Glycan Degradation (GlyDeR) Analysis Predicts Mammalian Gut Microbiota Abundance and Host Diet-Specific Adaptations

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ABSTRACT Glycans form the primary nutritional source for microbes in the human gut, and understanding their metabolism is a critical yet understudied aspect of microbiome research. Here, we present a novel computational pipeline for modeling glycan degradation (GlyDeR) which predicts the glycan degradation potency of 10,000 reference glycans based on either genomic or metagenomic data. We first validated GlyDeR by comparing degradation profiles for genomes in the Human Microbiome Project against KEGG reaction annotations. Next, we applied GlyDeR to the analysis of human and mammalian gut microbial communities, which revealed that the glycan degradation potential of a community is strongly linked to host diet and can be used to predict diet with higher accuracy than sequence data alone. Finally, we show that a microbe’s glycan degradation potential is significantly correlated ($R = 0.46$) with its abundance, with even higher correlations for potential pathogens such as the class Clostridia ($R = 0.76$). GlyDeR therefore represents an important tool for advancing our understanding of bacterial metabolism in the gut and for the future development of more effective prebiotics for microbial community manipulation.

IMPORTANCE The increased availability of high-throughput sequencing data has positioned the gut microbiota as a major new focal point for biomedical research. However, despite the expenditure of huge efforts and resources, sequencing-based analysis of the microbiome has uncovered mostly associative relationships between human health and diet, rather than a causal, mechanistic one. In order to utilize the full potential of systems biology approaches, one must first characterize the metabolic requirements of gut bacteria, specifically, the degradation of glycans, which are their primary nutritional source. We developed a computational framework called GlyDeR for integrating expert knowledge along with high-throughput data to uncover important new relationships within glycan metabolism. GlyDeR analyzes particular bacterial (meta)genomes and predicts the potency by which they degrade a variety of different glycans. Based on GlyDeR, we found a clear connection between microbial glycan degradation and human diet, and we suggest a method for the rational design of novel prebiotics.
branched-chain fatty acids (17), the detailed growth requirements for the majority of gut bacteria remain unknown (18). Characterizing these requirements will shed light on the different metabolic niches organisms fill and may enable the design of dietary interventions that promote growth of particular beneficial microbes, an approach collectively termed “prebiotics” (19). While several glycans are currently marketed around the world as prebiotics, few have been validated through high-quality human trials (19, 20). Furthermore, dietary enrichment of a specific prebiotic compound may permit preferential expansion of a microbial group that is well adapted to its use, but the outcomes for the gut community as a whole can be unpredictable (21).

In this study, we investigated the connections between diet and glycan metabolism of the human gut microbiota. Whereas the study of the metabolic activity conducted by gut microbiota has been the focal point of a wide range of computational studies (9, 11), current approaches have been highly limited in their ability to analyze glycan degradation. We present a novel algorithm (termed GlyDeR) for predicting the glycan degradation patterns of any bacterium with a sequenced genome. The algorithm is based on manual curation of nearly 150 carbohydrate-active enzymes (CAZymes) and is applied to a set of 10,000 glycan structures and 203 microbial genomes. Given a particular bacterial (meta)genome, GlyDeR can be used to reverse engineer the predicted potency by which a bacterium degrades a variety of different glycans. These predictions correlate with known KEGG reactions and expand upon the limited, previously available glycan degradation data 100-fold. We determined that the microbiota of herbivores and carnivores have stronger degradation affinities for plant-derived and animal-derived glycans, respectively, and that a Western diet in humans correlates more strongly with meat-derived glycans than a non-Western diet. Finally, we show that species-specific glycan degradation profiles are associated with and can be used to predict bacterial species abundance, making GlyDeR a valuable tool for the future rational design of novel prebiotics, by deliberately manipulating the microbiome based on nutrient availability.

RESULTS

Construction of the glycan degradation (GlyDeR) pipeline. Although the exact biochemistry of glycan degradation is missing from all currently available databases, considerable knowledge is embedded in the descriptions of the CAZymes that catalyze these degradation reactions and is typically represented by enzymatic commission (EC) numbers. We leveraged this knowledge to develop a new computational pipeline that uses enzymatic and structural data sources to predict the degradation of every glycan in the KEGG database (22) that has a sequenced (meta)genome. That is, given (meta)genomic data as input, GlyDeR yields phenotypic (glycan degradation) data as output. The construction of the pipeline comprises two steps. (i) The first step relies on a novel algorithm that we developed, which we term glycan degradation (GlyDeR). The algorithm takes as input a manually curated annotation of all the reactions that each known CAZyme is capable of performing (see Table S1 in the supplemental material) and a network representation of all the glycans in KEGG, in which the nodes are the monosaccharides and the edges are the glycosidic linkages (Fig. 1a; see also Table S6 in the supplemental material). We converted the CAZyme annotations to the computer-based rules that dictate their mechanism for breaking a given glycan into two subcomponents. The manual curation of this critical step was done using the help of experts with knowledge of the biochemistry of glycan metabolism. GlyDeR then executes these rules recursively on all the glycans, to generate 141,561 GlyDeR reactions, each linking a specific enzyme to a glycan substrate and its products (an example reaction is given in Fig. 1a, and a more detailed explanation is provided in Materials and Methods). (ii) In the second step, GlyDeR reactions are mapped back to CAZymes in order to produce a table where the rows are CAZymes, the columns are glycans, and each entry contains a CAZyme score for CAZyme i and glycan j. If CAZyme i is unable to break glycan j, then the score is 0, otherwise the score is calculated as follows:

$$\text{CAZyme score}_{ij} = \frac{1}{g_i}$$

where $g_i$ is the number of glycans that are broken by CAZyme i. The entire construction process is summarized in Fig. 1b. A CAZymes table that contains all of the CAZyme scores can be found in Table S5 in the supplemental material.

