An Ancient Pathway Combining Carbon Dioxide Fixation with the Generation and Utilization of a Sodium Ion Gradient for ATP Synthesis

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Abstract

Synthesis of acetate from carbon dioxide and molecular hydrogen is considered to be the first carbon assimilation pathway on earth. It combines carbon dioxide fixation into acetyl-CoA with the production of ATP *via* an energized cell membrane. How the pathway is coupled with the net synthesis of ATP has been an enigma. The anaerobic, acetogenic bacterium *Acetobacterium woodii* uses an ancient version of this pathway without cytochromes and quinones. It generates a sodium ion potential across the cell membrane by the sodium-motive ferredoxin:NAD oxidoreductase (Rnf). The genome sequence of *A. woodii* solves the enigma: it uncovers Rnf as the only ion-motive enzyme coupled to the pathway and unravels a metabolism designed to produce reduced ferredoxin and overcome energetic barriers by virtue of electron-bifurcating, soluble enzymes.

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Introduction

The atmosphere in the early days of our planet was highly reducing and did not contain oxygen but gases such as molecular hydrogen and carbon dioxide. How life evolved under these conditions is a matter of debate but some researchers favour a pathway that combines two essential features: carbon fixation into organic molecules and, at the same time, generation of ATP [1]. The Wood-Ljungdahl pathway of carbon dioxide fixation is such a pathway and, therefore, could be considered to have evolved on earth very early [2,3]. It is present in strictly anaerobic sulfate reducing bacteria and archaea, in methanogenic archaea and acetogenic bacteria [4]. Only in the latter it fulfills the dual function mentioned which is growth on molecular hydrogen and carbon dioxide according to:

$$4 H_2 + 2 CO_2 \rightarrow CH_3 COO^- + H^+ + 2 H_2 O$$

$$\Delta G^{o'} = -95 \text{ kJ/mol}$$
(1)

The pathway involves reduction of carbon dioxide to formate, binding of formate to the cofactor tetrahydrofolate (THF) and subsequent reduction of the formyl group to methyl tetrahydrofolate. Another mol of carbon dioxide is reduced to enzymetrapped carbon monoxide that is then condensed with a methyl group and coenzyme A to yield acetyl- CoA (Fig. 1). In the anabolic route, acetyl-CoA is the precursor molecule for biosynthetic reactions and in the catabolic route the precursor of acetate [5,6,7]. Acetyl-CoA allows for the synthesis of one mol ATP via substrate level phosphorylation. However, this ATP is required for formate activation to formyltetrahydrofolate leaving no ATP. Therefore, acetogenesis from 4 H₂+2 CO₂ must yield additional ATP by a mechanism other than substrate level phosphorylation. How this is achieved is unknown. The amount of energy gained from equation 1 would theoretically allow the generation of only 1.5 mol of ATP per mol acetate produced [8]. This calculation is based on a partial pressure of molecular hydrogen of 1 bar. However, at the low hydrogen partial pressures observed in the environment in which the acetogenic bacteria thrive the free energy change associated with reaction 1 is much more positive and sufficient to generate only a fraction of an ATP [9]. Therefore, acetogens clearly live at the thermodynamic limit of life.

Acetogens can be divided in two groups with respect to their energy metabolism [10]. One group contains cytochromes and quinones that are considered to be involved in a membrane-bound electron transport process that generates a proton potential across the membrane; they use a proton potential-driven ATP synthase. This can be considered as the evolutionary more advanced version of the pathway since it required the evolution of cytochromes [11].

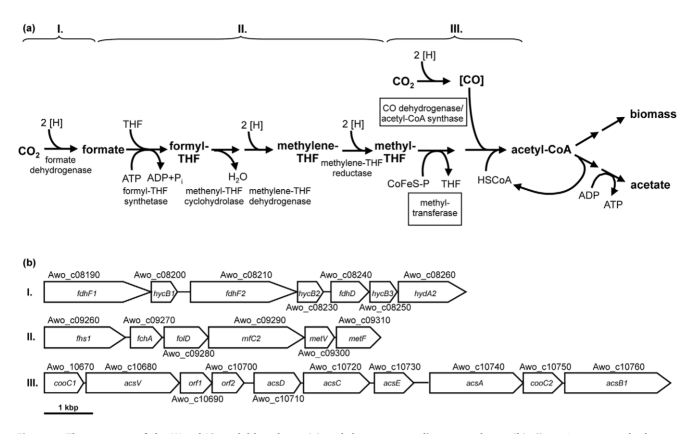


Figure 1. The enzymes of the Wood-Ljungdahl pathway (a) and the corresponding gene cluster (b). Cluster I: two genes for formate dehydrogenase (*fdhF*) are organized together with three copies of a FeS-containing subunit of a [FeFe]-hydrogenase (*hycB*), a formate dehydrogenase accessory protein (*fdhD*) and another subunit of the [FeFe]-hydrogenase (*hydA2*). Cluster II: genes for formyl-THF synthetase (*fhs1*), methenyl-THF cyclohydrolase (*fchA*), methylene-THF dehydrogenase (*folD*) and methylene-THF reductase (*metF*, *metV*) are organized together with a RnfC-similar protein (*rnfC2*). Cluster III: genes for the subunits of the CO dehydrogenase/acetyl CoA synthase complex consisting of CO dehydrogenase (*acsA*), acetyl-CoA synthase (*acsB1*), corrinoid-iron sulfur protein (*acsCD*) and methyltransferase (*acsV*) are organized together with two copies of a CODH nickel-insertion accessory protein (*cooC*), a corrinoid activation/regeneration protein (*acsV*) and two hypothetical proteins (Orf1 and Orf2). doi:10.1371/journal.pone.0033439.g001

The second group, to which the Gram positive bacterium Acetobacterium woodii belongs, represents a more ancient version of the pathway. It does not involve cytochromes and quinones but at least one ion pump, a sodium ion-translocating ferredoxin:NAD⁺ oxidoreductase (Rnf) [12]. The Rnf complex is a membranebound electron transfer complex containing iron-sulfur centers and flavins. Exergonic electron transfer from reduced ferredoxin (Fd²⁻) to NAD⁺ is found to be coupled with vectorial, electrogenic Na⁺ transport in inverted membrane vesicles of A. woodii [13]. Immunological studies revealed that the Rnf complex is present in cells grown on H₂+CO₂, underlining its importance in the bioenergetics of the Wood-Ljungdahl pathway [14]. The complex is also present in cells grown on formate, methanol, betaine or fructose and its synthesis was also independent of the electron acceptor used and, therefore, is constitutive [14]. The sodium ion potential across the membrane drives the synthesis of ATP via a membrane-integral Na⁺ F₁F_O ATP synthase [15,16].

