

1 **How a Formate Dehydrogenase Responds to Oxygen:**
2 **Unexpected O₂ Insensitivity of an Enzyme Harboring**
3 **Tungstopterin, Selenocysteine, and [4Fe-4S] Clusters**

4
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13 **KEYWORDS** Oxygen-insensitive metal-dependent formate dehydrogenase, Formate
14 oxidase, Tungsten, Pterin, Selenium, Iron-sulfur cluster, Electron Paramagnetic Resonance,
15 ¹³C NMR, Oximetry, Hydrogen peroxide, Electron bifurcation, Sulfate-reducing bacteria,
16 *Desulfovibrio vulgaris* Hildenborough.

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1 **ABSTRACT**

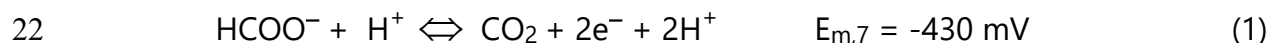
2 The reversible two-electron interconversion of formate and CO₂ is catalyzed by both non-
3 metallo and metallo-formate dehydrogenases (FDHs). The latter group comprises
4 molybdenum- or tungsten-containing enzymes with the metal coordinated by two
5 equivalents of a pyranopterin cofactor, a cysteinyl or selenocysteinyl ligand supplied by
6 the polypeptide, and a catalytically essential terminal sulfido ligand. In addition, these
7 biocatalysts incorporate one or more [4Fe-4S] clusters for facilitating long-distance
8 electron transfer. But an interesting dichotomy arises when attempting to understand how
9 the metallo-FDHs react with O₂. Whereas existing scholarship portrays these enzymes as
10 being unable to perform in air due to extreme O₂ lability of their metal centers, studies
11 dating as far back as the 1930s emphasize that some of these systems exhibit formate
12 oxidase (FOX) activity, coupling formate oxidation to O₂ reduction. Therefore, to reconcile
13 these conflicting views, we explored context-dependent functional linkages between
14 metallo-FDHs and their cognate electron acceptors within the same organism vis-à-vis
15 catalysis under atmospheric conditions. Here, we report the discovery and
16 characterization of an O₂-insensitive FDH2 from the sulfate-reducing bacterium
17 *Desulfovibrio vulgaris* Hildenborough that ligates tungsten, selenocysteine, and four [4Fe-
18 4S] clusters. Notably, we advance a robust expression platform for its recombinant
19 production, eliminating both the requirement of nitrate or azide during purification and
20 reductive activation with thiols and/or formate prior to catalysis. Because the distinctive
21 spectral signatures of formate-reduced DvH-FDH2 remain invariant under anaerobic and
22 aerobic conditions, we benchmarked the enzyme activity in air, identifying CO₂ as the
23 bona fide product of catalysis. Full reaction progress curve analysis uncovers a high
24 catalytic efficiency when probed with an artificial electron acceptor pair. Furthermore, we
25 show that DvH-FDH2 enables hydrogen peroxide production sans superoxide release to

1 achieve O₂ insensitivity. Direct electron transfer to cytochrome *c* in air also reveals that
2 electron bifurcation is operational in this system. Taken together, our work
3 unambiguously proves for the first time the coexistence of redox bifurcated FDH and FOX
4 activities within a metallo-FDH scaffold. These findings have important implications for
5 engineering O₂-tolerant FDHs and bio-inspired artificial metallocatalysts, as well as for the
6 development of authentic formate/air biofuel cells, modulation of catalytic bias, assessing
7 the limits of reversible catalysis, understanding directional electron transfer, and
8 discerning formate bioenergetics of gut microbiota.

9 INTRODUCTION

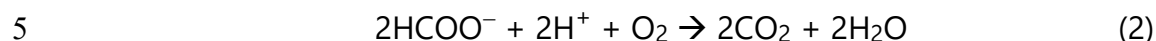
10 The simplest carboxylic acid (formic acid), and its conjugate base (formate) are
11 normal products of metabolic activity in living organisms, including bacteria and humans.¹
12 However, bacterial aerobic respiration of formate derived from human gut microbiota
13 drives inflammatory dysbiosis.² Although formic acid is primarily used as a food
14 preservative (E236) or as silage additive for maintaining the nutritive value of animal feed,³
15 it is a highly sought-after electron-mediator and feedstock in (electro)microbial
16 bioproduction,⁴ as well as a low carbon-footprint molecule that serves as a chemically
17 robust hydrogen storage medium.⁵ In addition to being a carbon and energy source for
18 the (an)aerobic growth of disparate bacteria,⁶ archaea,⁷ and syntrophic consortia,⁸
19 formate can be generated abiotically from CO₂ and renewable electricity.⁵

20 Formate oxidation and CO₂ reduction are interconvertible processes that are
21 carried out by prokaryotic formate dehydrogenases (FDHs) (Reaction 1).^{9, 10}



1 There are two phylogenetically distinct FDH families that can be distinguished by their
2 transition metal ion requirement for enzyme activity.¹¹ Metallo-FDHs are thought to be
3 highly sensitive to O₂,^{12, 13} necessitating catalytic measurements under anaerobic
4 conditions. However, available data in the primary literature are more confusing than
5 definitive. For example, DvH-FDH3 has been reproducibly shown to be O₂ sensitive^{14, 15}
6 while its ortholog from *D. desulfuricans* ATCC 27774 (Dd) can be purified in air.^{16, 17}
7 Similarly, *D. gigas* (Dg) FDH1 is readily isolated and stored under atmospheric conditions¹⁸
8 ¹⁹ but its counterpart from DvH requires the presence of 10 mM sodium nitrate to prevent
9 O₂ inactivation.²⁰ Regardless of the purification protocols, the resulting enzymes are
10 'dead on arrival' in that they must be resurrected by lengthy incubations with high
11 concentrations of thiols (10 – 50 mM dithiothreitol²⁰ for DvH-FDH1 and 130 mM β-
12 mercaptoethanol^{16, 17, 19} for Dd-FDH3 and Dg-FDH1) and/or formate^{17, 21} prior to catalytic
13 measurements under anaerobic conditions (a representative example can be found in
14 Figure S5a of Oliveira et al²⁰). A satisfactory molecular explanation for these
15 phenomenological observations has not been forthcoming for over three decades.¹⁸ The
16 situation is equally unclear with FDHs isolated from organisms other than sulfate-reducing
17 bacteria (SRB). *Escherichia coli* Fdh-H has been purified and characterized in the presence
18 of sodium azide to minimize O₂ inactivation.^{22, 23} 10 mM sodium nitrate^{24, 25} or azide^{26, 27}
19 has been added as stabilizers during the isolation of other bacterial metallo-FDHs as well.
20 Very little is known about how these small molecules protect the enzyme from O₂.
21 Because stability in air does not enable aerobic catalysis, the inhibitors are either removed
22 prior to measurements under anaerobic conditions^{20, 25} or allowed to remain while the
23 activity is probed anaerobically²³ or in air.²⁸ To our knowledge, no FDH has been shown
24 to reversibly interconvert formate and CO₂ in air. Claims to the contrary have been
25 considered as experimental artifacts.²⁵ Moreover, mechanistic details regarding how O₂
26 reacts with these metalloenzymes are not available.

1 Largely overlooked, however, is the fact that the present-day claims of FDH O₂
2 sensitivity fail to recognize or rationalize the findings reported between the late 1920s
3 and early-1990s vis-à-vis existence of metallo-FDHs capable of oxidizing formate with
4 oxygen (Reaction 2).

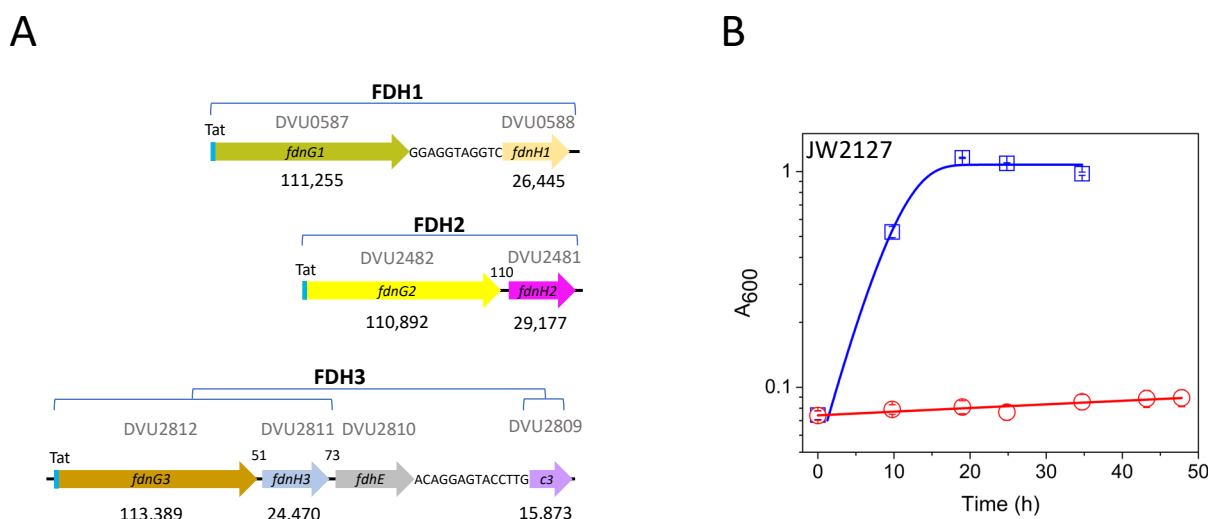


6 Starting with the first purification of a bacterial FDH by Stickland in 1929, O₂ uptake served
7 as a proxy for measuring enzyme activity.²⁹ Subsequently, Stephenson and Stickland
8 isolated *Escherichia coli* hydrogenlyase (Fdh-H), revealing that it was not responsible for
9 the FOX activity.³⁰ A key insight regarding the latter came from Gale's demonstration that
10 formate dependent O₂ consumption by *E. coli* was higher in aerobically grown cells.³¹
11 Pinsent's groundbreaking study relied on Gale's O₂ utilization assay to convincingly show
12 that not only *E. coli* FDH requires molybdenum and selenium for function, but more
13 importantly, that it retained robust FOX activity.³² This has since been independently
14 confirmed by several research laboratories.³³⁻³⁷ Pichinoty went on to show that FOX
15 activity was broadly distributed across bacteria.³⁸ Collectively, these findings paved the
16 way for Sawers to discover the third Fdh-O (O for oxidase; the remaining two being Fdh-
17 N (nitrate)³⁹ and Fdh-H (hydrogen)⁴⁰) in *E. coli*.^{9, 41, 42} Although others have confirmed the
18 presence of Fdh-O,^{43, 44} isolation and characterization of a metallo-FDH with FOX activity
19 has remained intractable.⁴⁵ And the possibility that coexistence of dehydrogenase and
20 oxidase activities would render a metallo-FDH insensitive to O₂ has not been entertained
21 thus far. To tackle this challenge, one of our laboratories (C.S.R.) advanced and tested the
22 hypothesis that FDHs capable of transferring electrons to natural high potential acceptors
23 are likely to be O₂-insensitive by virtue of their FOX activity, for such physiological
24 reactions are poised to occur under aerobic conditions. Despite the paucity of
25 information regarding redox partners (two well characterized systems exhibit low

1 reduction potentials.^{18, 46}), our central hypothesis was inspired by Yagi's 1969 observation
2 that an FDH from *D. vulgaris* Miyazaki (DvM) preferentially transfers electrons to a high-
3 potential cytochrome *c*₅₅₃.^{47, 48} Because the genetically tractable DvH^{49, 50} is closely related
4 to DvM⁵¹ and encodes a 73% identical cytochrome *c*₅₅₃ ($E_{m,7} = +62$ mV),⁵² we probed the
5 O₂ sensitivity of periplasmic FDHs from this SRB shown to thrive in microaerobic niches.⁵³⁻
6 ⁵⁵ We focused on the poorly characterized DvH-FDH2 (locus tag DVU2482-2481)^{56, 57} and
7 cytochrome *c*₅₅₃-reducing DvH-FDH3 (locus tag DVU2812-2809)^{14, 15} instead of the well-
8 studied DvH-FDH1 (locus tag DVU0587-0588)²⁰, which couples anaerobic formate
9 oxidation to sulfate reduction by initiating electron transfer to a low-potential cytochrome
10 *c*₃ ($E_{m,7} = -350$ mV).⁵⁰ Here, we describe our discovery and characterization of an O₂-
11 insensitive FDH that retains both dehydrogenase and oxidase activities.