Use of the GlyDeR pipeline. Microbial (meta)genomes were annotated for CAZymes by using BLAST analysis (23) against three reference databases: the Carbohydrate-Active Enzymes (CAZy) database (24), the Seed-RAST annotation (25), and KEGG (26) (see Materials and Methods). Then, CAZyme scores were assigned to genes, and a GlyDeR score was calculated for each glycan i and (meta-) genome j as follows:

$$\text{GlyDeR score}_{ij} = \sum \text{CAZyme score}_{ij}, \forall n_{ij}$$

where $n_{ij}$ is the number of genes in (meta)genome j which translate to a CAZyme that can break glycan i.

The GlyDeR score represents the predicted potency with which the glycan can be degraded by that (meta)genome, taking into account how many CAZymes can degrade the glycan and with decrements for the score of a promiscuous enzymes with low specificities (see Materials and Methods). For example, an organism that contains three enzymes that degrade maltotetraose, each of which also degrades four other glycans, would have a GlyDeR score of 3/5 (two examples are provided in Fig. 1c). The use of GlyDeR is captured in Fig. 1d.

Validating the GlyDeR pipeline. To assess the biological relevance of GlyDeR, we performed a cross-validation procedure that examined its consistency in capturing known degradation reactions in KEGG (see Materials and Methods). We found that the products of GlyDeR reactions were highly enriched with known rather than hypothetical glycans ($P = 10^{-19}$, hypergeometric test; see also Fig. S2a in the supplemental material). As further validation, we compared the predicted genome-specific GlyDeR scores of each bacterial strain with the glycans that, according to KEGG, the strain is able to break (KEGG glycans). Since the above information from KEGG was not used to construct the set of GlyDeR reactions, a circular argument was avoided. We found a significantly higher mean GlyDeR score for KEGG glycans across all strains compared to non-KEGG glycans (see Fig. S2a). Notably, our analysis produced GlyDeR scores for over 100 times the number of unique glycan degradation reactions that are currently reported in KEGG (116,388 versus 1,374), highlighting the limited scope of glycan metabolism information captured in the KEGG database.

Characterization of glycan degradation patterns across the major gut bacterial phyla. We first applied GlyDeR to a cohort of
203 reference gut microbial genomes retrieved from the Human Microbiome Project (HMP) (27). All of the available information on these strains is listed in Table S2 of the supplemental material. We used GlyDeR to study the extent to which different microbial phyla metabolize different glycans. Initially, we examined whether phylogenetic clusters are reflected in glycan degradation patterns. We therefore computed for each of the HMP genomes a GlyDeR profile, i.e., a vector of its GlyDeR scores for all glycans. The species-specific GlyDeR “signature” describes the potency by which a given species can catabolize each of the ~10,000 reference glycans in our database and hence provides an overall view of its glycan utilization capabilities. We then mapped each species to its respective phylum and performed principal coordinates analysis (PCoA) on the Bray-Curtis dissimilarities between the species GlyDeR profiles. This yielded clusters of phyla that were statistically distinct (multivariate analysis of variance [ANOVA] test, Wilke’s lambda < 0.001) (see Fig. S2b in the supplemental material). Still, there were apparent significant differences in the average glycan degradation capacities of genera belonging to a given phylum (see Fig. S2c).

**Bacteroides** species are highly effective degraders of animal-derived glycoproteins. Recently, it has been shown that human diets high in animal protein are associated with high levels of *Bacteroides*, whereas diets rich in plant-derived carbohydrates and very low in animal protein display enrichment for *Prevotella* (28–30). Among the phyla in the HMP data set, we found *Bacteroidetes* to be the most efficient degraders of animal-derived glycans (see Fig. S2e in the supplemental material). Notably, this trend was apparent for the *Bacteroides* genus but absent for *Prevotella*, which also belongs to that phylum (Fig. 2b). Furthermore, all 19 of the highest-scoring species with GlyDeR belonged to the *Bacteroidetes* genus (see Table S3 in the supplemental material), consistent with their known roles as primary glycan degraders in the gut (31–33).

Recent papers have also shown that glycans found in human milk, such as human milk oligosaccharides (HMOs), are utilized mainly by several *Bifidobacterium* and *Bacteroides* species (31, 34). According to GlyDeR, 21 out of the 23 HMP genomes that are capable of degrading HMOs belong to *Bacteroides* species (the other degraders were *Parabacteroides* sp. D13 and *Bifidobacterium bifidum*) (see Table S4 in the supplemental material). To further investigate this point, we examined two of the most abundant animal-derived glycoproteins in the human diet: ovalbumin (35) and casein (36). Indeed, we confirmed that members of *Bacteroides* degrade these glycoproteins more efficiently than any other genus (see Fig. S2f in the supplemental material). Given that many dietary animal glycans are derived from proteins (e.g., glycoproteins and proteoglycans), we propose that the high capability of *Bacteroides* to degrade animal glycans might explain why their abundance is increased in Westerners (30, 37).

**Glycan degradation patterns can be used to predict bacterial abundance.** We studied the relationship between the glycan degradation scores of a given bacterial taxon and its abundance in the gut. The GlyDeR platform. (a) A visual representation of the glycan degradation reaction performed for EC 3.2.1.115, breaking down Kojitriose into Kojibiose and glucose. (b) A schematic representation of the construction of the computational pipeline. Information is taken from multiple databases and analyzed as follows. Step 1 (lred arrow on left): by using CAZyme information and the GlyDeR algorithm, glycan degradation reactions are reconstructed. Step 2 (red arrow on right): a CAZyme table is constructed that represents the potency with which different CAZymes break different glycans. (c) GlyDeR score calculation. (Top) The organism has one enzyme (yellow PacMan) dedicated to the degradation of one glycan (purple); therefore, the GlyDeR score for the purple glycan equals 1. (Bottom) The organism has two enzymes capable of degrading 3 and 4 glycans, respectively, and therefore the GlyDeR score for the purple glycan equals 7/12. (d) GlyDeR utilization. (Meta)genomes are annotated for CAZymes by using CAZy, SEED, and KEGG databases, and with the CAZyme table a GlyDeR score can be calculated, reflecting the capacity of a (meta)genome to degrade a specific glycan.
We matched the abundance of 16S rRNA marker gene sequences from 325 human individual gut samples found in the HMP database with the aforementioned 203 microbial reference genomes (see Materials and Methods). For each taxon, we extracted 6 features that characterized its glycan degradation capacity, including plant-specific glycans, animal-specific glycans, disaccharides, oligosaccharides, short polysaccharides, and long polysaccharides (see Table S3 in the supplemental material). Each feature represents the sum of GlyDeR scores for the glycans that belong in the class. Based on these features, we built a linear regression model for the abundance of these taxa in the samples. In order to apply the linear regression model, we filtered out taxa that were not detected in any sample and taxa that were highly varied (see Materials and Methods for criteria), resulting in 48 predictable taxa for the analysis. This regression yielded a correlation coefficient of 0.46 (Fig. 2c), a score markedly higher than the correlation achieved using a model based on CAZyme abundances in a genome ($R = 0.11$). We next built similar regression models independently for each class of bacteria. Remarkably, the Clostridia class had the highest combination of $R$ (0.76) and $P$ (0.0001) values; other classes with significantly predictive models were Bacteroidia and Fusobacteria (Fig. 2d; see also Table S8 in the supplemental material). These results suggest that glycan supplements can be tailored to control certain species abundances, especially those of potentially pathogenic Clostridia.

