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The plant pathogen *Pectobacterium atrosepticum* contains a functional formate hydrogenlyase-2 complex

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Summary

*Pectobacterium atrosepticum* SCR1043 is a phytopathogenic Gram-negative enterobacterium. Genomic analysis has identified that genes required for both respiration and fermentation are expressed under anaerobic conditions. One set of anaerobically expressed genes is predicted to encode an important but poorly understood membrane-bound enzyme termed formate hydrogenlyase-2 (FHL-2), which has fascinating evolutionary links to the mitochondrial NADH dehydrogenase (Complex I). In this work, molecular genetic and biochemical approaches were taken to establish that FHL-2 is fully functional in *P. atrosepticum* and is the major source of molecular hydrogen gas generated by this bacterium. The FHL-2 complex was shown to comprise a rare example of an active [NiFe]-hydrogenase-4 (Hyd-4) isoenzyme, itself linked to an unusual selenium-free formate dehydrogenase in the final complex. In addition, further genetic dissection of the genes encoding the predicted membrane arm of FHL-2 established surprisingly that the majority of genes encoding this domain are not required for physiological hydrogen production activity. Overall, this study presents *P. atrosepticum* as a new model bacterial system for understanding anaerobic formate and hydrogen metabolism in general, and FHL-2 function and structure in particular.

Introduction

Many members of the γ-proteobacteria are facultative anaerobes with the ability to switch their metabolisms to exploit the prevailing environmental conditions. Aerobic or anaerobic respiration is generally preferred, depending on the availability of respiratory electron acceptors. In this phylum, and specifically under anaerobic conditions, the three-carbon product of glycolysis, pyruvate, is often further metabolised by the oxygen-sensitive pyruvate formate-lyase enzyme to generate acetyl CoA and the one-carbon compound formic acid (Pinske and Sawers, 2016). Studies of the model bacterium *Escherichia coli* have established that endogenously produced formate is initially excreted directly from the cell using a dedicated channel (Suppmann and Sawers, 1994; Hunger et al., 2014; Mukherjee et al., 2017). Under respiratory conditions this formate would be used as an electron donor through the activity of periplasmic enzymes, but under fermentative conditions the formate accumulates in the extracellular milieu until its rising concentration begins to cause a drop in extracellular pH. This is thought to trigger formate re-uptake, which in turn induces synthesis of formate hydrogenlyase (FHL) activity in the cell (Rossmann et al., 1991; McDowall et al., 2014; Sargent, 2016). FHL activity then proceeds to detoxify the formic acid by disproportionation to carbon dioxide and molecular hydrogen (H₂).

While FHL activity has been characterised in *E. coli* (Sargent, 2016), it is not confined to enteric bacteria and has been reported across the prokaryotic domains, including in hyperthermophilic archaea where it is not only involved in pH homeostasis but also in generating transmembrane ion gradients (Kim et al., 2010; Lim et al., 2014; Bae et al., 2015). The ion-pumping activity stems from an evolutionary link between FHL and the respiratory NADH dehydrogenase Complex I (Bohm et al., 1990; Friedrich and Scheide, 2000; Batista et al., 2013; Marreiros et al., 2013; Schut et al., 2016). Like Complex I, FHL comprises a cytoplasm-facing catalytic domain (termed the
Peripheral arm in Complex I terminology) linked to an integral membrane arm. In FHL, the peripheral arm contains a [NiFe]-hydrogenase of the ‘Group 4’ type, which is primarily dedicated to H₂ production (Greening et al., 2015), and is linked by [Fe-S]-cluster-containing proteins to a molybdenum-dependent formate dehydrogenase (Maia et al., 2015). The FHL membrane arm is predicted to take two different forms allowing the enzyme to be further sub-classified as either ‘FHL-1’ or ‘FHL-2’ (Friedrich and Scheide, 2000; Marreiros et al., 2013; Sargent, 2016; Finney and Sargent, 2019). The FHL-1 is the predominant archetypal FHL activity of E. coli (McDowall et al., 2014) and comprises [NiFe]-hydrogenase-3 (Hyd-3), formate dehydrogenase-H (FdhF), and a relatively small membrane arm compared to Complex I that contains only two proteins (Fig. 1A). Genes for the much less well-understood FHL-2 enzyme are also found in E. coli (Andrews et al., 1997). This isoenzyme is predicted to comprise a [NiFe]-hydrogenase-4 (Hyd-4), an as-yet undefined formate dehydrogenase, and a much larger membrane arm than FHL-1, containing at least five individual integral membrane subunits and more closely resembling the Complex I structure (Fig. 1B). Understanding the structure, function and physiological role of E. coli Hyd-4 and FHL-2 has been hindered by poor native expression levels (Skibinski et al., 2002; Self et al., 2004); a missing important accessory gene from the E. coli hyf cluster (Sargent, 2016); and a lack of consensus on the appropriate experimental conditions to test (Bagramyan et al., 2001; Mantsakanyan et al., 2004). Thus, in order to bring fresh impetus to understanding the physiology and biochemical characterisation of the FHL-2 complex, it was considered important that an appropriate alternative biological model system was established. In this work, Pectobacterium atrosepticumSCRI1043 was chosen (Bell et al., 2004; Babujee et al., 2012). However, very recently an operon encoding a Hyd-4 isoenzyme was cloned from Trabulsiella guamensis, which is a bacterium previously misidentified as a γ-proteobacterium that can grow under anaerobic conditions (Babujee et al., 2012). A plant pathogen with a functional FHL-2 complex (Babujee et al., 2012). Therefore, the initial goal of this study was to establish the growth conditions under which molecular hydrogen could be evolved. First, theSCRI1043 wild-type strain was grown under anaerobic fermentative conditions in a minimal medium supplemented with 0.8% (w/v) glucose. The culture headspace was sampled at periodic intervals and the amount of H₂ present quantified by gas chromatography (GC). Under these conditions, H₂ evolution activity was found to be temperature dependent, with H₂ accumulation in the headspace observed to be maximal when the phytopathogen was incubated at 20 or 24°C (Fig. 2A). Taking forward 24°C as standard incubation temperature, H₂ evolution was observed and found to level off after 40 h incubation (Fig. 2B).

