The thermophilic biomass-degrading bacterium *Caldicellulosiruptor bescii* utilizes two enzymes to oxidize glyceraldehyde 3-phosphate during glycolysis

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*Caldicellulosiruptor bescii* is an extremely thermophilic, cellulolytic bacterium with a growth optimum at 78 °C and is the most thermophilic cellulose degrader known. It is an attractive target for biotechnological applications, but metabolic engineering will require an in-depth understanding of its primary pathways. A previous analysis of its genome uncovered evidence that *C. bescii* may have a completely uncharacterized aspect to its redox metabolism, involving a tungsten-containing oxidoreductase of unknown function. Herein, we purified and characterized this new member of the aldehyde ferredoxin oxidoreductase family of tungsten enzymes. We show that it is a heterodimeric glyceraldehyde-3-phosphate (GAP) ferredoxin oxidoreductase (GOR) present not only in all known *Caldicellulosiruptor* species, but also in 44 mostly anaerobic bacterial genera. GOR is phylogenetically distinct from the monomeric GAP-oxidizing enzyme found previously in several Archaea. We found that its large subunit (GOR-L) contains a single tungstopterin site and one iron-sulfur [4Fe-4S] cluster, that the small subunit (GOR-S) contains four [4Fe-4S] clusters, and that GOR uses ferredoxin as an electron acceptor. Deletion of either subunit resulted in a distinct growth phenotype on both C₅ and C₆ sugars, with an increased lag phase, but higher cell densities. Using metabolomics and kinetic analyses, we show that GOR functions in parallel with the conventional GAP dehydrogenase, providing an alternative ferredoxin-dependent glycolytic pathway. These two pathways likely facilitate the recycling of reduced redox carriers (NADH and ferredoxin) in response to environmental H₂ concentrations. This metabolic flexibility has important implications for the future engineering of this and related species.

*Caldicellulosiruptor bescii* is an extremely thermophilic bacterium with an optimal growth temperature of 78 °C, making it the most thermophilic cellulose degrader identified to date (1–3). This organism is an attractive target for use in biotechnological applications due to its ability to ferment high concentrations of untreated plant biomass, thereby potentially mitigating the costly thermochemical pretreatment steps that are typically required for lignocellulose conversion (4–6). *C. bescii* and other species of the *Caldicellulosiruptor* genus do not contain the well-characterized cellulosomes described for other cellulolytic bacteria, such as *Clostridium thermocellum*, but rather secrete a suite of multidomain noncellulosomal glycosyl hydrolases that work synergistically to convert plant biomass into fermentable sugars (5, 7). *C. bescii* ferments C₅ and C₆ sugars derived from cellulose and hemicellulose primarily to hydrogen gas, lactate, acetate, and carbon dioxide (8, 9). The development of a genetic system for this organism has created the potential to metabolically-engineer it for consolidated biomass processing to produce liquid fuels and other bio-based products from renewable biomass. Engineered strains of *C. bescii* that can produce moderate amounts of ethanol directly from the conversion of plant biomass have already been achieved (10, 11). The recent developments of a high temperature kanamycin-resistance gene and of an inducible promoter have further increased the potential use of this organism as a platform for biotechnological applications (12–15). The further development of efficient metabolic engineering strategies will require an in-depth understanding of its primary metabolism.

We recently reported that *C. bescii* has a completely uncharacterized aspect to its primary redox metabolism involving the metal tungsten (16). This was very surprising; tungsten is an element that is rarely utilized in biological systems, as virtually all forms of life utilize the chemically analogous element molybdenum (17). Molybdoenzymes are ubiquitous, serving roles in carbon, nitrogen, and sulfur metabolism (18). In contrast, the number of microorganisms known to require tungsten is relatively few (17). Tungsten is chemically very similar to molybdenum with comparable coordination chemistries and physical properties. They are both coordinated to enzymes via pyranopterin cofactors and they both stabilize three redox states (IV, V, and VI) in the biological range, although for tungsten these are...
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at much lower reduction potentials than those of molybdenum (19). Despite this, there is a large bias toward molybdoenzymes in all three domains of life, whereas tungstoenzymes appear to be more abundant in thermophilic organisms (20).

Tungsten metabolism is best characterized in hyperthermophilic Archaea, represented by Pyrococcus furiosus, which grows optimally at 100 °C. It contains five members of the aldehyde oxidoreductase (AOR)3 family of tungstoenzymes, which are phylogenetically unrelated to any of the three families of molybdoenzymes (21). All five have been characterized and they are referred to as aldehyde ferredoxin oxidoreductase (AOR) (22), glyceraldehyde-3-phosphate ferredoxin oxidoreductase (GAPOR) (23), and formaldehyde ferredoxin oxidoreductase (FOR) (24), together with two tungstoenzymes of unknown function, WOR4 (25) and WOR5 (26). All of these enzymes oxidize aldehydes of various types. The prototypical AOR has a broad substrate specificity and is thought to be involved in peptide catabolism wherein it oxidizes amino acid-derived aldehydes (17). However, of all of these archaeal tungstoenzymes only the function of GAPOR is definitively understood (23). GAPOR replaces the conventional NAD-dependent glyceraldehyde-3-phosphate (GAP) dehydrogenase (GAPDH) and is part of a modified glycolytic pathway in Pyrococcus furiosus and several other archaeal species (23, 27, 28). In addition to using ferredoxin rather than NAD as an electron acceptor, the GAPOR reaction oxidizes GAP directly to 3-phosphoglycerate rather than to the 1,3-bisphosphoglycerate generated by GAPDH (23). Hence, this unusual GAPOR-containing glycolytic pathway found in these Archaea does not contain phosphoglycerate kinase and consequently GAP oxidation does not result in net ATP synthesis by substrate level phosphorylation.

