A widely distributed hydrogenase oxidises atmospheric H₂ during bacterial growth

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Abstract

Diverse aerobic bacteria persist by consuming atmospheric hydrogen (H₂) using group 1h [NiFe]-hydrogenases. However, other hydrogenase classes are also distributed in aerobes, including the group 2a [NiFe]-hydrogenase. Based on studies focused on Cyanobacteria, the reported physiological role of the group 2a [NiFe]-hydrogenase is to recycle H₂ produced by nitrogenase. However, given this hydrogenase is also present in various heterotrophs and lithoautotrophs lacking nitrogenases, it may play a wider role in bacterial metabolism. Here we investigated the role of this enzyme in three species from different phylogenetic lineages and ecological niches: Acidithiobacillus ferrooxidans (phylum Proteobacteria), Chloroflexus aggregans (phylum Chloroflexota), and Gemmatimonas aurantiaca (phylum Gemmatimonadota). qRT-PCR analysis revealed that the group 2a [NiFe]-hydrogenase of all three species is significantly upregulated during exponential growth compared to stationary phase, in contrast to the profile of the persistence-linked group 1h [NiFe]-hydrogenase. Whole-cell biochemical assays confirmed that all three strains aerobically respire H₂ to sub-atmospheric levels, and oxidation rates were much higher during growth. Moreover, the oxidation of H₂ supported mixotrophic growth of the carbon-fixing strains C. aggregans and A. ferrooxidans. Finally, we used phylogenomic analyses to show that this hydrogenase is widely distributed and is encoded by 13 bacterial phyla. These findings challenge the current persistence-centric model of the physiological role of atmospheric H₂ oxidation and extend this process to two more phyla, Proteobacteria and Gemmatimonadota. In turn, these findings have broader relevance for understanding how bacteria conserve energy in different environments and control the biogeochemical cycling of atmospheric trace gases.

Introduction

Aerobic bacteria mediate the biogeochemically and ecologically important process of atmospheric hydrogen (H₂) oxidation [1]. Terrestrial bacteria constitute the largest sink of this gas and mediate the net consumption of ~70 million tonnes of atmospheric H₂ per year [2, 3]. The energy derived from this process appears to be critical for sustaining the productivity and biodiversity of ecosystems with low organic carbon inputs [4–9]. Atmospheric H₂ oxidation is thought to be primarily mediated by group 1h [NiFe]-hydrogenases, a specialised oxygen-tolerant, high-affinity class of hydrogenases [4, 10–13]. To date, aerobic heterotrophic bacteria from four distinct bacterial phyla, the Actinobacteria (formerly Actinobacteria; [10, 12, 14, 15]), Acidobacteria (formerly Acidobacteria; [16, 17]), Chloroflexota (formerly Chloroflexi; [18]), and Verrucomicrobiota (formerly Verrucomicrobia; [19]), have been experimentally shown to consume atmospheric H₂ using...

Supplementary information

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this enzyme. This process has been primarily linked to energy conservation during persistence. Reflecting this, the expression and activity of the group 1h hydrogenase is induced by carbon starvation across a wide range of species [10, 12, 16, 18, 20–22]. Moreover, genetic deletion of hydrogenase structural genes results in impaired long-term survival of *Mycobacterium smegmatis* cells and *Streptomyces avermitilis* spores [20, 21, 23, 24].

Genomic and metagenomic surveys have suggested that other uptake hydrogenases are widely distributed among aerobic bacteria and potentially have a role in atmospheric H₂ uptake [4, 25]. These include the widely distributed group 2a [NiFe]-hydrogenases. This hydrogenase class has primarily been investigated in Cyanobacteria, where it is encoded by most diazotrophic strains; the enzyme recycles H₂ released as a by-product of the nitrogenase reaction and inputs the derived electrons into the respiratory chain [26–29]. However, according to HydDB, group 2a hydrogenases are also encoded by isolates from at least eight other phyla [25], spanning both obligate organoheterotrophs (e.g., *Mycobacterium*, *Runella*, *Gemmatimonas*) and obligate lithoautotrophs (e.g., *Acidithiobacillus*, *Nitrospira*, *Hydrogenobacter*) [12, 30, 31]. In *M. smegmatis*, this enzyme has a sufficiently high apparent affinity to oxidise H₂ even at sub-atmospheric levels [12, 22] and is maximally expressed during transitions between growth and persistence [22, 32]. In common with the group 1h hydrogenase also encoded by this bacterium, the group 2a hydrogenase requires potential electron-relaying iron-sulphur proteins for activity [33] and is obligately linked to the aerobic respiratory chain [22]. However, it remains unclear if atmospheric H₂ oxidation by the group 2a hydrogenase reflects a general feature of the enzyme or instead is a specific adaptation of the mycobacterial respiratory chain.