When anaerobic respiratory conditions were tested, comprising 0.5% (v/v) glycerol and 0.4% (w/v) nitrate, H₂ production was found to cease with no H₂ detectable after 48 h growth (Fig. 2C). However, replacement of nitrate with 0.4% (w/v) fumarate as a terminal electron acceptor allowed the generation of low, but detectable, levels of H₂ (Fig. 2C). Maximal H₂ production is observable under fermentative conditions (Fig. 2C).
Hyd-4 is the predominant hydrogen-producing enzyme in *P. atrosepticum*

To determine the molecular basis of the observed H₂ production activity, a molecular genetic approach was taken. Initially, the genes encoding the catalytic subunits of the [NiFe]-hydrogenases were targeted. First, a strain PH001 (Table 1) was constructed carrying an unmarked in-frame deletion of the *hyfG* gene, predicted to encode the catalytic subunit of a Hyd-4 isoenzyme. When cultured fermentatively in the presence of glucose, the PH001 (Δ*hyfG*) strain produced less than 5% of the total H₂ accumulated by the wild-type control under the same conditions (Fig. 3A). Next, the gene encoding the catalytic subunit of Hyd-2 (*hybC*) was tested. Mutant strain PH002 (Table 1) was prepared carrying only a Δ*hybC* allele and, in this case, H₂ evolution under fermentative conditions was essentially indistinguishable from the wild-type strain (Fig. 3A). Finally, a Δ*hybC* Δ*hyfG* double mutant (PH003,

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**Fig. 1.** Biochemistry and genetics of formate hydrogenlyase. Structural models of (A) formate hydrogenlyase-1 (FHL-1) from *Escherichia coli* and (B) formate hydrogenlyase-2 (FHL-2) from *Pectobacterium atrosepticum*. Subunits related at the primary and tertiary levels are coloured similarly. Structural modelling of the formate hydrogenases complexes was performed using Phyre² predictions of respective subunits (Kelley and Sternberg, 2009). Using Chimera (Pettersen et al., 2004) and the Cryo-EM structure of the *Pyrococcus furiosus* Membrane Bound Hydrogenase, MBH (PDB: 6CFW), individual FHL-2 subunits were manually assembled. FdhF, which is not present in *Pyrococcus furiosus* MBH, was positioned principally to align its [4Fe–4S] cluster with that of the surface-exposed [4Fe–4S] cluster from HyfA. C. The genetic organisation of the hydrogen metabolism gene cluster of *P. atrosepticum* (ECA1225–ECA1252). Predicted gene product functions are indicated and the operon for Hyd-4 is colour coded to match the structure model in panel (B).

