Hydrogenomics of the Extremely Thermophilic Bacterium 
Caldicellulosiruptor saccharolyticus

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Caldicellulosiruptor saccharolyticus is an extremely thermophilic, gram-positive anaerobe which ferments cellulose-, hemicellulose- and pectin-containing biomass to acetate, CO2, and hydrogen. Its broad substrate range, high hydrogen-producing capacity, and ability to coutilize glucose and xylose make this bacterium an attractive candidate for microbial bioenergy production. Here, the complete genome sequence of C. saccharolyticus, consisting of a 2,970,275-bp circular chromosome encoding 2,679 predicted proteins, is described. Analysis of the genome revealed that C. saccharolyticus has an extensive polysaccharide-hydrolyzing capacity for cellulose, hemicellulose, pectin, and starch, coupled to a large number of ABC transporters for monomeric and oligomeric sugar uptake. The components of the Embden-Meyerhof and nonoxidative pentose phosphate pathways are all present; however, there is no evidence that an Entner-Doudoroff pathway is present. Catabolic pathways for a range of sugars, including rhamnose, fucose, arabinose, glucuronate, fructose, and galactose, were identified. These pathways lead to the production of NADH and reduced ferredoxin. NADH and reduced ferredoxin are subsequently used by two distinct hydrogenases to generate hydrogen. Whole-genome transcriptome analysis revealed that there is significant upregulation of the glycolytic pathway and an ABC-type sugar transporter during growth on glucose and xylose, indicating that C. saccharolyticus coferments these sugars unimpeded by glucose-based catabolite repression. The capacity to simultaneously process and utilize a range of carbohydrates associated with biomass feedstocks is a highly desirable feature of this lignocellulose-utilizing, biofuel-producing bacterium.

Microbial hydrogen production from biomass has been recognized as an important source of renewable energy (13, 47). High-temperature microorganisms are well suited for production of biohydrogen from plant polysaccharides, as anaerobic fermentation is thermodynamically favored at elevated temperatures (17, 43). The extremely thermophilic bacterium Caldicellulosiruptor saccharolyticus DSM 8903, a fermentative anaerobe initially isolated from wood in the flow of a thermal spring in New Zealand, first received attention because of its capacity to utilize cellulose at its optimal growth temperature, 70°C (37). Further work showed that C. saccharolyticus (i) can utilize a wide range of plant materials, including cellulose, hemicellulose, starch, and pectin, (ii) has a very high hydrogen yield (almost 4 mol of H2 per mol of glucose) (14, 20, 48), and (iii) can ferment C5 and C6 sugars simultaneously. These features have led to the development of bioprocessing schemes based on C. saccharolyticus. For example, H2 production is now being investigated using a two-step process in which H2 and acetate are generated from biomass hydrolysates in one bioreactor and the acetate is fed to a second bioreactor and used by phototrophic organisms (Rhodobacter spp.) to produce additional H2 in the presence of light (10). To provide a basis for full exploitation of the biohydrogen-producing capacity of C. saccharolyticus, the complete genome of this organism was sequenced and analyzed in conjunction with transcriptome information for the bacterium grown on glucose and xylose. The comparative and functional genomics approach that was employed here is referred to as “hydrogenomics.” Insights arising from this effort revealed that C. saccharolyticus has the capacity to process and utilize a broad range of sugars and ultimately forms hydrogen from catabolism of these compounds.

MATERIALS AND METHODS

Cultivation and DNA isolation. C. saccharolyticus DSM 8903 (ATCC 43494) was cultured overnight at 70°C on DSMZ 640 medium with glucose (50 mM) as the carbon and energy source. Cells were harvested and genomic DNA was isolated by the method of Pitcher et al. (35) using guanidinium thiocyanate.
TABLE 1. General features of the C. saccharolyticus genome

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length of chromosome (bp)</td>
<td>2,970,275</td>
</tr>
<tr>
<td>G+C content (%)</td>
<td>35.3</td>
</tr>
<tr>
<td>Coding density (%)</td>
<td>86.3</td>
</tr>
<tr>
<td>Total no. of protein-encoding genes</td>
<td>2,679</td>
</tr>
<tr>
<td>Avg length of protein-encoding genes (bp)</td>
<td>958</td>
</tr>
<tr>
<td>Total no. of pseudogenes</td>
<td>92</td>
</tr>
<tr>
<td>Total no. of tRNA genes</td>
<td>46</td>
</tr>
<tr>
<td>Total no. of tRNA genes (no. of operons)</td>
<td>9 (3)</td>
</tr>
<tr>
<td>No. of CRISPR loci</td>
<td>9</td>
</tr>
</tbody>
</table>

Residual protein was removed in an additional purification step with sodium dodecyl sulfate and protease K, followed by chloroform-isomyl alcohol extraction and isopropanol precipitation.

**Genome sequencing and assembly.** High-molecular-weight genomic DNA was provided to the U.S. Department of Energy Joint Genome Institute (http://www.jgi.doe.gov/) for cloning and shotgun sequencing. A combination of small (average insert sizes, 3 and 8 kb) and large (40 kb, fosmid) insert libraries were prepared and used for analysis, as indicated at http://www.jgi.doe.gov/.

