



# Energy Conservation via Hydrogen Cycling in the Methanogenic Archaeon *Methanosarcina barkeri*

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**ABSTRACT** Energy conservation via hydrogen cycling, which generates proton motive force by intracellular H<sub>2</sub> production coupled to extracellular consumption, has been controversial since it was first proposed in 1981. It was hypothesized that the methanogenic archaeon *Methanosarcina barkeri* is capable of energy conservation via H<sub>2</sub> cycling, based on genetic data that suggest that H<sub>2</sub> is a preferred, but non-essential, intermediate in the electron transport chain of this organism. Here, we characterize a series of hydrogenase mutants to provide direct evidence of H<sub>2</sub> cycling. *M. barkeri* produces H<sub>2</sub> during growth on methanol, a phenotype that is lost upon mutation of the cytoplasmic hydrogenase encoded by *frhADGB*, although low levels of H<sub>2</sub>, attributable to the Ech hydrogenase, accumulate during stationary phase. In contrast, mutations that conditionally inactivate the extracellular Vht hydrogenase are lethal when expression of the *vhtGACD* operon is repressed. Under these conditions, H<sub>2</sub> accumulates, with concomitant cessation of methane production and subsequent cell lysis, suggesting that the inability to recapture extracellular H<sub>2</sub> is responsible for the lethal phenotype. Consistent with this interpretation, double mutants that lack both Vht and Frh are viable. Thus, when intracellular hydrogen production is abrogated, loss of extracellular H<sub>2</sub> consumption is no longer lethal. The common occurrence of both intracellular and extracellular hydrogenases in anaerobic microorganisms suggests that this unusual mechanism of energy conservation may be widespread in nature.

**IMPORTANCE** ATP is required by all living organisms to facilitate essential endergonic reactions required for growth and maintenance. Although synthesis of ATP by substrate-level phosphorylation is widespread and significant, most ATP is made via the enzyme ATP synthase, which is energized by transmembrane chemiosmotic gradients. Therefore, establishing this gradient across the membrane is of central importance to sustaining life. Experimental validation of H<sub>2</sub> cycling adds to a short list of mechanisms for generating a transmembrane electrochemical gradient that is likely to be widespread, especially among anaerobic microorganisms.

**KEYWORDS** *Methanosarcina*, energy conservation, hydrogenase, methanogenesis

An essential requirement for life is the ability to couple exergonic metabolism to the endergonic synthesis of ATP. While some ATP is made by direct phosphorylation of ADP using “high-energy” metabolites such as phosphoenolpyruvate or 1,3-diphosphoglycerate, the vast majority is produced via the enzyme ATP synthase using energy stored in a transmembrane proton (or sodium) gradient. These electrochemical gradients are typically established during the process of electron transport by membrane proteins that couple exergonic redox reactions to generation of an ion motive force by one of three general mechanisms: (i) vectorial proton pumping; (ii) scalar movement of protons across the membrane, as in the Q-cycle or Q-loop; or (iii) coupled

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reactions that consume protons within the cell and produce protons on the outside (1, 2). Given the importance of this process, it is not surprising that this central aspect of living systems has been the subject of intense study (and at least three Nobel Prizes). Indeed, we now possess a detailed, molecular-level understanding of chemiosmotic energy conservation as it applies to photosynthesis and aerobic respiration in a wide variety of organisms, including eukaryotes, bacteria, and archaea. Nevertheless, unique and sometimes surprising mechanisms for generation of chemiosmotic gradients continue to be found, including sodium-pumping methyltransferases in methanogenic archaea (3), electrogenic formate:oxalate antiporters in bacteria (4, 5), and light-driven, proton-pumping rhodopsins (6).

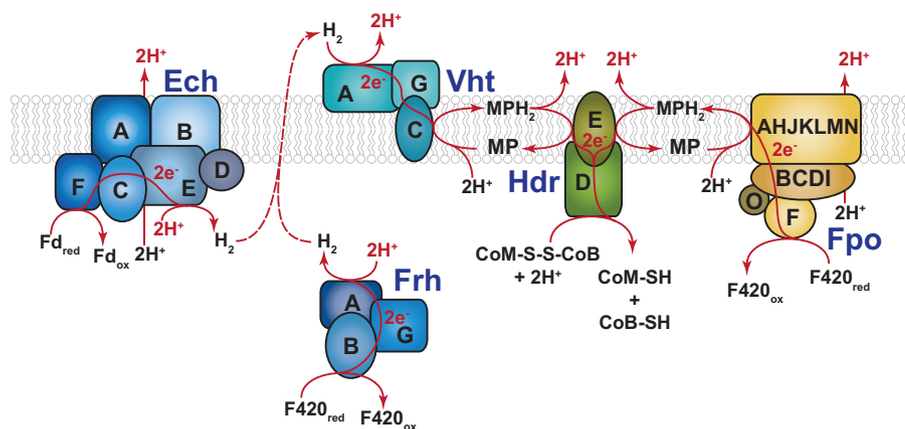
A controversial, and as yet unproven, mechanism for creating transmembrane proton gradients called H<sub>2</sub> cycling was proposed by Odom and Peck in 1981 to explain ATP synthesis in sulfate-reducing bacteria (7). In this proposed energy-conserving process, protons in the cytosol are reduced to molecular H<sub>2</sub> by enzymes known as hydrogenases. The H<sub>2</sub> so produced then diffuses across the membrane where it is reoxidized by extracellular hydrogenases, releasing protons that contribute to a transmembrane proton gradient that can be used to make ATP. The electrons produced by this reaction are returned to the cytoplasm via a membrane-bound electron transport chain, completing the redox process.

