</td>
<td>2.95 (0.04)</td>
<td>0.76 (0.02)$^a$</td>
</tr>
<tr>
<td>Sunn hemp (SH)</td>
<td>77.45 (1.23)</td>
<td>20.29 (0.32)$^a$</td>
<td>24.20 (0.40)</td>
<td>12.07 (0.20)$^a$</td>
<td>0.84 (0.05)$^b$</td>
<td>3.04 (0.06)</td>
<td>0.51 (0.03)$^b$</td>
</tr>
<tr>
<td>Soybean (SB)</td>
<td>72.73 (1.30)</td>
<td>18.82 (0.63)</td>
<td>22.51 (0.31)</td>
<td>11.07 (0.32)</td>
<td>0.90 (0.12)</td>
<td>2.93 (0.10)</td>
<td>0.59 (0.05)</td>
</tr>
<tr>
<td>Red clover (RC)</td>
<td>69.34 (1.55)</td>
<td>17.50 (0.62)</td>
<td>21.33 (0.51)</td>
<td>10.34 (0.68)</td>
<td>0.93 (0.11)</td>
<td>2.72 (0.08)</td>
<td>0.59 (0.06)</td>
</tr>
<tr>
<td>Hairy vetch (HV)</td>
<td>76.90 (2.90)</td>
<td>19.55 (0.87)</td>
<td>23.36 (1.00)</td>
<td>11.09 (0.55)</td>
<td>1.20 (0.14)</td>
<td>2.94 (0.14)</td>
<td>0.59 (0.10)</td>
</tr>
<tr>
<td>Forage radish (FR)</td>
<td>73.55 (2.27)</td>
<td>18.27 (0.56)</td>
<td>23.08 (0.80)</td>
<td>10.80 (0.32)</td>
<td>1.13 (0.20)</td>
<td>2.91 (0.11)</td>
<td>0.59 (0.06)</td>
</tr>
<tr>
<td>Oat (OA)</td>
<td>78.15 (2.16)</td>
<td>19.40 (0.76)</td>
<td>24.37 (0.80)</td>
<td>11.36 (0.64)</td>
<td>1.23 (0.15)</td>
<td>3.26 (0.14)</td>
<td>0.60 (0.11)</td>
</tr>
<tr>
<td>Canola (CA)</td>
<td>74.70 (1.92)</td>
<td>18.90 (0.56)</td>
<td>23.28 (0.77)</td>
<td>10.91 (0.55)</td>
<td>1.01 (0.11)</td>
<td>2.86 (0.08)</td>
<td>0.61 (0.06)</td>
</tr>
<tr>
<td>Cereal rye (CR)</td>
<td>74.03 (4.41)</td>
<td>18.38 (1.27)</td>
<td>23.04 (1.64)</td>
<td>10.63 (0.66)</td>
<td>1.11 (0.12)</td>
<td>2.96 (0.15)</td>
<td>0.63 (0.04)</td>
</tr>
<tr>
<td>FR + OA + CA + CR</td>
<td>74.53 (3.88)</td>
<td>18.31 (0.99)</td>
<td>23.21 (1.09)</td>
<td>10.80 (0.51)</td>
<td>0.95 (0.12)</td>
<td>3.10 (0.20)</td>
<td>0.65 (0.06)</td>
</tr>
<tr>
<td>SH + SB + FR + OA</td>
<td>76.25 (3.47)</td>
<td>19.03 (1.22)</td>
<td>23.97 (1.00)</td>
<td>11.25 (0.82)</td>
<td>1.09 (0.12)</td>
<td>3.11 (0.17)</td>
<td>0.67 (0.09)</td>
</tr>
<tr>
<td>RC + HV + CA + CR</td>
<td>76.30 (4.53)</td>
<td>19.40 (1.42)</td>
<td>23.31 (1.43)</td>
<td>11.10 (0.99)</td>
<td>1.15 (0.15)</td>
<td>3.11 (0.18)</td>
<td>0.67 (0.06)</td>
</tr>
<tr>
<td>RH + SB + CA + CR</td>
<td>73.82 (2.81)</td>
<td>18.73 (0.89)</td>
<td>22.44 (0.71)</td>
<td>10.97 (0.41)</td>
<td>0.96 (0.13)</td>
<td>2.94 (0.10)</td>
<td>0.70 (0.19)</td>
</tr>
<tr>
<td>RC + HV + FR + OA</td>
<td>76.52 (2.83)</td>
<td>19.42 (1.01)</td>
<td>23.47 (1.04)</td>
<td>11.28 (0.68)</td>
<td>0.97 (0.10)</td>
<td>3.20 (0.13)</td>
<td>0.75 (0.06)</td>
</tr>
<tr>
<td>8 species mix</td>
<td>78.10 (3.59)</td>
<td>19.35 (1.07)</td>
<td>24.67 (1.21)</td>
<td>11.19 (0.60)</td>
<td>1.16 (0.21)</td>
<td>3.14 (0.17)</td>
<td>0.76 (0.07)</td>
</tr>
</tbody>
</table>