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**Fig. 2.** *P. atrosepticum* produces molecular hydrogen gas.
A. Anaerobic hydrogen production is optimal at lower temperatures. The *P. atrosepticum* SCR11043 parent strain was incubated in M9 medium supplemented with 0.8% (w/v) glucose for 168 h at the temperatures indicated before gaseous H₂ accumulation was quantified.
B. A time course of H₂ accumulation. *P. atrosepticum* SCR11043 was incubated in M9 medium supplemented with 0.8% (w/v) glucose at 24°C and gaseous H₂ accumulation was measured every 24 h.
C. *P. atrosepticum* SCR11043 was incubated in M9 medium supplemented with either 0.5% (v/v) glycerol and 0.4% (w/v) nitrate (‘Gly Nit’); 0.5% (v/v) glycerol and 0.4% (w/v) fumarate (‘Gly Fum’); 0.5% (v/v) glycerol only (Gly); or 0.8% (w/v) glucose only (‘Glc’) at 24°C for 48 h. In all cases, the levels of molecular H₂ in the culture headspace were quantified by GC and normalised to OD₆₀₀ and culture volume. Error bars represent SD (*n* = 3).
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a wider FHL-2 complex (Fig. 1). First, formate depen-
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Having established that Hyd-4 was active, the next task
contains an active FHL-2 with a
seelenium-free formate dehydrogenase
P. atrosepticum
containing a cysteine residue at position 140 (Fig. S1),
evolution could be rescued in the mutant strains by
H2 production remained dependent upon the
Table 1) was constructed and was found to be completely
duced very low, but detectable, levels of H2 gas under fer-
To that end, the data presented in Figs 2 and 3
demonstrate that Hyd-4 is responsible for the majority of
P. atrosepticum
a plant pathogen with a functional FHL-2 complex
Table 1. P. atrosepticum strains and plasmids used in this study.

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Plasmid Description Source

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P. atrosepticum can be stably transformed and plas-
mids encoding either hyfG or hybC were constructed. In
the case of PH001 (ΔhyfG) and PH003 (ΔhybC ΔhyfG),
H2 evolution could be rescued in the mutant strains by
supplying extra copies of hyfG on a plasmid (Fig. 3B).

Taken altogether, the data presented in Figs 2 and 3
demonstrate that Hyd-4 is responsible for the majority of
physiological H2 production by P. atrosepticum, and that
this activity is present under fermentative conditions at
temperate growth temperatures ≤ 24°C.

P. atrosepticum contains an active FHL-2 with a
seelenium-free formate dehydrogenase

Having established that Hyd-4 was active, the next task
was to test the hypothesis that Hyd-4 could be part of
a wider FHL-2 complex (Fig. 1). First, formate depend-
ence on H2 production was tested by growing the wild-
type parental strain, the PH001 (ΔhyfG) strain, and the
PH002 (ΔhybC) strain anaerobically in the presence of
increasing amounts of exogenous formate (Fig. 4A). A
correlation was observed between the amount of H2 pro-
duced and the amount of formate added to the growth
medium, and this was particularly clear when the uptake
hydrogenase activity was inactivated (Fig. 4A). High
levels of H2 production remained dependent upon the
presence of an active Hyd-4 (Fig. 4A), providing initial
evidence for a link between formate and H2 metabolism.

The P. atrosepticum SCR1043 genome contains a
gene encoding a putative formate dehydrogenase close
to those for Hyd-4 (Fig. 1C). The gene product shares 85%
overall sequence identity with E. coli FdhF but interest-
ingly contains a cysteine residue at position 140 (Fig. S1),
which is occupied by a critical selenocysteine in the E. coli
version and other related enzymes (Axley et al., 1991). A
mutant strain was therefore constructed (PH004, Table 1)
carrying a ΔfdhF allele. The PH004 (ΔfdhF) strain pro-
duced very low, but detectable, levels of H2 gas under fer-
mentative conditions (Fig. 4B). Addition of a ΔhybC allele
to the ΔfdhF strain to generate a double mutant (PH005,
Table 1) had no further effect on the amount of H2 that
could be produced (Fig. 4B).
It is interesting, however, that a strain devoid of both hybC (Hyd-2 activity) and hyfG (Hyd-4 activity) could not produce any H2 gas (Fig. 3B). This suggests that the low levels of H2 evolved from the ΔhybC, ΔfdhF strain are derived from Hyd-4 but that an alternative electron donor may be operating. Notably, two further genes encoding homologs of FdhF are encoded on the P. atrosepticum SCRI1043 chromosome (Bell et al., 2004). The ECA1507 gene encodes a protein with 65% overall sequence identity with FdhF, and the ECA1964 gene encodes a protein with 22% overall sequence identity with FdhF. Deletion of the genes encoding ECA1507 or ECA1964 alone (Table 1) had no influence on the H2 production capability of the bacterium (Fig. 4C). Moreover, when the genes were supplied in multicopy on an expression vector, neither was able to rescue the phenotype of the ΔfdhF mutant back to native levels of H2 production (Fig. 4D). However, it is clear that extra levels of ECA1507 in the cell result in a slight increase in H2 accumulation over the time course of the experiment (Fig. 4D). The increase in H2 production is statistically significant (P < 0.0001) and suggests ECA1507 could function as an alternative electron donor subunit for P. atrosepticum Hyd-4. Interestingly, extra copies of ECA1964 on a plasmid had the opposite effect (Fig. 4D). In this case, H2 production in the ΔfdhF strain was pushed to a statistically significant (P < 0.0001) even lower level (Fig. 4D), suggesting ECA1964 was either largely inactive or interfering with the interaction of Hyd-4 with other redox partners.

