Glycoside Hydrolase Activities of Thermophilic Bacterial Consortia Adapted to Switchgrass††

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Industrial-scale biofuel production requires robust enzymatic cocktails to produce fermentable sugars from lignocellulosic biomass. Thermophilic bacterial consortia are a potential source of cellulases and hemicellulases adapted to harsher reaction conditions than commercial fungal enzymes. Compost-derived microbial consortia were adapted to switchgrass at 60°C to develop thermophilic biomass-degrading consortia for detailed studies. Microbial community analysis using small-subunit rRNA gene amplification followed by pyrosequencing and short-read metagenomic sequencing demonstrated that thermophilic adaptation to switchgrass resulted in low-diversity bacterial consortia with a high abundance of bacteria related to thermophilic paenibacilli, Rhodothermus marinus, and Thermus thermophilus. At lower abundance, thermophilic Chloroflexi and an uncultivated lineage of the Gemmatimonadetes phylum were observed. Supernatants isolated from these consortia had high levels of xylanase and endoglucanase activities. Compared to commercial enzyme preparations, the endoglucanase enzymes had a higher thermostolerance and were more stable in the presence of 1-ethyl-3-methylimidazolium acetate ([C2mim][OAc]), an ionic liquid used for biomass pretreatment. The supernatants were used to saccharify [C2mim][OAc]-pretreated switchgrass at elevated temperatures (up to 80°C), demonstrating that these consortia are an excellent source of enzymes for the development of enzymatic cocktails tailored to more extreme reaction conditions.

Enzyme cocktails that hydrolyze plant cell wall polysaccharides are a critical component of bioprocessing configurations designed to transform lignocellulosic biomass into biofuels (11, 23). The large variety of potential biomass feedstocks and pretreatments available require tailored glycoside hydrolase cocktails that function optimally under diverse conditions, including high temperatures, extreme pH, and the presence of residual pretreatment chemicals and inhibitors (12, 24). Current commercial cocktails consist of preparations of fungus-derived glycoside hydrolases, primarily cellulases and hemicellulases (3, 17). However, fungal enzymes are often deactivated by elevated temperatures or by residual chemicals from pretreatment (3). For example, ionic liquids (ILs), such as 1-ethyl-3-methylimidazolium acetate ([C2mim][OAc]), can dissolve lignocellulosic biomass and dramatically improve cellulose hydrolysis kinetics (10, 21, 27), yet multiple studies have shown that fungal endoglucanases are deactivated at low levels of ILs that may persist in the biomass after pretreatment (4, 32). In contrast, thermophilic bacterial and archaeal endoglucanases have been shown to be active in >20% [C2mim][OAc], suggesting that thermophilic prokaryotes may be an important source of enzymes for the development of more robust enzyme cocktails (5).

Aerobic, thermophilic bacteria are a potentially rich source of glycoside hydrolases for biomass decomposition. However, these bacteria generally secrete low levels of glycoside hydrolases, especially cellulolytic enzymes (23). Recent efforts to identify these enzymes have involved functional screening of expression libraries or bioinformatic homology-based searches of sequences derived from isolated organisms or environmental samples (1, 34). These approaches have significant limitations: expression libraries may miss relevant genes due to low representation within the library and poor expression in laboratory host strains, while bioinformatic approaches suffer mainly from the limited ability to predict the specific characteristics of an enzyme (activity, thermostability, etc.), which is often based on homology to known enzymes (28). The complexity of microbial communities in environmental samples also hampers bioinformatic enzyme discovery efforts, because it often prevents assembly of full-length genes from metagenomic databases (1, 6). Enrichment cultures on lignocellulosic biomass provide a method to reduce the complexity of the microbial communities and provide more tractable samples for
TABLE 1. Switchgrass-adapted communities

<table>
<thead>
<tr>
<th>Culture ID</th>
<th>Feedstock (% wt/vol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>JP-9, 6% SG</td>
<td>Switchgrass (6)</td>
</tr>
<tr>
<td>JP-9, 1% SG</td>
<td>Switchgrass (1)</td>
</tr>
<tr>
<td>JP-9, SGCMC</td>
<td>Switchgrass (1), carboxymethylcellulose (0.5%)</td>
</tr>
<tr>
<td>Z-9, 6% SG</td>
<td>Switchgrass (6)</td>
</tr>
<tr>
<td>Z-9, 1% SG</td>
<td>Switchgrass (1)</td>
</tr>
<tr>
<td>Z-9, SGCMC</td>
<td>Switchgrass (1), carboxymethylcellulose (0.5%)</td>
</tr>
</tbody>
</table>

* The Culture ID represents the final culture used for analysis of microbial community composition and glycoside hydrolase activities.
* Enrichments with 1% switchgrass were inoculated with 6% enrichment after the seventh 2-week passage of the enrichment.
* The enrichment with 1% switchgrass/0.5% switchgrass was inoculated with the 6% enrichment after the eighth passage.

