Diversity of Bacteria and Glycosyl Hydrolase Family 48 Genes in Cellulolytic Consortia Enriched from Thermophilic Biocompost

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The enrichment from nature of novel microbial communities with high cellulolytic activity is useful in the identification of novel organisms and novel functions that enhance the fundamental understanding of microbial cellulose degradation. In this work we identify predominant organisms in three cellulolytic enrichment cultures with thermophilic compost as an inoculum. Community structure based on 16S rRNA gene clone libraries featured extensive representation of clostridia from cluster III, with minor representation of clostridial clusters I and XIV and a novel Lutispora species cluster. Our studies reveal different levels of 16S rRNA gene diversity, ranging from 3 to 18 operational taxonomic units (OTUs), as well as variability in community membership across the three enrichment cultures. By comparison, glycosyl hydrolase family 48 (GHF48) diversity analyses revealed a narrower breadth of novel clostridial genes associated with cultured and uncultured cellulose degraders. The novel GHF48 genes identified in this study were related to the novel clostridia Clostridium straminisolvans and Clostridium clariflavum, with one cluster sharing as little as 73% sequence similarity with the closest known relative. In all, 14 new GHF48 gene sequences were added to the known diversity of 35 genes from cultured species.

The exploration and understanding of cellulose fermentation capabilities in nature could inform and enable industrial processes converting cellulosic biomass to fuels and other products. Enrichment of microbial communities that can utilize cellulose is useful in this context for the identification of novel organisms, novel metabolisms, and novel functions. Of particular interest are communities that can utilize cellulose at high temperatures and under anaerobic conditions, featuring high rates of solubilization under conditions where the energy and the reducing power of substrates are conserved in potentially useful fermentation products.

Some evidence indicates that cocultures may be able to utilize cellulose more fully and produce higher concentrations of ethanol than pure cultures of model cellulolytic organisms such as Clostridium thermocellum and Clostridium straminisolvans (16, 20, 34). An initial step toward understanding the functional roles of community members in cooperative cellulose degradation is answering the question of what organisms are present in cellulolytic consortia obtained from nature. Currently, diversity estimation methods applied to cellulolytic communities range from traditional methods targeting the 16S rRNA gene to complex metagenomic analyses targeting the breadth of functional genes present in genomes of mixed cultures and the environment (3).

From a functional gene standpoint, cellulase systems are complex assemblages of multifunctional glycosyl hydrolases, 9, tend to include hydrolyses with multiple substrate specificities, deep evolutionary roots, and extensive sequence diversity within the same organism (19). However, family 48 glycosyl hydrolases include a select group of cellulosomal and unbound cellulases thought to play an essential role in cellulose solubilization by model cellulolytic clostridia (5, 7, 15), actinobacteria (6, 13), and anaerobic fungi (31). One key feature of this family of glycosyl hydrolases (mostly exoglucanases) is their ability to enhance cellulose solubilization in synergistic interactions with family 9 glycosyl hydrolases (2, 13). But unlike the latter, and with the notable exception of CelS and CelY in Clostridium thermocellum, family 48 hydrolases are present mostly in single copies in the genomes of cellulosytic microbes, making family 48 hydrolyase genes a desirable target for primer design and molecular characterization.

In this paper we describe the enrichment of microbial communities from a thermophilic compost pile and provide an assessment of diversity in stable cellulolytic enrichments by addressing total bacterial diversity using the 16S rRNA gene as well as introducing a novel method to assess functional diversity in cellulosytic consortia by targeting glycosyl hydrolase family 48 (GHF48) genes.

MATERIALS AND METHODS

Sampling site. Compost samples were collected at the Middlebury College composting facility in May 2008. Samples were collected with a T-shaped steel coring tube (diameter, 2 cm) from specific locations in the compost pile where temperatures were determined to be the highest, 40 to 50 cm below the surface of the compost pile. Temperatures at each of these hot spots ranged from 52 to 72°C, and samples were designated CO-4, CO-5, and CO-6.