Another argument that this version of the pathway has evolved early is that it uses Na⁺ as coupling ion. Membranes are more tight for Na⁺ than for H⁺ [17] which is essential for organisms living in ecosystems that have a high concentration of organic acids that function as "proton ferries" allowing protons to backenter the cell and thus destroying part of the already very little power available to fuel the ATP synthase. Therefore, sodium bioenergetics using a Δ pNa is considered an early step in the evolution of cellular bioenergetics [18]. A. woodii employs the Wood-Ljungdahl pathway without cytochromes and is one of the very few organisms known to completely rely on a sodium ion potential for energetic reactions [10]. It grows well in the absence of a proton potential across its membrane, and apparently uses a primary Na⁺ potential to fuel ATP synthesis, transport processes and flagellar rotation [16]. Thus it is a prime candidate to unravel the bioenergetics of what may have been one of the first pathways in coupling exergonic and endergonic reactions in life. Therefore, we sequenced the entire genome of A. woodii and performed a few key experiments to fill the gaps in understanding how the pathway is coupled to the synthesis of ATP.

Results and Discussion

Genetic organization and predicted properties of enzymes of the Wood-Ljungdahl pathway

General genome features of *A. woodii* are listed in Table 1. The genome of *A. woodii* consists of a single circular 4,044,785 bp chromosome with a GC content of 39.3%. 3473 protein-encoding ORFs, 5 rRNA clusters and 60 tRNA genes account for 85.1% of the genomic DNA. 71.1% of the predicted ORFs are with assigned functions (NCBI database, accession number CP002987). The genes encoding proteins of the Wood-Ljungdahl pathway are mainly found in three clusters on the chromosome (Fig. 1).

In cluster I, two genes, each encoding a formate dehydrogenase (fdhF), are present. The formate dehydrogenase catalyzes the first

Table 1. General features of the A. woodii genome.

	A. woodii	
	A. Woodii	
Genome size (Mbp)	4.044785	
protein encoding orfs	3472	
percent coding (%)	85.1	
G+C content (mol%)	39.34	
rRNA cluster	5	
tRNA	59	
orfs with assigned functions	2469	
orfs without assigned functions	1004	

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step in the pathway, the conversion of CO₂ to formate. FdhF1 and FdhF2 are 79% identical and FdhF2 seems to be a selenocysteincontaining protein. Apparently, it is the only selenocysteincontaining protein encoded on the genome of A. woodii. The genes encoding selenocystein biosynthesis proteins are next to the cluster upstream of fdhF1. Furthermore, there are genes present that potentially encode a formate dehydrogenase accessory protein (fdhD), three copies of a putative FeS-containing electron transfer protein (hycB) and a subunit (hydA2) harbouring the active site characteristic of [FeFe]-hydrogenase. HycB1 and HycB2 are 82% identical and the encoding genes are downstream of *fdhF1* and fdhF2, respectively. Therefore, we suggest that HycB1 and HycB2 are electron transfer proteins specific for FdhF1 and FdhF2, respectively. HycB3 is only 34 and 33% identical to HycB1 and HycB2, respectively, and its localization in the gene cluster indicates that it is the electron output module for the hydrogenase HydA2.

Cluster II encodes the enzymes catalyzing the reduction of the formyl-group to the methyl-group, the formyl-tetrahydrofolate (THF) synthetase, methenyl-THF cyclohydrolase, methylene-THF dehydrogenase and methylene-THF reductase. The genetic organization is different from Moorella thermoacetica where these genes are spread all over the genome [19]. A second formyl-THF synthetase is found in A. woodii (Awo_c08040), whereas in M. thermoacetica and Clostridium ljungdahlii only one such enzyme is encoded. Interestingly, metF, the gene encoding the methylene-THF reductase is preceded in A. woodii by two genes (Awo_c09290, Awo_c09300) of unknown function. Homologues of Awo_c09300 are also found together with metF in many acetogens such as M. thermoacetica and C. ljungdahlii, but also in other bacteria and archaea like Alkaliphilus metalliredigens, Geobacter sulfurreducens or Methanosarcina acetivorans. In contrast, homologues of Awo c09290 are only found in this region in Ruminococcus obeum, Blautia hydrogenotrophica and Bryantella formatexigens, and three members of the genus *Clostridium*. The possible function of these proteins is discussed below.

Acetyl-CoA is formed from methyl-THF and a second CO₂ by the CO dehydrogenase/acetyl-CoA synthase complex. Genes coding for the subunits (*acsAB*) are found in cluster III (Awo_c10670-Awo_c10760) together with the genes coding for the corrinoid-iron sulfur protein (*acsCD*) and the methyltransferase (*acsE*). Additionally there are two genes in the cluster encoding for a CODH nickel-insertion accessory protein (*cooC*) and a corrinoid activation/regeneration protein (*acsV*) (Fig. 1.). The same genes are also clustered in *M. thermoacetica, Acethalobium arabaticum* and *C. ljungdahlii*. The genes for the phosphotransacetylase (Awo_c19620) and the acetate kinase (Awo_c21260) are located in other regions of the genome.