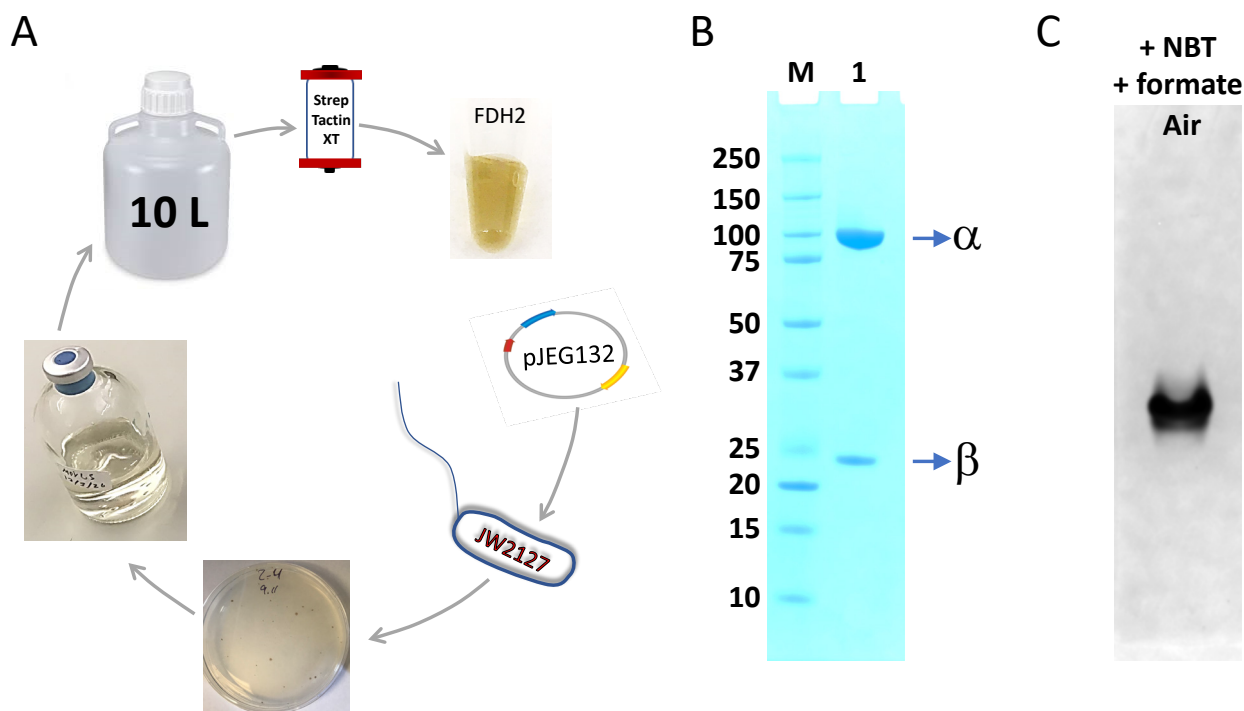
12 **RESULTS AND DISCUSSION**

13 **Robust Expression Platform for Facile Production of Highly Pure O₂-Insensitive**
14 **Metallo-FDHs.** There are three distinct *fdh* loci in the DvH genome⁵⁷ (Figure 1A). Only
15 FDH1 encoded by the first locus is essential for growth when sulfate and formate serve as
16 electron acceptor and donor, respectively.⁵⁰ The cellular function of FDH2 and FDH3 are
17 not well defined. Oliveira *et al*²⁰ expressed FDH1 in a $\Delta fdh1$ deletion strain using the
18 vectors and strategies developed in one of our laboratories (J.D.W.). We took a different
19 approach. We reasoned that construction of a markerless FDH-free strain could be
20 beneficial on three fronts: (a) Facilitate biochemical investigations of a native or foreign
21 FDH without potential interference from host counterparts, (b) Benchmark whole cell
22 biocatalysts, and (c) Uncover how synergy between enzyme catalysis and bioenergetics
23 modulates organismal dynamics. To that end, we generated a DvH strain (JW2127; see
24 Methods; Tables S1 and S2) that is devoid of all three *fdh* loci. Although JW2127 is unable
25 to grow on formate-acetate-sulfate, it maintains wild-type-like growth profile on lactate-



1
 2 *Figure 1. Structure and function of FDH operons in DvH: (A) Condensed map of the three fdh loci. fdnG2*
 3 *(yellow; large subunit) and fdnH2 (magenta; small subunit) encode FDH2 investigated in this work and are*
 4 *part of a five gene (not shown) operon. Short intergenic regions are illustrated at the nucleotide level while the*
 5 *length of their long counterparts is identified by two- or three-digit numbers. Periplasmic FDH localization is*
 6 *made possible by the twin-arginine translocation (Tat) signal peptide (cyan). Theoretical molecular masses of*
 7 *the encoded polypeptide in daltons are listed below each gene. (B) Growth curves of JW2127: formate-acetate-*
 8 *sulfate (red), lactate-sulfate (blue). The lines going through the points represent fits to Weibull⁵⁸ growth model.*
 9 *Error bars represent standard deviations from three independent measurements.*

10
 11 sulfate medium (Figure 1B). We also constructed deletion strains harboring different
 12 combinations of *fdh* genes for functional analyses, including JW2111 ($\Delta fdh3$) and JW2121
 13 ($\Delta fdh1$ and $\Delta fdh3$; see Tables S1 and S2). The latter two served as controls in this study
 14 (Figure S1). Subsequently, we used JW2127 for the homologous expression of FDH2.
 15 Introduction of a Strep-tag II at the C-terminus of the large subunit facilitated one-step
 16 affinity purification. Whereas Oliveira *et al*²⁰ used DvH cells derived from 300 L
 17 fermentation to purify FDH1, we have streamlined our workflow to produce 1.8 mg of
 18 highly-pure heterodimeric FDH2 from a gram of wet cell paste (Figures 2A and 2B). Thus,
 19 our 10 L culture (biomass yield of ~ 8 g) generates sufficient protein to tackle even the
 20 most demanding experiments. And our method can be readily scaled up.



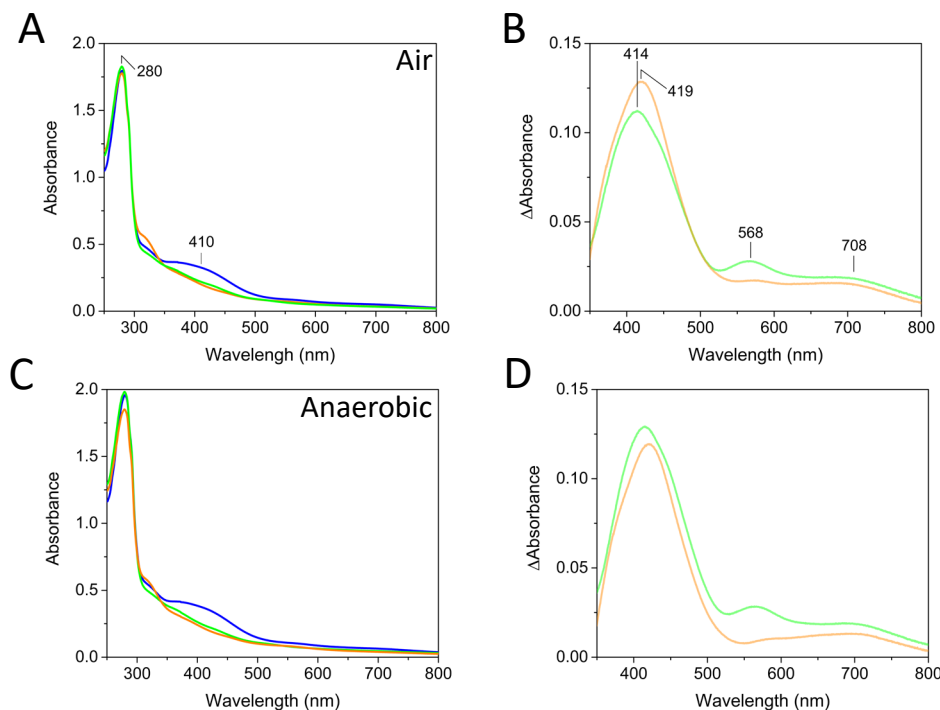
1
2 *Figure 2. Isolation of DvH-FDH2. (A) Streamlined expression and purification workflow. Single colonies*
3 *resulting from the transformation of *fdh2* plasmid into strain JW2127 were used to start a pre-culture that*
4 *served as the inoculum for a 10 L scaleup, cells from which were aerobically lysed and subjected to affinity*
5 *purification, yielding Strep-tagII FDH2. (B) SDS-PAGE of purified protein (lane 1) and molecular weight*
6 *markers (lane M). α and β represent the large and small subunits of FDH2, respectively. (C) Following non-*
7 *denaturing PAGE, FDH2 activity (dark single band) is detectable in air via NBT staining.*
8

9 Notably, there is a fundamental difference between prevailing strategies for metallo-FDH
10 isolation and what we have advanced. Our purification workflow (Figure 2) and
11 downstream handling steps (including storage) occur in air without involving nitrate,
12 azide, thiols, or formate at any stage of the process.

13 **Aerobic In-Gel Catalysis of Recombinant DvH-FDH2.** Literature precedents exist for
14 anaerobic activity staining of FDHs in native polyacrylamide gels using 2,3,5-
15 triphenyltetrazolium chloride^{56, 59} or phenazine methosulfate (PMS) / nitroblue
16 tetrazolium chloride (NBT).⁶⁰⁻⁶² However, this has not been achieved for any FDH in air.
17 Because O₂-insensitive group 5 [NiFe]-hydrogenases have been zymographically
18 visualized using redox dyes,⁶³ we asked whether a similar approach could work with DvH-

1 FDH2. When native polyacrylamide gel strips containing recombinant DvH-FDH2 were
2 incubated aerobically with NBT and formate, a single dark blue colored band appeared
3 within two minutes (Figure 2C). In the absence of formate, this band was not observed.
4 The same pattern was recapitulated in the spot assay where the blue color developed
5 within 15 s (Figure S2). These observations demonstrate that electrons released from
6 enzymatic aerobic formate oxidation are readily transferred to an artificial electron
7 acceptor with positive reduction potential ($E_{m,7} = +50 \text{ mV}^{24,63}$), resulting in the generation
8 of insoluble reduced NBT-formazan precipitates. Furthermore, our results establish that
9 both nitrate-assisted purification of FDH and/or reductive activation with high
10 concentration of thiols are not essential for maintaining redox activity under anaerobic or
11 atmospheric conditions.

12 **[4Fe-4S] Metalloclusters, Tungstopterin, and Selenocysteine Remain Unaffected by**
13 **O₂ During Catalytic Turnover.** As correctly pointed out by Hagen⁶⁴, the metal specificity
14 profiles of SRB FDHs remain incompletely described. Moreover, the nature of redox
15 centers in DvH-FDH2 is unknown.⁵⁶ Because DvH-FDH1 and DvH-FDH2 exhibit 61%
16 protein sequence identity (large subunit) and share all the metal coordination sites within
17 the two subunits (Figure S3), we surmised that a similar complement of redox centers
18 must exist in both systems. Since the DvH biomass was derived from a medium containing
19 Mo (1.24 μM) and W (0.15 μM), we predicted a metal ratio of 1Mo/W:16Fe:1Se. Consistent
20 with this, inductively coupled plasma mass spectrometry (ICP-MS) revealed that for every
21 mole of ¹⁸²W present, another 17 ± 1 moles of ⁵⁶Fe and 0.7 ± 0.1 moles of ⁷⁸Se were also
22 found (Table S3). Despite the nine-fold excess of molybdate (excluding contributions
23 from yeast extract) in the growth culture, we did not detect ⁹⁵Mo in our FDH2 samples.
24 These results underscore definitive tungsten selectivity of DvH-FDH2, distinguishing it
25 from Mo-specific^{14, 56} DvH-FDH3 and the promiscuous DvH-FDH1, which is capable of
26 incorporating both Mo and W.^{20, 56}



1
2 *Figure 3. Electronic spectra of DvH-FDH2. As-isolated (blue), formate-reduced (green), and dithionite-reduced*
3 *(orange) states are shown in panels A and C. Difference spectra are shown in panels B and D. As-isolated*
4 *minus formate-reduced and as-isolated minus dithionite-reduced are in green and orange, respectively.*
5

6 **Electronic and Electron Paramagnetic Resonance (EPR) Spectral Signatures of DvH-**

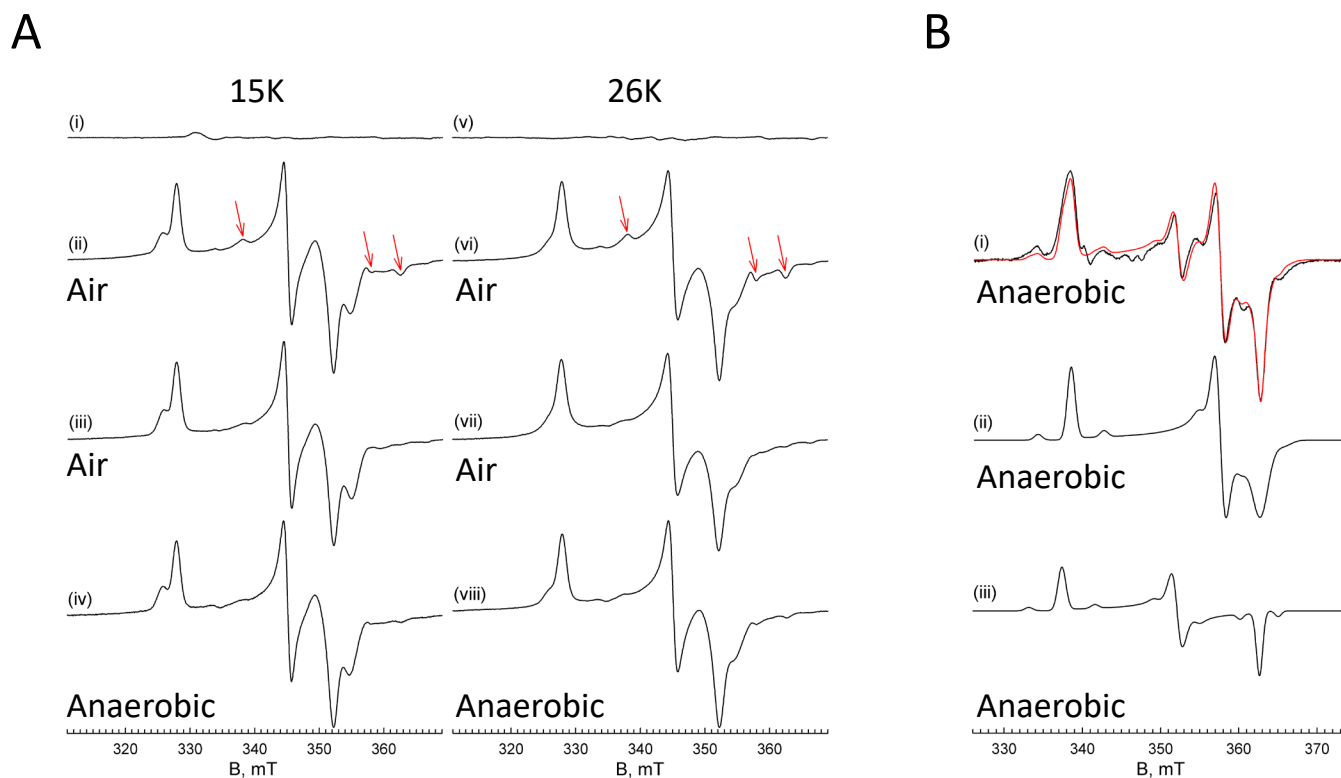
7 **FDH2 are Virtually Invariant in Air.**

8 The bulk of metallo-FDH electronic spectra in the
9 primary literature have been measured under anaerobic conditions to avoid inactivation
10 by molecular O₂.^{13, 16, 20, 24, 65} Although aerobic spectra exist for an O₂-tolerant Mo-Cys-
11 FDH stabilized by 10 mM nitrate,²⁸ their utility remains unclear, for the addition of formate
12 did not afford a characteristic spectral change. Here, we offer the first functional
13 validation of a W-Sec-FDH in air via electronic spectroscopy. Aerobically purified DvH-
14 FDH2 is brown in color and shows a broad S → Fe³⁺ charge transfer transition at 412 nm
15 (Figure 3A, blue trace), which is characteristic of [4Fe-4S]²⁺ clusters.⁶⁶ Addition of formate
16 leads to a substantial loss of this signal, indicating reduction to the [4Fe-4S]⁺ state (Figure
17 3A, green trace). Independently, reduction with dithionite yields a similar result (Figure
3A, orange trace). Employing anaerobic conditions makes no difference to the outcome

1 (Figure 3C). The virtually identical lineshape and amplitude of the difference spectra
2 (Figure 3B,D) illustrate that formate completely reduces (six reducing equivalents; four
3 $[4\text{Fe-4S}]^{2+}$ clusters and one W center) the majority of catalytically competent FDH2 in
4 solution. As dithionite would be expected to reduce both functional and non-functional
5 metal centers, we conclude that >94% of DvH-FDH2 is functionally fit. We have also
6 obtained the source DvH-FDH1 spectrum (Figure S4, pink trace, of Oliveira et al (2020))
7 and compared it with our as-isolated DvH-FDH2 counterpart acquired under anaerobic
8 conditions (Figure S4). The A_{400}/A_{280} ratio –an indicator of the extent of cluster loading⁶⁷–
9 estimated from these spectra are 0.18 (DvH-FDH2) and 0.17 (DvH-FDH1), affirming that
10 the two orthologs exhibit comparable protein purity and cofactor integrity.