Enrichment protocol and media. Three samples of approximately 15 g of compost were used as inocula in bottles with 100 ml of mineral medium containing 1 g of Avicel (PH-105; FMC Corp., Philadelphia, PA) as the carbon source. Bottles were flushed with nitrogen gas on site immediately after sampling and inoculation, thus ensuring strict anaerobic conditions from primary sampling. The primary enrichment (PE) medium consisted of the following, in grams
per liter: KH₂PO₄, 2.08; KH₂PO₄, 2.22; MgCl₂·6H₂O, 0.1; NH₄Cl, 0.4; CaCl₂·2H₂O, 0.05. Primary enrichment cultures were incubated at 55°C upon arrival at the laboratory and were transferred to maintenance (M) medium after 4 to 6 days of incubation. M medium consisted of 3 g of Avicel/liter, 1.04 g of KH₂PO₄/liter, 1.11 g of KH₂PO₄/liter, 2.5 g of NaHCO₃/liter, 0.2 g of MgCl₂·6H₂O/liter, 0.4 g of NH₄Cl/liter, 0.05 g of CaCl₂·2H₂O/liter, 0.05 g of FeCl₃·4H₂O/liter, 1 ml of SL-10 trace elements/liter (1), 0.5 g of l-cysteine HCl/liter, and 4 ml of 0.025% resazurin/liter. Vitamins were added in the following concentrations (in milligrams per liter): pyridoxine hydrochloride, 0.8; p-aminobenzoic acid (PABA), 0.4; biotin, 0.2; vitamin B₁₂, 0.2; thiamine-HCl, 0.05; folic acid, 0.05; pyridoxine-HCl, 0.1; thioctic acid, 0.5; riboflavin, 0.05. Phosphates and other minerals were prepared and autoclaved separately to avoid precipitation and chemical interactions during autoclaving. Vitamins were sterilized by filtration. Amino acids, tryptophan, and other growth factors were added separately after the autoclaving of fermentor cultures. Yeast extract (Bio-Rad, Hercules, CA) was added after autoclaving to a final concentration of 0.1 g/liter. The concentrations of fermentation products and soluble sugars (acetate, formate, lactate, succinate, cellobiose, and glucose) were analyzed by gas chromatography using an Aminex HPX-87H column on a high-performance liquid chromatography system and by dry weight measurements, using triplicate 3-ml samples after filtration (0.22-μm pore size Millipore Isopore GTTP filters), washing, and drying for 24 h at 60°C. The concentrations of fermentation products and soluble sugars (acetate, ethanol, formate, lactate, succinate, cellobiose, and glucose) were analyzed by high-performance liquid chromatography using an Aminex HPX-87H column (Bio-Rad, Hercules, CA). Pellet nitrogen was measured in centrifuged pellet samples by using a TOC-V combustion analyzer coupled with a TNM-1 total-nitrogen module (Shimadzu Corporation, Columbia, MD) and comparing the results to a 1-g liter⁻¹ nitrogen standard.

**Batch cultivation in fermentors.** After 10 consecutive transfers, anaerobic cultivation of each enrichment culture was carried out at 55°C in 1.6-liter Biostat Aplus fermentors (Sartorius Stedim, Gottingen, Germany), with a working volume of 1 liter and without a pH control. Fermentors were equipped with a non-prene tubing (Cole Palmer Instrument Company, Vernon Hills, IL) to minimize oxygen diffusion. Cultures were grown in the same maintenance (M) medium, with 3 g liter⁻¹ Avicel (PF-105; FMC Corp., Philadelphia, PA) and with reducing agents and vitamins added separately after the autoclaving of fermentor vessel contents. All cultures were inoculated with a 2% (vol/vol) inoculum.

**Analytical methods.** Samples were taken every 3 h from each fermentation experiment for determination of the residual dry weight of cellulose, pH, pellet nitrogen, and fermentation products. Residual cellulose levels were determined by dry weight measurements, using triplicate 3-ml samples after filtration (0.22-μm pore size Millipore Isopore GTTP filters), washing, and drying for 24 h at 60°C. The concentrations of fermentation products and soluble sugars (acetate, ethanol, formate, lactate, succinate, cellobiose, and glucose) were analyzed by high-performance liquid chromatography using an Aminex HPX-87H column (Bio-Rad, Hercules, CA). Pellet nitrogen was measured in centrifuged pellet samples by using a TOC-V combustion analyzer coupled with a TNM-1 total-nitrogen module (Shimadzu Corporation, Columbia, MD) and comparing the results to a 1-g liter⁻¹ nitrogen standard.