The notion of C. bescii utilizing tungsten came from the identification of a 12-gene cluster of highly expressed genes (Athe_0820-Athe_0831) (Fig. S1) that are completely conserved across all 14 members of the Caldicipellulosiruptor genus that have been characterized to date (29). These genes encode proteins involved in pyranopterin synthesis and transport, and a putative tungsopterin-containing enzyme (Athe_0821) belonging to the AOR family of tungstoenzymes that we termed XOR to signify its uncharacterized status. To demonstrate that C. bescii could utilize tungsten, the known tungstoenzyme AOR of P. furiosus was heterologously produced in an active form (16). Additionally, we showed that during growth on simple and complex C6 and C5 sugars, Athe_0821 is constitutively expressed and its protein product represents ∼2% of the cytoplasmic protein in C. bescii. XOR is phylogenetically distinct from the characterized AOR enzymes and was proposed to represent a new member of this family of tungstoenzymes (16). XOR is also unusual in that the gene encoding Athe_0821 is adjacent to another highly-expressed gene (Athe_0820) encoding a polyferredoxin-like protein predicted to contain four [4Fe-4S] clusters. It was not known if this functioned as an electron carrier for XOR or was an additional subunit.

It was therefore of importance to elucidate the function of this tungstoenzyme in the Caldicipellulosiruptor genus, both in terms of understanding its primary metabolism and in utilizing this information to engineer these organisms for biotechnological applications. In contrast to the archaebon P. furiosus, C. bescii does not grow on peptides and its genome encodes the standard Embden-Meyerhof (EM) pathway, including a classical phosphorylating GAPDH (Athe_1406). Moreover, cell-free extracts of C. bescii did not catalyze the oxidation of any of the aldehydes, including GAP, used by the AOR-type enzymes of P. furiosus (16). Herein we show that XOR is indeed a new member of the AOR family of tungstoenzymes and is a novel bacterial-type of glyceraldehyde-3-phosphate ferredoxin oxidoreductase. We will refer to it as GOR to differentiate it from the phylogenetically-distinct GAPOR found primarily in Archaea. Affinity-tagged GOR was purified and kinetically characterized. Utilizing a deletion mutant strain lacking GOR, metabolomics was used to show that GOR provides C. bescii with an alternative glycolytic pathway, generating reduced ferredoxin rather than NADH. This is the first report of a functional “GAPOR” in a bacterium.

Results

Purification of affinity-tagged GOR

The genes encoding XOR (Athe_0821) and the putative polyferredoxin (PFD, Athe_0820) were modified by the inclusion of a polyhistidine tag at the N terminus of XOR and the expression of both genes was placed under control of the high level promoter of the S-layer protein (slp) to give the overexpression strain (OE-XOR/PFD, Fig. S1). The slp gene is expressed at a level that is ∼10-fold higher than that of Athe_0821 and of Athe_0820 in WT C. bescii (16). XOR was purified from the cytoplasmic extract of the OE-XOR/PFD strain by ion exchange, affinity, and size exclusion chromatography and its properties are described below. The first priority was to determine a catalytic activity for the purified enzyme. XOR was assayed for its ability to oxidize a variety of aldehydes and reduce the artificial electron acceptor benzyl viologen (BV). These included the following compounds: 2-ethylhexanal, 2-methylbutyraldehyde, 2-methylpentanal, 2-methoxybenzaldehyde, 2-naphthaldehyde, 2-phenylpropionaldehyde, 3-phenylbutyraldehyde, 3,4-dihydroxybenzaldehyde, acetaldelyde, acetoaminobutyaldehyde, cinnamaldehyde, coniferaldehyde, dihydroxyacetone, galactose, glucose, glucose 6-phosphate, glyceraldehyde, glyceraldehyde, glyciclic acid, glyxol, glyoxylic acid, hexanal, isobutyaldehyde, mannose, methyl glyoxal, phenylpropionaldehyde, ribose, sinapaldehyde, syringaldehyde, vanillin, and xylose. However, none were utilized as a substrate by XOR. We had previously shown that a cytoplasmic extract of WT C. bescii was unable to oxidize several other aldehydes that are known substrates of the AOR family members in the Archaea (16). These included formaldehyde, propionalde-
hyde, crotonaldehyde, glutaraldehyde, isovaleraldehyde, benzaldehyde, and glyceraldehyde 3-phosphate (GAP). Similarly, purified XOR was unable to use these substrates, with the notable exception of GAP. Surprisingly, GAP (1 mM) was readily oxidized by purified XOR with a specific activity (at 70 °C) of 48.3 units/mg and the activity increased with increasing concentrations of XOR in the assay (Table S1). However, contrary to expectation, only a very low level of GAP oxidation activity was detected in the cytoplasmic extract of the OE-XOR/PFD strain (0.01 units/mg). As previously reported, the same was also observed with extracts from the WT strain (<0.01 unit/mg). Because the purified XOR enzyme clearly utilizes GAP as a substrate, we will now refer to XOR as a GAP oxidoreductase or GOR.

