The Complete Genome Sequence of *Fibrobacter succinogenes* S85 Reveals a Cellulolytic and Metabolic Specialist

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Abstract

*Fibrobacter succinogenes* is an important member of the rumen microbial community that converts plant biomass into nutrients usable by its host. This bacterium, which is also one of only two cultivated species in its phylum, is an efficient and prolific degrader of cellulose. Specifically, it has a particularly high activity against crystalline cellulose that requires close physical contact with this substrate. However, unlike other known cellulosic microbes, it does not degrade cellulose using a cellulosome or by producing high extracellular titers of cellulase enzymes. To better understand the biology of *F. succinogenes*, we sequenced the genome of the type strain S85 to completion. A total of 3,085 open reading frames were predicted from its 3.84 Mbp genome. Analysis of sequences predicted to encode for carbohydrate-degrading enzymes revealed an unusually high number of genes that were classified into 49 different families of glycoside hydrolases, carbohydrate binding modules (CBMs), carbohydrate esterases, and polysaccharide lyases. Of the 31 identified cellulases, none contain CBMs in families 1, 2, and 3, typically associated with crystalline cellulose degradation. Polysaccharide hydrolysis and utilization assays showed that *F. succinogenes* was able to hydrolyze a number of polysaccharides, but could only utilize the hydrolytic products of cellulose. This suggests that *F. succinogenes* uses its array of hemicellulose-degrading enzymes to remove hemicelluloses to gain access to cellulose. This is reflected in its genome, as *F. succinogenes* lacks many of the genes necessary to transport and metabolize the hydrolytic products of non-cellulose polysaccharides. The *F. succinogenes* genome reveals a bacterium that specializes in cellulose as its sole energy source, and provides insight into a novel strategy for cellulose degradation.

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Introduction

Herbivorous mammals are essential components of terrestrial ecosystems and are major participants in the global carbon cycle, as well as the foundation of animal agriculture. Much of the plant biomass consumed by herbivores is degraded by symbiotic microorganisms in the host digestive tract. This symbiotic interaction between plant-degrading microbial communities and their herbivorous hosts is perhaps best exemplified by ruminants such as domestic cattle. Plant biomass digestion occurs in the rumen, a specialized pregastric fermentative organ that can comprise up to one-sixth of the weight of the host animal [1]. The ruminal fermentation is characterized by an incomplete anaerobic digestion in which plant material is converted to a mixture of C2 to C6 volatile fatty acids (VFAs), some of which are produced via intermediates such as succinic and lactic acids. These VFAs are used by the host as its primary energy source. The ruminal microflora are also responsible for producing other metabolic products, including methane, carbon dioxide and microbial cells, the last of which are digested...
postriminually to supply a major portion of the protein requirements of the host.

An analysis of bacterial diversity in the rumen reveals many microbes capable of degrading plant cell components (cellulose, hemicelluloses, and starch), including members of the genera *Ruminococcus, Prevotella,* and *Butyrivibrio* [2]. One of the most highly cellulolytic of the ruminal microbes is *Fibrobacter succinogenes,* a Gram-negative bacterium originally classified in the phylum Bacteroidetes [3], but later resolved to its own unique phylum, Fibrobacteres [4]. Several studies employing quantitative PCR have revealed that in dairy cattle, *F. succinogenes* comprises several percent of the total bacterial 16 S rRNA genes, a proxy for relative population size within the prokaryotic community [5]. The microbial species composition of the rumen depends strongly on the animal model used and feed composition, and in some cases *F. succinogenes* is the predominant cellulolytic organism [reviewed in [6]].

Pure culture studies have shown that *F. succinogenes* is a highly cellulolytic mesophilic bacterium capable of growth on crystalline cellulose with a maximum specific growth rate constant of \( \sim 0.076 \text{ h}^{-1} \) [7]. Moreover, this species is a very effective competitor for cellodextrin products of cellulose hydrolysis [8,9], and its ability to efflux cellodextrins produced by intracellular cellodextrin phosphorylase may contribute to the cross-feeding of other ruminal bacteria, both cellulolytic and non-cellulolytic [10].

The substantial cellulolytic capabilities of *F. succinogenes* appear related to its unique mode of hydrolysis. Like most anaerobic cellulolytic bacteria, *F. succinogenes* does not excrete significant amounts of cellulosomes into its environment, and degradation requires attachment of cells to the cellulose surface. However, this species does not appear to contain surface-bound cellulosomes or their signature features such as scaffolds or dockerin-binding domains [11] that comprise the degradation apparatus of the most commonly studied cellulolytic anaerobes such as *Clostridium thermocellum* or the ruminococci [12]. In *F. succinogenes*, degradation of crystalline cellulose may be facilitated by a highly unusual orientation of the cells along the crystallographic axis of the degrading cellulose fiber [13,14].

To gain a better understanding of the biology of *F. succinogenes,* we sequenced the genome of the type strain *Fibrobacter succinogenes* subsp. *succinogenes* S85 ATCC 191699 (Henceforth *Fibrobacter succinogenes*) to completion. A metabolic reconstruction analysis of this genome reveals how *F. succinogenes* is physiologically adapted to the rumen environment. An analysis of its plant polysaccharide degrading machinery coupled with growth assays reveals that, while it has the ability to hydrolyze a wide variety of polysaccharides, it can only utilize cellulose and its hydrolytic products for growth. These data confirm that *F. succinogenes* is a metabolic specialist that mediates its cellulolytic lifestyle by removing plant cell wall hemicelluloses to gain access to cellulose. Given the recent interest in optimizing carbohydrate degradation for the production of biofuels, the genome sequence of *F. succinogenes* not only provides insight into its unique lifestyle, but is also a valuable resource for understanding microbial models of plant cell wall deconstruction.

**Results**

**General features of the *F. succinogenes* genome**

The *F. succinogenes* genome consists of a single, circular chromosome of 3,842,635 base pairs with a GC content of 48%, confirming a previous report on a *F. succinogenes* genomic map [15]. Gene prediction revealed 3,085 putative coding sequences, covering 90.76% of the genome, with an average length of 1,130 bp. A total of three rRNA operons were identified including three 5 S rRNAs, three 16 S rRNAs, and three 23 S rRNAs; in addition, 59 tRNAs covering all 20 protein amino acids were also recovered (GenBank accession CP001792.1). Comparison of the predicted proteins in *F. succinogenes* against a database containing the proteins from over 1,100 other sequenced microbial genomes revealed that 1,787 proteins (58%) could be assigned a putative function, 510 proteins (16.5%) were similar to those encoding hypothetical proteins, and the remaining 738 proteins (25.5%) had no significant similarity to any protein in the database, indicating that these may be genus- or species-specific proteins.

**Phylogenetic placement of *F. succinogenes***

*F. succinogenes* was originally placed within the phylum Bacteroidetes, and was included in the genus *Bacteroides* [3]. Subsequent 16 S rRNA sequencing revealed that *F. succinogenes* did not belong to the Bacteroidetes and was reclassified into the novel phylum Fibrobacteres [4]. Only the genus *Fibrobacter* has been described for this phylum, and this genus presently contains only one other formally described species, *F. intestinalis* [4]. Later work resolving the phylogenetic relationship of the Fibrobacteres relative to other phyla using housekeeping genes have shown that it is most closely related to the Bacteroidetes/Chlorobi [16,17] and either the Actinobacteria [17] or the Chlamydiae [18]. To help further resolve this phylogenetic placement, we performed a taxonomic distribution analysis [19] by comparing each predicted protein in the *F. succinogenes* genome against a database containing all proteins from the complete microbial genome collection. For each *F. succinogenes* protein with a match, we determined the taxonomic identity of its top match and counted the total number of proteins in *F. succinogenes* that have their closest match to microbes belonging to a given phylum as shown in Figure 1. If *F. succinogenes* is closely related to bacteria in a different phylum, it is expected that a majority of proteins in *F. succinogenes* would have close matches to proteins in bacteria belonging to that phylum. We found that *F. succinogenes* contains proteins similar to a variety of bacteria in the current sequenced genome database collection. The majority of these proteins had multiple closest matches to bacteria belonging to Firmicutes (19%), Deltaproteobacteria (16%), Bacteroidetes (15%), Gammaproteobacteria (11%), and the Chlorobi (5%). These data confirm, on a whole-genome scale, that *F. succinogenes* belongs to its own phylum.