**DNA extraction and PCR amplification of the 16S rRNA gene.** After 10 consecutive transfers using M medium with Avicel as the growth substrate, DNA was extracted from enrichment cultures using the ZR soil microbe DNA kit (Zymo Research Corp., Orange, CA) with minor modifications. Triplicate 1.5-ml samples from each enrichment were subjected to DNA extraction. Ballistic explosion times in a Mini-BeadBeater (Bio101, Vista, CA) were 1 min, and bead-beating speed was set to 2,500 rpm to maximize the DNA yield. The DNA obtained from each set of triplicate extractions was pooled for each enrichment. The extracted DNA was used in triplicate PCR amplification targeting the 16S rRNA gene, using the universal oligonucleotide primers 8F and 1492R, designed to anneal to conserved regions of bacterial 16S rRNA genes (18). PCR amplifications were prepared with 10% buffer, 0.25 mM each deoxynucleoside triphosphate (dNTP), 2 U Taq polymerase (all from New England Biolabs), and 0.5 μM each primer (Integrated DNA Technologies, Coralville, IA) for final volumes of 30 μl, using 30 cycles of 94°C (30 s), 50°C (30 s), and 72°C (30 s) with an initial denaturation at 95°C (5 min) and a final extension at 72°C (5 min). As positive controls, cultures of Clostridium thermocellum DSM 1313, Clostridium cellulolyticum H10, and Clostridium straminisolvens DSM 16021 (kindly provided by Mascoma Corporation), Clostridium clariflavum DSM 19732 (obtained from the DSMZ), and Clostridium thermocellum ATCC 27405 were subjected to DNA extraction using the GenElute bacterial genomic DNA kit (Sigma, St. Louis, MO) according to the manufacturer’s instructions.

**Design of GHF48 primers and PCR amplification of GHF48 genes.** A total of 35 sequences, representing all bacterial and fungal GH48 amino acid sequences, were aligned and edited using BioEdit, version 7.0.5 (11). An alignment of the eukaryotic sequences was further used in the primer design for amino acid positions 460 (t) to Ala443, surrounding the catalytic domain of C. thermocellum Cel5A (10). Four sets of primers were designed, and PCR amplification was optimized using a PCR gradient with genomic DNA of C. thermocellum ATCC 27405, C. thermocellum DSM 1313, C. straminisolvens, C. clariflavum, C. cellulolyticum, and C. acetobutylicum as individual positive controls. The final product utilized two degenerate primers, GH48F (5’GGAAATTCTTATAYTGT HATWGAAA-3’) and GH48R (5’-CATGCTTGYAVWCCRAACA-3’). Optimal PCR conditions utilized 30 cycles of 94°C (30 s), 52°C (30 s), and 72°C (30 s), with an initial denaturation at 95°C (5 min) and a final extension at 72°C (5 min), and the same concentrations of reagents used for 16S rRNA gene PCR.

**Clone library generation.** Clone libraries based on the 16S RNA gene and the clostridial GHF48 gene were constructed as previously described (14). Briefly, after the final round of PCR amplification, pooled amplified fragments were purified and cloned into a pGEM-T Easy vector (Promega, Madison, WI). Two vector-specific primers were used for the amplification of the DNA inserts from individual clones: pGemF (5’-GCA AGC CGA TTA AGT TGG G-3’) and pGemR (5’-ATG ACC ATG ATT CCA AG-3’). PCR amplification used the same cycles used for 16S rRNA gene amplification. Clones were screened by agarose gel electrophoresis in order to detect those with inserts of the correct size. After a clone library was created for each enrichment, and each library was screened for a total of 100 clones. Sequencing was performed by Agencourt Bioscience Corporation (Beverly, MA). The amplified clones 16S RNA and GH48 gene sequences were aligned with representative sequences obtained from GenBank using the ClustalX software package (33). BioEdit, version 7.0.5, was used for manual editing of the sequences and, in the case of GH48, their translation into amino acid sequences (11). Rarefaction curves and operational taxonomic unit (OTU) definition at 99% sequence similarity were determined using the DOTUR software package (26). For the construction of phylogenetic trees, distances between pairs of nucleotide sequences were calculated based on the minimum-evolution criterion in the MEGA software package, version 3.1 (17). Bootstrap confidence values were obtained based on 1,000 replications.

