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Abstract

We describe a novel approach for identifying microbial contributions to soil C-cycling dynamics using nucleic acid stable isotope probing coupled with next generation sequencing (SIP-NGS). In a series of parallel soil microcosms we amended soils with a complex mixture of model carbon (C) substrates and inorganic nutrients common to plant biomass, where a single C constituent is substituted for its ¹³C-labeled equivalent. Using this approach we assessed incorportation of ¹³C-xylose or ¹³C-cellulose as proxies for labile soluble C and polymeric insoluble C utilization, respectively. Using CsCl gradient fractionation, incorporation of ${}^{13}\mathrm{C}$ into DNA was measured over 30 days. The 16S rRNA gene sequences from CsCl gradient fractions were characterized by 454 pyrosequencing and classified into Operational Taxonomic Units (OTU). We describe specific patterns of C-assimilation by discrete OTUs as a function of substrate, time, and level of isotope incorporation. Incorporation of ¹³C from xylose into OTUs was observed at days 1, 3, and 7, while notable incorporation of ¹³C from cellulose was observed only after day 14. Of over 6,000 OTUs detected, a total of 43 and 35 unique OTUs significantly assimilated ¹³C from xylose and cellulose, respectively. We did not observe consistent C utilization at the phylum level although both xylose and cellulose utilization were observed across 7 phyla each revealing a high diversity of bacteria able to utilize these substrates. OTUs that assimilate xylose and those that assimilate cellulose are largely mutually exclusive. Xylose assimilating OTUs are more abundant in the microbial community than cellulose assimilating OTUs, while cellulose OTUs demonstrate a greater substrate specificity than xylose OTUs. Furthermore, the increased depth provided by SIP-NGS allowed us to identify several novel cellulose utilizing bacte-

monolayer | structure | x-ray reflectivity | molecular electronics

Abbreviations: SAM, self-assembled monolayer; OTS, octadecyltrichlorosilane

Introduction

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When have only a rudimentary understanding of carbon flow through soil microbial communities. This deficiency is driven by the staggering complexity of soil microbial food webs and the opacity of these biological systems to current methods for describing microbial metabolism in the environment. Relating community composition to overall soil processes, such as nitrification and denitrification, which are mediated by defined functional groups has been a useful approach. However, carbon-cycling processes have proven more recalcitrant to study due to the wide range of organisms participating in these reactions and our inability to discern diagnostic functional genetic markers.

Excluding plant biomass, there are 2,300 Pg of carbon (C) stored in soils worldwide which accounts for \sim 80% of the global terrestrial C pool BATJES, 1996; Amundson, 2001. When organic C from plants reaches soil it is degraded by fungi, archaea, and bacteria. This C is rapidly returned to the atmosphere as CO_2 or remains in the soil as humic substances that can persist up to 2000 years Yanagita, 1990. The majority of plant biomass C in soil is respired and produces 10 times more CO_2 than anthropogenic emissions on an annual basis Chapin, 2002. Global changes in

atmospheric CO₂, temperature, and ecosystem nitrogen inputs, are expected to impact primary production and C inputs to soils Groenigen *et al.*, 2006 but it remains difficult to predict the response of soil processes to anthropogenic change DAVIDSON *et al.*, 2006. Current climate change models concur on atmospheric and ocean C predictions but not terrestrial Friedlingstein *et al.*, 2006. These contrasting terrestrial ecosystem model predictions reflect how little is known about soil C cycling dynamics and it has been suggested that incosistencies in terrestial modeling could be improved by elucidating the relationship between dissolved organic carbon and microbial communities in soils Neff and Asner, 2001.

An estimated 80-90% of C cycling in soil is mediated by microorganisms Nannipieri et al., 2003a; n.d. Understanding microbial processing of nutrients in soils presents a special challenge due to the hetergeneous nature of soil ecosystems and methods limitations. Soils are biologically, chemically, and physically complex which affects microbial community composition, diversity, and structure Nannipieri et al., 2003a. Confounding factors such as physical protection/aggregation, moisture content, pH, temperature, frequency and type of land disturbance, soil history, mineralogy, N quality and availability, and litter quality have all been shown to affect the ability of the soil microbial community to access and metabolize C substrates Sollins et al., 1996; Kalbitz et al., 2000. Further, rates of metabolism are often measured without knowing the identity of the microbial species involved Nannipieri et al., 2003b leaving the importance of community membership towards maintaining ecosystem functions unknown Nannipieri et al., 2003b; Allison and Martiny, 2008; Schimel and Schaeffer, 2012. Litter bag experiments have shown that the community composition of soils can have quantitative and qualitative impacts on the breakdown of plant materials Schimel, 1995. Reciprocal exchange of litter type and microbial inocula under controlled environmental conditions reveals that differences in community composition can account for 85% of the variation in litter carbon mineralization Strickland et al., 2009. In addition, assembled communities of cellulose degraders reveal that the composition of the community has significant impacts on the rate of cellulose degradation Wohl et al., 2004.