In an effort to include taxa that were initially omitted in the analysis above due to their high levels of variation, we first clustered the HMP samples according to their 16S rRNA data into 2 main groups by using KMeans (see Materials and Methods), and we recalculated the average taxon abundance separately for each cluster. The same procedure for building predictors of bacterial taxon abundance based on their genome-specific GlyDeR features was then used, and this yielded 53 predictable taxa in the first cluster and 71 predictable taxa in the second cluster, with concomitant increases in the correlation coefficients (0.51 and 0.57, respectively). Based on the two clusters, we assembled a list of 25 strains with highly predictable abundances (see Table S9 in the supplemental material), and with only one exception, all the
strains belonged either to the *Bacteroidia* or *Clostridia* classes (see Discussion). Notably, based on the regression formulas of all three models, the degradation capacity of long polysaccharides had the highest effect on bacterial abundance. It is therefore likely that because most long polysaccharides are not digested by the human host prior to reaching the colon, an ability to degrade them provides a significant selective advantage for gut microbes.

**Glycan degradation profiles of mammalian species are associated with their diet.** Because diet is the prime determinant of colonic glycan composition and gut microbiota vary according to general dietary patterns (38), we expected that glycan degradation would systematically vary between the microbiota of different mammalian hosts based on their diet. To test this, we analyzed variation in diet and glycan degradation profiles across different mammalian species, using metagenomic sequencing data from 57 fecal samples across 34 different species, including 18 human samples (38). According to the host’s diet, each sample was characterized as being either herbivorous, carnivorous, or omnivorous. To correct for research biases arising from uneven annotations of CAZymes between species, we normalized each sample by the total number of CAZymes in it before running PCoA on the GlyDeR profiles. PCoA revealed a clear spectrum of samples over the first principal coordinate, from herbivores, through omnivores, to carnivores (see Fig. S3a in the supplemental material). With a subset of glycans that were categorized into either plant-derived or animal-derived glycans, we discovered a striking relationship between the diet of a host organism and the glycans predicted to be degraded by its gut microbiota: microbiota from herbivores tend to degrade plant-derived glycans (*P* = 0.04 compared to carnivores, Wilcoxon test) (Fig. 3a), while microbiota from carnivores prefer animal-derived glycans (*P* = 0.0001 compared to herbivores, Wilcoxon test) (Fig. 3a). Interestingly, the degradation efficiencies of omnivores and human gut microbiota place them as intermediates between herbivores and carnivores (Fig. 3a). Notably, the higher overall degradation efficiencies for animal-based glycans is probably due to the higher number of animal-derived glycans (1,898) compared to plant-derived glycans (594) in the database.

To further explore where humans stand with respect to the dietary spectrum, we trained a support vector machine (SVM) classifier to distinguish between herbivore and carnivore samples based on their inferred glycan degradation profiles (see Materials and Methods). The classifier predicted all but one sample correctly in a leave-one-out cross-validation (area under the curve

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**FIG 3** The connection between glycan degradation and diet. (a) GlyDeR profiling analysis of the Muegge data set. Bars showing the average sum of plant-specific and animal-specific GlyDeR scores of the samples grouped according to their host diet and normalized by the number of CAZymes in each sample. A fourth group was created to segregate humans from all other omnivores. The plant- and animal-specific GlyDeR scores of herbivores and carnivores are significantly different (*P* = 0.04 and *P* = 0.0001, respectively). (b) The Yatsunenko data set. A scatterplot showing the sample projections on the first principal coordinate and colored according to the country of origin. Samples from individuals younger than 2 years old were omitted (see text).
[AUC] = 0.93, \( F = 0.96 \) and notably outperformed a classifier based only on the abundance of CAZymes found in each sample (which had three misclassifications; AUC = 0.71, \( F = 0.84 \)), confirming the added predictive value of GlyDeR. We next applied this classifier to the 11 available nonhuman omnivore samples and classified 6 and 5 of the samples as herbivores and carnivores, respectively. These samples were missing direct dietary labeling; however, a comparison of these classifications versus the fiber index for these mammals (39) showed a nice correspondence with the predicted dietary regimens of the animals (Table 1). We next applied the classifier to predict the dietary habits for the human samples, which were unknown, resulting in 15 out of the 18 samples being labeled as carnivores. Thus, at least in the small population sample analyzed here, humans may be closer to carnivores in some functional aspects of their gut microbiota.