Although H<sub>2</sub> cycling has been suggested to occur in a number of anaerobic organisms (7–11), the hydrogen cycling hypothesis has not been widely accepted. A key argument against the idea is based on the high diffusion rate of molecular hydrogen. Thus, unless extracellular recapture is exceptionally efficient, hydrogen produced in the cytoplasm would be easily lost, resulting in redox imbalance and presumably cell death. Nevertheless, experimental demonstration of simultaneous production and consumption of H<sub>2</sub> by *Desulfovibrio vulgaris* supports the model (12), as does metabolic modeling (13). However, other data are inconsistent with the idea, including the ability of hydrogenase mutants to grow on lactate (14) and the inability of high external H<sub>2</sub> pressures to inhibit substrate catabolism (15). Thus, the H<sub>2</sub> cycling model for energy conservation remains unproven.

On the basis of a series of genetic experiments, we proposed that the methanogenic archaeon *Methanosarcina barkeri* employs H<sub>2</sub> cycling during growth on one-carbon (C<sub>1</sub>) substrates and acetate (16, 17). During growth on C<sub>1</sub> compounds such as methanol, the putative cycling pathway would produce H<sub>2</sub> using the cytoplasmic F420-dependent (Frh) and energy-converting ferredoxin-dependent (Ech) hydrogenases, while H<sub>2</sub> production during growth on acetate would be mediated solely by Ech. Both pathways would converge on the methanophenazine-dependent hydrogenase (Vht), which is thought to have an active site on the outer face of the cell membrane (18), to consume extracellular H<sub>2</sub> and deliver electrons to the membrane-bound electron transport chain, where they serve to reduce the coenzyme M-coenzyme B heterodisulfide (CoM-S-S-CoB) produced during the production of methane (Fig. 1). However, these genetic studies remain incomplete because neither the role of Vht nor the production and consumption of hydrogen were examined. Here we explicitly test both, providing strong experimental support for the role of H<sub>2</sub> cycling in energy conservation in *M. barkeri*.

## RESULTS AND DISCUSSION

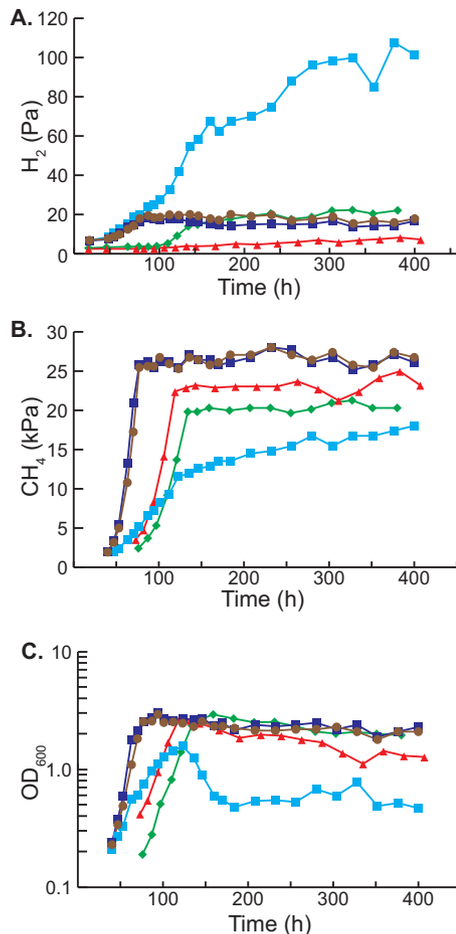
**Hydrogenases of *M. barkeri*.** Three distinct types of hydrogenases are encoded by *Methanosarcina barkeri* Fusaro (see Fig. S1 in the supplemental material) (19). The F420-reducing hydrogenase (Frh) is a cytoplasmic, three-subunit ( $\alpha$ ,  $\beta$ , and  $\gamma$ ) enzyme encoded by the *frhADGB* operon, which also includes a maturation protease, FrhD (20). This enzyme couples the oxidation/reduction of the deazaflavin cofactor F420 with production/consumption of H<sub>2</sub>. The membrane-bound Vht hydrogenase utilizes the quinone-like electron carrier, methanophenazine, as a cofactor (21). Like Frh, Vht is a three-subunit enzyme encoded by a four-gene operon (*vhtGACD*) that includes a maturation protease, VhtD (19). *M. barkeri* also contains genes that encode homologs