Taken altogether, these data establish that P. atrosepticum SCRI1043 has functional formate hydrogenlyase activity where molecular hydrogen production is clearly linked to both formate availability and the presence of a formate dehydrogenase gene. Importantly, the predominant electron donor for the reaction is an unusual version of formate dehydrogenase that does not require selenocysteine at its active site, and the enzyme responsible for proton reduction is a [NiFe]-hydrogenase-4.

The role of the FHL-2 membrane arm in hydrogen production

One clear defining structural difference between the FHL-1 type formate hydrogenlyase found in E. coli and the FHL-2 type of P. atrosepticum SCRI1043 is the number of genes encoding components of the membrane arms (Fig. 1). An FHL-1 enzyme is predicted to contain two different membrane proteins, HycC (related to HyfB in FHL-2) and HycD (related to HyfC) (Fig. 1A). Alternatively, an FHL-2 enzyme is predicted to contain three additional membrane proteins, including HyfE (not present in FHL-1) and two further homologs of HycC/HyfB, namely HyfD and HyfF (Fig. 1B).

To explore the roles of the extra hyfDEF genes located within the FHL-2 locus, mutant strains were constructed (Table 1). First, versions of the ΔhybC strain PH002, lacking either the genes encoding the entire FHL-2 membrane arm (PH007: ΔhybC, ΔhyfBCDEF – Table 1) or the extra membrane components not found in FHL-1 (PH008: ΔhybC, ΔhyfDEF – Table 1) were constructed. In addition, the ΔhybC strain PH002, producing Hyd-4 as the only active hydrogenase, was modified by addition of a 10-His sequence between codons 82 and

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83 of the hyfG gene. This new epitope-tagged strain was called PH009 (ΔhybC, hyfG^His – Table 1). Finally, versions of PH009 lacking either the genes encoding the entire FHL-2 membrane arm (PH020: ΔhybC, hyfG^His, ΔhyfBCDEF – Table 1) or lacking the extra membrane components not found in FHL-1 (PH021: ΔhybC, hyfG^His, ΔhyfDEF – Table 1) were constructed.

Deletion of the genes encoding the entire membrane arm reduced the FHL-2-dependent H₂ accumulation levels to around 5% of that observed in the parent strain (Fig. 5A). The addition of the 10-His tag to HyfG allowed the Hyd-4 catalytic subunit to be visualised in whole cell extracts by western immunoblotting (Fig. 5B). The polypeptide was clearly detectable when P. atrosepticum was cultured under anaerobic fermentative conditions (Fig. 5C). Interestingly, the amount of cellular HyfG^His was seen to increase when the genes encoding the membrane arm were removed (Fig. 5). This is particularly pertinent for the PH020 strain (ΔhybC, ΔhyfBCDEF), which is essentially devoid of FHL-2 activity (Fig. 5A), since it can be concluded that genetic removal of the complete membrane arm does not destabilise the Hyd-4 catalytic subunit, but instead leads to a physiologically inactive enzyme. It is also notable that in the absence of the genes...
encoding membrane proteins that the HyfG\(_{\text{His}}\) protein migrates as two electrophoretic species during SDS-PAGE (Fig. 5B). This is a common observation for catalytic subunits of [NiFe]-hydrogenases as they are synthesised as precursors that undergo proteolytic processing at the C-terminus following cofactor insertion (Bock et al., 2006). In this case, the faster migrating species was calculated as 56.4 kDa, while the slower migrating species was estimated as 62.6 kDa by SDS-PAGE. The predicted molecular mass of HyfG\(_{\text{His}}\) prior to proteolytic processing is 67,559 Da, and the predicted molecular weight of the 32-residue C-terminal tail that is removed is 3,821 Da.

Conversely, partial modification of the FHL-2 membrane arm to leave only those subunits present in FHL-1 (\(\Delta\text{hybC}\), \(\Delta\text{hyfDEF}\)) had no negative effect on hydrogen production levels (Fig. 5A), rather a slight increase was observed. This is consistent with a noticeable enhancement of HyfG\(_{\text{His}}\) levels in the cells upon removal of the hyfDEF genes (Fig. 5B). The available evidence suggests that HyfD, HyfE and HyfF have no essential roles.
in the biosynthesis and hydrogen production activity of FHL-2.

**A requirement for accessory genes in anaerobic hydrogen production**

FHL-2 is a multi-subunit metalloenzyme and assembly of such enzymes is often carefully co-ordinated by dedicated chaperones, sometimes called accessory proteins or ‘maturases’. Maturation of molybdenum-dependent formate dehydrogenases has been reported to require the action of an FdhD protein, which is believed to supply an essential sulfur ligand to the active site metal (Arnoux et al., 2015). In *P. atrosepticum* SCRI1043, *fdhD* (ECA0093) is not part of the FHL-2 locus but is located elsewhere on the chromosome next to a gene encoding superoxide dismutase (*sodA* or ECA0092) (Bell et al., 2004). Genetic modification of the PH002 strain, containing only Hyd-4 and FHL-2 activities, by the incorporation of a Δ*fdhD* allele (PH013: Δ*hybC*, Δ*fdhD* – Table 1) led to a defect in physiological H₂ production under fermentative conditions (Fig. 5D). This phenotype could be rescued by the provision of extra copies of *fdhD* in trans (Fig. 5D).