Notes: Gram + = Gram positive. Gram – = Gram negative. AM = arbuscular mycorrhizal. df = degrees of freedom. Est. = estimate. Values within a column with different letters were significantly different due to year or cover crop based on Tukey’s honestly significant difference ($p < 0.05$). The absence of letters in a column indicates that the effect of year or cover crop was not significant.
concur with work in grassland communities. This association may be evident by the increase in Gram-negative bacteria, both fungal groups, and protozoa were found in higher proportions of fungi was an important factor in discriminating the composition of microbial communities associated with individual treatments in both fall and spring (figures 1a and 1b). By spring, the relative proportion of AM fungi versus non-AM fungi also served to differentiate individual treatments, indicated by the significance of axis two ($p = 0.02$; figure 1b). Observed concentrations of fungi (both AM and non-AM) tended to be lower than those of bacterial groups (tables 3 and 4), which most likely reflects the difference in the number of biomarkers used to identify fungi compared to bacteria. The generally positive association between cover crops and fungal communities may be indicative of conditions favorable for the recovery of these communities following tillage disturbance in cover cropped systems (Kabir et al. 1997; Schipanski et al. 2014). Cover crops also led to higher concentrations of the protozoan biomarkers in spring compared to the control (estimated difference = 0.19 nmol g$^{-1}$, $p = 0.01$; table 4), and this microbial group