The Rnf complex is the only membrane-bound electron transfer system of the Wood-Ljungdahl pathway

Inspection of the genome sequence ruled out that any of the orfs encoding for enzymes involved in carbon flow in the Wood-Ljungdahl pathway are membrane-bound and potential Na⁺ pumps. This was a surprise since it was speculated for decades that the methyl-THF:corrinoid-iron sulfur protein methyltransferase is a membrane integral, Na⁺-translocating enzyme [10]. This assumption was based on the finding of membrane-bound corrinoids in *A. woodii* [20] and the finding of a corrinoid-containing, Na⁺translocating methyltransferase in methanogens [21]. *A. woodii* encodes 30 different methyltransferase systems involved in channelling methyl groups from various substrates into the central pathway, but none is predicted to be membrane-bound. The corrinoidcontaining proteins found in the membrane fraction of *A. woodii* [20] may thus stem from "sticky" cytoplasmic methyltransferases.

Thus, the genome sequence revealed that Rnf is probably the only ion-pumping enzyme active during litho-autotrophic growth of A. woodii. The use of Rnf as the only coupling site in the pathway leads to important consequences. The E_0' of the Fd_{ox}/Fd^2 couple is probably -500 mV and that of the NAD⁺/NADH couple is -320~mV (the $E_0{\,^\prime}$ of the Fd_{ox}/Fd^{2-} couple is assumed to be near -500 mV because in A. woodii reduced ferredoxin is used as electron donor for the reduction of CO_2 to CO, the E_0' of the CO_2/CO couple being -520 mV [20]). The higher the Fd^{2-}/NAD^{+} concentration ratio, the higher the energy that can be used for building up a sodium ion gradient. Therefore, the entire catabolic metabolism has to be optimized to produce a maximal Fd²⁻/NAD⁺ ratio. A maximal number of oxidation steps should produce reduced ferredoxin whereas a maximal number of reduction steps should use NADH as electron donor. This leads to a metabolism focused on ferredoxin.

Overcoming a first energy barrier in the electron pathway

In the first step of the electron transfer pathway, ferredoxin is reduced with H₂ as reductant. This reaction is highly endergonic with a $\Delta G^{o'} = +18.5 \text{ kJ/mol} (E_0' \text{ Fd}_{ox}/\text{Fd}_{red} = -500 \text{ mV}; E_0' 2$ $H^+/H_2 = -414 \text{ mV}$ and thus requires the input of energy. One possibility to overcome this steep energy barrier is to drive the endergonic electron transfer from molecular hydrogen to ferredoxin by the electrochemical ion potential across the membrane through ion influx into the cell. Such a kind of reverse electron flow is catalyzed by Ech-type or related hydrogenases [22] but genes encoding such enzymes are absent from the chromosome. Moreover, the genome of A. woodii does not harbour genes encoding [NiFe]-hydrogenases but a putative operon consisting of 5 genes (hydA1-hydE, Awo_c26970-Awo_c27010, Fig. 2) that potentially encode a cytoplasmic multimeric [FeFe]-hydrogenase. hvdA1 encodes a putative catalytic hydrogenase subunit with a complete H-cluster, one [2Fe2S] and three [4Fe4S] clusters, based on sequence similarities. hydB codes for a putative iron sulfur protein with one [2Fe2S], three [4Fe4S] and a flavin binding site. hydC is the first gene of the cluster and codes for a protein with one predicted [2Fe2S] binding site. The protein encoded by hydD has apparently no cofactor binding sites. HydE has no similarity to hydrogenase subunits but is similar to ATP - binding domains of histidine kinases. Proteomic analysis of cells grown with H2+CO2 vs. cells grown on fructose revealed that the [FeFe]-hydrogenase is upregulated ~3 fold during litho-autotrophic growth (Fig. S1, Table S1) which is consistent with the hypothesis that it catalyzes hydrogen activation.

The putative encoded multimeric hydrogenase (HydA-D) shares similarities with a trimeric hydrogen-evolving hydrogenase of

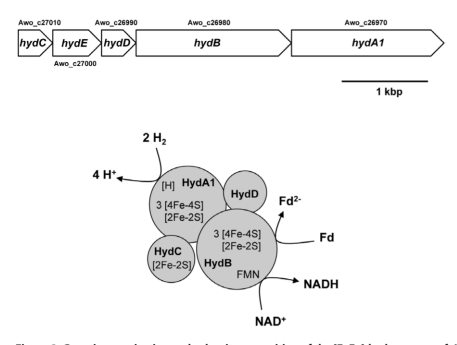


Figure 2. Genetic organization and subunit composition of the [FeFe]-hydrogenase of *A. woodii*. Awo_c27000 (HydE) is not a component of the active enzyme but may be involved in assembly or signalling. doi:10.1371/journal.pone.0033439.g002

Thermotoga maritima [23,24] that consists of the three proteins HydA, HydB and HydC. Production of hydrogen from NADH+H⁺ is thermodynamically highly unfavourable and Schut and Adams [23] proposed a flavin-based electron bifurcating mechanism to overcome the energetic barrier. Flavin-based electron bifurcation was first shown in anaerobic clostridia as a new coupling mechanism in bioenergetics: the energy liberated during energetic "downhill" transfer of one electron from a donor to an acceptor is used to transfer another electron "uphill" to an acceptor with a more negative redox potential. That was first experimentally proven for the exergonic NADH+H⁺-driven reduction of crotonyl-CoA that is coupled with endergonic ferredoxin reduction [25,26]. For the hydrogen-evolving hydrogenase of T. maritima, it is suggested that the energetic "downhill" transfer of two times one electron from ferredoxin $(E_0' \text{ approx.} -$ 500 mV) to two protons (2 H^+/H_2 ; E_0' -414 mV) drives the energetic "uphill" transport of two times one electron from NADH+H⁺ ($E_0' = -320$ mV) to two protons. In sum, the reaction is:

$$Fd^{2-} + NADH + H^+ \rightarrow 2 H_2 + Fd_{ox} + NAD^+$$
(2)

Due to the similarity of the enzymes from *A. woodii* and *T. maritima* we suggest that the [FeFe]-hydrogenase of *A. woodii* catalyzes a reversal of equation 2, namely the reduction of one ferredoxin and one NAD⁺ with two H₂ (Fig. 2). The finding that cell free extracts of *A. woodii* grown on fructose catalyze hydrogen-dependent NAD⁺ reduction only after the addition of FMN and ferredoxin purified from *Clostridium pasteurianum* is consistent with this hypothesis.

