Aerobic H₂ respiration enhances metabolic

flexibility of methanotrophic bacteria

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Abstract

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Methanotrophic bacteria are important soil biofilters for the climate-active gas 27 methane. The prevailing opinion is that these bacteria exclusively metabolise single-28 carbon, and in limited instances, short-chain hydrocarbons for growth. This specialist 29 lifestyle juxtaposes metabolic flexibility, a key strategy for environmental adaptation 30 of microorganisms. Here we show that a methanotrophic bacterium from the phylum 31 Verrucomicrobia oxidises hydrogen gas (H₂) during growth and persistence. 32 Methylacidiphilum sp. RTK17.1 expresses a membrane-bound hydrogenase to 33 aerobically respire molecular H₂ at environmentally significant concentrations. While 34 H₂ oxidation did not support growth as the sole electron source, it significantly 35 enhanced mixotrophic growth yields under both oxygen-replete and oxygen-limiting 36 conditions and was sustained in non-growing cultures starved for methane. We 37 propose that H2 is consumed by this bacterium for mixotrophic growth and 38 persistence in a manner similar to other non-methanotrophic soil microorganisms. 39 We have identified genes encoding oxygen-tolerant uptake hydrogenases in all 40 41 publicly-available methanotroph genomes, suggesting that H₂ oxidation serves a general strategy for methanotrophs to remain energised in chemically-limited 42 environments. 43

Introduction

- Methane (CH₄) is a potent greenhouse gas with a 100-year global warming potential
 34 times greater than carbon dioxide (1). Aerobic methane-oxidising bacteria
- 47 (methanotrophs) constitute the primary biological sink for atmospheric methane (~30
- Tg annum⁻¹) (2). Together with anaerobic methane-oxidising archaea (ANME), they
- also capture the majority of biologically- and geologically-produced methane before it
- enters the atmosphere (3). Relative to their global impact as greenhouse gas
- 51 mitigators, methanotrophic bacteria exhibit low phylogenetic diversity and are
- 52 presently limited to 22 genera in the Alphaproteobacteria and Gammaproteobacteria
- 53 (4), two candidate genera (Methylacidiphilum and Methylacidimicrobium) in the
- 54 phylum Verrucomicrobia (5, 6) and two representatives of candidate phylum NC10
- 55 (7, 8). Methanotrophic bacteria have been isolated from a variety of ecosystems,

particularly at the oxic-anoxic interfaces where the fluxes of methane gas are high, 56 including peat bogs, wetlands, rice paddies, forest soils and geothermal habitats (9). 57 Methanotrophic bacteria are classified as obligate aerobes and obligate 58 methylotrophs. They grow by oxidising single carbon or short-chain organic 59 compounds (10). All species oxidise methane to methanol via the soluble or 60 particulate form of methane monooxygenase (sMMO or pMMO respectively). 61 Methanol is then oxidised to carbon dioxide (CO₂), yielding reducing equivalents for 62 energisation of the respiratory chain. The proteobacterial methanotrophs generate 63 biomass by assimilating formaldehyde, a product of MxaFI type methanol 64 65 dehydrogenase activity, via the ribulose monophosphate or serine pathways (2). The verrucomicrobial methanotrophs, in contrast, express an XoxF type methanol 66 dehydrogenase that oxidises methanol directly to formate (11) and generate biomass 67 by fixing CO₂ via the Calvin-Benson-Bassham cycle (12). To remain viable, all 68 methanotrophic bacteria simultaneously require methane, an endogenous reductant 69 (NAD(P)H or quinol), and an exogenous terminal electron acceptor (O₂) to sustain 70 the highly challenging methane monooxygenase reaction (CH₄ + O₂ + [NAD(P)H + 71 $H^+I/QH_2 \rightarrow CH_3OH + NAD(P)^+/Q + H_2O)$ and to assimilate carbon. This apparently 72 specialist lifestyle is likely to make methanotrophic bacteria highly vulnerable to 73 74 environmental changes, such as methane-limitation, hypoxia or oxidative stress. But contrary to their apparent metabolic inflexibility, it is known that methanotrophic 75 bacteria are capable of surviving within environments where methane may be 76 limited, variable, or restricted to atmospheric concentrations (1.7 ppmv) (13). Under 77 these conditions low-affinity methane monooxygenase enzymes, which dominate 78 cultivated representatives of methanotrophic bacteria (14), are incapable of methane 79 oxidation and hence other energy sources are required for survival. Recent reports 80 have shown that some methanotrophic bacteria can grow on simple organic acids, 81 alcohols and short-chain alkane gases (15). While this expansion of metabolic ability 82 supersedes the long-held paradigm that methanotrophic bacteria are obligate 83 84 methylotrophs, these observations are restricted to three of the 24 described methanotrophic genera, Methylocella, Methylocapsa and Methylocystis (16). It 85 remains unknown whether methanotrophic bacteria more commonly exploit 86

alternative energy sources to survive periods of methane starvation.

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Recent physiological and ecological studies have collectively shown hydrogen is a widely utilised energy source for microbial growth and survival across a growing range of taxa and soil ecosystems (17-19). The discovery that the dominant soil phyla including Actinobacteria (20) and Acidobacteria (18) can switch from growing on heterotrophic substrates to persisting on atmospheric hydrogen has provided a new understanding of how microorganisms survive nutrient-limited conditions. Given its ubiquity and diffusivity, hydrogen gas is an ideal energy source to support the growth or non-replicative persistence of soil bacteria. Our recent survey of hydrogenase distribution (17), metalloenzymes that catalyse the reversible oxidation of hydrogen, noted that genes encoding hydrogenases were extremely widespread. This result led us to investigate the distribution of hydrogenases in methanotrophic bacteria, and we subsequently identified genes encoding these enzymes in all 31 publicly available genomes (Figure S1). This prevalence of hydrogenase genes was surprising given the apparent specialist lifestyle of methanotrophic bacteria. Reports of hydrogenase activity in these microorganisms are scarce. Formate-dependent hydrogen production, under anoxic conditions, has been shown in cultures of Methylomicrobium album BG8 and Methylosinus trichosporium OB3b (21). Methylococcus capsulatus (Bath) is known to express both cytosolic and membranebound hydrogenases, though their physiological function has not been resolved (22). The oxidation of hydrogen in methanotrophic bacteria has been predicted to contribute reducing energy for methane oxidation (22), to recycle endogenous hydrogen produced during nitrogen-fixation (23), and to drive the non-productive oxidation of chlorinated solvents (24). However, no studies have confirmed whether hydrogen metabolism contributes to the growth and persistence of these microorganisms. To investigate the physiological role of hydrogenases in methanotrophic bacteria, we undertook a geochemical, molecular and cultivation-based survey of a geothermal soil profile at Rotokawa, New Zealand with known methanotrophic activity (25). In this work, we isolated a thermoacidophilic methanotroph from the genus Methylacidiphilum and demonstrate that it oxidises hydrogen gas during both growth and persistence. Hydrogen oxidation occurred through an aerobic respiratory process mediated by a membrane-bound [NiFe]-hydrogenase. Hydrogen oxidation enhanced mixotrophic growth yields during methanotrophic growth, under both

oxygen-replete and oxygen-limiting conditions, and was sustained in the absence of methane oxidation. We propose aerobic hydrogen respiration serves as a dependable mechanism for this bacterium – and potentially methanotrophic bacteria in general – to remain energised in otherwise physically-challenging and energetically-variable environments.

