Multisubstrate Isotope Labeling and Metagenomic Analysis of Active Soil Bacterial Communities

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ABSTRACT Soil microbial diversity represents the largest global reservoir of novel microorganisms and enzymes. In this study, we coupled functional metagenomics and DNA stable-isotope probing (DNA-SIP) using multiple plant-derived carbon substrates and diverse soils to characterize active soil bacterial communities and their glycoside hydrolase genes, which have value for industrial applications. We incubated samples from three disparate Canadian soils (tundra, temperate rainforest, and agricultural) and diverse soils to characterize active soil bacterial communities and their glycoside hydrolase genes, which have value for industrial applications. We incubated samples from three disparate Canadian soils (tundra, temperate rainforest, and agricultural) and diverse soils to characterize active soil bacterial communities and their glycoside hydrolase genes, which have value for industrial applications. We incubated samples from three disparate Canadian soils (tundra, temperate rainforest, and agricultural) and diverse soils to characterize active soil bacterial communities and their glycoside hydrolase genes, which have value for industrial applications.

IMPORTANT The ability to identify genes based on function, instead of sequence homology, allows the discovery of genes that would not be identified through sequence alone. This is arguably the most powerful application of metagenomics for the recovery of novel genes and a natural partner of the stable-isotope-probing approach for targeting active-yet-uncultured microorganisms. We expanded on previous efforts to combine stable-isotope probing and metagenomics, enriching microorganisms from multiple soils that were active in degrading plant-derived carbohydrates, followed by construction of a cellulose-based metagenomic library and recovery of glycoside hydrolases through functional metagenomics. The major advance of our study was the discovery of active-yet-uncultivated soil microorganisms and enrichment of their glycoside hydrolases. We recovered positive cosmids clones in a higher frequency than would be expected with direct metagenomic analysis of soil DNA. This study has generated an invaluable metagenomic resource that future research will exploit for genetic and enzymatic potential.
TABLE 1 Location and physicochemical characteristics of the soil samples selected for DNA stable-isotope probing incubations

| Sample                      | Location                  | Latitude and longitude | Bulk density (g/cm³) | Amt of carbon (% dry wt) | Moisture (% dry wt) | Amt of nitrogen (% dry wt) | Soil type     |
|-----------------------------|---------------------------|------------------------|----------------------|--------------------------|---------------------|-----------------------------|---------------|
| Arctic tundra (1AT)         | Daring Lake, North-West Territories, Canada | 64°52’N, 111°35’W | 0.2                  | 46.9 BDL                 | 3.9                 | 1417.7                      | Organic       |
| Temperate rainforest (7TR)  | Pacific coastal rainforest, Vancouver Island, Canada | 48°36’N, 124°13’W | 0.6                  | 10.8 BDL                 | 4.9                 | 69.8 0.35                   | Coarse sandy loam |
| Agricultural soil-wheat (11AW) | Elora Research Station, Ontario, Canada | 43°38’N, 80°24’W | 1.1                  | 1.85 0.12 1.7            | 7.4                 | 17.9 0.19                   | Silt loam     |

a For more details, see http://www.cm2bl.org/.
b BDL, below detection limit.

8), supporting the use of cultivation-independent methods, such as metagenomics, as most strategic for the recovery of genes and enzymes from these microorganisms.

Metagenomics captures the genomes of environmental community microbes, circumventing the need for cultivation and enabling the exploration of microbial genetic diversity and biotechnological potential (9). Metagenomic analyses have exposed new microbial pathways and reactions, yielding novel enzymes and products of economic importance. Given that metagenomic studies demonstrate that the majority of total genetic diversity space remains unexplored, “it will be far more efficient and productive to seek new enzymes from metagenomes than from cultures” (10). Indeed, there are several recent examples of GHs (e.g., cellulases) recovered by functional screening of metagenomic libraries from terrestrial environments (e.g., see references 11, 12, 13, and 14). These studies reflect a laborious limitation of bulk DNA metagenomic library construction: in the absence of suitable selections for phenotype, many clones (e.g., tens of thousands) must be screened prior to recovering targets of interest. In addition, recovered clones are theoretically the most abundant target genes in the microbial community of interest. Targeted metagenomic approaches, such as those involving an enrichment culture step (15), thus offer the potential to filter for sequences specific to an activity of environmental or industrial relevance.

Stable-isotope probing (SIP) is a culture-independent method for targeting microorganisms that assimilate a particular growth substrate (16–18). For the analysis of genomic DNA of active organisms, a SIP substrate (e.g., 13C labeled or 15N labeled) is incorporated into the DNA (DNA-SIP) or RNA (RNA-SIP) of active organisms, and isopycnic ultracentrifugation can differentiate labeled nucleic acids from an abundant background of unlabeled community genomes. Combining SIP with metagenomics provides access to the genomes of less-abundant community members and offers insight into complex environmental processes, such as biodegradation (as reviewed in references 19, 20, and 21). Several studies have combined DNA-SIP and metagenomic sequencing to identify high proportions of genes from active microorganisms, such as those using glycerol (22), C5 compounds (23–26), and biphenyl (27, 28). Previous SIP studies reported that in an agricultural soil (clay loam soil, pH 6.6), cellulose was metabolized by *Bacteroidetes*, *Chloroflexi*, and *Planctomycetes*; cellobiose and glucose were degraded predominantly by *Actinobacteria* (8). The results also suggested that cellulolytic bacteria are different from saccharolytic bacteria and that oxygen availability defined the different taxonomic groups involved. Under anoxic conditions, cellulose was metabolized by *Actinobacteria*, *Bacteroidetes*, and *Firmicutes*; carbon from cellobiose and glucose were assimilated by *Firmicutes*. Others found that members of the *Burkholderiales*, *Caulobacteriales*, *Rhizobiales*, *Sphingobacteriales*, *Xanthomonadales*, and Group 1 *Acidobacteria* were associated with three different soils amended with cellulose (29). A recent survey of active bacteria in an Arctic tundra sample found *Clostridium* and *Sporolactobacillus* involved in 13C-glucose assimilation and *Betaproteobacteria*, *Bacteroidetes*, and *Gammaproteobacteria* involved in the assimilation of carbon derived from 13C-cellulose (30). Others have used SIP and labeled cellulose to identify *Dyella*, *Mesorhizobium* sp., *Sphingomonas* sp., and an uncultured deltaproteobacterium (affiliated with *Myxobacteria*) linked to cellulose degradation (6).