Overcoming a first energy barrier in the carbon pathway

The first step in CO_2 reduction is an endergonic reaction assuming that electrons are passed from molecular hydrogen to CO_2 via NADH+H⁺ as a reductant (the standard redox potential of the CO_2 /formate couple is -430 mV). Electrons may be delivered via NADPH generated from NADH+H⁺ in an energy-dependent reaction driven by either a membrane-integral transhydrogenase [27] or an electron bifurcating soluble transhydrogenase [28]. However, homologues of these enzymes are apparently not encoded in A. woodii. But how are the electrons transferred from molecular hydrogen to carbon dioxide? As mentioned above, the formate dehydrogenase gene cluster harbors one gene (Awo_c08260; hydA2) encoding a protein with high similarity (43%) to HydA1 (Awo_c26970) from the [FeFe]-hydrogenase of A. woodii. HydA2 is predicted to contain a complete H-Cluster. The genetic organization is similar to a gene cluster of *Treponema primitia* that was reported to belong to the same class as the hydrogenase-coupled formate dehydrogenase (FdhF) from *Escherichia coli*, which oxidizes formate and produces H_2 during sugar fermentation. Based on the similarity of FdhF_{Tp} to FdhF of *E. coli* and the presence of the hydrogenase genes it was suggested for the enzyme of T. primitia that it directly uses molecular hydrogen as reductant [29]. A related gene cluster was also found in Eubacterium acidaminophilum [30]. This gene cluster consists of genes encoding hydrogenase and formate dehydrogenase subunits but the encoded hydrogenase is more complex and similar to the multimeric hydrogenase from A. woodii. Graentzdoerffer et al. [30] proposed the most simple type of a formate hydrogen lyase system.

Based on the analogy we suggest that Awo_c08190-Awo_c08260 encode a hydrogenase-coupled formate deyhdrogenase that oxidizes hydrogen and donates the electrons *via* the electron transfer subunits HycB3 and HycB1/2 to CO_2 which is reduced to formate (Fig. 3).

Re-gaining reduced ferredoxin in the carbon pathway

The methylene-THF reductase catalyzed reaction is the only considerably exergonic reaction of the pathway and, therefore, was already considered in 1977 [8] as one of the energy conserving reactions in acetogens. The methylene-THF reductase was purified from the two acetogens *Clostridium formicoaceticum* [31] and *Blautia producta* [32]. The latter one consists of a single subunit with a molecular mass of 32 kDa, contains FAD as cofactor and is NADH-dependent. In contrast, the enzyme from *C. formicoaceticum*

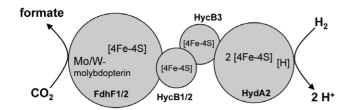


Figure 3. Subunit composition of the formate dehydrogenase of *A. woodii. fdhF1* and *fdhF2* code for sulfur and selenium containing isoenzymes, respectively. HycB1 and HycB2 may be specific for FdhF1 and FdhF2, respectively. The electron transfer subunits HycB1, HycB2 and HycB3 each have four conserved tertacysteine motifs. doi:10.1371/journal.pone.0033439.g003

consists of two subunits (26 and 35 kDa), has been proposed to be ferredoxin-dependent and contains FAD, FeS cluster and zinc.

In *A. woodii*, there is one gene (Awo_c09310) that encodes a soluble, 33 kDa protein with similarity to methylene-THF reductases (MetF) with a conserved flavin binding site. The preceding gene (Awo_c09300) encodes a protein with 205 amino acids and the region from amino acid 104 to 200 is similar to the C-terminal domain of methylene-THF reductases. This domain is encoded in many organisms next to *metF*, in several organisms it is even fused to *metF*. Awo_c09300 encodes for a 23 kDa protein that contains 8 conserved cysteine residues, which could coordinate two iron-sulfur clusters. Therefore it seems likely that this protein is the small subunit of the methylene-THF reductase of *A. woodii* and the entry point for electrons. Thus, the methylene-THF reductase is suggested to have a small (MetV) and large subunit (MetF) encoded by Awo_c09300 and Awo_c09310 with flavin and FeS cluster as cofactors.

The reduction of methylene-THF (E_0 ' methylene-THF/methyl-THF = -200 mV) with NADH is exergonic [33] and was suggested to drive the reduction of ferredoxin with NADH [34] by electron bifurcation [26,35]. This is supported by the predicted flavin and FeS centers (Fig. 4). Reduction of ferredoxin in this step is beneficial for the overall bioenergetics of the pathway (see below).

Interestingly, the genes encoding the small and large subunit of the methylene-THF reductase are cotranscribed with the preceding gene, Awo_c09290, indicating that it is also involved in the reduction of methylene-THF. Awo_c09290 encodes a protein with similarity (61%) to the RnfC-subunit of the Rnf complex, and the residues for binding flavin and the FeS-cluster are conserved. Thus, it is named RnfC2. In contrast to RnfC1 an extension of 220 amino acids is found at the N-terminus, which is surprisingly similar (59%) to the small subunit of the methylene-THF reductase. It is, therefore, conceivable that RnfC2 could substitute for the small subunit in the methylene-THF reductase and bind directly to the Rnf complex at the RnfC binding site. The electrons derived from the Rnf complex could then be transferred to the methylene-THF reductase without the intermediate step via NADH (Fig. 4, right panel). Reduced ferredoxin from the methylene-THF reductase could be reoxidized immediately at RnfB. This would fit to previous findings that this step may be membrane-associated and involved in ion translocation [36,37].