Results

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Verrucomicrobial methanotrophs are abundant in environments with high 127 levels of soil methane and hydrogen oxidation. A molecular survey of total 128 microbial taxa revealed bacterial and archaeal populations were consistent with 129 acidic soil ecosystems. Methanotrophic verrucomicrobial taxa (family 130 Methylacidiphilaceae) were the dominant OTUs identified at 10 cm depth, accounting 131 for 47 % of all bacterial 16S rRNA gene sequences retrieved (Figure 1a). 132 Methylacidiphilum spp. have been previously isolated in acidic geothermal soils in 133 134 Kamchatka (26), Italy (27, 28) and New Zealand (29) (pH optima of < 3.0), and along with *Methylacidimicrobium* spp. (6), are the only acidophilic methanotrophic bacteria 135 described. At depths between 20-50 cm, Proteobacteria, Actinobacteria and 136 Acidobacteria were present at greater abundance than Verrucomicrobia, none of 137 which include known acidophilic methanotrophs or hydrogenotrophs. At all depths, 138 total microbial taxa were dominated by archaeal OTUs, classified into the order 139 Thermoplasmatales, which is consistent with observations made previously of other 140 acidic geothermal environments (25, 30, 31). Methane and hydrogen soil gas 141 concentrations in this soil embankment were greatest at a depth of 50 cm and 142 steeply decreased below detectable limits by 10 cm depth, suggesting high levels of 143 methane and hydrogen gas oxidation was occurring in shallow depth soils (Figure 1). 144 We consistently observed rapid oxidation of hydrogen and methane in these surface 145 soils when incubated in vitro at both mesophilic and thermophilic temperatures 146 147 (Figure S2). Anticipating soil gas oxidation would primarily be driven by verrucomicrobial methanotrophs, we designed PCR primers targeting the methane 148 149 monooxygenase and hydrogenase genes encoded in the *Methylacidiphilum* infernorum V4 genome (32) (Table S1), qPCR analysis on soil DNA extracts 150 151 confirmed the genetic capacity of Methylacidiphilaceae to oxidise methane and hydrogen. The abundance of these genes was greatest in the top 20 cm of soil, 152

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which corresponded to the zones of the lowest methane and hydrogen soil gas concentrations (Figures 1b & 1c). Methylacidiphilum sp. RTK17.1 encodes membrane-bound and cytosolic [NiFe] hydrogenases. To gain further insight into the hydrogen-oxidising capacity of the Methylacidiphilaceae within the geothermal soil profile, we isolated, characterised, and sequenced a thermotolerant *Methylacidiphilum* sp. (strain RTK17.1; Table S2) from the Rotokawa soils. Characterisation of *Methylacidiphilum* sp. RTK17.1 showed that it optimally grew at pH 2.5 and 50 °C (T_{max} 58 °C) and could oxidise methane and fix carbon dioxide in common with the reported observations other Methylacidiphilum spp. (12). Glycogen accumulation, as previously described in this genus (33), was also observed (data not shown). Analysis of the Methylacidiphilum sp. RTK17.1 genome confirmed the basis for these processes (Table S2) and indicated that the metabolic strategy of this strain is consistent with the other verrucomicrobial methanotrophs. Amperometric assays were performed and demonstrated that the isolate oxidised hydrogen gas, providing the first confirmation that *Methylacidiphilum* spp., and indeed members of the dominant soil phylum Verrucomicrobia can metabolise hydrogen. The microorganism rapidly oxidised hydrogen at rates proportional to cell density (Figure 2a) and consumed hydrogen at concentrations as low as 55 ppmv (Figure S3). As with the closely related M. infernorum V4, we noted the presence of two gene clusters encoding [NiFe] hydrogenases in the sequenced genome (Table S2). These enzymes were classified into Groups 1d (hyaABC) and 3b (hyhBGSL) based on the criteria of our recent survey on environmental hydrogenase distribution (Figure S1) (17). Group 1d enzymes are membrane-bound, hydrogen-uptake [NiFe] hydrogenases that are believed to yield electrons for aerobic respiration via quinone carriers. In comparison, the identified Group 3b [NiFe] hydrogenase belongs to a class of cytosolic enzymes that couples the oxidation of NADPH to the production of hydrogen (34). Reverse transcription (RT) PCR confirmed Methylacidiphilum sp. RTK17.1 constitutively expresses both hydrogenases during methanotrophic growth in the presence of hydrogen (Figure S4). Methylacidiphilum sp. RTK17.1 oxidises hydrogen in an aerobic respiratorylinked process. Cell lysates targeting the Group 1d hydrogenase efficiently coupled

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the oxidation of hydrogen to the reduction of the artificial electron acceptor benzyl viologen ($E_0 = -374$ mV). This activity was localized in the membrane, as shown by the 31-fold increase in benzyl viologen reduction of the membrane fraction relative to the cytosol fraction (Figure 2b). The hyaC subunit of this enzyme encodes a predicted membrane-bound b-type cytochrome domain known to mediate electron transfer from hydrogen to guinone (35). To probe the physiological interactions of this hydrogenase, we measured real-time hydrogen consumption within cell suspensions. Hydrogen oxidation was stimulated by treatment with the ionophores nigericin and valinomycin respectively (Figures 2c & 2d), showing hydrogenase activity is sensitive to the magnitude of the proton and charge gradients of the proton-motive force. Consistent with being linked to the aerobic respiratory chain, such uncoupled hydrogen oxidation ceased rapidly and could be rescued by supplementation with the terminal electron acceptor, O2; under these conditions the onset of O₂-limitation was likely exacerbated by the catabolism of endogenous glycogen reserves (Figure S5). By contrast, rates of hydrogen oxidation were reduced following treatment with the protonophore carbonyl cyanide *m*-chlorophenyl hydrazone (CCCP), likely a consequence of secondary intracellular acidification (Figure S6). The Group 1d [NiFe] hydrogenase mediating this process appears oxygen-tolerant, a finding consistent with the sequenced small subunit gene encoding a 6Cys[4Fe3S] cluster, which rapidly reactivates hydrogenases following oxygen inhibition (36, 37). Collectively, these findings demonstrate that this hydrogenase is a strictly membrane-bound, respiratory-linked, oxygen-dependent enzyme that drives ATP synthesis through the Knallgas reaction, i.e. by chemiosmotically coupling hydrogen oxidation to oxygen reduction. Oxidation of hydrogen enhances mixotrophic growth. Pure culture experiments with *Methylacidiphilum* sp. RTK17.1 were performed to determine the physiological roles of the Group 1d hydrogenase. We observed that methane and hydrogen were co-oxidised in both batch bioreactor cultures (Figure S7) and chemostat experiments (Table 1). In these experiments, an exogenous energy source (i.e. hydrogen or methane) was always supplied in excess to suppress the induction of endogenous metabolism and prevent glycogen catabolism (38). Chemostat experiments revealed that hydrogen oxidation is stimulated under hypoxic growth conditions. Observed rates of hydrogen oxidation were 77-fold greater when cells were oxygen-limited

compared to oxygen excess conditions (Table 1). Further, under oxygen-limiting conditions with 1.9 % hydrogen addition, the specific consumption rate of hydrogen was 42 % greater than observed rates of methane oxidation. Hydrogen addition into the feedgas significantly (p-value < 0.05) increased growth yields (gcDW molcH4⁻¹) of Methylacidiphilum sp. RTK17.1 by 33 % under oxygen excess and 51 % under oxygen-limited conditions (Table 1). Additionally, cellular growth yields (as determined by total protein) of *Methylacidiphilum* sp. RTK17.1, following one week incubation in the presence of 1 % hydrogen and 10 % methane (v/v), were significantly greater than when grown exclusively on methane in batch experiments (Figure S3b). Hydrogen oxidation was also observed both in methane-starved cultures of Methylacidiphilum sp. RTK17.1 and following the addition of 4 % (v/v) acetylene (Figure S3a), an inhibitor of methane monooxygenase activity (39). Hence, while hydrogen oxidation alone cannot sustain growth of this methanotroph, it appears sufficient to maintain both the proton-motive force and ATP synthesis required for long-term survival in the absence of the methylotrophic growth substrates.

Discussion

This study shows a methanotrophic bacterium is able to constitutively consume methane and hydrogen, either together or separately depending on gas availability, to meet energy needs. We demonstrated that the genetic determinants of aerobic hydrogen respiration, specifically genes encoding a member of the Group 1d [NiFe] hydrogenases, are encoded and expressed in the verrucomicrobial methanotroph *Methylacidiphilum* sp. RTK17.1. The strain rapidly oxidised exogenous hydrogen with this enzyme in a membrane-bound aerobic respiratory process and likely provides hydrogen-derived reducing equivalents to the quinone pool for methane monooxygenase activity and/or cytochrome reduction, *via* activity of a recently described alternative (ACIII) cytochrome complex (40). Our model of how methane, methanol and hydrogen oxidation are integrated into the respiratory chain of verrucomicrobial methanotrophs is shown in Figure 3. Through this metabolic flexibility, we propose that *Methylacidiphilum* sp. RTK17.1 is able to balance the catabolic and anabolic requirements for reducing energy imposed by its challenging methanotrophic lifestyle and the methane monooxygenase reaction. This flexibility