**COG analysis**

To understand how *F. succinogenes* deploys genes in its genome, we performed a clusters of orthologous group (COGs) analysis [20]. This analysis places proteins into specific categories related to different aspects of cellular metabolism and physiology. A COG analysis of the *F. succinogenes* genome was able to classify 1,938 proteins into 17 different categories as shown in Table S1. From this analysis, over 30% of these proteins belong to four categories, including cell wall, membrane, and envelope biogenesis (category M, 11%); amino acid transport and metabolism (E, 9%); translation, ribosomal structure and biogenesis (J, 8%); and carbohydrate transport and metabolism (G, 8%). These results likely reflect the importance of these categories to the biology of *F. succinogenes,* as much of its lifestyle is dependent on the metabolism of carbohydrates for cellular processes. The high percentage of genes devoted to cell wall, membrane, and envelope biogenesis likely reflects *F. succinogenes* ability to adhere to plant biomass. In contrast, other categories have a paucity of proteins, including cell motility (N, 1%); secondary metabolites biosynthesis, transport and catabolism (Q, 2%); and defense mechanisms (V, 2%). The compositions of these categories are in accord with the lifestyle of...
**Fibrobacter succinogenes** as it is not actively motile and appears to have few defenses against antibiotics (see below). A key feature of the **F. succinogenes** COG analysis is the large number of proteins not identifiable: 1,147 out of 3,085 (37%) proteins could not be classified. Of the 1,938 proteins that could be classified, 357 (18%) were identified as either general function prediction only or function unknown. This indicates that over 50% of the **F. succinogenes** proteome could not be assigned to a known COG category. Although the COG database may not be extensive enough to capture all known functional categories, the large numbers of proteins with unassigned COGs, combined with our discovery that over 25% of the **F. succinogenes** proteome has no orthologs to microbes in the sequenced genome collection, may indicate numerous proteins unique to **F. succinogenes**.

### Polysaccharide degradation and utilization

A hallmark of **F. succinogenes** is its ability to efficiently hydrolyze the many plant polysaccharides it encounters in the rumen. We tested the ability of **F. succinogenes** to hydrolyze and/or utilize different polysaccharides by performing growth assays using a variety of substrates as shown in Table 1. Of the tested polysaccharides, which include cellulose, pectin, starch, gluco-mannan, arabinogalactan, and various forms of xylan, only cellulose was found to be both hydrolyzed and metabolized. A large number of other polysaccharides were found to be hydrolyzed without being metabolized, including all forms of xylan tested. The capacity to hydrolyze xylan but not utilize it for growth is reflected in the genome of **F. succinogenes**, as it does not have the genes necessary to either transport or metabolize xylose or xylodextrins (see below). Therefore, the hydrolytic capability of this cellulose-dependent bacterium to degrade other plant polysaccharides suggests that **F. succinogenes** removes xylose-rich hemicelluloses to gain access to cellulose, and likely uses this exposed cellulose as its major energy source. We describe below two separate aspects of this strategy of plant cell wall deconstruction including the enzymatic hydrolysis of plant polysaccharides by each of the major classes of polysaccharide hydrolases and associated proteins, and the subsequent utilization of the products of cellulose hydrolysis. Other metabolic capabilities encoded by the genome are discussed in Supplemental Text including glycerol biosynthesis and utilization (Text S1), amino acid synthesis and nitrogen assimilation (Text S2), fatty acid synthesis and catabolism (Text S3), vitamin biosynthesis (Text S4), transporters (Text S5), antibiotic production and resistance (Text S6), DNA repair mechanisms (Text S7), and CRISPRs, insertion sequences, and genomic islands (Text S8).

### Enzymatic hydrolysis of polysaccharides

A carbohydrate-active enzyme (CAZy) [21] analysis revealed a total of 134 genes encoding enzymes that were classified into 49 different families of glycosyl hydrolases (GHs), carbohydrate binding modules (CBMs), carbohydrate esterases (CEs), and polysaccharide lyases (PLs) (Table S2). Because some of these enzymes have multiple CAZy family memberships (i.e. contain both a GH and a CBM), 196 total CAZymes were identified using this ontology. Only a small number of these CAZy family members have been experimentally characterized and these studies agree well with their bioinformatically predicted annotations, with some discrepancies [22–37] (Table S2). In addition, the majority of the **F. succinogenes** CAZymes are predicted to have signal peptides, indicating that these enzymes are targeted outside of the cytoplasm.

#### Cellulases

**F. succinogenes** is predicted to possess 31 putative cellulase genes, including 10 members of GH5; 6 members of GH8; 9 members of GH9; 4 members of GH45; and 2 member of GH51 (Table 2). An analysis of the GHs predicted in the CAZy database for *C. thermocellum*, another prolific cellulose degrader, reveals a similar number of GH5 members, but only one member of GH8 and no members of GH45 or GH51. The GH45 members are especially significant, as these represent half of the predicted prokaryotic GH45 genes in the CAZy database. The most significant observation is that none of these 31 putative cellulase genes contain CBMs associated with binding to crystalline cellulose (CBM1, CBM2 or CBM3); in fact, the majority contains no recognizable CBM domains. A single xylanase-carboxymethyl cellulase has been reported to contain a GH5 and a CBM2 domain (AAC06197.1) [38], but BLAST analysis of this sequence reveals that it is not present in the **F. succinogenes** S85 genome. None of the cellulase genes contain domains that are homologous to known dockerin domains, and no genes with homology to scaffoldins were found in the genome, confirming the absence of cellulosomal structures in **F. succinogenes**. There was also no evidence of membrane anchoring or cell wall anchoring domains present in any of the cellulase genes. Further analysis of these...
activity is perhaps surprising, but is commensurate with the reports predicted to belong to family PL10 but had cellulolytic activity assays on other putative glycoside hydrolases as shown in Table 2 by performing cloning, expression, and carbohydrate hydrolysis experimentally [22,26–30] and we contributed to this growing list ‘atypical’ activities, including GH9 [29] and GH43 [41] enzymes. Cellulases may be targeted to one or more extra cytoplasmic produced by F. succinogenes this bacteria and the reported inability of the extracellular proteins adherence as an absolute requirement for cellulose degradation in peptide in many of these cellulases. Given the importance of cellulases using SignalP [39] reveals the presence of a signal component regulator (Fisuc_1517) found upstream of the GH5 cellulase Fisuc_0857, a sigma-24 extracellular cytoplasmic factor (ECF) transcriptional regulator (Fisuc_1804) found in other portions of the genome transcriptional regulators located in other portions of the genome includes two metal-dependent response regulators and a sugar transporter. Given the paucity of transcriptional regulators near known cellulase genes in F. succinogenes, it is likely that transcriptional regulators located in other portions of the genome contribute to the modulation of genes encoding for cellulases.

Table 1. Polysaccharide degradation and utilization by Fibrobacter succinogenes S85.

| Polysaccharide | Source | Linkage pattern* | mM succinate produced | mg reducing sugar accumulated (g added substrate) |
|---------------|--------|------------------|-----------------------|-----------------------------------------------|
| Hydrolyzed and utilized: | | | | |
| Cellulose | Wood | β-1,4-Glc | 20.6b | 4.9c |
| Hydrolyzed, but not utilized: | | | | |
| Homoxylan | Tobacco stalk | β-1,4-Xyl | 0.1 | 381.8 |
| Xylan | Larch | β-1,4-Xyl (with 4-O-MeGlcA substituents) | <0.1 | 328.6 |
| Glucomannan | Salep | Mixed β-1,4-Glc and –Man | 0.4 | 403.9 |
| Xyloglucan | Tamarind | β-1,4-Glc (with β-1,6-Xyl substituents) | 0 | 259.5 |
| Pectin | Citrus | α-1,4-GalUA (with substantial methylation) | 0.1 | 167.7 |
| Inulin | Chicory | β-2,1-Fru | 1.3 | 532 |
| Phlein (Fructan) | Orchardgrass | β-2,6-Fru | 0.3 | 13.8 |
| Not hydrolyzed or utilized: | | | | |
| Arabinogalactan (type II) | (not specified) | β-1,3 – Gal with some β-1,6 – Gal sidechains capped with various monosaccharides | 0.2 | 0 |
| Curdlan | Agrobacterium | β-1,3 – Glc | <0.1 | 0 |
| Laminarin | Brown algae | β-1,3 – Glc and β-1,6 – Glc (mixed linkage, 3:1) | <0.1 | 0 |
| Chitosan | Crab shells | β-1,4-GlcNH2 and N-AcGlcNH2 | <0.1 | 0 |
| Starch | Potato | α-1,4-Glc (with α-1,6 branches) | <0.1 | 0 |

*Monosaccharides and derivatives: Fru, fructose; Gal, galactose; GalUA, galacturonic acid, Glc, glucose; GlcNH2, glucosamine; Man, mannose; N-AcGlcNH2, N-acetylglucosamine; 4-O-MeGlcA, 4-methoxyglucuronic acid; Xyl, xylose.

bMean of Avicel (20.6 mM), Amorphous cellulose (21.5 mM) and Cellulose II (19.9 mM).

cMean of Avicel (6.2 mg/g), Amorphous cellulose (21.5 mM) and Cellulose II (0.8 mg/g).