The ability to identify genes based on function, instead of sequence homology, is arguably the most powerful application of metagenomics for the recovery of novel genes (31) and a natural partner of the SIP approach for targeting active-yet-uncultured microorganisms (21). Previous studies were focused on the analysis of single substrates or individual samples. In addition, only one previous study combined SIP and functional metagenomic screens, expressing labeled DNA within a surrogate *Escherichia coli* host for identification of enzyme activity (22). In this study, we expand on previous efforts to combine SIP and metagenomics (as reviewed in reference 21), enriching soil microorganisms active in degrading plant-derived carbohydrates and screening GHs through activity-based functional metagenomics. We combined SIP, high-throughput sequencing of labeled 16S rRNA genes and metagenomic DNA, multiple-displacement amplification (MDA), and functional metagenomics to identify active microorganisms and associated GH enzymes. We also isolated GH-positive clones from a cosmid library in a much higher frequency than would be expected with traditional efforts using conventional metagenomics.

RESULTS AND DISCUSSION

Characterization of active soil bacteria. We used DNA-SIP as a targeted approach for enriching active soil microorganisms involved in the metabolism of five plant-derived carbohydrates (glucose, cellobiose, xylose, arabinose, and cellulose). Three disparate soil samples were obtained from the CM/BL soil collection based on maximal physicochemical diversity (Table 1) (http://www.cm2bl.org/). In particular, soil pH was low for the Arctic tundra and temperate rainforest soil samples, suggesting that the microbial composition and diversity of these two samples would be fundamentally different from those in agricultural soil (32, 33).
The water-filled pore space (WFPS) was maintained between 50% and 60% to avoid decreased aerobic microbial activity at WFPS values of >60% (34, 35).

Because 13C-labeled cellulose was commercially unavailable at the time of this research, both native cellulose and 13C-labeled cellulose were produced as the substrates for SIP incubations by *Glusconacetobacter xylinus*, generating predominantly amorphous cellulose (36), which is more readily degraded than crystalline cellulose (37). To ensure detectable labeling, similar to a previous experimental approach (8), glucose, cellobiose, arabinose, and xylose were added weekly (1.5 mmol of C) for 3 weeks, reaching levels approximately 5 to 500 times higher than those normally detected in soils (38, 39). Although substrate concentrations were higher than typical bulk soil concentrations, higher polysaccharide substrate concentrations would be expected in the root rhizosphere and in areas of active plant matter decomposition (as reviewed in reference 39), suggesting that our incubation conditions would not be unrealistic for some naturally occurring soils. These concentrations were chosen to ensure that labeled isotope was more abundant than endogenous soil carbon sources for the success of DNA-SIP, enabling the separation and purification of labeled DNA for subsequent molecular analyses (16, 40). Similar substrate concentrations and incubation times with glucose and cellulose were used previously (30), demonstrating minimal-yet-detectable labeling of DNA in an Arctic tundra soil sample.

Metabolism of labeled substrates in DNA-SIP incubations was confirmed by higher headspace CO2 production in all substrate-amended serum vials compared to uninoculated controls for each of the three soils (Fig. 1). In all cases, cellulose-amended vials demonstrated reduced CO2 production compared to the other substrates, further justifying an extended incubation time for this comparably recalcitrant substrate. The average amount of CO2 released after 6 days was 13% of the headspace, which, after subtraction of the average CO2 produced in uninoculated vials, was approximately equivalent to 1.4 mmol of carbon. This represents 93% of the total weekly carbon added (~1.5 mmol of carbon).

In addition to monitoring CO2 production in all vials, separate soil incubations were prepared with a defined helium-oxygen headspace and glucose amendment in order to monitor O2 consumption. As expected, the addition of glucose stimulated O2 consumption, but the headspace remained oxic for each of the weekly incubation periods over the first 3 weeks (see Fig. S1 in the supplemental material), indicating that weekly aeration of experimental vials was sufficient to deplete CO2 and replenish O2. Maintaining oxic conditions was important to ensure that the DNA-SIP incubation recovered DNA from microorganisms involved in aerobic degradation of complex carbohydrates in addition to capturing DNA from microorganisms involved in anaerobic metabolism (41). Indeed, recent oxic incubations demonstrated activity of anaerobic clostridia (8, 30, 42), presumably because anoxic environments exist even within oxic experimental microcosms.

**Confirmation of isotope labeling.** At the two time points of all incubations (1 and 3 weeks for all substrates, except for cellulose, which was sampled at 3 and 6 weeks), DNA was retrieved for the analysis of bacterial community composition by agarose gel electrophoresis and denaturing gradient gel electrophoresis (DGGE) (43). All DNA extracts from microcosm soils were subjected to density gradient ultracentrifugation and recovered in 12 fractions, which were analyzed in agarose gels. The results demonstrated that all soils possessed more DNA in 13C-incubated heavy fractions (i.e., 1 to 7) than in 12C-control fractions (i.e., 8 to 12) from glucose, cellobiose, arabinose, and xylose SIP incubations (see Fig. S2 to S6 in the supplemental material). For cellulose, only temperate rainforest and agricultural soil incubations resulted in heavier DNA visible in agarose gels corresponding to 13C-labeled sample heavy DNA fractions (see Fig. S6) for the 6-week time point. Similar results were observed for all earlier time points but with less DNA associated with heavy fractions for 13C-incubated samples compared to the later time points (data not shown). Although extended incubation times were important, one caveat of extended incubation times for SIP incubations (e.g., for cellulose) is that labeled carbon might have been distributed more broadly within the microbial community, which may result in less-specific enrichment of substrate-degrading microbial genomes in the resulting data and libraries.

The presence of distinct fingerprint profiles in heavy fractions for 13C-incubated samples, but not for the corresponding 12C-control fractions, demonstrates isotopic enrichment of nucleic acids (16). Bacterial DGGE fingerprints corresponding to all late-
time-point fractions demonstrated unique patterns associated with the heavy fractions (e.g., fractions 1 to 7) for all 13C-incubated SIP microcosms (see Fig. S2 to S6 in the supplemental material). Although some cross-gradient fingerprint variations were associated with 13C-control DNA, these differences were likely GC content shifts because they were pronounced only in the lightest fractions (e.g., fractions 10 to 12) and were distinct from shifts associated with fractionated 13C-DNA. Substrate- and soil-specific heavy fraction patterns were consistent for early- and late-time-point samples (data not shown), which indicated that detected active bacteria were stable over time rather than changing due to food web dynamics (40).