Energy conservation during autotrophic growth: a quantitative model

Based on the experimental data and the predictions from the genome sequence, a model for the bioenergetics of acetogenesis from H_2+CO_2 is proposed (Fig. 5). Oxidation of hydrogen is catalyzed by the electron bifurcating [FeFe]-hydrogenase, six mol of hydrogen are used to reduce three mol of NAD⁺ and three mol of ferredoxin. The three mol of ferredoxin are oxidized by the Rnf complex to reduce another three mol of NAD. Assuming a Na⁺/ e⁻ stoichiometry of 1, electron transfer is accompanied by the

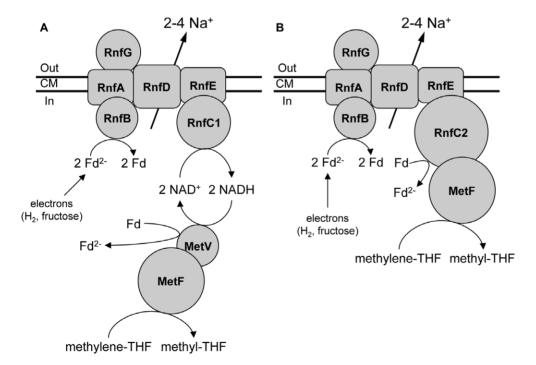


Figure 4. Subunit composition and model of the methylene-THF reductase of *A. woodii.* Panel A depicts an indirect coupling via NADH+H⁺ and the small subunit of the methylene-THF reductase (MetV) as electron input module, panel B a direct coupling of the methylene-THF reductase to the Rnf complex *via* RnfC2. doi:10.1371/journal.pone.0033439.q004

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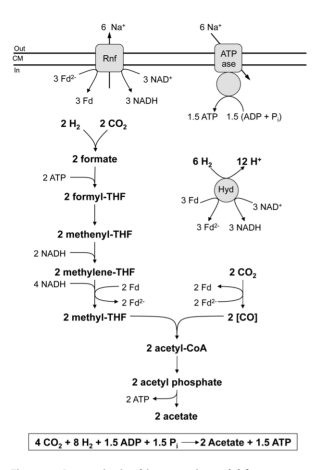


Figure 5. A quantitative bioenergetic model for acetogenesis from H₂+CO₂. The amount of ions translocated by the Rnf complex and the Na⁺ F₁F₀ ATP synthase are not exactly known. For sake of clarity, a Na⁺/e⁻ stochiometry of 1:1 is assumed (based on a $\Delta G^{O'}$ value of -37 kJ/mol; E₀' Fd^{2-/}Fd_{ox} = -500 mV; E₀' NADH+H⁺/NAD⁺ = -320 mV; and Δu_{Na+} (electrochemical Na⁺ potential across the cytoplasmic membrane) = -320 mV). For the ATP synthase, a Na⁺/ATP stoichiometry of 4 is assumed. The redox potentials (E₀') are: CO₂/formate = -430 mV, methenyl-THF/methylene-THF = -300 mV, methylene-THF/methyl-THF = -200 mV, CO₂/CO = -520 mV.

translocation of six mol of Na⁺. Reentry of the six mol of Na⁺ allows for the production of 1.5 mol ATP by the ATP synthase, assuming a stoichiometry of $4H^+/ATP$ for the ATP synthase. Two mol NADH+H⁺ are channelled into the methylene-THF dehydrogenase reaction, four mol into the methylene-THF reductase reaction. The latter may yield two mol of reduced ferredoxin by electron bifurcation that are used to reduce two mol of CO₂ to CO. The model gives an even redox balance and postulates an ATP gain of 0.75 ATP per mol of acetate formed.

The reduction of CO₂ to formate with H₂ is endergonic by +3 kJ/mol under physiological standard conditions (H₂ and CO₂ as gases at 10⁵ Pa partial pressure, formate at 1 M concentration and pH = 7.0). As will be shown below, the threshold concentration of *A. woodii* for H₂ at a CO₂ partial pressure of 0.2×10^5 Pa is 250 Pa. Under these conditions the equilibrium concentration of formate is about 0.1 mM which is in the range of the K_m for formate of formyl-THF synthethase, which catalyzes the second step in CO₂ reduction to acetate, namely the formation of formyl-THF from formate, THF and ATP. The reduction of CO₂ to formate with H₂ in *A. woodii* is therefore most likely not energy dependent. Consistent with this hypothesis, the reduction of CO₂ to formate with H_2 in *A. woodii* is catalyzed by a cytoplasmic enzyme complex composed of a [FeFe]-hydrogenase, two electron transfer iron-sulfur proteins and a molybdenum/tungsten-dependent formate dehydrogenase.

The free energy change associated with 2 CO_2 reduction with 4 H_2 to acetate under physiological conditions (CO_2 partial pressure of 0.2×10^5 Pa, a H_2 partial pressure of 250 Pa and an acetate concentration of 10 mM at pH 7) is about -40 kJ/mol which is sufficient to drive the phosphorylation of only less than 1 mol ADP considering that under the irreversible conditions in cells between 50 and 80 kJ/mol are required for the synthesis of 1 mol ATP [8].