Maturation of [NiFe]-hydrogenases requires the activity of a network of proteins involved in metal homoeostasis and cofactor maturation and insertion (Sargent, 2016). The *P. atrosepticum* FHL-2 locus (Fig. 1C) contains a *hoxN* gene (ECA1252) encoding a putative membrane-bound nickel ion transporter (Eitinger and Mandrand-Berthelot, 2000). Deletion of the *hoxN* gene in *P. atrosepticum* SCRI1043 (strain PH011, Table 1) reduced hydrogen evolution levels to around 50% of that observed for the parental strain (Fig. 5D). Note that there is no other homologue of *hoxN* encoded on the *P. atrosepticum* SCRI1043 genome, but there are several uncharacterised ABC transporters that could be related to the high-affinity *nkaA* system (Wu et al., 1991), which could account for the continued availability of nickel for hydrogenase biosynthesis in this experiment.

Once inside the cell, nickel is processed into the Ni–Fe–CO–2CN⁻ cofactor through the action of several enzymes and chaperones. One key step in the biosynthesis of the cofactor is the first step in the generation of CN⁻ from carbamoyl phosphate by HypF (Sargent, 2016). Deletion of the *hypF* gene from *P. atrosepticum* (PH010, Table 1), which is located in the hydrogen metabolism gene cluster under investigation here (Fig. 1C), led to the complete abolition of all detectable H₂ evolution (Fig. 5D). The mutant phenotype could be rescued by supply of *hypF* in trans, but note that full H₂ evolution levels were not restored (Fig. 5D).

Finally, it was observed that a member of the HyfR family of transcriptional regulators was encoded in the hydrogen metabolism gene cluster (Fig. 1C). The HyfR protein is predicted to be related to FhlA, which is a formate-sensing transcriptional activator (Skibinski et al., 2002). A Δ*hyfR* strain devoid of the HyfR protein has very low formate hydrogenlyase-2 activity (Fig. 5D).

Taken altogether, it can be concluded that all of the genes required for biosynthesis of FHL-2 are functional in *P. atrosepticum* SCRI1043, which is entirely consistent with the physiological data reported here.

**Discussion**

**Key differences between FHL-2 and FHL-1**

Formate hydrogenlyases can be classified into two structural classes, FHL-1 and FHL-2 (Finney and Sargent, 2019). The most obvious structural difference between an FHL-1, such as the best-characterised *E. coli* enzyme (McDowall et al., 2014; Pinske and Sargent, 2016), and an FHL-2, such as the *P. atrosepticum* enzyme characterised here, is the predicted size and composition of the membrane arm (Fig. 1B). Indeed, this large membrane arm is thought to be the ancient progenitor to the ion-pumping membrane arm of respiratory Complex I (Batista et al., 2013; Marreiros et al., 2013; Yu et al., 2018). Although eukaryotic Complex I, prokaryotic Complex I, and Group 4 hydrogenases such as FHL-1, FHL-2, Ech and MBH are clearly evolutionarily related, the gene and protein names for each type of enzyme are different. Some review articles contain useful tables to highlight the relatedness of the individual subunits (Friedrich and Scheide, 2000; Marreiros et al., 2013; Schut et al., 2016). FHL-1 includes only two membrane proteins, which are a single HycD/HyfC-type protein together with a single HycC/HyfB. This is sufficient to anchor the peripheral arm close to the membrane and, in the case of *Thermococcus onnurineus* FHL-1 (Lim et al., 2014) and the related Ech hydrogenase from *Methanosarcina mazei* (Welte et al., 2010), will also allow initial generation of a proton gradient (Yu et al., 2018). Operons encoding FHL-2 complexes encode at least three further integral membrane proteins. In *P. atrosepticum* these are HyfD and HyfF, which are extra versions of the HycC/HyfB putative ion channels, and the HyfE protein, which is more closely related to a region of NuoK in Complex I. Interestingly, if FHL-2 is modelled based on the Complex I structure (Fig. S2), the extra HyfDEF proteins would be placed between HyfBC, thus separating them and pushing HyfB to the most distal point in the peripheral arm (Marreiros et al., 2013). Alternatively, if FHL-2 is modelled based on the Hyd-4-like MBH structure from *Pyrococcus furiosus* (Yu et al., 2018), then HyfBC remain in contact with each other and HyfDEF form the distal region of the membrane arm (Figs 1 and S2). The experimental evidence presented in this work suggests *P. atrosepticum* FHL-2 adopts a membrane arm architecture similar to the *Pyrococcus furiosus*...
MBH hydrogenase (Fig. 1). This is because removal of all of the extra HyfDEF membrane proteins from FHL-2 had no discernible effect on the physiological activity of the P. atrosepticum system (Fig. 5A), suggesting an active FHL-1-like core enzyme remains. Clearly if HyfB was normally separated from HyfC by the extra proteins they would be unlikely to come together to form a complex when placed in a ΔhyfDEF background, and E. coli FHL-1, for instance, is completely inactive when lacking its HyfB homolog HycC (Pinske and Sargent, 2016). This highlights the principle of modularity in metalloenzyme evolution, since it is clear that the HyfDEF module may be added or removed depending on both selective pressure and also the, as yet undefined in terms of hydrogenases, biochemical function of these membrane proteins (Friedrich and Scheide, 2000). Indeed, it is notable that distal components of the Pyrococcus furiosus MBH membrane arm (MbhABC) could also be genetically removed with only minor effects on cellular hydrogenase activity (Yu et al., 2018). Taken together, this perhaps points to Hyd-3 from FHL-1 as the minimal module of a Group 4 hydrogenase.