Heavy DNA fingerprints were used to identify fractions containing 13C-labeled DNA for subsequent 16S rRNA gene sequencing, bulk DNA sequencing, and functional metagenomics. Based on DGGE patterns, we identified fraction 5 and/or 6 as being representative of heavy DNA and fraction 10 as representing light DNA for all soils, substrates, and incubation times (see Fig. S2 to S6 in the supplemental material). Although fractions 1 to 5 also may have captured DNA from labeled microorganisms, these fractions were not analyzed further because the vanishingly small proportions of DNA recovered from these gradient fractions would have made PCR and subsequent metagenomic library preparation problematic.

Taxonomic characterization of heavy DNA. We selected representative gradient fractions from all soils, substrates, and incubation times for profiling of the bacterial V3 region of 16S rRNA genes. Based on DGGE data, we selected fractions 5 (heavy) and 10 (light) for Arctic tundra and fractions 5 (heavy) and 10 (light) for temperate rainforest and the agricultural soil. In addition, we sequenced V3 regions of 16S rRNA genes from DNA extracted from the initial soil samples used to establish SIP incubations to determine whether light fractions resembled the original soil community as would be expected. Following paired-end-read assembly, we analyzed 630,000 assembled sequences (10,000 sequences per sample) using an AXIOME management of the QIIME pipeline and additional custom analyses (e.g., multiresponse permutation procedure [MRPP] and indicator species analysis). Good’s coverage (44) for the heavy fraction samples ranged from 84 to 92%, and light fraction samples ranged from 68 to 85%, which indicates that this level of sequencing captured the majority of bacterial taxa in these samples. β diversity was assessed by weighted UniFrac distances visualized within principal coordinate analysis (PCoA) plots. The results indicated that all samples from within each of the three soil treatments were grouped distinctly according to soil type (Fig. 2A), which was highly significant based on MRPP analysis (A = 0.18 [chance-corrected within-group agreement], T = −20.4 [test statistic], P < 0.001). Both the Arctic tundra and temperate rainforest soil profiles grouped more closely with one another, which is likely a result of both soils sharing low pH (Table 1), a major determinant of soil bacterial diversity and taxonomic composition (45, 46). In addition, all heavy and light fraction profiles for the three soils were clustered distinctly (Fig. 2A), which was also highly significant (A = 0.40, T = −28.3, P < 0.001). Native soil phylogenetic profiles clustered with their respective light fractions, indicating that the "background" bacterial community remained relatively constant throughout the SIP incubation. Although the two time points for some 13C-labeled substrates grouped together (Fig. 2B), the differences between heavy and light fractions were much greater than those observed between the five substrates used in this study.

Many operational taxonomic units (OTUs) were affiliated with SIP-derived heavy DNA, but multiple permutations of the analysis were required to summarize indicator OTUs for different sample subsets. We used indicator species analysis (47), with an indicator value (IV) threshold of 70% and a >250 minimum sequence sum threshold to identify the strongest significant OTUs (P < 0.01) associated with (i) all heavy DNA samples (versus all light
identified in heavy fractions from tundra soil (1AT) were soil type showed that the predominant genus-classified OTUs in the supplemental material). The indicator species analysis from and Asticcacaulis tinobacteria

DNA samples) (Fig. 3; see Table S1 in the supplemental material), (ii) all heavy DNA samples within each soil type (versus all light DNA samples for the same soil type) (see Table S2 in the supplemental material), (iii) each substrate across all heavy DNA samples from all soil types (versus the heavy DNA for the other substrates for the same soil type heavy DNA) (see Table S3 in the supplemental material), and (iii) each substrate from heavy DNA within each soil type (versus all light DNA samples from all soils, indicator species DNA) (see Tables S4 to S6 in the supplemental material).

When we compared OTUs associated with all heavy DNA samples versus all light DNA samples from all soils, indicator species analysis revealed multiple poorly classified indicators, in addition to genus-classified OTUs associated with the Salinibacterium (Actinobacteria), Devosia (Alphaproteobacteria), Telmatospirillum (Alphaproteobacteria), Phenylbacterium (Alphaproteobacteria), and Asticcacaulis (Alphaproteobacteria) genera (Fig. 3; see Table S1 in the supplemental material). The indicator species analysis from all heavy DNA samples versus all light DNA samples within each soil type showed that the predominant genus-classified OTUs identified in heavy fractions from tundra soil (1AT) were Salinibacterium (Actinobacteria), Rhodanobacter (Gammaproteobacteria), Conexibacter (Actinobacteria), Telmatospirillum (Alphaproteobacteria), Asticcacaulis (Alphaproteobacteria), and Burkholderia (Betaproteobacteria), in addition to OTUs within orders such as Sphingomonadales and Acidobacterales (see Table S2 in the supplemental material). The temperate rainforest soil (7TR) heavy DNA was dominated by OTUs classified to the genera Paucibacter (Betaproteobacteria), Burkholderia (Betaproteobacteria), Spirochaeta (Spirochaetes), Salinibacterium (Actinobacteria), Telmatospirillum (Alphaproteobacteria), Labrys (Alphaproteobacteria), Mesorhizobium (Alphaproteobacteria), and Phenylbacterium (Alphaproteobacteria), in addition to uncharacterized genera from other phyla, such as Verrucomicrobia (see Table S2).