The hydrogen threshold

To determine the minimal hydrogen concentration that can be used by *A. woodii* cell suspensions were incubated under different hydrogen partial pressures and the hydrogen consumption was monitored by gas chromatography. *A. woodii* was able to oxidize hydrogen down to 2500 ppm (250 Pa) which leads to a redox potential of about -340 mV (Fig. 6). Therefore, we can conclude that the minimal hydrogen concentration in the ecosystem must be above 250 Pa to allow litho-autotrophic growth on H₂+CO₂. Acetogenesis from H₂+CO₂ has a free energy change of -95 kJ per mol which leads to a theoretical hydrogen threshold ($\Delta G = 0$ kJ/mol) of about 10 Pa. If we assume an ATP gain of n ATP/acetate this leads to the equation:

$$4 H_2 + 2 CO_2 + n ADP + n P_i \rightarrow$$

$$CH_3COO^- + H^+ + n ATP + 2 H_2O$$
(3)

With a ΔG of -45 kJ/mol (ΔG for ADP phosphorylation ~ 50 kJ/mol) and n = 0.5 the theoretical threshold for H₂ increases to approximately 100 Pa. For n = 1 the threshold is reached at approximately 1100 Pa. The measured and theoretical hydrogen concentrations are in very good accord leading to an ATP gain in acetogenesis of 0.5 to 1 mol ATP/mol acetate. It has to be considered that the measured threshold is always lower than the theoretical one due to a partial uncoupling of metabolism and energy conservation. The threshold for H₂ of 250 Pa is very high compared to the thresholds of methanogens without cytochromes which is one order of magnitude lower [38]. Therefore *A. woodii* can not compete with those methanogens in its natural habitat for H₂+CO₂ as carbon and energy source.

Conclusions

The acetogenic bacterium A. woodii is a prime candidate for an organism that catalyzes what may be one of the first life sustaining processes on earth based on CO2 fixation. Acetate formation from hydrogen and carbon dioxide by the Wood-Ljungdahl pathway is catalyzed by cytoplasmic, soluble enzymes. Energy conservation is via a chemiosmotic mechanism of the simplest type with just two enzyme complexes: the ATP synthase and the Rnf complex. The Rnf complex couples oxidation of reduced ferredoxin with reduction of NAD⁺ and concomittant export of Na⁺ from the cells. The use of Na⁺ as coupling ion underlines current views that a sodium ion-based bioenergetics evolutionary predates a protonbased bioenergetics [39]. The energy barriers that have to be overcome in the pathways of electron and carbon flow are not driven by a membrane-bound, reverse electron flow but by soluble enzymes that use electron bifurcation. Moreover, even the little potential drop between NADH ($E_0' = -320$ mV) and methylene-THF ($E_0' = -200 \text{ mV}$) may be used bioenergetically, to reduce ferredoxin. A. woodii lives on the thermodynamic edge of life and uses a fascinating repertoire of enzymes to cope with energy

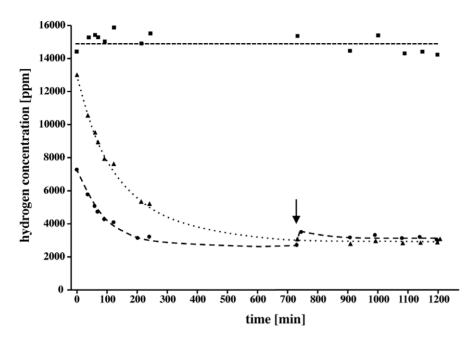


Figure 6. The hydrogen threshold concentration for acetogenesis from H_2+CO_2 . Time course of hydrogen oxidation by cell supensions of *A. woodii* with CO_2 as terminal electron acceptor. Assays received 1.5 (\blacktriangle) or 1 ml (\bigcirc) hydrogen. One assay did not contain cells (\blacksquare). At the indicated time point, hydrogen was added again to proof the viability of the cells. doi:10.1371/journal.pone.0033439.q006

limitations. The metabolism is optimized to (i) get as much as possible reduced ferredoxin to fuel the Rnf complex and (ii) not to use the sodium motive force to overcome energy barriers but the soluble electron bifurcating [FeFe]-hydrogenase and methylene-THF reductase and a hydrogen-coupled formate dehydrogenase. Altogether, this allows for the synthesis of about to 0.5–1 mol ATP per mol of acetate produced.

Materials and Methods

Growth of A. woodii

A. woodi was cultivated at 30°C. The medium was prepared as described [40]. Fructose was used as carbon source at a final concentration of 20 mM. Growth was followed by measuring the optical density at 600 nm (OD₆₀₀).

Sequencing strategy

Whole-genome sequencing was performed using the 454 FLX pryrosequencing system (Roche 454, Branford, USA). Total DNA of *A. woodii* was extracted by MasterPureTM complete DNA purification kit (Epicentre, Madison, USA). The isolated DNA was used to create 454-shotgun libraries following the GS FLX general library protocol (Roche 454, Branford, USA). One medium lane of a Titanium picotiter plate was used for sequencing of the library, resulting in 232121 shot gun reads. The reads were *de novo* assembled using the Roche Newbler assembly software 2.3 (Roche 454).

Gene prediction and annotation

Annotation was done by using the ERGO tool from Integrated Genomics with a two-step approach. Initially, all proteins were screened against Swiss-Prot and NCBI databases and available protein sequences from other completed genomes by using FASTA3. All predictions were verified and modified manually by comparing the protein sequences with Pfam, GenBank, ProDom, COG, and Prosite public databases. All coding sequences were searched for similarities to protein families and domains using CD-search [41].

Proteome analysis

A. woodii was grown either on fructose or on hydrogen and carbon dioxide. Cells were harvested during early exponential, late exponential and stationary growth. 2D gel electrophoresis (500 μ g soluble protein, pH 4–7) was done according to [42]. Staining and analysis of the gels and identification of proteins by mass spectrometry were done as described earlier [43] except that a Proteome Analyzer 4800 was used and the signal-to-noise ratio of the TOF-TOF measurement was raised to 10. An *A. woodii* data base was used for searching the peak lists.

Transcriptional organization

Cells were grown on fructose and harvested in the exponentiell growth phase. Isolation of RNA and analysis of transcriptional organization was performed as described previously [44].

