Mixotrophy drives niche expansion of verrucomicrobial methanotrophs

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Aerobic methanotrophic bacteria have evolved a specialist lifestyle dependent on consumption of methane and other short-chain carbon compounds. However, their apparent substrate specialism runs contrary to the high relative abundance of these microorganisms in dynamic environments, where the availability of methane and oxygen fluctuates. In this work, we provide in situ and ex situ evidence that verrucomicrobial methanotrophs are mixotrophs. Verrucomicrobia-dominated soil communities from an acidic geothermal field in Rotokawa, New Zealand rapidly oxidised methane and hydrogen simultaneously. We isolated and characterised a verrucomicrobial strain from these soils, Methylacidiphilum sp. RTK17.1, and showed that it constitutively oxidises molecular hydrogen. Genomic analysis confirmed that this strain encoded two [NiFe]-hydrogenases (group 1d and 3b), and biochemical assays revealed that it used hydrogen as an electron donor for aerobic respiration and carbon fixation. While the strain could grow heterotrophically on methane or autotrophically on hydrogen, it grew optimally by combining these metabolic strategies. Hydrogen oxidation was particularly important for adaptation to methane and oxygen limitation. Complementary to recent findings of hydrogenotrophic growth by Methylacidiphilum fumariolicum SolV, our findings illustrate that verrucomicrobial methanotrophs have evolved to simultaneously utilise hydrogen and methane from geothermal sources to meet energy and carbon demands where nutrient flux is dynamic. This mixotrophic lifestyle is likely to have facilitated expansion of the niche space occupied by these microorganisms, allowing them to become dominant in geothermally influenced surface soils. Genes encoding putative oxygen-tolerant uptake [NiFe]-hydrogenases were identified in all publicly available methanotroph genomes, suggesting hydrogen oxidation is a general metabolic strategy in this guild.

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Introduction

Aerobic methane-oxidising bacteria (methanotrophs) consume the potent greenhouse gas methane (CH4) (Kirschke et al., 2013). They serve as the primary biological sink of atmospheric methane (~30 Tg annum⁻¹) (Hanson and Hanson, 1996) and, together with anaerobic methane-oxidising archaea, also capture the majority of biologically and geologically produced CH4 before it enters the atmosphere (Oremland and Culbertson, 1992). Relative to their global impact as greenhouse gas mitigators, aerobic methanotrophs exhibit low phylogenetic diversity and are presently limited to 26 genera in the Alphaproteobacteria and Gammaproteobacteria (Euzéby, 1997), two candidate genera in the phylum Verrucomicrobia (Op den Camp et al., 2009; van Veen et al., 2014), and two representatives of candidate phylum NC10 (Ettwig et al., 2010; Haroon et al., 2013). Reflecting their aerobic methylotrophic lifestyle, methanotrophs thrive in oxic–anoxic interfaces where CH4 fluxes are high, including peat bogs, wetlands, rice paddies, forest soils and geothermal habitats (Singh et al., 2010; Knief, 2015). However, they also exist within soil and marine ecosystems where CH4 and oxygen (O2) are more variable (Knief et al., 2003; Tavormina et al., 2010; Knief, 2015).
Based on current paradigms, aerobic methanotrophs are thought to primarily grow on one-carbon (C1) compounds in the environment (Dedysh et al., 2005). All species can grow by oxidising CH4 to methanol via particulate or soluble methane monooxygenase. They subsequently oxidise methanol to carbon dioxide (CO2), yielding reducing equivalents (e.g. NADH) for respiration and biosynthesis. Proteobacterial methanotrophs generate biomass by assimilating the intermediate formaldehyde via the ribulose monophosphate or serine pathways (Hanson and Hanson, 1996). In contrast, verrucomicrobial methanotrophs oxidise methanol directly to formate (Keltjens et al., 2014) and generate biomass by fixing CO2 via the Calvin–Benson–Bassham cycle (Khadem et al., 2011). While these specialist C1-based metabolisms are thought to be the primary growth strategy under optimal conditions (i.e. CH4 and O2 replete conditions), they would presumably be less effective in dynamic environments where CH4 and oxidant availability are likely to fluctuate.

To add to this complexity, the methane monoxygenase reaction (CH4+O2+[NAD(P)H+H+]/QH2 → CH3OH+NAD(P)(+)/Q+H2O) (Hakemian and Roszenwieg, 2007) is metabolically demanding, given it requires simultaneous sources of CH4, endogenous reductant (NAD(P)H or quinol) and exogenous O2 to proceed. Methanotrophs therefore must carefully allocate resources to meet carbon, energy and reductant demands (Hanson and Hanson, 1996). This complex balancing act provokes that, in order to be viable in environments limited for CH4 and O2 gases (Knief et al., 2003; Tavormina et al., 2010), methanotrophs should be able to supplement C1 usage with other energy-yielding strategies.

