The Physiological Functions and Structural Determinants of Catalytic Bias in the [FeFe]-Hydrogenases CpI and CpII of Clostridium pasteurianum Strain W5

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The first generation of biochemical studies of complex, iron-sulfur-cluster-containing [FeFe]-hydrogenases and Mo-nitrogenase were carried out on enzymes purified from Clostridium pasteurianum (strain W5). Previous studies suggested that two distinct [FeFe]-hydrogenases are expressed differentially under nitrogen-fixing and non-nitrogen-fixing conditions. As a result, the first characterized [FeFe]-hydrogenase (CpI) is presumed to have a primary role in central metabolism, recycling reduced electron carriers that accumulate during fermentation via proton reduction. A role for capturing reducing equivalents released as hydrogen during nitrogen fixation has been proposed for the second hydrogenase, CpII. Biochemical characterization of CpI and CpII indicated CpI has extremely high hydrogen production activity in comparison to CpII, while CpII has elevated hydrogen oxidation activity in comparison to CpI when assayed under the same conditions. This suggests that these enzymes have evolved a catalytic bias to support their respective physiological functions. Using the published genome of C. pasteurianum (strain W5) hydrogenase sequences were identified, including the already known [NiFe]-hydrogenase, CpI, and CpII sequences, and a third hydrogenase, CpIII was identified in the genome as well. Quantitative real-time PCR experiments were performed in order to analyze transcript abundance of the hydrogenases under diazotrophic and non-diazotrophic growth conditions. There is a markedly reduced level of CpI gene expression together with concomitant increases in CpII gene expression under nitrogen-fixing conditions. The Physiological Functions and Structural Determinants of Catalytic Bias in the [FeFe]-Hydrogenases CpI and CpII of Clostridium pasteurianum Strain W5. Front. Microbiol. 8:1305. doi: 10.3389/fmicb.2017.01305
their catalytic activities in vitro, with CpII capturing excess reducing equivalents under nitrogen fixation conditions. Comparison of the primary sequences of CpI and CpII and their homologs provides an initial basis for identifying key structural determinants that modulate hydrogen production and hydrogen oxidation activities.

Keywords: hydrogenase, nitrogenase Clostridium pasteurianum, hydrogen metabolism, nitrogen metabolism, CpI, CpII

INTRODUCTION

The genus Clostridium includes a diverse group of Gram-positive, spore-forming anaerobes (Patakova et al., 2013). In general, clostridial fermentative metabolism functions by the conversion of hexose sugars to butyrate, acetate, and CO₂. During this process reduced electron carriers in the form of ferredoxin accumulate and must be recycled for sustained fermentative energy metabolism. C. pasteurianum recycles reduced ferredoxin by coupling electrons and protons to form hydrogen (H₂) through the activity of a hydrogenase. C. pasteurianum may also fix nitrogen during fermentative growth, a process that requires high amounts of both ATP and reducing equivalents (Mortenson, 1964).

Clostridium pasteurianum strain W5 has been a model for studying the biochemistry of nitrogen fixation and H₂ metabolism. The first preparations of a soluble hydrogenase (CpI) were obtained from this organism (Valentine et al., 1963), and subsequently, the presence of a second [FeFe]-hydrogenase (CpII) was revealed (Chen and Blanchard, 1978), and its physical and catalytic properties were studied along with those of CpI (Adams and Mortenson, 1984a).

[FeFe]-hydrogenase 1 from CpW5 was proposed to evolve H₂ to recycle electron carriers during fermentative growth in the presence of fixed nitrogen (Adams and Mortenson, 1984a). CpII was proposed to function under nitrogen-fixing conditions to capture reducing equivalents in the form of H₂ which is an obligate byproduct of nitrogenase-catalyzed reduction of nitrogen to ammonia. This is consistent with the observations that CpII accumulates at a higher cellular concentration during diazotrophic growth (Chen and Blanchard, 1978). Comparison of the rates of H₂ evolution and oxidation revealed that, while these two enzymes are both reversible in vitro, CpI produces H₂ 550-times faster than CpII (5,500 vs. 10 μmol of H₂/min-mg, respectively) while it oxidizes H₂ about 30% slower than CpII (24,000 vs. 34,000 μmol of H₂/min-mg, respectively) (Adams, 1990). Typically, [FeFe]-hydrogenases have characteristically high catalytic rates for H₂ production (Adams, 1990). Therefore, the two [FeFe]-hydrogenases exhibit a strong “catalytic bias,” which is manifested as the adaptation of CpII toward H₂ oxidation.