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likely contributes to the dominance of the Methylacidiphilaceae in aerated acidophilic geothermal soils at Rotokawa, New Zealand. As a dominant genus in the uppermost 20 cm of these soils, Methylacidiphilum sp. RTK17.1 can simultaneously oxidise geothermally-produced methane and hydrogen gases while consuming atmospheric oxygen and fixing carbon dioxide and nitrogen gases. In addition to conferring a survival advantage during periods of methane limitation, our data shows hydrogen oxidation may be favoured during hypoxia. When grown under oxygen-limiting conditions, rates of hydrogen consumption in cultures of Methylacidiphilum sp. RTK17.1 increased by 77-fold and exceeded observed rates of methane oxidation (Table 1). Similarly, a transcriptome study investigating nitrogen fixation reported that the Group 1d hydrogenase in Methylacidiphilum fumarolicum SolV was upregulated 20-fold in response to oxygen limitation (41). This preference for hydrogen suggests methanotrophic bacteria occupying the soil oxic/anoxic interface display a predominately mixotrophic lifestyle. In fact, under oxygen limited experimental conditions, up to 32 % of the total reducing energy supplied to the respiratory chain may be provided from hydrogen-derived electrons. Recalling that pMMO requires oxygen as a substrate to catalyse the conversion of methane to methanol, mixotrophy through hydrogen respiration both sustains the energy requirements for cell growth while reducing the overall demand for oxygen. Given the energetic requirement for cell maintenance is three orders of magnitude less than for growth (42, 43), it is probable that hydrogen oxidation serves as a reliable mechanism for the persistence of methanotrophic bacteria in these habitats. Genomic evidence supports that hydrogen oxidation is a commonly utilised strategy for the growth and survival of methanotrophic bacteria. Hydrogenases were identified in all 31 genomes of methanotrophic bacteria surveyed (Figure S1). Previous studies have shown two model methanotrophs, Methylosinus trichsporium OB3b (21) and Methylococcus capsulatus (Bath) (24), oxidise hydrogen but the physiological role of their encoded hydrogenases has never been fully investigated. Consistent with hydrogen contributing to the persistence of methanotrophic bacteria, genes encoding the Group 1h [NiFe] hydrogenases, while absent from *Methylacidiphilum* sp. RTK17.1, were identified in the genomes of selected alphaproteobacterial and verrucomicrobial methanotrophs. These high-affinity enzymes are capable of oxidising tropospheric concentrations of hydrogen and have been linked to

persistence within the dominant soil phyla Actinobacteria (20) and Acidobacteria (18). Our genomic analysis indicates methanotrophic bacteria may additionally evolve hydrogen fermentatively (via Group 3b and 3d [NiFe] hydrogenases), which by analogy with *Mycobacterium smegmatis*, may be important for surviving anoxia (34). They also encode the capacity to sense hydrogen (via Group 2b and 2c [NiFe] hydrogenases) which may serve to regulate expression of uptake hydrogenases and also potentially global cellular processes. Hydrogen is likely an attractive energy source for these microorganisms because of its high-diffusibility, relative dependability, and high energy content. Whereas biogenic methane is exclusively produced by methanogenic archaea, hydrogen is biologically produced by diverse organisms across the three domains of life as a result of fermentation, photobiological processes, and nitrogen fixation (17). Hence, the ability of methanotrophic bacteria to utilise this widespread energy source may enhance their capacity to colonise environments and adapt to environmental change. This hidden metabolic flexibility may explain why the range of low-affinity methanotrophic bacteria extends beyond ecosystems with high methane fluxes (44). By demonstrating that respiratory-linked hydrogen oxidation enhances growth, we reveal that verrucomicrobial methanotrophs are mixotrophs capable of growth and survival on hydrocarbon and non-hydrocarbon substrates alike. The narrative that methanotrophic bacteria are niche specialists, with obligately methylotrophic lifestyles, is based on studies under optimal growth conditions and ignores the requirement of these organisms to adapt to methane and/or oxygen limitation. Particularly within dynamic oxic/anoxic boundaries, where methanotrophic bacteria occur in high abundance (45), a specialist metabolism is likely to reduce the overall fitness of these microorganisms. We therefore propose that mixotrophic growth and survival through hydrogen metabolism provides a general mechanism by which this contradiction is resolved. Expanding the metabolic flexibility of methanotrophs will have broad ecological implications and result in an improved understanding of global methane and hydrogen cycles.

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Environmental sampling. Soil samples (~ 50 g) were collected every 10 cm from the surface of the sampling site (38°37'30.8"S, 176°11'55.3"E) to a maximum depth of 50 cm. This site, located within Rotokawa geothermal field was selected on the basis of previous research showing elevated soil gas concentrations (25). Soil temperatures were measured in the field using a 51II single input digital thermometer (Fluke). The pH was measured upon returning to the laboratory using a model HI11310 pH probe (Hanna Instruments). Soil gas samples were collected every 10 cm using a custom built gas-sampling probe equipped with a 1 I gas tight syringe (SGE Analytical Science). Gas samples were collected and stored at 25 °C in 50 ml Air & Gas Sampling Bags (Calibrated Instruments) and were processed within 48 h on a 490 MicroGC (Agilent Technologies, Santa Clara, CA) equipped with Molecular Sieve 5A with heated injector (50 °C, back-flush at 5.10 s, column at 90 °C, 150 kPa), a PoraPak Q column with heated injector (50 °C, no back-flush, column at 70 °C,150 kPa) and a 5CB column with heated injector (50 °C, no back-flush, column at 80 °C, 150 kPa). Methane and hydrogen gas consumption by soil microbial communities was determined by incubating 1 g of soils collected from the Rotokawa sampling site (depth < 10 cm) in 112 ml gas-tight serum bottles at 37 and 50 °C. The serum bottle headspace were air emended with 300 ppmv and 400 ppmv of hydrogen and methane gas respectively. Headspace methane and hydrogen gas concentrations were measured with a PeakPerformer gas chromatograph equipped with a flame ionising detector (FID: methane) and a PP1 Gas Analyzer equipped with a reducing compound photometer (RCP: hydrogen). **Isolation and cultivation of** *Methylacidiphilum* **sp. RTK17.1.** Soil samples (1 g) were inoculated into serum bottles containing 50 ml media (pH 2.5). All cultivations were performed in a V4 mineral medium as described previously (29) but with the addition of rare earth elements lanthanum and cerium (27). Methane (10 % v/v) and carbon dioxide (1 %) were added to an air headspace and samples were incubated at 60 °C with shaking 150 rpm. Methane in the headspace was monitored with a PeakPerformer gas chromatograph equipped with a flame ionising detector (FID). Following several passages (10 % v/v) into liquid media, enrichments were transferred onto solid media. Following several weeks incubation (60 °C) single

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colonies were re-streaked before being transferred back into liquid media. Isolate identity was confirmed via sequencing of the 16S rRNA gene (Macrogen) using bacterial 9f/1492r primers (46). Chemostat cultivation and batch experiments. Chemostat cultivation of *Methylacidiphilum* sp. RTK17.1 was performed to investigate the influence of hydrogen on growth under oxygen excess and oxygenlimiting conditions. A 1 I bioreactor (BioFlo 110, New Brunswick Scientific. Edison, NJ. USA) was used for these studies. Cultures were incubated at 50 °C and pH 3 with continuous stirring (800 rpm). Bioreactor volume was kept constant at 0.5 l by automatic regulation of the culture level. V4 mineral media was supplied at a constant flow rate of 10 ml h⁻¹ (D=0.02 h⁻¹). Dissolved oxygen was monitored using a InPro 6810 Polarographic Oxygen Sensor (Mettler-Toledo). Custom gas mixtures were prepared in a compressed gas cylinder and supplied to the chemostat at a rate of 10 ml min⁻¹ using a mass flow controller (EL-FLOW, Bronkhorst, Netherlands). Gas mixtures contained approximately (v/v) 3 % methane and 26 % carbon dioxide for all experiments. For oxygen excess and oxygen-limiting conditions, (v/v) 14.1 % and 3.5 % oxygen was supplied. High, medium and low hydrogen experiments consisted of (v/v) 1.9 %, 0.7 % and 0.4 % hydrogen additions. The balance of all gas mixtures was made up with nitrogen. Cell density in liquid samples was monitored by measuring turbidity at 600 nm using a Ultrospec 10 cell density meter (Amersham Bioscience). One unit of OD600 was found to be equivalent to 0.43 g l⁻¹ cell dry weight (CDW) for *Methylacidiphilum* sp. RTK17.1. After achieving a steady state condition as determined by OD₆₀₀, influent and effluent gas concentrations were monitored over several days using a 490 MicroGC (Agilent Technologies, Santa Clara, CA). Biomass cell dry weight was used to calculate growth rate and specific gas consumption rate. To investigate whether Methylacidiphilum sp. RTK17.1 oxidises ecologically significant concentrations of hydrogen, 350 ml cultures (in triplicate) were incubated in 1 I rubber-stoppered Schott bottles in a headspace of air supplemented with 100-250 ppmv hydrogen, 1 % (v/v) methane and 1 % carbon dioxide. Acetylene gas was added in some experiments (4 % v/v) to inhibit MMO activity as previously described