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Some of these predicted cellulases have been previously verified experimentally [22,26–30] and we contributed to this growing list by performing cloning, expression, and carbohydrate hydrolysis assays on other putative glycoside hydrolases as shown in Table 2 and S2. We experimentally characterized a total of six cellulases including four GH5s, one GH8, and Fisuc_0505, which is predicted to belong to family PL10 but had cellulolytic activity in our assays (Table S2). The discovery of a PL10 with cellulolytic activity is perhaps surprising, but is commensurate with the reports of other GHs in F. succinogenes that have been described as having ‘atypical’ activities, including GH9 [29] and GH43 [41] enzymes.

An unanswered question regarding cellulose degradation by F. succinogenes is how these cellulase genes are regulated at the transcriptional level. In many polysaccharide-degrading bacteria, clusters of cellulases are often accompanied by transcriptional regulators that modulate their gene expression [42–45]. We analyzed clusters of genes surrounding cellulase-encoding genes in F. succinogenes and found only a handful had recognizable transcriptional regulators. These include an ArsR-type one-component regulator (Fisuc_0782) found upstream of a GH5 cellulase Fisuc_0786, a BadM-type transcriptional regulator (Fisuc_0900) found downstream of the GH5 cellulase Fisuc_0857, a sigma-24 extracellular cytoplasmic factor (ECF) transcriptional regulator (Fisuc_1517) found upstream of the GH5 cellulase Fisuc_1523, and a two-component sigma54-like response regulator (Fisuc_0397) found downstream of the GH9 cellulase Fisuc_0393. Any or all of these response regulators could potentially play a role in cellulase gene expression in F. succinogenes. Finally, a large number of cellulase (and hemicellulase) genes are found clustered between Fisuc_1762 and Fisuc_1804, which includes two metal-dependent response regulators and a sugar transporter. This cluster, which has been previously characterized [46], is likely controlled by these two response regulators, modulating the expression of both the GHs and the sugar transporter. Given the paucity of transcriptional regulators near known cellulase genes in F. succinogenes, it is likely that transcriptional regulators located in other portions of the genome contribute to the modulation of genes encoding for cellulases.

endo-Hemicellulases

F. succinogenes possesses genes with a wide range of annotated hemicellulolytic activities including a large number of xylanases, arabinoxylanases, mannanases, curdlanases (β-1,3 glucanases), licheninases (β-1,4 glucanases), and xyloglucanases; these enzymes come from a range of GH families including GH10, GH11, GH18, and GH26 (Table 3). The annotated functions of these genes may not represent their true in vivo catalytic activities because of their low homology to characterized enzymes from other species. For example, when we cloned and expressed an annotated pectinase gene, Fisuc_0678, we found that it possessed cellulase and glucanase activity, but no pectinase activity (Table S2). To confirm the activity of other hemicellulases, we also characterized two GH26 β-mannanases (Fisuc_0727 and Fisuc_0729) as shown in Table 3.

Our hydrolysis and utilization assays reveal F. succinogenes cannot metabolize any products produced by these endo-hemicellulases
(Table 1), and this is further underscored by the lack of genes necessary to both transport and metabolize these carbohydrates. For example, *F. succinogenes* is missing the genes necessary for xylose and xylodextrin transport (i.e. a xylose permease) into the cell, and further lacks a xylose isomerase (EC 5.3.1.5) to convert xylose into xylulose. As a result, we surmise that the function of these *endo*-hemicellulases is to serve as a form of biomass pretreatment by removing hemicelluloses and making cellulose microfibrils accessible to attack by the organism. Unlike the cellulases, many of the hemicellulases (GH10 and GH26) contain CBM6 or CBM35 domains that are known to bind to single chain substrates. These may increase the activity of the enzymes on the insoluble hemicellulose substrates present in the cell walls of biomass. Recently a novel domain, designated *F. succinogenes*-specific paralogous module 1 (FPm-1) was identified in 24 genes of *F. succinogenes* [46]. The function of this positively charged, basic domain located at the C-terminus of the proteins is unclear, but may be related to transport and localization of these proteins to the outer surface of the cell, or be involved in binding of the proteins to the outer membrane.

In general, the *endo*-hemicellulases, like the cellulases, contain a majority of enzymes with signal peptides. This fits with a general model of *F. succinogenes* hemicellulases localized to the outer membrane, which would facilitate the removal of these carbohydrates to gain access to cellulose.

**exo-Hemicellulases**

One would expect that an organism that is unable to metabolize the monosaccharides released by *exo*-hemicellulases would possess few of these genes. Yet *F. succinogenes* possesses an unexpectedly large number of *exo*-acting hemicellulolytic enzymes: 2 members of GH2; 3 members of GH3; and 11 members of GH43 (Table 3). Like the *endo*-hemicellulases, many of the hemicellulases (GH43) contain CBM6 domains, known to bind to single chain substrates. These GH43 family members contain β-xylidosidases and α-arabinofuranosidases; the CBM6 regions may improve synergistic

| CAZy Family | Fisuc Locus | FSU Locus* | CBM Family | Signal Peptide | In Silico Prediction | Characterized Activity | Reference |
|-------------|-------------|------------|------------|----------------|----------------------|------------------------|-----------|
| GH5         | Fisuc_0786  | FSU_1228   | –          | Yes            | cellulase            | cellulase              | This work |
| GH5         | Fisuc_0897  | FSU_1346   | –          | Yes            | cellulase            | cellulase              | [24]      |
| GH5         | Fisuc_1224  | FSU_1685   | –          | Yes            | cellulase            | cellulase              | This work |
| GH5         | Fisuc_1523  | FSU_2005   | –          | Yes            | cellulase            | cellulase              | This work |
| GH5         | Fisuc_1584  | FSU_2070   | –          | No             | cellulase            | –                      | [23]      |
| GH5         | Fisuc_1661  | FSU_2150   | –          | Yes            | endoglucanase        | –                      | –         |
| GH5         | Fisuc_2011  | FSU_2534   | –          | Yes            | cellulase            | –                      | –         |
| GH5         | Fisuc_2230  | FSU_2772   | –          | Yes            | endoglucanase        | cellulase              | [26]      |
| GH5         | Fisuc_2364  | FSU_2914   | –          | Yes            | cellulase            | cellulase              | [27]      |
| GH5         | Fisuc_3081  | FSU_0347   | –          | Yes            | cellulase            | cellulase              | This work |
| GH8         | Fisuc_0207  | FSU_0613   | –          | No             | endoglucanase        | xylanase               | [22]      |
| GH8         | Fisuc_0241  | FSU_0651   | –          | No             | endoglucanase        | cellulase              | This work |
| GH8         | Fisuc_0471  | FSU_0889   | –          | Yes            | endoglucanase        | xylanase               | [22]      |
| GH8         | Fisuc_1219  | FSU_1680   | –          | Yes            | endoglucanase        | –                      | –         |
| GH8         | Fisuc_1802  | FSU_2303   | –          | No             | endoglucanase        | cellulase              | [27]      |
| GH8         | Fisuc_2579  | FSU_3149   | –          | Yes            | β-glucanase          | cellulase              | [22]      |
| GH9         | Fisuc_0057  | FSU_0451   | –          | No             | β-glucanase          | cellulase              | [25]      |
| GH9         | Fisuc_0393  | FSU_0809   | –          | No             | cellulase            | –                      | –         |
| GH9         | Fisuc_0394  | FSU_0810   | –          | No             | cellulase            | –                      | –         |
| GH9         | Fisuc_1531  | FSU_2013   | –          | Yes            | cellulase            | –                      | –         |
| GH9         | Fisuc_1859  | FSU_2361   | –          | Yes            | β-glucanase          | cellulase              | [28]      |
| GH9         | Fisuc_1860  | FSU_2362   | –          | Yes            | cellulase            | –                      | [28]      |
| GH9         | Fisuc_2033  | FSU_2558   | –          | Yes            | endoglucanase        | cellulase              | [29]      |
| GH9         | Fisuc_2362  | FSU_2912   | –          | Yes            | cellulase            | cellulase              | [30]      |
| GH9         | Fisuc_2876  | FSU_0134   | –          | No             | cellulase            | –                      | –         |
| GH45        | Fisuc_1425  | FSU_1893   | –          | No             | cellulase            | cellulase              | [22]      |
| GH45        | Fisuc_1426  | FSU_1894   | –          | No             | cellulase            | xylanase               | [22]      |
| GH45        | Fisuc_1473  | FSU_1947   | –          | Yes            | cellulase            | –                      | –         |
| GH45        | Fisuc_1933  | FSU_2443   | –          | Yes            | cellulase            | –                      | –         |
| GH51        | Fisuc_2081  | FSU_2610   | –          | Yes            | endoglucanase        | cellulase              | [22]      |
| GH51        | Fisuc_3111  | FSU_0382   | CBM11, CBM30| Yes            | β-glucanase          | cellulase              | [31]      |