Hydrogenases have been characterized as providing many functions in vivo, including disposal of excess reducing equivalents (Verhagen et al., 1999; Tamagnini et al., 2007; Schut and Adams, 2009), hydrogen sensing (Lenz et al., 2002; Brecht et al., 2003), ion translocation (Hedderich, 2004), and oxidizing H₂ (Dementin et al., 2004; Kanai et al., 2011), a key method by which reducing equivalents are generated. This diversity of function allows hydrogenases to perform a critical role in a variety of metabolisms, including those of methanogens (Thauer et al., 2010), sulfate reducers (Baltazar et al., 2011), nitrogen fixers (Zhang et al., 2014), and photosynthesizers (Melis et al., 2004; Ghirardi et al., 2006; Tamagnini et al., 2007).

In this work we provide experimental evidence that under nitrogen replete conditions (in the absence of nitrogenase), CpI functions to reduce protons during the recycling of electron carriers during fermentation, while CpII functions in H₂ oxidation under diazotrophic conditions. The genome of C. pasteurianum ATCC 6013 (strain W5) (Rotta et al., 2015) was subjected to homology searches using known hydrogenase sequences as queries to determine the complement of encoded hydrogenases, their sequences and their gene context. Using these data, we analyzed the transcript abundance of each hydrogenase under nitrogen-fixing and nitrogen-replete culture conditions to assign physiological roles for CpI and CpII. Furthermore, detailed primary amino acid structural-based comparison together with phylogenetic analysis provide insights into the determinants of the profound catalytic bias observed for these two related enzymes.

RESULTS AND DISCUSSION

Genome

The sequencing of the C. pasteurianum strain W5 (CpW5) genome was carried out independently of the recently published complete genome (Rotta et al., 2015). Our analysis resulted in a draft genome consisting of 14 contigs and 4.2 Mbp that shares 99.97% average nucleotide identity with the published genome (Supplementary Figure 1). The published complete genome contains 4.3 Mbp, which indicates that our genome is nearly complete. In particular, the sequences of the genes encoding all four hydrogenases discussed in the present study are identical to those in the complete genome (Rotta et al., 2015). Like the genomes of other clostridial species (Sakaguchi et al., 2005; Yutin and Galperin, 2013; Sedlar et al., 2015), the GC content of CpW5 was low at 30.0%. C. pasteurianum NRRL B-598, which is an oxygen-tolerant species, is also related to CpW5 and has a genome size that is ~50% larger, comprising 6.1 Mbp (Kolek et al., 2014). According to SEED Viewer (Overbeek et al., 2014), which does not include sequences from these C. pasteurianum genomes (i.e., ATCC 6013 DSM 525 and NRRL B-598), the closest

Abbreviations: CpI, [FeFe]-hydrogenase 1 from CpW5; CpII, [FeFe]-hydrogenase 2 from CpW5; CpIII, [FeFe]-hydrogenase 3 from CpW5; CpW5, Clostridium pasteurianum ATCC 6013 (strain W5); WT, wild type.
neighbors with completed genomes are *C. acetobutylicum* (3.94 Mbp) (Nolling et al., 2001), *C. botulinum* (3.89 Mbp) (Sebaihia et al., 2007), *C. novyi* NT (2.55 Mbp) (Bettegowda et al., 2006), and *C. sporogenes* ATCC 15579 (4.09 Mbp) (Poehelein et al., 2015).

**Hydrogenases**

The genome of CpW5 encodes the two characterized [FeFe]-hydrogenases, CpI and CpII, and an additional homolog designated CpIII, as well as one (previously annotated) [NiFe]-hydrogenase (Pyne et al., 2014), together with all of the necessary genes for hydrogenase maturation. These sequence data therefore allow us to carry out the first comparative analysis of the primary sequence of CpII since it was biochemically characterized more than two decades ago (Adams and Mortenson, 1984a).