![Cleveland plot of operational taxonomic unit (OTU) abundance for OTUs possessing the highest indicator values (i.e., >70%) for an association with DNA-SIP heavy DNA (black squares [average abundance]) for all substrates and soils combined, in comparison to light DNA (gray squares [average abundance]). Taxonomic affiliations are included for phyla, with additional classifications for order (o_), family (f_), and genus (g_). For additional details, see Table S1 in the supplemental material.](mbio.asm.org)
positive and negative controls. Substrates. MU, methylumbelliferone units based on equal volumes of sample for each assay.

$$\text{Activity (\muM MU released)}^a$$

| Clone   | Insert size (kb) | $\alpha$-1-Arabinofuranosidase pyranoside | $\beta$-1-Cellobiopyranoside | $\beta$-1-Glucopyranoside | $\beta$-1-Xylopyranoside | N-Acetyl-$\beta$-1-galactosaminide | CMC activity$^b$ |
|---------|-----------------|--------------------------------------|-------------------------------|--------------------------|--------------------------|-----------------------------------|-----------------|
| C122    | 21.6            | 0.4                                   | 0.2                           | 0.6                      | 0.7                      | 124.2                             | −               |
| C424    | 8.2             | 0.9                                   | 57.6                          | 109.4                    | 1.6                      | 0.7                               | −               |
| C762    | 13.5            | 2.4                                   | 5.4                           | 21.2                     | 0.7                      | 0.4                               | −               |
| C1024   | 16.8            | 123.8                                 | 6.5                           | 35.8                     | 1.7                      | 0.5                               | −               |
| C1088   | 11.9            | 0.5                                   | 25.6                          | 79.2                     | 1.2                      | 0.6                               | −               |
| C2194   | 12.9            | 0.5                                   | 0.3                           | 0.6                      | 0.4                      | 39.6                             | −               |
| C2380   | 14.9            | 0.38                                  | 0.46                          | 0.53                     | 0.41                     | 0.40                              | ++ ++           |
| C2044   | 14.7            | 0.40                                  | 0.40                          | 0.52                     | 0.39                     | 0.36                              | ++ ++           |

$^a$ Cellulase activity was scored by Congo red staining of clones on the LB-CMC plate. Other activities were measured in cell-free extracts using methylumbelliferone-based substrates. MU, methylumbelliferone units based on equal volumes of sample for each assay.

$^b$ CMC, carboxymethyl cellulose. Plate-based clearing (high, ++ +; medium, ++; negative, −) was detected by Congo red stain and activity based on comparison to those of positive and negative controls.

Xanthomonadales, Actinomycetales, Burkholderiales, and Bacillales (see Table S2), among others.

Ordered associated with the metabolism of cellulose were dominated by Actinomycetales and Caulobacterales (genus Phenylobacterium) (see Table S3 in the supplemental material). Members of the Alphaproteobacteria were associated with the metabolism of arabinose, and members of the order Rhizobiales were strongly associated with the metabolism of xylose. There were no specific indicator species associated with glucose or cellobiose across all soils (see Table S3), which might also suggest that abundant soil OTUs were also active in assimilating these substrates.

The predominant indicator species for the agricultural soil fed with [13C]glucose were associated with Paenibacillus (Bacillales) (see Table S4 in the supplemental material). The use of cellulose was associated with Mesorhizobium (Alphaproteobacteria), Devosia (Alphaproteobacteria), and Cellvibrio (Gammaproteobacteria), in addition to other poorly classified OTUs from the Sphingomonadales and Actinomycetales. The use of cellulose in temperate rainforest soil was associated with the Myxococcales (Deltaproteobacteria) (see Table S5 in the supplemental material). An OTU affiliated with Caulobacterales was associated with the metabolism of glucose in Arctic tundra. Nevskia (Gammaproteobacteria), and two OTUs affiliated with the Acidobacteria were associated with tundra cellulose assimilation (see Table S6 in the supplemental material). No other OTUs were significant indicators for the remaining substrates (i.e., cellobiose, arabinose, and xylose) for the three soils, which might indicate that active taxa were also abundant soil bacteria.

Although our DNA-SIP incubation revealed many poorly classified indicator taxa (see Tables S1 to S6 in the supplemental material), many of the indicator species associated with heavy DNA were expected based on previous studies. For example, Salinibacterium was associated with frozen soils from glaciers (48) and Arctic permafrost (49). This genus has been associated with the metabolism of a variety of carbon sources, including sucrose, glucose, cellobiose, mannose, melibiose, maltose, galactose, arabinose, and fructose (48, 50). In addition, Devosia species were isolated from greenhouse soil and beach sediments, testing positive for the hydrolysis of esculin, β-galactosidase, β-glucosidase, and N-acetyl-β-glucosaminidase, although unable to degrade carboxymethyl cellulose (CMC) (51, 52). Phenyllobacterium and Burkholderia are abundant in forest soils (53) and the genus Asticaulus was identified among aerobic chemoheterotrophs in tundra wetlands, able to use glucose, sucrose, xylose, maltose, galactose, arabinose, lactose, fructose, rhamnose, and trehalose, among other carbon sources (54). The genus Spirochaeta has some species that are free-living saccharolytic and obligate or facultative anaerobes and were isolated from diverse environments, mainly from extreme aquatic environments (55, 56). Spirochaeta americana was reported to be a consumer of D-glucose, fructose, maltose, sucrose, starch, and D-mannitol (56), and Spirochaeta thermophilic was reported to be a cellulolytic organism; the study of its genome revealed a high proportion of genes encoding more than 30 GHs (55).

**MG-RAST analysis and functional annotation.** We used next-generation sequence analysis of bulk [13C]-labeled DNA to survey the prevalence of annotated GHs within three pooled samples that were targeted for subsequent functional metagenomic screens. Guided by the UniFrac-based PCoA plot (Fig. 2), we pooled heavy

**TABLE 2** Substrate-specific activities of positive metagenomic clones from the [13C]cellulose DNA-SIP library

*FIG 4* Glycoside hydrolase (GH) families associated with pooled heavy DNA. Functional annotation of the metagenomic data revealed diverse GH gene representation within the pooled heavy DNA.
DNA samples representing all substrates (except cellulose) associated with low pH (i.e., temperate rainforest, Arctic tundra), heavy DNA for all substrates (except cellulose) from the agricultural soil, and the cellulose-enriched DNA from the three soils. Analysis of paired-end reads was performed by MG-RAST using annotations derived from the Swiss-Prot/Uniprot database. Only 19.4% (low-pH library), 19.6% (cellulose library), and 22.0% (agricultural library) of sequences were annotated by Swiss-Prot in MG-RAST using an E value threshold of 0.01, which is an important consideration for any subsequent analysis of annotation data based on this minority of sequences. Nonetheless, using a custom Perl script to convert Swiss-Prot annotations to CAZY GH identifiers, we detected 81 distinct GH families for the pooled-cellulose library and 80 GH families for each of the low-pH and agricultural soil composite libraries. The distribution of annotated GHs varied between samples, and the most abundant families in the three pooled samples were GH1, -2, -3, -5, -9, -13, -23, -28, and -35 (see Table S7 in the supplemental material). In addition, the three next-generation sequence data sets were very similar in their distributions (i.e., r > 0.99) for the three libraries (Fig. 4), and all had representation among GH families commonly associated with known cellulases (GH1, -2, -3, -4, -5, -9, -12, -45, -48, and -61), hemi-cellulases (GH8, -10 to -12, -26, -28, -53, and -74), and debranching enzymes (GH51, -54, -62, -67, -78, and -74) as reviewed elsewhere (57, 58). The GH families involved in the hydrolysis of cellulose that were most abundant in our data were GH families 3, 5, and 9 (Fig. 4; see Table S7). However, given that most GH family annotations were not represented by known CAZY identifiers and that only ~20% of our paired-end reads were annotated by Swiss-Prot, the abundance and distribution of functional GH families in our pooled DNA is underrepresented. As a result, we used functional screens of large-insert metagenomic libraries for the recovery of GHs to help circumvent the limitations of sequence-based analysis of our heavy DNA samples.