**Genome annotation and comparative analysis.** Critica (2) and Glimmer (13) software programs were used for coding region detection and gene identification. TMHMM 2.0 (23) was used to predict transmembrane helices in translated sequences. SignalP v2.0b2 (33) was used to predict the presence and location of N-terminal signal peptides. All automatic gene and function predictions were manually checked using BLAST programs (1), InterPro (31), and the Integrated Microbial Genomes system (28) and corrected if necessary. Protein functions were checked by using the Carbohydrate-Active enzymes (CAzy) (12; http://www.cazy.org) classification. A comparative analysis was performed based on assignment and classification of Clusters of Orthologous Groups (COG) of proteins (44) by using the Integrated Microbial Genomes system.

**Growth experiments and RNA isolation.** C. saccharolyticus was subcultured overnight three times using the substrate of interest in modified DSMZ 640 medium before it was inoculated into a pH-controlled (pH 7) 1-liter fermentor containing 4 g substrate per liter. Cells were grown at 70°C until mid-logarithmic phase (optical density at 660 nm, 0.3 to 0.4) and were harvested by centrifugation and rapid cooling to 4°C and then stored at −80°C. Total RNA was isolated using a modified Trizol (Invitrogen) protocol in combination with an RNAsesy kit (Qiagen). Quality was determined with an Experion bioanalyzer (Bio-Rad), and a modified Trizol (Invitrogen) protocol in combination with an RNAeasy kit (QIAGEN). Quality was determined with an Experion bioanalyzer (Bio-Rad), and the RNA was further purified with the RNaseasy Mini Kit (QIAGEN) according to the manufacturer's instructions. The cDNA samples were processed using a QIAquick purification kit (Qiagen, California), and they were eluted using phosphate buffer. The quantity and quality of the recovered cDNA samples were subsequently analyzed by using the ratio of absorbance at 260 to absorbance at 280 nm. Cyanine-3 and Cyanine-5 (Amersham, United Kingdom) dye labeling and sample hybridization were performed by following the instructions provided by TIGR (http://www.tigr.org/db/microarray/protocolsTIGR.shtml), with minor adjustments to accommodate long-oligonucleotide platforms. Samples were hybridized using a four-slide loop (see Fig. S1 in the supplemental material).

**Microarray hybridization.** The cDNA samples were processed using a QIAquick purification kit (Qiagen, California), and they were eluted using phosphate buffer. The quantity and quality of the recovered cDNA samples were subsequently analyzed by using the ratio of absorbance at 260 to absorbance at 280 nm. Cyanine-3 and Cyanine-5 (Amersham, United Kingdom) dye labeling and sample hybridization were performed by following the instructions provided by TIGR (http://www.tigr.org/db/microarray/protocolsTIGR.shtml), with minor adjustments to accommodate long-oligonucleotide platforms. Samples were hybridized using a four-slide loop (see Fig. S1 in the supplemental material).

**Data collection and analysis.** After incubation, slides were washed to remove nonspecifically bound material and scanned with a ScanArray Lite microarray scanner (Perkin Elmer, Massachusetts). Data acquisition and spot quantitation were performed with the ScanArray Express software. Once all the slides were quantitated, data from the loop were analyzed with JMP Genomics 3.0 (SAS, North Carolina), as described previously (36), using a mixed-effects analysis of variance model (49).

**Nucleotide sequence and gene expression data accession numbers.** The complete final genome assembly was released on 8 May 2007 and was deposited in the GenBank database under accession number CP000679 (http://genome.jgi-psf.org/finished_microbes/calsa/calsa.home.html). The gene expression data have been deposited in the Gene Expression Omnibus database (http://www.ncbi.nlm.nih.gov/projects/geo/) under accession number GSE11153.

**RESULTS**

**General features and comparative genomics of the genome of C. saccharolyticus.** The genome of C. saccharolyticus DSM 8903 (= ATCC 43494) consists of one 2,970,275-bp circular chromosome, which has a G+C content of 35.3% (Table 1). The locations of the 2,679 predicted coding sequences on the two strands reflect the correlation between the direction of transcription and replication and show that the chromosome has two unequal repliclores (Fig. 1). In addition, a GC skew analysis confirmed the huge size difference in the two replication arms, which might be attributed to a recent major inversion event. In addition to the protein-encoding genes, which were classified according to the COG system (Table 2), the chromosome harbors three rRNA operons and 46 tRNA genes with 41 different anticodons. These anticodons code for all 20 canonical amino acids. Like the chromosomes of many prokaryotes, the chromosome of C. saccharolyticus contains clustered regularly interspaced short palindromic repeats (CRISPR). CRISPR are DNA repeats that are separated by highly variable intervening sequences (spacers) and accompanied by CRISPR-associated (CAS) genes. The CRISPR and CAS protein products have been proposed to function as a defense mechanism against bacteriophages (4). With nine CRISPR loci and three different CAS genes, C. saccharolyticus is well equipped to fend off bacteriophages.