(39). Finally, to determine if growth was enhanced in the presence of hydrogen, 20 377 paired cultures (n = 40) of Methylacidiphilum sp. RTK17.1 were incubated with or 378 without 1 % (v/v) hydrogen in an air headspace supplemented with (v/v) 10 % 379 methane and 1 % carbon dioxide. Cultures were incubated in a custom Test Tube 380 Oscillator (TTO; agitation 1.2 Hz; Terratec). Following seven days incubation, total 381 protein was determined as described previously (47). Statistical significance of 382 observed differences of growth yields was determined using a *student's t-test* (α = 383 384 0.05). Headspace mixing ratios of hydrogen and methane were monitored throughout batch experiments as described above. 385 **Hydrogenase activity measurements.** Hydrogenase activity of *Methylacidiphilum* 386 sp. RTK17.1 was measured in stationary-phase cultures harvested from the 387 bioreactor. For amperometric measurements, whole-cells were concentrated 5-, 10-, 388 20-, and 30-fold by centrifugation followed by resuspension in phosphate-buffered 389 saline (pH 7.4). Rate of hydrogen oxidation was measured at 50 °C using a 390 391 hydrogen-MR hydrogen microsensor (Unisense) as previously described (18, 48). For colourimetric assays, 500 ml culture was harvested by centrifugation (15 min, 392 393 5.000 x g, 4 °C) and treated as previously described (18, 20) to prepare crude. cytosolic and membrane fractions for analysis. To test for hydrogenase activity, 394 samples (20 µg protein) from each cell fraction were incubated with 1 ml 50 mM 395 potassium phosphate buffer (pH 7.0) and 50 µM benzyl viologen for eight hours in an 396 anaerobic chamber (5 % hydrogen, 10 % carbon dioxide, 85 % nitrogen (v/v)). 397 Debris was removed by centrifugation (15 min, 10,000 x g, 4 °C) and the 398 absorbance of the supernatants was read at 604 nm in a Jenway 6300 399 spectrophotometer. 400 Additional methodologies for nucleic acid extraction, PCR, genome sequencing, 401 oxygen consumption experiments and soil microbial community determination can 402 be found in the supporting information. 403 404

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Author contributions

CRC, CG, MBS, GC, KMH, KH, CC, DJG, RS and BM contributed to experimental design. Bioreactor and wet lab experiments were conducted by CRC, BM, CC, DJG and KMH. CRC, CG and KH undertook cell respiratory analysis. CRC and MBS performed fieldwork and genomic analyses. JFP performed the microbial community analysis. CRC, MBS, CG, KH and GC wrote the manuscript.

Competing Interests

The authors declare no competing financial interests.

Figures and Tables

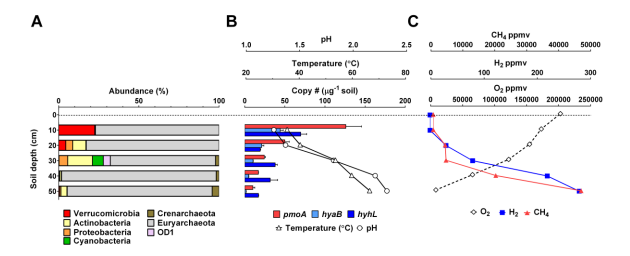


Figure 1. Methanotrophic bacteria are highly abundant in the uppermost soils in Rotokawa, New Zealand. OTUs classified belonging to phylum Verrucomicrobia were the dominant bacterial taxa identified in the top 10 cm. Consistent with a methanotrophic lifestyle, all verrucomicrobial OTUs were further classified into the family Methylacidiphilaceae. 16S rRNA gene sequencing was performed (illumina) on total DNA extracted at 10-50 cm soil depth. Non-rarefied abundance results (%) are shown for all OTUs (> 100 reads) from 130,289 total sequence reads (with an average of 26,058 per sample depth). The abundance (B) of the verrucomicrobial particulate methane monooxygenase (*pmoA*), Group 1d (*hyaB*) and Group 3b (*hyhL*) hydrogenase genes and corresponding (C) soil gas, temperature and pH profiles suggests both high levels of aerobic methane and hydrogen oxidation activity by these acidophilic taxa in the uppermost 20 cm. Error bars shown represent the standard deviation of triplicate measurements.

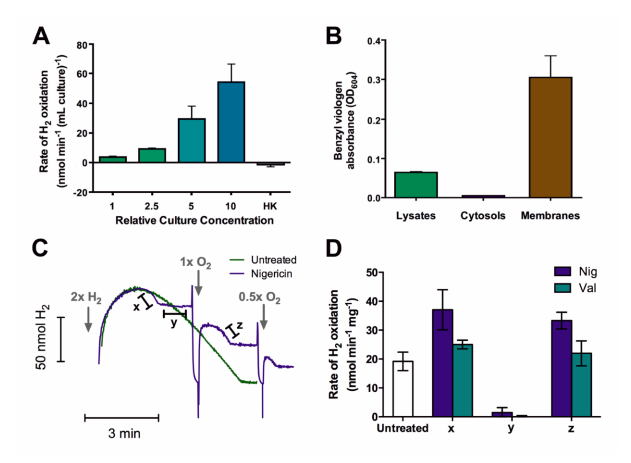


Figure 2. Hydrogenase activity of *Methylacidiphilum* sp. RTK17.1. (A) Whole-cell oxidation of hydrogen measured amperometrically using a hydrogen microsensor. The density-dependence of this process and its sensitivity to heat inactivation (HK) is shown. (B) Colourimetric detection of hydrogenase activity in protein concentration-normalized cell lysates, cytosols, and membranes anaerobically treated with the artificial electron acceptor benzyl viologen. (C) Real-time hydrogen oxidation by untreated cells and nigericin-treated cells. The relative amounts of hydrogen and oxygen added at specific time points is shown. (D) Rates of hydrogen oxidation of untreated, nigericin-treated, and valinomycin-treated cells. For the uncoupler-treated cultures, the initial (*x*), oxygen-limited (*y*), and oxygen restored (*z*) rates of oxidation are shown, which correspond to the rates highlighted in panel (C). Endogenous glycogen catabolism likely contributed to oxygen-limitation (y) observed in nigericin treated cells (Figure S5).

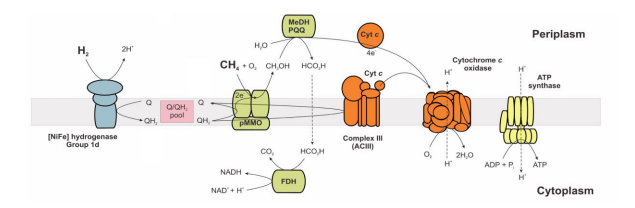


Figure 3. Proposed model of respiratory-linked methane and hydrogen oxidation in *Methylacidiphilum* sp. RTK17.1. During mixotrophic growth, the oxidation of both hydrogen (H₂) and methane (CH₄) yields reducing equivalents (QH₂) for the respiratory chain, a large proton-motive force is generated, and sufficient ATP is produced for growth via an H⁺-translocating F₁F₀-ATP synthase. Some of the quinol generated through H₂ oxidation provides the electrons necessary for pMMO catalysis. Following CH₄ oxidation by pMMO, ensuing reactions catalysed by an XoxF type methanol dehydrogenase (MeDH) and formate dehydrogenase (FDH) contribute additional reductant (Cyt *c* and NADH) into the respiratory chain for ATP production and growth (11). In methane-limiting conditions, the identified Group 1d [NiFe] hydrogenase generates a small proton-motive force, oxidised reductant (Q) is maintained via type ACIII cytochrome complex (40) and sufficient ATP is generated for maintenance. Respiratory complexes I and II are not shown but are encoded in the genome of *Methylacidiphilum* sp. RTK17.1 (Table S2).

Condition a,b	Growth rate H ₂ consumption		CH₄ consumption		Biomass Yield ^d	
	(mg l ⁻¹ h ⁻¹)	(mmol l ⁻¹ h ⁻¹)	(mmol g _{CDW} ⁻¹ h ⁻¹)	(mmol l ⁻¹ h ⁻¹)	(mmol g _{CDW} ⁻¹ h ⁻¹)	(g _{CDW} mol _{CH4} -1)
O ₂ excess, medium H ₂	9.33 (± 0.52)	0.01 (± 0.01)	0.01 (± 0.03)	1.06 (± 0.12)	2.40 (± 0.38)	$8.90^{\text{T}} (\pm 1.51)$
O ₂ excess, no H ₂	8.17 (± 0.82)	0.00	0.00	1.22 (± 0.05)	2.85 (± 0.18)	6.68 (± 0.41)
O ₂ -limiting, high H ₂	5.57 (± 0.16)	0.78 (± 0.02)	2.66 (± 0.11)	0.55 (± 0.03)	1.89 (± 0.11)	$10.11^{4} (\pm 0.64)$
O ₂ -limiting, medium H ₂	4.99 (± 0.17)	0.33 (± 0.01)	1.34 (± 0.06)	0.70 (± 0.02)	2.80 (± 0.13)	7.16 (± 0.34)
O_2 -limiting, low H_2	5.60 (± 0.31)	0.19 (± 0.00)	0.73 (± 0.04)	0.71 (± 0.01)	2.68 (± 0.16)	$7.87^{4} (\pm 0.45)$
O ₂ -limiting, no H ₂	5.40 (± 0.26)	0.00	0.00	0.80 (± 0.05)	3.11 (± 0.27)	6.79 (± 0.55)

^a Feedgas was continuously supplied at a rate of 10 ml min⁻¹ with the following composition % (v/v): O₂ Excess, 14.1 %; O₂ limiting, 3.5 %. High, medium and low H₂ experiments consisted of 1.9 %, 0.7 % and 0.4 % H₂ supply. For all experiments excess CH₄ was supplied at 3 %, CO₂ at 26 % with the balance made up with N₂.