The lack of significant benzyl viologen (BV)-linked GAP-oxidation activity in the cytoplasmic extract of the WT strain appeared to be due to the presence of an inhibitor of GOR activity. For example, as shown in Table 1, the total GAP-oxidizing units increased ~4-fold following the affinity purification of GOR. This was accompanied by a 76-fold purification yielding 2.1 mg of purified protein from 50 g (wet weight) of cells. A similar response to the partial purification from the cytoplasmic fraction was observed with GAPDH, where a greater than 100-fold increase in the NAD-dependent oxidation of GAP was observed after an anion exchange chromatography step (Fig. S2). The addition of the cytoplasmic extract to purified GOR resulted in a ~90% reduction in its GAP-oxidizing activity (Fig. 1), indicating that the cytosol contains an inhibitory factor. To investigate the nature of this inhibitory factor, the cytoplasmic extract from the OE-XOR/PFD strain was fractionated using a 30-kDa filter (and the protein fraction <30 kDa was washed with buffer). The washed extract (nominally >30 kDa) and the resulting filtrate (<30 kDa) were each tested for their capacity to inhibit the GAP-oxidizing activity of purified GOR (Fig. 1). The inhibitor appeared only in the high molecular mass fraction (>30 kDa) as the low molecular mass fraction (<30 kDa) did not have any inhibitory effect on GOR (Fig. 1). Moreover, heat treatment (95 °C for 60 min) of the washed cytoplasmic extract eliminated the inhibitory effect, indicating that the inhibitor is likely to be a protein.

Based on its ability to inhibit purified GOR activity, the inhibiting factor was purified from the flow-through fraction of the GOR affinity purification step by hydrophobic interaction chromatography. Analysis of the purified inhibitor fraction by LC-tandem MS (LC-MS/MS) (Proteomics and Mass Spectrometry Facility, University of Georgia) identified a 70-kDa bifunctional phosphoglycerate kinase/triose-phosphate isomerase (PGK/TPI) fusion protein (Athe_1405). Addition of the purified PGK/TPI protein to the GOR assay virtually abolished GAP-oxidizing activity (Fig. 1).

|         | Protein | Volume | Activity | Specific activity | Purification | Yield |
|---------|---------|--------|----------|------------------|--------------|-------|
| S100    | 3361.9  | 235    | 33.6     | 0.01             | 1            | 100   |
| Q-HP    | 518.1   | 370    | 339      | 0.7              | 1            | 1     |
| Nickel-nitrilotriacetic acid | 2.9     | 2.4    | 144      | 50.1             | 76           | 1     |
| Sephadex 200 | 2.1    | 2.1    | 101      | 48.3             | 74           | 30    |

Figure 1. The activity of purified GOR (0.006 mg/ml) was measured in the presence of cytoplasmic extract (S100, 0.174 mg/ml), heat-treated (95 °C for 60 min) cytoplasmic extract (HT-S100, 0.036 mg/ml), cytoplasmic extract washed using a 30-kDa membrane (HT-S100, 0.325 mg/ml), and the flow-through material that passed through the 30-kDa membrane (<30 kDa-S100, protein was not measured), or partially purified PGK/TPI (C. bescii PGK/TPI, 0.002 mg/ml). The concentrations given in parentheses are the final concentration of each component in the assay cuvette.

TPI catalyzes the interconversion of GAP and dihydroxyacetone phosphate (DHAP) with a strong bias in favor of DHAP (~95:5) (30). Hence, TPI activity is responsible for the inhibition of GOR and GAPDH in cytoplasmic extracts by a “substrate-stealing” mechanism and thus its separation from these enzymes during their purification results in a dramatic increase in their GAP-dependent activities. The presence of the PGK/TPI protein in the standard GOR assay enabled reduction of BV to be measured in the presence of DHAP (which generates GAP) in the absence of added GAP (Table S2). Likewise, the presence of PGK/TPI in the standard GAPDH assay enabled the reduction of NAD if DHAP (rather than GAP) was added (Table S2). Interestingly, DHAP itself partially inhibits GOR activity in the absence of PGK/TPI and, conversely, increases GAPDH activity in the absence of PGK/TPI (Table S2). GOR activity is reduced by 70% in the presence of 10 mM DHAP in the absence of TPI, but only decreases by 22% in the presence of 1 mM DHAP. In contrast, GAPDH activity increased 4-fold in the presence of 10 mM DHAP. Notably, TPI inhibition of both can be overcome by sufficiently high concentrations of GAP (5 and 15 mM) as measured by the initial activities (Table S2). However, in all cases where PGK/TPI was present, activity diminished within 15 s, compared with >30 s in the standard assays.