The complete genome sequence confirmed the phylogenetic position of C. saccharolyticus as a member of the class Clos-
and revealed that *Thermoanaerobacter tengcongensis* (whose genome sequence has also been completed) is the closest relative (see Table S1 in the supplemental material). According to recently updated small-subunit rRNA databases, *C. saccharolyticus* and *T. tengcongensis* are assigned to the order *Thermoanaerobacterales*. The diversity in this order is, however, very great, and thus the order has limited biological significance. Reclassification is expected in the near future (26). The genome of *C. saccharolyticus* was compared to the genomes of two thermophilic relatives, *Clostridium thermocellum* and *T. tengcongensis* (3), as well as to the genomes of distantly related hyperthermophiles, including the bacterium *Thermotoga maritima* (32) and the archaeon *Pyrococcus furiosus* (39). These microorganisms have small to moderate-size genomes and also produce hydrogen when they grow on a range of carbohydrates (see Table S2 in the supplemental material). The genomic distributions of proteins in COG categories are comparable for this group of species (Table 2).

### Table 2. Functional categories of predicted open reading frames in the genomes of hydrogen-producing organisms

<table>
<thead>
<tr>
<th>COG category</th>
<th>COG functional category</th>
<th>No. of proteins encoded in genomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Information storage and processing</td>
<td>Chromatin structure and dynamics</td>
<td>2 1 2 1 4</td>
</tr>
<tr>
<td>L</td>
<td>Replication, recombination, and repair</td>
<td>222 252 149 89 109</td>
</tr>
<tr>
<td>A</td>
<td>RNA processing and modification</td>
<td>0 0 0 0 2</td>
</tr>
<tr>
<td>K</td>
<td>Transcription</td>
<td>134 174 141 82 80</td>
</tr>
<tr>
<td>J</td>
<td>Translation, ribosomal structure, and biogenesis</td>
<td>147 165 150 135 166</td>
</tr>
<tr>
<td>Cellular processes and signaling</td>
<td>Cell cycle control, cell division, and chromosome partitioning</td>
<td>35 38 40 21 18</td>
</tr>
<tr>
<td>N</td>
<td>Cell motility</td>
<td>71 95 67 58 13</td>
</tr>
<tr>
<td>M</td>
<td>Cell wall, membrane, and envelope biogenesis</td>
<td>107 172 110 76 61</td>
</tr>
<tr>
<td>Z</td>
<td>Cytoskeleton</td>
<td>3 1 0 0 0</td>
</tr>
<tr>
<td>V</td>
<td>Defense mechanisms</td>
<td>48 40 42 27 29</td>
</tr>
<tr>
<td>U</td>
<td>Intracellular trafficking, secretion, and vesicular transport</td>
<td>42 60 46 39 20</td>
</tr>
<tr>
<td>O</td>
<td>Posttranslational modification, protein turnover, and chaperones</td>
<td>59 89 81 55 55</td>
</tr>
<tr>
<td>T</td>
<td>Signal transduction mechanisms</td>
<td>125 170 122 73 19</td>
</tr>
<tr>
<td>Metabolism</td>
<td>Amino acid transport and metabolism</td>
<td>166 166 206 181 158</td>
</tr>
<tr>
<td>G</td>
<td>Carbohydrate transport and metabolism</td>
<td>213 144 160 166 93</td>
</tr>
<tr>
<td>H</td>
<td>Coenzyme transport and metabolism</td>
<td>101 102 67 62 89</td>
</tr>
<tr>
<td>C</td>
<td>Energy production and conversion</td>
<td>111 115 130 119 128</td>
</tr>
<tr>
<td>P</td>
<td>Inorganic ion transport and metabolism</td>
<td>73 91 96 113 95</td>
</tr>
<tr>
<td>I</td>
<td>Lipid transport and metabolism</td>
<td>34 46 55 31 25</td>
</tr>
<tr>
<td>F</td>
<td>Nucleotide transport and metabolism</td>
<td>56 62 62 54 52</td>
</tr>
<tr>
<td>Q</td>
<td>Secondary metabolite biosynthesis, transport, and catabolism</td>
<td>14 18 27 15 11</td>
</tr>
<tr>
<td>Poorly characterized</td>
<td>Function unknown</td>
<td>177 187 173 137 189</td>
</tr>
<tr>
<td>R</td>
<td>General function prediction only</td>
<td>228 261 249 204 275</td>
</tr>
<tr>
<td>Not in COG</td>
<td></td>
<td>655 985 615 305 322</td>
</tr>
</tbody>
</table>

*a* Genes were classified with the Integrated Microbial Genomes system (28) using the COG functional classification for proteins (44).

*b* Number of protein-encoding genes in each category without pseudogenes.