11 To evaluate the predictions made via electronic spectroscopy, we pursued EPR
12 measurements. Figure 4 shows the EPR spectra seen with reduced DvH-FDH2 under a
13 variety of conditions, as well as the spectra for the as-isolated protein (panels (i), (v)). At
14 15K, we observed predominantly two distinct EPR signals (Figure 4A(ii)-(iv)), the relative
15 intensities of which are essentially independent of reductant (formate or dithionite) and
16 environment (anaerobic or air). The signals are consistent with the presence of reduced
17 iron sulfur clusters. At 26K, one of the signals is significantly broadened (Figure 4A(vi)-
18 (viii)) while by 40K both signals have disappeared (data not shown). This behavior is
19 consistent with fast relaxing $[4\text{Fe-4S}]$ clusters. There is also some indication of additional
20 signals (Figure 4A, red arrows), which are described in more detail below.

21 The simulated spectrum for the formate-reduced DvH-FDH2 prepared under
22 aerobic conditions and collected at 15K from Figure 4(iii) is shown in Figure S5, and the
23 simulation parameters given in Table S4. The g-values obtained are again consistent with
24 iron-sulfur clusters and the relative contribution of each cluster is approximately 1:0.75,
25 indicating almost a complete reduction of both centers. At higher microwave power, no
26 indication of additional signals was observed. The absence of additional signals in the
27 four-cluster FDH2 seen here is reminiscent of *D. gigas* W-FDH1 results,⁶⁸ where the two



1
2 *Figure 4: EPR spectra of DvH-Fdh2. (A). (i), (v) as isolated (45 μ M). (ii), (vi) dithionite-reduced (45 μ M). Red*
3 *arrows indicate approximate location of W^V g tensors. (iii), (vii) formate-reduced aerobic (45 μ M). (iv), (viii)*
4 *formate-reduced anaerobic (27 μ M). (i)-(iv) were collected with 10 Gauss modulation amplitude and 0.2*
5 *mW power at 15K. (v)-(viii) were collected with 10 Gauss modulation amplitude and 4 mW power at 26K.*
6 *(B) Simulation of the W center of dithionite-reduced sample. (i) W^V EPR spectrum (black trace) and*
7 *simulation (red trace) of 150 μ M Fdh2 collected with 8 Gauss modulation amplitude and 10 mW power at*
8 *108K. (ii), (iii) Scaled individual contributions to the simulation of the composite spectrum in (i). Simulation*
9 *includes hyperfine splittings originating from the 14.3% naturally occurring ^{183}W ($I = 1/2$). Simulation*
10 *parameters are presented in Table S4.*

11
12 observed signals represent pairs of Fe/S clusters with similar g-values. Alternatively, the
13 remaining two clusters present in DvH-FDH2 may have reduction potentials that are too
14 low to be reduced by formate or dithionite to any appreciable degree.⁶⁹

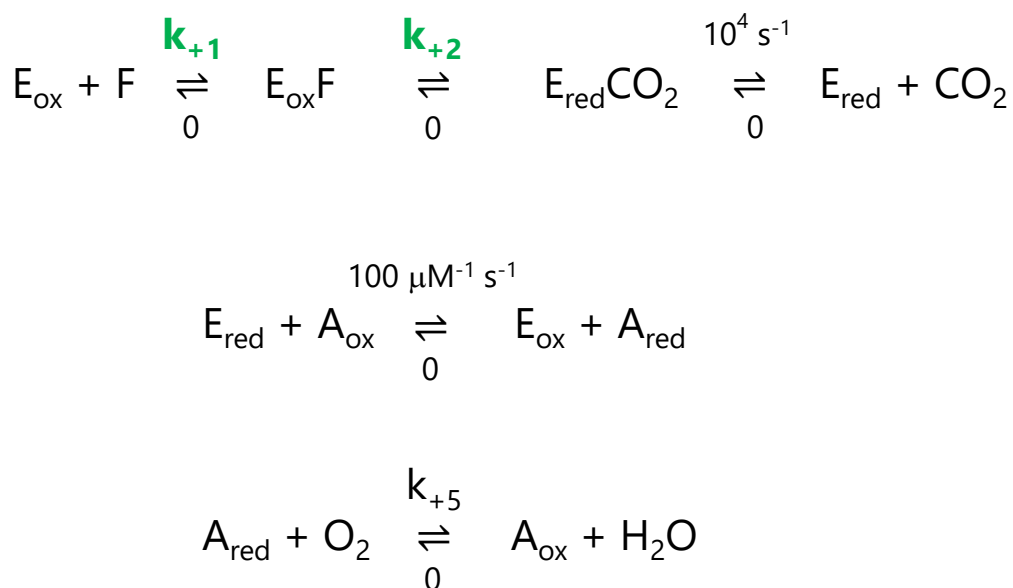
15 When 150 μ M enzyme is incubated with dithionite under anaerobic conditions for
16 an extended amount of time (\sim 12 hours or more) and the spectrum is collected at 108K,

1 an additional pair of signals are obtained (Figure 4B(i)); there is no evidence of the Fe/S
2 signals described in Figures 4A and S5 at this temperature. The new signals persist from
3 15K all the way to 108K without considerable line broadening, consistent with their arising
4 from slowly relaxing W(V) species. Their simulations are superimposed on the
5 experimental spectra. Figure 4B(ii) and (iii) present the component spectra scaled to their
6 contribution to the composite simulation in Figure 4B(i). The simulation parameters are
7 presented in Table S4 and include the well-resolved tungsten $I=1/2$ hyperfine splittings
8 originating from the 14.3% natural abundance ^{183}W isotope. The presence of the $I=1/2$
9 hyperfine splitting is further evidence that these signals arise from the tungsten center
10 rather than additional Fe/S clusters. The simulations indicate that the two species are in
11 an approximate ratio of 1:0.54 and the g -values ($g_{1-3} = 1.982, 1.876, 1.849$ and $1.904, 1.849,$
12 1.914 , respectively) are in good agreement with those seen from other W-containing
13 enzymes. Somewhat surprisingly, the large anisotropy of the W(V) g -values more closely
14 resembles the "low potential" signal for the *P. furiosus* aldehyde ferredoxin
15 oxidoreductase (AOR), which is a member of a different family of tungsten-containing
16 enzyme than the FDHs.⁷⁰ The presence of multiple W(V) signals in a single sample has
17 been seen with a number of W-containing enzymes and may be due to the presence of
18 inactive species in addition to the catalytically competent one, which is a rather common
19 feature of W-containing enzymes.⁶⁴

20 **Full Progress Curves Reveal High Catalytic Efficiency Under Atmospheric Conditions**
21 **and Lack of Enzyme Inactivation or Product Inhibition.** To the best of our knowledge,
22 solution enzyme kinetics investigations of metallo-FDHs have not directly probed formate
23 depletion or CO_2 production. Instead, low-potential artificial electron acceptors, most
24 commonly benzyl viologen (BV; $E_{m,7} = -360 \text{ mV}^{24}$) and methyl viologen (MV; $E_{m,7} = -446$
25 mV^{24}) for the forward and reverse reactions, respectively, have been routinely used as
26 surrogates to report on catalytic robustness. Although cautions have been raised against
27 trusting kinetic parameters derived from the use of these "inefficient and slow redox

1 *mediators*^{71,72}, they continue to be favored. Mo-Cys-FDHs offer an alternative by making
2 it possible to track NAD⁺ reduction or NADH oxidation.^{24, 28} Unfortunately, this strategy
3 can be extended only to select metallo-FDHs and it is prone to yield false-positive results
4 when interrogating aerobic CO₂ reduction with aerotolerant enzymes.²⁵ To further
5 complicate matters, FDHs from SRB are in a class of their own (Table S5). Unlike other
6 bacterial FDHs, these retain little to no activity after purification, requiring lengthy
7 reductive activation with high concentration of thiols and/or formate.^{17, 20} Finally, it is
8 impossible to assess the validity or robustness of the published results when experimental
9 data remains inaccessible – we are not aware of a report on metallo-FDH enzymology that
10 has disclosed a complete set of raw absorbance versus time data used to extract kinetics
11 parameters.

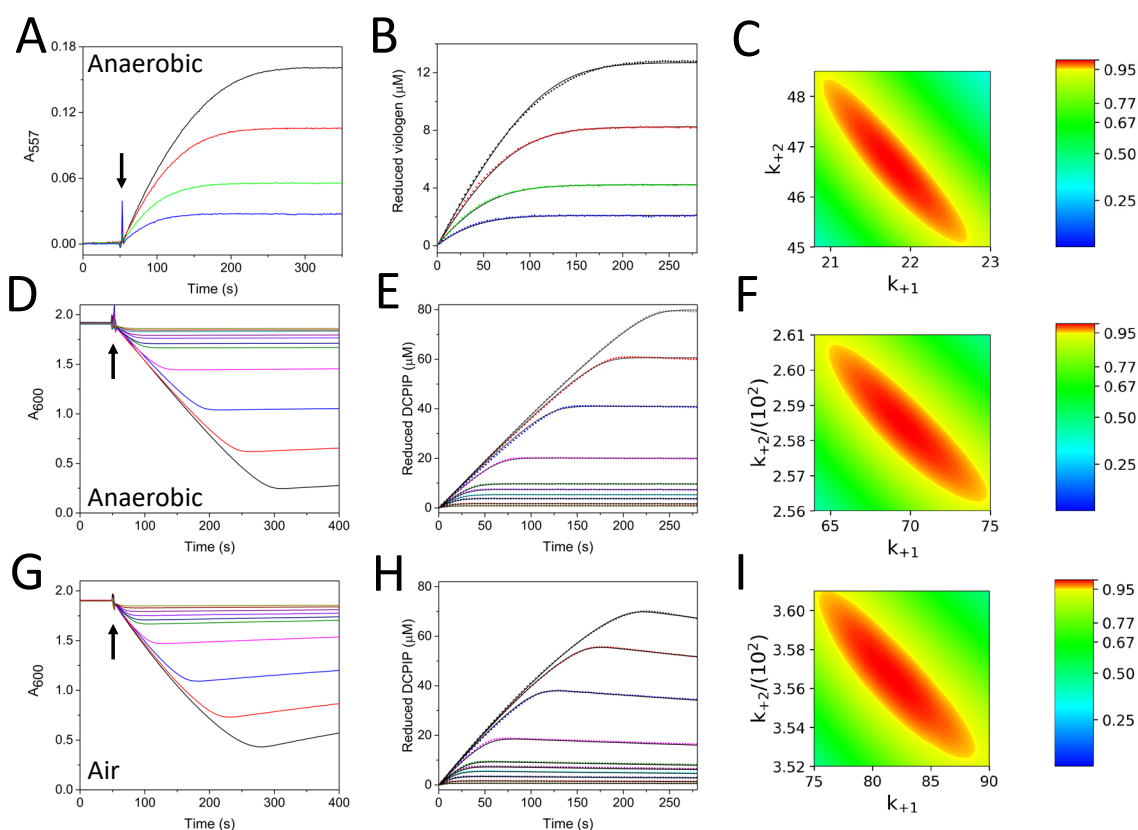
12 To resolve these uncertainties, we explored rigorous and reproducible solution
13 enzyme kinetics approaches capable of yielding results with functional information
14 content. First, we assessed catalytic efficiency with two chemically distinct artificial
15 electron acceptors, one each from the low- (BV) and high-potential (PES/DCPIP) $E_{m,7} =$
16 $+217$ mV²⁴) categories. Only the latter afforded the ability to acquire kinetics data both
17 in air and under argon. Second, we identified conditions under which full progress curves
18 could be measured. Such an approach is only possible for stable enzymes that catalyze a
19 single-substrate irreversible reaction in the absence of enzyme inactivation or product
20 inhibition (see below).⁷³⁻⁷⁶ And third, we have simultaneously analyzed several full
21 progress curves using dynamic simulation-based global fitting⁷⁵ to extract k_{cat} and
22 k_{cat}/K_m . This strategy overcomes the limitations of classical steady-state analysis, such as
23 the use of only the first few seconds of data, unreliable initial velocity values, and
24 overparameterization.⁷⁷ To benchmark our models (Schemes 1 and S1), we obtained
25 source BV enzyme kinetics data that formed the basis of Figure S3 and Figure 1C of Maia
26



1
 2 *Scheme 1. Minimal catalytic model used solely for estimating k_{cat}/K_m (k_{+1}) and k_{cat} (k_{+2}) from full progress*
 3 *curves via dynamic simulation-based global fitting. E_{ox} and E_{red} represent oxidized and reduced enzyme,*
 4 *respectively. A_{ox} and A_{red} denote oxidized and reduced forms of the electron acceptor, respectively. F , formate.*
 5 *The rate of product release (k_{+3}) is set to a high value, facilitating the reaction chemistry to define k_{cat} .*
 6 *Reaction of E_{red} with A_{ox} (k_{+4}) is assumed to be diffusion limited. Rationale for setting reverse rate constants*
 7 *to zero has been explained by Johnson.⁷⁷ The last equation is relevant only under aerobic conditions. Also, see*
 8 *Scheme S1.*

9
 10 *et al*¹⁷ (*Desulfovibrio desulfuricans* FDH3) and Oliveira *et al*²⁰ (DvH-FDH1), respectively.
 11 Global fitting of the steady-state progress curves from the former work allowed us to
 12 recapitulate the published values (Figure S6). Although Oliveira *et al*²⁰ only reported
 13 results from the initial velocity data, we were able to extract five full progress curves from
 14 their source data and perform *de novo* analysis (Figure S7A-C). In addition to finding k_{cat}
 15 values in the reported range, our method redefines the K_m of DvH-FDH1 to be 4.6 ± 0.3
 16 μM rather than $17 \mu\text{M}$ (Figure S7D-F). These observations illustrate that our catalytic
 17 models are poised to extract reliable kinetic parameters from DvH-FDH2 progress curves.