<table>
<thead>
<tr>
<th>Year/crop</th>
<th>Total PLFA</th>
<th>Gram + bacteria</th>
<th>Gram – bacteria</th>
<th>Actinomycetes</th>
<th>Non-AM fungi</th>
<th>AM fungi</th>
<th>Protozoa</th>
</tr>
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<tbody>
<tr>
<td>Year 1</td>
<td>100.14 (1.28)a</td>
<td>25.14 (0.31)a</td>
<td>31.01 (0.44)a</td>
<td>13.32 (0.16)a</td>
<td>2.43 (0.10)a</td>
<td>4.19 (0.08)a</td>
<td>1.05 (0.04)a</td>
</tr>
<tr>
<td>Year 2</td>
<td>73.53 (1.43)b</td>
<td>18.54 (0.33)b</td>
<td>23.38 (0.49)b</td>
<td>10.92 (0.23)b</td>
<td>1.21 (0.09)b</td>
<td>2.72 (0.07)b</td>
<td>0.47 (0.03)b</td>
</tr>
<tr>
<td>No cover crop</td>
<td>77.32 (6.29)</td>
<td>20.31 (1.64)</td>
<td>23.48 (1.96)b</td>
<td>11.67 (0.72)</td>
<td>1.42 (0.31)bc</td>
<td>2.96 (0.33)b</td>
<td>0.58 (0.08)</td>
</tr>
<tr>
<td>Sunn hemp (SH)</td>
<td>85.89 (5.44)</td>
<td>22.77 (1.48)</td>
<td>26.11 (1.87)ab</td>
<td>12.47 (0.80)</td>
<td>1.81 (0.58)abc</td>
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<td>0.54 (0.04)</td>
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<tr>
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<td>82.78 (5.36)</td>
<td>21.64 (1.28)</td>
<td>25.24 (1.58)abc</td>
<td>12.06 (0.77)</td>
<td>1.70 (0.28)abc</td>
<td>3.26 (0.35)ab</td>
<td>0.68 (0.11)</td>
</tr>
<tr>
<td>Red clover (RC)</td>
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<td>21.24 (1.93)</td>
<td>27.47 (2.52)abc</td>
<td>11.78 (0.80)</td>
<td>1.90 (0.35)abc</td>
<td>3.40 (0.45)ab</td>
<td>0.67 (0.11)</td>
</tr>
<tr>
<td>Hairy vetch (HV)</td>
<td>87.33 (6.31)</td>
<td>20.98 (1.56)</td>
<td>27.73 (1.93)abc</td>
<td>11.54 (0.68)</td>
<td>2.06 (0.35)abc</td>
<td>3.25 (0.32)ab</td>
<td>0.86 (0.14)</td>
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<tr>
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<td>84.98 (5.74)</td>
<td>22.17 (1.56)</td>
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<td>12.04 (0.62)</td>
<td>1.69 (0.37)abc</td>
<td>3.22 (0.28)ab</td>
<td>0.80 (0.22)</td>
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<tr>
<td>Oat (OA)</td>
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<td>22.84 (1.52)</td>
<td>27.37 (1.71)abc</td>
<td>12.60 (0.73)</td>
<td>1.28 (0.20)c</td>
<td>3.66 (0.33)ab</td>
<td>0.77 (0.13)</td>
</tr>
<tr>
<td>Canola (CA)</td>
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<td>22.96 (1.52)</td>
<td>28.23 (1.74)abc</td>
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<td>3.38 (0.29)ab</td>
<td>0.79 (0.13)</td>
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<tr>
<td>Cereal rye (CR)</td>
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<td>21.73 (1.18)</td>
<td>27.27 (1.36)abc</td>
<td>12.16 (0.56)</td>
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<td>3.64 (0.29)ab</td>
<td>0.82 (0.14)</td>
</tr>
<tr>
<td>FR + OA + CA + CR</td>
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<td>12.11 (0.54)</td>
<td>1.40 (0.16)abc</td>
<td>3.48 (0.26)ab</td>
<td>0.75 (0.11)</td>
</tr>
<tr>
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<td>27.83 (1.63)abc</td>
<td>12.57 (0.70)</td>
<td>1.91 (0.29)abc</td>
<td>3.48 (0.29)ab</td>
<td>0.78 (0.10)</td>
</tr>
<tr>
<td>RC + HV + CA + CR</td>
<td>87.97 (6.59)</td>
<td>20.66 (1.47)</td>
<td>28.38 (2.03)abc</td>
<td>11.55 (0.75)</td>
<td>2.41 (0.43)abc</td>
<td>3.54 (0.36)ab</td>
<td>0.77 (0.11)</td>
</tr>
<tr>
<td>SH + SB + CA + CR</td>
<td>86.26 (6.15)</td>
<td>21.54 (1.56)</td>
<td>27.19 (1.77)abc</td>
<td>11.98 (0.72)</td>
<td>1.52 (0.22)abc</td>
<td>3.56 (0.36)ab</td>
<td>0.74 (0.13)</td>
</tr>
<tr>
<td>RC + HV + FR + OA</td>
<td>89.26 (7.22)</td>
<td>21.76 (1.56)</td>
<td>28.56 (2.40)abc</td>
<td>11.99 (0.61)</td>
<td>1.95 (0.42)abc</td>
<td>3.68 (0.39)ab</td>
<td>0.90 (0.19)</td>
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<tr>
<td>8 species mix</td>
<td>94.47 (7.61)</td>
<td>22.95 (1.63)</td>
<td>30.20 (2.39)abc</td>
<td>12.67 (0.75)</td>
<td>2.11 (0.43)abc</td>
<td>3.92 (0.44)ab</td>
<td>0.94 (0.19)</td>
</tr>
</tbody>
</table>