*tridia* and revealed that *Thermoanaerobacter tengcongensis* (whose genome sequence has also been completed) is the closest relative (see Table S1 in the supplemental material). According to recently updated small-subunit rRNA databases, *C. saccharolyticus* and *T. tengcongensis* are assigned to the order *Thermoanaerobacterales*. The diversity in this order is, however, very great, and thus the order has limited biological significance. Reclassification is expected in the near future (26). The genome of *C. saccharolyticus* was compared to the genomes of two thermophilic relatives, *Clostridium thermocellum* and *T. tengcongensis* (3), as well as to the genomes of distantly related hyperthermophiles, including the bacterium *Thermotoga maritima* (32) and the archaeon *Pyrococcus furiosus* (39). These microorganisms have small to moderate-size genomes and also produce hydrogen when they grow on a range of carbohydrates (see Table S2 in the supplemental material). The genomic distributions of proteins in COG categories are comparable for this group of species (Table 2). However, one major difference is that both *C. thermocellum* and *C. saccharolyticus* have many more transposases and trans-
posase derivatives than the other species (99 and 93 such molecules, respectively). The genomes of *T. maritima*, *P. furiosus*, and *T. tengcongensis* encode 11, 17, and 43 transposases and transposase derivatives, respectively. Therefore, *C. saccharolyticus* and *C. thermocellum* have high numbers of proteins in the replication, recombination, and repair category (category L) (222 and 252 proteins, respectively). Furthermore, the *C. saccharolyticus* genome contains the largest number of carbohydrate transport and metabolism genes in this group. In fact, the *C. saccharolyticus* genome contains at least 177 ABC transporter genes, which is more than the 165 ABC transporter genes identified in *T. maritima* (11, 32). The *C. saccharolyticus* genome contains a reverse gyrase gene (*Csac_1580*), the product of which induces positive supercoiling of DNA (22). Reverse gyrase is considered a molecular marker of hyperthermophilicity and therefore distinguishes *C. saccharolyticus* from *C. thermocellum*, which lacks reverse gyrase. Despite the fact that *C. saccharolyticus* was described as a nonmotile organism, a set of flagellum structure, biogenesis, and chemotaxis genes was detected; it is not clear whether these genes are functional, since one of them is interrupted by a stop codon (pseudogene *Csac_1277*). *C. saccharolyticus* has a nitrogen fixation cluster (*Csac_2461* to *Csac_2466*) and many sporulation genes; neither of these properties has been described previously for this bacterium.

**Central carbon metabolism.** *C. saccharolyticus* is able to metabolize a wide variety of carbohydrates, including the monosaccharides D-glucose, D-xylose, D-fructose, D-galactose, D-ribose, L-rhamnose, and L-fucose, as well as α- and β-linked di- and polysaccharides, including maltose, starch, pullulan, sucrose, trehalose, amorphous and microcrystalline cellulose, xylan, locust bean gum, and pectin (37). Once they are hydrolyzed, sugars are channeled to the central catabolic pathways (Fig. 2). Analysis of the genome sequence re-
revealed components of a complete Embden-Meyerhof (EM) pathway, including a ROK family glucokinase (Csac_0778), 6-phosphofructokinase, (Csac_2366 and Csac_1830), a bifunctional phosphoglucomutase/phosphomannose isomerase (Csac_1187), fructose-1,6-bisphosphate aldolase (Csac_1189), and pyruvate kinase (Csac_1831), as well as pyruvate-phosphate dikinase (PPDK) (Csac_1955). Also, a gapA operon was found, consisting of genes encoding glyceraldehyde-3-phosphate dehydrogenase (Csac_1953), the fusion protein phosphoglycerate kinase/triose-phosphate isomerase (Csac_1952), phosphoglycerate mutase (Csac_1951), and enolase (Csac_1950) (Fig. 2). However, the oxidative branch of the pentose phosphate pathway (PPP) and the Entner-Doudoroff pathway were not detected, which is consistent with previous findings obtained using 13C nuclear magnetic resonance (14). The absence of the oxidative branch of the PPP, however, raises questions about how NADPH is produced for biosynthesis. The only other obvious NADPH-producing reaction is the isocitrate dehydrogenase (Csac_0751) reaction. However, based on sequence homology, the isocitrate dehydrogenase is likely to produce NADH instead of NADPH. Also, no obvious homolog of an NADPH-producing glyceraldehyde-3-phosphate dehydrogenase could be identified, as has been reported previously for Streptococcus species and some Clostridia (8). NADPH can also be synthesized from NADH by a transhydrogenase, which is either membrane bound or soluble. In the genome there are no genes encoding orthologs of subunits of the membrane-bound type, but several homologous genes coding for soluble flavin adenine dinucleotide-dependent pyridine nucleotide-disulfide oxidoreductases are present (Csac_0759, Csac_2199, and Csac_0402). The exact physiological roles of these enzymes have not been determined. Furthermore, no ferredoxin: NADPH reductase homolog is present, although such activity has been found in some Thermotoga maritima spp. (19).

Xylose, a major constituent of hemicellulose, is funneled by a putative xylose isomerase (Csac_1154) and a xylulokinase (Csac_0798) into the nonoxidative branch of the PPP. The nonoxidative branch of the PPP uses ribulose phosphate 3-epimerase (Csac_2074), ribose-5-phosphate isomerase (Csac_1200), the N-terminal (Csac_1351) and C-terminal (Csac_1352) transketolases, and transaldolase (Csac_2036) to produce the EM intermediates fructose-6-phosphate and glyceraldehyde-3-phosphate. Galactose also enters the EM via the Leloir pathway, which includes galactokinase (Csac_1511), galactose-1-phosphate uridylyltransferase (Csac_1510), UDP-glucose 4-epimerase (Csac_1512), and phosphoglucomutase (Csac_2295). Strikingly, genes encoding the established types of fructose bisphosphatase (classes I to IV [41]) are not evident in the C. saccharolyticus genome. Since fructose bisphosphatase is an essential enzyme in gluconeogenesis, C. saccharolyticus presumably uses a novel phosphatase. Moreover, a gene encoding the gluconegenic phosphoenolpyruvate (PEP) synthetase is also not present, although conversion of pyruvate to PEP could occur via the reversible PPDK (Csac_1955) or via oxaloacetate.