Recent pure culture studies have provided evidence that CH4-oxidising bacteria are indeed more metabolically versatile than previously thought. A minority of conventional methanotrophs can meet energy demands by oxidising the trace concentrations of CH4 (1.8 ppmv) found in the atmosphere (Kolb et al., 2005; Ho et al., 2013; Cai et al., 2016). Contrary to the long-held paradigm that methanotrophs are obligate methylotrophs, species from three alphaproteobacterial genera have been shown to grow on simple organic acids, alcohols and short-chain alkane gases (Dedysh et al., 2005; Crombie and Murrell, 2014). Most recently, it has been shown that some methanotrophs are not exclusive heterotrophs: the verrucomicrobium Methylociacidiphilum fumariolicum SolV can sustain chemolithoautotrophic growth on molecular hydrogen (H2) through the activity of two [NiFe]-hydrogenases (Mohammadi et al., 2016). Proteobacterial methanotrophs can also consume H2, though to date this process has only been reported as providing reductant to supplement methanotrophic growth (Chen and Yoch, 1987; Shah et al., 1995; Hanczár et al., 2002). Our recent findings demonstrating a widespread distribution and diversity of hydrogenases in aerobic bacteria, in specific methanotrophs (Greening et al., 2014a,b, 2015), led us to surmise that H2 metabolism could serve a multifaceted role in adaptation of methanotrophic bacteria to their environment. Specifically, H2 may serve as an important electron donor for the organism to meet carbon, energy and reductant demands in response to fluctuations in CH4 and oxidant availability.

In this work, we addressed this hypothesis by conducting an interdisciplinary investigation of the role of H2 in defining the physiology and ecology of verrucomicrobial methanotrophs. Evidence obtained from in situ field studies indicate that Verrucomicrobia simultaneously oxidised CH4 and H2 in geothermally heated soils in Rotokawa, New Zealand, suggesting they are mixotrophic with respect to energy metabolism. Pure culture studies on a verrucomicrobium representative isolated from this site confirmed that the microorganism grew most efficiently through a mixotrophic lifestyle and depended on H2 consumption to acclimate to fluctuations in CH4 and O2 availability. Integrating these findings with genome surveys, we propose that H2 oxidation expands the ecological niche of methanotrophs, enabling them to meet energy and biomass demands in dynamic environments where O2 and CH4 concentrations are variable. We provide evidence that, while methanotrophic bacteria are often pervasively viewed as C1 specialists, their niche space is likely broader than previously recognised. Combining heterotrophic and lithotrophic electron donors allows for a more flexible growth/survival strategy with clear ecological benefits (Semrau et al., 2011).

**Materials and methods**

**Environmental sampling**

Soil samples (~50 g) were collected every 10 cm from the surface of the Rotokawa sampling site (38°37′30.8″ S, 176°11′55.3″E) to a maximum depth of 50 cm. Soil temperatures were measured in the field using a 51II single input digital thermometer (Fluke, Everett, WA, USA). The pH of the soil (1 g in 10 ml of dH2O) was measured upon returning to the laboratory using a model HI11310 pH probe (Hanna Instruments, Woonsocket, RI, USA). Soil gas samples were collected every 10 cm using a custom-built gas-sampling probe equipped with a 1 l gas-tight syringe (SGE Analytical Science, Melbourne, VIC, Australia). Gas samples were collected and stored at 25 °C in 50 ml Air & Gas Sampling Bags (Calibrated Instruments, McHenry, MD, USA) and were processed within 48 h on a 490 MicroGC (Agilent Technologies, Santa Clara, CA, USA) equipped with Molecular Sieve 5A with a heated injector (50 °C, back-flush at 5.10 s, column at 90 °C, 150 kPa), a PoraPak Q column with a heated injector (50 °C, no back-flush, column at 70 °C, 50 kPa) and a 5CB column with a heated injector (50 °C, no back-flush, column at 80 °C, 150 kPa). CH4 and H2 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 headspaces were amended with 300 ppmv H₂ or 400 ppmv CH₄. Headspaces CH₄ and H₂ mixing ratios were measured with a PeakPerformer gas chromatograph (Peak Laboratories, Mountain View, CA, USA) equipped with a flame ionising detector (FID: CH₄) and a PP1 Gas Analyzer (Peak Laboratories, Mountain View, CA, USA) equipped with a reducing compound photometer (RCP: H₂).

Isolation and cultivation of Methylacidiphilum sp.

RTK17.1

Soil samples (1 g) from the first 10 cm of soil at the Rotokawa sampling site were inoculated into serum bottles containing 50 ml media (pH 2.5). All cultivations were performed in a V4 mineral medium as described previously (Dunfield et al., 2007) but with the addition (0.2 μm) of rare earth elements lanthanum and cerium (Pol et al., 2014). CH₄ (10% v/v) and CO₂ (1%) were added to an air headspace and samples were incubated at 60°C with shaking 150 r.p.m. CH₄ in the headspace was monitored with a PeakPerformer gas chromatograph (Peak Laboratories) equipped with an FID. Following several passages (10% v/v) into liquid media, enrichments were transferred onto solid media. Following several weeks incubation (60°C) single colonies were re-streaked before being transferred back into liquid media. Isolate identity was confirmed via sequencing of the 16S rRNA gene (Macrogen, South Korea) using bacterial 9f/1492r primers (Weisburg et al., 1991).