Materials and Methods

Environmental samples. The compost inocula for switchgrass-adapted cultures were collected from two municipal green waste composting facilities. The green waste consisted of yard trimmings and discarded food waste. The first facility was Grover Soil Solutions located in Zamora, CA. Samples collected from this site are referred to as Zamora (Z) and were collected as previously described (1). The second facility was Jepson Prairie Organics located in Vacaville, CA. Samples collected from this site are referred to as Jepson Prairie (JP). This facility composes municipal green waste in watered and turned windrows. Compost was collected from windrows in the mesophilic (7 days) and thermophilic (30 and 60 days) composting stages. A spade was used to remove the top 12 in. of each windrow, and the exposed biomass was placed into 50-ml Falcon tubes, transported at room temperature, and frozen at −80°C within 2 h.

Cultivation conditions. Compost microbial communities were adapted to ground switchgrass (Panicum virgatum L.) for their sole carbon source by passing the community through nine liquid cultures (Table 1). Chemical characteristics of the switchgrass cultivar has been previously described (21). The switchgrass was exhaustively extracted with water and ethanol in a Soxhlet apparatus to remove soluble sugars and other nutrients and then dried at 50°C prior to use. Detailed procedures for the enrichments are described in the supplemental material.

Isolation of culture supernatants. The final cultures (passage 9) were used for glycoside hydrolase assays, and all passages were prepared for DNA isolation. The culture supernatant was clarified by decanting 30 ml of the culture supernatant into several 2-ml centrifuge tubes and spinning at 21,000 × g for 10 min. The supernatant was removed, and the pellets from four tubes consisting of the switchgrass-adapted microbial community and particles of switchgrass were combined, transferred into 2-ml lysis matrix D tubes (Qiogene, Montreal, QC, Canada), and frozen at −80°C for DNA extraction. The clarified supernatants were pooled in 50-ml Falcon tubes and passed through a 0.2-μm filter; this supernatant was used directly for measuring glycoside hydrolysis activity. For zymography, contaminating lignin-derived phenolic compounds were removed from the clarified supernatant by adding polyethyleneimine to a final concentration of 0.1% to 1 ml of supernatant, shaking for 2 h at 4°C, and centrifuging at 10,000 × g for 20 min at 4°C. Small-subunit (SSU) rRNA amplicon pyrosequencing. DNA was isolated from the pellets generated during isolation of culture supernatants described above. DNA isolation and sequencing were performed as previously described (6). Sequencing tags were quality trimmed and analyzed using the pyroclust version of the software tool PyroTagger (http://pyrotagger.rgj-pf.org) with a 220-bp sequence length threshold and an accuracy of 10% for low-quality bases (9, 19). All singleton operational taxonomic units (OTUs) were removed from the data set to reduce noise in the statistical analysis. The raw pyrotags reads are available in the NCBI Short Read Archive (SRA030499, SRA030513, and SRA030539).

Illumina sequencing and EMIRGE reconstruction of SSU rRNA genes. Illumina libraries were constructed and sequenced from DNA isolated from Z-9 1% SG and JP-9 1% SG using previously described protocols (25). Libraries of paired-end 76-bp reads were created for JP-9 1% SG (4.8 GB) and Z-9 1% SG (1.3 GB). Near-full-length ribosomal small-subunit genes were reconstructed from Illumina sequencing reads by using EMIRGE (25). For both communities, all reads were trimmed from the 3′ end until a base with a quality score of ≥3 was encountered. Paired-end reads where both reads were at least 60 nucleotides in length after trimming were used as inputs to EMIRGE, with the reference database previously described (25). Data from each community were processed separately, for 80 iterations, and SSU sequences, 23 in total, with relative abundances of >1% were kept for further analysis. Chimera check with Bellerophon (greengenes.lbl.gov) and manual analysis of EMIRGE-derived sequences excluded two of these sequences at 1.1% and 2.7% abundance, respectively.