In this study, we investigated whether group 2a [NiFe]-hydrogenases play a general role in atmospheric H₂ consumption. To do so, we studied this enzyme in three species, *Gemmatimonas aurantiaca*, *Acidithiobacillus ferrooxidans*, and *Chloroflexus aggregans*, that differ in their phylogenetic affiliation, ecological niches, and metabolic strategies. The obligate chemoorganoheterotroph *G. aurantiaca* (phylum Gemmatimonadota; formerly Gemmatimonadetes) was originally isolated from a wastewater treatment plant and to date has not been shown to utilise H₂ [34, 35]. The obligate chemolithoautotroph *A. ferrooxidans* (phylum Proteobacteria) was originally isolated from acidic coal mine effluent, and has been extensively studied for its energetic flexibility, including the ability to grow exclusively on H₂ [31, 36, 37]. The metabolically flexible *C. aggregans* (phylum Chloroflexota), a facultative chemolithoautotroph and anoxygenic phototrophetrotroph, was originally isolated from a Japanese hot spring and is capable of hydrogenotrophic growth [38–40]. The organisms differ in their carbon dioxide fixation pathways, with *A. ferrooxidans* mediating the Calvin-Benson cycle via two RuBisCO enzymes, *C. aggregans* encoding the 3-hydroxypropionate cycle [37, 41, 42], and *G. aurantiaca* unable to fix carbon dioxide [34]. While all three species have previously been shown to encode group 2a [NiFe]-hydrogenases [4, 37], it is unknown whether they can oxidise atmospheric H₂. To resolve this, we investigated the expression, activity, and role of this enzyme in axenic cultures of the three species.

**Materials and methods**

**Bacterial growth conditions**

*Gemmatimonas aurantiaca* (DSM 14586), *Acidithiobacillus ferrooxidans* (DSM 14882), and *Chloroflexus aggregans* (DSM 9486) were imported from DSMZ. All cultures were maintained in 120 mL glass serum vials containing a headspace of ambient air (H₂ mixing ratio ~0.5 ppmv) sealed with lab-grade butyl rubber stoppers. Prior to use, starters were treated to prevent H₂ release by boiling twice in 0.1 M sodium hydroxide for 2 h, and twice in deionised water for 2 h, prior to baking in a 70 °C overnight. Broth cultures of *G. aurantiaca* were grown in 30 mL of NM1 media as previously described [43] and incubated at 30 °C at an agitation speed of 180 rpm in a New Brunswick Scientific Excella E24 incubator. Cultures of *C. aggregans* were maintained chemoheterotrophically in 30 mL of 1/5 PE media, as previously described [38], and incubated at 55 °C at an agitation speed of 150 rpm in an Eppendorf 40 Incubator in the dark. Cultures of *A. ferrooxidans* were maintained in 30 mL DSMZ medium 882 supplemented with an additional 13 g L⁻¹ of FeSO₄·7H₂O (pH 1.2) and incubated at 30 °C at an agitation speed of 180 rpm in a New Brunswick Scientific Excella E24 incubator. To assess whether bacterial growth was enhanced by the presence of H₂ for each species, ambient air headspaces were amended with either 1 or 10% H₂ (via 99.999% pure H₂ gas cylinder). Growth was monitored by determining the optical density (OD₆₀₀) of periodically sampled 1 mL extracts using an Eppendorf BioSpectrophotometer.