The sequences of CpI and CpII are 33% identical, with 45% identity and 61% similarity over the conserved region (Figure 1), which suggests that these two enzymes have generally conserved protein architectures. A homology model of CpII (Figure 2) based on the solved crystal structure of CpI (Peters et al., 1998) and generated using SwissModel (Arnold et al., 2006), as well as amino acid sequence alignment, indicate the absence of accessory domains in CpII that are present in CpI. The CpI sequence contains conserved cysteine residues for each Fe/S cluster that sequentially bind clusters [2Fe-2S] (FS2), the distal [4Fe-4S] cluster (FS4C), the medial [4Fe-4S] cluster (FS4B) and the proximal [4Fe-4S] cluster (FS4A). In contrast, the N-terminus of CpII lacks the cysteine residues responsible for binding accessory Fe/S clusters FS2 and FS4C (Figure 2).

However, conserved regions binding the two Fe atoms of the catalytic site, known as the H-cluster, and two [4Fe-4S] accessory clusters were identified in CpII. CpIII, which has thus far not been biochemically characterized, has a unique N-terminal arrangement of cysteines. Sequence alignment reveals that the FS4A binding motif is conserved, while the FS4B motif lacks two of the four cysteine residues that typically ligate this cluster (Figure 1). CpIII therefore has significant sequence differences from other biochemically characterized hydrogenases, which may provide it with intriguing properties.

The [FeFe]-hydrogenase sequences encoded in the CpW5 genome contain the evolutionarily conserved H-cluster motifs; TSCCPxW (L1), MPCxxKxxE (L2), and ExMxCxxGCxxG (L3) (Figure 1; Vignais and Billoud, 2007; Schmidt et al., 2010).
These motifs include all of the H-cluster binding cysteines, as well as residues for ligating accessory clusters (Figure 1). Phylogenetic clustering of the H-cluster domains of clostridial [FeFe]-hydrogenases has shown a variety of distinct clusters, designated A1-A5, A7, A8, and B1-B3 (Calusinska et al., 2011). Group A2, which includes CpI, is comprised of monomeric, soluble, H₂-producing enzymes. CpII, which lacks the [2Fe-2S] cluster as well as one of the [4Fe-4S] clusters, is classified as an A3 hydrogenase. Members of group B2, that includes CpIII, have an average size of 450 amino acids and an additional characteristic cysteine residue in the P1 motif (TSCCPART) of the H-cluster (Calusinska et al., 2010). No hydrogenases of this type have been biochemically characterized to date, and it is unclear if these sequences produce active hydrogenases. CpI, CpII, and CpIII are predicted to be monomeric, group 1 enzymes based on flanking gene analysis according to the classification system of Poudel et al., 2016, and are thus not expected to bifurcate (Poudel et al., 2016).

The [NiFe]-hydrogenase gene cluster of CpW5 contains the required accessory genes (hypABCEFD and hoxN) downstream of the structural genes, hyaAB, which encode the large and small subunits, respectively. The predicted protein sequence of the large subunit contains previously described (Vignais and Billoud, 2007) L1 and L2 motifs characteristic of membrane-bound, uptake hydrogenases. The L1 and L2 motifs encompass the highly conserved cysteine pairs (CxxC) near each terminus that ligate the NiFe center. Unlike [FeFe]-hydrogenases, maturases for the [NiFe]-hydrogenase are often found in a single gene cluster with the structural genes (Casalot and Rousset, 2001), as is the case for the CpW5 genome. This gene cluster is not co-localized with any other hydrogenase or nitrogenase genes. The gene for the hydrogenase large subunit (hyaB) clusters phylogenetically with other clostridia in group 1 (Calusinska et al., 2010) (data not shown), which comprises membrane-associated uptake hydrogenases (Vignais and Billoud, 2007).

To understand hydrogenase metabolism in CpW5 fully, it was necessary to include transcriptional information for all of the encoded hydrogenases, including the [NiFe]-hydrogenase.