Notes: Gram + = Gram positive. Gram – = Gram negative. AM = arbuscular mycorrhizal. Est. = estimate. Values within a column with different letters were significantly different due to year or cover crop based on Tukey’s honestly significant difference ($p < 0.05$). The absence of letters in a column indicates that the effect of year or cover crop was not significant.
also contributed to the distinction of microbial communities among treatments in both fall and spring (figures 1a and 1b). Previous studies have similarly reported increases in protozoan abundance associated with cover cropping (Schutter et al. 2001; Carrera et al. 2007). Protozoans are bacterial grazers, therefore their increased abundance under cover crops may be due to the increased microbial biomass generally associated with cover cropped treatments in this study.

We also assessed microbial community composition using the ratio of fungal to bacterial PLFA biomarkers (Joergensen and Wichern 2008). In both fall and spring, the fungal:bacterial ratio averaged across treatments was higher in Year 1 than in Year 2 (table 5). Fungal:bacterial ratios did not vary among treatments in fall; however, cover crops led to an increase of 0.01 in the index value compared to the no cover crop control (p = 0.01; table 5). We observed the same increase in fungal:bacterial ratio from cover cropped treatments in spring (p = 0.03; table 5). Additionally, in spring, the fungal:bacterial ratio was higher under the four species cover crop mixture that included hairy vetch (Vicia villosa L.), red clover (Trifolium pratense L.), cereal rye, and canola (Brassica napus L.) than the no cover crop control (table 5). The db-RDA had shown that most cover cropped treatments were associated with higher proportions of fungi compared to the no cover crop control (figures 1a and 1b); therefore, we expected that at least some cover cropped treatments would exhibit a higher fungal:bacterial ratio than the control. One explanation for the fact that compositional changes indicated by db-RDA did not manifest as differences in the fungal:bacterial ratio among treatments is that cover crops were associated not only with increased proportions of fungi, but also Gram-negative bacteria, which could offset increases in the overall ratio. Nonetheless, evidence of an increase in the relative abundance of fungi associated with cover crops provided by the db-RDA and the increase in fungal:bacterial ratio observed in cover cropped treatments relative to the control suggest that cover crops can lead to greater fungal dominance, which in turn may promote SOM stabilization and enhance soil health (Six et al. 1999; Lehman et al. 2015b). As a whole, our analyses of microbial community composition clearly demonstrate that cover crops influenced the structure of microbial communities. Differences in the absolute and relative abundances of distinct microbial groups observed among treatments beg the question, “Do different cover crops influence different microbial groups?”—a question we addressed with a second redundancy analysis.