**RNA extraction**

Triplicate 30 mL cultures of *G. aurantiaca*, *A. ferrooxidans* and *C. aggregans* were grown synchronously in 120 mL sealed serum vials. Whereas one set of triplicate cultures were grown in an ambient air headspace, another set was grown in an ambient air headspace supplemented with H₂ to a final concentration of 10% v/v (via a 99.999% pure H₂ cylinder). Cultures were grown to either exponential phase (OD₆₀₀ 0.05 for *G. aurantiaca*; OD₆₀₀ 0.1 for *C. aggregans*;
OD$_{600}$ 0.05 for *A. ferrooxidans* or stationary phase (Day 10 for *G. aurantiaca*; Day 4 for *C. aggregans*; Day 14 for *A. ferrooxidans*). For *G. aurantiaca* and *C. aggregans*, cells were then quenched using a glycerol-saline solution (−20°C, 3:2 v/v), harvested by centrifugation (20,000 × g, 30 min, −9°C), resuspended in 1 mL cold 1:1 glycerol:saline solution (−20°C), and further centrifuged (20,000 × g, 30 min, −9°C). Briefly, resultant cell pellets were resuspended in 1 mL TRIzol Reagent (Thermo Fisher Scientific), mixed with 0.1 mm zircon beads (0.3 g), and subject to bead-beating (five cycles, 4000 rpm, 30 s) in a Mini-Beadbeater 96 (Biospec) prior to centrifugation (12,000 × g, 10 min, 4°C). Total RNA was extracted using the phenol-chloroform method as per manufacturer’s instructions (TRIzol Reagent User Guide, Thermo Fisher Scientific) and resuspended in diethylpyrocarbonate-treated water. RNA was treated using the TURBO DNA-free kit (Thermo Fisher Scientific) as per manufacturer’s instructions. RNA from *A. ferrooxidans* was extracted using a previously described extraction method optimised for acid mine drainage microorganisms [44]. RNA concentration and purity were confirmed using a NanoDrop ND-1000 spectrophotometer.

**Quantitative RT-PCR**

Quantitative reverse transcription PCR (qRT-PCR) was used to determine the expression profile of all hydrogenase genes present in each species during different growth phases with and without supplemental H$_2$. cDNA was synthesised using a SuperScript III First-Strand Synthesis System kit for qRT-PCR (Thermo Fisher Scientific) with random hexamer primers, as per manufacturer’s instructions. For all three species, the catalytic subunit gene of the group 2a [NiFe]-hydrogenase (*hucL*) was targeted. In addition, transcript levels of the catalytic subunit of all additional [NiFe]-hydrogenases present in these strains were analysed, i.e., group 3d (*hoxH*) for *C. aggregans* and both group 1e (*hyiB*) and group 3b (*hyhL*) for *A. ferrooxidans*. Quantitative RT-PCR was performed using a LightCycler 480 SYBR Green I Master Mix (Roche) as per manufacturer’s instructions in 96-well plates and conducted in a LightCycler 480 Instrument II (Roche). Primers used in the study (Table S1) were designed using Primer3 [45]. Copy numbers of each gene were interpolated from standard curves of each gene created from threshold cycle (C$_T$) values of amplicons that were serially diluted from $10^9$ to 10 copies ($R^2 > 0.95$). Hydrogenase expression data were then normalised to housekeeping genes in exponential phase under ambient air conditions for each species (16S rRNA gene for *G. aurantiaca* and *C. aggregans*; DNA-directed RNA polymerase subunit beta gene *rpoC* for *A. ferrooxidans*). All biological triplicate samples, standards, and negative controls were run in technical duplicate.