Phylogenetic tree construction. Maximum likelihood trees were built with RAxML (30), using the GTR+ model of nucleotide substitution and 100 bootstrap-replicated sequences. Sequences were first aligned with Muscle (8) using default parameters, and columns in the full alignment with gaps were removed. An alignment for tree construction and pairwise percent identity calculations. For Fig. S1 in the supplemental material, the alignment was manually edited to span the region covered by pyrotag sequencing, and columns in the alignment with a majority of gaps were removed. Methanolalcaloicosus jannaschii (GenBank M59126.1) was used as the outgroup to root the trees.

Glycoside hydrolase activity assays. The glycoside hydrolase activities present in each switchgrass-adapted supernatant were measured using the following enzymes: p-nitrophenyl (pNP) (β-D-glucosidase, cellobiohydrolase, β-D-xylanase, and α-L-arabinofuranosidase) and the dinitrosalicylic acid (DNS) reducing sugar assay (endoglucanase, xylanase) and the p-nitrophenol (pNP) assay (β-D-glucosidase, cellobiohydrolase, β-D-xylanase, and α-L-arabinofuranosidase) (5, 29). Heat-killed samples generated by heating the supernatant to 95°C for 16 h were used as blanks. Activity units for all assays were calculated as μmol of sugar liberated min⁻¹ ml⁻¹ and reported as U/ml. The endoglucanase and xylanase activities of the JP/Z-9 1% SG and JP/Z-9 SGCMC supernatants were also measured using zymography with SDS-PAGE gels embedded with carboxymethylcellulose (CMC) and olate spelt sylan. Detailed procedures are described in the supplemental material.

Saccharification of IL-pretreated switchgrass. The supernatant from the JP-9 1% SG switchgrass-adapted community was tested for its ability to saccharify oniconic liquid-pretreated switchgrass. The switchgrass was pretreated by dissolution in [C2min][OAc] at 140°C for 3 h and precipitation with an acetone-electrolyte mixture. The precipitated material was successively washed with ethanol and water to remove residual lignin (D. C. Davidson, unpublished observation). Duplicate 10-ml saccharification reaction mixtures were set up with 250 mg of IL-pretreated switchgrass. The JP-9 1% SG reaction mixture consisted of 9.5 ml of 1% NaOAc mixture with 0.5 ml of 1 M NaOAc (pH 5.0). For comparison, commercial enzyme preparations containing both cellulase and xylanase activities were mixed using recommended enzyme/glucon content of biomass (wt/wt) loadings: 8.32 μl of NS85013 (1% wt/wt) and 0.832 μl each of NSS0100 and NSS0300 (0.1% [wt/wt]) were added to 10 ml pretreated 100 mM NaOAc (pH 5.0) buffer. Smaller amounts of enzymes were used to test thermotolerance and IL tolerance due to limited amounts of the JP-9 1% SG supernatant. Lyophilized supernatant (1 ml) was resuspended in 1 ml of 100 mM NaOAc (pH 5.0) and added to 9 ml of the same prewarmed buffer. The Novozymes enzyme preparations were adjusted to match more closely the endoglucanase and xylanase activities of the JP-9 1% SG supernatant: 0.25 μl of the cellulase mix (8.32 μl of NS85013 and 0.832 μl of NSS0100) and 2.5 μl of the xylanase NSS0300 were added to 10 ml prewarmed buffer. For IL tolerance, the reaction buffer used was 100 mM NaOAc−15% [C2min][OAc] at pH 5.0. All samples were incubated at 72 h at 70°C or 180°C, and 150 μl was withdrawn for each time point and frozen at −20°C. Time point samples were then spun at 21,000 × g for 5 min at 4°C, and 5 μl was added to 55 μl water and 60 μl DNS reagent. Samples were incubated at 95°C for 5 min, and the absorbance at 540 nm was taken. A background subtraction blank was made by adding 5 μl of each sample to 115 μl of water. The total sugars were calculated by comparing to a standard curve of glucose. The percent total sugar was calculated using the estimated glucon and xylan contents of switchgrass (24% and 33%, respectively).