An important step in understanding soil C cycling dynamics is to identify individual contributions of discrete microorganisms and to investigate the relationship between genetic diversity, community structure, and function O'Donnell *et al.*, 2002. The vast majority of microorganisms continue to resist cultivation in the laboratory, and even when cultivation is achieved, the traits expressed by

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a microorganism in culture may not be representative of those expressed when in its natural habitat. Stable-isotope probing (SIP) provides a unique opportunity to link microbial identity to activity and has been utilized to expand our knowledge of a myriad of important biogeochemical processes Chen and Murrell, 2010. The most successful applications of this technique have identified organisms which mediate processes performed by a narrow set of functional guilds such as methanogens Lu, 2005. The technique has been less applicable to the study of soil C cycling because of limitations in resolving power as a result of simultaneous labeling of many different organisms in the community. Additionally, molecular applications such as TRFLP, DGGE, and cloning that are frequently used in conjunction with SIP provide insufficient resolution of taxon identity and depth of coverage. We have developed an approach that employs a complex mixture of substrates added to soil at a low concentration relative to soil organic matter pools along with massively parallel DNA sequencing. This greatly expands the ability of nucleic acid SIP to explore complex patterns of C-cycling in microbial communities with increased resolution.

A temporal cascade occurs in natural microbial communities during the plant biomass degradation in which labile C degradation preceeds polymeric C Hu and Bruggen, 1997; Rui et al., 2009. The aim of this study is to track the temporal dynamics of C assimilation through discrete individuals of the soil microbial community to provide greater insight into soil C-cycling. Our experimental approach employs the addition of a soil organic matter (SOM) simulant (a complex mixture of model carbon sources and inorganic nutrients common to plant biomass), where a single C constituent is substituted for its ¹³C-labeled equivalent, to soil. Parallel incubations of soils amended with this complex C mixture allows us to test how different C substrates cascade through discrete taxa within the soil microbial community. In this study we use ¹³C-xylose and ¹³Ccellulose as a proxy for labile and polymeric C, respectively. Using a novel approach we couple nucleic acid stable isotope probing with next generation sequencing (SIP-NGS) to elucidating soil microbial community members responsible for specific C transformations. Amplicon sequencing of 16S rRNA gene fragments from many gradient fractions and multiple gradients make it possible to track C assimilation by hundreds of different taxa. Ultimately we identify discrete microorganisms responsible for the cycling of specific C substrates.

Results and Discussion

In this study, we couple nucleic acid SIP with next generation sequencing (SIP-NGS) to observe C use dynamics by the soil microbial community. A series of parallel soil microcosms all amended with a C substrate mixture were incubated for 30 days. The substrate mixture was identical for each bottle except in one series of bottles the cellulose was $^{13}\mathrm{C}$ -labeled in another the xylose was $^{13}\mathrm{C}$ -labeld and in the last no sustrattes were labeled. The C substrate mixture was designed to approximate freshly degrading plant biomass. Xylose or cellulose carried the isotopic label so we could examine C assimilation dynamics for labile, soluble C versus insoluble, polymeric C. 5.3 mg total mass of C substrate mixture per gram soil (including 0.42 mg xylose-C and 0.88 mg cellulose-C g soil⁻¹) was added to each microcosm representing 18% of the total soil C. Microcosms were harvested at several time points during the incubation period and ¹³C assimilation was observed by sequencing 16S rRNA gene amplicons from bulk soil DNA and CsCl gradient fractions. Assimilation of ¹³-C from Xylose degradation peaked immediately, while cellulose ¹³C assimilation peaked after two weeks of incubation (Figure 1).

 $^{13}\mathrm{C}$ from cellulose assimilated by canonical cellulose-degrading and uncharacterized microbial lineages in many

phyla including Chloroflexi and Verrucomicrobia. Only 2 and 5 OTUs were found to have incorporated \$^{13}\$C from labeled cellulose at days 3 and 7, respectively. At days 14 and 30, however, 42 and 39 OTUs were found to incorporate $^{13}\mathrm{C}$ from cellulose into biomass. An average 16% of the ¹³C-cellulose added was respired within the first 7 days, 38% by day 14, and 60% by day 30. A Cel-Ivibrio and Sandaracinaceae OTU assimilated 13C from cellulose at day 3. Day 7 responders included the same Cellvibrio responder as day 3, a Verrucomicrobia OTU and three Chloroflexi OTUs. 50% of Day 14 responders belong to Proteobacteria (66% Alpha-, 19% Gamma-, and 14% Beta-) followed by 17% Planctomycetes, 14% Verrucomicrobia, 10% Chloroflexi, 7% Actinobacteria and 2% Cyanobacteria. Bacteroidetes OTUs begin to incoporate 13C from cellulose at day 30 (13% of day 30 responders). Other day 30 responding phyla include Proteobacteria (30% of day 30 responders; 42% Alpha-, 42% Delta, 8% Gamma-, and 8% Beta-), Planctomycetes (20%), Verrucomicrobia (20%), Chloroflexi (13%) and Cyanobacteria (3%). Proteobacteria, Verrucomicrobia, and Chloroflexi had relatively high numbers of responders with heavy response across multiple time points (Figure 2).