We next explored whether humans who live in different geographical areas with markedly different diets exhibit different GlyDeR profiles. We analyzed the Yatsunenko et al. data set (29), which contains metagenomic sequences from fecal samples of 110 humans who live in Venezuela, Malawi, and the United States. Malawian and Venezuelan diets are dominated by plant-derived polysaccharides, while typical U.S. diets contain large quantities of meat (29). As before, we ran GlyDeR and performed PCoA on all the GlyDeR profiles. Because infants display large variabilities over the first coordinate (see Fig. S3b in the supplemental material) and have an unusual diet relative to adults, we filtered out all samples from individuals younger than 2 years old. This led to a clear separation over the first coordinate between low meat consumers (Malawians and Venezuelans) and high meat consumers (Americans) (Fig. 3b). The ratio of animal- to plant-specific GlyDeR scores revealed significant differences between samples from different countries of origin (ANOVA; \( F = 6.56, P < 0.005 \)), with a higher animal/plant ratio in the United States (\( P < 0.003 \)) and Venezuela (\( P < 0.03 \)) than in Malawi. The ratio for the United States was slightly but not significantly higher than for Venezuela (Tuky–Kramer honestly significant difference test; see Fig. S3c).

**DISCUSSION**

In this analysis, we aimed to determine the association between human diet and microbial metabolism in the gut. We maintain that in order to properly study this relationship, one must incorporate the degradation of glycans into the equation. We detected diet-driven adaptations at both the level of single species (Fig. 2b) and of communities (Fig. 3a). We found species of the Bacteroidetes phylum to be the most efficient degraders of animal-derived glycans and human milk oligosaccharides. While this trend was apparent for Bacteroides, it was absent for Prevotella, another key member of that phylum. Diets that are high in animal protein have been associated with high levels of Bacteroides, whereas enrichment of Prevotella has been associated with diets rich in plant-derived carbohydrates and very low in animal protein (28–30). Given that many dietary animal glycans are derived from proteins (i.e., glycoproteins and proteoglycans), we propose that the high capabilities of Bacteroides and Parabacteroides to degrade animal glycans explains why their abundance is increased in Westerners (30, 37).

The plethora of novel glycans and their predicted glycans degradation efficiencies supplied by our method may prove to be highly important for designing prebiotic interventions. For example, the inability of some prebiotics to result in significant changes to the gut microbiota may be due to the use of glycans that are utilized too universally by the communities and thus do not provide a competitive advantage to individual, beneficial microbiota community members. GlyDeR provides researchers and clinicians with the ability to predict exactly what glycans are best metabolized by a given desirable taxon and excluded by others in a very specific way. As a striking example, a linear regression model based on GlyDeR-related features was capable of accurately predicting the abundance of bacterial strains that displayed low intersample variance. Degradation of long polysaccharides was the most predictive feature in the model, an unsurprising result considering the importance of these glycans as the main carbon and energy sources for colonic bacteria. Finally, our results were improved significantly by dividing the HMP samples into two clusters and reanalyzing each cluster individually. This supports the notion that microbiome analysis should not be general, but should rather be based carefully on the background community structure.

Our GlyDeR profiling revealed that the relative abundances of many taxa, especially those of Clostridium, are significantly correlated with their ability to degrade glycans. It was recently shown that Clostridium difficile and other pathogenic gut bacteria rely on microbiota-liberated mucosal glycans during their expansion in
the gut following antibiotic treatment (40). Thus, it may be possible to design prebiotics that help increase the levels of beneficial Clostridia and prevent the expansion of pathogenic strains. More generally, since the breakdown of a given substrate can be highly species specific (18), the prediction of bacterial glycan degradation efficiencies may prove to be an important tool for designing nutritional interventions to help alter microbial communities.

In analyzing the mammalian fecal samples data reported by Muegge et al. (38), we demonstrated that differences in microbial community compositions carry functional importance—that is, the microbiota of herbivores and carnivores have stronger affinities to plant- and animal-derived glycans, respectively. To the best of our knowledge, this is the first time that a computational framework has been able to provide such observations. The lack of large-scale in vitro glycan utilization assays makes straightforward validation of many of our predictions difficult at present. Nevertheless, our ability to train an accurate classifier to predict the diet of a host based on its microbiota glycan degradation profile, and the correspondence between the classifier’s predictions and animal nutrition (Table 1), both provide a strong supportive testimony to the veracity and utility of GlyDeR.

While humans are generally thought of as omnivores, there is an ongoing debate on the subject of our dietary history and adaptations. Tackling this question through the lens of our microbiota, we used the aforementioned binary herbivore-carnivore classifier in order to classify humans. Remarkably, the classifier predicted 15 out of 18 human subjects to be carnivores. This result is less surprising considering that all of the human subjects were U.S. residents and that the United States is the highest meat-consuming country per capita in the world (41). In contrast to the U.S. population, the diets of individuals from Malawi and Venezuela mainly include plant-derived polysaccharides (these populations consume, on average, 8.3 and 76.8 kg of meat per year, as opposed to 120.2 kg per year in the United States [41]). We consequently found a lower ratio for animal versus plant degradation potency in the microbiota of individuals from these countries (see Fig. S4c in the supplemental material). Notably, GlyDeR does not predict a reduced potency of plant degradation within the U.S. population. Therefore, it seems that the capacity of Western individuals to degrade glycans has not diminished over the course of evolution, but merely shifted toward the direction of carnivores.

Taken together, these results further advance our understanding of human diet-specific adaptations, but conclusions must be drawn with caution. First, the data upon which GlyDeR relies are often incomplete. For instance, only 74 out of the 146 CAZymes mapped to at least one HMP genome (see Fig. S2c in the supplemental material). Furthermore, 31 CAZymes were not capable of breaking any glycan, either because some glycan structures are missing from the database or because of inaccurate enzymatic annotation (see Fig. S2c). Second, several biases may arise from the existence of nonproportional representation of different glycan categories in the KEGG Glycan database. For example, the animal-derived GlyDeR scores are always higher than plant-derived scores because there are three times more animal-derived glycans in the database. We therefore suggest looking only at relative scores and conducting comparisons within a given glycan category (e.g., comparing the degradation of animal-derived glycans in Malawi and the United States) and not between different glycan categories. Finally, the GlyDeR platform does not take into account many important factors, such as enzyme transcription levels, kinetic parameters, and downstream biochemical pathways for glycan utilization. Nevertheless, GlyDeR is the first computational analysis framework that successfully enables one to directly model how the microbiota can respond to dietary glycans from a mechanistic point of view.