**FIG 1** Putative H<sub>2</sub> cycling electron transport chain of *M. barkeri*. Growth on C<sub>1</sub> substrates generates reduced cofactor F420 (F420<sub>red</sub>), which is a hydride carrying cofactor analogous to NADH, and the reduced form of the small electron-carrying protein ferredoxin (Fd<sub>red</sub>). During acetitlastic methanogenesis, only Fd<sub>red</sub> is produced. These reduced electron carriers are reoxidized in the cytoplasm by the Frh and Ech hydrogenases, respectively, with concomitant consumption of protons to produce molecular H<sub>2</sub>. H<sub>2</sub> subsequently diffuses out of the cell where it is reoxidized by the Vht hydrogenase, which has an active site located on the outer face of the cell membrane. This reaction releases protons on the outside of the cell and produces reduced methanophenazine (MPH<sub>2</sub>), a membrane-bound electron carrier analogous to ubiquinone. MPH<sub>2</sub> subsequently delivers electrons to the enzyme heterodisulfide reductase (Hdr), which serves as the terminal step in the *Methanosarcina* electron transport chain. This final reaction regenerates coenzyme B (CoB-SH) and coenzyme M (CoM-SH) from the mixed disulfide (CoM-S-S-CoB), which is produced from the free thiol cofactors during methanogenic metabolism. Electron (e<sup>-</sup>) flow and scalar protons (H<sup>+</sup>) are shown in red. It should be noted that *M. barkeri* can also reoxidize F420<sub>red</sub> using the membrane-bound, proton-pumping F420-dehydrogenase (Fpo). Thus, the cell has a branched electron transport chain, and therefore, it is not dependent on H<sub>2</sub> cycling during growth on methylotrophic substrates (16); however, both pathways for electron transport from F420 have identical levels of energy conservation: namely, 4 H<sup>+</sup>/2e<sup>-</sup>. It should also be noted that the Ech hydrogenase acts as a proton pump in addition to its role in H<sub>2</sub> cycling, thus electron transport from Fd<sub>red</sub> during methylotrophic and acetitlastic methanogenesis conserves 6H<sup>+</sup>/2e<sup>-</sup>. Individual subunits of the various enzymes are indicated by capital letters (e.g., A, B, C...).

of both the *frh* and *vht* operons (the *freAEGB* and *vhxGAC* operons, respectively); however, multiple lines of evidence suggest that these genes are incapable of producing active hydrogenases (16, 22). Thus, the presence of these genes has no bearing on the results presented herein. The final hydrogenase encoded by *M. barkeri* is a membrane-bound, energy-converting hydrogenase (Ech), which couples the oxidation/reduction of ferredoxin and H<sub>2</sub> to the production/consumption of a proton motive force (23, 24). Thus, the enzyme can use proton motive force to drive the endergonic reduction of ferredoxin by H<sub>2</sub>, which is required for CO<sub>2</sub> reduction during hydrogenotrophic methanogenesis and for biosynthesis during growth by H<sub>2</sub>-dependent reduction of C<sub>1</sub> compounds (methyl-reducing methanogenesis). During both methylotrophic and acetitlastic methanogenesis, Ech is believed to couple oxidation of reduced ferredoxin to production of proton motive force and H<sub>2</sub>. The hydrogen thus produced would need to be recaptured by Vht in a putative H<sub>2</sub> cycling process that contributes to proton motive force (Fig. 1) (17).

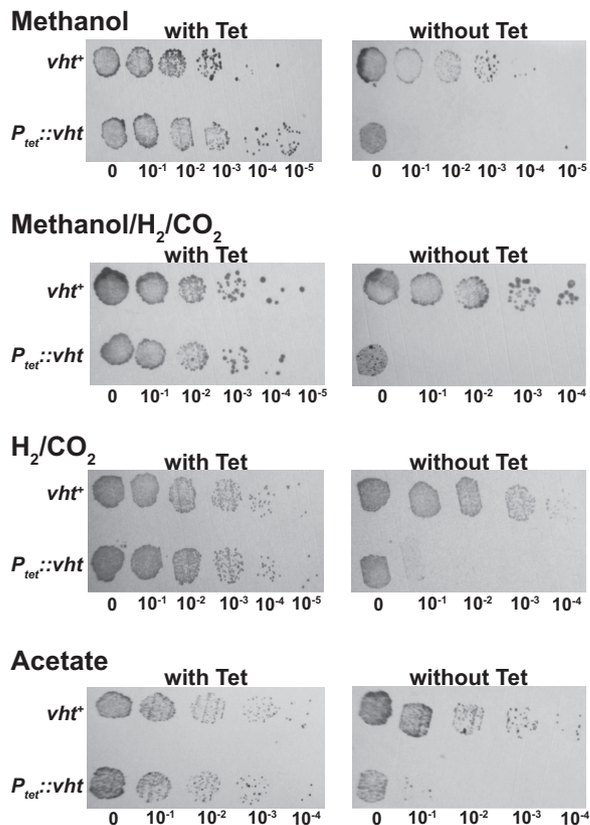
**The cytoplasmic Frh hydrogenase is responsible for production of H<sub>2</sub> during growth on methanol.** A number of studies have shown that assorted *Methanosarcina* strains produce H<sub>2</sub> during growth on methylotrophic and acetitlastic substrates (9, 25–30); however, to our knowledge, this has never been assessed in *M. barkeri* strain Fusaro. To test this, we quantified the accumulation of CH<sub>4</sub> and H<sub>2</sub> during growth on methanol-containing medium (Fig. 2). Consistent with the hydrogen cycling hypothesis, we observed significant H<sub>2</sub> production, which reached a maximum partial pressure of ca. 20 Pa near the end of exponential growth. As expected, the culture also produced substantial levels of methane. As previously observed (16), a mutant lacking Frh (strain WWM115 [Table S1]) grew at a lower rate than its isogenic parent and produced somewhat smaller amounts of methane. Very little H<sub>2</sub> (<4 Pa) was produced during