Proteobacteria represent 46% of all cellulose responding OTUs identified. Cellvibrio accounted for 3% of all Proteobacterial responding OTUs detected. Cellvibrio was one of the first identified cellulose degrading bacteria and was originally described by Winogradsky in 1929 who named it for its cellulose degrading abilities (Boone, 2001). All ¹³C-cellulose responding Proteobacteria share high sequence identity with 16S rRNA genes from sequenced type straints (Table XX) except for OTU.442 (best type strain match 92% sequence identity in the Chrondomyces genus) and OTU.663 (best type strain match outside Proteobacteria entirely, Clostridium genus, 89% sequence identity). Some Proteobacteria responders share high sequence identity with type strains for genera known to posess cellulose degraders including Rhizobium, Devosia, Stenotrophomonas and Cellvibrio. One Proteobacteria OTU shares high sequence identity with the Brevundimonas type strain. Brevundimonas has not previously been identified as a cellulose degrader, but has been shown to degrade cellouronic acid, an oxidized form of cellulose (Tavernier et al., 2008).

Verrucomicrobia, a cosmopolitan soil phylum often found in high abundance (Fierer et al., 2013), are implicated in polysaccaride degradation in many environments (Fierer et al., 2013; Herlemann et al., 2013; Chin et al., n.d.). Verrucomicrobia comprise 16% of the total cellulose responder OTUs detected. 40% of Verrucomicrobia responders belong to the uncultured FukuN18 family originally identified in freshwater lakes (Parveen et al., 2013). The Verrucomicrobia OTU with the strongest Verrucomicrobial response to ¹³C-cellulose shared high sequence identity (97%) with an isolate from Norway tundra soil (Jiang et al., 2011) although growth on cellulose was not assessed for this isolate. Only one other $^{13}\mathrm{C}\text{-cellulose}$ responding verrucomicrobium shared high DNA sequence identity with a sequenced type strain, OTU.638 with Roseimicrobium gellanilyticum (100% sequence identity) and Roseimicrobium gellanilyticum grows on soluble celluose (Otsuka et al., 2012). The remaining 13C-cellulose Verrucomicrobia responders did not share high sequence identity with any type strains (maximum sequence identity with any type strain 93%).

Chloroflexi are traditionally known for their metabolically dynamic lifestyles ranging from anoxygenic phototropy to organohalide respiration (Hug *et al.*, 2013). Recent studies have focused on *Chloroflexi* roles in C cycling (Goldfarb *et al.*, 2011; Cole *et al.*, 2013; Hug *et al.*, 2013) and several members of this phylum demonstrated cellulose utilization (Goldfarb *et al.*, 2011; Cole *et al.*, 2013; Hug *et al.*, 2013). Four closely related OTUs in an undescribed *Chloroflexi* lineage (closest matching type strain for all four OTUs: *Herpetosiphon geysericola*, 89% sequence identity) responded to ¹³C-cellulose in this microcosm experiment. One additional OTU also from a poorly characterized lineage (clos-

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est type strain match a proteobacterium at 78% sequence identity) responded to 13 C-cellulose (Figure 3).

Other notable 13C cellulose responders include a Bacteroidetes OTU that shares high sequence identity (99%) to Sporocytophaga myxococcoides a known cellulose degrader (Vance et al., 1980), and three Actinobacteria OTUs that share high sequence identity (100%) with sequenced type strains. One of the three *Actinobacteria* ¹³C-cellulose responders is in the *Strepto*myces, a genus known to possess cellulose degraders, while the other two closely match the type strains Allokutzneriz albata (Tomita et al., 1993; Labeda and Kroppenstedt, 2008) and Lentzea waywayandensis (LABEDA and LYONS, 1989; Labeda et al., 2001) that do not decompose cellulose in culture. Nine Plantomycetes OTUs responded to ¹³C-cellulose but none are within described genera (closest type strain match 91% sequence identity) (Figure 3). Interestingly, one responder is annotated as belonging in the Cyanobacteria. The phylum annotation is misleading as the OTU is not closely related to any oxygenic phototrophs (closest type strain match Vampirovibrio chlorellavorus, 95% sequence identity). A sister clade to the oxygenic phototrophs that does not itself possess known phototrophs has recently been proposed to constitute its own phylum ("Melainabacteria", (Rienzi et al., 2013)) although its phylogenetic position is debated (Soo et al., 2014). The catalog of metabolic capabilities associated with Cyanobacteria (or candidate phyla previously annotated as Cyanobacteria) are quickly expanding (Rienzi et al., 2013; Soo et al., 2014). Our findings provide evidence of cellulose degradation within a lineage closely related to but apart from oxygenic phototrophs. Notably, polysaccharide degradation is suggested by the analysis of a Melainabacterial genome (Rienzi et al., 2013). Although we highlight $^{13}\mathrm{C}\text{-cellulose}$ responders that share high sequence identity with described genera, by and large $^{13}\mathrm{C}\text{-cellulose}$ responders uncovered in this experiment are not closely related to isolates (Table XX).