18 Because the original characterization of native DvH-FDH2 –by the same laboratory
 19 that has reported extensively on DvH-FDH1– was done using 2 mM BV (see Table S5),⁵⁶



1
 2 *Figure 5. Full progress curve probing of DvH-FDH2 catalysis. Raw experimental traces are shown in A (BV), D*
 3 *(PES/DCPIP), and G (PES/DCPIP) panels. Arrows represent the point at which the experiments were started by*
 4 *either the addition of formate (panel A) or FDH2 (panels D and G). Data normalized for electron acceptor*
 5 *concentration (B, E, and H panels) were globally fit (solid lines) using Kintek Explorer. Whereas panel B was fit*
 6 *to model shown in Scheme S1, panels E and H were fit to the counterpart in Scheme 1. Confidence contour*
 7 *analysis (panels C, F, and I) illustrates that both k_{+1} (k_{cat}/K_m) and k_{+2} (k_{cat}) are well constrained by the kinetic*
 8 *data. Upper and lower bounds of each rate constant is reflected by the axes labels. When plotted as a function*
 9 *of one another, red ovals signify the extent of variability in k_{+1} and k_{+2} while still being constrained by the*
 10 *model. Consequently, both parameters display a defined boundary (red), χ^2 of which is 0.95 (side bar). Table*
 11 *1 lists rate constants, as well as best fit parameters derived from this analysis.*

12
 13 we attempted to reproduce the published results with aerobically purified recombinant
 14 DvH-FDH2. However, our enzyme was added to the reaction mix without any pre-
 15 activation with thiols or formate. Although DvH-FDH2 displays redox activity in air (Figure
 16 2C), abiotic reaction of reduced BV⁺ with O₂ made us employ strict anaerobic conditions.
 17 Since our measurements were not made inside an anaerobic chamber, we ensured anoxic
 18 conditions by adding 1 unit/mL of glucose oxidase (GO). Catalase was also

Table 1. Parameters gleaned from full progress curve analysis[#]

Electron acceptor	k_{cat} (s^{-1})	K_m (μM) [‡]	k_{cat}/K_m ($\mu\text{M}^{-1} \text{s}^{-1}$)	Product stoichiometry [§]	Enzyme inactivation	Product inhibition
2 mM BV (anaerobic)	47 ± 2	2.1 ± 0.1	22 ± 1.1	2BV ⁺ :1F	No	No
DCPIP (anaerobic)	258 ± 3	3.7 ± 0.3	69 ± 6	1DCPIP:1F	No	No
DCPIP (air)	354 ± 5	4.5 ± 0.4	79 ± 6	1DCPIP:1F	No	No

[#]Errors for k_{cat} and k_{cat}/K_m via confidence contour analysis implemented in FitSpace Explorer

[‡]Standard error values for K_m (calculated from k_{cat} and k_{cat}/K_m) estimated according to Johnson (2019)

[§]F, formate; BV⁺, reduced benzylviologen

1
2 included to eliminate H₂O₂ that may arise from GO activity. Global fitting of full progress
3 curves (1,2,4, and 6 μM formate) yielded values comparable to those reported by da Silva
4 *et al*⁵⁶ (Figure 5A-C, Table 1). Furthermore, our progress curves revealed that two
5 molecules of BV⁺ are generated for every formate molecule oxidized by DvH-FDH2.
6 Despite elimination of the reductive activation step, the enzymatic parameters derived
7 from our standard steady-state kinetics analysis were virtually identical to the published
8 values (Table S6), suggesting that the pre-activation step introduced by da Silva *et al*
9 (2011) had no effect on the outcome. In our hands, DvH-FDH2 exhibits catalytic
10 parameters that are roughly an order of magnitude smaller than their DvH-FDH1
11 counterparts (Table S6 and Figure S7F). A closer inspection of product stoichiometry at
12 high formate concentrations suggested that BV concentration could be limiting (Figure
13 S8). Therefore, we repeated the experiments at 20 mM BV. This restored 2BV⁺:1F

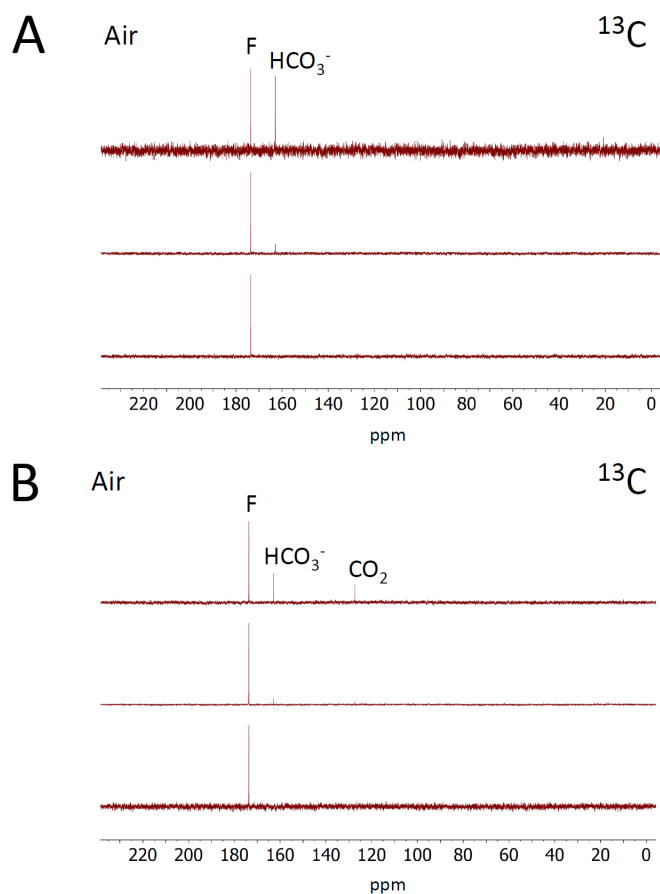
1 stoichiometry across the board but catalytic parameters did not change appreciably
2 (Figure S9, Table S6). We have also confirmed that addition of GO and catalase do not
3 interfere with the results of activity measurements (Figure S10).

4 Collectively, the observations above suggest that BV is not a good electron
5 acceptor for DvH-FDH2. To test this hypothesis, we independently pursued activity assays
6 with PES/DCPIP. Consistent with literature precedents on dehydrogenases,⁷⁸ phenazine
7 ethosulfate (PES; $E_{m,7} = +65 \text{ mV}^{79}$) was required for transferring electrons from FDH to
8 DCPIP (Figure S11A). By varying the concentrations of DCPIP and PES, we were able to
9 identify optimal conditions that would support activity measurements both in air (Figure
10 S11B-F and Figure S12) argon (Figure S13). At DCPIP concentrations below $100 \mu\text{M}$, global
11 fitting of anaerobic full progress curves (Scheme 1 and Figure 5D-F) resulted in roughly
12 five-fold higher turnover number (TN) than what we obtained with 2 mM BV (Table 1). K_m
13 values did not show a significant difference, however. The same pattern was reproduced
14 when the measurements were made in air (Figure 5G-I). A key difference between the
15 two conditions is that the slow reoxidation ($k_{+5} = 3 \pm 0.5 \times 10^{-6} \mu\text{M}^{-1} \text{ s}^{-1}$; see last equation
16 in Scheme 1) of reduced DCPIP by O_2 caused the post reaction region to slope slightly
17 upward (Figure 5G, Figure S11D, and Figure S12B). However, this should not be confused
18 with alterations to the shape of progress curves stemming from product inhibition,⁷⁵
19 which we did not observe when DCPIP (Figure 5G) or BV (Figure 5A) served as electron
20 acceptors. In fact, a product stoichiometry of one reduced DCPIP for every formate
21 oxidized was reproducibly found in our measurements (Table 1). Since addition of fresh
22 substrate at the end of a progress curve cleanly reproduced the original trace (Figure
23 S12F), we can further ascertain that the enzyme was stable and fully active during catalysis
24 in air. Taken together, PES/DCPIP-dependent catalytic parameters obtained from global
25 fits are in excellent agreement with those from our initial velocity calculations (Table S6).
26 And the TN with PES/DCPIP remains virtually the same under anaerobic and aerobic

1 conditions. Notably, the catalytic efficiency of DvH-FDH2 in air is in the range of 7×10^7
2 $M^{-1} s^{-1}$ (Table 1), which is comparable to that reported for DvH-FDH1²⁰ ($\sim 8 \times 10^7 M^{-1} s^{-1}$)
3 when BV serves as the electron acceptor under anaerobic conditions. Moreover, our
4 PES/DCPIP-based TN and k_{cat}/K_m for formate oxidation are roughly an order of magnitude
5 and 500-fold higher, respectively, than what has been reported for the aerotolerant Mo-
6 Cys-FDH from *Rhodobacter capsulatus* using the natural (NAD^+) electron acceptor.²⁸
7 Finally, we have found that the inhibition profiles of DvH-FDH2 in the presence of azide
8 or nitrate are not significantly impacted by O_2 (Figures S14 and S15). Whereas azide
9 blocks the enzyme with an IC_{50} of about 0.8 mM, nitrate is far less effective.

10 **Catalytic Redundancy or Gain of a New Enzyme Function? Exploiting the Peck-Gest** 11 **Paradigm to Seek Insights into How FDHs May Have Evolved to Achieve Aerobic**

12 **Catalysis.** Our full progress curve analysis establishes that both k_{cat} and k_{cat}/K_m are
13 severely underestimated when BV is used as the electron acceptor. It also reveals a
14 preference for the latter viz., whereas DvH-FDH2 favors high-potential acceptors, such as
15 PES/DCPIP or NBT, DvH-FDH1 is highly active with BV²⁰. Although DCPIP data is not
16 available for the latter enzyme, a close ortholog (Dg-FDH1) only shows 10% of BV activity
17 with the high-potential acceptor.¹⁸ Such linkages take on special significance when
18 multiple FDHs encoded by the same organism are compared. In 1957, Peck and Gest⁸⁰
19 discovered two types of FDH in *Escherichia coli* solely based on their preference for
20 artificial electron acceptors – one was linked to PMS/DCPIP and its expression was
21 confined to O_2 /nitrate-respiring cells while the other was BV-linked and unique to non-
22 respiring cells (reviewed by Stewart³⁹). It is now clear that Fdh-N is DCPIP-linked,⁶¹ and
23 Fdh-H is BV-linked. The third poorly characterized variant of *E. coli*, Fdh-O, is also DCPIP-
24 linked.⁴² Extending the Peck-Gest paradigm to DvH –only the second microbe for which
25 all three FDHs have now been characterized– we would predict that the BV-linked FDH1

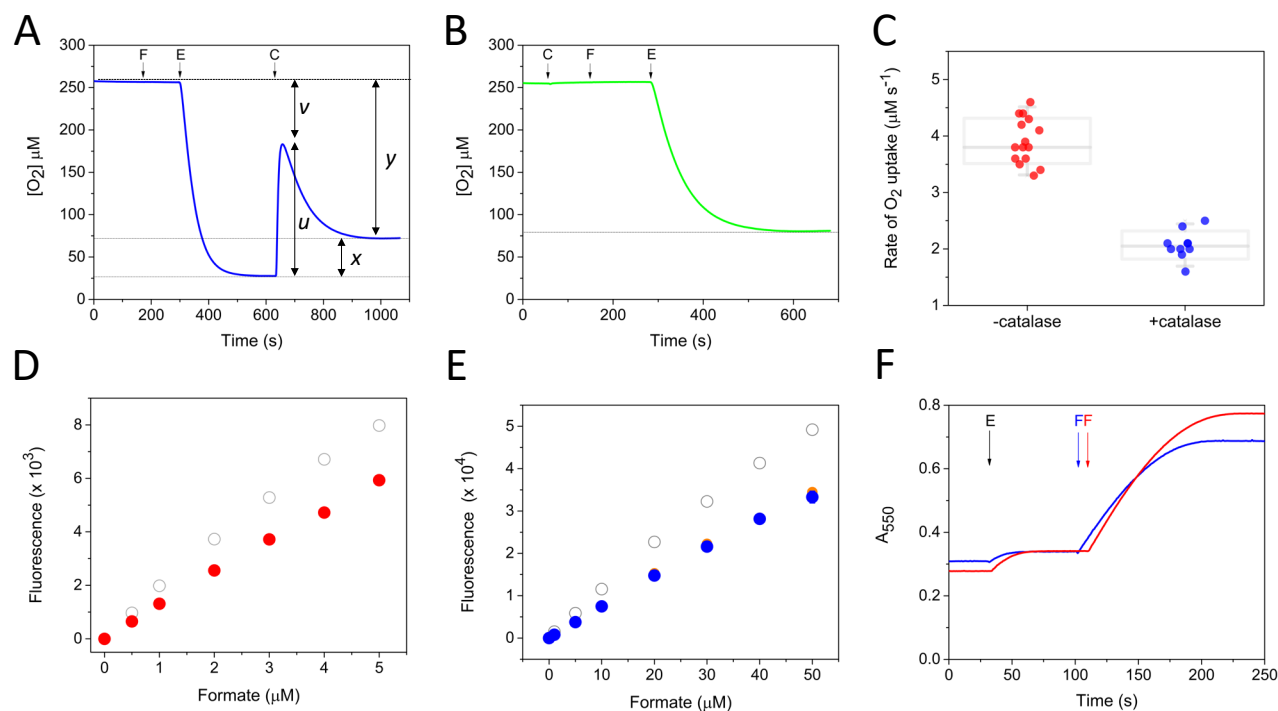


1
2 *Figure 6. Product generated by DvH-FDH2 catalyzed reaction. ^{13}C NMR spectra at pH 7.5 (A) and pH 6 (B).*
3 *^{13}C -formate (bottom), ^{13}C -formate + enzyme (middle), ^{13}C -formate + enzyme + PES (top).*
4

5 is involved in anaerobic respiration and that the PES/DCPIP-linked FDH2 plays a role in
6 aerobic respiration. It has already been established that FDH1 is essential for anaerobic
7 sulfate respiration when formate serves as the electron donor.⁵⁰ Biological function of
8 FDH2 remains to be elucidated. Our study proves that catalytic parameters derived from
9 viologen-based measurements lack functional information content to make predictions
10 about how well a given FDH would perform under aerobic conditions. Instead, high
11 catalytic performance on BV only guarantees activity under anaerobic conditions.
12 Confirmation bias has boosted reliance on viologen-based kinetics and stymied efforts to
13 uncover O_2 -immune FDHs that can reversibly function in air. This is best exemplified by
14 DvH-FDH2, which exhibits the lowest TN with BV (Table S5) and yet is the most O_2 -

1 insensitive of all metallo-FDHs characterized to date from any bacterium. Therefore,
2 biological context must factor critically into future search efforts aimed at discovering air
3 insensitive FDHs.