*FSU locus tags refer to the equivalent ORF call in the *F. succinogenes* genome sequence project described by GenBank accession: CP002158.
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interactions of the GH43 enzymes with the endo-hemicellulases for degradation of insoluble substrates. The function of these enzymes may be to provide growth substrates for the microbial consortia present in the rumen that supplies the essential branched and odd-chain length volatile fatty acids needed by F. succinogenes. Surprisingly, F. succinogenes appears not to contain any putative alpha-glucuronidase genes. This suggests that the specificities of F. succinogenes endo-hemicellulases and exo-hemicellulases may be such that they are able to bypass alpha-glucuronic acid residues in substituted xylans. Alternately, the organism may utilize an unrecognized gene product with alpha-glucuronidase to perform the hydrolysis. In addition to cataloguing these exo-hemicellulases, we characterized and verified the activity of two GH43 enzymes, including Fisuc_1769 as an arabinoxylanase and Fisuc_1994 as an arabinase (Table 3).

Table 3. Known endo- and exo-hemicellulases predicted from the Fibrobacter succinogenes S85 genome.

| CAZy Family | Fisuc Locus | FSU Locus* | CBM Family | Basic Terminal domaina | Signal Peptide | In Silico Prediction | Characterized Activity | Reference |
|-------------|-------------|------------|------------|------------------------|---------------|----------------------|-----------------------|-----------|
| GH2         | Fisuc_1788  | FSU_2288   | –          | BTD                   | Yes           | β-galactosidase       | –                     | –         |
| GH2         | Fisuc_3049  | FSU_0315   | –          | –                     | No            | β-galactosidase β-galactosidase | This work |
| GH3         | Fisuc_1751  | FSU_2249   | –          | –                     | Yes           | β-xylosidase          | –                     | –         |
| GH3         | Fisuc_1985  | FSU_2508   | –          | –                     | Yes           | β-xylosidase          | –                     | –         |
| GH3         | Fisuc_2065  | FSU_2592   | –          | –                     | Yes           | β-xylosidase cellulobiosidase | This work |
| GH10        | Fisuc_0754  | FSU_1192   | –          | –                     | Yes           | xylanase              | –                     | –         |
| GH10        | Fisuc_0757  | FSU_1195   | –          | –                     | Yes           | xylanase              | –                     | –         |
| GH10        | Fisuc_1791  | FSU_2292   | CBM6       | BTD                   | Yes           | xylanase              | –                     | [32]      |
| GH10        | Fisuc_1793  | FSU_2293   | CBM6       | BTD                   | Yes           | xylanase xylanase     | [32]      |
| GH10        | Fisuc_1794  | FSU_2294   | CBM6       | BTD                   | Yes           | xylanase xylanase     | [32]      |
| GH10        | Fisuc_2303  | FSU_2851   | –          | BTD                   | Yes           | endoglucanase         | –                     | –         |
| GH10        | Fisuc_2992  | FSU_0257   | –          | –                     | Yes           | cellulase             | –                     | [33]      |
| GH11        | Fisuc_0362  | FSU_0777   | –          | BTD                   | Yes           | xylanase xylanase     | [34]      |
| GH11        | Fisuc_2201  | FSU_2741   | –          | BTD                   | Yes           | xylanase xylanase     | [22]      |
| GH11        | Fisuc_2442  | FSU_3006   | –          | –                     | Yes           | xylanase xylanase     | [22]      |
| GH18        | Fisuc_1530  | FSU_2012   | –          | –                     | Yes           | chitinase cellulose binding | –                     | [35]      |
| GH18        | Fisuc_2465  | FSU_0303   | –          | –                     | Yes           | chitinase             | –                     | –         |
| GH26        | Fisuc_0727  | FSU_1165   | CBM35      | –                     | Yes           | mannanase β-mannanase | This work |
| GH26        | Fisuc_0729  | FSU_1167   | CBM35      | –                     | Yes           | mannanase β-mannanase | This work |
| GH26        | Fisuc_0730  | FSU_1168   | –          | –                     | Yes           | mannanase β-mannanase | [22]      |
| GH26        | Fisuc_1266  | FSU_1729   | –          | –                     | Yes           | mannanase             | –                     | –         |
| GH26        | Fisuc_1688  | FSU_2181   | CBM35      | –                     | Yes           | mannanase             | –                     | –         |
| GH43        | Fisuc_1762  | FSU_2262   | CBM6       | BTD                   | No            | β-xylosidase          | –                     | –         |
| GH43        | Fisuc_1763  | FSU_2263   | CBM6       | BTD                   | Yes           | β-xylosidase          | –                     | –         |
| GH43        | Fisuc_1764  | FSU_2264   | CBM6       | BTD                   | Yes           | β-xylosidase          | –                     | [22]      |
| GH43        | Fisuc_1769  | FSU_2269   | CBM6       | BTD                   | No            | β-xylosidase arabinoylanase | This work |
| GH43        | Fisuc_1775  | FSU_2274   | CBM6       | BTD                   | Yes           | β-xylosidase          | –                     | –         |
| GH43        | Fisuc_1994  | FSU_2517   | CBM35, CBM61 | –                     | Yes           | β-xylosidase arabinase | This work |
| GH43        | Fisuc_1997  | FSU_2520   | –          | –                     | No            | xylanase              | –                     | –         |
| GH43        | Fisuc_1998  | FSU_2521   | –          | –                     | Yes           | β-xylosidase          | –                     | –         |
| GH43        | Fisuc_1999  | FSU_2522   | CBM35      | –                     | No            | β-xylosidase          | –                     | –         |
| GH43        | Fisuc_2621  | FSU_3190   | –          | –                     | Yes           | β-xylosidase          | –                     | –         |
| GH43        | Fisuc_2622  | FSU_3191   | CBM35      | –                     | Yes           | β-xylosidase          | –                     | –         |
| GH43        | Fisuc_2623  | FSU_3192   | CBM35      | –                     | Yes           | β-xylosidase          | –                     | –         |
| GH43        | Fisuc_2886  | FSU_0145   | –          | –                     | Yes           | β-xylosidase          | –                     | –         |
| GH43        | Fisuc_2929  | FSU_0192   | CBM6       | BTD                   | No            | xylanase              | –                     | –         |

*FSU locus tags refer to the equivalent ORF call in the F. succinogenes genome sequence project described by GenBank accession: CP002158.

aBasic Terminal Domain (BTD) is also known as F. succinogenes-specific paralogous module 1 (FPm-1) [46].

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The Fibrobacter succinogenes S85 Genome

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with direct access to cellulose, which it actively degrades and metabolizes.