Bioreactor and batch cultivation

Methylacidiphilum sp. RTK17.1 was cultivated to the stationary phase for subsequent hydrogenase activity and oxygen respiration measurements in a semi-continuous static-liquid fed-batch bioreactor (New Brunswick; volume 1 l, pH control 2.5, temp 50°C, agitation 100 r.p.m.) in an artificial headspace composed of 10% CH₄, 10% H₂, 20% O₂, 40% CO₂ (v/v), balance N₂; flow rate 60 ml min⁻¹) equipped with headspace recirculation and automated sampling via a 490 MicroGC (Agilent Technologies). Gas mixtures were supplied for 2 min every hour at a rate of 60 ml min⁻¹. Gas feeds for the bioreactor, batch and chemostat cultivation-based experimental work are presented as headspace compositions. For batch culture experiments, 350 ml cultures (in triplicate) were incubated in 1 l rubber-stoppered Schott bottles. The headspace of the bottles was amended with different mixing ratios of H₂, CH₄, O₂ and CO₂ gas, as described in figure legends. Acetylene gas was added in some experiments (4% v/v) to inhibit MMO activity as previously described (Bédard and Knowles, 1989). Finally, to determine if growth was enhanced in the presence of H₂, 20 paired cultures (n = 40) of Methylacidiphilum sp. RTK17.1 were incubated with or without 1% (v/v) H₂ in an air headspace supplemented with (v/v) 10% CH₄ and 1% CO₂. Cultures were incubated in a custom test tube oscillator (agitation 1.2 Hz; Terratec, Hobart, TAS, Australia). Following 7 days incubation, total protein was determined by the Bradford assay (Bradford, 1976). Statistical significance of observed differences of growth yields was determined using a Student’s t-test (α = 0.05). Headspace mixing ratios of H₂ and CH₄ were monitored throughout batch experiments by GC as described above.

Chemostat cultivation

Chemostat cultivation of Methylacidiphilum sp. RTK17.1 was performed to investigate the influence of H₂ on growth under O₂-replete and O₂-limiting conditions. A 1 l bioreactor (BioFlo 110; New Brunswick Scientific, Edison, NJ, USA) was used for these studies. Cultures were incubated at 50°C and pH 2.5 with continuous stirring (800 r.p.m.). 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 O₂ was monitored using an InPro 6810 Polarographic Oxygen Sensor (Mettler-Toledo, Columbus, OH, USA). 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% CH₄ and 26% CO₂ for all experiments. For O₂-replete and O₂-limiting conditions, influent O₂ was supplied at (v/v) 14.1% and 3.5%, respectively. Within respiring cultures, these values corresponded to 57.5% and 0.17% oxygen saturation. High, medium and low H₂ experiments consisted of (v/v) 1.9%, 0.7% and 0.4% H₂ additions. The balance of all gas mixtures was made up with N₂. Cell density in liquid samples was monitored by measuring turbidity at 600 nm using a Ultrospec 10 cell density meter (Amersham Bioscience, UK). One unit of OD₆₀₀ was found to be equivalent to 0.43 g l⁻¹ cell dry weight 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). Biomass cell dry weight was used to calculate growth rate and specific gas consumption rate.

Whole-cell biochemical assays

Hydrogenase activity of Methylacidiphilum sp. RTK17.1 was measured in stationary-phase cultures harvested from the bioreactor. For amperometric measurements, whole cells were concentrated 5-, 10-, 20- and 30-fold by centrifugation followed by resuspension in V4 mineral medium (pH 3.0). Rate of H₂ oxidation was measured at 50°C using an H₂-MR
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Results and discussion

Verrucomicrobia-dominated surface soils serve as a sink of geothermally derived H₂ and CH₄ in Rotokawa geothermal field

We performed a geochemical, molecular and biochemical survey of CH₄ and H₂ metabolism in an acidic geothermal soil in Rotokawa, New Zealand. The Rotokawa geothermal field is a predominately steam-driven system dominated by acidic and sulphurous springs and heated soil features. We selected a geothermally heated and acidic soil where previous studies have indicated methanotrophic activity (Sharp et al., 2014). Substantial vertical gradients in temperature, pH and mixing ratios of CH₄, H₂ and O₂ were observed in the soil profile (Figure 1a). Consistent with the geothermal activity at the site, high soil mixing ratios of CH₄ (47 000 ppmv) and H₂ (280 ppmv) were detectable at the deepest soil depths sampled. The levels of both gases decreased in the upper 30 cm of soil and, in the case of H₂, dropped towards atmospheric levels by 10 cm depth (Figure 1b). These sharp decreases suggested that there were active methanotrophs and hydrogenotrophs in the oxic zone of the soil that consume most geothermally derived gas before it is emitted into the atmosphere. Indeed, microcosm incubations containing surface soils and associated communities rapidly consumed H₂ and CH₄ introduced into ambient air headspaces. Rates of H₂ oxidation exceeded that of CH₄, suggesting that H₂ serves as a major energy source for this geothermal soil community (Figures 1c and d).