Putative spore-formers in the Firmicutes assimilate ¹³C from xylose within first day after soil amendment. Within the first 7 days of incubation an average 63% of ¹³C-xylose was respired and only an additional 6% more was respired between days 7 and 30 (Table S1). At the end of the 30 day experiment 30% of the original ¹³C from xylose remained in the soils. The ¹³C remaining in the soil from ¹³C-xylose addition has likely been stabilized by assimilation into microbial biomass and/or microbial conversion into other forms of organic matter, though it is possible that some ¹³C-xylose remains unavailable to microbes due to abiotic interactions in soil (Kalbitz *et al.*, 2000). Of the 60 total xylose responders 53 were responsive within the first 7 days and only 7 responders detected for days 14 and 30 (Table S1).

At day 1 (d1), 57% of xylose responsive OTUs belong to Firmicutes (Paenibacillaceae, Planococcaceae, and Bacillaceae) and the remaining 43% of responders were comprised of 19% Bacteroidetes (Flavobacteriaceae), 14% Proteobacteria (Enterobacteriaceae, Comamonadaceae, and uncultured Gammaproteobacteria), and 10% Actinobacteria (Micrococcaceae and Microbacteriaceae) (Fig. 2). Of the xylose responders detected at day 3 (d3), Firmicutes responders decreased to 8% (from 12 OTUs to 1) and an increase in Bacteroidetes (67%) and Proteobacteria (25%) from d1. Cellvibrio, a canonical cellulose degrader, was notably one of the three Proteobacterial xylose responders at d3 (discussed later). Day 7 (d7) responders were 50% Actinobacteria, 35% Proteobacteria, and 5% of each Bacteroidetes, Firmicutes, and Planctomycetes. A substantial amount (75%) of xylose responders for day 7 had not previously been identified as responders at earlier time points which attests to the wide number of taxa able to use xylose. Each of the 10 Actinobacteria responders at day 7 belonged

to a different family making it the phylum with the most wide-spread use of xylose. However, it should be noted that they were confined within two Actinobacterial Orders; Frankiales and Micrococcales.

We observe dynamic changes in ¹³C-xylose assimilation with time; dominant xylose responders shift from Firmicutes (d1) to Bacteroidetes (d3) then finally Actinobacteria (d7). At any given time soils harbor microorganisms at varying degrees of dormancy depending on nutrient availability (Jones and Lennon, 2010). The sudden addition of our complex C mixture would most certainly prompt dormant and non-dormant microbes back into metabolic activity, with those exhibiting higher rRNA operon copy numbers responding the fastest. The responders identified at d1 for xylose utilization have all been noted for exhibiting some form of dormancy strategy (Finkel, 2006; Sachidanandham and Gin, 2008; Mulyukin et al., 2009; Jones and Lennon, 2010; Darcy et al., 2011; Lay et al., 2013; Rittershaus et al., 2013; Tada and Grossart, 2013), though the only spore-forming responders at d1 are Firmicutes. Additionally, d1 responders exhibit 6-14 rRNA operon copies with the exception of the Betaproteobacteria Comamonadaceae and the Actinobacterial OTUs which exhibit 1-2 copies according to representative taxa in the rrnDB v. 3.1.227 (Klappenbach et al., n.d.; Lee et al., n.d.). With the exception of the single Firmicutes (Paenibacillaceae), responders on d3 possess 3-6 rRNA operon copies, less than responders on d1. Similar to d3, 85% of d7 responders exhibit between 1-5 rRNA operon copies with the remaining 15% (Flavobacteriaceae, Enterobacteriaceae, and Paenibacillaceae) containing between 6-14 rRNA operon copies. Paenibacillus (100% identity) was the only OTU to be identified at every time point (up to day 30) as a xylose responder. This result suggests that large numbers of cells from Paeni-bacillus sporulated after ¹³C-labeling of their DNA and that these spores remained throughout the experiment. It was the the second most enriched xylose responder, log₂ fold change (l2fc) of 3.5, measured in the time series second only to a Gammaproteobacteria (Xanthomonadaceae; I2fc = 3.7).

Cellulose degrader DNA exhibits greater bouyant density shifts upon $^{13}\mathrm{C}$ incorporation than xylose degrader DNA.

DNA BD shifts due to ¹³C-asimilation differ across phylogenetic types. Temporal dynamics of C-assimilation in soil. The dynamics of ¹³C-xylose and ¹³C-cellulose assimilation varied dramatically. Isotope incorporation increases the bouyant density (BD) of DNA and labeled DNA is enriched in 'heavy' fractions of the density gradient. Isotope incorporation by an OTU is revealed by enrichment of the OTU in heavy CsCl gradient fractions containing ¹³C labeled DNA relative to heavy fractions from control gradients (no ¹³C labeled DNA). Variation in 16S rRNA gene amplicon pool composition in fractions of ¹³C-labeled samples and their corresponding controls is readily observed in 'heavy' gradient fractions (partitioning along axis 2, Fig. 1). The amplicon pool composition of 'heavy' fractions of ¹³C-xylose and ¹³C-cellulose samples vary from corresponding controls and from each other, indicating that the substrates were assimilated by different members of the microbial community(Fig. 1A).