Western immunoblotting pointed towards either stabilisation or upregulation of the catalytic subunit HyfG in the absence of hyfDEF or hyfBCDEF (Fig. 5B). This is unlikely to be caused by an accumulation of formate in the cells, perhaps leading to maximal transcription, because the ΔhyfDEF strain retained normal levels of formate hydrogenlyase activity (Fig. 5A). It is more likely that the removal of genes encoding large membrane proteins from immediately upstream of hyfG relaxes some restrictions on the rates of transcription and translation. In bacteria, transcription, translation and membrane insertion of the nascent chain are thought to be coupled together in a process called transertion (Roggiani and Goulian, 2015), and removal of some or all of the elaborate membrane integration step could have an effect on translation of downstream genes.

At native levels, the HyfGHis protein can be detected as a single species migrating at 56.4 kDa in SDS-PAGE (Fig. 5B), and occasionally a slower migrating form is as a single species migrating at 56.4 kDa in SDS-PAGE (Fig. 5B and C); however, only the mature form of HyfG could contribute to physiological formate hydrogenlyase activity.

The P. atrosepticum HyfG catalytic subunit from the Hydrogenase-4 component of FHL-2 shares 74% overall sequence identity (85% similarity) with the E. coli HycE protein from Hydrogenase-3/FHL-1. The sequence variation between these two Group 4A hydrogenases is therefore small with only subtle notable differences. For instance, each protein is known or predicted to undergo cleavage during cofactor insertion and maturation leaving a C-terminal arginine residue in the mature form of the proteins. The cleavage sites themselves are slightly differently conserved in an FHL-1-type enzyme compared to an FHL-2, for example, …R*MTVV… for HycE-like proteins compared to …R*VTLV… for HyfG. This may reflect the need for a different maturation protease for each type of hydrogenase, however, this remains to be tested experimentally. In addition, it is notable that both E. coli and P. atrosepticum hyfG initiate translation with a GUG start codon, which may have a role in controlling cellular levels of the enzyme (Belinky et al., 2017).

Phylogenetic analysis of the Group 4A [NiFe]-hydrogenase subunits, including HycE and HyfG, shows that the enzymes associated with FHL-1 separate into a clearly distinct evolutionary clade from those associated with FHL-2, which form their own distinct clade (Fig. S3). Examples of species that encode both FHL-1 and FHL-2 are rare (Fig. S3).

A selenium-free formate dehydrogenase

Arguably one of the best-studied FdhF enzymes is the E. coli version, which contains selenocysteine at its active site (Axley et al., 1991; Gladyshev et al., 1994; Boyington et al., 1997). Selenocysteine is incorporated co-translationally at a special UGA ‘nonsense’ codon within the coding sequence (Zinoni et al., 1987), and replacement of selenocysteine with cysteine in the E. coli enzyme resulted in a dramatically reduced turnover number (Axley et al., 1991). One surprising aspect of P. atrosepticum SCR1043 is that it contains none of the biosynthetic machinery to synthesise selenocysteine (Babujee et al., 2012) and the fdhF gene studied in this work contains a cysteine codon where selenocysteine would be encoded in the E. coli enzyme (Fig. S1). Certainly, the discovery of an active FHL-2 with no need for selenocysteine would benefit scientists interested in engineering this activity into other biological systems. Indeed, an in-frame deletion in the fdhF gene located in the FHL-2 gene cluster (Fig. 1C) resulted in a ~ 500 times reduction in H2 production (Fig. 4), indicating the majority of H2 production from P. atrosepticum is dependent on this formate dehydrogenase engaging with Hyd-4 to
form an FHL-2 complex. However, the ΔhybC ΔfdhF double mutant still produced low, but quantifiable, levels of H₂ (Fig. 4). Compare that with the behaviours of the ΔhybC ΔfdhF strain (Fig. 3B) and the ΔhypF mutant (Fig. 5C), neither of which produced any detectable H₂ gas. This genetic approach points to the residual H₂ emitting from Hyd-4, perhaps through the activity of alternative electron donors. Certainly for the E. coli FHL, it is known that FdhF is only loosely attached (Boyington et al., 1997) and this may be because the enzyme is ‘moonlighting’ in other biochemical pathways (Iwadate and Kato, 2019). It raises the possibility that other FdhF-like enzymes in particular could ‘plug in’ to Hyd-4 and pass excess reducing electrons on to protons. In this work, ECA1507 was found to partially rescue the phenotype of a ΔfdhF strain (Fig. 4D) suggesting it could be an alternative redox partner: note well, however, that the potential substrates and kinetics of ECA1507 cannot be reliably predicted and should be determined empirically.