Cover Crop Specific Influences on Soil Microbial Communities. Differences in microbial community composition among treatments suggest that there are species-specific relationships between cover crops and microbial communities. The second db-RDA demonstrated that the relative abundance of specific microbial groups within a season was related to quantity of biomass produced by specific cover crops and arable weeds (figures 2a and 2b), supporting this hypothesis. In this context, aboveground biomass of a particular species reflects the abundance of particular plant traits that may influence microbial abundance (e.g., root biomass, surface area, and exudates) and mediate the species-specific associations we observed. In both fall and spring, microbial communities were distinguished from one another based on whether they were dominated by Gram-positive bacteria and actinomycetes or by protozoa, AM-fungi, non-AM fungi, and Gram-negative bacteria (fall: p < 0.01, spring: p < 0.01; figures 2a and 2b, axis 1 in both panels). Cover crops contributing to high proportions of the latter groups in the fall were oat and forage radish (Raphanus sativus L.) (figure 2a). In contrast, red clover, hairy vetch, and sunn hemp cover crops were associated with Gram-positive bacteria and actinomycetes during the fall growing season (figure 2a). This differentiation may have
been due to lower biomass production in the fall by the legume cover crops compared to oat and forage radish (Finney et al. 2016) or unique influences of legume versus nonlegume cover crops on microbial communities. By spring, cover crops generally associated with higher proportions of AM fungi, protozoa, Gram-negative bacteria, and non-AM fungi were cereal rye, red clover, and hairy vetch (figure 2b). The winter-killed cover crops, namely soybean, sunn hemp, and forage radish, were associated with higher proportions of actinomycetes and Gram-positive bacteria (figure 2b). Again differences in biomass production by winter-killed versus winter-hardy cover crop species (Finney et al. 2016) may have led to this result in spring. In other words, cover crops that provided C inputs to the microbial community in both fall and spring led to higher proportions of AM fungi, protozoa, Gram-negative bacteria, and non-AM fungi in the soil microbial community by spring. By this logic, canola would have also been expected to promote increased proportions of these microbial groups, particularly in the spring, but, in fact, canola did not contribute to the differentiation of microbial communities along the first db-RDA axis in either season (figures 2a and 2b). One contributing factor to this result is the fact that AM fungi are obligate plant symbionts (Brundrett 2002) and members of the Brassicaceae family such as canola and forage radish are not AM fungal hosts (Vierheilig et al. 2000). Previous research on standing crops and oilseed meals derived from glucosinolate-containing brassicas have indicated that members of this family can also alter bacterial community structure (Rumberger and Marschner 2003; Hollister et al. 2013), and a recent study comparing PLFA profiles of oat and rape (Brassica napus L.) cover crops found that rape was associated with larger populations of Gram-positive bacteria (Mackie et al. 2014). One hypothesis to explain the lack of a canola effect on microbial community composition in our study is that any increases in microbial groups generally associated with cover crops (i.e., Gram-negative bacteria, non-AM fungi, and protozoa, but not AM fungi) were offset by increases in Gram-positive bacteria specific to this species.

There were several cover crop–microbial group associations indicated by the second db-RDA (figures 2a and 2b) and supported by ANOVA results (tables 3 and 4). In fall, increasing oat biomass was associated with increasing proportions of AM fungi (figure 2a). Concentrations of the AM fungal biomarker were also significantly higher in the oat monoculture (3.26 ± 0.14 nmol g⁻¹) than the no cover crop control (2.71 ± 0.08 nmol g⁻¹) and tended to be higher in the oat monoculture than in other cover crop treatments (table 3). This finding supports previous work by Lehman et al. (2012) demonstrating that oat and oat-containing cover crop mixtures increased AM fungi in the fall. The current study also provided a means to evaluate the longevity of this effect by sampling in the spring. Similar to the results of the db-RDA on treatment-level effects, the second axis of the db-RDA on species-specific effects was significant in spring and differentiated microbial communities based on the relative proportions of AM to non-AM fungi (p = 0.01; figure 2b). Two cover crops were associated with higher proportions of AM fungi—oat and cereal rye (figure 2b). Oat and cereal rye monocultures also had higher concentrations of the AM fungal biomarker in spring (3.66 ± 0.33 and 3.64 ± 0.29 nmol g⁻¹, respectively) compared to the no cover crop control (2.96 ± 0.33 nmol g⁻¹), as did two cover crop mixtures, a four species mixture combining oats with forage radish, hairy vetch, and red clover (3.68 ± 0.39 nmol g⁻¹), and the eight species mixture (3.92 ± 0.44 nmol g⁻¹) that contained both oat and cereal rye (table 4). These results provide evidence that the positive effect of oat on AM fungi in the fall is not ephemeral, but persists into the spring. There is also evidence that the oat benefit, quantified as the density of mycorrhizal fungi, carries over into the subsequent crop in no-till sweet corn production (Kabir and Koide 2002). The same study also found that the density of mycorrhizal hyphae sampled in corn was higher following not only oat monocultures, but also cereal rye grown in monoculture or as an oat–cereal rye biculture compared to a fallow control (Kabir and Koide 2002). A similar positive effect on mycorrhizal colonization of corn roots following a cereal rye cover crop has also been reported (White and Weil 2010), as has a positive effect of oats on AM-fungi propagules enumerated approximately three months after cover crop planting (Lehman et
al. 2012). The fact that similar effects of oats and rye on AM fungal abundance have been detected using both microscopic (Kabir and Koide 2002; Lehman et al. 2012; White and Weil 2010) and biochemical (Lehman et al. 2012 and this study) quantification provides strong evidence that use of these cover crops is an effective strategy to increase AM fungi in agricultural systems. Unlike previous studies, however, we did not find that cover crop mixtures outperformed monocultures in the promotion of AM fungi (Kabir and Koide 2002; Lehman et al. 2012), though by spring, concentrations of the AM fungal biomarker tended to be highest in the most diverse (eight-species) cover crop mixture (table 4).