A kinetic analysis of purified GOR with varying concentrations of GAP revealed a linear increase in activity, with no saturation observed up to 20 mM GAP with BV (1 mM) as the electron acceptor. Although these data precludes the determin-
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Utilization of NAD (1 mM) as the electron acceptor, with no saturation observed at GAP concentrations up to 20 mM (Fig. S3B).

Phylogenetic analyses of GOR

A phylogenetic tree of the AOR family of enzymes was constructed using 4,063 homologs of the GOR-L sequence based on the InterPro domains (IPR013983 and IPR001203). Although the five previously characterized types of tungsten-containing oxidoreductases, AOR, FOR, GAPOR, WOR4, and WOR5, are all from Archaea, such as *P. furiosus*, the AOR family is also well-represented in the bacterial domain with 64% of the sequences from Bacteria and 30% from Archaea, with the remaining (6%) from uncharacterized metagenomes or environmental samples (Fig. S9). The five types of previously characterized members of the AOR family from *P. furiosus* form distinct clades that are well-separated from each other and *C. bescii* GOR is clearly more closely related to GAPOR than it is to any of the other enzymes, sharing a more recent common ancestor. Nevertheless, the GOR and GAPOR branches also subdivide into bacterial and archaeal subclades (Fig. S9B).

Discussion

Herein we describe the first glyceraldehyde-3-phosphate ferredoxin oxidoreductase from a bacterium and designate it GOR to show that it is distinct from the previously characterized GAPOR from Archaea (16). We also demonstrate that GOR is directly involved in the glycolytic pathway of *C. bescii*. This result was unexpected because the genome of *C. bescii* encodes the standard pathway that includes the classical phos-
phorylating GAPDH. This catalyzes the conversion of GAP to 1,3-bisphosphoglycerate, which is then used to conserve energy in the form of ATP by PGK generating 3-phosphoglycerate. Note that GOR oxidizes GAP directly to 3-phosphoglycerate and so ATP is not synthesized in the GOR-dependent glycolytic pathway. Both GAPDH and PGK/TPI enzymes were partially purified in this study, indicating that the enzymes are present at substantial concentrations in the cell. However, the results of our genetic and metabolomics-based studies clearly indicate an important glycolytic role for GOR under the usual growth conditions. Moreover, deletion of either of the genes encoding GOR affects the efficient oxidation of GAP, as evident from the accumulation of metabolic intermediates upstream of this step in the glycolytic pathway and depletion of intermediates downstream (Fig. 3).

Lack of saturation kinetics for both GOR and GAPDH is a highly unusual feature for enzymes of this type. Although there are other examples of nonsaturable enzymes (e.g. AhpC from Helicobacter pylori) this phenomenon is rare (32). In the case of GOR, the order of $k_{cat}/K_m$ (4.7 × 10^4 s⁻¹ M⁻¹) indicates that the lack of saturation is not because the enzyme is diffusion limited. Instead, it likely has an extremely low affinity for GAP but with a high turnover rate, such that formation of the [ES] complex is slow, but turnover occurs rapidly upon complex formation. Nonsaturation kinetics is even more surprising for GAPDH. This enzyme has been extremely well-studied from many organisms with $K_m$ values for GAP ranging from 20 μM to 1.4 mM (33, 34). However, in an organism like C. bescii with two competing primary metabolic pathways for GAP oxidation, it is not unreasonable that the two GAP-oxidizing enzymes have low affinities for the substrate as a means of regulating substrate flux. It is also possible that another component is necessary in the assay for these enzymes to be saturable with GAP, as in the case with human ADH5 (class III alcohol dehydrogenase), which cannot be saturated with ethanol (up to 3 M) in the absence of activation by fatty acids (35).

We show here that both GAPDH and GOR activities are masked by the presence of a highly active TPI, which is biased toward DHAP formation thereby maintaining only a relatively low concentration of GAP that is available to both enzymes. The high TPI activity might play a unique thermoprotective role by minimizing the pool of the thermally-labile glycolytic intermediate GAP by reversibly converting it to the more thermostable DHAP, thus regulating the glycolytic flux through GOR and GAPDH (36). This is in contrast to the situation in some Archaea where GAPOR is the sole GAP-oxidizing enzyme in a modified EM pathway generating 3-phosphoglycerate in a single irreversible step, where GAPDH serves solely in a gluconeogenic role (23, 37, 38). The concurrent presence of GOR and GAPDH activity suggest that in C. bescii both enzymes operate in parallel. The operation of parallel pathways in glucose degradation is unusual, although not without precedent in other thermophilic microorganisms. For example, the archaeon Thermoproteus tenax has been demonstrated to utilize modified versions of both EMP and ED pathways (39–42). The inhibition of the activity of GOR by DHAP, although GAPDH appears to be activated by DHAP (Table S2), was unanticipated. However, it does explain our inability to measure significant GAP-dependent oxidoreductase activity (BV-dependent) in cytoplasmic extracts previously. That
DHAP is a regulator of both GOR and GAPDH cannot be ruled out as it could potentially activate GAPDH, whereas inhibiting GOR, but further study will be required to clarify this, especially because high and likely nonphysiological concentrations of DHAP (10 mM) were required to observe significant effects.