**Nucleotide sequence accession numbers.** The nucleotide sequences of both the 16S rRNA genes and glycosyl hydrolase family 48 genes from the clone libraries have been deposited in the GenBank database under accession numbers QG265308 to QG265351 and GQ487568 to GQ487569.

**RESULTS**

**Dynamics of cellulose utilization and fermentation.** After 8 to 10 consecutive transfers, anaerobic compost-derived enrichments CO-4, CO-5, and CO-6 exhibited consistent fermentation rates at 3 and 5 g liter⁻¹ Avicel at 55°C. In batch fermentor experiments using 3 g liter⁻¹ Avicel, between 86% and 97% of the cellulose was utilized in 60-h fermentations (Fig. 1A, C, and E). Cellulose utilization slowed down in all fermentations as the pH dropped from 8.4 to 6.5 in CO-4 and to 7.0 in CO-5 and CO6. Fermentation products accumulated between 20 and 40 h of incubation and consisted mostly of acetate, ethanol, and formate, with no residual sugars detected. Acetate and ethanol were the dominant products in the CO-5 and CO-6 fermentations, yielding between 0.4 and 0.5 g liter⁻¹. However, formate was equally dominant in CO-4 enrichment fermentations, at around 0.5 g liter⁻¹, with higher acetate and ethanol production. Biomass production, as evidenced by changes in pellet nitrogen levels, peaked between 30 and 40 h of incubation for the CO-5 and CO-6 enrichments at around 65 mg N liter⁻¹, while the CO-4 enrichment reached a maximum of 40 mg N liter⁻¹ at around 30 h of incubation (Fig. 1B, D, and F). Based on the products measured, we were able to account for most of the carbon in fermented carbohydrates: at least 94.1% was accounted for in CO-4, 80.2% in CO-5, and 80.5% in CO-6.

**16S rRNA gene diversity.** Different levels of diversity were detected across enrichments at a sequence similarity cutoff of 99%. The CO-4 enrichment had the lowest diversity, with a total of 3 representative operational taxonomic units (OTUs) after 100 clones were surveyed. The CO-5 enrichment displayed slightly higher diversity, with a total of 9 OTUs, while CO-6 had the highest, with a total of 18 OTUs. Rarefaction analysis of 16S rRNA gene libraries (Fig. 2) showed that while...
the CO-4 and CO-5 enrichments had both attained a plateau in the number of retrievable OTUs, the CO-6 enrichment was still yielding new OTUs after 100 clones. However, additional sampling of 50 more clonal sequences from the CO-6 enrichment contributed only two new sequences as singletons (occurrence of 1 clone per OTU) and, given the repetition of previously found OTUs, reduced the slope of the curve (Fig. 2).

All of the 16S rRNA gene sequences retrieved from the CO-4, CO-5, and CO-6 enrichments belonged to the family Clostridiaceae (Fig. 3). The great majority of these sequences belonged to cluster III clostridia and were closely related to Clostridium straminisolvens, Clostridium stercorarium, and Clostridium clariflavum. However, differences in the distribution of sequences across clostridial clusters were also observed between enrichments. All OTUs from the CO-4 enrichment grouped within cluster III, as did 7 out of 8 OTUs from the CO-5 enrichment. However, not only did the CO-6 enrichment have the largest number of detected OTUs, but they were spread across clusters I, III, and XIV and a novel cluster of which the novel clostridium Lutispora thermophila (29) is the main representative.

FIG. 1. Dynamics of residual dry weight of cellulose and product formation (A, C, and E) and changes in pH and total pellet N levels (B, D, and F) for the CO-4 (A and B), CO-5 (C and D), and CO-6 (E and F) enrichment cultures during 3-g liter⁻¹ Avicel fermentations. (A, C, and E) Symbols: open squares, dry weight of cellulose; filled squares, acetate production; open circles, ethanol production; filled triangles, formate production. (B, D, and F) Symbols: open circles, changes in pH; filled squares, total pellet N. For the dry weight (of cellulose) and pellet N measurements, each data point represents the mean ± standard deviation calculated from triplicate samples.
*Clostridium straminisolvens*-like sequences were consistently found in all three enrichments, although in different proportions. The most abundant 16S rRNA gene sequences retrieved in the CO-5 and CO-6 enrichments accounted for 68% and 45% of each clone library, respectively, and were almost identical to *Clostridium straminisolvens*. On the other hand, the most abundant OTU in the CO-4 enrichment was most similar to *Clostridium clariflavum* (previously sp. EBR-02E-0045 [GenBank accession no. AB186359]), obtained from a thermophilic methanogenic reactor (28, 30). Underlining the differences between the CO-4 enrichment and the others, this particular sequence was not retrieved in the libraries constructed for the CO-5 and CO-6 enrichments.