The ¹³C-incorporation reveals temporal dynamics of C degradation demonstrated by ¹³C-xylose incorporation at days 1, 3, and 7 and ¹³C-cellulose incorporation at days 14 and 30 (Fig. 1B), as expected (Amelung *et al.*, 2008). The microbial community changed significantly (pval) with time in the bulk community supporting the temporal dynamics observed in the gradient fraction amplicons (Fig. S2). Although within a single time point, the bulk community demonstrated no significant difference between treatments.

'Heavy' fraction amplicon pools from samples that received ¹³C-xylose diverged from corresponding controls on days 1

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through 7 (Fig. 1). Furthermore, amplicon pool composition varied across these days (Fig. 1B) indicating dynamic changes in ¹³C-xylose assimilation with time. At days 14 and 30 heavy fractions from ¹³C-xylose labeled samples are no longer differentiated from corresponding controls indicating that ¹³C is no longer detectable in DNA. The decline in ¹³C-labelling of DNA is likely due to isotopic dilution resulting from assimilation of unlabeled C and/or due to cell turnover resulting from mortality.

¹³C-cellulose incorporation isn't detected until day 14 and amplicon composition is consistent for both days 14 and 30 (Fig. 1). The consistency of amplicon composition for cellulose degradation over time compared to xylose suggests a wider array of microorganisms utilize xylose, whereas, cellulose utilization occurs in a select few. This is consistent with long standing notions that more microorganisms are capable of utilizing simple carbohydrates than complex C substrates.

Differential C utilization by taxa. Individual OTUs that assimilated ¹³C-substrates were identified using the DESeq framework (Anders and Huber, 2010) to analyze differential representation in 'heavy' fractions (Fig. 2). There were 43 and 35 unique OTUs that significantly (false discovery rate corrected *P*-values <0.10, SI) assimilated ¹³C-xylose and ¹³C-cellulose, respectively; herein called 'responders' (Figs. S3, S4,S5).

There were 6 shared responders among all unique responders identified in both the xylose and cellulose treatments (n = 72); Stenotrophomonas, Planctomyces, two Rhizobiaceae, Comamonadaceae, and Cellvibrio. Of these, Stenotrophomonas and Comamonadaceae are the only taxa that are among the top ten l2fc responses measured in both treatments. On the other hand, the only shared responder that is not among the top ten responders for either the cellulose or xylose treatment is Rhizobiaceae. Two of the shared responders corresponded in time between the two treatments (Table S1); Cellvibrio (d3) and Planctomyces (d14).

Responder Characteristics. We found xylose responders were from higher rank abundances than cellulose responders, however, cellulose responders exhibited a greater shift in BD (i.e. assimilated more ¹³C) than xylose responders in response to isotope incorporation (Fig. 3).

The kernel density estimate (KDE) of BD shifts resulting from ^{13}C -assimilation reveal that cellulose responders exhibit a significantly ($p{<}0.01$) greater BD shift than xylose responders (Fig. 3A). A density profile for each responder is generated for the experimental and control treatment at each of the sampling time points using relative abundances from sequence libraries (Figs. S5, S6). The difference in center of mass for each set of density profiles (control and experimental) is measured (supp. MM) and each KDE curve represents the collection of density shifts calculated for all responders in the ^{13}C -cellulose or ^{13}C -xylose treatment (Fig. 3A). We observe xylose utilizers having a smaller density shift (0.008 \pm 0.008 g mL $^{-1}$) than cellulose utilizers (0.015 \pm 0.009 g mL $^{-1}$), with few exceptions.

Most xylose responders are found at higher rank abundances than cellulose responders (0.01 < p <0.05), which fall among the rarer taxa in the tail of the RA curve (Fig 3B). This demonstrates that many taxa important to cellulose cycling are present in the rarer fraction of the overall microbial community. Yet, the transitions in abundances of responders is difficult to discern in the bulk community abundances (Figs. S5 , S6) or may not be detected with bulk community sequencing efforts. For example, the increase in Bacteroidetes in the xylose treatment at d3 is not observed in the bulk community abundances. Other instances may result in subtle changes in bulk community abundance that would be difficult to differentiate from natural variation or methodological noise.

Patterns of carbon use vary dramatically within phylum. Dynamic patterns of ¹³C-assimilation from xylose and cellulose oc-

cur at discrete, fine-scale taxonomic units (Fig. 4). Responders for xylose and cellulose are widespread across 6 and 7 phyla, respectively (Fig. 4). There are 5 phyla containing responders for both treatments; of all the responder OTUS detected within those phyla for either xylose or cellulose, there are only six OTUs that respond to both xylose and cellulose (discussed previously). This result suggests that phyla do not represent coherent ecological units with respect to the soil C-cycle, that is, taxa within phyla exhibit differences in substrate use, level of substrate specialization, and dynamics of incorporation.