If the inhibition of GAPDH activity in cytosolic extracts results from PGK/TPI activity, it would be expected that this effect should have been observed in the purification of GAPDH from other organisms where TPI is present. This situation in *C. bescii* is extremely unusual as shown by comparison with a related anaerobic fermentative thermophile, *Thermotoga maritima* (T,opt 80 °C). Like *C. bescii*, *T. maritima* also contains a PGK/TPI fusion protein but its GAPDH behaves like a conventional enzyme. It shows no inhibition in cytoplasmic extracts and has been purified by conventional chromatography with no significant increases in activity (43). Notably, *T. maritima* lacks GOR. Thus, we hypothesize that the PGK/TPI effect is related to the presence of GOR in *C. bescii*. Similarly, this effect may be related to the lack of saturation kinetics discussed above as a means of regulating the dynamic pathways for GOR oxidation present in this organism.

We speculate that the role of GOR in *C. bescii* and other *Caldicellulosiruptor* species is to serve as an alternative glycolytic pathway (Fig. 4) that is used by the cell to regulate carbon and electron flux during growth under conditions in which *C. bescii* is unable to efficiently recycle its electron carriers. This might be particularly important when *C. bescii* is exposed to excess carbon loading, such as during growth on unprocessed biomass (9). For example, the genome of *C. bescii* encodes two hydrogenases, one using only reduced ferredoxin as the electron donor, whereas the other simultaneously uses both reduced ferredoxin and NADH. The latter is a cytoplasmic ferredoxin/NADH-dependent bifurcating [FeFe] hydrogenase that is the primary source of H₂ production during growth on glucose-based substrates (44). The former is a membrane-bound ferredoxin-dependent [NiFe] hydrogenase (Ech) that
can generate a proton gradient across the cell membrane (45). The standard EM pathway utilizing the classic phosphorylating GAPDH is likely the primary glycolytic route used by *C. bescii* as this pathway produces ATP. Although ferredoxin-dependent conversion of GAP to 3-phosphoglycerate by GOR yields less ATP than the standard EM pathway, it allows glycolysis to proceed with only ferredoxin as the redox carrier. This can be recycled via H₂ production by Ech, avoiding the potential generation of excess NADH. This hypothesis is supported by the finding that genes necessary for the biosynthesis of GOR are up-regulated in the closely-related *Caldicellulosiruptor saccharolyticus* in the presence of high H₂ concentrations (46), along with the recent discovery that GOR (XOR) is regulated by Rex, a redox-sensing transcription factor that responds to the intracellular NADH/NAD ratio (47). These findings suggest that high concentrations of NADH or H₂, either of which could interfere with the regeneration of oxidized cofactors via the bifurcating hydrogenase (13, 48), steer carbon flux toward GOR to utilize the more thermodynamically favorable Ech pathway, thereby maintaining a high rate of H₂ production (13, 48). However, it is not possible at present to reconcile such an analysis with the observed growth phenotype of the Δgor mutant strains, which exhibit a longer lag phase but higher cell densities compared with the parent strain (Fig. 2 and Fig. S8).

Phylogenetic analyses of the AOR family of enzymes (Fig. S9) show that each of the five characterized members of the AOR family of enzymes from *P. furiosus* can be assigned to distinct clusters, which is in accordance with their function in terms of both their substrate specificities and oligomeric structures. Hence AOR from *P. furiosus* and also from *Pyrobaculum aerophilum* are homodimeric enzymes and catalyze the oxidation of a wide range of both aliphatic and aromatic aldehydes (31, 49). FOR from *Thermococcus litoralis* and *P. furiosus* are homotetrameric enzymes but with a more limited substrate range than AOR, oxidizing short chain aliphatic aldehydes (24, 50). WOR4 and WOR5, both from *P. furiosus*, are also homodimeric enzymes. WOR5, like AOR, can oxidize a wide range of aldehydes but the substrate for WOR4 is not known and this enzyme was isolated purely based on its tungsten content (26, 51). GAPORs from *Methanococcus maripaludis, P. furiosus*, and *Pyrobaculum aerophilum* are monomeric enzymes and are absolutely specific for glyceraldehyde 3-phosphate as they are not known to oxidize any other aldehyde (23, 27, 28). We show here that *C. bescii* GOR-L is part of a distinct clade, separate from other members of the AOR family, including GAPOR (Fig. S9), its closest characterized homolog. The GOR-L clade contains all of the *Caldicellulosiruptor* species plus 44 other anaerobic genera, representing 9 taxonomic classes: Archaeoglobi, Bacilli, Clostridia, Deltaproteobacteria, Fusobacteria, Methanomicrobia, Nitrospira, Thermodesulfbacteria, and Thermoprotei. Although many of the genera are thermophilic or hyperthermophilic some are mesophilic; however, almost all species with GOR-L also have a co-located GOR-S homolog (SynTax web server) (52). Moreover, this analysis shows that, unlike GAPOR, which is found primarily in Archaea, GOR is found primarily in Bacteria. For example, the GOR-L clade could be divided into three subclades representing a bacterial-only clade, making up 54% of all GOR-L clade sequences, an archaeal-only making up 26%, and a mixed clade (split 50/50) making up 20% (Fig. S9).