**Transcriptional and Physiological Analyses**

Quantitative real-time PCR assays were performed to compare the transcript abundance of each CpW5 hydrogenase under both non-nitrogen-fixing and nitrogen-fixing conditions. Transcript levels for CpI decreased under nitrogen-fixing conditions (2.9-fold change) while CpII transcripts increased (7.5-fold change). The transcript levels of CpIII and the [NiFe]-hydrogenase were low with and without fixed nitrogen. For CpIII, relative transcript abundances were approximately 1% of those for CpI or CpII, based on the amplification threshold for each gene, and slightly lower under diazotrophic conditions (50% decrease). The [NiFe]-hydrogenase transcript levels were approximately 1% those of CpII under non-nitrogen-fixing conditions, but underwent an 8.7-fold increase during diazotrophic growth. Despite this increase in transcript abundance for the [NiFe]-hydrogenase, the relative transcript abundance of the [NiFe]-hydrogenase was approximately 10% of the CpII transcript abundance under nitrogen-fixing conditions. Thus, low levels of [NiFe]-hydrogenase transcripts are present under both nitrogen replete and nitrogen-fixing conditions.
Collectively, the abundance of hydrogenase transcripts agrees with previously established protein expression conditions and supports a rational model of hydrogenase usage by CpW5: CpI, which is known to be abundantly expressed under standard, non-nitrogen-fixing fermentative conditions (Adams et al., 1989), functions to dispose of excess reducing equivalents as H₂, whereas under diazotrophic conditions, dinitrogen reduction by Mo-nitrogenase consumes a large amount of electrons and therefore subverts the need for an electron-consuming, proton-reducing enzyme. While CpI and CpII are ATP-independent, ATPases are necessary for nitrogenase to reduce dinitrogen to ammonia (Taylor, 1969).

In contrast, CpII has an exceptionally low H₂ production activity (Adams and Mortenson, 1984b) and thus it is unlikely that it is capable of removing excess reducing equivalents. Rather, its high H₂ oxidation activity and almost negligible proton reduction capacity is consistent with this hydrogenase functioning in the uptake direction. This H₂ oxidation is of particular importance for recycling electrons from the nitrogenase-produced H₂ and feeding those electrons back into the reductant-consuming, nitrogen-fixing metabolism. In this model, CpII thereby acts to recycle reducing equivalents, mitigating loss of electrons from H₂ produced by nitrogenase (Figure 3). The specific catalytic abilities of CpI and CpII, along with the changes in transcript abundance, suggest that there is little interaction between these hydrogenases. The [NiFe]-hydrogenase has a similar transcriptional profile to CpII, which is up-regulated under nitrogen-fixing conditions. This suggests that the [NiFe]-hydrogenase may also have a role in recapturing reducing equivalents, as has been previously demonstrated for [NiFe]-hydrogenases in aerobic, nitrogen-fixing organisms (Walker and Yates, 1978; Laane et al., 1979; Walker et al., 1981; Hamilton et al., 2011), but our data suggest a much lower abundance of [NiFe]-hydrogenase transcripts compared to CpII transcripts. In vitro data has shown that C. pasteurianum hydrogenase donates electrons to nitrogenase (Mortenson, 1964), similar to the observed roles of hydrogenases across a diversity of species (Robson and Postgate, 1980; Bothe et al., 2010). The qRT-PCR data are consistent with previous observations that CpII is expressed primarily under nitrogen-fixing conditions and that CpI and CpII account for the majority of the total hydrogenase activity observed during protein purification (Chen and Blanchard, 1978; Adams and Mortenson, 1984b). The proficiency of CpII at H₂ oxidation precludes the necessity of the [NiFe]-hydrogenase functioning to recycle reducing equivalents during nitrogen fixation, however, it may have its own metabolic niche, such as in Fe-limited circumstances. In the case of CpIII, the miniscule changes in transcript abundance, and apparent lack of protein expression, suggest that this enzyme does not significantly contribute to C. pasteurianum metabolism under conditions such as those measured in this study.

The different catalytic rates and metabolic functions of CpI and CpII provide a unique system for the study of catalytic bias.
The specific catalytic abilities of these hydrogenases function to enhance one direction of the reaction while minimizing the other; they demonstrate the complex interplay between the metabolic niche of a protein and the structural fine-tuning it must possess to perform a particular function. There is a large selective advantage to having an enzyme with very low rates of proton reduction, such as CpII, because such enzymes are likely to be operating near equilibrium under nitrogen-fixing conditions, and the slow rates of proton reduction would reduce the loss of precious reducing equivalents.