In addition, a number of sequences grouped most closely to novel clostridia. Two OTUs from the CO-6 enrichment, representing 1% to 3% of this clone library each, were determined to be most similar to *Clostridium clariflavum* (previously sp. EBR-02E-0045 [GenBank accession no. AB186359]), obtained from a thermophilic methanogenic reactor (28, 30). Underlining the differences between the CO-4 enrichment and the others, this particular sequence was not retrieved in the libraries constructed for the CO-5 and CO-6 enrichments.

In addition, a number of sequences grouped most closely to novel clostridia. Two OTUs from the CO-6 enrichment, representing 1% to 3% of this clone library each, were determined to be most similar to the newly isolated *Lutispora thermophila*, which is a noncellulolytic organism from a thermophilic methanogenic reactor (29). Likewise, a minor component of the CO-5 enrichment was most closely related to a proposed novel clostridium within cluster XIV, *Clostridium islandicum* (GenBank accession no. EF088328). A larger number of sequences in the CO-6 enrichment (representing 1% to 24% of this particular clone library) did not have any close cultured representatives, although they fell within cluster III and seemed most similar to *C. stercorarium*.

**PCR amplification of glycosyl hydrolase family 48 gene sequences.** Primers targeting a structural region essential to the function of family 48 glycosyl hydrolases were prepared based on translated amino acid sequences of *C. thermocellum* and *C. cellulolyticum* controls. This region includes the acidic residue responsible for protonating the glycosidic oxygen, identified as Glu87 in *C. thermocellum* CelS (10) and Glu55 in *Clostridium cellulolyticum* CelF (21); the Trp184, Asp255, and Tyr351 residues, associated with the water nucleophile involved in the catalytic mechanism of *Clostridium thermocellum* CelS (10); and the Asp230, Tyr299, Trp310, and Trp312 residues in the catalytic tunnel of *C. cellulolyticum* CelF (21).

An alignment of the PCR-amplified fragment with all the other sequences of GHF48 genes available in GenBank as of August 2009 and subsequent phylogenetic analyses revealed a grouping similar to the clustering observed in whole-gene alignments (Fig. 4). Sequences from the *Fungi* and the classes *Clostridia*, *Actinobacteria*, and *Bacilli* formed clear clusters, with the class *Clostridia* as the most divergent taxon. Known representatives of the *Gammaproteobacteria* (Hahella chejuensis) and the *Chloroflexi* (Herpetosiphon aurantiacus) each formed distinct, separate branches adjacent to the *Actinobacteria* cluster. A single exception to the class-based clustering was presented by the Cel48 sequence from Myxobacter sp. AL-1 (a deltaproteobacterium), which grouped with sequences from the *Bacilli*. It should also be noted that the GHF48 gene of *Ruminococcus albus* (cel48A) is distinctly different from those of all other glycosyl hydrolases from this family, including fellow clostridia and fellow member of the *Ruminococcaceae family Bacteroides cellulosolvens*.