4 **CO₂ is the Product of Aerobic Formate Oxidation by DvH-FDH2.** Although several
5 metallo-FDHs have been investigated, there is just one report in the literature describing
6 the product resulting from enzymatic oxidation of formate under anaerobic conditions.²³
7 In all remaining works, product formation is implied based on the reduction of a natural
8 (NAD⁺) or artificial electron acceptor, which is often BV. Although we have used two
9 different artificial electron acceptors in this study, we made sure to leave no stone
10 unturned where product analysis in air is concerned. At pH 7.5, combining DvH-FDH2
11 with isotopically labeled ¹³C-formate readily yields a discernible H¹³CO₃⁻ resonance at
12 162.93 ppm (Figure 6A, middle and Figure S16) via ¹³C NMR spectroscopy. Addition of
13 PES to the mix substantially enhances the resonance intensity, indicating that the number
14 of turnovers has increased in the presence of an artificial electron acceptor (Figure 6A,
15 top; Figure S17). This hints at the likelihood that O₂ may not be a good electron acceptor
16 for DvH-FDH2 despite its abundance under our experimental conditions (257 μM at 25
17 °C). Similarly, at pH 6, both H¹³CO₃⁻ (162.88 ppm) and ¹³CO₂ (127.29 ppm) resonances
18 appear, and their intensities amplify when PES is included (Figure 6B, middle and top,
19 respectively; Figures S18 and S19). As a positive control, we have confirmed that
20 NaH¹³CO₃, in isolation at pH 6, generates H¹³CO₃⁻ and ¹³CO₂ resonances at positions
21 identical to those found with enzymatic formate oxidation (Figure S20). These chemical

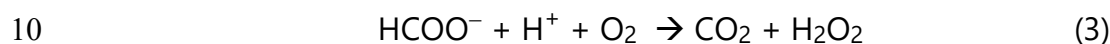


1
 2 *Figure 7. Mechanistic basis of DvH-FDH2 O₂ insensitivity. (A) Oximetry reveals coupling of formate oxidation*
 3 *to O₂ reduction. (B) Oximetry in the presence of catalase. (C) Comparison of O₂ uptake rates in the absence*
 4 *(n = 15) and presence (n = 10) of catalase. Both dot and box plots are shown with the latter deemphasized*
 5 *due to medium sample size.⁸¹ (D) Enzymatic H₂O₂ generation monitored via Amplex Red assay (filled red*
 6 *circles) (n=3). (E) Enzymatic H₂O₂ production evaluated using the CBA assay (filled blue circles) (n=3). Filled*
 7 *orange circles represent data obtained at 10 nM FDH2. In both D and E panels, open grey circles represent*
 8 *H₂O₂ standard curve determined in the absence of formate or FDH2. (F) Reduction of equine cytochrome c*
 9 *under aerobic (blue) and anaerobic (red) conditions (n = 3). Points of addition of formate (F), enzyme (E =*
 10 *FDH2), and catalase (C), are identified by down arrows. u, v, x, and y are defined in the text.*
 11

12 shifts agree well with those reported in the literature.⁸² ¹³C-formate was independently
 13 validated by both ¹H and ¹³C NMR spectra. Whereas the latter generates a single
 14 resonance at 173.65 ppm (Figures 6A and 6B, bottom; Figures S21 and S22), *J*-coupling
 15 (~ 195 Hz) between the ¹H and ¹³C atoms splits ¹H spectrum of ¹³C-formate into two
 16 resonances (8.66 ppm and 8.27 ppm) (Figure S23). Despite the poor sensitivity of ¹³C

1 NMR spectroscopy⁸³ and not optimizing data collection for $5 \times T_1$ (relaxation time), we
2 succeeded in demonstrating that CO₂ is the true product of aerobic catalysis.

3 **FOX Activity Generates H₂O₂, Enabling Oxygen Insensitivity of DvH-FDH2.** To
4 understand how DvH-FDH2 deals with O₂, we used a Clark-type O₂ electrode to ask
5 whether formate oxidation under atmospheric conditions is coupled to O₂ reduction.
6 Addition of enzyme to formate-containing aerobic buffer led to robust O₂ consumption
7 (Figure 7A). Once the signal plateaued, catalase was added. This led to O₂ evolution
8 followed by O₂ uptake until a plateau was reached, suggesting the production of H₂O₂
9 via 2e⁻ reduction of O₂ (reaction 3).



11 We attempted a calculation of the electron flux that led to H₂O₂ formation.^{84, 85} The x/y
12 value in Figure 7A would imply that roughly one quarter of the electrons from formate
13 were ending up in H₂O₂. However, this is likely to be an underestimate because formate
14 was in large excess and was continuing to be oxidized post dismutation of H₂O₂ by
15 catalase, manifesting as the second O₂ uptake step. Instead, if we considered the u/v
16 value, ~ 80% of the electron flux goes towards peroxide generation. To resolve this
17 uncertainty, we pursued the kinetics approach developed by Lu and colleagues.⁸⁶ By
18 comparing the initial velocities of O₂ uptake in the absence (Figure 7A) and presence
19 (Figure 7B) of catalase, we found that it was 50% lower in the latter (Figure 7C). And the
20 rates did not vary significantly between pH 6.0 and 8. This outcome suggested that H₂O₂
21 was the major product of O₂ reduction during aerobic formate oxidation. Appropriate
22 controls were built into our experimental design for evaluating alternate endpoints. H₂O₂
23 addition in the absence of exogenous catalase showed that DvH-FDH2 lacks catalase
24 activity (Figure S24A). To rule out the possibility of abiotic O₂ consumption, DvH-FDH2

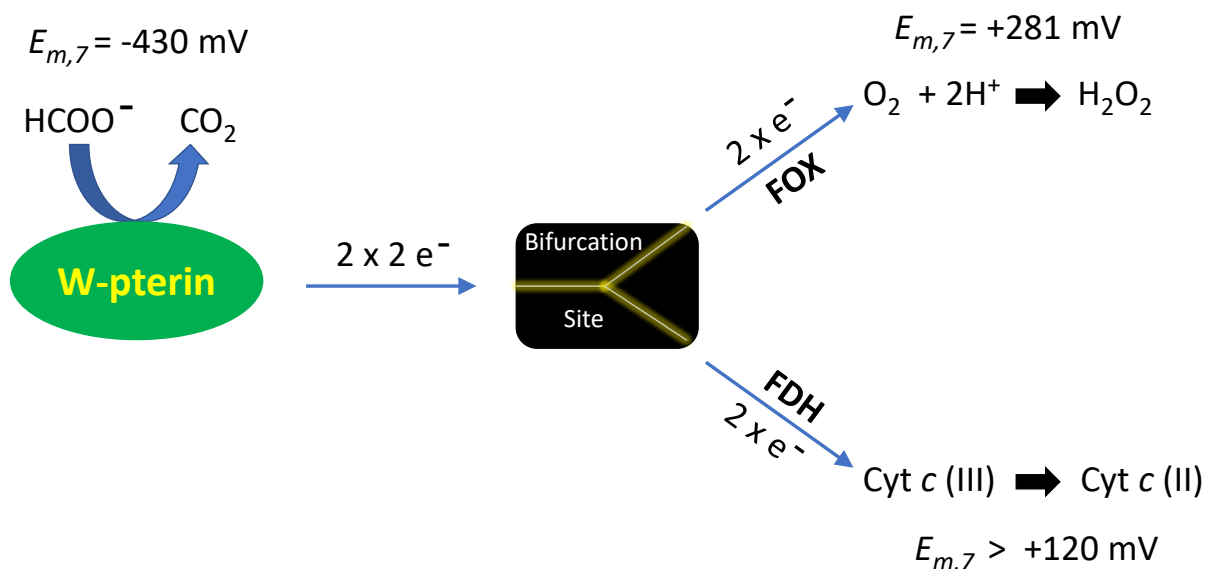
1 was heat denatured and subjected to oximetry. Neither O₂ uptake nor H₂O₂ generation
2 was found (Figure S24B). Moreover, inclusion of 1 mM EDTA minimized artifacts arising
3 from transition metal contaminants. Even with these controls in place, we could not
4 eliminate the possibility that atmospheric O₂ was completely excluded during oximetry.
5 Therefore, we pursued two orthogonal approaches to directly quantify H₂O₂. In the first
6 method, horseradish peroxidase (HRP) catalyzed formation of fluorescent resorufin from
7 H₂O₂ and amplex red was monitored. We observed that for every mole of formate
8 oxidized, roughly 0.75 mole of H₂O₂ was produced during aerobic DvH-FDH2 catalysis
9 (Figure 7D). Inclusion of catalase abolished the fluorescence signal and denatured
10 enzyme failed to yield H₂O₂ (Figure S25). However, the inability of amplex red assay to
11 quantify H₂O₂ beyond 5 μM made it impossible for us to investigate the consequences of
12 O₂ reduction at formate concentrations approaching 10 – 20K_m. Furthermore, despite
13 being considered the gold standard, this assay is prone to artifacts.^{87, 88} For example,
14 interferences stem from interaction between redox enzymes and resorufin⁸⁹. To
15 overcome these limitations, we resorted to a method independent of HRP and amplex
16 red. Here, we followed the direct reaction of non-fluorescent coumarin-7-boronic acid
17 (CBA) with H₂O₂, leading to the production of fluorescent 7-hydroxycoumarin (COH).⁹⁰
18 Although this reaction is slow⁹⁰ (k_{on} 1.5 M⁻¹ s⁻¹), the assay is linear over a much broader
19 range of H₂O₂. Therefore, we quantified H₂O₂ production during aerobic DvH-FDH2
20 catalysis, varying formate concentration between 1 - 10K_m. It amounted to 64 ± 6% and
21 did not show significant variation when higher enzyme concentrations were used.
22 Catalase addition eliminated the signal completely, confirming that H₂O₂ is indeed the
23 major product of O₂ reduction by DvH-FDH2 (Figure S26).

24 Next, we tried to assess superoxide (O₂^{•-}) generation⁹¹ by DvH-FDH2. Because
25 addition of SOD had negligible effect on both direct quantification (Figures S25 and S26)

1 and oximetry (Figure S27), we probed the reduction of partially acetylated cytochrome *c*.
2 The advantage of using the latter is that it is still reducible by $O_2^{\bullet-}$ but not susceptible to
3 interferences arising from oxidase or reductase activities when unmodified cytochrome *c*
4 serves as the substrate.⁹² Significant reduction was not observed, implying that $O_2^{\bullet-}$ was
5 not released by DvH-FDH2 (Figure S28).

6 Taken together, our results establish FOX activity of a highly pure metallo-FDH. To
7 the best of our knowledge, this has never been demonstrated before. Based on IUPAC-
8 IUB nomenclature, the term “*oxidase*” is reserved for enzymes, which utilize O_2 as the
9 electron acceptor. In our case, formate oxidation is coupled to $2e^-$ reduction of O_2 by
10 DvH-FDH2, resulting in 65 to 75% H_2O_2 production (reaction 3). Hence, we project that
11 roughly 25 – 35% O_2 is reduced to H_2O by a $4e^-$ process (reaction 2). From a mechanistic
12 perspective, this is reminiscent of how O_2 -tolerance is achieved in some [NiFe]-
13 hydrogenases (see below)⁹³. However, given the high level of difficulty associated with
14 detecting and quantifying H_2O , only a handful of studies have been performed on this
15 front using redox enzymes.^{86, 89, 94, 95}

16 **Redox Bifurcation Facilitates the Coexistence of FOX and FDH Activities:** When DvH-
17 FDH2 couples formate oxidation to the reduction of electron acceptors other than O_2 , it
18 functions as a dehydrogenase. Our kinetic (Figure 5) and product (Figure 6) analysis
19 illustrate that FOX activity does not interfere with the latter function. Nevertheless, it is
20 unknown if this pattern holds when a macromolecule serves as an electron acceptor.
21 Because our central hypothesis rests on the assumption that FOX activity is likely to
22 preserve electron transfer to natural high potential acceptors (see Introduction), we tested
23 it directly. Since an eleven-heme cytochrome *c* (*uhc*) is located immediately adjacent to
24 *fdh2* in DvH⁵⁷, we asked whether DvH-FDH2 reduces native equine cytochrome *c* ($E_{m,7} =$
25 $+260$ mV) in the presence of formate. Whereas stoichiometric reduction occurred under