**FIG 2** Hydrogen and methane production during methylotrophic growth. (A to C) The partial pressures of H<sub>2</sub> (A) and methane (B) were monitored during the course of growth (as indicated by optical density [C]) in methanol-containing medium for various *M. barkeri* strains. Strains used were *M. barkeri* isogenic parental strain (WWM85 [brown circles]), tetracycline-regulated  $P_{tet::vht}$  mutant (WWM157) with tetracycline (dark blue squares) and without tetracycline (light blue squares),  $\Delta frh$  mutant (WWM115 [red triangles]), and  $\Delta frh \Delta vht$  double mutant (WWM351 [green diamonds]). Measurements were performed in triplicates as described in Materials and Methods. Complete strain genotypes can be found in Table S1 in the supplemental material.

growth of the  $\Delta frh$  mutant; however, after growth ceased, the H<sub>2</sub> concentration slowly rose, reaching a maximum level of 7 Pa. Thus, Frh is responsible for most hydrogen production during growth of *M. barkeri* Fusaro on methanol, although some hydrogen is still produced in the  $\Delta frh$  mutant. As will be shown below, Ech is probably responsible for the low levels of H<sub>2</sub> seen in the  $\Delta frh$  mutant.

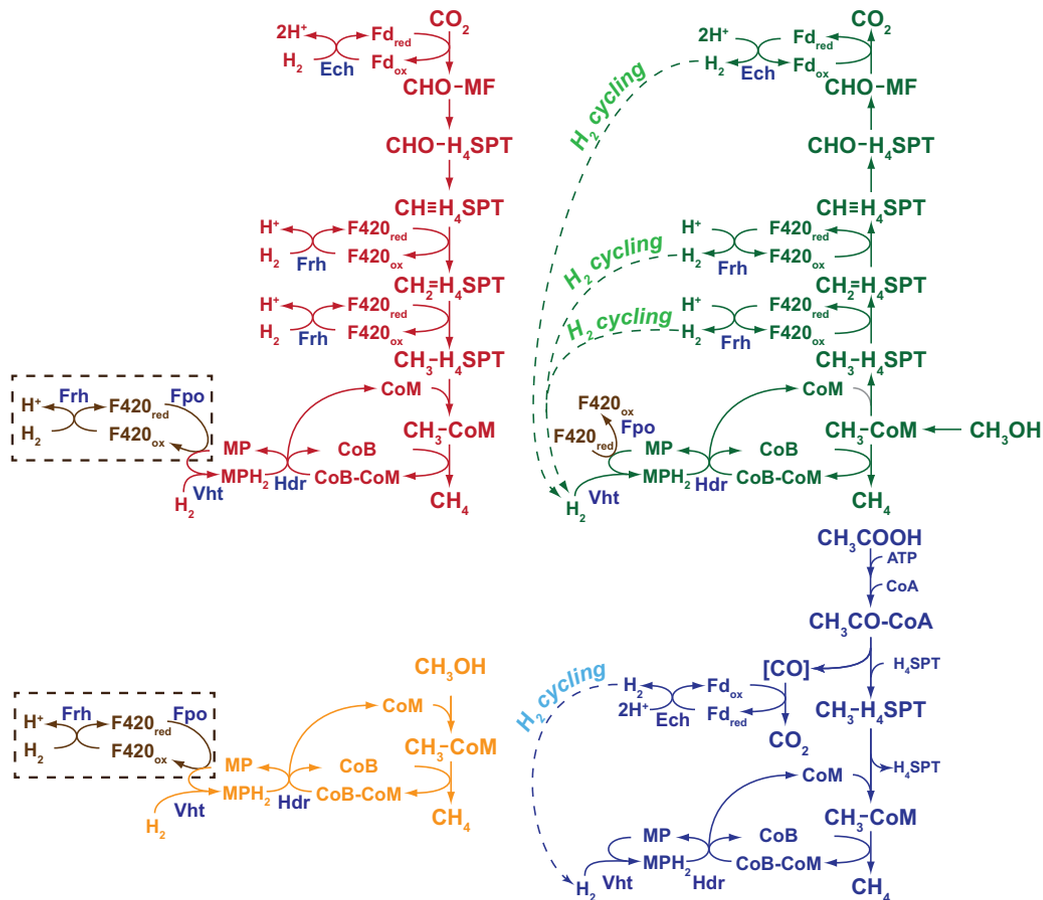
**Vht activity is required for viability of *M. barkeri*.** To investigate the role of Vht during growth of *M. barkeri*, we attempted to delete the *vhtGACD* operon via homologous gene replacement (31, 32). However, despite numerous attempts, including selection on a variety of media, with and without supplementation of potential biosynthetic intermediates, no mutant colonies were obtained. We also attempted to delete the *vht* operon using the markerless deletion method of genetic exchange (33). This method relies on construction of a merodiploid strain with both mutant and wild-type alleles. Upon segregation of the merodiploid, 50% of the recombinants are expected to be mutants if there is no selective pressure against the mutant allele. However, if the mutation causes a reduction in growth rate (with lethality being the most extreme case), the probability of obtaining recombinants with the mutant allele is severely reduced. We tested 101 haploid recombinants obtained from a *vhtGACD*<sup>+</sup>/ $\Delta vhtGACD$  merodiploid; all carried the wild-type *vht* allele. Taken together, these data suggest that the *vhtGACD* operon is critical for normal growth of *M. barkeri*.