^b a dilution rate of 0.02 h⁻¹ was maintained for all experiments

° gram cell dry weight (g_{CDW}) measurements were determined by correlating OD_{600} measurements to an existing standard curve ($OD_{600} = 1$ corresponds to 0.43 g_{CDW}/I).

^d Standard deviation shown in brackets are calculated from a minimum of triplicate measurements

*denotes significant increase (p-value < 0.05) in yield from O₂-limiting, no H₂ result; unpaired t-test, one tailed.

[‡] denotes significant increase (*p*-value < 0.05) in yield from O₂ excess, no H₂ result; unpaired *t*-test, one tailed.

Supporting Information

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Supplementary Methods

Nucleic acid extraction Methylacidiphilum sp. RTK17.1 genomic DNA was extracted using the NucleoSpin Tissue kit (Macherey-Nagel) as per the manufacturer's instructions for difficult to lyse Gram-positive bacteria. DNA was extracted from Rotokawa soil samples using the NucleoSpin Soil DNA extraction kit (Macherey-Nagel) as per the manufacturers recommended protocol. For RNA isolation from *Methylacidiphilum* sp. RTK17.1, stationary phase (Abs_{600nm} = 4.0, 10 ml cultures) cells were harvested by centrifugation (27,000 x g, 20 min, 4 °C), resuspended in 1 ml RNA Later (ThermoFisher Scientific) and stored at -20 °C as recommended. Cell lysis, total RNA extractions and cDNA synthesis were performed as previously described (18). PCR, RT-PCR, Quantitative PCR and genome sequencing Methylacidiphilum sp. RTK17.1 genomic DNA and cDNA (10 ng) was (RT)PCR amplified using i-Taq DNA polymerase (2.5 U; Intron Biotechnology) in 50 µl reactions containing dNTPs (1 mM), 1X PCR Buffer, MgCl₂ (1.5 mM), forward primer (0.5 μM) and reverse primer (0.5 μM). Amplifications conditions using an Mx3000p thermocycler (Strategene) were as follows; an initial 94 °C for 5 min; followed by 30 cycles of 94 °C for 45 s; 55 °C for 45 s; 72 °C for 90 s; and a final extension at 72 °C for 5 min. PCRs were run through electrophoresis on a 1 % (w/v) TAE agarose gel with 1X RedSafe nucleic acid staining solution and visualised on a GeneGenius Bio Imaging System (Syngene). Quantitative-PCR (gPCR) was performed to investigate the abundance of Verrucomicrobial methanotrophs type by probing for MMO (pmoA) and hydrogenase genes (Group 1d and 3b) within the Rotokawa soil profile. Total DNA extracted from 1 g soil was amplified using a Probe Fast Universal qPCR kit (KAPA Biosystems) according to the manufacturer's instructions. Primer sequences for all qPCR

amplifications are provided in Table S2. qPCR reactions were optimised and

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conducted on an Mx3000p thermocycler (Strategene). Abundance (expressed per µg Soil) of MMO and hydrogenase genes in DNA samples was calculated from cycle threshold values (Ct) and standard curves. To prepare qPCR standard curves, amplicons were cloned following the recommended protocol for the TOPO-TA Cloning kit (ThermoFisher Scientific). Transformation of competent *Eschericia coli* cells was achieved using a MicroPulse electroporator (BioRad) at setting EC1. Standard curves for qPCR were prepared from the serial dilution of plasmid DNA recovered using the Zippy Plasmid Miniprep kit (Zymo Research) following Qubit fluorometirc quantification (ThermoFisher Scientific). The genome of Methylacidiphilum sp. RTK17.1 was sequenced using the PacBio RS platform (Macrogen). Genome assembly was performed, de novo, via the SMRT Portal using hierarchical genome-assembly process (HGAP) pipeline (49). A high quality de novo assembly was generated from three SMRT sequencing runs the HGAP.3 protocol. A pre-assembly genome construct was assembled using Celera Assembler and then polished using Quiver (49). Quality assessment of the de novo assembly was conducted using BridgeMapper. Gene prediction was performed using Quiver and genome annotation was completed using the integrated microbial genomes database pipeline (50). Soil microbial community determination The V4 region of the 16S rRNA gene was sequenced using the Illumina MiSeq® System. Sequences were analysed using UPARSE pipeline for quality filtering and clustering (51). Briefly, paired end reads were merged and those less than 200 or 350 bp were discarded, for 515f/806r and 341/785r primer amplicons respectively (Table S1). Quality filtering was applied with a maximum expected error value of 1. Retained sequences were dereplicated and unique sequences removed. Next, reads were clustered to 97 % similarity, which includes a chimera check, and a de novo database was created of representative operational taxonomic units (OTUs). Original pre-filtered sequences were mapped to these OTUs, and taxonomy was assigned using the Ribosomal Database Project classifier (with a minimum confidence score of 0.5) (52) against the GreenGenes 16S rRNA database (13_8 release) (53). Final

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count number was 130,289 reads across all depths for the 515f/806r primer set, with a total of 48 differential OTUs (492,588 for 341/785r, with 107 OTUs). Static liquid fed-batch bioreactor cultivations Methylacidiphilum sp. RTK17.1 was cultivated to stationary phase for subsequent hydrogenase activity and oxygen respiration measurements in a semi-continuous fed-batch bioreactor (New Brunswick; volume 11, pH control 2.5, temp 50 °C, agitation 100 rpm) in an artificial headspace composed of 10 % methane, 10 % hydrogen, 2 0 % O₂, 40 % carbon dioxide (v/v, balance N₂; flow rate 60 ml min⁻¹) equipped with headspace recirculation and automated sampling via a 490 MicroGC (Agilent Technologies, Santa Clara, CA). Gas mixtures were supplied for 2 min every hour at a rate of 60 ml min⁻¹. Oxygen consumption experiments. Oxygen consumption experiments were performed on cell suspensions of Methylacidiphilum sp. RTK17.1 to determine the influence of endogenous glycogen catabolism on O₂ dependent hydrogenase measurements. For these experiments, 2 mL cells (OD₆₀₀ 1.0) were added to a Clarke-type oxygen electrode and incubated at 50 °C for up to 12 min without the addition of exogenous energy sources. Cell suspensions were treated with the protonophore (1 µM) CCCP, 1 mM lodoacetamide (an inhibitor of glycolysis) and 1 mM potassium cyanide (KCN) to determine whether observed rates oxygen consumption were a consequence of glycogen catabolism.

Figure S1. The classification and phylogeny of the [NiFe] hydrogenase large subunit identified in the genomes of methanotrophic bacteria.

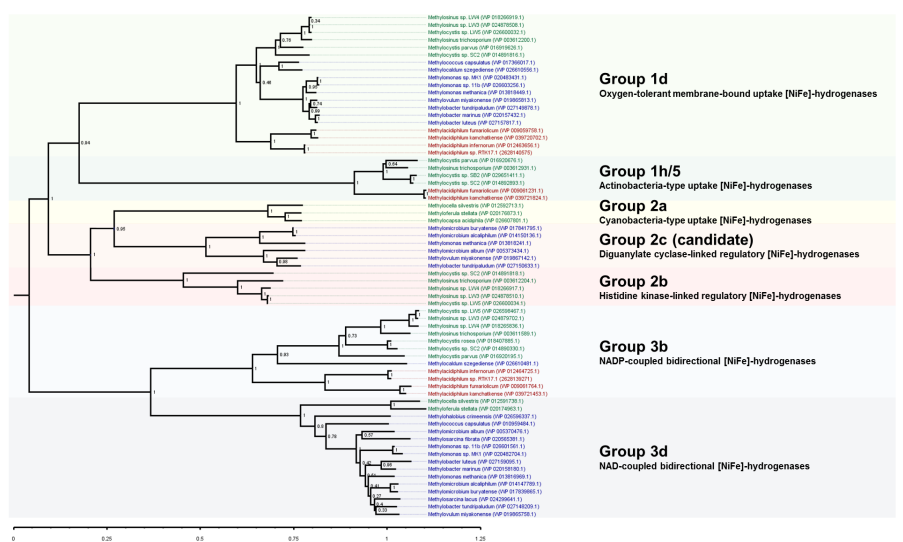


Figure S2. Oxidation of CH₄ and H₂ by surface Rotokawa soils (< 10 cm depth) incubated at mesophilic (37 °C) and thermophilic (50 °C) temperatures. Soils samples (1 g) were collected and incubated following addition of A) \sim 400 ppmv CH₄ and B) \sim 300 ppmv H₂ into the headspace. The average and standard deviation of triplicate samples is shown.