Pyruvate, the end product of the EM pathway, is decarboxylated to acetyl coenzyme A (acetyl-CoA) by pyruvate/ferredoxin oxidoreductase (POR). C. saccharolyticus contains three 2-oxoacid:ferredoxin oxidoreductase enzyme complexes (Csac_2248 and Csac_2249, Csac_1458 to Csac_1461, and Csac_1548 to Csac_1551). According to transcriptional response data (see below), the true POR is probably encoded by the Csac_1458 to Csac_1461 genes. Acetyl-CoA is used to generate acetate and ATP (Csac_2040 and Csac_2041), or it enters the tricarboxylic acid (TCA) cycle for biosynthetic purposes. The TCA cycle in C. saccharolyticus is incomplete; it has an oxidative branch to succinyl-CoA catalyzed by a citrate (Re) synthase (Csac_0746), aconitate hydratase (Csac_0750), isocitrate dehydrogenase (Csac_0751), and the 2-oxoglutarate:ferredoxin oxidoreductase complex (Csac_1548 to Csac_1551). In the reductive direction, only orthologs of the subunits of fumarate hydratase were detected with a high level of confidence (Csac_2759/ Csac_0738). Malate dehydrogenase (oxaloacetate decarboxylating) (Csac_2059) may be used to generate malate directly from pyruvate instead of from oxaloacetate. Fumarate reductase, however, could not be identified, which is in agreement with the absence of this enzyme in related clostridia. In addition to the activity of malate dehydrogenase, TCA metabolites could be replenished by a putative sodium pump oxaloacetate decarboxylase enzyme complex (Csac_2482 to Csac_2485).

**Polysaccharide-degrading enzymes.** The capacity of C. saccharolyticus to hydrolyze a broad range of polysaccharides prior to fermentation differentiates this bacterium from many thermophilic anaerobes. Indeed, the genome of C. saccharolyticus encodes a wide range of enzymes active with carbohydrates (see Table S3 in the supplemental material). The genes encoding these carbohydrate-utilizing enzymes are often clustered on the chromosome, and the enzymes can be assigned to substrate-specific catabolic pathways for cellulose, hemicellulose, and, to a lesser extent, starch and pectin. α,1,4-Glucan polymers, for instance, can be transported into the cell using the maltodextrin ABC transport system proteins (Csac_0427 to Csac_0428/Csac_0431). An intracellular α-amylase (Csac_0426) and a 1,4-α-glucan phosphorylase (Csac_0429) further degrade the intracellular maltodextrins, releasing glucose-1-phosphate. Remarkably, a transcriptional regulator of the LacI family (Csac_0430) is also in this maltodextrin cluster and therefore is a good candidate for controlling expression of this maltodextrin-degrading pathway at the transcriptional level. In addition, a GCAACGTTTGC consensus sequence was found in sequences upstream of genes in this transport cluster and genes encoding several starch-degrading enzymes, such as an α-amylase precursor (Csac_0408), an oligo-1,6-glucosidase (Csac_2428), a pullulanase (Csac_0689), a 4-α-glucanotransferase (Csac_0203), and a putative glucon 1,4-α-glucosidase (Csac_0130). The consensus sequence resembles the binding site (CGCAACGTTTGC) of the maltose/maltodextrin transcriptional repressor MalR from the gram-positive organism Streptococcus pneumoniae (34). Besides this putative starch-degrading regulon, C. saccharolyticus has genes encoding a glycogen metabolic cluster (Csac_0780 to Csac_0784), a malate ABC transport system (Csac_2491 to Csac_2493), and a second pullulanase (Csac_0671). Thus, C. saccharolyticus is well equipped for starch utilization.

An important feature of C. saccharolyticus is its ability to produce H2 not only from α-linked polymers but also from complex β-linked glycans, such as cellulose, hemicellulose, laminarin, and galactomannan. Growth on cellulolic substrates is rare among (hyper)thermophilic microorganisms. C. saccharolyticus does not metabolize cellulose by means of a cellulase (45). For example, genes encoding the typical molecular
components of a cellulosome (i.e., dockerin domains and scaffolding proteins) were not identified in the genome. Nevertheless, a gene cluster (Csac_1076 to Csac_1081) containing genes encoding cellulase precursors is present. The cellulases are potentially capable of degrading cellulose, a plant polysaccharide (6) (see Table S3 in the supplemental material). Moreover, the product of another gene cluster (Csac_1089 to Csac_1091) and an extracellular cellulase (Csac_0678) may assist in completely hydrolyzing cellulose to glucose. A remarkable aspect of the \( \beta \)-glycanases in Caldicellulosiruptor species is their bifunctional domain architecture consisting of central cellulose binding domains which are bordered by distinct catalytic domains (6). The \textit{C. saccharolyticus} genome sequence confirmed the presence of genes encoding these multidomain hydrolases (Csac_1076 to Csac_1079 and Csac_2411), which comprise cellulase, mannanase, xylanase and arabinofuranosidase domains (6, 27, 30, 45). No genes encoding additional bifunctional \( \beta \)-glycanases were found in the genome. The bifunctional Csac_2411 gene is part of another large gene cluster (Csac_2404 to Csac_2411) coding for enzymes involved in hydrolysis of the plant polysaccharide xylan (hemicellulose). The genes encoding these mostly extracellular enzymes might be coexpressed with genes in a smaller putative xylan-utilizing cluster (Csac_0203 to Csac_0205). The genes in the latter cluster were not significantly upregulated during growth on xylose, in contrast to the genes in the former cluster. Furthermore, putative genes that encode enzymes that degrade galactomannan (Csac_0663 and Csac_0664), galactoarabinan (Csac_1560 to Csac_1562), and laminarin (Csac_2548) were identified.