Of more than 40 fully sequenced genomes of *Clostridium* species, all but two (*C. kluyveri* and *C. butyricum*) encode nitrogenase and a [NiFe]-hydrogenase. The two exceptions have genes encoding three or more [FeFe]-hydrogenases, suggesting that one or more of the [FeFe]-hydrogenases serves to consume H₂ and thereby supplants the need for a [NiFe]-hydrogenase during nitrogen fixation.

**Structural Basis for Catalytic Bias in [FeFe]-Hydrogenases**

[FeFe]-hydrogenase I from CpW5 and CpII display substantial differences in the ability to reduce protons and oxidize H₂, and we hypothesize that these differences are due to structural differences and thus to their amino acid sequences. Comparison of activities among hydrogenases does not reveal a trend in H₂ oxidation to production ratios based on the differing FeS cluster binding motifs (Mulder et al., 2011). It is currently unclear to what extent the presence of additional Fe/S clusters contributes to differences in catalytic biases amongst these hydrogenases. Most likely, a suite of structural features is responsible for tuning the directionality of a given hydrogenase. The particular amino acids involved in gas channel lining (Liebgott et al., 2011), proton transfer (Morra et al., 2015), electron transfer, and H-cluster ligand environment (Knörzer et al., 2012) may all play a role.

Although the motifs coordinating H-clusters are conserved across [FeFe]-hydrogenases, amino acids in the second coordination sphere are not conserved. For example, three residues near the 2Fe subcluster, A230, I268, and M353 in CpI, are S99, T137, and T223 in CpII (Figure 4). These differences highlight how variation in the second coordination sphere may play a role in modulating catalytic bias. Previous work by Knörzer et al. (2012) showed that Thr (T137 in CpII) is the most frequent substitution for a Met (residue 353 in CpI) that is adjacent to the μ-CO of the 2Fe subcluster, in 409 CpI homologs. These authors used site-directed substitution to change M353 to L353 in CpI and observed a significant decrease in H₂ production (to 15% of WT enzyme) and a small decrease in H₂ oxidation (to 74% of WT), which they attributed to a lower turnover rate (Knörzer et al., 2012). This suggests that close proximity of this residue to 2Fe influences the enzymatic preference for H₂ oxidation or production, and that Leu results in an enzyme that favors oxidation to a greater degree relative to one that has Met.

A further comparison of the crystal structure of CpI with the CpII homology model revealed 14 potential sites (four at the FS4B, one at the FS4A, and nine in the H-cluster region) that may influence the redox potential of the Fe/S centers and thereby alter the catalytic bias (Figure 5). These sites were identified by examining amino acids that differed between CpI and CpII and were found within 5 Å of the FeS clusters. By cross-checking these residues with their conservation percentage and phylogenetic signal (K-statistic) among [FeFe]-hydrogenases, it is evident that most of these 14 residues are under strong selective pressure, demonstrating the functional importance of these residues and positions. These findings agree with evidence suggesting the importance of accessory clusters on the catalytic properties of hydrogenases (Abou Hamdan et al., 2012; Winkler et al., 2013). Electrochemical investigations provide evidence that the terminal cluster of the electron transfer pathway within

![FIGURE 4](https://www.frontiersin.org) A zoomed in comparison of several possible key residues in the active site that influence catalytic bias, including alanine 230, isoleucine 268, and methionine 353 in CpI, which correspond to a serine, threonine, and threonine in CpII, respectively.
enzymes influence catalytic bias (Hexter et al., 2012), for example, in *E. coli* [NiFe]-hydrogenase 1 (Armstrong et al., 2016). Most likely, the determinants of catalytic bias are not found at a single site, but are rather a suite of residues that act in concert with one another. The work presented here suggests 14 specific amino acids that may influence the electronic properties of the accessory FeS clusters as well as the active site, and provides a platform for future studies using a site-directed mutagenesis approach.