In this study, we have identified Actinobacteria responders for both substrates (Figs. S5, S6). Although there were no shared Actinobacteria OTUs that responded to both xylose (Microbacteriaceae, Micrococcaceae, Cellulomonadaceae, Nakamurellaceae, Promicromonosporaceae, and Geodermatophilaceae) and cellulose (Streptomycetaceae and Pseudonocardiaceae). This information may suggest that while Actinobacteria exhibit an ability to utilize an array of carbon substrates, substrate use may be more clade specific and not widespread throughout the phylum (Fig. 4). Similarly, Bacteroidetes responders were identified for both substrates, yet, at a finer taxonomic resolution there is a clear differential response for xylose (Flavobacteriaceae and Chitinophagaceae) and cellulose (Cytophagaceae).

Whole phylum responses were not detected for xylose or cellulose yet utilization of these substrates spanned many phylogenetically diverse groups. Within each phylum we observed substrate utilization at the clade or single taxa level with each exhibiting a unique pattern of ¹³C-assimilation over time (Fig. 4, heatmap). It has previously been suggested that all taxa within a phylum are unlikely to share ecological characteristics (Fierer et al., 2007), and furthermore, within a species population (Hunt et al., 2008; Preheim et al., 2011; Choudoir et al., 2012). Habitat traits of coastal Vibrio isolates were mapped onto microbial phylogeny revealing discrete ecological populations based on seasonal occurrence and particulate size fractionation (Hunt et al., 2008; Preheim et al., 2011). Yet, it has been proposed that the microbial community functionality responsible for soil C cycling appear at the level of phlya rather than species/genera (Schimel and Schaeffer, 2012). The traditional phylum level assignment conventions could in part be due to limitations in finer scale taxonomic identifications or methodological limitations (i.e. sequencing depth). Our data in concert with others (Fierer et al., 2007; Hunt et al., 2008; Goldfarb et al., 2011; Preheim et al., 2011; Choudoir et al., 2012) would suggest that assigning substrate utilization of a few OTUs or clades as a phylum level response is not accurate.

Conclusions. We have demonstrated how next generation sequencing-enabled SIP gives an OTU level resolution for substrate utilization. Using this technique, we are able to resolve discrete OTUs that would otherwise be missed using bulk community sequencing efforts. Additionally, this technique provides greater taxonomic resolution than previous techniques (cloning, TRFLP, ARISA) used to determine substrate utilizing community members. While we are currently able to resolve highly responsive OTUs, there is still a need to resolve taxa that are partially responsive which we cannot differentiate from noise with confidence at this time. Although, if we could identify partially responsive taxa, their contributions to the C-cycle would still be difficult to discern. For example, a generalist utilizing many substrates including ¹²C substrates and the ¹³C-labeled substrate may exhibit the same partial labeling that a specialist utilizing both the 13C-substrate and the same substrate (unlabeled) that is inherent in the soil. Additionally, partially labeled taxa could be further down the trophic cascade including predators or secondary consumers of waste products from primary consumer microbes that were highly labeled.

OTUs that assimilate xylose and those that assimilate cellulose are largely mutually exclusive. Those OTUs that assimilate xylose are labeled within 1-7 days, while those that assimilate cel-

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lulose are labeled primarily after 2-4 weeks. The xylose responders demonstrate a smaller change in BD than the cellulose responders suggesting that xylose responders assimilate multiple C sources (labeled and unlabeled) consistent with a generalist response, while cellulose responders are more heavily labeled suggesting that cellulose is their main source of C, a response more consistent with a specialist lifestyle. Xylose responders include many taxa, such as spore-fomers, known for the ability to respond rapidly to an influx of new nutrients while cellulose responders include many OTUs that are common uncultivated soil organisms. Finally, xylose responders are more abundant in the community while cellulose responders are, on average, more rare as indicated by their rank abundance within the soil community. These results indicate that different bacteria in soil have distinct physiological and ecological responses which govern their interactions with soil C pools.

We did not observe consistent C utilization at the phylum level although both xylose and cellulose utilization were observed across 7 phyla each revealing a high diversity of bacteria able to utilize these substrates. The high taxonomic diversity may enable substrate metabolism under a broad range of environmental conditions (Goldfarb et al., 2011). Other studies of microbial communities have observed a positive correlation with taxonomic or phylogenetic diversity and functional diversity (Tringe, 2005; Gilbert et al., 2010; Philippot et al., 2010; Bryant et al., 2012; Fierer et al., 2012, 2013). The data presented here supports that specific functional attributes can be shared among diverse, yet distinct, taxa while closely related taxa may have very different physiologies (Philippot et al., 2010; Fierer et al., 2012). This information adds to the growing collection of data suggesting that community membership is important to biogeochemical processes. Furthermore, it highlights a need to examine substrate utilization by discrete microbial taxa within a whole community context to better understand how specific community members function within the