**Enriched metagenomic library.** Pooled high-molecular-weight DNA from the 13C-cellulose-enriched SIP incubations for the three soils were captured in cosmid libraries and screened for GHs involved in the degradation of cellulose and other plant-derived polymers based on activity in *E. coli*. Multiple-displacement amplification (MDA) increased the amount of nucleic acids obtained from pooled cellulose DNA-SIP incubations prior to the isolation of 25- to 75-kb DNA fragments via pulsed-field gel electrophoresis (PFGE). The cellulose-SIP metagenomic library contained ~83,000 clones with an average insert size of 31 kb based on restriction digestion of a subset of 40 random clones (data not shown). These results compare favorably to a library of ~10,500 clones generated from MDA-amplified SIP-enriched seawater DNA, which had an average insert size of 27 kb, ranging from 17 to 40 kb (26).

We used a combined parallel approach for functional screening of 2,876 randomly selected clones from the cellulose-enriched metagenomic library. Growth of colonies on LB supplemented with carboxymethyl cellulose (CMC), followed by poststaining with Congo red (59), facilitated identification of clones expressing either endoglucanase (EC 3.2.1.X) or glucosidase (EC 3.2.1.X) activities (60). From the 2,876 clones screened, we identified eight positive clones, two of which (C2380 and C2044) were capable of hydrolyzing CMC (Table 2). Restriction mapping showed that these two clones were distinct (Fig. 5). Clones C122 and C2194 carried dissimilar DNAs encoding β-N-acetyl-galactosaminidases (EC 3.2.1.53). β-Glucosidase activities (EC 3.2.1.21) were detected in clones C424, C762, and C1088. Clones C424 and C1088 contained overlapping DNA—probably from the same organism—consistent with the substrate activity profiles. Restriction pattern of clone C1024 was similar to C1088 and C424 (Fig. 5), but C1024 had both α-L-arabinofuranosidase (EC 3.2.1.55) and β-glucosidase (EC 3.2.1.21). The open reading frame (ORF) en-

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**Table 3** Analysis of cosmid insert end sequences

| Clone | Description | Forward read | Reverse read |
|-------|-------------|--------------|--------------|
| C122  | *Porphyromonas gingivalis* | 4e–5 (29 [40/139]) | 8e–136 (82 [131/162]) |
| C424  | *Cellvibrio* sp. strain BR | 1e–28 (69 [66/80]) | 2e–103 (91 [157/163]) |
| C762  | *Chitonibacter flavus* | 1e–86 (78 [151/171]) | 2e–28 (54 [83/125]) |
| C1024 | *Cellvibrio* sp. strain BR | 2e–17 (95 [34/40]) | 2e–46 (80 [85/96]) |
| C1088 | *Saccharophagus degradans* | 6e–61 (68 [123/150]) | 5e–44 (75 [101/114]) |
| C2194 | *Dyadobacter fermentans* | 1e–91 (95% [140/142]) | Failed sequencing reaction |
| C2380 | *Alcyclobacillus acidocaldarius* | 2e–15 (52 [51/69]) | 3e–105 (96 [162/163]) |
| C2044 | *Cellvibrio* sp. strain BR | 1e–71 (96 [116/118]) | 9e–129 (97 [181/184]) |

*a Cosmids were end sequenced with M13 forward and reverse primers flanking the site of metagenomic DNA insertion. For each clone, two end sequences were obtained and are referred to as “reverse” and “forward” reads. Top matches for BLASTx analyses are shown. Positive results are the number of amino acids from the query that match the amino acids from the subject sequence. The total number of amino acids from the subject is shown.
Acidocaldarius

Saccharophagus degradans

69 to 95% identity (Table 3). Other top BLAST matches included a lytic member of the Gammaproteobacteria, Cellvibrio sp. (61), with 98-99% identity. Although these bacteria are not well characterized to date, other researchers have reported that they use cellulose and other carbohydrates as a carbon source (Table 3), with 29 to 97% identity. Although these bacteria are not well characterized to date, other researchers have reported that they use cellulose and other carbohydrates as a carbon source (62–65). As predicted, the end sequence identities for CA2 and C1088 were very similar taxonomically (i.e., Cellvibrio sp.). On the other hand, end sequence data for C122 and C2194 did not suggest a similar genomic origin (Table 3), consistent with the restriction pattern of these cosmids (Fig. 5). Posterior analysis of reverse and forward end sequences of the positive clones was done by comparing end sequences to Illumina forward and reverse reads from whole-genome sequencing of the three SIP libraries (see Table S8 in the supplemental material). The results showed that the majority of end sequences were represented in the cellulose library, as expected, and only a few sequence matches were found in other libraries using the selected threshold. The high frequency of positive clones after screening of DNA-SIP-derived clones compares favorably to those from previous soil functional metagenomic studies reporting the recovery of single positive cellulose hits from screening of tens of thousands of clones. For example, a single cellulose-encoding clone and two xylanase-encoding clones were recovered from functional screening of 13,800 clones from three fosmid metagenomic libraries derived from grassland in Germany, with an insert size range of between 19 and 30 kb (11). Also, one cellulase-encoding clone was retrieved from the functional screening of 3,024 clones from a bacterial artificial chromosome metagenomic library derived from red soil in China, with insert sizes ranging from 25 to 165 kb (12). In another study, one cellulose-encoding clone was recovered from functional screening of 14,000 clones with an average insert size of 5 kb from a metagenomic phagemid library from a forest soil in China (13). Finally, a CMC-positive clone was retrieved from a metagenomic fosmid library derived from wetland soil in South Korea, after screening of 70,000 clones with an average insert of 40 kb (14). Although not conducted here, a well-replicated direct comparison of GH gene recovery from metagenomic libraries prepared from SIP-derived heavy DNA, light DNA, and the original soil DNA would be necessary to confirm the effectiveness of DNA-SIP. In addition, the ability to recover GH genes in high proportions using cultivation-based enrichment approaches is a well-established alternative to direct metagenomics (15). DNA-SIP incubations are designed to be less dependent on rapid growth of a readily cultivated subset of the microbial community (40). Indeed, our labeled DNA contained many OTUs that were classified poorly within described bacterial taxonomies (see Tables S1 to S6 in the supplemental material). Direct DNA-SIP and enrichment culture comparisons would be valuable but have not yet been conducted to our knowledge.