Chemicals. All chemicals were reagent grade and were obtained from Sigma (St. Louis, MO) unless otherwise noted and used as received.
TABLE 2. Comparison of estimated abundances for pyrotag and metagenomic sequencing for JP-9 1% SG and Z-9 1% SG

<table>
<thead>
<tr>
<th>Group</th>
<th>JP-9 1% SG estimated abundance (%)</th>
<th>Z-9 1% SG estimated abundance (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyrotags</td>
<td>40.4</td>
<td>20.4</td>
</tr>
<tr>
<td>EMIRGE</td>
<td>20.4</td>
<td>69.7</td>
</tr>
<tr>
<td>Thermophilic paenibacillib</td>
<td>40.4</td>
<td>20.4</td>
</tr>
<tr>
<td>Thermus thermophilus</td>
<td>39.3</td>
<td>49.0</td>
</tr>
<tr>
<td>Rhodothermus marinus</td>
<td>9.62</td>
<td>10.4</td>
</tr>
<tr>
<td>Gemm-5</td>
<td>2.05</td>
<td>7.70</td>
</tr>
<tr>
<td>Thermorhodospirillum sp.</td>
<td>1.18</td>
<td>&lt;1.00</td>
</tr>
<tr>
<td>Thermobacillus terraeunum</td>
<td>1.43</td>
<td>1.40</td>
</tr>
<tr>
<td>Sphaerobacter thermophilus</td>
<td>&lt;1.00</td>
<td>1.16</td>
</tr>
<tr>
<td>Trueperacae</td>
<td>&lt;1.00</td>
<td>&lt;1.00</td>
</tr>
</tbody>
</table>

| a Groups with >1% estimated abundance in pyrotag- or EMIRGE-derived sequences are depicted in the table. Total estimated abundances of these depicted groups were 90 to 95% of the total community. |
| b Pyrotag abundances are represented by the sums of the abundances for pyrotag clusters related to Paenibacillus sp. strain SAEN-407, Paenibacillus kobensis DSM 10249, Paenibacillus D273a, and Thermobacillus sp. strain KCW4. EMIRGE-derived sequence abundance values are the sums of individual sequences presented in Fig. 3 that clustered with the Paenibacillaceae family (JP 211, JP 261, JP 2339, JP 2453, JP 2539, Z 19, Z51, Z 137, Z 146, Z 261, Z 484, Z 1266, and Z 1300). The individual estimated abundances of each of these sequences are listed in the phylogenetic tree in Fig. S1 of the supplemental material. |

RESULTS

Enrichment of thermophilic bacterial consortia on switchgrass. Enrichment cultures with inocula sourced from two compost facilities in northern California were adapted to switchgrass (6% [wt/vol]) through multiple passages at 60°C. The compost inocula were complex communities composed of 259 taxa for the JP enrichment and 593 taxa for the Z enrichment. A significant reduction in taxonomic diversity was observed in the initial adaptation to switchgrass at 60°C. This reduction in complexity was demonstrated by a reduction in the Shannon diversity index from 3.87 in the initial JP inoculum to 2.04 in the first JP enrichment and from 4.42 to 2.28 for the first Z enrichment (data not shown). These low-diversity communities were maintained through nine passages on switchgrass. Additional cultures were generated at later passages by lowering switchgrass loadings to 1% or by amending 1% switchgrass cultures with 0.5% CMC, affording six independent samples for microbial community analysis and glycoside hydrolyase measurements (Table 1).

Comparison of SSU rRNA amplicons recovered from the final cultures at the level of individual OTUs (97% identity) demonstrated that the enrichment cultures were dominated by only a few phylotypes (Table 2; Fig. 1A and B) (9). Despite the different inocula from which they were derived, the JP and Z enrichment cultures had remarkably similar community profiles. The most abundant sequences recovered from both the high- and low-biomass JP and Z enrichments were closely related to thermophilic Gram-positive Firmicutes of the Paenibacillaceae family (Thermobacillus, Paenibacillus) and Rhodothermus marinus, a known biomass-deconstructing member of the Bacteroidetes. Sequences related to thermophilic Chloroflexi (Thermobaculum, Sphaerobacter, Thermomicrobium) were recovered at lower abundances, as were sequences related to an uncultivated lineage, Gemm-5, in the Gemmatimonadetes phylum. A significant difference between the JP and Z enrichments was the prominence of bacteria closely related to Thermus thermophilus in the JP enrichments that was nearly absent in the Z enrichments. However, in the Z enrichments, sequences related to the family Trueperaceae in the Deinococcus/ Thermi phylum were recovered. When CMC was added as a cosubstrate, the proportion of amplicons related to Rhodothermus marinus increased in both the JP and Z cultures (Fig. 1A and B).