Amplicons obtained from *C. thermocellum* ATCC 27405 and *C. thermocellum* DSM 1313 required further cloning in order to separate coamplified copies of the celY and celS genes prior to sequencing. For both strains, PCR amplification produced an even mixture (approximately 50:50) of the two genes, as observed in randomly selected clones used for amplification,
FIG. 3. Phylogenetic tree of 16S rRNA gene diversity in cellulolytic enrichments. Phylogenetic relationships were inferred by minimum evolution analysis of nucleotide sequences obtained from clone libraries from the CO-4 (open circles), CO-5 (filled squares), and CO6 (open triangles) enrichments. Bootstrap values are shown for 1,000 replicates. To indicate specific abundance, the percentage of each clone’s contribution to its library is specified in parentheses. The most abundant clones in each library are marked with asterisks.
FIG. 4. Phylogenetic tree of glycosyl hydrolase family 48 (GHF48) genes in cellulolytic enrichments. Phylogenetic relationships were inferred by minimum-evolution analysis of nucleotide sequences obtained from clone libraries from the CO-4 (filled circles), CO-5 (filled squares), and CO-6 (open triangles) enrichments. Controls using type cultures sequenced in-house are labeled with filled diamonds. Bootstrap values are shown for 1,000 replicates. To indicate specific abundance, the percentage of each clone’s contribution to its library is specified in parentheses.
thus demonstrating the ability of these primers to amplify both cellulosomal (celS) and noncellulosomal (celY) copies of GHF48 genes present in the same organism.

**Diversity of family 48 glycosyl hydrolases in enrichments.** All sequences retrieved from GHF48 gene libraries from enrichments belonged to the clostridia (Fig. 4). As a reference, we amplified the GHF48 genes in *Clostridium straminisolvens* and *Clostridium clariflavum*, given their prevalence in 16S rRNA libraries. Two distinctly different sequences obtained from *C. straminisolvens* were novel GHF48 gene fragment sequences, with 84% sequence similarity to *Clostridium thermocellum* celY and 89% sequence similarity to *Clostridium thermocellum* celS, respectively. Unlike *C. thermocellum*, where the celS/celY sequence retrieval ratio was close to 50:50, in *C. straminisolvens* the ratio favored celY-like sequences, with a celS/celY sequence retrieval ratio close to 30:70. Similarly, dominant sequences in the CO-5 and CO-6 enrichments (representing 70% to 84% of the clone libraries) were most similar to the novel *Clostridium straminisolvens* celY-like GHF48 gene reported here. The remaining GHF48 gene sequences from the CO-5 and CO6 enrichments, accounting for 16% and 30% of each clone library, were almost identical to the celS-like gene identified in *C. straminisolvens*. The dominant sequences from the CO-4 enrichment and a novel GHF48 gene sequence from *Clostridium clariflavum* grouped very closely together (>99% sequence similarity), with only 78% amino acid sequence similarity to their closest match found in *C. thermocellum* celY. Although they were retrieved in much lesser proportions, one sequence from the CO-4 enrichment was identical to *C. cellulolyticum* celF, and one was most closely related to *C. stercorarium* celY. Aside from the CO-4 GHF48 gene sequence that was identical to *C. cellulolyticum* celF, all the other GHF48 gene sequences reported here (including those of the *Clostridium straminisolvens* and *C. clariflavum* type cultures) represent novel cellulase genes.

**DISCUSSION**

Anaerobic enrichments from compost with cellulose as the sole carbon source facilitate the growth of consortia with various levels of community complexity and fermentation capabilities. By examining both the 16S rRNA and the glycosyl hydrolase family 48 gene composition, we have been able to characterize these communities and their dominant members at both the taxonomic and the functional level.

The breadth of 16S rRNA gene diversity in our enrichments, beyond the dominant sequences closely related to *Clostridium straminisolvens* and *C. clariflavum*, opens the question of what other functional roles are played by other clostridia within these communities. This is particularly true for enrichments resulting from multiple transfers such as those described here, where the presence of noncellulosolytic clostridia has not been diluted after several transfers with cellulose as the sole carbon source. Of particular interest are the uncultivated members with moderately high abundances (such as 16S rRNA gene clone CO6-40), where the isolation of pure cultures from a mixed consortium would enable further understanding of the different functional roles within both simple and complex cellulosolytic communities. Obtaining stable cellulosolytic enrichments with various levels of diversity and their taxonomic characterization are the first steps toward this goal. Additionally, the development of a molecular marker targeting an essential functional gene for these enrichments, as is the case for glycosyl hydrolase family 48 genes, further expands the possibilities for future comparisons between communities and community members from a transcriptomic point of view. However, it should be noted that fermentation experiments revealed subtle yet important differences in product and cell biomass formation. The CO-4 enrichment, dominated by *C. clariflavum*-like sequences, produces comparatively larger amounts of formate with lower overall cell biomass production and is able to continue cellulose utilization beyond pH 7.0 to a final pH of 6.5. This is consistent with previous observations for *C. clariflavum* DSM 19732, where formate production is evident, in contrast to the findings for its closest relatives, *C. thermocellum* and *C. straminisolvens* (30). On the other hand, the more diverse CO-5 (9 OTUs) and CO-6 (18 OTUs) enrichments are characterized by similar product profiles dominated by acetate and ethanol, higher cell biomass formation, and a final pH of 7.0, indicating that the *C. straminisolvens* dominance, and not the total diversity or community structure, may be an important factor in the ability of these communities to utilize cellulose.