We also show here that GOR is part of a small minority of AOR family members that are heteromeric. These include a homolog of AOR (termed Aa AorB) that is a heterotrimer containing the equivalent of the GOR-S subunit with a string of iron-sulfur clusters (53). In addition, a second heteromeric member of the AOR family described recently is part of an enzyme complex that reduces benzoyl-CoA (54). Termed BamB, the AOR homolog also has a partner subunit (BamC) with multiple iron-sulfur clusters. Furthermore, of the characterized AOR family members from hyperthermophilic Archaea, the gene encoding WOR5 has an adjacent gene homologous to that encoding GOR-S, suggesting that WOR5, like GOR, has an additional iron-sulfur containing subunit that was likely not detected during standard SDS analysis. Finally, the heteromeric AOR family group also includes an AOR from *Escherichia coli* termed EcYdhV that contains a polyferredoxin-like subunit similar to GOR-S. Interestingly, YdhV contains molybdenum rather than tungsten as *E. coli* does not metabolize tungsten (55). It is not clear why GOR and these other AOR family members require an additional subunit with four [4Fe-4S] clusters, which in GOR are proposed to transfer electrons between the catalytic subunit (GOR-L) and ferredoxin. GOR has a high affinity for *C. bescii* ferredoxin (Kₘ ~ 40 μM), consistent with this being the physiological electron acceptor, although it is possible that GOR could provide reductant to the Ech hydrogenase directly, whereas ferredoxin is required to interact with the bifurcating hydrogenase.

In any event, the results presented here lay the foundation for a more comprehensive understanding of the two glycolytic pathways that are present in the cellulolytic genus *Caldicellulosiruptor* and for more effective metabolic engineering projects in the future. Moreover, this work suggests that alternative glycolytic pathways using either GOR or GAPOR may be more widespread than previously thought in both the bacterial and archaeal worlds.

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**Experimental procedures**

**Growth of *C. bescii***

*C. bescii* strains used or constructed in the present study are listed in Table S3. Strains were transformed as previously described (16, 56). For growth of uracil auxotrophic strains, uracil was added at a concentration of 20 μM. When required, kanamycin and 5-fluoroorotic acid were used at concentrations of 50 μg/ml and 4 mM, respectively, unless otherwise noted. Growth experiments were performed with 5-ml cultures in screw cap Hungate tubes sealed with butyl rubber stoppers in modified DSMZ 516 medium with the following composition per liter: 1X salt solution, 1X vitamin solution, 1X trace element solution, 0.16 μM sodium tungstate, 0.25 mg of resazurin, 5 g of xyllose (unless otherwise noted), 1 g of cysteine hydrochloride, 1 g of sodium bicarbonate, and 1 mM potassium phosphate buffer (pH 7.2). The ×50 stock salt solution contained the following per liter: 16.5 g of NH₄Cl, 16.5 g of KCl, 16.5 g of MgCl₂·6H₂O, and 7 g of CaCl₂·2H₂O. Stock solutions of ×200 vitamins and ×1,000 trace elements were prepared as previ...
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GorL activity is described (1). Cultures were also grown in a 15-liter fermenter at 75 °C with 20 g/liter of xylose as the growth substrate. The culture was sparged at 2 liters/h with 80% (v/v) nitrogen and 20% (v/v) carbon dioxide and stirred at 150 rpm with a flat blade impeller. The pH was maintained at 7.0 (25 °C) by the addition of 10% (w/v) sodium bicarbonate. Cells were harvested in the late-exponential phase by centrifugation at 6,000 × g for 10 min (Beckman Avanti J-30I JLA 10.500 rotor) and the resulting cell pellets were flash frozen in liquid nitrogen and stored at −80 °C. The cell yield from the fermenter was ~12 g (wet weight).

**Vector construction**

Plasmids were constructed from PCR products using NEBuilder® HiFi DNA Assembly Master Mix (New England BioLabs). PCR was performed using Takara PrimeSTAR Max DNA polymerase (Takara R045A). Genomic DNA, plasmid DNA, and PCR products were purified with kits from Zymo Research and Stratagene. The templates for PCR were either previously sequenced plasmids or genomic DNA from *C. bescii*. All primers used in this study can be found in Table S5 and plasmids constructed for this study can be found in File S10. Plasmids were cloned into New England Biolabs 10-β competent *E. coli* (New England BioLabs C30191I), followed by Sanger sequencing through GeneWiz.