In summary, the combination of DNA-SIP and metagenomics helped recover soil GHs in higher proportions than all previously reported efforts via direct metagenomics, which demonstrates the power of using DNA-SIP as an activity-based prefilter for targeted metagenomic approaches. Our study demonstrated the capability of scaling DNA-SIP analysis for the interrogation of multiple environmental samples with multiple substrates, with sampling at multiple time points. A high-quality cosmid library with >31-kb inserts was constructed from heavy DNA originating from a 13C-cellulose-incubated sample, and highly efficient screening of GHs from a small set of clones (0.3% positive hits) showed strong potential of the techniques combined in this study for functional metagenomics. Identification of the genes encoding GHs and characterization of these enzymes are ongoing and further functional screening of the 13C-cellulose DNA-SIP library clones in other surrogate hosts will be assessed to identify additional GH representation.

MATERIALS AND METHODS

Soil samples. Three soil samples from the Canadian MetaMicroBiome Library (http://www.cm2bl.org/) were used: Arctic tundra 1 (1AT), temperate rainforest (7TR), and agricultural soil-wheat (11AW). Triplicate surface soils from the top 10 cm below the litter layer were combined to prepare a single composite for each site. Composite soil samples were sieved (2 mm), and subsamples were sent to the Agriculture and Food Laboratory (University of Guelph, Guelph, Ontario, Canada) for analysis of physicochemical properties (Table 1).

SIP. D-Glucose was obtained from Bio Basic (Markham, Ontario, Canada), (U-13C5)-D-glucose (99%) was supplied by Cambridge Isotope Laboratories (Cambridge, Ontario, Canada), D-(-)-cellulbiose, D-(-)-arabinose, and D-(-)-xylose were purchased from Sigma-Aldrich. D-(UL-13C3)-arabinose, D-(UL-13C12)-xylose, and (UL-13C12)-cellulbiose were obtained from Omicron Biochemicals (South Bend, IN). To minimize carbon available for competition with labeled substrates, composite soil samples were preincubated for 2 weeks in darkness at 15°C for 1AT and at 24°C for 7TR and 11AW. Ten grams of soil samples was added to 120-ml serum vials, which were sealed with butyl septa. Incubations were conducted with stable-isotope (13C) and native (12C) sub-
strates, as well as no-substrate controls, for each of the three soils. Finely
shredded cellulose was prepared from Gluconacetobacter xylinus grown
with 13C- or 12C-glucose (30) as the sole carbon source. Purified bacterial
 cellulose (200 mg, 6.6 mmol C) was mixed into serum vials in a single
cost. Labeled (13C) and unlabeled (12C) substrates were added to soil
samples in multiple dosages over periods of 1 week and 3 weeks for glu-
cose, cellobiose, xylose, and arabinose incubations or 3 weeks and 6 weeks
for the cellulose incubations. Serum vials were aerated once per week for
1 h in a fume hood. The weight of incubation vials was assessed weekly,
and water-filled pore space (WFPS) was maintained between 50 and 60%
by adding distilled water and/or substrate for each experiment according
to the following formula (34): WFPS = \( w \left( \rho_p / \rho_s - 1 \right) \), where \( w \) is
the gravimetric water content (%), \( \rho_p \) is the soil bulk density (g/cm\(^3\)),
and \( \rho_s \) is the soil particle density (2.65 g/cm\(^3\)).

GC. CO\(_2\) accumulation in the headspaces of serum vials was deter-
mained using a GC-2014 gas chromatograph (Shimadzu) equipped with a
thermal conductivity detector (TCD), methanizer, and a flame ionization
detector (FID). The gas chromatography (GC) temperatures were main-
tained for the oven (80°C), TCD (280°C), methanizer (380°C), and FID
(250°C). No-carbon control incubations and separate serum vials
amended with 12C-glucose were used as surrogates for experimental vials
because an N\(_2\)-free headspace was required for measurement of O2 with
the gas chromatograph. The headspaces of these separate vials were
flushed with helium and supplemented with oxygen (20%) at the start of
the experiment. Headspace CO2 and O2 were measured every 3 days by
direct injection of 0.5 ml of headspace gas through a packed Poropak Q
column with a helium flow of 20 ml/min.

DNA extraction and isopycnic centrifugation. Two grams of soil was
sampled from each vial at the time points described above. DNA was
extracted with a PowerSoil DNA Isolation kit (MO BIO Laboratories,
Carlsbad, CA) according to the manufacturer’s instructions. Extracted
DNA was quantified using a NanoDrop 2000 UV-Vis spectrophotometer
(Thermo Scientific; Montreal, Quebec, Canada) and a 1% agarose gel with
a 1-kb DNA ladder (Invitrogen) for comparison. Cesium chloride (CsCl)
gradients were processed by ultracentrifugation, and 12 fractions were
collected for each sample as described previously (16, 66).