Near-full-length SSU rRNA sequences (1,200 to 1,600 bp) were reconstructed at 97% identity from metagenomic data obtained for JP-9 1% SG and Z-9 1% SG enrichment cultures. These sequences were reconstructed by the EMIRGE method, which uses an expectation maximization algorithm to reconstruct SSU rRNA gene sequences from Illumina metagenomic sequencing data (25). Comparison of the near-full-length SSU sequences with a maximum likelihood tree confirmed that the two enrichments contained closely related communities (Fig. 2). For most members of the enrichment cultures, the reconstructed full-length SSU sequences contained identical ~200-bp segments to a corresponding representative pyrotag cluster sequence, validating the ability of the EMIRGE method to reconstruct SSU genes (see Fig. S1 in the supplemental material).

The EMIRGE method also predicts the relative abundances of microbes in mixed consortia by probabilistically measuring the relative proportions of reads recruited to each reconstructed SSU rRNA gene sequence (25). For the JP and Z enrichments, EMIRGE abundance estimates showed good general concordance with abundance estimates made from pyrotag sequencing (Table 2). However, the estimated abundance of the thermophilic paenibacilli was lower for the full-length sequences than the pyrotags, and for the JP enrichment, the abundance of the sequences related to Thermus thermophilus was higher than with the pyrotags. For both enrichments, the estimated abundance of the Gemmatimonadetes was higher than predicted by pyrotag abundance, and Trueperaceae were predicted at higher abundance in the Z enrichment by EMIRGE.

Glycoside hydrolyase activities in switchgrass-adapted cultures. Comparison of endoglucanase and xylanase activities at 70°C and pH 5.0 in supernatants obtained from the JP/Z 6% SG and JP/Z 1% SG cultures indicated that more activity was recovered from the 1% switchgrass cultures, so the activity profiles of these supernatants were studied in detail. The supernatants displayed significant enzymatic activities in standard assays, including endo-xoglucanase, β-glucosidase, endoxylanase, β-xylanidase, and α-1,5-arabinofuranosidase activities (Fig. 3A and B; see also Fig. S2 in the supplemental material). Endoglucanase and xylanase activities were compared to commercial cellulase and xylanase cocktails pro-
duced by Novozymes that were diluted to comparable activity levels.

Temperature profiles of endoglucanase activity demonstrate that these supernatants possess more thermostable/active enzymes than the commercial cocktails from Novozymes (Fig. 4A). Higher optimum temperatures (T_{opt}) were observed for the endoglucanases recovered from the 1% switchgrass enrichments than the Novozymes enzyme preparations (T_{opt}, 80°C versus 60°C). These supernatants even retained endoglucanase activity (15 to 50%) at 99°C. Their endoglucanase enzymes also exhibited high levels of tolerance to the ionic liquid [C2mim][OAc], retaining ~50% activity in the presence of 30% [C2mim][OAc], while the Novozymes endoglucanase cocktail was essentially inactive at 10% [C2mim][OAc] (Fig. 4B)

FIG. 1. Plot of the relative abundance of dominant taxa (≥1%) based on SSU rRNA amplicon pyrosequencing for the switchgrass-adapted enrichments. (A) Jepson Prairie enrichments (JP 6% SG, JP 1% SG, and JP SGCMC); (B) Zamora enrichments (Z 6% SG, Z 1% SG, Z SGCMC). A cutoff of 1% abundance was chosen to highlight the most abundant organisms present in the community. OTUs that were assigned the same name by Pyrotagger are distinguished by adding the pyrotagger cluster number after the name. GenBank accessions numbers for each cluster assigned by Pyrotagger are included in Table S1A (JP) and B (Z) in the supplemental material. Amplicon library sizes: JP 6% SG, 2,004; JP 1% SG, 2,107; JP SGCMC, 1,496; Z 6% SG, 3,095; Z 1% SG, 3,095; Z SGCMC, 2,310.
In contrast, xylanase activities of the culture supernatants exhibited only a slightly higher thermostolerance and tolerance to [C2mim][OAc] than the Novozymes xylanase preparation (see Fig. S3 and S4 in the supplemental material).