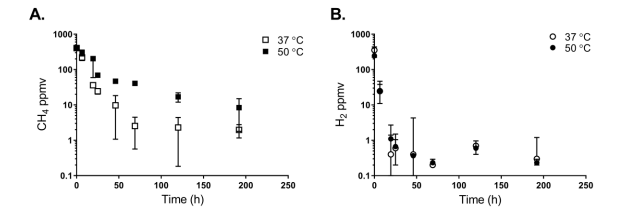


Figure S3. Batch culture growth experiments and observations of methane and hydrogen oxidation for *Methylacidiphilum* sp. RTK17.1 following treatment with acetylene. (A) Rates of hydrogen oxidation are greatest during methanotrophic growth (red circle) but are also measurable in the presence of 4 % (v/v) acetylene gas (an inhibitor of pMMO; green triangle) and in the absence of methane (blue square). Black triangles with dashed lines denote negative controls. Observed rates of methane and hydrogen oxidation decline as substrates are exhausted. Error bars shown represent the standard deviation of biological triplicate samples. (B) Methanotrophic growth of *Methylacidiphilum* sp. RTK17.1 is enhanced by the addition of 1 % H₂ (v/v). Differences in growth observed (as determined by total protein) with and without hydrogen addition were analysed following 1 week incubation (50 °C) in 20 paired samples by one-tailed t-test assuming Gaussian distribution and determined to be significant (*p-value* 0.007, *df*= 19, SEM error bars are shown).

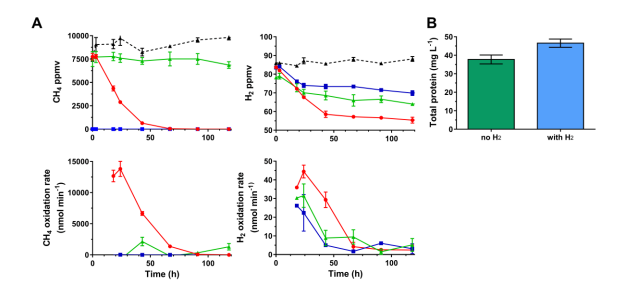


Figure S4. Reverse Transcriptase PCR of total mRNA isolated from stationary phase cultures of *Methylacidiphilum* RTK17.1 confirms the expression of the membrane bound Group 1d [NiFe] hydrogenase. Only faint amplification of cDNA encoding the cytosolic Group 3b enzyme was observed.



Figure S5. Treatment of cell suspensions of *Methylacidiphilum sp.* RTK17.1 with the uncoupler carbonyl cyanide *m*-chlorophenyl hydrazone (CCCP), the glycolytic inhibitor iodoacetamide (IAA) and the respiratory chain inhibitor potassium cyanide (KCN). This indicates that observed oxygen consumption, in the absence of exogenous energy sources (H₂ or CH₄), is likely a consequence of endogenous glycogen catabolism.

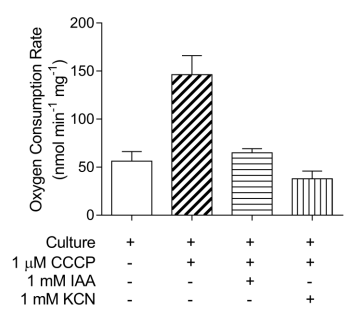


Figure S6. Treatment of concentrated cell suspensions of *Methylacidiphilum sp.* RTK17.1 with the ionophores nigericin (Nig) and valinomycin (Val) indicates hydrogenase activity is sensitive to the magnitude of the proton- and chargegradients of the proton-motive force. The reduction in rate of hydrogen oxidation observed when treated with carbonyl cyanide *m*-chlorophenyl hydrazone (CCCP) is likely a consequence of secondary intracellular acidification.

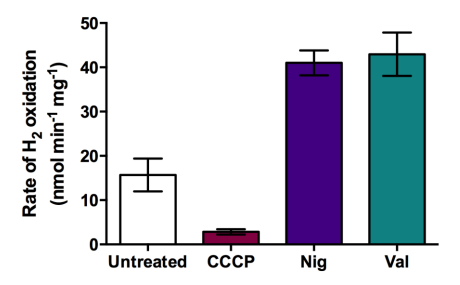
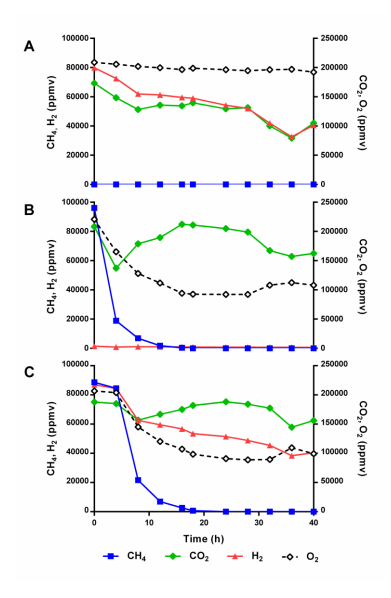


Figure S7. *Methylacidiphilum* sp. RTK17.1 oxidises hydrogen (H₂) and fixes carbon dioxide (CO₂) in the absence of methane (CH₄). In static liquid fed-batch bioreactor experiments, stationary-phase cultures were incubated (50 °C) in a nitrogen headspace with mixing ratios (v/v) as follows: (A) 0 % CH₄, 9 % H₂, 19 % CO₂, 20 % O₂; (B): 9 % CH₄, 0 % H₂, 19 % CO₂, 20 % O₂; (C): 9 % CH₄, 9 % H₂, 19 % CO₂, 20 % O₂. In the absence of CH₄ or H₂ addition, Argon (10 %) supplementation was used to maintain equivalent headspace mixing ratios of the other gases between experiments. We did not determine whether CO₂ fixation in the absence of CH₄ was driven by H₂ oxidation or instead metabolism of endogenous glycogen reserves.



Target Locus		Amplicon size (bp)	Primer name	Sequence	References
hyd1d	Minf_1320	187	qMinf_1320_fwd	TGCACCTGACACAAATGGTT	This study
	Minf_1320		qMinf_1320_rvs	TGCTTCAGTCGATTTTGCAC	
hyd3b	Minf_2390	241	qMinf_2390_fwd	CCTGGAAGAAGAGCTGTTGG	This study
	Minf_2390		qMinf_2390_rvs	ACTTTCCAGGGAGGCAAAAT	
ртоА	Minf_1507	208	qMinf_1507_fwd	ACTTCGTTGTGACCGCTCTT	This study
	Minf_1507		qMinf_1507_rvs	GCAAAGCTTCTCATCGTTCC	
ртоА	Minf_1510	172	qMinf_1510_fwd	AGAACTTTGGGGTTGGCTTT	This study
	Minf_1510		qMinf_1510_rvs	CCTCATCGTTCCCGTTTCTA	
ртоА	Minf_1590	202	qMinf_1590_fwd	CGCATATGCTTTTGCTTTGA	This study
	Minf_1590		qMinf_1590_rvs	CGGTCCTAAAGCTCCTCCTT	
16S rRNA		1483	9f	GAGTTTGATCMTGGCTCAG	(46)
			1492r	TACCTTGTTACGACTT	
16S rRNA		291	Uni515f	GTGYCAGCMGCCGCGGTAA	(54)
			Uni806r	GGACTACNVGGGTWTCTAAT	
16S rRNA		444	Bac341f	CCTACGGGNGGCWGCAG	(55)
			Bac785r	GACTACHVGGGTATCTA ATCC	

Table S2. Genes involved in one-carbon and hydrogen metabolism, as well as glycogen storage and utilization in *Methylacidiphilum* sp. RTK17.1. Amino acid orthologs to *Methylacidiphilum infernorum* V4 are shown.