Carbohydrate esterases

In addition to the large number of \textit{endo-} and \textit{exo-}hemicellulases present in the \textit{Fibrobacter succinogenes} genome, the genome sequence also predicts genes for 17 CEs from families CE1, CE2, CE6, CE8, CE12, and CE15 (Table 4). All but 3 of the 17 CEs are predicted to have signal peptides, indicating their secretion outside the cytoplasm. Like the \textit{exo-}hemicellulases, many of these genes (8 of 17) contain CBM modules that may assist in binding to and degrading of insoluble hemicelluloses and pectins. The \textit{in vivo} substrates of these individual CE enzymes are unclear, because many of these genes show low homology to known and characterized esterases. The presence of CBM modules and signal peptides suggest these proteins may act synergistically with \textit{endo-}hemicellulases and \textit{exo-}hemicellulases in the degradation of xylan and pectin by cleaving acetic and ferulic acid esters linkages. A number of these enzymes have been characterized [47–49], and \textit{in vitro} activity has been demonstrated on synthetic substrates such as p-nitrophenyl acetate, xylose tetraacetate, or glucose pentacetate. In addition to cataloguing these CEs, we characterized and verified the activity of three putative esterases including a CE1 (Fisuc\_1771), a CE2 (Fisuc\_1641), and a CE6 (Fisuc\_2534). All were found to possess acetyl esterase activity on p-nitrophenyl acetate (Table 4).

Non-catalytic CBM-containing proteins

The genome of \textit{Fibrobacter succinogenes} contains seven genes that appear to encode for CBM domains with no apparent catalytic function (Table 5). These include members of CBM4, CBM6, CBM30 and CBM51. Members of CBM4 and CBM30 domains typically bind to single chain cellulose, CBM6 to single chain hemicelluloses, and family CBM51 to galactose. The function of these proteins is unclear.

Non-CBM-containing adherence proteins

In addition to these non-catalytic CBM proteins, \textit{Fibrobacter succinogenes} is known to produce a number of cellulose-binding proteins localized to the cell surface that facilitates its adherence to crystalline cellulose [33,50]. Previous proteomic analyses of \textit{Fibrobacter succinogenes} outer membrane proteins identified a cellulose-binding protein potentially involved in this process [50], and inhibition of this protein using antibodies significantly reduced the ability of \textit{Fibrobacter succinogenes} to bind to cellulose [50]. Structure-based analysis of this protein revealed a specific domain with strong homology to a domain in the slime mold \textit{Dictyostelium discoideum}, and this protein was annotated as a fibro-slime-domain-containing protein [33]. A total of 10 fibro-slime proteins were identified in that analysis, and we searched the \textit{Fibrobacter succinogenes} genome for evidence of the underlying genes encoding these proteins. We confirmed the presence of these 10 paralogs and mapped them to 10 different genes in the \textit{Fibrobacter succinogenes} genome [Fisuc\_0377, Fisuc\_1326, Fisuc\_1327, Fisuc\_1474, Fisuc\_1475, Fisuc\_1979, Fisuc\_2031, Fisuc\_2293, Fisuc\_2471, Fisuc\_2484]. The large number of paralogous fibro-slime genes in the \textit{Fibrobacter succinogenes} genome suggests that they may play an important role in the ability of \textit{Fibrobacter succinogenes} to bind to cellulose.

An analysis of these genes on the \textit{Fibrobacter succinogenes} genome shows that they are typically surrounded by groups of hypothetical proteins with the exception of the gene encoding fibro-slime

\begin{table}
\centering
\begin{tabular}{|c|c|c|c|c|c|c|c|}
\hline
CAZy Family & Fisuc Locus & FSU Locus\textsuperscript{a} & CBM Family & Basic Terminal domain\textsuperscript{b} & Signal Peptide & In Silico Prediction & Characterized Activity & Reference \\
\hline
CE1 & Fisuc\_1771 & FSU\_2270 & CBM4 & BTD & Yes & esterase & esterase & This work \\
\hline
CE1 & Fisuc\_1568 & FSU\_2052 & – & – & Yes & esterase & – & – \\
\hline
CE1 & Fisuc\_1569 & FSU\_2054 & – & – & Yes & esterase & – & – \\
\hline
CE1 & Fisuc\_1768 & FSU\_2268 & – & – & Yes & feruloyl esterase & – & – \\
\hline
CE1 & Fisuc\_1948 & FSU\_2468 & – & – & Yes & esterase & – & – \\
\hline
CE1 & Fisuc\_2396 & – & CBM4 & – & Yes & esterase & – & – \\
\hline
CE2 & Fisuc\_1641 & FSU\_2130 & – & – & Yes & esterase & esterase & This work \\
\hline
CE6 & Fisuc\_1766 & FSU\_2266 & CBM6 & BTD & Yes & acetyl xylan esterase & acetyl xylan esterase & [36] \\
\hline
CE6 & Fisuc\_1767 & FSU\_2267 & CBM6 & BTD & Yes & acetyl xylan esterase & acetyl xylan esterase & [36,37] \\
\hline
CE6 & Fisuc\_2315 & FSU\_2864 & – & – & Yes & esterase & – & – \\
\hline
CE6 & Fisuc\_2534 & FSU\_3103 & CBM6 & BTD & Yes & acetyl xylan esterase & xylan esterase & This work \\
\hline
CE6 & Fisuc\_2800 & FSU\_0054 & – & – & Yes & acetyl xylan esterase & – & – \\
\hline
CE8 & Fisuc\_0679 & FSU\_1115 & CBM35 & – & No & pectin esterase & – & – \\
\hline
CE12 & Fisuc\_1995 & FSU\_2518 & – & – & Yes & esterase & – & – \\
\hline
CE12 & Fisuc\_2478 & FSU\_3044 & CBM35 & – & Yes & esterase & – & – \\
\hline
CE12 & Fisuc\_2479 & FSU\_3045 & CBM35 & – & No & esterase & – & – \\
\hline
CE15 & Fisuc\_2348 & FSU\_2898 & – & – & No & esterase & – & – \\
\hline
\end{tabular}
\caption{Known carbohydrate esterases predicted from the \textit{Fibrobacter succinogenes} S85 genome.}
\textsuperscript{a}FSU locus tags refer to the equivalent ORF call in the \textit{Fibrobacter succinogenes} S85 genome. \textsuperscript{b}Basic Terminal Domain (BTD) is also known as \textit{Fibrobacter succinogenes}-specific paralogous module 1 (FPm-1) [46].
\label{table4}
\end{table}
protein Fisuc_2484, which appears in a putative operon (Fisuc_2484-Fisuc_2487) containing a CBM6, a hypothetical protein, and an ABC-transporter related protein. Given that CBM6 domains bind to hemicelluloses, this fibro-slime protein may play a role in hemicellulose degradation. Finally, there are two clusters of fibro-slime proteins that have putative cellulases located near by. These include the clustered genes encoding fibro-slime proteins Fisuc_1474 and Fisuc_1475, which has a GH45 cellulase immediately upstream (Fisuc_1473) and the gene encoding fibro-slime protein Fisuc_2031, which has a GH9 cellulase (Fisuc_2033) located downstream. The close proximity of these genes to two cellulases may indicate a role in their activity.

Recently, type IV pilin proteins have been implicated in F. succinogenes adherence [33], as adherence deficient mutants lack some of these proteins. A survey of the F. succinogenes genome reveals the presence of a small number of type IV pilin genes (Fisuc_0251-Fisuc_0253; Fisuc_0956; and Fisuc_1016-Fisuc_1018). Although a complete set of genes necessary to produce pilin are not present in the genome, these pilin genes may be used by F. succinogenes to adhere to cellulose in a manner similar to how Escherichia coli and other gram negative bacteria adhere to solid substrates [51].

The reliance of F. succinogenes on membrane-bound adherence proteins may also be correlated to its protein secretion systems. F. succinogenes appears to contain the Sec-dependent and type II secretion pathways. In other bacteria such as the plant pathogen Xanthomonas campestris, the Sec-dependent and type II secretion systems are used to secrete hydrolytic enzymes like cellulases, chitinases, and pectate lyases to facilitate its pathogenic lifestyle [52]. Type II secretion pathways have also been implicated in the adherence of human pathogenic bacterial cells to the host epithelium [52], and given that F. succinogenes requires adherence to cellulose for efficient hydrolysis, this pathway may play a role in cell adherence.