To infer the microorganisms responsible for CH₄ and H₂ uptake, we determined the microbial community structure of the soil profile. Consistent with findings in other acidic soil ecosystems (Golyshina, 2011; Sharp et al., 2014; Lee et al., 2016), the euryarchaeotal order Thermoplasmatales was dominant at all depths. Methanotrophic verrucomicrobial genera, specifically Methylacidiphilum spp. and Methylacidimicrobium spp., were the dominant bacterial OTUs in surface soils and accounted for 47% of all bacterial 16S rRNA gene sequences in the soil profile (Figure 1e). Bacteria from these genera have been previously isolated in acidic geothermal soils in New Zealand (Dunfield et al., 2007; Sharp et al., 2014), Kamchatka (Islam et al., 2008) and Italy (Pol et al., 2007; van Teeseling et al., 2014). As the only known acidophilic methanotrophs (Op den Camp et al., 2009), it is probable that the verrucomicrobial phylotypes were solely responsible for CH₄ consumption in this ecosystem. Moreover, given putative uptake [NiFe]-hydrogenases have been detected in the genomes of Methylacidiphilales but not in Thermoplasmatales (Hou et al., 2008; Khadem et al., 2012b; Greening et al., 2016), it was likely that the Verrucomicrobia detected in these soils serve as major sinks of H₂ in this ecosystem. To test this possibility, we designed PCR primers to detect the presence of genes encoding the large

Details of nucleic acid extraction, amplification, genome sequencing, environmental quantitative PCR and soil microbial community composition determination methodologies are presented in the Supplementary Information.

microsensor (Unisense, Denmark) as previously described (Berney et al., 2014b; Greening et al., 2015). For colourimetric assays, 500 ml culture was harvested by centrifugation (15 min, 5000 g, 4 °C) and treated as previously described (Greening et al., 2014a, 2015) to prepare crude, cytosolic and membrane fractions for analysis. To test for hydrogenase activity, samples (20 μg protein) from each cell fraction were incubated with 1 ml 50 mM potassium phosphate buffer (pH 7.0) and 50 μM benzyl viologen for 8 h in an anaerobic chamber (5% H₂, 10% CO₂, 85% N₂ (v/v)). Debris was removed by centrifugation (15 min, 10 000 × g, 4 °C) and the absorbance of the supernatants was read at 604 nm in a Jenway 6300 spectrophotometer (Cole Palmer, UK). O₃ consumption experiments were performed on cell suspensions of Methylacidiphilum sp. RTK17.1 to determine the influence of endogenous oxygen catabolism (Khadem et al., 2012a) on O₃-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, carbonyl cyanide m-chlorophenyl hydrazine (CCCP), 1 mM iodoacetamide (an inhibitor of glycolysis) and 1 mM potassium cyanide (KCN) to determine whether observed rates of O₂ consumption were a consequence of oxygen catabolism. An oxygen solubility value of 220 nmol ml⁻¹ was used for calculations. Values were expressed as nmol O₂ min⁻¹ (mg protein)⁻¹. Protein was determined, from lysed cell pellets, using the BCA assay (Thermo Fisher Scientific, Waltham, MA, USA), with bovine serum albumin as a standard.

CO₂ fixation was measured by incubating cultures with 14C-labelled HCO₃⁻ (as CO₂ at medium pH of 2.3). Triplicate cultures were initially grown at 50 °C with a headspace of CH₄ (20%), CO₂ (10%), H₂ (10%) in air. At late exponential stage growth, the headspaces of cultures (and heat-killed controls) were replaced by sparging with N₂ (10 min) and then amended with H₂ (8%), CO₂ (8%) and O₂ (1%). 15 μCi of 14C-HCO₃⁻ (51 mCi mmol⁻¹; American Radiolabeled Chemical, St Louis, MO, USA) were added to each culture. Given that the pK₅ for HCO₃⁻ at 50 °C is ~6.8 (Amend and Shock, 2001), it was assumed that all added radiolabelled substrate equilibrated with unlabelled CO₂ pools immediately. Subsamples (1 ml) were harvested by centrifugation at regular time points, washed with sterile medium and subjected to liquid scintillation counting using Cytoscint cocktail as previously described (Urschel et al., 2015). Disintegrations per minute were converted to rates of CO₂ fixed using previously described approaches (Urschel et al., 2015).

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subunits of the three particulate methane monooxygenases (pmoA) and a single oxygen-tolerant uptake hydrogenase (hyaB) encoded in the genome of *Methylacidiphilum infernorum* V4 (Hou *et al.*, 2008) (Supplementary Table S1). These primers were applied in qPCRs on DNA extracts from the Rotokawa soil profile. Both the hydrogenase and methane monooxygenase genes were detected at all depths, with the most abundant templates detected in the top 10 cm of soil (Figure 1f), corresponding to the zones with the highest relative abundance of Verrucomicrobia-affiliated sequences and where the lowest CH₄ and H₂ soil gas concentrations were detected (Figure 1b).
A verrucomicrobial strain isolated from Rotokawa constitutively oxidises CH$_4$ and H$_2$ gas