**Gas chromatography**

Gas chromatography measurements were used to determine the capacity of the three species to use sub-atmospheric concentrations of H$_2$. For initial experiments, H$_2$ consumption by triplicate cultures in vials containing an ambient air headspace was monitored during growth; H$_2$ mixing ratios were measured immediately following inoculation (mixing ratio = 440 ± 34 ppbv), then at mid-exponential and late stationary phase. In subsequent experiments, to determine H$_2$ oxidation rate constants, biological triplicate cultures of each species were opened, equilibrated with ambient air (1 h), and resealed. These aerated vials were then amended with H$_2$ (*via* 1% v/v H$_2$ in N$_2$ gas cylinder, 99.999% pure) to achieve final headspace concentrations of ~10 ppmv. Headspace mixing ratios were measured immediately after closure and at regular intervals thereafter for 200 h or until the limit of quantification of the gas chromatograph was reached (42 ppbv H$_2$). This analysis was performed for both exponential phase and stationary phase cultures. For H$_2$ quantification, 2 mL headspace samples were measured using a pulsed discharge helium ionisation detector (model TGA-6791-W-4U-2, Valco Instruments Company Inc.) calibrated against standard H$_2$ gas mixtures of known concentrations (0.1, 0.5, 1, 5, 20, 50, 100, 150, 500, 1000, 2500, 5000, and 7000 ppmv), prepared by diluting either 99.999% pure H$_2$ gas cylinder in synthetic air (20.5% v/v O$_2$ in N$_2$) or 1% v/v H$_2$ in N$_2$ gas into He (99.999% pure) as described previously [18]. The vials for each species were maintained at their respective growth temperatures and agitation speeds for the entire incubation period to facilitate H$_2$ and O$_2$ transfer between the headspace and the culture. Concurrently, headspace mixing ratios from media-only negative controls (30 mL of media for each species) were measured to confirm that observed decreases in gas concentrations were biological in nature. First order rate constants (k values) for exponential and stationary phase H$_2$ consumption were determined using the exponential function in GraphPad Prism (version 8.0.2).

**Phylogenetic analysis**

A phylogenetic tree was constructed to investigate the distribution and evolutionary history of group 2a [NiFe]-hydrogenases across bacterial phyla. Amino acid sequences of the catalytic subunit of the group 2a [NiFe]-hydrogenase (*HucL*) and related enzymes were retrieved from the National Center for Biotechnology Information (NCBI) Reference Sequence database by protein BLAST in February 2020. The resultant sequences were then classified using HydDB [25], with sequences matching group 2a [NiFe]-hydrogenases retained and any duplicate and multispecies sequences removed. The 207 amino acid
sequences representative of genus-level diversity were aligned with reference sequences using Clustal W in MEGA X [46]. Evolutionary relationships were visualised by constructing a maximum-likelihood phylogenetic tree, with Neighbour-Joining and BioNJ algorithms applied to a matrix of pairwise distances that were estimated using a JTT model and topology selected by superior log-likelihood value. Gaps were treated with partial deletion, the tree was midpoint rooted. Sequences used in this analysis are listed in Table S2. In addition, 20 annotated reference genomes (representative of order-level diversity) were retrieved from the NCBI GenBank database and manually analysed for putative group 2a [NiFe]-hydrogenase gene clusters. The web-based software Properon (doi.org/10.5281/zenodo.3519494) was used to generate to-scale gene organisation diagrams of these group 2a [NiFe]-hydrogenases. All species names and taxonomic assignments follow the Genome Taxonomy Database [47, 48].

Results

The expression profile of group 2a [NiFe]-hydrogenases is antithetical to group 1h [NiFe]-hydrogenases

We used qRT-PCR to quantify the expression of the large subunit of the group 2a [NiFe]-hydrogenase (hucL). The gene was expressed at moderate to high levels in all three strains during aerobic growth on preferred energy sources (organic carbon for G. aurantiaca and C. aggregans, ferrous iron for A. ferrooxidans) (Fig. 1). Expression levels did not significantly differ between strains grown in an ambient air headspace containing atmospheric H2 or supplemented with 10% H2 (Fig. 1). This suggests hydrogenase expression is constitutive and occurs even when atmospheric concentrations of the substrate are available.

Across all three strains, hydrogenase expression significantly decreased during the transition from growth to persistence. For G. aurantiaca, high expression was observed during exponential phase under both H2-supplemented and H2-unamended conditions (av. 8.4 × 10^6 copies per gdw) and decreased 51-fold during stationary phase (av. 1.6 × 10^5 copies gdw⁻¹; p = 0.012) (Fig. 1a). Hydrogenase expression of A. ferrooxidans was moderate during growth (av. 1.8 × 10^6 copies per gdw) and dropped 3.9-fold in stationary phase cultures (av. 4.5 × 10^5 copies per gdw; p = 0.013) (Fig. 1b), whereas expression in C. aggregans was very high during exponential growth (av. 2.9 × 10^8 copies gdw⁻¹) and fell 15,000-fold during persistance (av. 1.9 × 10^5 copies gdw⁻¹; 0.003) (Fig. 1c). Overall, while expression levels greatly vary between species, these results clearly show the group 2a [NiFe]-hydrogenase is expressed primarily in growing cells. These expression profiles contrast with the group 1h [NiFe]-hydrogenase, which is induced during long-term persistence in a range of species [10, 16, 18, 20–22].