**FIG 3** Essentiality of the Vht hydrogenase in *M. barkeri*. Cultures of the *P<sub>tet</sub>::vht* mutant (WWM157) and its isogenic parent (WWM154) were adapted to four different substrates of interest (and in the presence of tetracycline for *P<sub>tet</sub>::vht*), then washed, serially diluted, and incubated with each substrate with and without tetracycline (Tet). The media used indicate the ability to grow via each of the four known methanogenic pathways: (i) methylotrophic (methanol), (ii) methyl reduction (methanol/H<sub>2</sub>/CO<sub>2</sub>), (iii) hydrogenotrophic (H<sub>2</sub>/CO<sub>2</sub>), and (iv) aceticlastic (acetate).

To test whether Vht is essential, we constructed a mutant in which the *vht* operon was placed under control of a tightly regulated, tetracycline-dependent promoter (34). We then examined the viability of the mutant and its isogenic parent by spotting serial dilutions on a variety of media, with and without tetracycline. As shown in Fig. 3, the *P<sub>tet</sub>::vht* mutant is unable to grow in the absence of the inducer but grew well when tetracycline was added, whereas the isogenic parent grew with or without the addition of tetracycline. These phenotypes were observed on a variety of media, including media containing (i) methanol, (ii) methanol plus H<sub>2</sub>, (iii) H<sub>2</sub>/CO<sub>2</sub>, and (iv) acetate, which were chosen because they encompass growth conditions that require each of the four known methanogenic pathways used by *M. barkeri* (Fig. 4). It should be stressed that the *P<sub>tet</sub>::vht* mutant used in this experiment was pregrown in the presence of inducer. Thus, at the start of the experiment, all cells have active Vht. However, during cultivation in the absence of tetracycline, preexisting Vht is depleted by protein turnover and cell division, thereby allowing characterization of the Vht-deficient phenotype. The absence of growth of the diluted cultures in all media shows that Vht is essential for growth via the methylotrophic (methanol), methyl-reducing (methanol plus H<sub>2</sub>), hydrogenotrophic (H<sub>2</sub>/CO<sub>2</sub>), and aceticlastic (acetate) methanogenic pathways.

**Depletion of Vht results in H<sub>2</sub> accumulation and cell lysis.** To help understand why Vht is essential, we quantified production of H<sub>2</sub> and CH<sub>4</sub> in cultures of the *P<sub>tet</sub>::vht* strain with and without tetracycline (Fig. 2). When the strains were grown in methanol-containing medium in the presence of tetracycline, the accumulation of H<sub>2</sub> and CH<sub>4</sub> was essentially identical to that of the isogenic parent. Cultures in which *vht* is not expressed (i.e., without tetracycline) grew initially but growth rapidly slowed and



**FIG 4** Role of  $H_2$  cycling in the four methanogenic pathways of *M. barkeri*. *M. barkeri* utilizes four distinct methanogenic pathways to allow growth on a variety of substrates. In the hydrogenotrophic pathway (shown in red),  $CO_2$  is reduced to methane using electrons derived from  $H_2$ , while in the methyl-reducing pathway (shown in orange),  $H_2$  is used to reduce  $C_1$  compounds, such as methanol, directly to  $CH_4$ . During methylotrophic methanogenesis (shown in green),  $C_1$  compounds are disproportionated to  $CO_2$  and methane, with one molecule of the  $C_1$  compound oxidized to provide electrons for reduction of three additional molecules to methane. Finally, in the acetoclastic pathway (shown in blue), acetate is split into a methyl group and an enzyme-bound carbonyl moiety. The latter is oxidized to  $CO_2$  to provide electrons required for reduction of the methyl group to methane. The steps catalyzed by Fpo, Frh, Vht, Ech, and Hdr proteins are indicated. Steps involving  $H_2$  cycling are shown by labeled dashed arrows. An alternate,  $H_2$ -independent electron transport pathway is shown in brown. Experimental data support the function of this alternate pathway during methylotrophic methanogenesis, but not in hydrogenotrophic or methyl-reducing methanogenesis (as indicated by the dashed brown box). Abbreviations; Fpo, F420 dehydrogenase; Frh, F420-reducing hydrogenase; Vht, methanophenazine-dependent hydrogenase; Ech, energy-converting ferredoxin-dependent hydrogenase; Hdr, heterodisulfide reductase; CoM, coenzyme M; CoB, coenzyme B; CoB-CoM, mixed disulfide of CoB and CoM; MP/MPH<sub>2</sub>, oxidized and reduced methanophenazine; F420<sub>ox</sub>/F420<sub>red</sub>, oxidized and reduced cofactor F420, respectively; Fd<sub>ox</sub>/Fd<sub>red</sub>, oxidized and reduced ferredoxin, respectively; CHO-MF, formyl-methanofuran; H<sub>4</sub>SPT, tetrahydrosarcinapterin; CHO-H<sub>4</sub>SPT, formyl-H<sub>4</sub>SPT; CH=H<sub>4</sub>SPT, methenyl-H<sub>4</sub>SPT; CH<sub>2</sub>=H<sub>4</sub>SPT, methylene-H<sub>4</sub>SPT; CH<sub>3</sub>-H<sub>4</sub>SPT, methyl-H<sub>4</sub>SPT; CH<sub>3</sub>-CoM, methyl-coenzyme M; CoA, coenzyme A; CH<sub>3</sub>CO-CoA, acetyl-coenzyme A; [CO], enzyme-bound carbonyl moiety.