The current study was focused on developing a novel method for the study of glycan degradation processes and establishing this method’s value in assessing a wide spectrum of diet-related trends. Future studies will examine whether GlyDeR could capture subtler differences in microbiomes that are derived from individuals with more homogeneous backgrounds (e.g., only meat-eaters). Another open question is the identification of strain differences within a given bacterial species. Since the current study involved only a selected set of 203 microbial genomes, we did not have sufficient coverage of any single species in order to establish significant strain differences in glycan degradation. Since the number of sequenced genomes is constantly increasing, we expect that a larger analysis would be able to provide insight into the glycan degradation capabilities of different strains within species. Finally, an important issue to tackle is the aspect of microbial cross-feeding. GlyDeR is fully capable of analyzing metabolic interactions in the degradation of complex substrates that demand more than one strain for breakdown, and it therefore provides a golden opportunity to examine glycan degradation processes that occur in synthetic communities (and ultimately in natural ones).

While metagenomics are still the gold standard for high-throughput functional analysis of microbial communities, 16S rRNA sequencing is a strong alternative in the many circumstances where metagenomic data are not available, or prohibitively expensive. To this end, we plan to extend and integrate GlyDeR into routine 16S rRNA analyses (e.g., with the help of PICRUSt [42]), as well as incorporate GlyDeR within the larger framework of genome-scale metabolic modeling (9, 43–46). Within this integrated framework, glycan analysis will hopefully become a standard tool in the arsenal of microbial researchers.

**MATERIALS AND METHODS**

We developed various tools for glycan degradation data integration, manipulation, and analysis, and these can be divided into 3 categories: (i) data retrieval, for which we describe here the sources of information for this study and how they were used; (ii) reconstruction of glycan degradation reactions and cross-validation, by which we reconstructed novel glycan degradation reactions by implementing our GlyDeR algorithm; (iii) data analysis, that is, the steps we performed in order to analyze single taxa and 16S rRNA and metagenomics sequence data and generated microbial (meta)genome glycan degradation potency predictions.

**Data retrieval.** Information about glycans and the enzymes that might break them down is spread across many databases and tools. The types of data sources used to infer genome-based glycan degradation capacities included bacterial taxa, genome annotations, and glycans.

(i) **Bacterial taxa.** A catalog of 281 taxa was downloaded on 8 October 2011 from The Human Microbiome Project website (http://www.hmpdacc.org/) using the following filters: NCBI superkingdom *Bacteria*; HMP isolation body site: gastrointestinal tract; project status complete; NCBI submission status annotation (and sequence) public on NCBI site. The catalog contains the following annotation fields: HMP ID, GOLD ID, organism name, domain, NCBI taxon ID, NCBI superkingdom, NCBI phylum, NCBI class, NCBI order, NCBI family, NCBI genus, NCBI species, all body sites, all body subsites, current finishing level, NCBI project ID, Genbank ID, Gene count, size (kb), GC content, Greengenes ID, NCBI 16S accession, strain repository ID, oxygen requirement, cell shape,
motility, sporulation, temperature range, optimum temperature, Gram stain, type strain.

(ii) Genome annotations. All of the HMP taxa were searched against the Seed database (http://pubseed.theseed.org/) using the key of NCBI Taxon Id number as a cross-reference. A total of 204 matches were detected, and their RAST genome annotations were extracted by using the web services of API.

(iii) Glycans. The entire KEGG Glycan database (http://www.genome.jp/kegg/glycan/) was downloaded on 7 January 2011. The database contains information on 10,978 glycans. We used the following annotation fields from the annotations: G number, name, KCF file, and class. An additional biological origin field was retrieved from an external source, as described below.

The KCF file for each glycan describes a graphical representation of its two-dimensional structure. This representation takes into account the monomeric building blocks (nodes) and the glycosidic linkages (edges) of the glycan. Textual and visual representations of the KCF graph for glycan G00010 are given in Fig. S1a and b in the supplemental material.

The glycans database was subsequently filtered according to the following criteria related to nodes, edges connecting two nodes, and synonym glycans (i.e., glycans with identical structures but different ID numbers). Since the database contains more than 800 nodes denoting glycan-related building blocks, many of which are extremely rare, we chose to focus on a subset of 35 nodes that corresponded to the most prominent sugar monosaccharides, prevalent modifications, amino acids found in the glycans, database, or it has more than 10 nodes.

The reactions from KEGG that map to the EC number. Linkages Broken—In the case of glycosidic linkages the value is a string representing two nodes and an edge that connects them based on the KCF graph representation of the glycan structure. For example, in the case of deacetylation reactions, this value is “Ac-R,” denoting the removal of an acetyl group from node R. Contained Subglycan—A G# identifier of a glycan structure contained within the structure of a larger glycan, e.g., “Glc b1-4 Glc” is a subglycan of Man a1-3 Glc b1-4 Glc. Contains Only (nodes)—A G# identifier for one or more nodes that the glycan must contain and only contain. Glycan Released—A G# identifier that defines a glycan that must be one of the products after the reaction with the enzyme takes place. Endo vs. Exo—endo enzymes, which remove only terminal sugars (the edges of the terminal nodes), and “exo” enzymes, which break all glycosidic bonds except for terminal ones (remove all edges except the ones of the terminal nodes), and both enzymes, which can break any bond (remove any edge).

Endo glycan degradation reactions). These fields can be further described as follows:

- **Enzyme Name**—The accepted name of the enzyme. KEGG Reactions—The reactions from KEGG that map to the EC number. 
- **Linkages Broken**—In the case of glycosidic linkages the value is a string representing two nodes and an edge that connects them based on the KCF graph representation of the glycan structure. For example, in the case of deacetylation reactions, this value is “Ac-R,” denoting the removal of an acetyl group from node R. 
- **Contained Subglycan**—A G# identifier of a glycan structure contained within the structure of a larger glycan, e.g., “Glc b1-4 Glc” is a subglycan of Man a1-3 Glc b1-4 Glc. 
- **Contains Only (nodes)**—A G# identifier for one or more nodes that the glycan must contain and only contain. 
- **Glycan Released**—A G# identifier that defines a glycan that must be one of the products after the reaction with the enzyme takes place. 
- **Endo vs. Exo**—endo enzymes, which remove only terminal sugars (the edges of the terminal nodes), and “exo” enzymes, which break all glycosidic bonds except for terminal ones (remove all edges except the ones of the terminal nodes), and both enzymes, which can break any bond (remove any edge).