**CONCLUSION**

In this work the complement of hydrogenases from CpW5 was analyzed to provide better insights into the H\(_2\) metabolism of this organism. The genome revealed sequences of three [FeFe]-hydrogenases and a [NiFe]-hydrogenase that allowed, for the first time, a comparison of the deduced amino acid sequences of the two biochemically characterized [FeFe]-hydrogenases, CpI and CpII, which have a sequence similarity of 61%. Targeted transcriptional analyses are consistent with a role for CpII in recapturing the reducing equivalents that are produced as H\(_2\) as part of Mo-nitrogenase catalysis during growth under nitrogen-fixing conditions. A role for CpII in H\(_2\) uptake is also consistent with the previously observed catalytic bias of CpII toward H\(_2\) oxidation. CpII probably evolved to be a poor proton-reducing enzyme, thereby limiting potential loss of H\(_2\) under nitrogen-fixing conditions when the availability of reducing equivalents may be growth-limiting. A comparison of the CpI and CpII sequences in the context of their respective phylogenetic and structural relationships reveal several likely determinants of catalytic bias, which can be studied by characterization of site-specific variants of these [FeFe]-hydrogenases.

**MATERIALS AND METHODS**

**Growth Conditions**

Freeze-dried *C. pasteurianum* strain W5 (ATCC\textsuperscript{\textregistered} 6013\textsuperscript{TM}) was obtained from ATCC and rehydrated with Difco\textsuperscript{\textregistered} Reinforced Clostridial (DRC) Medium following the ATCC protocol. Sealed 25 ml glass serum vials (Wheaton) containing 10 ml of DRC medium under a headspace of 10% H\(_2\)-10% CO\(_2\)-80% N\(_2\) were then inoculated with the rehydrated culture and incubated at 37°C following ATCC propagation procedures for this organism. Agar plates prepared with DRC medium were used to store *C. pasteurianum* strain W5 for further use. Plates were incubated at 37°C for 24–48 h and were then stored at room temperature in an anaerobic chamber.
For genome sequencing, cultures were inoculated from a single colony from a DRC agar plate that was inoculated into a sealed 25-ml serum vials containing 10 ml of the same medium. Cells were grown under a headspace of 10% H\textsubscript{2}-10% CO\textsubscript{2}-80% N\textsubscript{2} by overnight incubation at 37°C. An aliquot (1 ml) of the culture was spun down at 14,000 \times g at room temperature before extracting DNA.

**Genome Sequencing**

Total genomic DNA of *C. pasteurianum* strain W5 was extracted using a Promega Wizard\textsuperscript{®} Plus SV miniprep DNA purification system. The concentration (220 ng/\mu l) was determined by a NanoDrop 1000 Spectrophotometer (A\textsubscript{\text{260/280}} = 2.04). Genomic DNA was submitted to the Genomics Core Facility at The Pennsylvania State University for 454 pyrosequencing (Mardis, 2008). Reads were assembled with the Newbler assembler (ver. 2.6; Roche) into 145 contigs of at least 500 bp with 116 of those contigs predicted to form six large scaffolds. The read depth was about 19X.

Gaps were closed by PCR using primers designed approximately 200 bases from the end of each contig. GoTaq\textsuperscript{®} 2x Master Mix (Promega, Madison, WI, United States) was used for the amplification reactions in a Techne Touchgene Gradient Thermal Cycler (Techne, Bulington, NJ, United States). Amplicons were then purified either directly using QIAquick PCR Purification Kit or from agarose gels using the Qiaex II Gel Extraction Kit (Qiagen, Valencia, CA, United States). Purified PCR products were sequenced by Davis Sequencing in Davis, CA, and sequence data were assembled using BioEdit (v7). The final assembled reads were compared to the recently published closed genome [9] using the ANI calculator tool with default settings (Rodriguez-R and Konstantinidis, 2016) using previously described methods (Goris et al., 2007; Rodriguez-R and Konstantinidis, 2014). This Whole Genome Shotgun project has been deposited at DDBJ/EMBL/GenBank under the accession LFYL01000000. The version described here is version LFYL01000000.

The genome was annotated using the RAST (Rapid Annotation using Subsystem Technology) genome annotation server (Overbeek et al., 2014). The amino acid sequence of the H-cluster domain of Cpl was used as a BLAST query against a database of the amino acid sequences encoded by the draft genome. All matches with an expect value (e-value) below 1.0 were aligned using ClustalW to determine whether they contained previously published signature motifs L1, L2, and L3 found in [FeFe]-hydrogenases.