The plant cell wall component pectin consists of an \( \alpha \)-1,4-linked \( \beta \)-galacturonic acid backbone, sometimes with \( \beta \)-rhamnose interspersed, and side chains consisting of monosaccharides, such as \( \beta \)-galactose, \( \alpha \)-xylose, and \( \beta \)-arabinose (38). Degradation of the main pectin component, \( \beta \)-galacturonate, requires a galacturonate isomerase, a tagaturonate reductase, and an altronate dehydratase to form 2-keto-3-deoxygluconate (KDG). Galacturonate isomerization may be catalyzed by glucuronate isomerase (Csac_1949). However, genes encoding tagaturonate reductase and altronate dehydratase were not detected in the genome of \textit{C. saccharolyticus}. Apparently, novel enzymes or a novel pathway is responsible for the degradation of galacturonate. In contrast, a gene cluster encoding enzymes involved in the conversion of glucuronic acid to KDG (Csac_2686 to Csac_2689) was identified, and it includes genes encoding fructuronate reductase, mannonate dehydratase, a putative \( \beta \)-galactosidase/\( \beta \)-glucuronidase, and an \( \alpha \)-glucuronidase. Glucuronic acid is a common substituent of xylan. Enzymes involved in the conversion of KDG to pyruvate and glyceraldehyde-3-phosphate, including KDG kinase (Csac_0355 or Csac_2720) and KDG-6-phosphate aldolase (Csac_0354), are present as well. The genes encoding enzymes involved in the last two steps are clustered with genes (Csac_0356 and Csac_0357, as well as Csac_2718 and Csac_2719) that encode enzymes involved in the conversion of 5-keto-4-deoxyxuronate, an unsaturated cleavage product of pectate, to KDG. The genes encoding enzymes that are able to hydrolyze the pectate backbone and the side chains (e.g., unsaturated rhamnogalacturonyl hydrolase [Csac_0360], galacturan 1,4-\( \alpha \)-galacturonidase [Csac_0361], \( \beta \)-galactosidase [Csac_0362], and a glycoside hydrolase with unknown substrate specificity [Csac_0363]) are close to the genes encoding the KDG metabolic enzymes as well. However, neither a pectate lyase gene nor a methylesterase gene could be definitively identified in the genome, although Csac_2721 and Csac_2728 might be candidates for a pectate lyase gene based on distant homology to known lyase genes.

\textit{C. saccharolyticus} is also able to grow on \( \beta \)-rhamnose and on \( \beta \)-fucose, producing 1,2-propanediol as the end product (unpublished data). A putative rhamnose catabolic pathway was identified, in which generation of dihydroxycetone phosphate and 1,2-propanediol is catalyzed by an \( \alpha \)-rhamnose isomerase (Csac_0876), a putative \( \alpha \)-rhamnulose-1-phosphate aldolase (Csac_0865), and a putative lactaldehyde reductase (Csac_0407). Fucose can be processed by a similar pathway, using the lactaldehyde reductase mentioned above and as-yet-identified versions of \( \alpha \)-fuculokinase, a bifunctional \( \beta \)-fucose isomerase/\( \beta \)-arabinose isomerase (Csac_1339), and fuculose-1-phosphate aldolase (Csac_0425).

\textbf{Fermentation products.} Reducing equivalents are produced at the level of NAD and ferredoxin (Csac_0737). Since \textit{C. saccharolyticus} can produce almost 4 mol of \( \text{H}_2 \) per mol of glucose (14), both NADH and reduced ferredoxin should ultimately be able to transfer their reducing equivalents to protons to form hydrogen. In the genome, two hydrogenase gene clusters were identified, which are very similar to the two related clusters in \textit{T. tengcongensis} (42). The first cluster includes genes encoding subunits of an Ni-Fe hydrogenase (EchA to EchF) (Csac_1534 to Csac_1539) and various genes required for maturation of the hydrogenase complex (HypA to HypF) (Csac_1540 to Csac_1545). In \textit{T. tengcongensis}, the Ni-Fe hydrogenase is ferredoxin dependent and membrane bound and may act as a proton pump to generate a proton motive force. The second cluster (Csac_1860 to Csac_1864) codes for an Fe-only hydrogenase (HydA to HydD), which is NAD dependent and located in the cytoplasm, similar to the situation in \textit{T. tengcongensis} (42). Hydrogenases that form \( \text{H}_2 \) directly from NADH are unusual and make an NAD:ferredoxin oxidoreductase (Nfo) redundant. Nfos (also known as Rnfs) are membrane-bound multisubunit complexes that use or create an \( \text{Na}^+ \) gradient coupled to the transfer of reducing equivalents between NADH and ferredoxin (7). An Nfo gene cluster has been identified in the genomes of \textit{C. thermocellum}, \textit{T. maritima}, and \textit{Thermotoga ethanolicus}, but not in the genomes of \textit{T. tengcongensis} and \textit{C. saccharolyticus}. The absence of an Nfo in \textit{C. saccharolyticus} also implies either that in the presence of elevated levels of \( \text{H}_2 \) reduced ferredoxin may not be used to produce NADH or that a novel type of enzyme (complex) performs this reaction. Altogether, the information available suggests that \textit{C. saccharolyticus} is able to produce hydrogen from ferredoxin but can also produce hydrogen directly from NADH. Production of hydrogen seems to be preferable, because under these conditions all pyruvate is converted to acetate (and \( \text{CO}_2 \)), which is coupled to the synthesis of ATP.