**DP preference**—This field reflects the degree of polymerization of the glycans upon which the enzyme works. This number is also the exact/ minimal (+)/maximal (−) number of nodes that the KCF graph for the glycan structure must contain. Terminal Side Preference—This field is unique for exo-acting enzymes and describes their specificity toward the reducing or nonreducing end. The KCF graph is directional, hence “reducing” means only the removal of the right-most node is allowed, “nonreducing” refers to the left-most, and “both” allows the removal of both edges. Notice that the terminal node of the reducing end is always at position 1 in the KCF graph, except for repeating glycans. Enzymatic Reaction—A textual description of the enzymatic reaction performed by a given enzyme (EC number) was obtained from http://enzyme.expasy.org/. Comments—Specific comments about an enzyme (EC number) were also taken from http://enzyme.expasy.org/. Special characters used in the above descriptions included the following. S signifies any number. R signifies any type of the following sugars [noder abbreviations are those used for the http://enzyme.expasy.org/ website]: ara, Araf, D/LAraf, LAnAraf, D/LAraf, LAn, Ape, D/LAp, D/LAp, 3,6-Anh4-Gal, L3,6-Anh4-Gal, 3,6-Anh4-Gal, Gal, D/LGal, GalNac, GalN, GlcNAc, D/LGlcNAc, GlcNAc, D/LGlcNAc, GlcN, D/LGlcN, Glc, D/LGlc, Fru, Frul, D/LFrul, D/ Frul, Man, Manf, D/LMan, ManA, Rha, D/LRha, Rha, D/LRha, Gal, GaL, D/LGal, D/LGalf, Fuc, D/LFuc, Fuc, LFuc, D/LFuc, Xyl, D/LXyl, Xylf, Neu, Neu5Ac, Neu5Gc, MurNAc. # indicates an “or” association, and & denotes an “and” association.) Overall, this workflow resulted in 141,361 glycan degradation reactions, of which 9,325 were reactions that degrade KEGG glycans and newly reconstructed glycans and 132,236 were intermediate glycan-degrading reactions.

(v) CAZY annotation. We used sequence similarity to match the genes which belong to the HMP taxa with specific CAZymes. We therefore
performed BLAST analysis for all of the genomes of the HMP taxa against
the bacterial protein sequences found in the CAZy database. Each enzyme
family in CAZy contains a set of manually curated enzymes that have been
determined to execute a specific catalytic function. We used the NCBI
BLAST utility and filtered errors at the level of 10e−10 and matches below
97% identity. At that point, we had a mapping between genes in the HMP
taxa and CAZyme families. Because many families contain a one-to-many
mapping between a family and its associated EC numbers, we had to refine
this annotation. We therefore extracted the genes predicted for enzymatic
annotations from the SEED and KEGG databases. While the CAZyme
annotations are more comprehensive, they are sometimes not as accurate
as the manually curated ones. Thus, we integrated the information ob-
tained from all of these sources by using the following logic: for proteins
that were mapped to families of multiple EC numbers, we first checked if they had an available annotation in SEED or
KEGG, and if that was the case, then we checked if this annotation be-
longed to one of the multiple annotations in CAZy. If it did, then we
accepted the KEGG/SEED annotation.

(vi) Subcellular localization annotation. To define the subcellular
localization (SCL) of reactions, we used the RAST genome annotation as
a first proxy. We mined the function and subsystem fields of the annota-
tion for special keywords. For our purposes, we were only interested in
whether the enzyme exerts its function inside the cell or outside. Enzymes
were defined as intracellular if their associated genes contained the key-
words cytoplasm, cytosol, or cytoplasmic. Enzymes were defined as cross-
membrane if their associated genes contained one of the keywords
periplasm, periplasmic, inner membrane, or cytoplasmic membrane. Fi-


ally, enzymes were defined as extracellular if their associated genes con-
tained one of the keywords cellulose, outer membrane, secreted, cell
wall, or extracellular. For enzymes that were not associated with any
meaningful keyword, we took advantage of the LOCtree localization pre-
diction software (https://rostlab.org/owiki/index.php/LOCtree). LOC-
tree uses a protein amino acid sequence to predict the SCL. It supplies five
possible SCLs: cytosol, inner membrane, periplasmatic, outer membrane,
and secreted. Enzymes with the value cytosol were classified as intracellu-
lar, enzymes with the values secreted or outer membrane were classified
as extracellular, and enzymes with the values periplasmatic or inner mem-
brane were classified as cross-membrane, i.e., enzymes that exert their
function on the cross-membrane between the cell and its environment.
To fix possible erroneous annotations, we refined our localization selection based on specific knowledge of the glycan degradation biochemistry. A
literature survey suggested that there are no polysaccharides within the
bacterial cytoplasm (with glycomycin being the only exception). Thus, en-
zymes that were predicted as extracellular or cross-membrane were fil-
tered out if the glycan that they processed was either repeating, defined as
a polysaccharide, or had more than 10 subcomponents.

(vii) Biological origin of glycans. We accessed the CarbBank database
(47) and mapped the KEGG glycans to it using the KEGG Glycan ID (G
number) as a cross-reference. CarbBank contains detailed descriptions of
where a specific glycans can be found in nature. We parsed these data in
order to define certain glycans as either plant-derived or animal-derived.

(viii) Degree of polymerization of glycans. Glycans are routinely cat-
ergorized into one of four possible degrees or classes of polymerization.
With respect to classes, glycans were defined as disaccharides if they con-
tained 2 nodes, oligosaccharides if they contained 3 to 10 nodes, short
polysaccharides if they contained >10 nodes, and long polysaccharides if
they had a repeating structure.