To gain insight into the metabolic strategies that verrucomicrobia use to dominate bacterial assemblages in geothermally heated acidic soil ecosystems, we isolated a thermotolerant methanotroph from surface soils. The strain, *Methylacidiphilum* sp. RTK17.1 (Supplementary Table S2), grew optimally at pH 2.5, 50 °C ($T_{\text{max}}$ 60 °C) and shared 99% 16S rRNA gene sequence identity with *Methylacidiphilum infernorum* V4 (Dunfield et al., 2007). Bacteriological characterisation confirmed that the strain, in common with other verrucomicrobial methanotrophs (Khadem et al., 2012a, 2011), oxidised CH$_4$, fixed CO$_2$ and accumulated glycogen. In addition, cultures rapidly consumed exogenous H$_2$ (Supplementary Figures S2 and S3). Real-time amperometric measurements confirmed that the strain oxidised H$_2$ under oxic conditions at rates proportional to increases in cell density (Figure 2a). H$_2$ oxidation occurred in all batch culture conditions tested, including when CH$_4$ was absent (Supplementary Figure S1A), when CH$_4$ was in excess (Supplementary Figure S1C), and following inhibition of CH$_4$ oxidation with acetylene (Supplementary Figure S2). This suggests that H$_2$ and CH$_4$ are oxidised independently serving to energise the respiratory chain through the reduction of the quinone pool. Moreover, this expands the role of hydrogenases in aerobic methanotrophs beyond their previously suggested role of providing reductant for pMMO (Shah et al., 1995; Hanczár et al., 2002). The observation that RTK17.1 can constitutively oxidise H$_2$ and CH$_4$ parallels results from the soil study showing that both H$_2$ and CH$_4$ are simultaneously oxidised (Figure 1). Considering that verrucomicrobia are dominant among taxa putatively capable of oxidising H$_2$ or CH$_4$, this provides further evidence that verrucomicrobial methanotrophs adopt a mixotrophic lifestyle with respect to their energy metabolism.
We sequenced the genome of Methylacidiphilum sp. RTK17.1 to obtain further insights into the potential functionality of this taxon (Supplementary Table S2). Genes encoding key enzymes and pathways for CH₄ oxidation to CO₂, CO₂ fixation through the Calvin–Benson–Bassham pathway, and aerobic respiration (Figure 3) were highly conserved with those identified in other Methylacidiphilum strains (Hou et al., 2008; Khadem et al., 2012b; Erikstad and Birkeland, 2015). We also detected two [NiFe]-hydrogenase-encoding gene clusters in the genome (Supplementary Figure S3) and confirmed their expression during aerobic growth with CH₄ and H₂ by RT-PCR (Supplementary Figure S4). The gene clusters were classified as groups 1d (hyaABC) and 3b (hyhBGSL) [NiFe]-hydrogenases based on phylogenetic affiliation with biochemically characterised enzymes (Supplementary Figure S3; Greening et al., 2016; Søndergaard et al., 2016). Biochemically characterised group 1d [NiFe]-hydrogenase are H₂-uptake multimeric proteins that are membrane-bound via their cytochrome b subunit (hyaC) and function by transferring electrons into the respiratory chain via the quinone pool (Fritsch et al., 2011). Consistent with the observed activity of the [NiFe]-hydrogenase in the presence of O₂ (Figure 2a), enzymes of this class are predicted to be O₂-tolerant due to the presence of a novel [4Fe3S] cluster that protects the O₂-sensitive active site from oxidative damage (Fritsch et al., 2011; Shomura et al., 2012b; Erikstad and Birkeland, 2015). Indeed, the six cysteine residues required to ligate such a cluster were conserved in the deduced RTK17.1 HyaB protein sequence. In comparison, biochemically characterised group 3b hydrogenases are reversible cytosolic enzymes that are relatively O₂-sensitive (Kwan et al., 2015); they directly couple NAD(P)H oxidation to H₂ formation during fermentation (Berney et al., 2014a) and, in some cases, H₂ oxidation by these enzymes supports CO₂ fixation through the production of reduced electron carriers (Yoon et al., 1996). The RTK17.1 [NiFe]-hydrogenase combination differs from Methylacidiphilum fumaroliicum SolV, where group 1 h/5 (hhy LH; a putative high-affinity H₂ uptake hydrogenase) and group 1d [NiFe]-hydrogenases (annotated as hupSLZ) were reported (Mohammadi et al., 2016), with our previous survey (Greening et al., 2016) also showing SolV encodes a group 3b enzyme (Supplementary Figure S3).

H₂ oxidation supports aerobic respiration and CO₂ fixation in the verrucomicrobial isolate

The observation that RTK17.1 encodes and utilises [NiFe]-hydrogenases prompted biochemical studies to investigate the role of H₂ in the metabolism of this bacterium. Biochemical assays targeting the group 1d [NiFe]-hydrogenase demonstrated that it is a membrane-bound uptake hydrogenase linked to the aerobic respiratory chain, consistent with our genome-based predictions. Fractionation experiments confirmed the activity was membrane-localised, as shown by the 31-fold increase in activity in membranes when compared to the cytosolic fraction (Figure 2b). We next tested the effect of the ionophores nigericin and valinomycin on rates of H₂ oxidation on whole cells (in the absence of CH₄). A large proton-motive force is generated and sufficient ATP is produced for growth via an H⁺-translocating F₁Fₒ-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 a cytochrome c oxidase complex (Complex III, Figure 3) were highly conserved with those identified in other Methylacidiphilum strains (Hou et al., 2008; Kwan et al., 2010; Shomura et al., 2012b; Erikstad and Birkeland, 2015). We also detected two [NiFe]-hydrogenase-encoding gene clusters in the genome (Supplementary Table S1). The observation that RTK17.1 encodes and utilises [NiFe]-hydrogenases based on phylogenetic affiliation with biochemically characterised enzymes (Supplementary Figure S3; Greening et al., 2016; Søndergaard et al., 2016) also showing SolV encodes a group 3b enzyme (Supplementary Figure S3).
with these ionophores (Figures 2c and d), showing hydrogenase activity behaves as expected from a component of the energy-conserving respiratory chain. This uncoupled activity rapidly ceased, due to O$_2$ consumption by the cells suspended in the sealed chamber, but could be restored by further supplementation with O$_2$. These results show hydrogenase is a *bona fide* component of this microorganism’s respiratory chain and is coupled to the activity of terminal cytochrome oxidases. Under these conditions, the onset of O$_2$-limitation was likely exacerbated by the catabolism of endogenous glycogen reserves (Supplementary Figure S5). Collectively, these findings demonstrate that this group 1d [NiFe]-hydrogenase is a membrane-bound, respiratory-linked, O$_2$-tolerant/dependent enzyme that drives ATP synthesis as has been observed in other aerobic hydrogenotrophs.