There was also an association between non-AM fungi and hairy vetch in the spring indicated by both db-RDA (figure 2b) and ANOVA (table 4). By spring, concentrations of the non-AM fungal biomarker were higher in the hairy vetch monoculture (2.60 ± 0.35 nmol g⁻¹) than the no cover crop control (1.42 ± 0.31 nmol g⁻¹), oat monoculture (1.28 ± 0.20 nmol g⁻¹), and two mixtures. It is important to recognize that interpretation of PLFA biomarkers, particularly those used to identify AM and non-AM fungi, is challenged by the fact that biomarkers may be present in several microbial and nonmicrobial groups (Frostegård et al. 2011). Recent work by Sharma and Buyer (2015) has shown that among three different lipid analyses (PLFA analysis [used in this study], neutral lipid fatty acid analysis [NLFA], and ester-linked fatty acid analysis [ELFA]), the PLFA biomarker did not correlate with two microscopic methods of quantifying AM-fungi in soil (spore counts) and plant roots (percentage colonization). Therefore, if the aim of a study is to estimate AM fungal spores or root colonization, NLFA or ELFA is advised. The PLFA biomarker for AM fungi used in this study has been found to indicate a viable fungal hyphal density and is a relevant indicator of abundance. The caveat associated with this biomarker, however, is that it is also found in Gram-negative bacteria and may overestimate AM fungal abundance (Buyer et al. 2010; Olson 1999). The fact that Gram-negative and AM-fungi responded similarly to cover crop treatments and biomass in our study may reflect this overlap. In spite of this potential limitation of the PLFA biomarker, the species-specific associations we identified between AM fungi and oats and cereal rye are consistent with previous studies as noted earlier. The PLFA biomarker used to identify non-AM fungi is also present in plants; therefore, fine roots could contribute to the measured concentration of non-AM fungi. In a study in which it was found that the fungal PLFA biomarker increased in the presence of hairy vetch shoots in a tomato (Solanum lycopersicum L.) cropping system, Buyer et al. (2010) also used PLFA on cover crop plant tissues to demonstrate that plant roots contributed little to measured concentrations of the fungal biomarker in soil. This gives us confidence that the results we observed do indicate an association between hairy vetch and non-AM fungi. Comparison of non-AM fungi concentrations among treatments further indicates that this association was present not only in hairy vetch monocultures, but also in mixtures containing hairy vetch (table 4).

Based on our analyses, we conclude that cover crops generally serve to increase proportions of AM fungi, non-AM fungi, Gram-negative bacteria, and protozoa in the soil microbial community, but specific cover crops are associated with increases in specific microbial groups. The most notable associations are those of AM fungi with oats and with cereal rye and the association between non-AM fungi and hairy vetch. These cover crop-specific associations led to microbial communities that could be distinguished from one another based on their proportions.
of AM and non-AM fungi by spring (figures 1b and 2b). Therefore, while cover crops appear to have a generally positive effect on fungal communities, cover crop species selection is a management strategy that can be used to increase targeted fungal groups.