When the hydrogen partial pressure is too high, formation of hydrogen from NADH is no longer thermodynamically favorable. In such a case, NADH is oxidized through the formation of lactate or ethanol. A gene for a lactate dehydrogenase was identified (Csac_1027), but genes for acetaldehyde dehydrogenase and alcohol dehydrogenase were not obvious. In \textit{T. teng-
congesis and \( T. \) ethanolicus, ethanol formation is NADPH dependent and catalyzed by a bifunctional alcohol dehydrogenase/acetyl-CoA thioesterase; this enzyme also has a homolog in \( C. \) saccharolyticus (Csac_0395).

A third small hydrogenase-like cluster was detected in the \( C. \) saccharolyticus genome and was composed of four genes encoding two NADH-binding proteins (Csac_0619 and Csac_0620), a molybdopterin oxidoreductase containing NAD and 4Fe-4S binding regions (Csac_0621), and an iron-containing alcohol dehydrogenase (Csac_0622). The function of this cluster is not known.

**Transport systems.** As mentioned above, a number of genes encoding ABC transporters have been found in the \( C. \) saccharolyticus genome, including genes encoding the previously described carbohydrate-specific maltodextrin ABC transport system (Csac_0427, Csac_0428, and Csac_0431) and the maltose ABC transport system (Csac_2491 to Csac_2493). As observed for certain \( T. \) maritima maltose transporters, both sets of transport proteins lack ATP-binding subunits. In many bacteria, the intracellular ATPase used in the system is not encoded in the same operon. The genes encoding both of these sets of ABC transporters are located downstream from the genes encoding a two-component system consisting of a sensor histidine protein kinase and a response regulator. In \( C. \) saccharolyticus ~50% of the genes encoding ABC carbohydrate transport systems are located near the genes encoding the two-component system on the chromosome.

Comparative analysis of \( C. \) saccharolyticus sugar binding proteins (SBPs) revealed that about two-thirds of them belong to the COG1653 category. This category includes the CUT1 subfamily members TM0432, TM0595, and TM1855, which transport a variety of \( \alpha \)-and \( \beta \)-oligosaccharides, such as maltose. In genomes, the genes encoding more than one-half of COG1653 members are close to genes encoding glycoside hydrolases, supporting the identification of these proteins as ABC transporters involved in carbohydrate utilization. Putative SBPs encoded by Csac_0242, Csac_0391, Csac_2326, and Csac_2507 belong to COG1879. Csac_2506 and Csac_2510 are associated with the xylose transport-specific COG4213 category. As in \( T. \) maritima, a few putative SBPs (Csac_0261 and Csac_4166) are annotated as peptide transporters, although their actual functions are unknown. Components of phosphotransferase systems have been identified in \( C. \) saccharolyticus (although only one set of carbohydrate-specific EIII has been identified), as have a few putative members of the major facilitator superfamily (Csac_0685, Csac_0786, Csac_1100, Csac_2326, and Csac_2298). However, it is likely that carbohydrate utilization proceeds mainly through ABC transporters.

**Transcriptional regulation.** The ability of \( C. \) saccharolyticus to utilize many different carbohydrates suggests that there is tight regulation in the pathways. The genes encoding many enzymes involved in carbohydrate utilization pathways appear to be regulated at the transcriptional level. Apart from the RNA polymerase core enzyme subunits (Csac_2259, Csac_2085, Csac_0951, and Csac_0952), this gram-positive species has 12 different \( \sigma \) factors for construction of the RNA polymerase holoenzyme. In addition, multiple copies of genes encoding many sugar transcriptional regulators are present in the genome; for example, there are nine proteins belonging to the LacI family, six proteins belonging to the DeoR family, and eight proteins belonging to the GntR family, as well as 19 receiver proteins involved in a two-component system with a helix-turn-helix AraC domain. The genes encoding the latter proteins are always clustered with genes encoding sugar transporters and sugar hydrolytic enzymes.

Carbon catabolite repression (CCR) by glucose was not observed in \( C. \) saccharolyticus (Fig. 3). Nevertheless, genes encoding some indicators of the presence of carbon control protein A (CcpA)-dependent CCR of gram-positive bacteria are present in the genome, including (i) a gene encoding a histidine-containing phosphocarrier (HPr) (Csac_2438) that is close to the gene encoding the only PEP-dependent phosphotransferase system, which is fructose specific; (ii) a gene encoding an HPr(Ser) kinase (Csac_1186); and (iii) genes encoding a catabolite repression HPr (CrH) (Csac_1163) and nine members of the CcpA-containing LacI family. Binding sites for a putative CcpA, the catabolite-responsive element (cre), could not be identified. The \( B. \) subtilis consensus sequence WW TGNAARCGNNWWCAWW (29), for instance, was detected only twice, once in the upstream region of the gene encoding the \( \alpha \)-amylase precursor mentioned above (Csac_0408), where it overlaps the putative MalR binding site, and once in the middle of the gene encoding fumarate hydratase subunit \( \alpha \) (Csac_2759). Nevertheless, CCR probably occurs in \( C. \) saccharolyticus, although the metabolite that induces this repression is not known.