reached an optical density that was less than half of that obtained when *vht* was expressed. The optical density subsequently dropped, suggesting cell death and lysis. Similarly, methane accumulation in cultures not expressing *vht* was much slower than in induced cultures and only reached half of that seen under inducing conditions. In contrast,  $H_2$  accumulation was much higher in the absence of Vht, with final levels nearly sixfold higher than those seen in cultures that express Vht. These data clearly show that Vht is required for efficient recapture of  $H_2$  produced by Frh and Ech. Moreover, they suggest that  $H_2$  loss is responsible for the lethal consequences of *vht* repression.

**Vht is not essential in  $\Delta frh$  mutants.** If the inability to recapture  $H_2$  is responsible for the essentiality of Vht, then it should be possible to delete the *vht* operon in strains

that do not produce hydrogen. As described above, Frh is responsible for the majority of H<sub>2</sub> production during growth. Thus, we attempted to introduce a  $\Delta vht$  allele into the  $\Delta frh$  host. In contrast to our prior unsuccessful attempts to create a  $\Delta vht$  single mutant, the  $\Delta vht \Delta frh$  double mutant was isolated in the first attempt. Therefore, Vht is not required when Frh is absent. Like the  $\Delta frh$  single mutant, the  $\Delta vht \Delta frh$  double mutant grows slowly on methanol and produces lower levels of methane (Fig. 2). Significantly, the double mutant does not produce the excessive level of H<sub>2</sub> seen in the uninduced P<sub>tet</sub>::*vht* strain, instead accumulating H<sub>2</sub> at levels similar to those of the parental strain (ca. 20 Pa). Because Ech is the only active hydrogenase remaining in the  $\Delta vht \Delta frh$  double mutant, it must be responsible for H<sub>2</sub> production in this strain. This begs the question of why H<sub>2</sub> accumulation stops at 20 Pa in the double mutant, while the uninduced P<sub>tet</sub>::*vht* strain produces much higher levels. We suggest that the coupling of Ech activity to generation of proton motive force thermodynamically restrains excessive H<sub>2</sub> production, even in the absence of H<sub>2</sub> uptake by Vht. This would also explain the viability of the  $\Delta vht \Delta frh$  double mutant. This situation is in stark contrast to that seen in the *vht*-depleted strain, where the F420-dependent Frh is responsible for most of the H<sub>2</sub> production (see above). Accordingly, at the low H<sub>2</sub> partial pressures observed in our experiments, reduction of protons with F420 is strongly exergonic, allowing excessive hydrogen accumulation. This is also consistent with the observation that the redox state of F420 is in rapid equilibrium with H<sub>2</sub> (35). Interestingly, the smaller amount of H<sub>2</sub> accumulation in the  $\Delta frh$  mutant relative to that seen in the  $\Delta vht \Delta frh$  double mutant shows that Vht also consumes H<sub>2</sub> produced by Ech. This supports previous studies indicating potential energy conservation via Ech/Vht H<sub>2</sub> cycling during acetate metabolism (17, 23).

***M. barkeri* has a bifurcated electron transport chain with H<sub>2</sub>-dependent and -independent branches.** We previously showed that *M. barkeri* has a branched electron transport chain, with Frh- and F420 dehydrogenase (Fpo)-dependent branches (16). The data reported here extend our understanding of the Frh-dependent branch and are fully consistent with the model depicted in Fig. 1. Thus, during growth on methylophilic substrates such as methanol, reduced F420 is preferentially oxidized via an energy-conserving, H<sub>2</sub> cycling electron transport chain that requires Frh. However, in the absence of Frh, reduced F420 is channeled into the Fpo-dependent electron transport chain, which supports growth at a significantly lower rate (Fig. 1 and 2). This alternate pathway accounts for the viability of the  $\Delta frh$  mutant, which is lost when both *frh* and *fpo* are deleted (16). Similar but less severe phenotypes have been observed in *fpo* and *frh* mutants of *Methanosarcina mazei*, thus it seems likely that H<sub>2</sub> cycling also occurs in this closely related species (36). However, many *Methanosarcina* species, especially those that inhabit marine environments, are devoid of hydrogenase activity, despite the presence of hydrogenase-encoding genes. We, and others, have interpreted this to be an adaptation to the marine environment, where H<sub>2</sub>-utilizing sulfate reducers are likely to disrupt H<sub>2</sub> cycling due to the superior thermodynamics of H<sub>2</sub> oxidation coupled to sulfate reduction (19, 37).