The results of this study also raise a compelling question regarding the effects of multispecies cover crop mixtures on microbial diversity. If specific species promote distinct microbial communities, it would follow that microbial communities associated with cover crop mixtures would retain characteristics of each of the component species and, therefore, demonstrate greater diversity (Qiao et al. 2012; Zak et al. 2003). This idea is supported in our results by the fact that species-specific associations were manifest not only in monocultures, but also multispecies cover crop mixtures. For example, the location of cover crop mixtures combining hairy vetch and cereal rye along the second db-RDA axis in spring (figure 1b) was intermediate between hairy vetch and cereal rye monocultures, suggesting that the mixtures retained the fungal group associations specific to each of these species. Due to the limitations of PLFA (Frostégård et al. 2011), the question of microbial community diversity associated with cover crop mixtures will need to be addressed with molecular analytical tools; however, this study offers preliminary evidence that further investigation of the linkages between above and belowground diversity in cover cropping systems is warranted and relevant to improving soil health (Lehman et al. 2015a).

Soil Biological Activity. Soil biological activity, indicated by the daily respiration rate during a seven day incubation, differed among treatments in both fall and spring (table 6). In the fall, grass monocultures (oat and cereal rye) increased respiration by 50% relative to the control, and several mixtures also led to increases in activity ranging from 33% to 52%. In spring, only three cover crop treatments exhibited increased respiration compared to the control: hairy vetch (17.06 ± 2.13 µg CO₂-C g⁻¹ soil d⁻¹) and cereal rye (16.99 ± 2.08 µg CO₂-C g⁻¹ soil d⁻¹) monocultures and the four species mixture that contained both of these species (17.80 ± 2.74 µg CO₂-C g⁻¹ soil d⁻¹; table 6), with an average increase of 36%.

While there was a positive correlation between aboveground plant biomass and daily respiration (figure 3a; r = 0.39, p < 0.01), there was a stronger correlation between soil microbial biomass, indicated by total PLFA concentration, and biological activity (figure 3b; r = 0.50, p < 0.01). These results suggest that cover crops influence soil C utilization by providing labile C inputs to fuel respiration and the accumulation of soil microbial biomass, which, in turn, leads to increased respiration. There was, however, an even stronger positive correlation between respiration and fungal:bacterial ratio (figure 3c; r = 0.82, p < 0.01), an indicator of microbial community composition. This relationship was reflected in the fact that the cover crop treatment with the highest respiration rate in the spring, a four species mixture of hairy vetch, red clover, canola, and cereal rye, also exhibited the highest fungal:bacterial ratio. Other treatments exhibiting high daily respiration rates in spring, hairy vetch and cereal rye monocultures, also tended to have higher fungal:bacterial ratios than the control. An important finding of this study, therefore, is that the influence of cover crops on C cycling is not confined to simply the quantity of C contributed to the soil, but also depends on species-specific plant-microbial associations like those we identified.

Arable Weeds and Microbial Community Composition. In addition to providing information on species-specific cover crop influences on microbial community composition, this study also advances our knowledge of the effects of arable weed communities on soil biology. In both fall and spring, increasing arable weed biomass was associated with higher proportions of Gram-positive bacteria and actinomycetes (figures 2a and 2b), a result consistent with the treatment-level db-RDA (figures 1a and 1b). While the positive correlation between microbial biomass and plant productivity observed in this study would suggest that the arable weed community could serve a purpose similar to cover crops (i.e., providing sufficient C via exudates and fine root turnover to support microbial growth and activity), arable weeds and cover crops functioned differently with regard to the microbial groups with which they associated. The correlation between soil biological activity and greater fungal dominance indicates that these associations are, in fact, consequential to microbial community function and offers a mechanistic explanation for lower rates of activity observed in control treatments versus many of the cover cropped treatments in both fall and spring (table 6).