Utilization of cellulose hydrolysis products

Cellodextrins liberated by the cellulolytic action of F. succinogenes can be transported into the cell for further processing via a cellodextrin-specific transporter. For example, the F. succinogenes genome encodes for extracellular-solute binding proteins that are known to participate in the active transport of solutes through the ABC active transport mechanism. These proteins, which are found attached to the cell surface and help recognize and bind specific solutes, could be used for cellodextrin transport. This type of transporter specificity is known in other cellulolytic bacteria including C. thermocellum [33], and an alignment of the known cellodextrin-specific extracellular-solute binding protein from this bacterium shows strong homology with an extracellular-solute binding protein (Fisuc_0617) in F. succinogenes (data not shown) [10]. Once incorporated into the cell, cellodextrins can be sequentially converted to glucose-1-phosphate by the action of cellodextrin phosphorylase, whose presence has been demonstrated in cell-free extracts. The F. succinogenes genome appears to contain only a single cellodextrin phosphorylase gene (Fisuc_2900), and physiological studies suggest that this enzyme can reversibly catalyze both degradation and synthesis of cellodextrins. This differs from the case of some cellulolytic bacteria such as C. thermocellum, in which cellodextrin is metabolized via a phospho-lyase and longer cellodextrins are metabolized by a second phosphorylase [54–56], but cellodextrins are not produced in significant quantities by either phosphorylase. An alternative pathway would be degradation of the cellodextrins in either the cell wall or periplasmic space by the chlorine-stimulated cellodextrinases (CLCB, Fisuc_2992) known to be required for degradation of crystalline cellulose [57]. Cellodextrin would then be transported via a similar system to that proposed for cellodextrins.

The catabolism of cellulose hydrolysis products by F. succinogenes was characterized further by performing a metabolic reconstruction analysis using the software program PRIAM [58] to generate KEGG pathway maps [59]. F. succinogenes is a strict anaerobe that produces a mixture of succinate, acetate and formate as fermentation end products [3,7]. The metabolic reconstruction shows this strain contains both a complete Embden-Meyerhof-Parnas pathway and an incomplete citric acid cycle, lacking both an α-ketoglutarate dehydrogenase and a succinyl-CoA-synthase (Figure 2). This agrees with enzymatic evidence [60] that F. succinogenes utilizes its incomplete citric acid cycle for the production of succinate, the bacterium’s major fermentative end product. Specifically, the reconstruction shows that F. succinogenes contains both a phosphoenolpyruvate carboxykinase (Fisuc_2949) and a pyruvate carboxylase (Fisuc_0845) that can reversibly convert PEP and pyruvate, respectively, to oxaloacetate, which is sequentially converted to malate, fumarate, and succinate. The reconstruction also suggests that energy production is likely facilitated through the transfer of electrons from a carrier such as menaquinone to fumarate through the action of a membrane-bound fumarate reductase (Fisuc_2993 and Fisuc_2994), resulting in the production of succinate. Previously, it was reported that fumarate reduction can be

| CAZY Family | Fisuc Locus | FSU Locus* | Basic Terminal domaina | Signal Peptide | In Silico Prediction | Characterized Activity | Reference |
|------------|------------|------------|------------------------|----------------|----------------------|------------------------|----------|
| CBM4       | Fisuc_1931 | FSU_2441   | –                      | No             | carbohydrate binding module | –                     | –        |
| CBM6       | Fisuc_1774 | FSU_2273   | –                      | Yes            | carbohydrate binding module | –                     | –        |
| CBM6       | Fisuc_2485 | FSU_3051   | BTD                    | Yes            | carbohydrate binding module | none found            | This work |
| CBM30      | Fisuc_1525 | FSU_2007   | –                      | Yes            | carbohydrate binding module | –                     | –        |
| CBM51      | Fisuc_0215 | FSU_0622   | –                      | No             | carbohydrate binding module | –                     | –        |
| CBM51      | Fisuc_0401 | FSU_0816   | –                      | No             | carbohydrate binding module | –                     | –        |
| CBM51      | Fisuc_1656 | FSU_2145   | –                      | Yes            | carbohydrate binding module | –                     | –        |

*FSU locus tags refer to the equivalent ORF call in the F. succinogenes genome sequence project described by GenBank accession: CP002158.

Basic Terminal Domain (BTD) is also known as F. succinogenes-specific paralogous module (FPm) [46].

Table 5. Known genes predicted to encode only carbohydrate binding modules from the Fibrobacter succinogenes S85 genome.
coupled to hydrogen oxidation in *F. succinogenes* [61]. However, a search of the *F. succinogenes* genome for the presence of a hydrogenase using over 700 previously identified enzymes [62], did not find any evidence for a hydrogenase in *F. succinogenes*. *F. succinogenes* can also produce formate and acetyl-CoA through the action of formate C-acetyltransferase (Fisuc_1044) with the assistance of a pyruvate-formate lyase activating enzyme (Fisuc_1047).

The metabolic reconstruction predicts that *F. succinogenes* does not have an Entner-Doudoroff pathway and that the glyoxylate shunt is also absent. This bacterium also appears to have incomplete pathways for the utilization of galactose, mannose, fructose and pentose sugars. These predictions are in agreement with our finding that *F. succinogenes* is unable to grow on any saccharides other than cellulose and its soluble components (glucose, cellobiose, and cello dextrans) (Table 1).

![Figure 2](https://www.plosone.org/figure/Fibrobacter-succinogenes-S85-Genome-1440x1080.png)

**Figure 2. An overview of metabolism and transport in *Fibrobacter succinogenes* S85.** Enzymes missing from metabolic pathways are indicated with a red cross. The major fermentative products succinate, acetate, and formate are shown with gray arrows indicating their export out of the cell. Predicted transporters are also shown, including sodium ion channel protein transporters in purple, ABC transporters in red, sec-dependent protein export in green, and other substrates in blue. Export or import of solutes is shown through the direction of the arrow through the transporter. Energy coupling mechanisms are also shown, including solutes transported by channel proteins; secondary transporters with two arrows into the cell indicating the solute and coupling ion; ATP-driven transporters with an ATP hydrolysis reaction; and transporters with an unknown energy-coupling mechanism, shown with a single arrow. Some multi-step pathways are not fully-represented, and are denoted with orange arrowheads. Abbreviations: AA, amino acids; Ala, alanine; Met, methionine; Pro, proline; PRPP, 5′-phospho-α-D-ribose 1-diphosphate; PEP, phosphoenolpyruvate; Meso 2,6 DAP, meso-2,6- diaminopimelic acid.

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Comparison of polysaccharide-degrading strategies with those of other ruminal bacteria

In addition to F. succinogenes, several other well-known polysaccharide-degrading bacteria have been isolated from the rumen including Butyryrobovibrio proteoclasticus B316 [63], Prevotella ruminicola 23 [64] and Ruminococcus flavefaciens FD-1 [65]. These species are thought to work synergistically with F. succinogenes to degrade plant biomass in the rumen, and utilize different strategies to deconstruct polysaccharides [2]. To gain an understanding of these systems we compared the carbohydrate-degrading potential of these bacteria as shown in Table S3. In general, these bacteria have an apparent specificity in their carbohydrate-active enzymes with some overlap. F. succinogenes and R. flavefaciens FD-1 are known to be prolific cellulose-degrading specialists, whereas P. ruminicola 23 is a generalist capable of degrading and utilizing many different polysaccharides, but not cellulose [64]. B. proteoclasticus B316 also cannot degrade cellulose, but can degrade and utilize xylan, starch, and pectin [63]. These capabilities are reflected in their carbohydrate-active enzyme profiles.

Both F. succinogenes S85 and R. flavefaciens FD-1 contain a number of cellulases from the same families including members of the GH5 and GH9. F. succinogenes also contains a number of enzymes in the GH8, GH45, and GH51 families classified as cellulases that are not found in R. flavefaciens, whereas a GH48 exoglucanase is present in R. flavefaciens FD-1 but not in F. succinogenes. The higher diversity of cellulases in F. succinogenes, relative to R. flavefaciens FD-1 may account for its ability to degrade all known allomorphs of cellulose, including the highly stable, chemically regenerated cellulose II. However, the relatively weak hydrolytic capacity of these enzymes to degrade crystalline cellulose in vitro and the in vivo requirement for direct adherence to cellulose, as discussed above, indicates that F. succinogenes must use other mechanisms such as adherence molecules or “atypical” cellulases such as the GH9s, which are thought to synergistically work with other cellulases [29].

B. proteoclasticus B316 and P. ruminicola 23 also have a number of enzymes within these GH families, but in much smaller numbers than are found in F. succinogenes and R. flavefaciens FD-1. These GH family enzymes appear to be xylanases and other hemicellulases, which corresponds to the reported inability of the former two organisms to degrade cellulose. These are complemented by other endo- and exo-hemicellulases in the families GH2, GH3, GH10, GH16, GH43, and GH53, with the largest numbers of enzymes within the GH43s; each bacterium is predicted to have at least 10 copies of GH43. In addition, B. proteoclasticus B316 and P. ruminicola 23 contain xylanases in the family GH11 and GH44 that are not found in either F. succinogenes or R. flavefaciens FD-1. The large diversity of hemicellulases within these 4 bacteria corresponds well to their known polysaccharide-degrading and utilization mechanisms, although F. succinogenes is the only one that degrades xylan without using its hydrolytic products.