Glycosyl hydrolase family 48 genes appear to be promising targets as functional marker genes for environments in which active anaerobic cellulose degradation occurs. Our work confirms the consistency in GHF48 gene taxonomic grouping and demonstrates substantial scope for expanding known exoglucanase diversity. The phylogenetic clustering observed with the primers used in this study, as with whole GHF48 gene sequences, revealed clear taxonomic clustering for representative members of the Clostridia, Actinobacteria, Bacilli, Gammaproteobacteria, Chloroflexi, and Bacteroidetes (Fig. 4). The exceptional grouping of Myxobacter sp. AL-1, a deltaproteobacterium, with sequences from the Bacilli is interesting from an evolutionary point of view but is not unexpected. Similarities between Myxobacter sp. AL-1 and different members of the Bacilli (in particular *Bacillus licheniformis*), in terms of unique structural similarities in family 48 glycosyl hydrolases (25) but also in terms of sequence similarities of other types of cellulases (22), have been reported previously.

Analysis based on both 16S rRNA and GHF48 gene sequences indicated that *Clostridium straminisolvens* and *Clostridium clariflavum*-like organisms were major components in the microbial communities present in the enrichments studied. This paper is the first report of *C. straminisolvens* and *C. clariflavum* possessing glycosyl hydrolase family 48 genes, and *C. straminisolvens* is the first organism after *C. thermocellum* to possess two distinctly different glycosyl hydrolase family 48 genes (NCBI accession no. GQ265349 and GQ487568). We also found that these genes are closely associated with the noncellulosomal celY gene and the cellulosomal celS gene in *C. thermocellum*. Given their close association with *C. thermocellum* counterparts, we speculate that the CelY-like cellulase in *C. straminisolvens* may act as a noncellulosomal exoglucanase in synergy with glycosyl hydrolase family 9 endoglucanases, as has been reported for *C. thermocellum* and *C. stercorarium* (2). Likewise, the presence of a CelY-like gene in *C. clariflavum* may indicate a similar mechanism. Further exploration of adjacent domains will be useful not only for determining the presence of cellulose-binding
domains and modes of action but also for gaining a better understanding of how conserved this family might actually be, and whether horizontal gene transfer plays a role in the observed diversity in GHF48 genes from environmental strains.

A variety of novel GHF48 gene sequences have been retrieved from the cellulolytic enrichments described in this study. Aside from the dominant C. straminislovens and C. clari-flavum sequences, two sequences associated with C. cellulosolyticum and C. stercorarium were also retrieved from cellulolytic enrichments and may belong to minor cellulolytic members of the community. It should, however, be noted that the mere presence of this gene does not necessarily confer functionality. For example, a GHF48 gene present in C. acetobutylicum, although it is expressed, does not confer cellulolytic activity on this organism (24).

With a variety of metagenomic studies addressing the diversity of glycolyse hydrodases in different environments (3, 9, 27), only one previous effort has targeted the diversity of cellulases beyond cultivated representatives by focusing on a specific family, describing the diversity and abundance of GHF5 in aquatic environments (8). More quantitative analyses of the relevance of GHF48 genes in cellulolytic microbial communities are needed, given the low abundance of these particular cellulases reported in metagenomic studies of complex cellulolytic microbial communities (3), although it is possible that their presence in mostly single copies per genome has contributed to the lack of detection in metagenomic studies. Given their prevalence in transcriptomic and proteomic analyses of model organisms (23, 32), it should be of interest to observe what metatranscriptomic analyses of both complex and simple cellulolytic communities reveal. The development of molecular tools for the detection of GHF48 genes in enrichments from environmental samples, as described here, offers an initial step toward this goal.

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