To test whether H$_2$ oxidation coupled to O$_2$ reduction could support CO$_2$ fixation in RTK17.1, we transferred mixotrophically grown, log phase cultures into a new microoxic headspace (O$_2$ 1% v/v) as the sole carbon source. Trace $^{14}$CO$_2$ (0.1% of total CO$_2$ supplied) was added and the amount fixed into biomass sampled over time was determined by measuring disintegrations per minute (DPM) via liquid scintillation counting. We observed systematic increases in DPMs associated with cells sampled over a 20 hour period relative to controls, indicating that CO$_2$ was rapidly incorporated into biomass in a time-dependent manner. The number of DPMs associated with cells after 20 hours of incubation were 200-fold greater in live than heat-killed cells (Figure 2e), showing that biological CO$_2$ fixation occurs in cultures supplied with H$_2$ as the sole reductant and O$_2$ as the sole oxidant. This activity was not observed in the absence of exogenous H$_2$, indicating that H$_2$ serves as the source of reductant for CO$_2$ fixation under these conditions.

H$_2$ oxidation supports adaptation of verrucomicrobial methanotrophs to CH$_4$ and O$_2$ limitation

The finding that *Methylacidiphilum* sp. RTK17.1 couples H$_2$ oxidation to aerobic respiration and carbon fixation suggests that it can grow chemolithoautotrophically. Consistent with this hypothesis, we observed a small but significant increase in biomass in cultures grown under microoxic conditions when H$_2$, CO$_2$ and O$_2$ (1% headspace concentration) were supplied as the sole electron donor, carbon source and electron acceptor, respectively (Supplementary Figure S2A). This biomass increase was concomitant with increased amounts of CO$_2$ fixed (Figure 2e), and the amount of carbon fixed per cell (40 to 80 fmol cell$^{-1}$) produced over this time period was consistent with biomass yields from other studies (Maestrini *et al.*, 2000). However, the observed growth rates (0.005 h$^{-1}$) were substantially lower than observed when RTK17.1 was supplied with CH$_4$, CO$_2$ and H$_2$ (0.037 h$^{-1}$) and when *Methylacidiphilum funariolicum* SolV was grown autotrophically under similar conditions (0.047 h$^{-1}$) (Mohammadi *et al.*, 2016). We also observed that autotrophic growth was only sustained when RTK17.1 was incubated under microoxic conditions (1% O$_2$) rather than in anoxic (20% O$_2$) headspace. We speculate that, under such microoxic conditions, sufficient O$_2$ is available to drive hydrogenotrophic aerobic respiration through activity of the group 1d [NiFe]-hydrogenase. Simultaneously, it is likely that the O$_2$-sensitive group 3b [NiFe]-hydrogenase can remain active and is able to supply reducing equivalents required for CO$_2$ fixation. In support of this notion, H$_2$ oxidation sustained energy-conservation via the group 1d hydrogenase of non-growing CH$_4$-limited cultures in the presence of ambient O$_2$ (Supplementary Figure S2C) and enhanced growth yields in CH$_4$-replete cultures (Supplementary Figure S2D).

To better understand the role of H$_2$ and CH$_4$ oxidation during mixotrophic growth, we compared growth and gas consumption kinetics of the cells cultivated in a chemostat under six different conditions (Table 1). We observed that H$_2$ addition into the feedgas of *Methylacidiphilum* sp. RTK17.1 increased growth yields under O$_2$-replete and O$_2$-limiting conditions. Whereas CH$_4$ oxidation predominated under O$_2$-replete conditions, the specific consumption rate of H$_2$ increased 80-fold and exceeded rates of CH$_4$ oxidation under O$_2$-limiting conditions. In combination, these results show that *Methylacidiphilum* sp. RTK17.1 grows mixotrophically and modulates rates of H$_2$ and CH$_4$ consumption in response to the availability of O$_2$ in order to balance energy-generation and carbon fixation.