Finally, B. proteoclasticus B316 and P. ruminicola 23 contain enzymes in the GH28, GH29, GH32, GH35, and GH38 families that are not found in either F. succinogenes or R. flavefaciens FD-1. These GHs include pectinases, fucosidases, fructanases, and mannoglycosidases, which correspond to their more generalist lifestyle of degrading and utilizing a wider variety of polysaccharides.

In addition to the GHs, the CBMs found in these four ruminal bacteria also support their polysaccharide degrading strategies. For example, F. succinogenes contains members of CBM11, CBM30, and CBM51 whereas B. proteoclasticus B316, P. ruminicola 23, R. flavefaciens FD-1 do not. Both CBM11 and CBM30 are known to bind to cellulose in F. succinogenes [31], providing further evidence for its ability to degrade cellulose. The only shared cellulose-binding CBMs between F. succinogenes and R. flavefaciens FD-1 is CBM4, which is known to bind to single chain cellulose. Furthermore, F. succinogenes has 20 copies of CBM6, more than 4-fold greater than any of the other three ruminal bacteria. These domains, which are found associated with hemicellulases in F. succinogenes, may indicate that they are the preferred modules for facilitating hemicellulose deconstruction. R. flavefaciens FD-1 and B. proteoclasticus B316 may also utilize organism-specific CBMs for hemicellulose and cellulose degradation, as CBM13 and nine copies of CBM2 are found in these bacteria, respectively. The apparent specificity of these CBMs in their respective organisms may indicate different strategies for binding to hemicelluloses such as xylan, or perhaps may reflect different preferences for different plant tissue types. P. ruminicola does not appear to have any organism-specific CBMs, when compared to the three other ruminal bacteria, but utilizes a variety of CBMs reflecting its more general polysaccharide-degrading lifestyle.

Discussion

Here we report for the first time, the complete genome sequence for the cellulolytic ruminal bacterium Fibrobacter succinogenes. This also represents the first complete genome of a bacterium belonging to the phylum Fibrobacteres. The rumen is an environment tailored for the conversion of plant biomass into volatile fatty acids usable by its host and it is apparent that ruminal microbes like F. succinogenes are specialized for this process. F. succinogenes, as its name implies, is a producer of succinate, and our metabolic reconstruction analysis confirms that, along with acetate and formate, are major fermentative end products.

However, unlike other ruminal bacteria that derive their energy from many different polysaccharide sources, F. succinogenes is specialized for using only cellulose. Our physiological assays and analysis of the F. succinogenes carbohydrate-degrading machinery reveals this property, and further suggests a specific model of plant polysaccharide deconstruction. The F. succinogenes genome encodes for a number of enzymes capable of degrading a wide array of polysaccharides and it is likely that it uses these to remove carbohydrates like xylan in order to gain access to cellulose. This is supported by our finding that while F. succinogenes can hydrolyze these substrates, it can not metabolize the end products as carbon sources. For example, F. succinogenes can hydrolyze xylan into xylose, but can not utilize this as an energy source because it lacks both a xylose permease for transport into the cell and a xylose isomerase.

The polysaccharide degrading strategy of F. succinogenes is markedly different from other cellulolytic bacteria, not only in its specialization on cellulose, but also in its method of cellulose degradation. Adherence of the bacterium to solid cellulose substrates appears to be a requirement for the degradation of cellulose [13,14], and little cellulase activity is detected in the culture medium [40]. F. succinogenes is one of only a few organisms reported to rapidly degrade all allomorphs of cellulose, including cellulose II [66]. Previous work has shown that the organism does not possess extensive secreted cellulases or cellulosomal structures like other ruminal organisms such as Ruminococcus flavefaciens (reviewed in [56]). Identified cellulolytic enzymes show low homology to cellulases from other organisms [29] and the F. succinogenes cellulases that have been cloned to date show poor performance on crystalline cellulose both alone and in combinations [29,67]. The lack of identifiable crystalline cellulose-binding domains and the poor performance of the identified enzymes on crystalline cellulose – despite active degradation of this substrate by whole cells – has led to the hypothesis that cellulose degradation
by this species relies on an unusual degree of cell-envelope synergy [56], and perhaps even utilizes a cell-based, non-enzymatic process [22]. A number of benefits appear to accrue from the novel lifestyle of F. succinogenes [68], but it is not yet clear how the genes identified in the organism contribute to its success. It is likely that the unconventional mode of cellulose degradation by F. succinogenes accrues at least in part from an unusual combination of cellulases distributed into certain families which are relatively poorly represented among microbes that employ more conventional modes of cellulose degradation.

Models of polysaccharide degradation for F. succinogenes

Several models can be proposed for cellulose degradation by F. succinogenes. Tenable models must be in accord with observations that F. succinogenes: i) produces trace amounts of measurable soluble cellulase activity in fermentations actively degrading cellulose [7]; ii) has an absolute requirement for adherence to effect cellulose degradation [13,14]; iii) is unable to degrade cellulose as cell-free extracts [40]; and iv) cannot bind to or degrade crystalline cellulose when certain non-cellulase genes are mutated [33]. Arguably, the two most studied systems for cellulose degradation are the soluble enzyme systems of *Trichoderma reesei* and the cellulolysosomal systems of bacteria like *C. thermocellum*. Both models can be eliminated for F. succinogenes because it does not produce soluble cellulases and our genomic analysis reveals no homologs to known cellulolysosomal proteins. Therefore, these models do not successfully meet all of the criteria required for F. succinogenes cellulase degradation, and we consider other possible models here.

One proposed model describes the growth of F. succinogenes as a biofilm on the cellulose surface, similar to the case of ruminococci [69]. F. succinogenes is thought to rely on fibro-slime proteins to attach to its substrate and initiate or support cellulose deconstruction. Proteins such as the fibro-slime [33,50] and type IV pilin structures [70] would facilitate cell-surface attachment to the substrate and mediate close contact of both GH and CBM-containing enzymes to polysaccharides. Other unidentified modules in the F. succinogenes genome may be expected to play similar roles to CBMs or the dockerins and scaffoldins that facilitate cellulolysosomal degradation in ruminal bacteria like *Ruminococcus*. The specific mechanism of substrate degradation has been suggested to proceed through the localization of cellulases and hemicellulases on the cell membrane. This may indeed be the case for hemicellulases, many of which contain the F. succinogenes-specific paralogous module 1. However, sequence analysis indicates that none of the cellulases possess this domain or any other previously-reported anchoring domain, making it unlike that they are attached to the membrane. This localization is also not supported by results from isolation of individual membrane fractions of the organism [71].

*F. succinogenes* also appears to employ ‘atypical’ cellulases that may obviate the need for extensive CBMs [29]. In particular, a GH9 cellulase has been characterized that synergistically increases the hydrolytic ability of other cellulases like Ce51 and Ce8B [27,29]. The large number of GH9s in the genome, and their potential localization to the cell membrane (Table 2) could act to increase the efficiency of many of *F. succinogenes* cellulases. The combination of adherence molecules, carbohydrate-binding molecules, and interacting cellulases may display a hydrolytic synergy that mediates efficient cellulose degradation as has been previously suggested [70,72]. However, it should be noted that cloned, expressed, purified and characterized *F. succinogenes* cellulases cannot significantly degrade crystalline cellulose, either alone or in combination, making it difficult to understand how they might function to quickly and effectively depolymerize all the allomorphs of cellulose.