DGGE. The V3 regions of bacterial 16S rRNA genes were PCR ampli-
fied using primers 341F-GC and 518r (67). Each reaction mixture con-
tained 19.75 μl of UV-treated water, 2.5 μl of 10× Thermolopho reaction
buffer (New England BioLabs), 0.05 μl of deoxynucleoside triphosphates
dNTPs) (100 mM), 0.05 μl of forward primer 341F-GC (100 μM), 0.05 μl
of reverse primer 518r (100 μM), 1.5 μl of bovine serum albumin (BSA)
(10 mg/ml), 0.25 μl of Taq DNA polymerase (5 U/μl) (New England
BioLabs), and 1 μl of DNA template purified from each gradient fraction.
The PCR conditions were initial denaturation at 95°C for 5 min, followed
by 30 cycles of denaturation at 95°C for 1 min, annealing at 55°C for 1 min,
and extension at 72°C for 1 min, followed by a final extension at 72°C for
7 min. All PCR products were analyzed on 1% agarose gels prior to DGGE.
Five micro liters of each PCR product was loaded onto a 10% poly-
acrylamide gel with a denaturing gradient of 30 to 70%. Gels were run at
60°C for 1 h at 85 V (DGGEK-2001-110; C.B.S. Scientific, San Diego,
CA) as described previously (43). A custom DGGE ladder was loaded into
the two outside wells of the gel for subsequent normalization. Gels were
stained for 45 min with SYBR green I nucleic acid gel stain (Thermo
Fisher) and rinsed once in water prior to imaging. Gel images were taken
with a Pharus Plus molecular imager system (Bio-Rad).

Next-generation sequencing. High-throughput sequencing of the 16S
rRNA gene (V3 region) and paired-end-read assembly were con-
ducted as described previously (68, 69). Based on DGGE data, we se-
lected gradient fractions 6 (heavy) and 10 (light) for IAT and fractions
3 (heavy) and 10 (light) for 7TR and 11AW (60 samples in total). Three
25-μl PCR amplifications per sample were conducted, each containing
5 μl of the 5′ Phusion HF buffer (Finnzyme, Finland), 0.125 μl of the
V3F-modified primer (100 μM), 1.25 μl of an indexed reverse primer
(10 μM) (V3-1R to V3-60R), 0.2 μl of dNTPs (100 mM), 0.25 μl of the
Phusion high-fidelity DNA polymerase (2 U/μl) (Finnzyme), and 1 μl of
DNA template (1 to 10 ng). The PCR conditions were as follows: initial
denaturation at 98°C for 2 min, followed by 20 cycles of denaturation at
98°C for 10 s, annealing at 50°C for 30 s, and extension at 72°C for 15 s. A
final extension was performed at 72°C for 7 min. The triplicate 330-bp
PCR products were pooled and analyzed on a 2% agarose gel. Individually
indexed composites were combined in equal nanogram amounts and then
resolved on a 2% agarose gel. The amplicon fragment was excised and
purified using Wizard SV gel and PCR cleanup system (Promega, Madi-
son, WI). Libraries were subjected to 108-bp end sequencing on the Ge-
name Analyzer IIX (Illumina, Inc., San Diego, CA) at the Plant Biotechnol-
yogy Institute (Saskatoon, Saskatchewan, Canada).

Shotgun metagenomic sequencing was performed on DNA from three
pooled fractions of the 13C-labeled DNA from each treatment. Pooling of
heavy DNA resulted in three composite samples for sequencing: (i) “low
pH” (fractions 5 and 6, and 7 of IAT and fractions 4, 5, and 6 of 7TR) for
week 3 incubations with glucose, cellobiose, arabinose, and xylose; (ii)
“agricultural” (fractions 4, 5, and 6 for 22AW) for week 3 incubations with
glucose, cellobiose, arabinose, and xylose; and (iii) “cellulose” (fractions 5, 6,
and 7 for 1AT and fractions 4, 5, and 6 for 22TR and 11AW) for week 6
incubations with cellulose. Shotgun sequencing samples of metagenomic
DNA were prepared using the Nextera DNA sample preparation kit (Illu-
mina). Pooled heavy DNA (25 to 50 ng) was fragmented using the tag-
mentation reaction (–200 to 5,000 bp), according to the manufacturer’s
instructions and purified using the DNA Clean & Concentrator kit (Zymo
Research Corporation, Irvine, CA). Purified fragments were used as the
template for a five-cycle PCR amplification; indexed sequencing adapters
(Epicenter, Madison, WI) were used for the PCR. Each amplified sample
was purified and subjected to size selection (400 to 800 bp) using a Pippin
Precipitate device (Sage Science, Beverly, MA). Afterward, each library was
quantified using the KAPA library quantification kit (KAPA Biosystems
Woburn, MA). Equimolar samples were pooled, concentrated, and quan-
tified. Final concentrations were adjusted to 10 nM. Libraries were se-
quenced using the HiSeq2000 sequencing system (Illumina) by the
Institute for Genomic Biology Core Facility (University of Illinois). Se-
quencing was performed using a TruSeq SBS kit (version 3), and data were
analyzed using the Cassava 1.8 pipeline. Error rates were estimated at
below 0.3%. Each sample yielded 42 to 90 million 100-bp end reads of 62 to
63% average GC content.

Statistical analysis. Taxonomic classification with RDP v2.2 (confi-
dence 0.8 and GreenGenes Oct 2012 revision), principal coordinates anal-
ysis (PCoA) with weighted UniFrac distances, multivariate permutation
procedures (MRPP), and indicator species (IS) analyses of 16S rRNA gene
sequences generated by assembled paired-end reads were performed us-
ing automated exploration of microbial diversity (AXIOME) automation of
PANDAseq (69), the QIIME pipeline (70), and custom AXIOME anal-
yses (71).

MG-RAST analysis and CAZy annotation. Paired-end shotgun se-
quencing from the pooled heavy DNA samples were analyzed for GHs
using the MG-RAST pipeline (72). Reads were annotated by comparison
to sequences in the UniProt database (73), with no maximum E value
cutoff, a 54% minimum percentage identity cutoff, and a 30-bp
minimum-alignment-length cutoff. Using custom Perl scripts (see Algo-
rithms S1 and S2 in the supplemental material), Swissprot and Trembl
database (UniProt release 2012 to 2014) hits were paired with matching
GH family CAZy identifiers by comparing an extracted database of acces-
sion numbers to CAZy identifiers (see Texts S1 and S2 in the supplemental
material).