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Gene	Predicted Function	Gene ID *	M. infernorum ortholog	% ID
Methane	oxidation to methanol			
pmoC1	particulate methane monooxygenase, C subunit	KU509367	WP_012463847.1	99
pmoA1	particulate methane monooxygenase, A subunit	KU509368	ABX56601.1	100
pmoB1	particulate methane monooxygenase, B subunit	KU509369	WP_012463845.1	99
pmoC2	particulate methane monooxygenase, C subunit	KU509373	WP_048810233.1	99
pmoA2	particulate methane monooxygenase, A subunit	KU509371	WP_012463843.1	99
pmoB2	particulate methane monooxygenase, B subunit	KU509372	WP_012463842.1	99
pmoC2	particulate methane monooxygenase, C subunit	KU509370	WP_012463844.1	99
ртоС3	particulate methane monooxygenase, C subunit	KU509318	WP_012463927.1	99
pmoA3	particulate methane monooxygenase, A subunit	KU509319	WP_012463926.1	99
ртоВ3	particulate methane monooxygenase, B subunit	KU509320	WP_012463925.1	97
Methano	ol oxidation to formate (11)			
xoxF	methanol dehydrogenase, large subunit	KU509410	WP_012463329.1	99
xoxJ [#]	unknown function, involved in methanol oxidation	Ga0079954 _12828	WP_012463332.1	99
тхаВ	transcriptional regulator	KU509376	WP_012463785.1	100
moxY	methanol utilization control sensor protein	KU509377	WP_012463784.1	100
mxaR	ATPase involved in methanol oxidation	KU509402	WP_012463503.1	97
no homo	logs of mxal mxaS mxaA mxaC mxaK mxaL mx	xaD mxaE mxal	H mxaW mxaG	
Tetrahyd	Irofolate synthesis			
folK	2-amino-4-hydroxy-6- hydroxymethyldihydropteridine diphosphokinase	KU509446	WP_048810035.1	97
foIP	dihydropteroate synthase	KU509345	WP_012464234.1	99
foIP	dihydropteroate synthase	KU509478	WP_048810563.1	98
foIC	bifunctional folylpolyglutamate synthase/dihydrofolate synthase	KU509444	WP_012462725.1	99
Formate	oxidation to CO ₂			
fdsG	formate dehydrogenase, gamma subunit	KU509398	WP_048810187.1	97
fdsB	formate dehydrogenase, beta subunit	KU509399	WP_012463566.1	100
fdsA	formate dehydrogenase, alpha subunit	KU509400	WP_048810186.1	99

fdsD	formate dehydrogenase delta subunit	KU509401	WP_048810185.1	94
hpr	formate dehydrogenase	KU509384	WP_012463659.1	99
ehrB	putative formate-dependent oxidoreductase complex subunit	KU509439	WP_012462850.1	99
ehrD	putative formate-dependent oxidoreductase complex subunit	KU509438	WP_012462851.1	99
ehrF	putative formate-dependent oxidoreductase complex subunit	KU509436	WP_012462853.1	99
ehrA	putative formate-dependent oxidoreductase complex subunit	KU509437	WP_012462852.1	99
ehrE	putative formate-dependent oxidoreductase complex subunit	KU509449	WP_012462854.1	100
ehrG	putative formate-dependent oxidoreductase complex subunit	KU509435	WP_048810433.1	97
Methylar	nine oxidation genes			
таиВ	methylamine dehydrogenase large subunit	KU509332	WP_012464333.1	98
mauE	methylamine utilization protein	KU509331	WP_012464334.1	99
mauD	methylamine utilization protein	KU509330	WP_012464335.1	100
mauA [#]	methylamine dehydrogenase small subunit	Ga0079954 _121718	WP_048810312.1	99
mauG	methylamine utilization protein, cytochrome c peroxidase	KU509329	ACD84055.1	99
no homo	logs of mauF mauC mauJ mauL mauM mauN			
Hydroger	n metabolism			
hyaF	hydrogenase-1 operon protein	KU509388	WP_012463654.1	99
hyaC	oxygen-tolerant membrane-bound [NiFe] - hydrogenase, cytochrome <i>b</i> subunit	KU509387	WP_012463655.1	99
hyaB	oxygen-tolerant membrane-bound [NiFe] hydrogenase, large subunit	KU509386	WP_012463656.1	99
hyaA	oxygen-tolerant membrane-bound [NiFe] hydrogenase, small subunit	KU509385	WP_048810203.1	99
hyhB	NADP-coupled cytosolic bidirectional hydrogenase, FeS subunit	KU509477	WP_012464721.1	99
crp	cyclic nucleotide-binding domain- containing protein	KU509476	WP_012464722.1	97
hyhG	NADP-coupled cytosolic bidirectional hydrogenase, diaphorase subunit NAD(P)H-flavin reductase	KU509475	WP_012464723.1	98
hyhS	NADP-coupled cytosolic bidirectional hydrogenase, small subunit	KU509474	WP_012464724.1	98
hyhL	NADP-coupled cytosolic bidirectional hydrogenase, large subunit	KU509473	WP_012464725.1	98
hypB	hydrogenase maturase protein, Ni storage	KU509355	WP_012464135.1	99
hypC	hydrogenase maturase protein, Fe- $(CN^{-})_{2}$ - CO insertion	KU509357	WP_012464132.1	100
hypD	hydrogenase maturase protein, Fe- $(CN^{-})_{2}$ - CO insertion	KU509358	WP_012464131.1	99
hypE	hydrogenase maturase protein, CN ⁻ ligand	KU509359	WP_012464130.1	99

hypF	hydrogenase maturase protein, CN ⁻ ligand biosynthesis	KU509356	WP_012464133.1	99
Copper h	omeostasis			
cueR	copper resistance operon regulatory protein	KU509440	WP_012462789.1	99
copA	Cu(I)-translocating P-type ATPase/ multicopper oxidase, copper-binding site	KU509375	gb ABX56608.1	99
cueO	multicopper oxidase family protein	KU509374	WP_048810228.1	99
cusA	copper efflux pump	KU509461	WP_012462437.1	99
cusB [#]	copper efflux pump membrane fusion protein	Ga0079954 _11259	WP_012462436.1	98
cusS	copper sensor histidine kinase	KU509433	WP_012462941.1	99
cusR	DNA binding copper response regulator	KU509432	WP_048810437.1	99
Coenzym	e PQQ synthesis			
pqqB	coenzyme PQQ synthesis protein B	KU509397	WP_012463570.1	99
pqqC	coenzyme PQQ synthesis protein C	KU509396	WP_012463571.1	99
pqqD	coenzyme PQQ synthesis protein D	KU509395	WP_048810188.1	98
pqqD	coenzyme PQQ synthesis protein D	KU509434	WP_012462864.1	97
pqqE	coenzyme PQQ synthesis protein E	KU509394	WP_048810189.1	99
No homo	logs of pqqG pqqF pqqA			
Glycogen	synthesis			
mdoG	glycans biosynthesis protein	KU509364	WP_012463952.1	98
mdoG	glycans biosynthesis protein	KU509365	WP_012463951.1	99
	glucan elongaton module (mdoH-like)	KU509366	WP_048810251.1	99
GH57	glycosyl hydrolase family 57	KU509382	WP_012463665.1	99
amyA	alpha amylase, catalytic domain	KU509383	WP_012463664.1	98
glgA	glycogen synthase	KU509360	WP_012464092.1	99
glgA	glycogen synthase (ADP-glucose)	KU509479	WP_048810560.1	98
glgB	1,4-alpha-glucan branching enzyme	KU509409	WP_012463357.1	99
glgC	glucose-1-phosphate adenylyltransferase	KU509460	WP_048810004.1	99
glgP	starch phosphorylase	KU509451	WP_012462597.1	99
glgP	starch phosphorylase	KU509431	WP_012462995.1	99
glgP	starch phosphorylase	KU509429	WP_012463128.1	99
gdb1	glycogen debranching enzyme (alpha-1,6-glucosidase)	KU509417	WP_012463227.1	99
rfaG	glycosyltransferase	KU509418	WP_012463226.1	99
glcD	glycolate oxidase	KU509414	WP_048810142.1	98
glcD	glycolate oxidase FAD binding subunit	KU509413	WP_012463233.1	98
асеВ	malate synthase	KU509415	WP_012463229.1	98
aceA	isocitrate lyase	KU509416	WP_048810457.1	98
manB	phosphoglucomutase	KU509328	WP_012464463.1	99
Calvin-Be	enson Cycle			
cbbS	ribulose 1,5-bisphosphate carboxylase, small subunit	KU509390	WP_012463599.1	98
cbbL	ribulose 1,5-bisphosphate carboxylase, large subunit	KU509389	WP_012463600.1	99
cbbX	probable Rubsico expression protein CbbX	KU509391	WP_012463598.1	99