The sensitivity of SIP-NGS provides a means to elucidate substrate utilization by discrete microbial taxa with the hope that we can begin to construct a belowground C food web. We obtained enough information to conclusively determine isotope incorporation for 61% of the more than 6,000 OTUs detected. For those OTUs with enough information (n = 3,825), approximately 2% (n = 72) significantly assimilated ¹³C from either xylose or cellulose. In the future deeper sequencing will enable us to increase coverage and assess C use by more community members. Using the informations we gain from SIP-NGS, we can expand our knowledge of specific C-cycling OTUs by taking a targeted metagenomic approach in the nucleic acid pools of 'heavy' fractions. Furthermore, we can now expand our knowledge of soil C use dynamics to a wide array of C substrates and increase our grasp on specific community member contributions. Illuminating these microbial contributions associated with decomposition in soil are important because as environments change, there are measurable and functional changes in soil C (Grandy and Neff, 2008) which could cumulatively have large impacts at a global scale.

Methods

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Additional information on sample collection and analytical methods is provided in SI Materials and Methods.

Twelve soil cores (5 cm diameter x 10 cm depth) were collected from six random sampling locations within an organically managed agricultural field in Penn Yan, New York. Soils were pretreated by sieving (2 mm), homogenizing sieved soil, and preincubating 10 g of dry soil weight in flasks for 2 weeks. Soils were amended with a 5.3 mg g soil⁻¹ carbon mixture; representative of natural concentrations Schneckenberger *et al.*, 2008. Mixture con-

tained 38% cellulose, 23% lignin, 20% xylose, 3% arabinose, 1% galactose, 1% glucose, and 0.5% mannose by mass, with the remaining 13.5% mass composed of an amino acid (in-house made replica of Teknova C0705) and basal salt mixture (Murashige and Skoog, Sigma Aldrich M5524). Three parallel treatments were performed; (1) unlabeled control,(2) 13 C-cellulose, (3) 13 C-xylose (98 atom% 13 C, Sigma Aldrich). Each treatment had 2 replicates per time point (n = 4) except day 30 which had 4 replicates; total microcosms per treatment n = 12, except 13 C-cellulose which was not sampled at day 1, n = 10. Other details relating to substrate addition can be found in SI. Microcosms were sampled destructively (stored at -80°C until nucleic acid processing) at days 1 (control and xylose only), 3, 7, 14, and 30.

Nucleic acids were extracted using a modified Griffiths procotol Griffiths et al., 2000. To prepare nucleic acid extracts for isopycnic centrifugation as previously described Buckley et al., 2007, DNA was size selected (>4kb) using 1% low melt agarose gel and β -agarase I enzyme extraction per manufacturers protocol (New England Biolab, M0392S). For each time point in the series isopycnic gradients were setup using a modified protocol Neufeld et al., 2007 for a total of five ¹²C-control, five ¹³C-xylose, and four ¹³Ccellulose microcosms. A density gradient (average density 1.69 g mL⁻¹) solution of 1.762 g cesium chloride (CsCl) ml⁻¹ in gradient buffer solution (pH 8.0 15 mM Tris-HCl, 15 mM EDTA, 15 mM KCl) was used to separate ¹³C-enriched and ¹²C-nonenriched DNA. Each gradient was loaded with approximately 5 $\mu \mathrm{g}$ of DNA and ultracentrifuged for 66 h at 55,000 rpm and room temperature (RT). Fractions of \sim 100 μ L were collected from below by displacing the DNA-CsCl-gradient buffer solution in the centrifugation tube with water using a syringe pump at a flow rate of 3.3 μ L s⁻¹ Manefield *et al.*, 2002 into AcroprepTM 96 filter plate (Pall Life Sciences 5035). The refractive index of each fraction was measured using a Reichart AR200 digital refractometer modified as previously described Buckley et al., 2007 to measure a volume of 5 μ L. Then buoyant density was calculated from the refractive index as previously described Buckley et al., 2007 (see also SI). The collected DNA fractions were purified by repetitive washing of Acroprep filter wells with TE. Finally, 50 μ L TE was added to each fraction then resuspended DNA was pipetted off the filter into a new microfuge tube.

For every gradient, 20 fractions were chosen for sequencing between the density range 1.67-1.75 g mL⁻¹. Barcoded 454 primers were designed using 454-specific adapter B, 10 bp barcodes Hamady et al., 2008, a 2 bp linker (5'-CA-3'), and 806R primer for reverse primer (BA806R); and 454-specific adapter A, a 2 bp linker (5'-TC-3'), and 515F primer for forward primer (BA515F). Each fraction was PCR amplified using 0.25 μ L 5 U μ l⁻¹ AmpliTag Gold (Life Technologies, Grand Island, NY; N8080243), 2.5 μ L 10X Buffer II (100 mM Tris-HCl, pH 8.3, 500 mM KCl), 2.5 μ L 25 mM MgCl $_2$, 4 μ L 5 mM dNTP, 1.25 μ L 10 mg mL BSA, 0.5 μ L 10 μ M BA515F, 1 μ L 5 μ M BA806R, 3 μ L H₂O, 10 μ L 1:30 DNA template) in triplicate. Samples were normalized either using Pico green quantification and manual calculation or by SequalPrepTM normalization plates (Invitrogen, Carlsbad, CA; A10510), then pooled in equimolar concentrations. Pooled DNA was gel extracted from a 1% agarose gel using Wizard SV gel and PCR clean-up system (Promega, Madison, WI; A9281) per manufacturer's protocol. Amplicons were sequenced on Roche 454 FLX system using titanium chemistry at Selah Genomics (formerly EnGenCore, Columbia, SC)