Construction of the CAZyme table (a key step in the GlyDeR pipe-
line). We manually curated all of the CAZymes (146 EC numbers) and
mapped each one to a set of computer-based rules dictating the mecha-
nism by which it can break a given glycan (i.e., split its graph into two
separate components). These rules account for structural features such as
the glycosidic linkages the enzyme can break, the cleavage mechanism, the
chemical neighborhood, and the degree of polymerization of the glycan
(see Table S3a and b for a list of the rules). We then executed these rules on
all the glycans that appear in the KEGG Glycan database, which yielded
141,561 glycan degradation (GlyDeR) reactions. In the following section we
describe the logic behind the reconstruction of glycan degradation
reactions by identifying for each glycan which enzymes are able to break it
and how the degradation reaction will look. GlyDeR reactions are then
mapped back to CAZymes in order to produce a table where the rows are
CAZymes, the columns are glycans, and each entry contains a CAZyme
score, calculated as follows:

\[ \text{CAZyme score}_ij = \sum \frac{1}{\text{Gk}_j} \quad \forall \text{e}_i \]

where \( \text{e}_i \) is an enzyme that can degrade glycan \( j \) and \( \text{Gk}_j \) is the number of
glycans that \( \text{e}_i \) breaks down. The entire construction process is summa-
rized in Fig. 1b. A CAZyme table which contains all of the CAZyme scores


(i) Glycan degradation (GlyDeR) rules. A reaction is represented by
its substrate(s), product(s), the enzyme(s) responsible for the catalysis, and
a subcellular localization. For the GlyDeR reaction generation process,
we used all the computer-based glycan breaking rules described in
Table S1 of the supplemental material. For an EC number-related enzyme
(rule) to break a glycan, the glycan and the resultant reaction must comply
with all the limitations defined in the fields of the given rule, namely, the
glycan must contain at least one of the glycosidic linkages (or nodes con-
taining an acetyl group in case of deacetylation reactions) described in the
Linkages Broken field; the glycosidic linkage hydrolyzed must appear in
the terminal edges of the glycan if the value in the Endo Vs. Exo is set to
Endo, and vice versa; the number of nodes the glycan contains must con-
form to the value described in the DP Preference field; in case the Endo
versus Exo field is set to Exo, the terminal side of the glycosidic linkage
hydrolyzed must be located on the right side of the graph of the glycan
if the value in the Terminal Side Preference field is set to Reducing and on
the left side if this field is set to Nonreducing. If this field is set to “Both,”
then location of this linkage on both sides is allowed. The glycan must
contain the structure of a glycan (nodes and edges) described in the Con-
tained Subglycan field, and the linking break must be more part of this
subglycan. The reaction must contain the glycan described in the
Glycan Released field as one of its products. Figure 1a gives an example of
an Exo-acting enzyme breaking a regular glycan.

(ii) Deacetylation rules. Some of the EC numbers (enzymes) we ana-
lyzed have a deacetylation activity, i.e., they have the capability to remove
acetyl groups. In the KEGG Glycan database, monosaccharides contain-
ing an acetyl group are described as a single unique node, e.g., the node
GlcNAc corresponds to N-acetyl-glucosamine. Therefore, if an enzyme
has the capability to remove an acetyl group, we simply remove the sub-
string “Ac” from the label of the node and make it the product of the
reaction, e.g., GlcNAc ↔ GlcN + Ac.

(iii) Reconstruction of new glycans. We manually constructed a set of
107 glycans which we determined were important but that were missing
from the KEGG Glycan database. To distinguish these glycans from the
ones previously available in the database, we gave these new glycans the
prefix TAU instead of G, which is assigned by KEGG. A list of all the TAU
glycans can be found in the supplemental material. Furthermore, in most
cases the products of the degradation reactions did not have a preexisting
G number, meaning they currently do not exist in the KEGG Glycan
database. Working under the assumption that most glycans in nature are
still uncharacterized in databases, we decided to add these new glycans
automatically. Thus, whenever a reaction produced a new glycan, we gave
this glycan a unique ID beginning with “TAUS” (to distinguish it from
original glycans, designations for which begin with G or TAU).

Data analysis. (i) Single-taxon data analysis. To define microbial
genome-specific GlyDeR scores, after building the CAZyme table we asso-
ciated the CAZymes with the genomes of the HMP gut taxa. For every
taxon-specific gene we calculated, based on its enzymatic annotation and
the enzyme’s subcellular localization, a GlyDeR score. Given a bacterial
taxon \( i \) and glycan \( j \), the GlyDeR score was calculated as follows:

\[ \text{GlyDeR}_{ij} = \frac{1}{\text{Gk}_j} \quad \forall \text{e}_i \]
GlyDeR score _c_ = \sum \frac{n_{j,k}}{g_{k}}, \quad \forall c \in C

where _c_ is an enzyme that can degrade glycan _j_, _n_ is the number of genes in its genome which translate to enzyme _c_, and _g_ is the number of glycans broken by enzyme _c_. This metric decrements the contribution of CAZymes that are more promiscuous versus those specifically geared to degrade the glycan in question (Fig. 1b). For some categories of glycans such as long polysaccharides and plant-specific glycans, we defined a category-specific score, _GS_ _c_, which is the sum of GlyDeR scores for glycans that belong in that group:

GlyDeR score _c_ = \sum GS_{j,p} \quad \forall j \in c

where _GS_ _j, p_ is the GlyDeR score for genome _i_ and glycan _j_ and _c_ is the collection of glycans that belong to category _C_. This scoring system has the feature that summing the GlyDeR scores over all the glycans in a given genome gives the total number of CAZymes in the genome, according to the following equation:

Total CAZymes _i_ = \sum GS_{j,p} \quad \forall j \in j

where _j_ \in _j_ is the set of all glycans and _i_ is the index of a specific taxon. Subsequently, we defined the GlyDeR profile of a bacterial taxon as follows:

GlyDeR profile _i_ = \{GS_{i,1}, GS_{i,2}, ..., GS_{i,j-1}, GS_{i,j}\}.