Cellulose-enriched metagenomic library construction. High-
molecular-weight DNA was extracted from all three soil samples that were
amended with 13C-labeled bacterial cellulose (week 6 time point), using a
gentle enzymatic lysis (74). Humic acids were removed from crude DNA
as described previously (75), using the SCODA device (Aurora, Boreal
Genomics; Vancouver, BC, Canada) with one wash cycle (70 V/cm, 10°C,
90 min) and two concentration cycles (70 V/cm, 10°C, 60 min). DNA was

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analyzed using a 1% agarose gel and quantified with the NanoDrop 2000 spectrophotometer. Samples were subjected to cesium chloride density gradient ultracentrifugation and fraction collection as described previously with minor modifications. Gradient fractions were diluted with 1 volume of water and then, following addition of 2 volumes of ethanol, the DNA was precipitated overnight at −20°C. DNA was collected by centrifugation for 30 min at 15,000 × g. The DNA was air-dried, dissolved in 300 μl of water, and then precipitated by adding 1/10 vol of 3 M sodium acetate (pH 5.3) and 2 volumes of ethanol. After confirming that the fingerprints generated from an alternative lysis protocol were the same as those observed by DGGE, pooled samples and fractions for large-insert cosmids cloning were mixed in the same equal nanogram ratio used to prepare template for sequence-based metagenomics.

To obtain a sufficient amount of DNA for 13C-cellulose-enriched metagenomic library construction, triplicate multiple displacement amplification (MDA) reactions were conducted using the illustra GenomiPhi V2 DNA amplification kit (GE Healthcare, Mississauga, Ontario, Canada), according to the manufacturer’s instructions. Each reaction mixture consisted of ~7 ng of DNA template in order to minimize potential amplification bias (26, 30, 76), yielding 3 to 4 μg of amplified DNA. Positive-control DNA from the kit and negative controls without DNA were run in parallel. MDA products were quantified on a 1% agarose gel and then pooled.

To inactivate 29 DNA polymerase, MDA-amplified DNA (100 μl) was mixed with 613 μl of Tris-EDTA (TE), 73 μl of 10× gel loading buffer, and 6.8 μl of 20% SDS. After being heated at 65°C for 10 min, the sample was left on ice for 5 min and then centrifuged at 15,900 × g for 5 min. The DNA-containing supernatant was loaded onto a 1% pulsed-field agarose gel (with Tris-acetate-EDTA [TAE] buffer) in order to size select DNA. Pulsed-field gel electrophoresis (PFGE) (CHEF Mapper; Bio-Rad) was run at 14°C, 5.5 V/cm, 120° angle, and an initial 1.0-s to final 6.0-s switch time for 20 h. The outer lanes were loaded with a size marker, and following electrophoresis, these lanes were sliced off, poststained with SYBR green I nucleic acid gel stain, and visualized with a Clare Chemical Research Dark Reader. After reassembly of the gel, a gel slice corresponding to 25 to 75 kb of sample DNA was excised, electroeluted, and concentrated as described previously (77). DNA end repair, ligation with cosmids pCB3, packaging, and transduction into E. coli HB101 were performed as reported previously (77). Resulting recombinant cosmid clones were pooled and saved in 75% dimethyl sulfoxide (DMSO) in 1-ml aliquots at −75°C. Prior to pooling, 40 random E. coli clones from the plates were selected for analysis of cosmid DNA restriction patterns. The average sizes of cloned metagenomic DNA and coverage of bacterial genomes were calculated based on sites of EcoRI-HindIII-BamHI fragments and the number of recombinant library clones. Additionally, 2,876 random clones were inoculated into LB-Tc in 96-well plates and then grown overnight at 37°C for functional screening.

**Functional screening.** Clones were randomly selected and subjected to activity-based screening of GHSs in E. coli HB101. These clones were grown in 96-well microtiter plates and were replicated onto 150-mm LB-Tc agar plates supplemented with carboxymethyl cellulose (CMC) (0.2%). The plates were incubated at 37°C for 7 day. Following removal of colonies from the plates by washing with water, 0.1% Congo red was used for poststaining.

These clones were also tested for activity on a host of methylumbelliferone-based fluorogenic proxy substrates. Clones were first grown in LB broth containing 15 μg/ml tetracycline at 37°C in microtiter plates. Each well contained one glass bead, and plates were incubated with orbital shaking. After 24 h, 70 μl of pre-culture was transferred to a deep-well plate (96 wells) and cultured in Terrific Broth containing 15 μg/ml tetracycline for a further 24 h at 37°C with a glass bead and orbital shaking. Cells were collected by centrifugation and frozen. For lysis, cell pellets were thawed and chemically lysed using the BugBuster protein extraction reagent (Novagen). GH activities in cell-free extracts were measured using α-1-arabino-furanoside/pyranoside, β-ν-cellobiopiranoside, β-ν-glucopyranoside, β-ν-xylpyranoside, and N-acetyl-β-ν-galactosaminide. Reactions were carried out in 384-well microplates. Library lysates were incubated with 0.1 mM each substrate for 1 h at 50°C in a 40-μl sodium citrate-buffered (50 mM, pH 5) reaction mixture. Reactions were stopped by the addition of 40 μl of 0.2 M glycine (pH 10). Fluorescence was detected at 445 nm following excitation at 370 nm. Clones that demonstrated activity on one or more substrates were subcultured and re-screened on appropriate substrates to eliminate false-positive reactions. Protein concentrations were measured by the Bradford method with bovine serum albumin (BSA) used as a standard. End sequences of positive cosmid clones were obtained by Sanger sequencing using M13 forward and reverse primers at TCGAG (Toronto, Ontario, Canada). We used BLASTx searches of translated nucleotide sequences against the NCBI protein database. End sequences were deposited in GenBank. Posterior BLAST analysis was done searching for sequence similarities in the three libraries: low pH, agricultural, and cellulose (forward and reverse). Sequences with ≥95% similarity and >30 bp were recorded as positive matches.

**Nucleotide sequence accession numbers.** Paired-end reads have been deposited in MG-RAST under identification no. 4482593.3 (low-pH forward), 4483544.3 (low-pH reverse), 4482599.3 (cellulose forward), 4483820.3 (cellulose reverse), 4482600.3 (agricultural forward), and 4483819.3 (agricultural reverse). End sequences of cosmid clones have been deposited in GenBank under accession no. KG771718 to KG771732.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at http://mbio.asm.org/lookup/suppl/doi:10.1128/mBio.01157-14/-/DCSupplemental.

Text S1, TXT file, 0.3 MB.
Text S2, TXT file, 3.3 MB.
Algorithm S1, TXT file, 0.1 MB.
Algorithm S2, TXT file, 0.1 MB.
Figures S1–S6, PDF file, 39.4 MB.
Tables S1–S6, XLSX file, 0.1 MB.
Table S7, XLSX file, 0.1 MB.
Table S8, XLSX file, 0.1 MB.

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