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Figures

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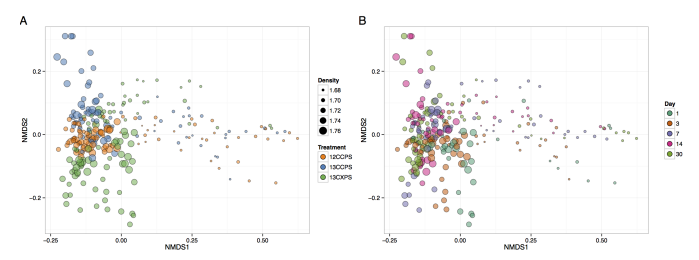


Fig. 1. NMDS analysis from weighted unifrac distances of 454 sequence data from SIP fractions of each treatment over time. Twenty fractions from a CsCl gradient fractionation for each treatment at each time point were sequenced (Fig. S1). Each point on the NMDS represents the bacterial composition based on 16S sequencing for a single fraction where the size of the point is representative of the density of that fraction and the colors represent the treatments (A) or days (B).

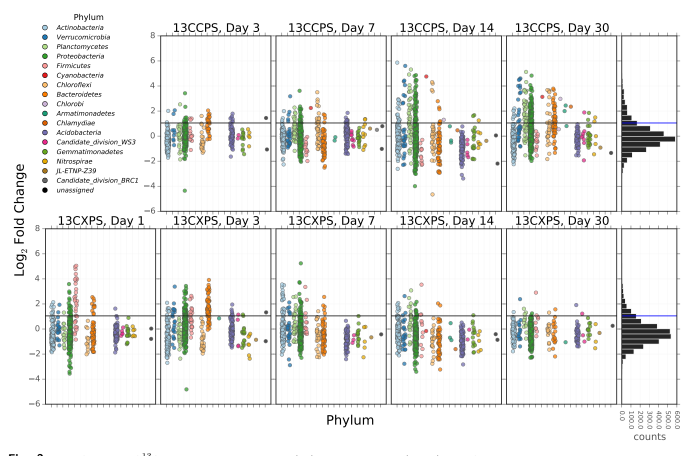


Fig. 2. Log₂ fold change of 13 C-responders in cellulose treatment (top) and xylose treatment (bottom). Log₂ fold change is based on the relative abundance in the experimental treatment compared to the control within the density range 1.7125-1.755 g ml⁻¹. Taxa are colored by phylum. 'Counts' is a histogram of number of sequences for each log₂ fold change value.

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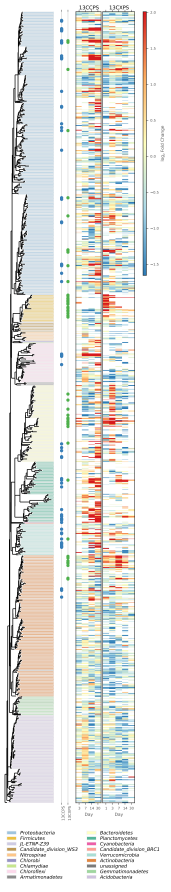


Fig. 3. Phylogenetic tree of sequences passing a user defined sparcity threshold (0.6) for at least one day of the time series. Branches are colored by phylum. ¹³C-responders for cellulose (blue) and xylose (green) are indicated by a point beside the respective branch. Heatmap demonstrates log₂ fold change of each taxa through the full time series for both treatments (cellulose, left; xylose, right).

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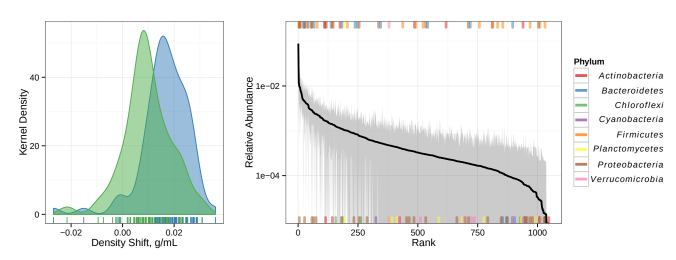


Fig. 4. 13 C-responder characteristics based on density shift (A) and rank (B). Kernel density estimation of 13 C-responder's density shift in cellulose treatment (blue) and xylose treatment (green) demonstrates degree of labeling for responders for each respective substrate. 13 C-responders in rank abundance are labeled by substrate (cellulose, blue; xylose, green) and the phylum which it belongs to.