For qRT-PCR, cultures were inoculated from a single colony from a DRC agar plate into a sealed 25-ml serum vial containing 10 ml of the same medium. Cells were grown under a headspace of 10% H\textsubscript{2}-10% CO\textsubscript{2}-80% N\textsubscript{2} by overnight incubation at 37°C. One to five ml of overnight culture was used to inoculate 50 ml of both DM-11 (Mallette et al., 1974) (N+; containing NH\textsubscript{4}Cl and (NH\textsubscript{4})\textsubscript{2}SO\textsubscript{4} and DM-11-N (N–; fixed nitrogen free) media to an OD\textsubscript{550} = 0.020 and were then sparged with 10% H\textsubscript{2}-10% CO\textsubscript{2}-80% N\textsubscript{2} for 10 min and incubated overnight. This was repeated after which point a C\textsubscript{2}H\textsubscript{2} reduction assay (Hardy et al., 1968) was performed to ensure that the N- culture contained nitrogenase activity. In sealed 120-ml serum vials, 1 ml (N+) cell culture was added to 50 ml of N+ media, and 6 ml of (N–) cell culture were added to 50 ml of N– media, to give an initial OD\textsubscript{550} = 0.020 for each culture. The vials were once again sparged with the gas mix and incubated overnight at 37°C. The following day, another C\textsubscript{2}H\textsubscript{2} reduction assay was performed to verify nitrogenase activity. Samples (500 \mu l) were mixed with 1 ml of RNAProtect Bacterial Reagent\textsuperscript{®} (Qiagen, Valencia, CA, United States) and either subjected to RNA extraction immediately or frozen at –20°C until later RNA extraction.

**Preparation of Total RNA**

Total RNA was extracted from *C. pasteurianum* strain W5 using a RNeasy\textsuperscript{®} Mini Kit (Qiagen, Valencia, CA, United States) according to the manufacturer’s protocol. For N+ conditions, the OD\textsubscript{550} of the cell culture was 0.9, while for N– cultures it was 0.4. Note that N– cultures reach stationary phase at a density of nearly half the N+ cultures. In both cases, cells were in the exponential phase at the time of harvesting. The DNase treatment step used the RQ1 RNase-Free DNase (Promega, Madison, WI, United States) and a re-purification using the RNeasy\textsuperscript{®} Mini Kit. The RNA concentration was determined using Qubit\textsuperscript{®} RNA Assay Kit (Life Technologies, Carlsbad, CA, United States), and the RNA solution was stored at –20°C until further analysis.

**Integrated DNA Technologies SciTools qPCR online primer designing software was used to design primers for the four hydrogenases (Cpl, CplII, CplIII and the large subunit of the [NiFe]-hydrogenase, Supplementary Table 1). The nitrogenase α-subunit, nifD, and the 16S rRNA small subunit (Supplementary Table 1), served as controls, to which expression was normalized. qRT-PCR was performed on a Rotor-Gene-Q real-time PCR detection system (Qiagen, Valencia, CA, United States) using the Power SYBR\textsuperscript{®} green RNA-to-C\textsuperscript{TM} 1-Step Kit (Life Technologies, Carlsbad, CA, United States) according to the specified protocol. Briefly, each reaction consisted of 10 \mu l Power SYBR\textsuperscript{®} Green PCR master mix, 100 nM each of the forward and reverse primers, 100 ng of RNA template, and nuclease-free H\textsubscript{2}O to a final volume of 20 \mu l. Cycling conditions were 40 min at 48°C, 10 min at 95°C, and then 40 cycles of 15 s at 95°C, and then 1 min at 60°C. Reactions were performed in triplicate with control reaction mixtures containing no reverse transcriptase. Each transcriptional experiment was repeated a minimum of three times using RNA isolated from separate cultures. Primer efficiencies for DNA were 0.98 for 16S rRNA gene, 0.97 for the Cpl gene, 0.92 for the CplII gene, 0.84 for the CplIII gene, and 0.88 for nifD, using a DNA concentration of 145 ng \mu l\textsuperscript{-1} to 0.145 ng \mu l\textsuperscript{-1}.