A similar branched electron transport chain may also explain the contradictory evidence regarding H<sub>2</sub> cycling in *Desulfovibrio* species. Thus, the viability of *Desulfovibrio* hydrogenase mutants and the inability of excess H<sub>2</sub> to suppress substrate catabolism can both be explained by the presence of alternative electron transport mechanisms. Indeed, metabolic modeling of *Desulfovibrio vulgaris* strongly supports this interpretation (13). Thus, it is critical that experiments designed to test the H<sub>2</sub> cycling mechanism be interpreted within a framework that includes the possibility of branched electron transport chains. With this in mind, it seems likely that many anaerobic organisms might use H<sub>2</sub> cycling for energy conservation. Indeed, since it was originally proposed, H<sub>2</sub> cycling has been suggested to occur in the acetogen *Acetobacterium woodii* (10) and in the Fe(III) respiring *Geobacter sulfurreducens* (8).

**Why are Vht mutants inviable during growth on methanol/H<sub>2</sub> or H<sub>2</sub>/CO<sub>2</sub>?** Although the data presented here strongly support the H<sub>2</sub> cycling model, they raise additional questions regarding H<sub>2</sub>-dependent methanogenesis that are not easily

explained. In particular, it is not readily apparent why the uninduced  $P_{tet}::vht$  mutants are inviable during hydrogenotrophic or methyl-reducing growth. As shown in Fig. 4, it should be possible to channel electrons from  $H_2$  oxidation into the electron transport chain via Frh and Fpo. Indeed, Thauer et al. have proposed that this alternate pathway is functional in *Methanosarcina* (38). Nevertheless, the  $P_{tet}::vht$  mutant does not grow under repressing conditions on either  $H_2/CO_2$  or methanol plus  $H_2$ . It should be stressed that we use high concentrations of hydrogen during growth on these substrates. Thus, it is expected that reduction of F420 via Frh should be exergonic in our experiments, which would favor this pathway. (This is in contrast to the methylotrophic or acetoclastic growth conditions described above, under which the reverse reaction [i.e., hydrogen production] is favored.) Thus, a thermodynamic argument cannot easily explain the results. Further, based on available evidence (16, 39, 40), energy conservation via the Vht-dependent pathway should be identical to that of the alternate Frh/Fpo-dependent pathway. Thus, an energy conservation argument also cannot explain the phenomenon. One might argue that faster kinetics of the Vht-dependent pathway could be responsible, but in our opinion, the growth (albeit slower than wild type) of the  $\Delta frh$  and  $\Delta vht \Delta frh$  mutants during methylotrophic growth, which depends on Fpo, argues against this explanation. Therefore, as yet unknown regulatory and/or biochemical constraints on hydrogen metabolism in *Methanosarcina* await discovery.

## MATERIALS AND METHODS

**Strains, media, and growth conditions.** The construction and genotypes of all *Methanosarcina* strains are presented in Table S1 in the supplemental material. *Methanosarcina* strains were grown as single cells (41) at 37°C in high-salt (HS) broth medium (42) or on agar-solidified medium as described previously (43). Growth substrates provided were methanol (125 mM in broth medium and 50 mM in agar-solidified medium) or sodium acetate (120 mM) under a headspace of either  $N_2/CO_2$  (80/20%) at 50 kPa over ambient pressure or  $H_2/CO_2$  (80/20%) at 300 kPa over ambient pressure. Cultures were supplemented as indicated with 0.1% yeast extract, 0.1% Casamino Acids, 10 mM sodium acetate, or 10 mM pyruvate. Puromycin (CalBioChem, San Diego, CA) was added at 2  $\mu$ g/ml for selection of the puromycin transacetylase (*pac*) gene (33). 8-Aza-2,6-diaminopurine (8-ADP) (Sigma, St. Louis, MO) was added at 20  $\mu$ g/ml for selection against the presence of *hpt* (33). Tetracycline was added at 100  $\mu$ g/ml to induce the tetracycline-regulated *PmcrB(tetO3)* promoter (34). Standard conditions were used for growth of *Escherichia coli* strains (44) DH5 $\alpha/\lambda$ -*pir* (45) and DH10B (Stratagene, La Jolla, CA), which were used as hosts for plasmid constructions.

**DNA methods and plasmid construction.** Standard methods were used for plasmid DNA isolation and manipulation using *E. coli* hosts (46). Liposome-mediated transformation was used for *Methanosarcina* as described previously (47). Genomic DNA isolation and DNA hybridization were performed as described previously (32, 42, 43). DNA sequences were determined from double-stranded templates by the W. M. Keck Center for Comparative and Functional Genomics, University of Illinois. Plasmid constructions are described in the supporting information (Tables S2 and S3).