Group 2a [NiFe]-hydrogenases oxidise H2 to sub-atmospheric levels

Hydrogenase activity of the three strains was inferred from monitoring changes in headspace H2 mixing ratios over time by gas chromatography. In line with the expression profiles (Fig. 1), we observed that all three strains oxidised atmospheric H2 during growth in an ambient air headspace (Fig. S1). These observations extend the trait of trace gas scavenging to three more species and suggest that group 2a [NiFe]-hydrogenases broadly have the capacity to oxidise H2 at atmospheric levels.

Fig. 1 Expression of the group 2a [NiFe]-hydrogenase in three bacterial strains during growth and survival. The normalised transcript copy number of the large subunit gene (hucL) are plotted for (a) Gemmatimonas aurantiaca (locus GAU_0412), (b) Acidithiobacillus ferrooxidans (locus AFE_0702), and (c) Chloroflexus aggregans (locus CAGG_0471). Copy number was analysed by qRT-PCR in cultures harvested during exponential phase and stationary phase, in the presence of either ambient H2 or 10% H2. Error bars show standard deviations of three biological replicates (averaged from two technical duplicates) per condition. Values denoted by different letters were determined to be statistically significant based on a one-way ANOVA with post-hoc Tukey’s multiple comparison (p < 0.05).
We subsequently monitored the consumption of H₂ by exponential and stationary phase cultures in ambient air supplemented with 10 ppmv H₂. For *G. aurantiaca* and *A. ferrooxidans*, H₂ was oxidised to sub-atmospheric levels under both conditions in an apparent first-order kinetic process (Fig. 2a, b). However, biomass-normalised first-order rate constants were higher in exponential than stationary phase cells by 23-fold ($p = 0.0029$) and 120-fold ($p < 0.0001$), respectively (Fig. 2d). For *C. aggregans*, H₂ was oxidised at rapid rates in exponentially growing cells, but occurred at extremely slow rates in stationary cells (Fig. 2c, d). These observations support the qRT-PCR results by showing hydrogenase activity predominantly occurs during growth. It should be noted that additional [NiFe]-hydrogenases are encoded by both *C. aggregans* (group 3d) and *A. ferrooxidans* (group 1e and 3b), but not for *G. aurantiaca*. These additional hydrogenases are expressed at tenfold lower levels for *C. aggregans*, but at similar levels for *A. ferrooxidans*, and hence may contribute to H₂ uptake (Fig. S2). It is nevertheless likely that the group 2a [NiFe]-hydrogenases mediate atmospheric H₂ uptake given (i) the H₂ uptake activities of *C. aggregans* and *A. ferrooxidans* mimic that of *G. aurantiaca*, which lacks additional hydrogenases; (ii) previous genetic studies show group 2a enzymes mediate high-affinity aerobic H₂ uptake in mycobacteria [12, 22]; and (iii) group 1e and 3b/3d enzymes are likely incapable of atmospheric H₂ oxidation given their respective characterised roles in anaerobic respiration and fermentation [25].

**H₂ consumption enhances mixotrophic growth in carbon-fixing strains**

The observation that expression and activity of the group 2a [NiFe]-hydrogenase is optimal during growth suggests this enzyme supports mixotrophic growth. To test this, we monitored growth by optical density of the three strains in headspaces containing H₂ at either ambient, 1%, or 10% mixing ratios. No growth differences in the obligate heterotroph *G. aurantiaca* were observed between the conditions ($p = 0.30$) (Fig. 3a). In contrast, H₂-dependent growth stimulation was observed for the obligate autotroph *A. ferrooxidans* (1.4-fold increase; $p = 0.0003$) (Fig. 3b) and facultative autotroph *C. aggregans* (1.2-fold increase; $p = 0.029$) (Fig. 3c). This suggests that reductant derived from H₂ oxidation can be used by these bacteria to fix CO₂ through the Calvin-Benson and 3-hydroxypropionate cycles, respectively.