**Metabolomics**

For the MS-based metabolomics, parent and ΔgorL strains of *C. bescii* were grown in duplicate in 16-mm sealed hange tubes in modified DSMZ 516 medium as described above with glucose as the carbon source. Cultures were grown to an OD<sub>600</sub> of ~0.15 and each culture was deposited by vacuum filtration onto a 0.2-μm nylon membrane (47 mm diameter). The membrane was then placed (cells down) into 1.5-ml cold (−20 °C) dry ice extraction solvent (20:20:10 (v/v/v), acetonitrile:methanol:water) in a small Petri dish and swirled. After a few moments the filter was inverted (cells up) and solvent was passed over the surface of the membrane several times to maximize extraction. Finally, the cell extract was stored at −80 °C. Cell extracts from *E. coli* (K-12 substrain MG1655 rph<sup>/H11001</sup> ilvG<sup>/H11001</sup>) grown in [13C]glucose were used as internal standards for quantitation of metabolites in *C. bescii* strains. *E. coli* cells were grown aerobically to an OD<sub>600</sub> of ~0.45 in M9 minimal medium containing 0.4% universally labeled [1<sup>13</sup>C]<sub>glucose</sub> as the sole carbon source. To assure complete labeling of metabolites in *E. coli*, inoculation was performed using a 1:50 dilution from an overnight culture that was also grown on [13C]-labeled glucose. *E. coli* metabolites were extracted in a similar manner to the procedure used for *C. bescii*.

*C. bescii* and [13C]-labeled *E. coli* extracts were thawed, centrifuged at 14,000 rpm for 10 min to remove cell debris, and the *C. bescii* supernatants were mixed with [13C]-labeled *E. coli* supernatant in ratios of 1:1, 5:1, and 1:0 (no [13C]-labeled internal standards). These mixed samples were then fully dried under a stream of nitrogen and subsequently re-suspended with HPLC running solvent (97:3, water:methanol containing 10 mM tributylamine and 9.8 mM acetate, pH 8.2) in half the initial volume. Samples were then centrifuged to remove particulates, transferred to HPLC vials, and analyzed by HPLC/MS for quantification as previously described (57).

**Enzyme assays**

All assays were carried out in serum-stopped cuvettes under anaerobic conditions at 70 °C, unless noted otherwise. To remove trace amounts of O<sub>2</sub>, sodium dithionite was added to the assay mixture to give an A<sub>0</sub> of ~0.2. The extract was added and after a 1-min incubation period the reaction was initiated by addition of the substrate. Aldehyde-oxidizing activity of cytoplasmic extracts of *C. bescii* strains and purified GOR were determined by following the reduction of ferredoxin or BV (1 mM) in 50 mM EPPS buffer (pH 8.0), at 425 or 600 nm, respectively, using various aldehydes (1 mM) as the substrate. With GAP as the substrate the assay temperature was 70 °C. GAPDH activity was measured by following GAP-dependent reduction of NAD (1 mM) at 320 nm. Extinction coefficients of 8.2, 6.2, and 8.1 mM<sup>−1</sup> cm<sup>−1</sup> were used for reduced benzyl viologen, NADH, and reduced ferredoxin, respectively. Specific enzyme activities are expressed as units/mg of protein, where 1 unit represents 1 μmol of substrate oxidized per min.

**Purification of affinity-tagged GOR**

All purification steps were performed under strictly anaerobic conditions. Frozen cells (50 g, wet weight) of *C. bescii* strain OE XOR/PFD were thawed and suspended in 25 mM Tris-HCl (pH 8.0), containing 1 mg ml<sup>−1</sup> lysozyme (Sigma) in a ratio of 3 ml/g of cells. The suspended cells were incubated at room temperature for 15 min followed by four 15-s intervals of sonication (amplitude 40; Qsonica Q55) interspaced by at least 30 s. Cell lysates were clarified by ultracentrifugation at 100,000 × g for 1 h (Beckman L90K ultracentrifuge 70.1Ti rotor) yielding clarified cell lysate (235 ml, 14.3 mg of protein ml<sup>−1</sup>). This was loaded onto a 250-ml QHP column equilibrated with buffer A (25 mM Tris, pH 8.0, containing 2 mM cysteine), at a rate of 10 ml min<sup>−1</sup> diluted 1:5 with buffer A. Protein was eluted with a linear gradient (2500 ml) from 0 to 500 mM NaCl at 10 ml min<sup>−1</sup> and collected as 75-ml fractions. Fractions were assayed for GOR and GAPDH activity. Fractions containing GAPDH activity were reserved for further purification, whereas fractions containing GOR activity were pooled and applied to a 5-ml HisTrap Excel column (GE Healthcare) equilibrated in buffer B (50 mM phosphate, pH 7.2, containing 300 mM NaCl). The column was washed with 5 volumes of buffer B, and GOR was eluted with a gradient of 0 to 500 mM imidazole in buffer C (buffer B containing 500 mM imidazole) over 20 column volumes. Fractions were collected in sealed serum bottles. Fractions containing GOR activity were pooled and concentrated (15-kDa Amicon Ultra centrifugal filter) to yield 2.9 ml of GOR (1.2 mg/ml). This was loaded onto a HiLoad 16/60 Superdex 200 column (GE Healthcare) equilibrated with 50 mM Tris-HCl (pH 8.2), containing 200 mM NaCl at a rate of 1.25 ml min<sup>−1</sup> and collected in 4-ml fractions in sealed serum bottles. Protein-containing fractions were analyzed for purity using denaturing SDS-PAGE gradient electrophoresis using precast 4–12% BisTris gels; Novex NuPAGE gels (Invitrogen) and stained with Imperial protein stain (Novex). Protein bands were identified via a Mascot MS/MS Ions Search of a custom
C. bescii database (version 2.6) using LC-tandem MS (LC-MS/MS) data (Proteomics and Mass Spectrometry Facility, University of Georgia).