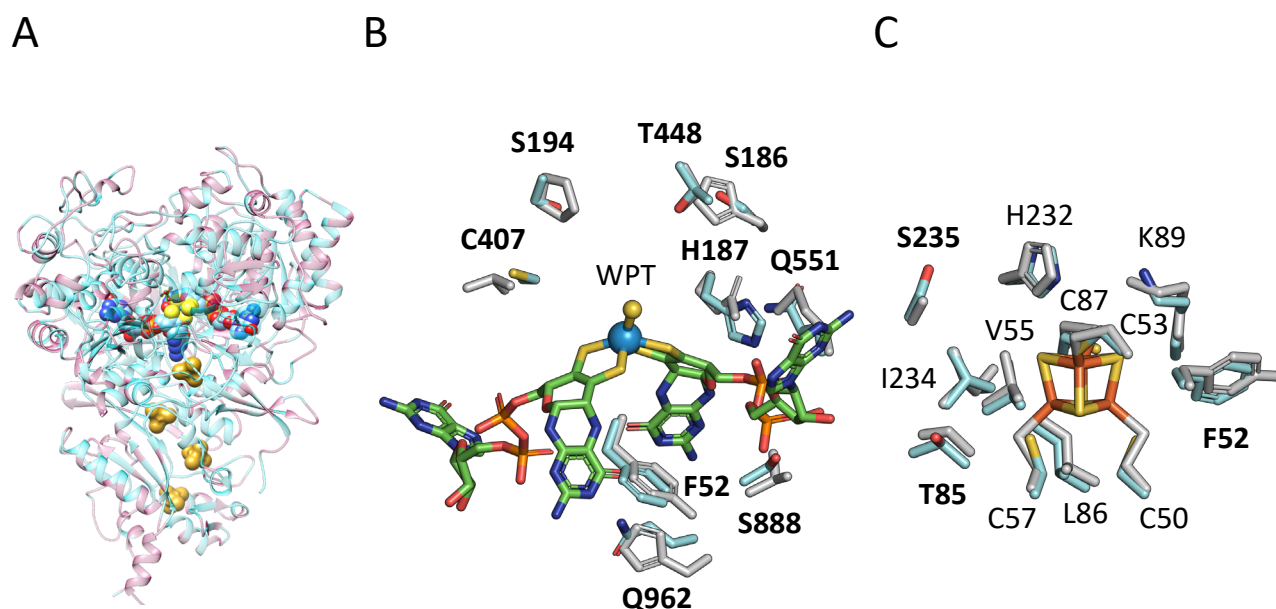


1
2 *Figure 8. Working model of redox bifurcation by DvH-FDH2. W-pterin denotes the tungstopterin active site.*
3 *The nature of bifurcation site (black box) is unknown. Since a multiheme cytochrome c (Cyt c) is likely the*
4 *natural redox partner of DvH-FDH2, it is shown at the end of the bottom branch. However, its reduction*
5 *potential is a predicted value based on our results with equine Cyt c ($E_{m,7} = +260 \text{ mV}$).*
6
7 anaerobic conditions, only ~80% underwent reduction in air (Figure 7F). However, the
8 initial rates remained invariant (Figure S29). Doubling the formate concentration resulted
9 in near stoichiometric reduction in air (Figure S30). Moreover, inclusion of SOD or catalase
10 did not result in a noticeable difference (Figure S31). Under the conditions employed,
11 dissolved O_2 (257 μM) is at a much higher concentration than cytochrome c. Yet, electrons
12 are readily delivered to the latter. These results prove for the first time that oxidase and
13 dehydrogenase activities coexist within the DvH-FDH2 scaffold. We advance a non-
14 energy-conserving electron bifurcation (EB)⁹⁶ mechanism to rationalize how this is
15 accomplished (Figure 8). Here, electrons resulting from the oxidation of a $2e^-$ donor
16 (formate) traverse two independent thermodynamically favorable electron transfer paths

1 (FOX and FDH) to reduce two different electron acceptors (O_2 and cytochrome c).
2 Although we do not know the identity of the cofactor engaged in EB, both the
3 tungstopterin and the proximal [4Fe-4S] cluster of the large subunit are strong candidates.
4 Apropos, arsenite oxidase is thought to achieve EB by cycling between the Mo(VI)-dioxo
5 and Mo(IV)-oxo states.⁹⁷ Whether tungsten can assume this role in DvH-FDH2 remains
6 to be investigated. Based on our observation that superoxide was not generated during
7 aerobic catalysis, we hypothesize that the bifurcating site likely favors normally ordered
8 over inverted potentials.^{98, 99} Even though the interconversion of formate to CO_2 is
9 reversible, the bifurcation events in either direction (especially O_2 reduction) are
10 potentially irreversible.¹⁰⁰ Finally, EB is thought to be the dominant source of reactive
11 oxygen species in biological systems.¹⁰⁰ We have shown that redox metalloenzymes can
12 capitalize on it to achieve O_2 insensitivity.

13 **Molecular Basis of DvH-FDH2 O_2 Insensitivity.** In their phylogenetic analysis, Oliveira
14 et al²⁰ began with over 6000 FDH sequences and reduced it by an order of magnitude in
15 an effort to understand how the variability impacts catalytic mechanism and O_2 stability.
16 Here, we have narrowed the sequence space to just two closely related paralogs – one of
17 these (DvH-FDH1) is unable to achieve catalysis in the presence of O_2 while the other
18 (DvH-FDH2) thrives in air. To gain atomic insights, we constructed a highly accurate
19 heterodimeric structure of the latter using AlphaFold2.¹⁰¹ (Figure 9A). We did not utilize
20 preexisting structure templates to model the structure. The resulting atomic coordinates
21 (sans cofactors) include confidence metrics (predicted Local Distance Difference Test,
22 pLDDT) at the single residue level wherein higher scores on a scale of 1 – 100 represent
23 greater confidence. Figure S32A shows that the heterodimeric structure of DvH-FDH2 is
24 modeled with high confidence; bulk of the polypeptide chain displays pLDDT scores >90.
25 Similarly, the predicted aligned error (PAE; color saturation found at any x,y coordinate in
26 Figure S32B) is a metric of how well a residue is positioned and oriented.

1



2

3 *Figure 9. Molecular insights into the O₂ insensitivity of DvH-FDH2. (A) AlphaFold2-based tertiary topology*
4 *and quaternary structure. Shading in pink represents sites that are unique to FDH2. The four [4Fe-4S] clusters*
5 *are depicted by a combination of yellow and orange spheres. The two pterin moieties coordinating a W (WPT)*
6 *are shown at the top using blue and red spheres. (B) Variability within a 10 Å radius of W. Side chains*
7 *belonging to DvH-FDH1 are shown in grey. W and sulfide are identified as blue and yellow spheres,*
8 *respectively. For clarity, conserved sites are not shown (see Figure S36). (C) Environment of the large subunit*
9 *[4Fe-4S] cluster (yellow-orange sticks). In both panels B and C bolded labels signify variations.*

10

11 Here too, very low PAE values are seen for both subunits. Independently, we predicted
12 the structures of the two subunits and compared them with the heterodimeric
13 counterpart. Both approaches yielded very similar results. DALI¹⁰² confirms that the
14 tertiary folds of large and small subunits are superimposable on their DvH-FDH1
15 counterparts with a root-mean-square-deviation (RMSD) of 1.4 Å (953 Ca atoms; Z-score
16 58.2; 61% identity) and 1.0 Å (213 C α atoms; Z-score 35.9; 63% identity), respectively.
17 Backbone RMSD variations are shown in Figures S33 and S34). Similarly, the RMSD

1 between DvH-FDH1 and DvH-FDH2 heterodimers is 1.37 Å (952 C α atoms). After
2 completing these validations, we computed a residue-residue (RR) distance map to
3 identify the regions of major variation between FDH2 and FDH1 (Figure S35).

4 Although the active site residues are largely conserved between FDH2 and FDH1,
5 there are several differences in the vicinity of the tungsten center (compare Figure 9B and
6 S36B). Specifically, two highly conserved residues²⁰ have been substituted in FDH2. The
7 introduction of S186 and H187 are noteworthy because they replace H187 and Q188,
8 respectively, in FDH1. These changes are likely to influence Sec reactivity. Nonetheless,
9 the relevance of Sec to O₂ insensitivity is unclear. Whereas it would be expected to confer
10 O₂ tolerance,^{103, 104} both FDH1 and FDH2 incorporate it and only the latter is capable of
11 aerobic catalysis. Consequently, additional investigations are required to ascertain how
12 Sec contributes differentially to stability and catalysis in enzymatic systems.

13 [4Fe-4S] clusters are prone to oxidative damage^{105, 106} and enzymes harboring
14 them would be expected to be inactivated by H₂O₂ generated during aerobic catalysis.¹⁰⁷
15 However, this does not happen with DvH-FDH2. As a corollary, cellular experiments with
16 *Campylobacter* group of bacteria have shown that they harbor an FDH capable of
17 producing H₂O₂.¹⁰⁸⁻¹¹⁰ In these organisms, H₂O₂ functions as a terminal electron acceptor
18 in respiration.^{84, 85, 111} *E. coli* capitalizes on H₂O₂ in a similar fashion.¹¹² It is unknown
19 whether DvH exploits this strategy. Therefore, we examined the environment of the active
20 site proximal [4Fe-4S] cluster to glean insights. Strikingly, there are three substitutions
21 (Y52F, S85T, A236 → S235) within a 5 Å radius (Figure 9C). This is relevant because in
22 some O₂-tolerant [Ni-Fe]-hydrogenases, introduction of a polar residue near catalytic
23 metal clusters has been shown to improve O₂ resistance. Because the predicted structure
24 of DvH-FDH2 requires us to dock the cofactors, we are unable to say more about their
25 conformation or reactivities. Nevertheless, one or more of the following precedents may
26 inform how O₂ insensitivity and / or EB evolved in our system: (1) group 1d O₂-tolerant

1 [Ni-Fe]-hydrogenases, which also reduce O₂ to H₂O₂ and H₂O^{89, 113}, generate an unusual
2 [4Fe-3S] cluster; (2) a canonical [4Fe-4S] cluster in another [NiFe] hydrogenase undergoes
3 redox-dependent structural changes,¹¹⁴ poisoning it in a protected state until the next
4 catalytic cycle; (3) second coordination sphere effects; and (4) local or remote
5 conformational fluctuations at the protein level that could offer protection from attack by
6 oxidants. Ongoing structural and biochemical studies are likely to clarify the nature of
7 innovations at work in DvH-FDH2.

8 **Impact** The findings reported here have broad utility in disparate fields of research. (1)
9 The current state of the art in formate/air biofuel cells is limited to mediated electron
10 transfer because O₂-sensitive metallo-FDHs need protection from redox polymer films to
11 operate.¹¹⁵ Consequently, a “true” formate/air BFC is yet to be fabricated. DvH-FDH2
12 should be able to work in the absence of protective matrices and power BFCs via direct
13 electron transfer. (2) There is no known biocatalyst that accomplishes CO₂ reduction in
14 air.¹¹⁶ And it is not possible to ground truth this reaction in a chemical environment
15 because low-potential artificial electron donors (e.g., methyl viologen) are futile under
16 atmospheric conditions.¹¹⁷ Since the tungstopterin active site and other redox centers of
17 DvH-FDH2 are O₂ insensitive, there is a high probability of exploiting bioelectrocatalysis
18 to run the reverse reaction aerobically. In so far as the O₂-tolerant [NiFe] hydrogenase
19 has been successfully shown to perform aerobic bioelectrocatalysis¹¹⁸⁻¹²⁰ it should be
20 possible to achieve the same for DvH-FDH2. (3) Insights gleaned from DvH-FDH2 should
21 inform tunability of catalytic bias,¹²¹ limits to electrocatalytic reversibility,¹²² and design of
22 biomimetic metallosynthetics.¹²³ (4) Virtually nothing is known about the redox partners
23 of metallo-FDHs in the context of aerobic respiration.¹²⁴ The ability to aerially manipulate
24 DvH-FDH2 will enable strategies for solving problems previously considered intractable
25 in bioenergetics.

26

1 **METHODS**

2 **Genetics**

3 **Construction of *fdh* Deletion Strains** The naming schemes for the *fdh* genes follow the
4 convention established previously⁵⁶. DvH deletion strains (Table S1) were constructed
5 using methods already described^{125, 126}. Briefly, for the deletion of each predicted operon,
6 two plasmids were constructed: one to create a marker-exchange deletion and another
7 to remove the marker. Both plasmids are suicide vectors and require at least one
8 homologous recombination event to occur to provide the selectable phenotypes. A
9 phenotypic screen was performed to determine if a double recombination event took
10 place, thereby increasing the likelihood of choosing isolates that had the desired
11 genotype. Each vector contained a cloned copy of at least 300 bp upstream and a similar
12 DNA region downstream of the operon targeted for deletion that were captured in a
13 vector backbone containing the pUC origin of replication and a gene conferring
14 spectinomycin-resistance. The plasmids were constructed by the sequence and ligation
15 independent cloning (SLIC) technique¹²⁷ with amplicons obtained from PCR using the
16 primers found in Table S2 (Integrated DNA Technologies, Coralville, IA) and the Herculase
17 II DNA polymerase (Life Technologies, Grand Island, NY). For the marker-exchange
18 plasmids, the two DNA regions up- and down-stream are separated by an artificial, two-
19 gene operon consisting of *aph(3')-IIa* (conferring antibiotic resistance to 50 µg
20 kanamycin/mL in *E. coli* and 400 µg G418/mL in DvH) and the counter-selectable marker
21 uracil phosphoribosyltransferase (*upp*, DVU1025) genes. The marker-exchange plasmids
22 were introduced by electroporation into a strain containing a deletion of the *upp* gene
23 and the operon to be deleted. The transformed DvH cells were allowed to recover
24 overnight at 34 °C, as previously described¹²⁶. The cells were then grown for 3-5 days on
25 solidified MOYLS4 medium¹²⁸ containing G418 to select for transformants. Single isolates
26 were screened for sensitivity to 100 µg spectinomycin/mL (consistent with the double
27 homologous recombination event), sensitivity to 40 µg 5-fluorouracil/mL (5FU^S; to ensure

1 the counter-selection of 5FU resistance (5FU^R) would be effective) and maintenance of
2 resistance to G418. A putative marker-exchange deletion isolate was then chosen and
3 transformed with the marker-less deletion plasmid, as described above. The transformed
4 cells were recovered, plated on medium containing 5FU and the three phenotypic markers
5 again screened. For the marker-less deletion isolates, however, isolates were selected that
6 were 5FU-resistant and G418-sensitive showing that the marker exchange cassette had
7 been removed from the cell by double homologous recombination. Up to three isolates
8 with the desired antibiotic-resistance phenotype were further analyzed by Southern blot.
9 Once confirmed, one of these isolates was chosen as the marker-less deletion mutant.

10 For operon deletions, the upstream and downstream regions, respectively,
11 consisted of 858 bp and 806 bp (*fdh1*; DVU0586-0588), 795 bp and 878 bp (*fdh2*;
12 DVU2485-2481), and 976 bp and 970 bp (*fdh3*; DVU2809-2812). Parental strain JW710¹²⁶
13 was used for the deletion of *fdh1* and *fdh3*. Confirmation by Southern blot was
14 accomplished by digesting the genomic DNA of the parental and putative deletion strains
15 with AgeI (NEB, Ipswich, MA), separating the DNA fragments by gel electrophoresis, and
16 probing with the upstream region.