H$_2$ oxidation may be a general ecological strategy for verrucomicrobial and proteobacterial methanotrophs

In this work, we demonstrated that a verrucomicrobial methanotroph adopts a mixotrophic lifestyle both in situ and ex situ. The environmental isolate *Methylacidiphilum* sp. RTK17.1 sustains aerobic respiration and carbon fixation by using organic (CH$_4$) and inorganic (H$_2$) electron donors either in concert or separately depending on substrate availability. Through the dual use of both electron donors, the bacterium is able to more flexibly adjust its metabolism to meet energy and carbon demands in response to simulated environmental change. A model of how CH$_4$ and H$_2$ metabolism is integrated into the physiology of this microorganism, based in part on genomic information (Supplementary Table S2), is shown in Figure 3. Integrating our genomic, physiological and biochemical findings, we conclude that the group 1d [NiFe]-hydrogenase is a membrane-bound uptake hydrogenase that is directly linked to the aerobic respiratory chain and supplements the quinone pool to power the methane monoxygenase reaction or feed directly into Complex
III. The organism is also capable of using H$_2$ as a reductant to support CO$_2$ fixation through the Calvin–Benson-Bassham pathway, likely through the cytosolic NAD(P)-coupled group 3b [NiFe]-hydrogenase. The [NiFe]-hydrogenase combination (group 1d and group 3b) in this RTK17.1 only supports weak autotrophic growth, but provides multiple layers of support for a mixotrophic lifestyle. In addition to supporting growth and survival during periods of CH$_4$ limitation, our data show that H$_2$ is the preferred electron donor during O$_2$-limiting conditions. Under these conditions, rates of H$_2$ consumption increased by 77-fold and exceeded observed rates of CH$_4$ oxidation (Table 1). This is likely to be a consequence of two factors. Firstly, some hydrogenases such as the group 3b [NiFe]-hydrogenase are inhibited at high O$_2$ concentrations (Kwan et al., 2015). Secondly, methanotrophy is more resource-intensive than canonical aerobic hydrogenotrophy, given it requires O$_2$ both as a substrate for methane monooxygenase and as the terminal electron acceptor for respiration (Hakemian and Rosenzweig, 2007).

Comparison of our independent findings with those made by Mohammadi et al. (2016) suggests that verrucomicrobial methanotrophs have evolved a range of strategies to integrate H$_2$ metabolism into their physiology. Both *Methylacidiphilum* sp. RTK17.1 and *Methylacidiphilum fumarophilum* SolV are capable of sustaining chemolithoautotrophic growth on H$_2$/CO$_2$ under microoxic conditions. However, the strains grow at drastically different rates of 0.005 and 0.047 h$^{-1}$ (Mohammadi et al., 2016), respectively, under optimal conditions. These differences may reflect that, while both strains possess group 1d (*hyaABC/hupLSZ*) and group 3b (*hybBGSZ*) hydrogenases, SolV has also acquired a group 1h enzyme (*hyhLH*) (Greening et al., 2016) with surprisingly fast whole-cell kinetics (Mohammadi et al., 2016). It is possible that, with this enhanced hydrogenase suite, SolV may be able to more efficiently partition electrons derived from H$_2$ oxidation between respiration and carbon fixation. In addition to supporting hydrogenotrophic growth, both organisms also modulate hydrogenase expression and H$_2$ oxidation rates in response to simulated environmental change, such as CH$_4$ and O$_2$ availability (Mohammadi et al., 2016). While the physiological significance of this regulation was not explored in SolV, our studies inferred that H$_2$ co-oxidation with CH$_4$ enhanced yields during CH$_4$ surplus and sustained survival during CH$_4$ limitation in RTK17.1. Further differences between the strains are reflected in the regulatory profile, with the group 1d enzyme constitutively expressed in RTK17.1, but repressed in favour of the group 1h enzyme under oxic conditions in SolV (Mohammadi et al., 2016). Overall, our findings suggest that SolV may fulfil a similar ecological niche to classical Knallgas bacteria (e.g. *Ralstonia eutropha*), switching between efficient heterotrophic and autotrophic growth dependent on energy availability. In contrast, H$_2$ metabolism appears to be more important for optimising growth and survival of RTK17.1 in response to energy and O$_2$ availability. In this regard, this organism’s metabolism more closely resembles the mixotrophic strategy employed by *Mycobacterium smegmatis* (Berney et al., 2014a; Greening et al., 2014b). Future studies would benefit from side-by-side comparisons of these strains under equivalent conditions and further exploration of the physiological role and biochemical features of the group 3b and group 1h [NiFe]-hydrogenase enzymes.

More generally, we predict that H$_2$ oxidation is likely to support the majority of aerobic methanotrophic bacteria. Whereas only a few methanotrophic genera appear to be capable of heterotrophic generalism (Crombie and Murrell, 2014), genomic surveys (Peters et al., 2015; Greening et al., 2016) show that all 31 publicly available aerobic methanotrophic genomes harbour the capacity to metabolise H$_2$ (Supplementary Figure S3). As with SolV (Mohammadi et al., 2016) and RTK17.1, most of the surveyed methanotroph genomes that were found to

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### Table 1 H$_2$ oxidation by *Methylacidiphilum* sp. RTK17.1 during chemostat cultivation

| Condition | Growth rate$^a$ | H$_2$ consumption | CH$_4$ consumption | Biomass yield$^a$ |
|-----------|-----------------|-------------------|--------------------|------------------|
| O$_2$-replete, medium H$_2$ | 9.33 (±0.52) | 0.01 (±0.01) | 0.03 (±0.03) | 1.06 (±0.12) |
| O$_2$-replete, no H$_2$ | 8.17 (±0.82) | 0.00 | 0.00 | 1.22 (±0.05) |
| O$_2$-limiting, high H$_2$ | 5.57 (±0.16) | 0.78 (±0.02) | 2.66 (±0.11) | 0.55 (±0.03) |
| O$_2$-limiting, medium H$_2$ | 4.99 (±0.17) | 0.53 (±0.01) | 1.34 (±0.06) | 0.70 (±0.02) |
| O$_2$-limiting, low H$_2$ | 5.60 (±0.31) | 0.19 (±0.00) | 0.73 (±0.04) | 0.71 (±0.01) |
| O$_2$-limiting, no H$_2$ | 5.40 (±0.26) | 0.00 | 0.00 | 0.80 (±0.05) |