Partial purification of GAPDH

To further purify GAPDH, the fraction containing the highest GAPDH activity from the QHP column was concentrated (10-kDa Amicon Ultra centrifugal filter) to yield 4.1 ml (14.2 mg/ml) and loaded onto a HiLoad 16/60 Superdex 200 and eluted in the same manner as above. Fractions containing GAPDH activity were pooled and applied to a 1-ml HiTrap Blue HP column (GE Healthcare) equilibrated in 20 mM EPPS (pH 8). The column was washed with 5 column volumes of the equilibration buffer. Proteins were eluted with 2 ml of the equilibration buffer containing 20 mM NAD. The eluate was concentrated and buffer exchanged (10-kDa Amicon Ultra centrifugal filter) into 50 mM EPPS (pH 8) to remove NAD, yielding 0.4 ml of partially purified GAPDH (1.6 mg/ml).

Estimation of molecular mass and metal analysis

For size determination of the purified GOR, the elution volume ($V_e$) were determined by measuring the absorbance at 280 nm and the total void volume ($V_0$) was determined by measuring the elution of blue dextran. The known molecular mass standards thyroglobulin (669 kDa), apoferritin (443 kDa), amylase (200 kDa), alcohol dehydrogenase (150 kDa), albumin (66 kDa), and carbonic anhydrase (29 kDa) were used as references. The molecular mass of the purified protein in each peak was estimated from the plot of $V_e/V_0$ against the logarithm of the molecular mass (kDa). Metal concentrations of the final chromatography fractions were measured using an Agilent 7900 inductively coupled plasma mass spectrometer (ICP-MS) fitted with MicroMist nebulizer, UHMI-spray chamber, Pt cones, and an Octopole Reaction System collision cell (Agilent Technologies, Santa Clara, CA) as described previously (58).

Purification of PGK/TPI

To isolate the GOR inhibiting factor, the flow-through from the affinity purification of GOR described above was applied to a 60-ml Phenyl-Sepharose column equilibrated with buffer A containing 1 M (NH₄)₂SO₄. Protein was eluted with a linear gradient (800 ml) from 1.0 to 0 M (NH₄)₂SO₄ at 7.5 ml/min and loaded onto a HiLoad 16/60 Superdex 200 and eluted in the same manner as above. Fractions containing GAPDH activity were pooled and applied to a 1-ml HiTrap Blue HP column (GE Healthcare) equilibrated in 20 mM EPPS (pH 8). The column was washed with 5 column volumes of the equilibration buffer. Proteins were eluted with 2 ml of the equilibration buffer containing 20 mM NAD. The eluate was concentrated and buffer exchanged (10-kDa Amicon Ultra centrifugal filter) into 50 mM EPPS (pH 8) to remove NAD, yielding 0.4 ml of partially purified GAPDH (1.6 mg/ml).

Phylogenetic analyses

In InterPro (version 63.0), GOR-L (Atthe_0821; UniProt ID: B9MQ12_CALBD) is annotated to have one aldehyde ferredoxin oxidoreductase N-terminal (IPR013983) domain and one aldehyde ferredoxin oxidoreductase C-terminal (IPR001203) domain (59). One of each domain was entered into the InterPro Domain Architecture search tool, yielding 4,107 GOR-L sequence homologs. The 4,107 sequences were aligned using Clustal Omega, version 1.2.1, with the default parameters (60). TrimAl was used to remove multiple alignment positions with greater than 99.7% gaps (i.e. –gt 0.003) (61), and the resulting alignment was used to construct a maximum likelihood phylogenetic tree using IQ-Tree (version 1.5.5) (62). The IQ-TREE standard model selection test was used to automatically determine the best-fit model (i.e. LG + F + R10) (63). This model tree was refined using ultrafast bootstrap approximation (UFBoot) and Shimodaira-Hasegawa–like approximate likelihood ratio tests with 3,000 bootstrap replicates for each. Only sequences that were not marked deleted by Uniprot were used to display the tree, leaving 4,063 sequences. iTOL (version 4.2.1) was used for analysis and display of the phylogenetic tree (64).

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