17 **Molecular Biology and Microbiology**

18 **Plasmid Construction** pMO9075 backbone was amplified via Phusion polymerase (New
19 England Biolabs #E0553S) using the primers pMo9075 slic_F and pMo9075 slic_R,
20 separated on 0.6% TAE (BioRad QBI 351-008-131) agarose gel (BioRad 161-3102) and
21 purified via gel extraction (Qiagen #28704). Inserts were amplified with Phusion
22 polymerase via standard reaction conditions. Primers 2482_pmo_F and 2481_pmo_R were
23 used to amplify FDH2 for cloning into pMO9075. Primers 2482_pmo_F and 2482_strII_R
24 were used to amplify DVU2482, introducing the upstream vector flank to DVU2482 and a
25 StrepII tag to the 3'-end of DVU2482. Primers strII_2481_F and 2481_pmo_R were used to
26 amplify DVU2481 with StrepII-tag overlap (while maintaining native intergenic spacer)
27 and downstream vector flank. Amplicons were separated in 0.6 % TAE agarose gels and

1 purified by gel extraction. Inserts were assembled with vector backbone via overlap
2 assembly using Gibson cloning (New England Biolabs #M5510A). Assembly reactions were
3 used to transform *E. coli* α -select chemically competent cells (Bioline BIO-85026) and
4 colonies were selected on YT glucose plus 50 mg/mL spectinomycin HCl (Sigma-Aldrich
5 S9007). For positive clones, 50 mL of transformant was grown in MDAG-11 formulated in
6 house¹²⁹ supplemented with spectinomycin, and the plasmid was purified using a Qiagen
7 Plasmid Midi Kit (Qiagen 12943).

8 **Bacterial growth.** DvH strains were grown on MOYLS4 medium (see Protocol S1), which
9 was adjusted to pH 7.2 with NaOH. Thioglycolate was added before bottling and after
10 equilibration to dinitrogen (Airgas NI NF200 or research grade). For generating inocula,
11 media were bottled anaerobically under dinitrogen (5 psi), 50 mL per 100 mL serum bottle
12 (Duran Wheaton Kimble 223747) with butyl rubber stopper (Chemglass CLS-4209-14) and
13 aluminum crimp seal (Wheaton 20-0000AS). For larger volumes, glass media bottles
14 (Pyrex 1395500, 13951L; Fisher 06-414-1C/06-41401D) were sealed with no. 6 neoprene
15 stoppers (RPI-259100-6) and capped with media bottle lid with a center bore to access
16 the stopper. Bottles were autoclaved and vitamins were syringed in from a filter sterilized
17 (RPI 256131) 1X stock just before inoculation.

18 **Transformation** One of our laboratories (C.S.R.) formulated a protocol that is different
19 from what has been reported earlier.⁴⁹ DvH strains were grown in 50 mL MOYLS4 in 100
20 mL serum bottle at 37 C with nitrogen headspace to near stationary phase and chilled on
21 ice. Cells were aerobically spun down in a 50 mL conical centrifuge tube (Corning 430828)
22 at 7,500 x g for 5 min, then washed twice in 17 mL of ice cold 15 mM Tris pH 7.2, 10%
23 glycerol supplemented to 1 mM with dithiothreitol. The final pellet was resuspended to 1
24 mL in the same buffer. A 100 μ L aliquot of cells was aerobically mixed on ice with 7.5 μ L
25 from plasmid midi prep (~2-3 μ g plasmid) and electroporated at 1.5 kV (1 mm gap
26 cuvette; MBP #5510) in an Eppendorf electroporator 2510. 1 mL of sterile anaerobic
27 MOYLS4 was immediately added and the entire volume was transferred to a bottle of

1 MOYLS4. The bottle was incubated at 37 C (Glascol, Micro-expression Vertiga shaker).
2 Once the culture recovered and became densely turbid, transfers were made to fresh
3 medium containing 100 µg/mL spectinomycin HCl. After two rounds of growth with
4 spectinomycin selection, freezer stocks in 10% glycerol were made. For colony selection
5 the same medium supplemented with 1.5% agar, 5 mM cysteine, 1 mM sodium sulfide,
6 and 100 µg/mL spectinomycin was used and kept in gas tight jars with an AnaeroGen 3.5
7 L Gas generating system pack (Oxoid). Colonies were picked into selective medium using
8 a sterile 1 mL syringe (Becton Dickinson 309659) fitted with an 18-gauge needle.

9 **10 L carboy growth of DvH-FDH2 producing strain CSR21271** For each carboy, the
10 strain was transferred from 10% glycerol freezer stock in MOYLS4 medium; ~0.5 mL of
11 stock added to a 50 mL bottle of anaerobic MOYLS4 medium, supplemented with vitamins
12 and 100 µg/mL spectinomycin hydrochloride. Transfers were made by nitrogen purged
13 syringe with 23-gauge needles (Becton Dickison 305190). The culture was incubated
14 overnight at 37 C or until mid-exponential phase of growth. 20 mL of the overnight culture
15 was used to inoculate a 500 mL bottle of MOYLS4 medium, containing vitamins and 100
16 µg/mL Spectinomycin HCl. The 500 mL culture was incubated overnight at 37 C. 10 Liters
17 of MOYLS4 medium in 2 L bottles, prewarmed, sterile, aerobic, with iron and EDTA
18 withheld, was poured into a sterile 10 L polypropylene carboy (Thermo 2250-0020). The
19 medium was completed by addition of filter sterilized vitamins, spectinomycin
20 hydrochloride (1g dissolved in 15 mL water; 100 µg/mL final) and 4.8 mL of iron chloride
21 (125 mM; Acros 423705000) / EDTA (250 mM; Fisher BP120-1) solution. The carboy was
22 closed and purged with nitrogen via a butyl rubber stopper port (Chemglass CLS-4209-
23 14) affixed to the lid (Figure S37A). 5 mL of 25% sterile neutral sodium sulfide (Alfa Aesar
24 36622) was then injected through the port and the carboy was mixed by shaking (Figure
25 S37B). Subsequently, the carboy was incubated at 37 C until resazurin indicator turned
26 colorless. The 500 mL culture (OD₅₅₀ ~0.6) was then transferred into the carboy via sterile
27 rubber transfer line (VWR-62993-726), 18-gauge needles (Becton Dickinson 305196) and

1 under nitrogen pressure (Figure S37C). The carboy was placed in an incubator (Sanyo
2 MCO-17A1C) at 37 C and the optical density (OD) was monitored at 550 nm (Beckman
3 DU-800 spectrophotometer) via 1 mL samples removed from the same port. Once OD₅₅₀
4 nm plateaued, the carboy was chilled in the cold room overnight (Figure S37D) and then
5 harvested by centrifugation at 8,000 x g (Beckman Avanti HP-26 XPI) in 1 L bottles. Cell
6 pellets were transferred to 50 mL conical centrifuge tubes, respun (7,500 x g; Corning
7 430828), and then froze at -80 C.

8 **Biochemistry**

9 **Protein Expression and Purification.** Strep-tag II DvH-FDH2 was purified from strain
10 CSR21271 (see Table S1). Unless specified otherwise all the following steps were done at
11 4 C and under atmospheric conditions. Nitrate, azide, thiols, or formate were not used at
12 any step of the purification or storage. Cells (~ 18 g) were suspended in six volumes of
13 50 mM sodium phosphate (Fluka 71505, Sigma-Aldrich S0786), pH 7.4, containing 150
14 mM sodium chloride and 1 mL of 50 x Complete Proteinase inhibitor (Roche 45582400; 1
15 tablet in 1 mL of MilliQ water), by gentle pipetting in cold buffer. Cells were lysed using
16 an Avestin C3 homogenizer and cell debris spun down at 4500 x g (Beckman Avanti HP-
17 26 XPI) for 15 min. Membrane vesicles were removed by centrifugation at 285,000 x g 1
18 hr (Beckman Optima L100XP). The clarified lysate was then fractionated by ammonium
19 sulfate precipitation with fractions pelleted at 10,000 x g for 10 min and the 40-70%
20 saturating fraction was retained and exchanged via centrifugal concentrator (Amicon 30
21 kDA molecular weight cutoff) into 100 mM Tris-HCL buffer pH 8, containing 150 mM NaCl
22 and 1 mM EDTA. The sample was loaded on to streptactin-XT superflow resin (IBA-
23 LifeSciences) and the column was washed with 40 volumes of the same buffer. StrepII-
24 tagged protein was eluted by several column volumes of 100 mM Tris-HCl buffer, pH 8,
25 containing 150 mM NaCl, 1mM EDTA, and 50 mM biotin (IBA-LifeSciences 2-1016-005).
26 The protein was concentrated via centrifugal concentrator (Amicon 30 kDA MWCO) and
27 exchanged into 20 mM Tris-HCl buffer pH 8.0, with or without 10% glycerol (Sigma-

1 Aldrich 49770) and stored at -80 C for future use. The protein concentration was estimated
2 by BCA assay (Thermo Fisher) versus a BSA standard.

3 **Gel Electrophoresis** DvH-FDH2 was separated on a Nupage 4-12% Bis-Tris Gel (Thermo
4 Fisher). The running buffer was 1x MES-SDS. The sample was loaded as 5 μ L of 12 μ M
5 DvH-FDH2 in 62.5 mM Tris-HCl buffer, containing 1.5% SDS, 10% sucrose, 0.0075%
6 bromphenol blue, pre-incubated at room temp (23 °C) for 30 minutes and then heated 5
7 minutes at 50 C. The protein was run alongside Precision plus Kaleidoscope prestained
8 standards (Bio-Rad #1610375) for 100 minutes at 100 V (Invitrogen mini gel tank A25977).
9 The gel was fixed in 40% methanol, 10% acetic acid, stained in 30% methanol, 10% acetic
10 acid, and 0.05% Coomassie blue G-250, and destained in 8% acetic acid. Gels were
11 scanned with a gel doc imager (Bio-Rad).

12 **Chromogenic Visualization (In-Gel Assay).** DvH-FDH2 was separated on a standard Tris
13 buffered 5% polyacrylamide gel, 2.6% crosslinker gel supplemented with 0.05% triton X-
14 100 (Fisher BP151-100). The running buffer was 25 mM Tris, 192 mM glycine and 0.05%
15 triton X-100 (v/v). Every other lane was loaded with 7.5 μ L of 3 μ M FDH2 in 20% sucrose,
16 0.25 M Tris pH 6.9, 0.05% triton, 0.0125% bromphenol blue. Electrophoresis was
17 conducted at 100 V for 209 minutes, 10 mA limit at 4 C. Lanes were cut into strips then
18 assayed in 10 mL of 0.24 mg/mL NBT (Invitrogen N6495) in 20 mM Tris pH 8.0 with or
19 without 10 mM formate (added as 100 μ L of 1 M). Strips were incubated with shaking for
20 15 minutes and then washed with MilliQ water for 3 x 5 min, and then scanned on a Bio-
21 Rad Gel Doc imager.

22 **NBT strip assay** 3 mm strips of chromatography paper (Whatman, 3MM CHR), cut with a
23 paper cutter, were soaked with a solution consisting of 300 μ M NBT (1 mL of 3 mM in
24 water) , 30 μ M PMS (Sigma P9625) (1 mL of 300 μ M in ethanol) in 8 mL Tris-buffered
25 saline pH 7.5, with or without 10 mM formate (Aldrich 798630-500g) (100 μ L of 1 M in
26 water). Strips were then spotted with 5 μ L of 12 μ M enzyme or control solution (buffer
27 only) and monitored for color development. PMS was added to the NBT solutions to 30

1 μM final from a 30 mM stock in water before soaking the strips. For the NBT only
2 experiment, strips of Whatman chromatography paper ($\sim 8.8 \text{ cm}^2$) were soaked with 140
3 μL of 293 μM NBT in 20 mM Tris pH 8.0 with or without 10 mM sodium formate. FDH2
4 was spotted onto strips as a 5 μL drop of 12 μM enzyme in 20 mM Tris pH 8.0.

5 **Metal analysis** Inductively coupled plasma mass spectrometry (ICP-MS; iCAP-RQ,
6 Thermo Fisher Scientific, Bannockburn, IL) was used in the KED mode to assess the metal
7 stoichiometry of DvH-FDH2. 25 μL of protein samples were placed in 15 mL conical tubes
8 followed by the addition of 200 μL of Optima grade HNO_3 . Samples were digested for 20
9 minutes at room temperature followed by the addition of 9.775 mL of Millipore H_2O to
10 obtain a final acid matrix of 2% HNO_3 (v/v). QCS27 high-purity standards (Delta Scientific
11 Laboratory Products) were used as a multi-element standard, as well as tungsten
12 individual standard. The following isotopes were analyzed: $^{56,57}\text{Fe}$, $^{63,65}\text{Cu}$, $^{77,78,82}\text{Se}$,
13 $^{95,96,98}\text{Mo}$, and $^{182,183,184}\text{W}$. All sample were run with one survey run and three main peak
14 jumping runs.

15 **Enzymology**

16 **Benzyl viologen assay** In this assay BV (colorless) is one-electron reduced to BV+
17 (purple)^{130, 131} by DvH-FDH2. Our workflow is described in Figure S38. FDH2 was assayed
18 in an argon (Airgas AR-300) atmosphere with 2 mM or 20 mM Benzyl viologen (Alfa Aesar
19 H66836) in 50 mM Tris pH 8.0 with 20 mM glucose and supplemented with glucose
20 oxidase (Sigma- G7141-50ku) 1U/mL, and catalase (Sigma C1345-G) 1 $\mu\text{g}/\text{mL}$, in a
21 stoppered quartz cuvette (Helma 110-10-40) fitted with a Suba seal silicone septum
22 (Sigma Z279730-25ea) and an 8 mm x 3 mm pivoted spin bar (VWR 37119-6183). Assays
23 were performed under argon in a UV-2600i spectrophotometer (Shimadzu Life Sciences)
24 equipped with a T2 Peltier/stirring unit (Quantum; T = 25 $^\circ\text{C}$; stirring speed 650 rpm) and
25 monitored at 557 nm. Measurements were started by the addition of 10 μL formate or
26 buffer blank using a 10 μL Hamilton syringe (under an argon headspace) to a 2.5 mL
27 reaction mix containing pre-equilibrated 1.6 nM FDH2.