**Bioinformatics**

Homologs of Cpl were compiled from the Integrated Microbial Genomes (IMG) database (Markowitz et al., 2012) using BLASTp, resulting in 829 protein sequences. Cpl and CplII, along with 829 homologs, were subjected to a multiple sequence alignment (MSA) using the Muscle algorithm as implemented in MEGA (vers. 6) (Tamura et al., 2013) with default settings.

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Residues at each aligned position were removed if they were found to be identical in both CpI and CpII. For each of the remaining residues the degree of conservation (as a percent) among the 829 homologs was calculated using the MSA. SWISS-MODEL (Schwede et al., 2003) was used to generate a homology model of CpII, based upon the structure of CpI (Peters et al., 1998). Pymol (Delano, 2002) was then used to superimpose CpII onto CpI with a structure-based alignment. Based on this superimposition, functionally important residues that differed between CpI and CpII were selected. Sites examined included the proton transfer channel (Cornish et al., 2011) and the protein sphere around the H-cluster, as well as the proximal and medial [4Fe-4S] clusters. Using this set of criteria, a total of 14 residues were identified in the Fe/S cluster regions that potentially differentiate the functionality of these enzymes.

The large subunit of [FeFe]-hydrogenase (HydA) contains an H-cluster domain containing at least ∼350 residues (Meyer, 2007; Vignais and Billard, 2007). In addition to the H cluster, hydA often encodes diverse N-terminal (F-cluster) and C-terminal (C-cluster) domains. To minimize bias in phylogenetic reconstruction, the alignment containing the 829 homologous hydrogenases, as well as CpI and CpII, was trimmed to contain just the H-cluster domain, as previously described (Mulder et al., 2010). A phylogenetic tree of the H-cluster alignment block was constructed using a maximum likelihood method, i.e., RaxML, specifying the following parameters: gamma distribution, base frequencies, and the BLOSUM62 substitution matrix (Stamatakis, 2014). The tree was rooted with Nar1 proteins from Homo sapiens (NP_036468, NP_071938) and Danio rerio (A2RRV9). The phylogenetic signal (K-statistic) associated with the distribution of the 14 individual amino acids at each of the identified alignment positions, as they are distributed on the H-cluster phylogenetic tree, was quantified using the program multiphylosignal within the Picante package (Kembel, 2010) as implemented with the base package R. The K statistic compares the observed signal in the distribution of a trait (e.g., particular amino acid usage at a specific alignment position) on a phylogeny to the signal under a Brownian motion model of evolution (Blomberg et al., 2003). Values of K that are close to 1 imply a Brownian motion for the evolution of that trait (or some degree of phylogenetic signal) while values greater than 1 indicate strong phylogenetic signal. K values closer to zero or which are negative correspond to a random or convergent pattern of evolution for that trait.

Based on the H-cluster phylogenetic tree constructed above, 39 hydrogenase homologs that grouped phylogenetically with CpI and 39 homologous hydrogenases that grouped with CpII were identified. These 78 hydrogenases along with CpI and CpII were aligned and subjected to phylogenetic reconstruction as described above. The F- and C-cluster domains of the hydrogenases were identified using BLASTp against the Conserved Domain Database (CDD) and the CDSEARCH/cdd v3.13 algorithm (Marchler-Bauer et al., 2015) (version 3.13) using an e-value of 0.01 as previously described (Calusinska et al., 2010). These CpI- and CpII-like hydrogenases were categorized into modular structures based on the presence of identified F- and C- clusters as described previously (Calusinska et al., 2010). The modular structure was then overlaid onto the respective tree to determine the extent to which phylogeny predicts the distribution of F- and C-clusters. The genomes of organisms with the previously identified 78 hydrogenase homologs were screened for NiFH using BLASTp and the NiFH sequence from Cp as a search query. The distribution of niFH in the genomes of the respective taxa was also mapped onto the respective phylogenetic trees (Supplementary Figures 2, 3). Interactive Tree Of Life (ITOL) was used to project the phylogenetic trees (Letunic and Bork, 2007).

**AUTHOR CONTRIBUTIONS**

JT performed Clostridial growth and qRT-PCR experiments, JA and SP carried out bioinformatics studies. JT and JA wrote the document. EB, TH, and JP contributed to experimental design and manuscript revision. ZL, SN, and DB sequenced the genome. PK and MA contributed to data interpretation. All authors read and approved the final manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2017.01305/full#supplementary-material

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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