The FdhF formate dehydrogenase from P. atrosepticum shares 65% overall sequence identity (and 85% similarity) with the well-known E. coli enzyme (Fig. S1). Interestingly, phylogenetic analysis suggests that > 50% of bacterial species that contain FHL genes utilise a cysteine-dependent, rather than selenocysteine-dependent, formate dehydrogenase (Fig. S4). P. atrosepticum ECA1507 and ECA1964 were identified here as two FdhF-like proteins that could potentially interact with Hydrogenase-4 to generate novel FHL-like complexes. Sequence analysis revealed ECA1507 and ECA1964 share 65% and 22% overall sequence identity with FdhF, respectively, and phylogenetic analysis determined that ECA1964 is more similar to E. coli YdeP than any other predicted molybdenum dependent oxidoreductases in P. atrosepticum (Fig. S5). YdeP has a putative role in acid resistance in E. coli (Masuda and Church, 2003).

A role for formate metabolism in a plant pathogen

In the potato pathogen P. atrosepticum, FHL-2 activity was found to be expressed at lower growth temperatures (Fig. 2). This suggests that FHL-2 may be produced in planta during the infection or colonisation event. Formate is produced endogenously by enteric bacteria under fermentative conditions, but plants and tubers have multiple metabolic pathways that generate and consume formate. Potato tubers produce a NAD⁺-dependent formate dehydrogenase (FDH), and the levels of this enzyme are boosted under stress conditions (Hourton-Cabassa et al., 1998). Indeed, proteomic experiments have identified FDH as a differentially produced protein during wound healing in potato tuber slices, with order of magnitude level changes in protein during this process (Chaves et al., 2009). It could be hypothesised that the expression of FDH in the potato tuber could be coordinated with the initial secretion of formate by a fermenting pathogen. Potentially, this would generate NADH from formate in stressed or damaged plant tissues. Recently, it was shown that FDH co-ordinates cell death and defence responses to phytopathogens in Capsicum annum (Bell pepper) (Choi et al., 2014). There is also indication that formate and other molecules that lead to the generation of formate, such as methanol and formaldehyde, induce the production of the NAD⁺-dependent FDH, perhaps suggesting there is a signalling response to these C1 compounds in plants (Hourton-Cabassa et al., 1998).

Concluding remarks

In this work, P. atrosepticumSCRI1043 has been established as a tractable new model organism for studying hydrogen metabolism in general and FHL function in particular. The organism is a rare example of a bacterium with an active hydrogenase-4-containing FHL-2 complex, however, in the course of this work, hydrogenase-4 activity was reported in T. guamensis, another γ-proteobacterium (Lindenstraß and Pinske, 2019). Interesting, the T. guamensis Hyd-4 was found to be active in vivo but very poorly reactive in vitro in standard enzymatic assays with redox-active dyes (Lindenstraß and Pinske, 2019). This again highlights the need for development of new approaches to characterise FHL-2 and its component parts. In P. atrosepticum, the active hydrogenase-4 enzyme operates in tandem with an unusual selenium-free formate dehydrogenase, which may be more amenable to biotechnological engineering than selenium-dependent isoenzymes. In evolutionary terms, the FHL-2 complex has been discussed as a key intermediate in the evolution of the NADH dehydrogenase (Complex I) from a structurally simpler membrane-bound hydrogenase (Friedrich and Scheide, 2000; Marreiros et al., 2013; Schut et al., 2016). The most obvious difference in the predicted quaternary structures inferred from the genetics is the large membrane arm present in FHL-2 compared to FHL-1, and data presented here points to the extra membrane proteins being not essential for formate-dependent hydrogen evolution in vivo. The role of the FHL membrane arm in generating a transmembrane ion gradient remains to be fully explored in enteric bacteria.