pgk	phosphoglycerate kinase	KU509406	WP_048810158.1	98
gapA	glyceraldehyde-3-phosphate dehydrogenase	KU509405	WP_012463370.1	99
fbaA	putative fructose-bisphosphate aldolase	KU509327	WP_048810542.1	98
fbaA2	fructose-bisphosphate aldolase, class II	KU509448	WP_012462629.1	100
fbp	fructose-1,6-bisphosphatase I	KU509361	WP_012464020.1	99
glpX	fructose-1,6-bisphosphatase II	KU509381	WP_012463695.1	100
tktB	transketolase	KU509393	WP_012463596.1	99
xfp	xylulose-5-phosphate/fructose-6- phosphate phosphoketolase	KU509412	WP_012463240.1	99
tpiA	triosephosphate isomerase	KU509407	WP_012463368.1	98
rpiB	ribose 5-phosphate isomerase B	KU509459	WP_012462539.1	99
prkB	phosphoribulokinase	KU509392	WP_012463597.1	100
rpe	ribulose-phosphate 3-epimerase	KU509354	WP_012464174.1	99
TCA cycle				
gltA	citrate synthase	KU509447	WP_012462674.1	99
acnA	aconitate hydratase	KU509408	WP_012463364.1	99
icd	isocitrate dehydrogenase	KU509411	WP_012463276.1	99
sucA	2-oxoglutarate dehydrogenase, E1 component	KU509464	WP_048810403.1	98
sucB	2-oxoglutarate dehydrogenase, E2 component, dihydrolipoamide succinyltransferase	KU509463	WP_012462410.1	98
lpdA	pyruvate/2-oxoglutarate dehydrogenase, E3 component, dihydrolipoamide dehydrogenase	KU509462	WP_012462411.1	99
lpdA	pyruvate/2-oxoglutarate dehydrogenase, E3 component, dihydrolipoamide dehydrogenase	KU509362	WP_012463971.1	98
sucD	succinyl-CoA synthetase subunit alpha	KU509403	WP_012463383.1	99
sucC	succinyl-CoA synthetase subunit beta	KU509404	WP_048810162.1	99
sdhB	succinate dehydrogenase, catalytic subunit	KU509442	WP_012462728.1	99
sdhA	succinate dehydrogenase, flavoprotein subunit	KU509441	WP_012462729.1	98
sdhC	succinate dehydrogenase, cytochrome b subunit	KU509443	WP_012462727.1	98
fumC	fumarase	KU509428	WP_012463130.1	100
mdh	malate dehydrogenase	KU509380	WP_048810212.1	99
ATP Synth	nase			
atpD	F-type H ⁺ -transporting ATPase subunit beta	KU509426	WP_048810122.1	99
atpC	F-type H ⁺ -transporting ATPase subunit epsilon	KU509425	WP_012463177.1	99
	ATP synthase protein I	KU509424	WP_012463178.1	98
atpB	ATP synthase F0 subcomplex A subunit	KU509423	WP_012463179.1	98
atpE	F-type H ⁺ -transporting ATPase subunit c	KU509422	WP_012463180.1	98
atpF	ATP synthase F0 subcomplex B subunit	KU509421	WP_012463181.1	99

atpA	F-type H+-transporting ATPase subunit	KU509420	WP_012463182.1	99
	alpha		W.D. 0494594994	400
atpG	F-type H+-transporting ATPase subunit gamma	KU509419	WP_012463183.1	100
atpH	F-type H⁺-transporting ATPase subunit delta	KU509469	WP_012464754.1	99
atpB	F-type H ⁺ -transporting ATPase subunit a	KU509472	WP_012464751.1	99
atpE	F-type H ⁺ -transporting ATPase subunit c	KU509471	WP_012464752.1	100
atpF	F-type H ⁺ -transporting ATPase subunit b	KU509470	WP_048810391.1	99
atpA	F-type H ⁺ -transporting ATPase subunit alpha	KU509468	WP_012464755.1	99
atpG	F-type H⁺-transporting ATPase subunit gamma	KU509467	WP_012464756.1	99
atpD	F-type H⁺-transporting ATPase subunit beta	KU509466	WP_012464757.1	100
atpC	F-type H*-transporting ATPase subunit epsilon	KU509465	WP_012464758.1	100
NADH de	hydrogenase			
пиоВ	NADH-quinone oxidoreductase subunit B	KU509458	WP_012462551.1	99
nuoA	NADH-quinone oxidoreductase subunit A	KU509480	WP_012464634.1	99
nuoC	NADH-quinone oxidoreductase subunit C	KU509457	WP_012462552.1	100
nuoD	NADH-quinone oxidoreductase subunit D	KU509456	WP_012462553.1	99
nuoE	NADH-quinone oxidoreductase subunit E	KU509455	WP_012462554.1	99
nuoF	NADH-quinone oxidoreductase subunit F	KU509454	WP_012462555.1	99
nuoG	NADH-quinone oxidoreductase subunit G	KU509453	WP_012462556.1	99
nuoH	NADH-quinone oxidoreductase subunit H	KU509452	WP_012462557.1	99
nuol	NADH-quinone oxidoreductase subunit I	KU509321	WP_048810348.1	99
nuoJ	NADH-quinone oxidoreductase subunit J	KU509322	WP_048810347.1	99
nuoK	NADH-quinone oxidoreductase subunit K	KU509323	WP_012464548.1	100
nuoL	NADH dehydrogenase subunit L	KU509324	WP_012464547.1	98
nuoM	NADH-quinone oxidoreductase subunit M	KU509325	WP_012464546.1	99
nuoN	NADH dehydrogenase subunit N	KU509326	WP_012464545.1	99
Cytochro	me C oxidase and Complex III (ACIII) (40)			
cyoE	protoheme IX farnesyltransferase	KU509378	WP_012463763.1	99
	cytochrome c oxidase assembly protein			
ctaA	subunit 15	KU509379	WP_012463762.1	99
суоС	cytochrome c oxidase subunit 3	KU509342	WP_012464286.1	99
cyoA	cytochrome c oxidase subunit 2	KU509344	WP_012464280.1	100
суоВ	cytochrome c oxidase subunit 1	KU509341	WP_012464287.1	99
	caa₃-type oxidase, subunit IV	KU509343	WP_048810305.1	98
4	cytochrome c oxidase, <i>cbb</i> ₃ -type, subunit	VI IE 002 40	WD 0424642004	100
cccA	III	KU509340	WP_012464288.1	100
cytC	cytochrome c	KU509481	WP_012463333.1	97
ccoO	cytochrome c oxidase <i>cbb</i> ₃ -type subunit 2	KU509339	WP_012464289.1	99
ccoN	cytochrome c oxidase <i>cbb</i> ₃ -type subunit 1	KU509338	WP_048810306.1	99
	cytochrome c & quinol oxidase polypeptide I	KU509482	WP 012462607.1	91
	heme/copper-type cytochrome/quinol		_: == ::3001	
	oxidase, subunit 1	KU509483	WP_012462607.1	99

ActG*	Hypothetical membrane-associated protein	Ga0079954 _121677	WP_012464292.1	99
ActF*	quinol:cytochrome c oxidoreductase	KUE00227	WD 012464202 1	99
	quinone-binding subunit 2	KU509337	WP_012464293.1	99
cccA –	cytochrome c oxidase, <i>cbb</i> ₃ -type, subunit	W.1500336	WD 0424642044	00
ActE*	III	KU509336	WP_012464294.1	99
	Hypothetical membrane-associated	Ga0079954		
ActD*	protein	_121680	WP_012464295.1	100
	prokaryotic molybdopterin-containing			
ActC*	oxidoreductase family, membrane subunit	KU509335	WP_012464296.1	99
	prokaryotic molybdopterin-containing			
hybA –	oxidoreductase family, iron-sulfur binding			
ActB*	subunit	KU509334	WP_012464297.1	99
ActA*	cytochrome c7	KU509333	WP_012464298.1	99
Nitrogen	fixation			
nifH	Mo-nitrogenase iron protein subunit NifH	KU509347	WP_012464212.1	99
nifV	homocitrate synthase	KU509346	WP_012464213.1	99
nifD	Mo-nitrogenase MoFe protein subunit NifD precursor	KU509348	WP_012464210.1	99
nifK	Mo-nitrogenase MoFe protein subunit NifK	KU509349	WP_048810527.1	99
nifE	nitrogenase molybdenum-cofactor synthesis protein	KU509350	WP_012464208.1	99
nifN	nitrogenase molybdenum-iron protein NifN	KU509351	WP_048810526.1	99
nifX	nitrogen fixation protein NifX	KU509352	WP 012464206.1	100

[#] nucleotide gene sequences contain nucleotide do not have accession numbers at the time of submission. * Alternative Complex III genes - see reference (40)