Zymograms with CMC and xylan as substrates were performed on the JP and Z 1% SG supernatants (see Fig. S5 in the supplemental material). More than 10 bands were observed for each activity, indicating that the enriched consortia each produced multiple endoglucanase and xylanase enzymes. Comparison of the JP and Z samples indicated that the active enzyme complements in both samples were remarkably similar, suggesting that similar communities secrete the endoglucanase and xylanase enzymes. Increased intensities of some of the endoglucanase bands were observed in the cultures amended with CMC, consistent with the observed increase in endoglucanase activity (Fig. 3).

Glycoside hydrolase activities on ionic liquid-pretreated switchgrass. Although the supernatants obtained from the switchgrass-adapted communities performed well on model biomass substrates, a more stringent test of their utility is their ability to efficiently deconstruct pretreated biomass. The supernatant obtained from the JP-9 1% SG enrichment performed well on model substrates and was selected to saccharify switchgrass pretreated with [C2mim][OAc]. For comparison, Novozymes enzyme preparations were mixed at the recommended enzyme/biomass loadings (wt/wt), which amounted to approximately 15× the endoglucanase and 0.25× the xylanase activity compared to the JP-9 1% SG supernatant at 70°C. Despite the comparatively low endoglucanase activity in the JP-9 1% SG supernatant, both cocktails liberated virtually all the sugars from the [C2mim][OAc]-pretreated switchgrass at 70°C after 72 h (Fig. 5A).
The enzyme cocktails were then compared at a higher temperature (80°C) or in 15% [C2mim][OAc] at 70°C (Fig. 5B and C). Due to limited amounts of sample, the JP-9 1% SG supernatant was diluted 10-fold in these saccharifications. The Novozymes cellulase and xylanase cocktails were used as positive controls. The Novozymes NS5003-10 and NS50030 enzymes were diluted 1:1,000 and 1:10,000, respectively, concentrations that are similar to those used to saccharify biomass at 2.5% (wt/vol) loadings (see Fig. 5A). Endoxylanase values reported were extrapolated from 1:10 dilutions of supernatant samples.

**DISCUSSION**

We found that growing compost microbial communities on extracted switchgrass under thermophilic conditions generated simplified bacterial consortia that produced glycoside hydrolase enzymes that are more stable than commercial fungal cocktails when assayed on pretreated switchgrass at high temperature and in the presence of pretreatment chemicals. Adaptive cultivation on other feedstocks, such as lignin, cellulose, and CMC, has also generated simplified communities and, along with this study, indicates that this method is a useful tool for developing simplified biomass-degrading consortia tailored to deconstruct a designated feedstock under defined conditions, such as temperature or pH (6, 22, 26).

Both the JP and Z enrichments converged to closely related consortia with similar microbial community compositions and levels of secreted glycoside hydrolase activities. Perturbation of the enriched communities by inoculation into cultures with low biomass (1% switchgrass) and amendment with CMC did not significantly alter the community composition of the enrichments, although the glycoside hydrolase activities measured in the supernatant were higher. This observation indicates that biomass loading is a critical component of enrichment cultiva-
tion studies and can be adjusted to maximize titers of enzymes of interest without dramatically affecting the community composition.

The success of this enrichment strategy was demonstrated by the selection of biomass-deconstructing bacteria. Analysis of amplicon pyrosequencing data and reconstructed SSU rRNA genes from metagenomic sequencing demonstrated that thermophilic paenibacilli and *Rhodothermus* groups were abundant in both enrichments. Both of these groups have cultured relatives known to degrade biomass, suggesting that they are the...
source of many of the glycoside hydrolases present in the culture supernatants (2, 31, 35). Additionally, compost and swine waste-derived microbial communities enriched on microcrystalline cellulose at elevated temperatures under aerobic conditions also contain thermophilic paenibacilli, suggesting that these microbes are critical components of biomass deconstruction under thermophilic aerobic conditions (22, 26). Bacteria closely related to *Thermus thermophilus* were dominant members of the JP enrichment but were absent in the Z enrichment. This observation suggests that *T. thermophilus* may not be involved in biomass deconstruction, as the glycoside hydrolase activities and the CMC/xylan zymography are very similar in both enrichments. However, *T. thermophilus* strains have been isolated from hot composts (60 to 80°C) and shown to express highly active xylanases (14). Detailed proteomic analysis of the secreted proteins in the supernatant and isolation of *T. thermophilus* strains from the enrichment cultures will resolve its role.