Experimental procedures

Bacterial strains

The parental P. atrosepticum strain used in this study wasSCRI1043 (Bell et al., 2004). In-frame deletion and insertion mutants were constructed using pKNG101 suicide vector.
in *E. coli* strain CC118:pIR (Kaniga et al., 1991; Coulthurst et al., 2006). Briefly, upstream and downstream regions (≥ 600 bp) of the target gene(s) was amplified and inserted into pKNG101 using a three fragment Gibson assembly reaction (HiFi Assembly, NEB). For the insertion of a deca-His encoding sequence into hyfG, primers were designed using the NEBuilder online tool to include the deca-His encoding sequence in the overlapping region of the two fragments containing the respective 3’ and 5’ sequences of hyfG. After successfully assembly and sequencing of pKNG101 plasmids, the CC118:pIR strain with desired plasmid, a HH26 pNJS5000 helper strain, and the desired *P. atrosepticum* strain were grown in rich media, with antibiotics as necessary. Equal volumes of the stationary phase cultures were mixed and 30 µl was spotted on a non-selective rich media plate for 24 h at 24°C. *P. atrosepticum* cells with the pKNG101 plasmid were initially selected for on minimal media agar plate for 24 h at 24°C. *P. atrosepticum* cells with the pKNG101 plasmid were initially selected for on minimal media agar plate and were then transferred to stationery phase in rich medium with no selection before the culture was diluted 1/500 with phosphate buffer. Then, 30 µl of this diluted culture was plated on minimal media agar with sucrose. These colonies were then screened for sensitivity to streptomycin before PCR screens were performed to check for presence of the desired mutation(s).

### Plasmids and complementation

All plasmids were cloned using Gibson assembly (HiFi Assembly, NEB) using DNA amplified from *P. atrosepticum* SCR11043 genomic DNA (Table 1). Genes were cloned into pSU-PROM (KanR), which includes the constitutive tatA promoter from *E. coli* (Jack et al., 2004). Complementation plasmids were used to transform electrocompetent *P. atrosepticum* cells using a 2 mm electroporation cuvette (Molecular BioProduct) with application of an electrical pulse (2.5 kV voltage, 25 µF capacitance, 200 Ω resistance and 2 mm cuvette length) via a Gene Pulsar Xcell electroporator (BioRad). Post recovery, cells were plated on LB Lennox agar plates with sucrose and 50 µg ml⁻¹ kanamycin.

### Hydrogen quantification

Hydrogen was directly quantified from 5 ml cultures grown in sealed Hungate tubes (Pinske and Sargent, 2016). Gas headspace samples were collected using a syringe with Luer lock valve (SGE). Samples were analysed using gas chromatography (Shimadzu GC-2014, capillary column, TCD detector). A hydrogen standard curve was used to quantify sample hydrogen content, this was then normalised to optical density (OD₆₀₀) and culture volume (Pinske and Sargent, 2016).

### Western immunoblotting

Proteins samples were first separated by SDS-PAGE using the method of Laemmli (1970) before transfer to nitrocellulose (Dunn, 1986). Nitrocellulose membranes were challenged with an anti-His-HRP antibody (Alpha Diagnostics) and a GeneGnome instrument (SynGene) was used to visualise immunoreactive bands following addition of ECL reagent (Bio-Rad).

### Structure modelling and phylogenetic analysis

Structural modelling of the formate hydrogenlyases complexes was performed using Phyre² predictions of respective subunits (Kelley and Sternberg, 2009). Using Chimera (Petterson et al., 2004), the X-ray crystal structure of *Thermus thermophilus* Respiratory Complex I (4HEA) and the Cryo-EM structure of membrane-bound hydrogenase (6CFW), the individual FHL-2 subunits were manually assembled into a putative complex organisation for FHL-1 and FHL-2. Phylogenetic analysis of *E. coli* FdhF-like proteins from organisms possessing a Group 4A [NiFe]-hydrogenase utilised the HydDB database (Greening et al., 2015) to collect accession numbers for all [NiFe]-hydrogenase subunits. In each organism, the FdhF orthologs were identified before MUSCLE multiple sequence alignment in Jalview (Waterhouse et al., 2009). Through percentage identity tree generation and manual inspection, the closest FdhF-like proteins in each organism were identified. FigTree (http://tree.bio.ed.ac.uk/software/figtree) was used to visualise the finalised phylogenetic trees.

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### Conflict of interest

The authors declare no conflict of interest.

### Author contributions

AJF Was a PhD student who designed experiments, analysed data, prepared figures for publication and wrote the paper. RL and MF were undergraduate project students who performed experiments and analysed data. MA was a Marie Skłodowska-Curie Independent Fellow who supervised the research, performed experiments and analysed data. SJC was a Wellcome Trust Senior Research Fellow who designed the research, supervised the research, analysed data and wrote the paper. FS conceived the project, assembled the research team, designed the research, supervised the research, analysed data and wrote the paper.

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