**Hydrogenases with common phylogeny and genetic organisation are widely distributed across 13 bacterial phyla**

Finally, we surveyed the distribution of group 2a [NiFe]-hydrogenases to infer which other bacteria may oxidise atmospheric H₂. We detected the large subunit of this hydrogenase (HuCL) across 171 genera and 13 phyla (Table S2; Fig. S3); this constitutes a 3.2-fold increase in the number of genera and 1.4-fold increase in the number of
phyla previously reported to encode this enzyme [4, 25]. This increase in HucL distribution is due to the increase in available genome sequences since previous analyses were performed. The HucL-encoding bacteria include various known hydrogenotrophic aerobes, such as *Nitrospira moscoviensis* (Nitrospira) [30], *Hydrogenobacter thermophilus* (Aquificota) [49], *Kyrpida tusciae* (Firmicutes) [50, 51], *Sulfobacillus acidophilus* (Firmicutes) [52], and *Pseudonocardia dioxanivorans* (Actinobacteria) [53], suggesting these strains may also consume atmospheric H₂. The hydrogenase was also distributed in various lineages of Bacteroidota, Alphaproteobacteria, Gammaproteobacteria, and Deinococcota for which H₂ oxidation has not, to our knowledge, been reported.

A maximum-likelihood phylogenetic tree showed the retrieved HucL sequences form a well-supported monophyletic clade. Most sequences clustered into four major radiations, Bacteroidota-associated, Cyanobacteria-associated, Proteobacteria-associated (including *A. ferrooxidans*), and a mixed clade containing sequences from seven phyla (including *G. aurantiaca* and *C. aggregans*) (Fig. 4). Several genes were commonly genomically associated with *hucL* genes in putative operons, including the hydrogenase small subunit (*hucS*), a Rieske-type iron-sulphur protein (*hucE*) [33], hypothetical proteins (including NHI-repeat proteins) [32], and various maturation factors (Fig. S4). The group 2a [NiFe]-hydrogenases are distinct in both phylogeny and genetic organisation to the two most closely related hydrogenase subgroups, the previously described group 2e [NiFe]-hydrogenases of aerobic hydrogenotrophic Crenarchaeota [25, 54] and the novel group 2f [NiFe]-hydrogenases that are distributed sporadically in bacteria and archaea (Fig. 4).

**Discussion**

Overall, these findings demonstrate that atmospheric H₂ oxidation is not solely a persistence-linked trait. We infer that group 2a [NiFe]-hydrogenases are optimally expressed and active during exponential phase, consume H₂ at sub-atmospheric concentrations, and support mixotrophic growth. Largely concordant findings were made in three phylogenetically, physiologically, and ecologically distinct bacterial species. These findings contrast with multiple pure culture studies that have linked expression, activity, and phenotypes associated with group 1h [NiFe]-hydrogenases to survival rather than growth [10, 12, 16, 18, 20, 23, 24]. However, a growth-supporting role of atmospheric H₂ oxidation is nevertheless consistent with several surprising recent reports: the measurement of atmospheric H₂ oxidation during growth of several strains [12, 19, 23, 55]; the discovery of an Antarctic desert community driven by trace gas oxidation [9]; and the isolation of a proteobacterial methanotroph thought to grow on air alone [56]. Together, these findings suggest that the current persistence-centric model of atmospheric H₂ utilisation is overly generalised and that this process also supports mixotrophic growth.