**Construction of the  $\Delta frh$  and  $\Delta vht \Delta frh$  mutants.** The markerless genetic exchange method (33) using plasmid pGK4 was employed to delete *frhADGB* ( $\Delta frh$ ) in the  $\Delta hpt$  background of *M. barkeri* Fusaro (Tables S1, S2, and S3) using methanol/ $H_2/CO_2$  as the growth substrate. The  $\Delta vht \Delta frh$  mutant was constructed by deleting *vhtGACD* in the  $\Delta frh$  markerless mutant by the homologous recombination-mediated gene replacement method (32). To do this, the 5.6-kb XhoI/NotI fragment of pGK82B was used to transform the  $\Delta frh$  mutant to puromycin resistance on methanol-containing medium. The mutants were confirmed by PCR and DNA hybridization (data not shown).

**Construction of the tetracycline-regulated *vht* mutant ( $P_{tet}::vht$ ).** The tetracycline-regulated *PmcrB(tetO3)* promoter was employed to drive conditional expression of the *vht* operon in *M. barkeri* WWM157 (34). This strain was constructed by transforming strain WWM154 to puromycin resistance using the 7-kb NcoI/SpeI fragment of pGK61A (Tables S1, S2, and S3). The transformants were selected on methanol plus  $H_2/CO_2$  medium in the presence of puromycin and tetracycline. The  $P_{tet}::vht$  strain was confirmed by DNA hybridization (data not shown). To ensure that the native *vht* promoter (*Pvht*) did not interfere with expression from *PmcrB(tetO3)*, 382 bp upstream of *vhtG* were deleted in  $P_{tet}::vht$ . This left 1,038 bp intact for the expression of the *hyp* operon, which is upstream of the *vht* operon and expressed in the opposite direction.

**Determination of Vht essentiality during growth on all substrate types.** Growth of strains WWM157 ( $P_{tet}::vht$ ) and WWM154 (isogenic parent) on methanol, methanol/ $H_2/CO_2$ ,  $H_2/CO_2$ , and acetate were analyzed by the spot-plate method (48). Cultures were first adapted for at least 15 generations to the substrate of interest; tetracycline was added to each medium for growth of strain WWM157. Upon reaching stationary phase, 10 ml of culture was washed three times and resuspended in 5 ml HS medium that lacked growth substrate. Subsequently, 10  $\mu$ l of 10-fold serial dilutions was spotted onto the following: three layers of GB004 paper (Whatman, NJ), two layers of GB002 paper (Schleicher & Schuell BioScience, NH), one layer of 3 MM paper (Whatman, NJ), and a 0.22 mM nylon membrane (GE Water and Process Technologies, PA) soaked in 43 ml of HS medium containing the substrate of interest with and

without tetracycline. The plates were sealed and incubated at 37°C for at least 2 weeks in an intrachamber anoxic incubator (49). Growth on acetate and methanol was tested under an atmosphere of N<sub>2</sub>/CO<sub>2</sub>/H<sub>2</sub>S (80/19.9/0.1 ratio), while growth on methanol/H<sub>2</sub>/CO<sub>2</sub> or H<sub>2</sub>/CO<sub>2</sub> was tested under an atmosphere of H<sub>2</sub>/CO<sub>2</sub>/H<sub>2</sub>S (80/19.9/0.1 ratio).

**Measurement of H<sub>2</sub>, CH<sub>4</sub>, and OD<sub>600</sub> during growth on methanol.** *M. barkeri* WWM85 (isogenic parent), WWM157 (*P<sub>ret</sub>::vht*; grown in the presence of tetracycline), WWM115 (*Δfrh*), and WWM351 (*Δvht Δfrh*) were grown on methanol until mid-exponential phase (optical density at 600 nm [OD<sub>600</sub>] of ca. 0.5) and then 1 ml (WWM85 and WWM157) or 5 ml (WWM115 and WWM351) was inoculated into 100 ml HS-methanol in a 500-ml serum bottle. For WWM157, the culture was washed once prior to inoculation with or without tetracycline. To measure H<sub>2</sub> and CH<sub>4</sub>, ca. 1-ml or 2-ml headspace sample was withdrawn aseptically from the culture at various time points with a syringe that had been flushed with sterile, anaerobic N<sub>2</sub>. The gas sample was then diluted into 70 ml helium. A gas-tight syringe flushed with helium was subsequently used to withdraw 3 ml of the diluted sample, which was then injected into an SRI gas chromatograph, equipped with a reduction gas detector (RGD) and a thermal conductivity detector (TCD) at 52°C. The RGD column was a three-foot-long 13× molecule sieve, whereas the TCD column was a six-foot HayeSep D porous polymer column. The RGD column was used to detect H<sub>2</sub> by peak height, and the TCD column was used to detect CH<sub>4</sub> by peak area. Helium was used as the carrier gas. OD<sub>600</sub> was also measured during the growth curve.

## SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <https://doi.org/10.1128/mBio.01256-18>.

**FIG S1**, PDF file, 0.7 MB.

**TABLE S1**, DOC file, 0.04 MB.

**TABLE S2**, DOC file, 0.05 MB.

**TABLE S3**, DOC file, 0.03 MB.

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