$^a$ Feedgas was continuously supplied at a rate of 10 ml min$^{-1}$ with the following composition (v/v): O$_2$-replete, 14.1%; O$_2$-limiting, 3.5%. High, medium and low H$_2$ experiments consisted of 1.9%, 0.7% and 0.4% H$_2$ supply. For all experiments excess CH$_4$ was supplied at 3%, CO$_2$ at 26% with the balance made up with N$_2$. O$_2$ saturations in the medium were 57.5% and 0.17% for the O$_2$-replete and O$_2$-limiting conditions, respectively.

$^b$ A dilution rate of 0.02 h$^{-1}$ was maintained for all experiments.

$^c$ Denotes significant increase ($P$-value < 0.05) in yield from O$_2$-replete, no H$_2$ result; unpaired t-test, one tailed.

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

$^e$ Denotes significant increase ($P$-value < 0.05) in yield from O$_2$-limiting, no H$_2$ result; unpaired t-test, one tailed.

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encode for [NiFe]-hydrogenases have been shown to support aerobic respiration (groups 1d, 1h, 2a) and carbon fixation (groups 3b, 3d, 1h) (Greening et al., 2016). Reports showing H₂ oxidation by several proteobacterial strains further support the classification of these enzymes as uptake hydrogenases (Chen and Yoch, 1987; Shah et al., 1995; Hanczár et al., 2002). H₂ is likely to be a particularly attractive energy source for methanotrophs because of its relative ubiquity when compared to C₁ compounds. H₂ is biologically produced by diverse organisms across the three domains of life as a result of fermentation, photobiological processes and nitrogen fixation (Peters et al., 2013; Schwartz et al., 2013; Poudel et al., 2016). Moreover, verrucomicrobial and proteobacterial methanotrophs harbouring the recently described group 1h [NiFe]-hydrogenases (Greening et al., 2014a, 2015) may be capable of scavenging atmospheric H₂ to survive CH₄ starvation. Considering these observations in concert, it seems likely that hydrogenases in aerobic methanotrophs function to supplement the energetic and reductant requirements in environments where CH₄ and O₂ gases are limiting or variable. Thus, while methanotrophic bacteria are often pervasively viewed as C₁ specialists, we propose that, via the utilisation of hydrogenases as part of a mixotrophic strategy, the niche space of methanotrophs is much broader than previously recognised. Combining heterotrophic and lithotrophic electron donors allows for a more flexible growth/survival strategy, with clear ecological benefits (Semrau et al., 2011). We therefore predict that most methanotrophs will be able to use H₂ to support either autotrophic growth, mixotrophic growth or long-term persistence/maintainence.

Finally, our geochemical and microbial community diversity investigation of the Rotokawa geothermal field provides ecological support to our assertion that the metabolic flexibility of methanotrophs enhances niche expansion in situ. We provide genetic and biochemical evidence that methanotrophic Verrucomicrobia inhabiting the near-surface soils co-metabolised CH₄ and H₂ gas (Figure 1) and in doing so adopt a clear mixotrophic strategy. In acidic geothermal soils, we demonstrated that verrucomicobial methanotrophs have grown to be the dominant bacterial taxon by simultaneously consuming gases primarily of geothermal and atmospheric origin, that is, CH₄ and H₂ as energy sources, respectively, CO₂ as a carbon source and O₂ as oxidant. Their metabolic flexibility also ensures resilience to temporal and spatial variations in the availability of key substrates allowing for CH₄ oxidation via the monooxygenase reaction. More generally, the prevalent narrative that methanotrophic bacteria are methylotrophic specialists is based on studies under optimal growth conditions and ignores the requirement of these organisms to adapt to environmental variations requiring a certain level of metabolic versatility. Intimate evolutionary and ecological interactions are likely to have selected for a spectrum of different lifestyles across methanotrophic lineages, ranging from strict C₁ specialism to broad substrate generalism, depending on the environment. However, based on the presence of [NiFe]-hydrogenase in numerous methanotroph genomes (Supplementary Figure S3) and the data presented here, we contend it is likely that most methanotrophs depend on H₂ oxidation to some extent to support either growth and/or survival. This finding has broad implications for future investigations on the ecology of methanotrophs as well as the biogeochemical cycles of H₂ and CH₄.

Conflict of Interest
The authors declare no conflict of interest.

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Author contributions
CRC, CG and MBS conceived the study. CRC, MBS, CG, GC, KMH, KH, CC, DJG, RS, ESB and BM contributed to experimental design. CRC, BM, CC, DJG and MHK conducted bioreactor and wet lab experiments. CRC and MBS performed fieldwork, JFP determined microbial community structure, and CRC, MBS and CG conducted genomic analyses. CRC, KH and CG undertook cell respiratory analysis, and ESB and MBS performed the radioactive tracing experiments. CRC, CG, MBS, KH and GC wrote the manuscript.

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