Determination of the H₂ threshold concentration

H₂-threshold concentrations were determined with 1 g cells (wet mass) harvested in the exponential growth phase and resuspended in 1 ml medium. The 2 ml cell suspensions were transferred to 150 ml H₂-free Müller–Krempel flasks (Bülach, Germany) sealed by Perbunan rubber stoppers (Deutsch & Naumann, Berlin, Germany) and repeatedly evacuated and filled with $80\% N_2/20\%$ CO₂ to remove remaining traces of H₂. Then H₂ was added up to 2000 Pa. The flasks were then rapidly shaken at 30°C. Gas samples were taken with a gas-tight syringe (Hamilton, Bonaduz, Switzerland) at first every 10 min and then later every hour. After the H₂ concentration remained constant, H₂ was added again to ascertain that the same H₂ threshold was reached a second time. The samples were analyzed by gas chromatography using a stainless steel separation column (0.5 nm molecular sieve: 80/ 100 mesh; length = 2 m×4 mm) kept at 120°C. H₂ was detected with a thermal conductivity detector (GC-8A; Shimadzu) (carrier

gas, N₂; flow rate, 40 mL/min) [45]. The peak heights/areas were proportional to the H_2 concentration. Calibrations were done with a calibration gas mixture (1,000 ppmv) from Messer Industriegase, Sulzbach, Germany.

Supporting Information

Figure S1 The soluble proteome of *A. woodii* grown either on fructose (greenimage) or on H_2+CO_2 (redimage). The dual channel image was created with the Delta 2D software (Decodon GmbH, Greifswald, Germany). Proteins were prepared during early exponential growth, separated in a pH gradient 4–7 and stained with colloidal Coomassie Brillant Blue. (TIF)

References

- Martin W, Russell MJ (2007) On the origin of biochemistry at an alkaline hydrothermal vent. Philos Trans R Soc Lond B Biol Sci 362: 1887–1925.
- Martin W (2011) Hydrogen, metals, bifurcating electrons, and proton gradients: The early evolution of biological energy conservation. FEBS Lett in press.
- Takami H, Noguchi H, Takaki Y, Uchiyama I, Toyoda A, et al. (2012) A deeply branching thermophilic bacterium with an ancient acetyl-CoA pathway dominates a subsurface ecosystem. PLoS One 7: e30559.
- 4. Fuchs G (1986) CO_2 fixation in acetogenic bacteria: variations on a theme. FEMS Microbiol Rev 39: 181–213.
- Wood HG, Ragsdale SW, Pezacka E (1986) The acetyl-CoA pathway of autotrophic growth. FEMS Microbiol Rev 39: 345–362.
- Ljungdahl LG (1986) The autotrophic pathway of acetate synthesis in acetogenic bacteria. Ann Rev Microbiol 40: 415–450.
- Ragsdale SW (2008) Enzymology of the Wood-Ljungdahl pathway of acetogenesis. Ann N Y Acad Sci 1125: 129–136.
- Thauer RK, Jungermann K, Decker K (1977) Energy conservation in chemotrophic anaerobic bacteria. Bact Reviews 41: 100–180.
- Cordruwisch R, Seitz HJ, Conrad R (1988) The capacity of hydrogenotrophic anaerobic bacteria to compete for traces of hydrogen depends on the redox potential of the terminal electron acceptor. Arch Microbiol 149: 350–357.
- Müller V (2003) Energy conservation in acetogenic bacteria. Appl Environ Microbiol 69: 6345–6353.
- Hugenholtz J, Ljungdahl LG (1990) Metabolism and energy generation in homoacetogenic Clostridia. FEMS Microbiol Rev 87: 383–389.
- Biegel E, Müller V (2010) Bacterial Na⁺-translocating ferredoxin:NAD⁺ oxidoreductase. Proc Natl Acad Sci U S A 107: 18138–18142.
- Biegel E, Schmidt S, González JM, Müller V (2011) Biochemistry, evolution and physiological function of the Rnf complex, a novel ion-motive electron transport complex in prokaryotes. Cell Mol Life Sci 68: 613–634.
- Biegel E, Schmidt S, Müller V (2009) Genetic, immunological and biochemical evidence of a Rnf complex in the acetogen *Acetobacterium woodü*. Environ Microbiol 11: 1438–1443.
- 15. Fritz M, Müller V (2007) An intermediate step in the evolution of ATPases the F_1F_0 -ATPase from Acetobacterium woodii contains F-type and V-type rotor subunits and is capable of ATP synthesis. FEBS J 274: 3421–3428.
- Schmidt S, Biegel E, Müller V (2009) The ins and outs of Na⁺ bioenergetics in Acetobacterium woodii. Biochim Biophys Acta 1787: 691–696.
- van de Vossenberg JLCM, Ubbink-Kok T, Elferink MGL, Driessen AJM, Konings WN (1995) Ion permeability of the cytoplasmic membrane limits the maximum growth temperature of bacteria and archaea. Mol Microbiol 18: 925–932.
- Mulkidjanian AY, Galperin MY, Makarova KS, Wolf YI, Koonin EV (2008) Evolutionary primacy of sodium bioenergetics. Biology direct 3: 13.
- Pierce E, Xie G, Barabote RD, Saunders E, Han CS, et al. (2008) The complete genome sequence of *Moorella thermoacetica* (f. *Clostridium thermoaceticum*). Environ Microbiol 10: 2550–2573.
- Dangel W, Schulz H, Diekert G, König H, Fuchs G (1987) Occurence of corrinoid-containing membrane proteins in anaerobic bacteria. Arch Microbiol 148: 52–56.
- Gottschalk G, Thauer RK (2001) The Na⁺-translocating methyltransferase complex from methanogenic archaea. Biochim Biophys Acta 1505: 28–36.
- Hedderich R, Forzi L (2005) Energy-converting [NiFe] hydrogenases: more than just H₂ activation. J Mol Microbiol Biotechnol 10: 92–104.
- Schut GJ, Adams MW (2009) The iron-hydrogenase of *Thermotoga maritima* utilizes ferredoxin and NADH synergistically: a new perspective on anaerobic hydrogen production. J Bacteriol 191: 4451–4457.
- Verhagen MFJM, O'Rourke T, Adams MWW (1999) The hyperthermophilic bacterium, *Thermotoga maritima*, contains an unusually complex iron-hydrogenase: amino acid sequence analyses versus biochemical characterization. Biochim Biophys Acta 1412: 212–229.
- Fuli L, Hinderberger J, Seedorf H, Zhang J, Buckel W, et al. (2008) Coupled ferredoxin and crotonyl coenzyme A (CoA) reduction with NADH catalyzed by

Table S1 Differentially regulated proteins duringgrowth on fructose versus growth on hydrogen andcarbon dioxide.