For the GlyDeR reaction consistency check and cross-validation, when applied to all the glycans available in KEGG, the GlyDeR pipeline produced a list of 114,573 intermediate glycan products, most of which were novel and thus do not appear in the original KEGG database. We refer to these as hypothetical glycans in the test. To test the consistency of GlyDeR, we performed a cross-validation process where we picked a random subset of 1,000 glycans from KEGG and applied GlyDeR to degrade them. We then tested whether the products obtained from these 1,000 glycans were enriched for known glycan degradation potential. These features were the GlyDeR scores for plant-specific glycans, animal-specific glycans, disaccharides, oligosaccharides, short polysaccharides, and long polysaccharides. Each feature represented the sum of the GlyDeR scores for the glycans that belong in the class.

(ii) 16S rRNA sequence data analysis. We retrieved the 16S rRNA sequence data and metadata from fecal samples belonging to 325 healthy human individuals from the HMP Data Analysis and Coordination Center (DACC) (48). Because we were not interested in time series data, we only used samples from the initial time point. 16S rRNA sequences were mapped to the HMP genomes based on sequence similarity and further used to build an operational taxonomic unit table that described the abundances of the HMP taxa in each sample. For this purpose, we used the QIIME software (49) with the following exact commands:

pick_otus.py -i HMP_samples_seqs-r HMP_taxa_ref_seqs -m uclust_ref-C make_otu_table.py -i pick_otus_output

(iii) Metagenomics sequence data analysis. We retrieved the Muegge et al. (38) and Yatsunenko et al. (29) data sets from MG-RAST (50). For both data sets, we downloaded the FragGeneScan gene-calling output, which maps each original read to 0, 1, or more open reading frames (ORFs). This way, many reads could be assigned to a single ORF and so the abundance of each ORF was taken into account. Next, ORFs were assigned to CAZymes and to specific subcellular localizations, as described above. The percent identity used for the BLAST search was changed to 60%, a value commonly used in metagenomic annotation projects (51). We then constructed a CAZymes abundance table to describe the abundances of all the CAZymes in each sample. In order to calculate a sample-specific GlyDeR score, we used the sample CAZymes abundance table and precalculated GlyDeR scores. Thus, the GlyDeR score, _GS_ _j_ for glycan _j_ in sample _k_ was defined as follows:

_\sum n_{j,k} D_{max}/D_{k} \quad \forall c_\in C

where _c_ is an enzyme that can degrade glycan _j_, _n_ is the number of genes in sample _k_ that map to enzyme _c_, and _g_ is the number of glycans broken by enzyme _c_. _D_{max}/D_{k} is a normalization factor that denotes the ratio between the depth (i.e., the total number of sequenced reads) of the sample with the maximum depth (D_{max}) and the depth of the current sample, D_{k}. Next, we defined the GlyDeR profile of sample _k_ as follows:

G_P_k = \{GS_{j,1}, GS_{j,2}, ..., GS_{j,j-1}, GS_{j,j}\}

(iv) Multivariate regression between GlyDeR features and bacterial abundance. We analyzed the 16S rRNA sequences from the HMP fecal samples in order to determine the abundance of our 203 bacterial taxa in each sample. To increase the signal-to-noise ratio (SNR), we excluded species with a high abundance variability based on the following criterion:

\text{SNR} = \frac{\text{mean abundance}}{\text{standard deviation}} > 0.3

The 6 features described in the previous section were used to build a linear regression model with bacterial abundance as the dependent variable: bacterial abundance = \{(-13.248 \times \text{plant-specific GlyDeR score}) + (28.1822 \times \text{disaccharides score}) + (9.6206 \times \text{oligosaccharides score}) + (32.701 \times \text{long polysaccharides score}) + 42.7354\}. To eliminate the possibility of overfitting the data, we used a standard 10-fold cross-validation method. All calculations were performed using WEKA (51).

To assess the added value of using the GlyDeR features over genomic information alone, we defined for each HMP taxon a vector containing the genomic copy number of the CAZymes used in the analysis. We then built a similar linear regression model with these 82 CAZymes used as features and bacterial abundance in the HMP samples as the dependent variable.

Because of the high variability in bacterial taxa abundance across the samples, we used the KMeans algorithm to cluster the samples. We chose to use 3 clusters because this option resulted in the lowest cubic clustering criterion (CCC). However, one cluster was composed of only one outlier taxon, so we omitted it from further analysis. Indeed, after removing the outlier point, the lowest CCC was achieved for _k_ = 2. We built a cluster-specific linear regression model, as described above: cluster 1 bacterial abundance = \{(-40.6116 \times \text{plant-specific GlyDeR score}) + (33.283 \times \text{long polysaccharides score}) + 31.6149\}; cluster 2 bacterial abundance = \{(-15.9773 \times \text{oligosaccharides score}) + (69.8184 \times \text{long polysaccharides score}) + 70.697\}. We defined a list of 25 bacterial taxa with highly predictable accuracy. A taxon was included in the list if the standard error of its predicted abundance obeyed the following rule for either cluster 1 or cluster 2:

\text{predicted abundance} - \text{actual abundance} < 1 \text{ actual abundance}

(v) Classification of dietary patterns based on GlyDeR scores. The GlyDeR scores of all herbivore and carnivore mammals from the Muegge et al. data set were used to train a binary SVM classifier. The SMO implementation of the LIBSVM package was used to do the classification.
mentation of this classification algorithm in WEKA (51) was used for the computation. To estimate the accuracy of the classifier, we used a standard leave-one-out cross-validation. To apply this classifier to the remaining human and nonhuman omnivore samples, we hid the labels of the samples and classified them as either carnivore or herbivore.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at http://mbio.asm.org/lookup/suppl/doi:10.1128/mBio.01526-14/-/DCSupplemental.

Table S1, XLSX file, 0.1 MB.
Table S2, XLSX file, 0.1 MB.
Table S3, XLSX file, 0.1 MB.
Table S4, XLSX file, 0.1 MB.
Table S5, XLSX file, 4.1 MB.
Table S6, XLSX file, 2.1 MB.
Table S7, XLSX file, 0.3 MB.
Table S8, XLSX file, 0.1 MB.
Table S9, XLSX file, 0.1 MB.

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