Atmospheric H₂ oxidation during growth is likely to primarily benefit bacteria that adopt a mixotrophic lifestyle. While atmospheric H₂ alone can sustain bacterial maintenance, theoretical modelling suggests this energy source is insufficiently concentrated to permit growth as the sole energy source [1, 57]. Instead, bacteria that co-oxidise this dependable gas with other organic or inorganic energy sources may have significant selective advantages, especially in environments where resource availability is very low or variable. Likewise, it is probable that many bacteria in natural environments supplement growth by taking advantage of transient increases in H₂ availability. For example, the metabolic generalist *C. aggregans* may facilitate its expansion in geothermal mats by simultaneously utilising geothermal and atmospheric sources of H₂, in addition to sunlight and organic compounds [38, 39, 58]. Similarly, in the dynamic environment of wastewater treatment plants, *G. aurantiaca* may be well-suited to take...
Fig. 4 Radial phylogenetic tree showing the distribution and evolutionary history of the group 2a [NiFe]-hydrogenase. Amino acid sequences of the catalytic subunit of the group 2a [NiFe]-hydrogenase (hueL) are shown for 171 bacterial genera. The taxon names of the three study species, *G. aurantiaca*, *A. ferrooxidans*, and *C. aggregans*, are coloured in blue. The tree was constructed using the maximum-likelihood method (gaps treated with partial deletion), bootstrapped with 500 replicates, and rooted at the mid-point. Accession numbers and amino acid sequences used to construct the tree are listed in Table S2. The total number of genomes identified per phylum are as follows: Acidobacteriota (1), Actinobacteriota (27), Aquificota (4), Bacteroidota (34), Chloroflexota (5), Cyanobacteria (61), Deinococcota (2), Firmicutes (19), Gemmatimonadota (1), Myxococcota (1), Nitrospirota (2), Planctomycetota (1), Proteobacteria (49).
advantage of fermentatively-produced H₂ released during transitions between oxic and anoxic states [35, 59].

The ability to consume atmospheric H₂ may also be particularly advantageous during early stages of ecological succession. Indeed, *A. ferrooxidans* may initially rely on this atmospheric energy source as it colonises barren tailings and establishes an acidic microenvironment conducive for iron oxidation [60]. Hydrogen synthesis in tailings can further benefit *A. ferrooxidans* as acid conditions and more complex bacterial consortia develop. Specifically, acetate-dependent growth of dissimilatory sulphate reducing bacteria in tailings [58] will initiate endogenous geochemical production of trace hydrogen (FeS + H₂S $\rightarrow$ FeS₂ + H₂). As tailings cycle between aerobic (vadose) and anaerobic (water-saturating) conditions, the H₂ available from atmospheric and geochemical sources respectively may provide a continuous energy source for *A. ferrooxidans*. In addition, any environments possessing sulphate and iron, i.e., “downstream” from acid-generating ecosystems (including marine sediments), can generate hydrogen through bacterial organotrophic sulphate reduction.

This study also identifies key microbial and enzymatic players in the global hydrogen cycle. The group 2a [NiFe]-hydrogenase is the second hydrogenase lineage shown to have a role in atmospheric H₂ oxidation across multiple bacterial phyla. The group 1h enzyme is probably the main sink of the H₂ cycle given it is the predominant hydrogenase in most soils [4, 11, 61]. However, the group 2a enzyme is moderately to highly abundant in many soil, marine, and geothermal environments [62], among others, and hence is also likely to be a key regulator of H₂ fluxes. This study also reports atmospheric H₂ oxidation for the first time in two globally dominant phyla, Proteobacteria and Gemmatimonadota, and uncovers *A. ferrooxidans* as the first H₂-scavenging autotroph. Until recently, atmospheric H₂ oxidation was thought to be primarily mediated by heterotrophic Actinobacteriota [1, 10–12], but it is increasingly apparent that multiple aerobic lineages are responsible [4, 16, 17, 19, 33]. Some six phyla have now been described that are capable of atmospheric H₂ oxidation and, given the group 2a [NiFe]-hydrogenase is encoded by at least eight other phyla, others will likely soon be described. It is possible that atmospheric H₂ oxidation extends to other important groups, such as nitrite-oxidising Nitrospirota [30], methane-oxidising Proteobacteria [53], and potentially even oxygenic phototrophs; while Cyanobacteria are known to recycle endogenously-produced H₂ [26, 63, 64], it should be tested whether they can also scavenge exogenous H₂. Indeed, while atmospheric H₂ oxidisers were only recently discovered [10, 14, 65], it is now plausible that these bacteria may represent the rule rather than the exception among aerobic H₂ oxidisers.

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**Author contributions** CG and ZFI conceived this study. CG, ZFI, and RG supervised this study. CG, ZFI, and CW designed experiments. ZFI, CW, and KB performed experiments. ZFI, CW, and CG analysed data. EJG and GS contributed to study conception and experimental development. ZFI, CG, and CW wrote the paper with input from all authors.

**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

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