Members of the *Chloroflexi* phylum are ubiquitous in the enrichments and are of interest, as the *Chloroflexi* have relatively few cultivated and sequenced representatives (7, 15, 16, 36). Inspection of genomes available for three thermophilic type strains belonging to this phylum (*T. terrenum, Sphaerobacter thermophilus*, and *Thermomicrobium roseum*) indicates that they possess a number of cellulases and hemicellulases (20). Another prominent group present in both the JP and Z enrichments is a member of an uncultivated lineage (*Gemm-5*), in the *Gemmatinimonadetes* phylum (37), that is distantly related to the sole cultivated representative of this phylum, *Gemmatinonas aurantica* (SSU rRNA genes have 88% identity). Since both the *Chloroflexi* and *Gemmatinimonadetes* are commonly found in soils and sediments, sequencing these consortia will illuminate how they function in biomass deconstruction.

A significant amount of xylanase activity was recovered from supernatants isolated from the switchgrass-adapted consortia. The xylanase activity was comparable to highly productive *Bacillus* and *Paenibacillus* strains cultivated under optimized conditions, suggesting that mixed consortial cultivation on switchgrass is an effective method to generate high levels of xylanase enzymes (18). Cellulolytic activity was relatively low, and cultivations on pretreated switchgrass are being explored to enhance the cellulolytic activity of the recovered supernatants. Comparison of the endoglucanase activities between the culture supernatants and the Novozymes preparations suggests that the thermophilic bacteria secrete enzymes that are both more thermotolerant and more active in the presence of the ionic liquid [C2mim][OAc], a property that has been observed for purified enzymes (5). This correlation will be confirmed by purifying endoglucanases from the supernatant and performing assays on the purified proteins in the presence of [C2mim][OAc]. Interestingly, the thermotolerance and the [C2mim][OAc] tolerance of the xylanase activities of the culture supernatants and the Novozymes preparations were comparable, suggesting that the properties of the xylanases in the culture supernatants and the Novozymes xylanase preparation (NS50030) may be similar. Zymography revealed that the thermophilic consortia produce multiple enzymes for at least two of the glycoside hydrolase activities profiled. These enzymes are currently being identified by a combination of metagenomic sequencing and mass spectrometry-based proteomic measurements.

Both the Novozymes enzyme preparations and the culture supernatants were able to saccharify [C2mim][OAc]-pretreated switchgrass at 70°C. These observations indicate that both systems possess the necessary complement of glycoside hydrolase enzymes to release sugars from complex biomass. Surprisingly, the Novozymes preparations efficiently released sugars from pretreated switchgrass at 70°C, despite that temperature being 10°C above the cellulase optimum temperature and 20°C higher than the recommended reaction temperature (13). Perhaps this was due to the greatly enhanced hydrolysis kinetics of [C2mim][OAc]-pretreated switchgrass compared to acid-pretreated switchgrass, allowing for rapid polysaccharide hydrolysis before enzyme denaturation (21). However, at 80°C or in the presence of ionic liquids, the enhanced stability of the glycoside hydrolase enzymes from the thermophilic bacterial cultures was evident (Fig. 5B and C), demonstrating their value as sources of enzymes for enzymatic cocktails adapted to more extreme reaction conditions than fungal enzymes can tolerate.

In conclusion, this study demonstrates that thermophilic, aerobic microbial consortia cultivated from compost produce significant titers of glycoside hydrolase enzymes that can be directly compared to commercial fungal biomass-deconstructing cocktails. Microbial community analysis has demonstrated that these consortia are composed of a few dominant phyotypes that consist of both well-studied and novel biomass-deconstructing bacteria. Therefore, these consortia are amenable to detailed genomic and proteomic investigations, which will reveal the suite of bacterial glycoside hydrolases used to deconstruct complex biomass (33). This approach will allow characterization of new bacterial glycoside hydrolases and accessory proteins from uncultivated organisms that can enhance biomass deconstruction.

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