(DOC)

Author Contributions

Conceived and designed the experiments: AP AK KS BV MH RD GG RKT VM. Performed the experiments: JV AT KS BV AP. Analyzed the data: MG JB RD RKT GG VM SS AK MH. Wrote the paper: VM GG RKT AP.

the butyryl-CoA dehydrogenase/Etf complex from *Clostridium klyuveri*. J Bacteriol 190: 843–850.

- Herrmann G, Jayamani E, Mai G, Buckel W (2008) Energy conservation via electron-transferring flavoprotein in anaerobic bacteria. J Bacteriol 190: 784–791.
- Pedersen A, Karlsson GB, Rydström J (2008) Proton-translocating transhydrogenase: an update of unsolved and controversial issues. J Bioenerg Biomembr 40: 463–473.
- Wang S, Huang H, Moll J, Thauer RK (2010) NADP⁺ reduction with reduced ferredoxin and NADP⁺ reduction with NADH are coupled *via* an electron bifurcating enzyme complex in *Clostridium kluyveri*. J Bacteriol 192: 5115–5123.
- Matson EG, Zhang X, Leadbetter JR (2010) Selenium controls transcription of paralogous formate dehydrogenase genes in the termite gut acetogen, *Treponema* primitia. Environ Microbiol Rep 12: 2245–2258.
- 30. Graentzdoerffer A, Rauh D, Pich A, Andreesen JR (2003) Molecular and biochemical characterization of two tungsten- and selenium-containing formate dehydrogenases from *Eubacterium acidaminophilum* that are associated with components of an iron-only hydrogenase. Arch Microbiol 179: 116–130.
- Clark JE, Ljungdahl LG (1984) Purification and properties of 5,10-methylenetetrahydrofolate reductase, an iron-sulfur flavoprotein from *Clostridium formicoaceticum.* J Biol Chem 259: 10845–10849.
- Wohlfarth G, Geerligs G, Diekert G (1990) Purification and properties of a NADH-dependent 5,10-methylenetetrahydrofolate reductase from *Peptostrepto*coccus productus. Eur J Biochem 192: 411–417.
- Wohlfarth G, Diekert G (1991) Thermodynamics of methylenetetrahydrofolate reduction to methyltetrahydrofolate and its implications for the energy metabolism of homoacetogenic bacteria. Arch Microbiol 155: 378–381.
- Köpke M, Held C, Hujer S, Liesegang H, Wiezer A, et al. (2010) *Clostridium ljungdahlü* represents a microbial production platform based on syngas. Proc Natl Acad Sci USA 107: 13087–13092.
- Li F, Hinderberger J, Seedorf H, Zhang J, Buckel W, et al. (2008) Coupled ferredoxin and crotonyl coenzyme A (CoA) reduction with NADH catalyzed by the butyryl-CoA dehydrogenase/Etf complex from *Clostridium kluyveri*. J Bacteriol 190: 843–850.
- Hugenholtz J, Ivey DM, Ljungdahl LG (1987) Carbon monoxide-driven electron transport in *Clostridium thermoautotrophicum* membranes. J Bacteriol 169: 5845–5847.
- Heise R, Müller V, Gottschalk G (1989) Sodium dependence of acetate formation by the acetogenic bacterium Acetobacterium woodii. J Bacteriol 171: 5473–5478.
- Kaster AK, Moll J, Parey K, Thauer RK (2011) Coupling of ferredoxin and heterodisulfide reduction via electron bifurcation in hydrogenotrophic methanogenic archaea. Proc Natl Acad Sci U S A 108: 2981–2986.
- Mulkidjanian AY, Dibrov P, Galperin MY (2008) The past and present of sodium energetics: may the sodium-motive force be with you. Biochim Biophys Acta 1777: 985–992.
- Imkamp F, Biegel E, Jayamani E, Buckel W, Müller V (2007) Dissection of the caffeate respiratory chain in the acetogen *Acetobacterium woodii*: indications for a Rnf-type NADH dehydrogenase as coupling site. J Bacteriol 189: 8145–8153.
- Marchler-Bauer A, Anderson JB, Derbyshire MK, DeWeese-Scott C, Gonzales NR, et al. (2007) CDD: a conserved domain database for interactive domain family analysis. Nucleic Acids Res 35: D237–240.
- Büttner K, Bernhardt J, Scharf C, Schmid R, Mader U, et al. (2001) A comprehensive two-dimensional map of cytosolic proteins of *Bacillus subtilis*. Electrophoresis 22: 2908–2935.
- Voigt B, Schweder T, Sibbald MJ, Albrecht D, Ehrenreich A, et al. (2006) The extracellular proteome of *Bacillus licheniformis* grown in different media and under different nutrient starvation conditions. Proteomics 6.
- Hess V, Vitt S, Müller V (2011) A caffeyl-coenzyme A synthetase initiates caffeate activation prior to caffeate reduction in the acetogenic bacterium *Acetobacterium woodii*. J Bacteriol 193: 971–978.
- Schuler S, Conrad R (1990) Soils Contain two different activities for oxidation of hydrogen. Microbiol Ecol 73: 77–83.