Summary and Conclusions
An important and relevant justification for the use of cover crops to improve soil health is that they provide resources to nourish soil microbial communities during periods when cash crops are not present. Arguably, arable weed communities that emerge in fallowed fields could perform this same function. This research demonstrates that cover crops perform an important function beyond simply providing resources to increase microbial biomass. Our work has shown that specific cover crops promote the abundance of specific groups of microorganisms and, therefore, can be used to direct microbial community composition toward assemblages that promote soil health. For example, rebuilding fungal populations that are typically depauperate in agricultural systems can promote SOM accumulation and enhance aggregate stability. The results of this study show that cover crops generally lead to increased concentrations of soil fungi compared to arable weeds, with the greatest gains associated with oats, hairy vetch, and cereal rye. Notably, our results also offer evidence that different fungal communities respond to different cover crop species. While cereal rye and oats were associated with increases in AM fungi, hairy vetch, and to a lesser extent red clover, led to increased abundance of non-AM fungi. Knowledge of these species-specific cover crop influences will enable land managers to design cover cropping systems that can efficiently and effectively manipulate soil biology to enhance soil health.

Acknowledgements
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Table 6
Main effects of year and cover crop treatment on least square means and standard errors (standard errors are values in parentheses) of daily respiration rate (μg CO$_2$-C g$^{-1}$ soil d$^{-1}$) during a seven day incubation of fresh soil collected in fall and spring, approximately two and nine months, respectively, following cover crop planting in August of 2011 (Year 1) and August of 2012 (Year 2) in central Pennsylvania.

<table>
<thead>
<tr>
<th>Year/crop</th>
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</tr>
</thead>
<tbody>
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<td></td>
<td></td>
</tr>
<tr>
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<td>17.97 (0.64)a</td>
</tr>
<tr>
<td>Sunn hemp (SH)</td>
<td>8.06 (1.15)d</td>
<td>11.14 (3.39)cd</td>
</tr>
<tr>
<td>Soybean (SB)</td>
<td>8.59 (1.37)cd</td>
<td>9.98 (2.05)e</td>
</tr>
<tr>
<td>Red clover (RC)</td>
<td>9.22 (1.31)abcd</td>
<td>9.44 (1.77)e</td>
</tr>
<tr>
<td>Hairy vetch (HV)</td>
<td>8.67 (0.98)cd</td>
<td>11.50 (1.63)cd</td>
</tr>
<tr>
<td>Forage radish (FR)</td>
<td>9.73 (1.33)abcd</td>
<td>17.06 (2.13)ab</td>
</tr>
<tr>
<td>Oat (OA)</td>
<td>10.21 (0.79)abcd</td>
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</tr>
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<td>Canola (CA)</td>
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<td>12.67 (2.77)abcd</td>
</tr>
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<td>Cereal rye (CR)</td>
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<td>11.23 (1.83)cd</td>
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<td>11.31 (2.38)cd</td>
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<tr>
<td>SH + SB + CA + CR</td>
<td>11.89 (1.50)ab</td>
<td>17.80 (2.74)a</td>
</tr>
<tr>
<td>RC + HV + FR + OA</td>
<td>10.12 (1.08)abcd</td>
<td>14.00 (1.84)abcd</td>
</tr>
<tr>
<td>8 species mix</td>
<td>10.73 (1.37)abcd</td>
<td>14.56 (2.65)abcd</td>
</tr>
<tr>
<td>8 species mix</td>
<td>11.61 (1.32)bcd</td>
<td>15.92 (3.06)abc</td>
</tr>
</tbody>
</table>

Notes: Values within a column with different letters were significantly different due to year or cover crop based on Tukey’s HSD (< 0.05). The absence of letters in a column indicates that the effect of year or cover crop was not significant.

Figure 3
Correlation of soil biological activity to (a) plant biomass, (b) microbial community size, and (c) microbial community composition in soils collected two and nine months after cover crop planting in 2011 and 2012 in central Pennsylvania. Soil biological activity was measured as the daily respiration in a seven day laboratory incubation of fresh soil. (a) Plant biomass is the sum of aboveground cover crop and weed biomass produced in the season (fall or spring) in which soil was collected. (b) Total phospholipid fatty acid (PLFA) concentration represents the size of the microbial community, and (c) the fungal:bacterial ratio of the microbial community represents community composition.
structure in tomato cropping systems. Soil Biology and Biochemistry 42(5):381-41.