Besides global regulation through CCR, many local transcriptional regulators control the expression of carbohydrate metabolic pathways. Several orthologous transcriptional regulators were identified in \( C. \) saccharolyticus. The central glycolytic gene regulator (CggR) (Csac_1954), for instance, represses transcription of the \( gapP \) operon (25), while FruR (Csac_2442) controls the fructose operon (5). Based on the fact that many transcriptional regulator genes are close to their target operons, putative functions could be assigned to an \( \alpha \)-linked glucan transcriptional regulator (Csac_0430), a regul-
lator of the oxidative branch of the TCA cycle to oxoglutarate (Csac_0752), a repressor of L-arabinose metabolism (Csac_0722), and a putative response regulator receiver protein involved in glucuronate degradation (Csac_2690).

Transcriptome analysis of C. saccharolyticus. One of the features of C. saccharolyticus that is beneficial for hydrogen production is its ability to degrade cellulosic substrates as well as hemicellulose. Moreover, mixtures of glucose and xylose can be fermented simultaneously (Fig. 3), suggesting that classical CCR by glucose does not occur. To elucidate the central carbon metabolic pathways and their regulation, a transcriptome analysis was performed after growth on glucose, xylose, and a 1:1 mixture of these two substrates. L-Rhamnose, which was likely to be metabolized by another pathway, was used as a reference substrate. The transcriptional data clearly show that glucose, xylose, and the glucose-xylose mixture all triggered upregulation of genes in the EM pathway, compared to the results for rhamnose (Fig. 4; see Table S4 in the supplemental material). In particular, fructose bisphosphate aldolase, glyceraldehyde-3-phosphate dehydrogenase, PPDK, and POR were significantly stimulated. The ultimate acetate-forming acetate kinase was also highly upregulated. A catabolic role for PPDK is intriguing, since this enzyme normally is associated with gluconeogenesis (as it is in propionic acid bacteria and plants) and PEP is usually converted by pyruvate kinase. However, homologs of PPDK are also present in related clostridia and Thermoanaerobacter species.

It is worth noting that growth on glucose, growth on xylose, and growth on both sugars all triggered transcription of the genes encoding a xylose-specific ABC transport system (Csac_2504 to Csac_2506) (Fig. 4), suggesting that glucose and xylose are transported by the same uptake system. Moreover, none of the identified putative CCR genes (see below) were differentially transcribed, confirming that catabolite repression by glucose was not a factor.

The transcriptional response to growth on monosaccharides enabled identification of genes and groups of adjacent genes (gene clusters) that were specifically upregulated in response to either glucose or xylose. On glucose, several genes coding for α-glucan hydrolases responded. The most striking observation, however, was the upregulation of an entire gene cluster (Csac_1991 to Csac_2000) involved in purine synthesis, which was not observed with xylose. Upregulation of purine biosynthesis genes was also detected in the transcriptome of Esche-
richia coli growing on glucose compared to the results obtained with xylose (16). On xylose, several gene clusters required for xylan or xylose conversion were upregulated (Csac0692 to Csac0696, Csac0240 to Csac0242, and Csac2416 to Csac2419). These clusters encode ABC transport systems, transcriptional regulators, and endoxylanases. In addition, genes specifically required for growth on rhamnose were highly upregulated during growth on rhamnose, indicating the pathway for utilization of this sugar.

**DISCUSSION**

*C. saccharolyticus* has been shown to be an excellent candidate for biohydrogen production (14, 20, 48). In contrast to mesophilic fermentative anaerobes, it produces almost no reduced end products, such as lactate or ethanol, and the amount of hydrogen produced approaches the “Thauer limit” of 4 mol of H₂/mmol of glucose (17, 46). Moreover, *C. saccharolyticus* hydrolyzes various biomass-derived polymers, such as cellulose, hemicellulose, starch, and pectin and ferments corresponding sugar monomers, including glucose and xylose. The complete genome sequence of *C. saccharolyticus* provides new insights into the exceptional capacity of this bacterium to degrade a variety of plant polysaccharides and further reveals its high level of plasticity with many transposases, CRISPRs, and two uneven replication arms. A large number of sugar hydrodolases and transferases were identified, and the number was far greater than the number in the hyperthermophile *T. maritima* (9). Metabolic pathways for the degradation of residual components of cellulose, hemicellulose, starch, and pectin could be assigned. Reducing equivalents are produced as NADH or reduced ferredoxin, which are apparently used directly to produce hydrogen by a soluble NADH-dependent Fe-only hydrogenase and a membrane-bound ferredoxin-dependent Ni-Fe hydrogenase. The ability to produce hydrogen directly from NADH has not been observed for mesophilic anaerobes and is being developed, which initially will target the dehydrogenases involved in lactate and ethanol formation. Alternatively, genes could be introduced to form an oxidative branch of the PPP in order to obtain H₂ yields greater than 4 mol per mol of glucose (50). In any case, the *C. saccharolyticus* genome provides new insights into the metabolic features of a versatile biohydrogen producer, which could inspire efforts to optimize microbial bioenergy systems.

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