In addition, two other cellulolytic mechanisms have been suggested. *Cytophaga hutchinsonii*, a bacterium within the *Cytophaga-Flavobacterium-Bacteroides* phylum that is phylogenetically-related to *F. succinogenes*, is known to associate tightly with cellulose. This bacterium has been proposed to degrade cellulose by disrupting cellulose fibers and taking up individual cellulose chains through the outer membrane [73]. Upon reaching the periplasmic space, these chains would be cleaved by endoglucanases. This presents an intriguing model for *F. succinogenes*, which could thus gain direct access to the hydrolytic products of cellulose (glucose, cellobiose, and cellooligosaccerides). However, *F. succinogenes* and *C. hutchinsonii* share few cellulase homologs and this model of cellulase degradation by *C. hutchinsonii* is thought to be facilitated by its ability to move in parallel across cellulose fibers using gliding motility [74], during which cellulose chains could be stripped from fibers as it glides across its substrate. In contrast, motility in *F. succinogenes* has not been demonstrated, nor did we find any known motility genes in its genome. If *F. succinogenes* were to employ an approach similar to *C. hutchinsonii*, it may be accomplished using the previously described fibro-slime proteins. In the slime mold *Dictyostelium discoideum*, slugs produce a cellulose sheath that enables it to move. The related fibro-slime genes in *F. succinogenes* may play a similar role, enabling it to move in an analogous manner without apparent motility in solution. In this regard, it is interesting that degradation of crystalline cellulose by *F. succinogenes* appears to occur along the crystallographic axis, suggesting a directionality of the hydrolytic process [13,14].

Alternatively, *F. succinogenes* may follow a model proposed for the α-proteobacterium *Sphingomonas* sp. A1 [75]. This bacterium appears to form “pits” across its cell membrane that act as channels that can import and depolymerize macromolecules like alginate [76]. Import of macromolecules through the membrane is thought to occur through two high-affinity periplasmic proteins facilitated by an ABC transporter. The degradation of macromolecules would occur in the cytoplasm. Comparison of the genes implicated in *Sphingomonas* sp. A1 pit formation with the *F. succinogenes* genome reveals a number of homologs including type IV pilin molecules and ABC transporters (data not shown). However, electron microscopic analysis of *F. succinogenes* [13] does not reveal the presence of pits as has been shown for *Sphingomonas* sp. A1. While it is possible that these pits may be too small to be detected using electron microscopy, one consideration is that alginate and cellulose are very different molecules with respect to their higher order structure, and this model would require the disassembly (decrystallization and delamination) of individual cellulose fibers or microfibrils before they could be imported through these pits.

Conclusion

The mechanism by which *F. succinogenes* degrades cellulose is not obvious and stands in stark contrast to the strategies used by other cellulolytic microbes. The availability of the *F. succinogenes* genome will serve to increase our understanding of its unique cellulose degrading properties and provide insight into the peculiar biology of this bacterium and its phylum, given the association of different *Fibrobacter* species in ruminants and other animals [4,77,78]. Furthermore, with the increasing number of ruminal bacterial genomes becoming available, we will be able to leverage this data to begin understanding how these microbes interact within the rumen and their impact on ruminant health and animal performance. Finally, from a biotechnological perspective, understanding how *F. succinogenes* accomplishes polysaccharide hydrolysis
will help inform our own efforts to optimally convert cellulosic material for the production of biofuels.

Materials and Methods

DNA extraction, genome sequencing and finishing

The type strain Fibrobacter succinogenes S85 (ATCC 19169T) was obtained from Dr. Cecil Forsberg (University of Guelph). We grew cultures of F. succinogenes in a modified Dehority medium [14] supplemented with 4 g cellulose/L for 48 h at 39 °C. Genomic DNA was then prepared as described by Stevenson and Weimer [5].

The genome of F. succinogenes S85 was sequenced at the DOE Joint Genome Institute (JGI) using a combination of Illumina [79] and 454 technologies [80]. An Illumina GAii shotgun library with reads of 477 Mb, a 454 Titanium draft library with average read length of 243 bases, and a paired end 454 library with average insert size of 20.5 Kb were generated for this genome. All general aspects of library construction and sequencing performed at the JGI can be found at http://www.jgi.doe.gov/. Illumina sequencing data was assembled with VELVET [81], and 103 additional reactions were necessary to close the final GenBank submitted annotation contains only 3,085 ORFs, which we report here. This difference is due to the GenBank standard submission process which removed two predicted ORFs considered to be spurious.

Whole-genome analysis

A protein comparison analysis was performed for the F. succinogenes genome by using BLAST [94] to query against a local database composed of proteins from 1,172 microbial genomes (http://www.ncbi.nlm.nih.gov/genomes/lproks.cgi?view=1, accessed: 06/17/2010). Those proteins that had a BLAST hit (e-value: 1e-05) were recorded and divided into two different categories: proteins that had a putative assigned function, and those that matched to hypothetical proteins. Proteins that did not have a significant BLAST hit were designated as species- or genus-specific proteins. These data were then used to generate a taxonomic distribution analysis as shown in Figure 1. This was done by counting the phyla/group membership for the top hit of each protein in the F. succinogenes genome against the local database of microbial proteomes and expressing this as a percentage of the total number of F. succinogenes proteins that could be assigned a closest taxonomic member.

A clusters of orthologous groups (COG) [20] analysis was performed using the F. succinogenes genome as follows. Predicted proteins from F. succinogenes were queried against a COG database retrieved from NCBI (ftp://ftp.ncbi.nih.gov/pub/rmndb/cdd/, accessed: 06/19/2010) using RPSBLAST [95] (e-value: 1e-05). The top RPSBLAST hit for each protein was tabulated and placed into each COGs respective category as shown in Table 2.

We also analyzed the predicted carbohydrate-active enzymes (CAZymes) [21] for F. succinogenes available through the CAZY database (http://CAZy.org, accessed: 03/01/2011). However, because there are many legacy enzymes in this database that have not been correlated to the current annotation of the genome, we used a custom Perl script to tabulate a single set of CAZymes that corresponds to the genome annotation as shown in Table 2.

Polysaccharide hydrolysis and growth measurements

Experiments were conducted under a CO2 gas phase in triplicate 60 mL glass serum vials that contained 10 mL of modified Dehority medium [96] supplemented with the indicated polysaccharide. Cultures were incubated without shaking at 39 °C for 72 h. Hydrolysis of polysaccharides was measured as release of reducing sugars by the dinitrosalicylic acid method [97], using glucose as standard. Growth on polysaccharides was not measured directly, but was instead measured as succinate production [98], using HPLC [96].

Enzyme cloning, expression, and characterization

Genes were cloned, amplified, and expressed as previously described [22]. Briefly, the presence of signal sequences within target genes was determined using the SignalP software [39]. Primers were then selected (Table S2) and these genes were amplified without signal sequences from DNA extracted from F. succinogenes. The resulting amplicons were cloned, expressed and characterized as described previously [22]. The range of substrates used for evaluation was expanded to include the following synthetic substrates: AZCL-Arabinian, AZCL-Arabinoxylan, AZCL-beta-Glucan, AZCL-Galactomannan, AZCL-HE Cellulose, AZCL-Xyloglucan, 4-methylumbelliferyl-β-D-celllobioside (MUC), 4-methylumbelliferyl-β-D-xylopyranoside (MIX), 4-methylumbelliferyl-β-D-glucopyranoside (MUG), pNP-β-glucoside, pNP-β-celllobioside, 4-methylumbelliferyl-β-D-arabinofuranoside (MUA), 4-methylumbelliferyl-β-D-lactopyranoside (MUL), 5-Bromo-4-chloro-3-indolyl-β-D-galactopyranoside (X-β-Gal, XAG), and 5-Bromo-4-chloro-3-indolyl-β-D-galactopyranoside
A total of 14 new GH family members, 3 new CE family members, 1 PL family member, and 1 CBM family member with no identified enzymatic activity investigated in this manner, as shown in Table S2. The observed activities of the cloned proteins are shown in Table S4.

Supporting Information

Table S1 Clusters of orthologous group (COG) analysis for Fibrobacter succinogenes S85.

Table S2 Carbohydrate-active enzymes encoded by the Fibrobacter succinogenes S85 genome.

Table S3 Comparison of carbohydrate-degrading enzymes (CAZymes) encoded by the genomes of the 4 ruminant bacteria Fibrobacter succinogenes S85 (Fsuc), Butyrivibrio proteoclasticus B316 (Bpro), Prevotella ruminicola 23 (Prum), and Ruminococcus flavefaciens FD-1 (Rfla).

Table S4 Primers used to clone selected CAZymes in F. succinogenes and their tested polysaccharide activity.

Text S1 Glycogen biosynthesis and utilization.

Text S2 Amino acid synthesis and nitrogen assimilation.

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Author Contributions

Conceived and designed the experiments: PJW GS DM PJB. Performed the experiments: GS PJW DMS FOA DM PJB. Analyzed the data: GS PJW DMS FOA DM PJB. Contributed reagents/materials/analysis tools: PJW GS DM PJB. Wrote the paper: GS PJW FOA DM PJB.
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