1 **Phenazine Ethosulfate (PES) / Dichlorophenolindophenol (DCPIP) Assay** Here, PES
2 reacts directly with the enzyme, gets reduced and then transfers electrons to DCPIP non-
3 enzymatically.⁷⁸ Our workflows are described in Figures S39 and S40 We chose PES over
4 PMS because of its long-term stability.¹³² Furthermore, we took the necessary precautions
5 in handling and preparing the reagents.¹³³ DvH-FDH2 was assayed in 1 mM PES (Sigma-
6 Aldrich P4544-1G) / 93 μ M DCPIP (98% pure; Acros 152870100) in 50 mM Tris pH 8.0.
7 Reactions, 2.5 mL volume, were set up to the desired formate concentration and then
8 started by addition of 10 μ L of 400 nM FDH2 (1.6 nM final). Assays were performed using
9 a UV-2600i spectrophotometer (Shimadzu Life Sciences) equipped with T2 Peltier/stirrer
10 accessory (Quantum) and monitored at 600 nm. For aerobic measurements (Figure S39),
11 reactions were performed in open top styrene disposable cuvettes (Brand 75907D) with 6
12 mm x 9 mm cuvette stirrer (Cowie 001.1609) spun at 400 rpm, and reactions started by
13 addition of 10 μ L of 400 nM FDH2 via micropipette. Some aerobic experiments were
14 conducted using a Cary 300 UV-Vis spectrophotometer (Agilent) at a volume of 3.0 mL
15 with a 9 mm x 8 mm spin bar (Sigma Z363545) cuvette stirrer. For the anaerobic
16 conditions (Figure S40), reactions were set up under argon in screw cap quartz cuvettes
17 (Starna 1-SOG-10_GL14-S) sealed with Suba-Seal 13 white rubber septa (Sigma Z167258).
18 The stirrer was an 8 mm x 3 mm pivot bar (VWR 37119-6183), stirring at 650 rpm.
19 Reactions were started by addition of FDH2 via 10 μ L syringe (Hamilton 701N 80300)
20 under an argon atmosphere.

21 **Data analysis** For classical steady-state kinetics, initial velocities (guided by residual plots)
22 were obtained using ICEKAT.¹³⁴ KinTek Explorer^{75, 77} (version 10.1.6, KinTek Corporation)
23 was used to perform global fitting of enzyme kinetics data to Schemes 1 and S1. This is
24 based on numerical integration of rate equations. Confidence contour analysis was
25 performed to assess whether the parameters were properly constrained by the data.

26
27

1 **Spectroscopy**

2 **Electronic** FDH2 spectra were collected at 23 °C in 50 mM Tris pH 8.0 using a screw cap
3 1 cm pathlength quartz cuvette (Starna; 1-SOG_10_GL14s). Under aerobic conditions, the
4 spectrum of air equilibrated enzyme was collected first. Then, 10 mM formate was added,
5 and the spectrum was measured again. The cuvette was then capped with silicone septa
6 (Starna GL14/SI) and 10 µL of 2 mM of dithionite was added under argon before collecting
7 a spectrum. For anaerobic measurements, FDH2 was gassed with argon in the sealed
8 cuvette before addition of formate or dithionite. Reduced spectra were also measured
9 using dithionite as the sole reductant (in the absence of formate).

10 **Electron Paramagnetic Resonance (EPR)** All samples were prepared in 20 mM Tris-HCl,
11 pH 7.6 containing 10% glycerol (v/v). Anaerobic samples were first purged with Ar and
12 then transferred to septum-sealed, Ar-flushed EPR tubes and reduced with either 20 mM
13 anaerobic sodium formate or ~4 mM anaerobic sodium dithionite. Aerobic samples were
14 reduced directly in open EPR tubes with either 20 mM sodium formate or ~2 mM sodium
15 dithionite. All samples were subsequently frozen in a dry ice/ethanol bath, then
16 transferred to liquid nitrogen for storage. The anaerobic sample reduced with sodium
17 dithionite was incubated for 12 h prior to freezing.

18 EPR spectra were recorded using a Bruker EMX spectrometer operating WinEPR version
19 4.33 acquisition software and equipped with a Bruker ER 4119HS high sensitivity X-band
20 cavity and gaussmeter. Temperature was controlled with a Bruker variable temperature
21 unit and a liquid nitrogen or liquid helium cryostat. For purposes of comparison, all
22 spectra were calibrated to a microwave frequency of 9.385 GHz. Detailed instrument
23 settings are included in the figure captions. Simulations were performed using the
24 EasySpin 4.5.5 software package.¹³⁵ Simulations included a “weight” term, which was used
25 to estimate the relative contribution of each component to the composite spectrum.

26 **Nuclear Magnetic Resonance.** NMR data were recorded on an Agilent DD2 500 MHz
27 spectrometer equipped with a 5 mm quadruple (¹H, ¹³C, ¹⁵N, ³¹P) PFG Penta Probe, which

1 was maintained at 25 °C. ¹³C data were acquired with 70332 points with a spectral width
2 of 30478 Hz, 242 ppm centered at 110 ppm, with proton-decoupling on throughout the
3 experiment (1 s delay between transients and 1.15s of acquisition time) and the number
4 of transients collected ranged from 64 to 1024. The fids were zero-filled and multiplied
5 with a 3-Hz line-broadening function prior to Fourier-transformation; the final size of the
6 spectrum was 65536 points. Proton data were recorded with 16384 points with a spectral
7 width of 7530 Hz (15 ppm centered at 4.7 ppm) with pre-saturation (2 s) to suppress the
8 water peak; 1 s delay between transients were used. Additional parameters are detailed
9 in Figures S15 – S23. Reference spectra of ¹³C-formate (9.5 – 10.5 mM in 100 mM sodium
10 phosphate buffer, pH 6 or 7.5) and ¹³C-sodium bicarbonate (4.8 mM in 100 mM sodium
11 phosphate buffer, pH 6) were first collected using standard 5 mm thin-walled NMR tubes
12 (Wilmad). 10% D₂O was used to obtain internal signal lock. Subsequently, 1.3 μM DvH-
13 FDH2 was added to the tube containing ¹³C-formate (pH 7.5), mixed, and spectra were
14 recollected. Upon completion, 2 mM PES was added to the same tube, mixed, and
15 remeasured. Independently, this process was repeated with ¹³C-formate at pH 6. NMR
16 data were processed with MestReNova NMR suite version 14.2.1-27684.

17 **Quantification of O₂ Reduction**

18 **Oximetry** A Clark-type O₂ electrode (Oxygraph Plus System from Hansatech Instruments,
19 UK) was used to monitor changes to the dissolved O₂ concentration, which corresponds
20 to 267 μM at 23 °C. O₂ saturation under these conditions would be equivalent to 1.27
21 mM. A decrease in O₂ level would indicate that O₂ was being consumed during aerobic
22 catalysis. Conversely, O₂ evolution would be diagnostic of catalase activity. The
23 electrode was calibrated each time before use with air saturated Milli-Q water and
24 dithionite per manufacturer's instructions. Freshly made reagent stocks and buffer
25 solutions were used throughout. 1 mL reactions were performed at 23 °C in a closed cell
26 using air-saturated 100 mM Tris-HCl, pH 8, containing 1 mM EDTA (Fisher BP120-1). The

1 latter was added to limit adventitious metal ions from mediating O₂ consumption. After
2 obtaining a stable baseline with the buffer, 10 mM formate was added, and the baseline
3 was allowed to stabilize. The reaction was started by the addition of 50 nM DvH-FDH2.
4 Once the O₂ consumption plateaued, 2 μM catalase (Sigma C1345-G) was added.
5 Catalase breaks down H₂O₂ to water and dioxygen (2H₂O₂ → 2H₂O + O₂) To test the effect
6 of additives, the order of addition was changed. For example, to test whether DvH-FDH2
7 had catalase activity, H₂O₂ was added to buffer first, followed by the enzyme. Similarly,
8 to assess the effect of SOD on O₂ uptake, SOD was the first component to be added. O₂
9 consumption rates were calculated as described before.¹³⁶ Initial velocities were
10 determined from the gradients of [O₂] versus time traces after subtracting O₂
11 consumption under the same experimental conditions without FDH2.

12 **Quantification of Hydrogen Peroxide**

13 **Amplex Red method:** 5 mg of Amplex Red (AR; Invitrogen A12222) was dissolved in
14 0.9725 mL neat DMSO to yield a 20 mM solution. Horseradish peroxidase (HRP;
15 Sigma P8250-5ku) was prepared at a concentration of 10 U/mL (45.5 μg/mL) in sodium
16 phosphate pH 7.4. Prior to assay, a 2x working solution was prepared from 10.6 μL of 20
17 mM AR, 80 μL of HRP, 1.6 μL of 0.5 M Diethylenetriaminepentaacetic acid (DTPA;
18 TCI D0504) and 3.9 mL of 50 mM sodium phosphate pH 7.4 and kept in the dark.
19 Production of H₂O₂ was measured by preparing reaction mixtures in a Costar 3915 black
20 flat bottom 96 well plate. Reactions consisted of 50 mM sodium phosphate pH 7.4, with
21 desired amounts of sodium formate added from a 50 μM stock and initiated by addition
22 of 5 μL of 32 nM DvH-FDH2 in the same buffer to a volume of 50 μL. This approach
23 allowed H₂O₂ generation to commence prior to the introduction of AR/HRP mixture.
24 H₂O₂ (Sigma-Aldrich H1009-100mL) standard curve was generated in the same buffer to
25 a volume of 50 μL. Detection was initiated by addition of 50 μL of the 2x Amplex Red/HRP

1 working solution and fluorescence was scanned in top read mode at medium sensitivity
2 on a SpectraMax M2 (Molecular Devices) plate reader (excitation 530 nm and emission
3 590 nm) every four minutes for twelve minutes (23 °C). Independently, we assessed
4 whether outcomes differed when order of addition was varied. To that end, in one set of
5 assays, 5 μ L of 32 nM FDH2 was added after AR/HRP. Here, 0.5 mM DTPA was used
6 instead of 0.1 mM.

7 **Coumarin boronic acid (CBA) assay:** 10 mg of coumarin-7-boronic acid (CBA) Cayman
8 Chemicals 14051) was dissolved in 3.33 mL of DMSO. 101 μ L of the CBA stock and 1.6 μ L
9 0.5 M DTPA were added to 3.9 mL 50 mM sodium phosphate pH 7.4 to produce a 2x
10 working solution. The remaining steps essential identical to those used in the Amplex
11 Red assay except that the plate was shaken at 400 rpm in an incubator (23 °C) for 15 min
12 prior to CBA addition. Fluorescence detection was initiated by addition of 50 μ L of the
13 CBA 2x working solution, and the plate was scanned in fluorescence mode (excitation 332
14 nm and emission 470 nm).

15 The following controls were common to both assays. (1) Heat denatured DvH-FDH2, (2)
16 Addition of catalase (100 U/mL), (3) Addition of superoxide dismutase (10 U / mL), (4)
17 Sans FDH2, (5) Fluorogenic substrate excluded, (6) Formate excluded, and (7) Buffer alone.

18 **Superoxide Generation / Redox Bifurcation.** Native (Sigma C2506) and partially
19 acetylated (Sigma C4186) equine heart cytochrome c were used to assess superoxide
20 production by DvH-FDH2. The integrity of oxidized cytochrome c was validated by
21 establishing the presence of 695 nm transition. The reduction of 30 μ M (native) or 60 μ M
22 (partially acetylated) cytochrome c by DvH-FDH2 was followed at 550 nm in a 1 cm
23 pathlength cell (Shimadzu UV-2600i spectrophotometer). We used two-fold higher
24 concentration of acetylated cytochrome c to offset its slightly weaker reactivity with
25 superoxide.⁹² The reaction mix (total volume 2.5 mL) was stirred (Cowie 001.1609) at 300
26 rpm (Quantum T2/Peltier unit) and maintained at 25 °C. For aerobic experiments, open
27 top styrene disposable cuvettes (Brand 75907D) were used. For anaerobic measurements

1 under argon, screw cap quartz cuvettes (Starna 1-SOG-10_GL14-S) sealed with Suba-Seal
2 13 white rubber septa (Sigma Z167258) were used. To 50 mM Tris-HCl buffer, pH 8,
3 containing cytochrome c, enzyme (1.6 nM final) was added first to obtain the background
4 signal. Subsequently, 10 μ M formate was added to start the reaction. Upon completion,
5 ~ 2 mM dithionite was spiked into the mix to estimate the amount of remaining oxidized
6 protein. Controls devoid of formate, enzyme, and cytochrome c were also employed.
7 The effect of superoxide dismutase (10 – 100 U / mL) or catalase (100 – 400 U / mL) was
8 tested independently.

9 **Structural analysis**

10 Protein alignments were constructed using MUSCLE or MAFFT. Structural alignments
11 were performed using Chimera v1.16. Amino acid sequences of the large (DVU2482) and
12 small (DVU2481) subunits of DvH-FDH2 were input together for running structure
13 predictions using a modified version of AlphaFold2.1.¹⁰¹ Because this algorithm does not
14 recognize Sec, a Cys was substituted and that Tat signal peptide (see Fig. 3A) was not
15 included. A dedicated Google Colab notebook ([AlphaFold.ipynb-Colaboratory](#)
16 [\(google.com\)](#)), which does not utilize homologous structures for making predictions was
17 used with default settings. We also independently predicted the structures of DVU2482
18 and DVU2481 using a full version of AlphaFold2.1. The resulting heterodimeric structures
19 were superposed on the DvH-FDH1 counterpart determined via X-ray crystallography
20 (PDB ID: 6SDV²⁰) to assess similarities and differences. Difference distance matrices were
21 computed using Chimera v1.16. Structure visualizations and manipulations were done via
22 Pymol.

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8 **Author Contributions**

9 C.S.R. conceived the work, designed the experiments, acquired funding, and directed the
10 project. J.E.G. created new DvH strains (identified by the CSR prefix in Table S1),
11 established a novel scaleup strategy and streamlined workflow for generating DvH
12 biomass, optimized DvH transformation protocol, developed a DvH expression platform
13 (cloning, homologous expression, and purification), characterized DvH-FDH2 via
14 electronic spectroscopy, demonstrated in-gel activity, measured enzyme kinetics data,
15 confirmed cytochrome *c* reduction, and validated formate oxidase activity under the
16 supervision of C.S.R. Oximetric measurements were made by C.S.R. and J.E.G. D.N.
17 performed EPR measurements and interpreted the results under the supervision of R.H.
18 DvH deletion strains (with a JW prefix in Table S1) were generated by Q.G., and G.M.Z.
19 measured their growth characteristics under the supervision of J.W. K.H. acquired NMR
20 spectra. C.S.R. analyzed data from enzyme kinetics, oximetry, and NMR, ran the AlphaFold
21 predictions and interpreted the structural differences, and wrote the manuscript. All
22 authors have given approval to the final version of the manuscript.

23 **Notes**

24 The authors declare the following competing financial interest(s): C.S.R. and J.E.G. are
25 inventors on a patent application related to this work filed by the University of Maryland
26 Baltimore. The authors declare no other competing interests.

27

1 **Associated Content**

2 Supporting Information

3 Strains and plasmids (Table S1), Primers (Table S2), Growth curves of JW2111 and JW2121
4 (Figure S1), NBT strip assay (Figure S2), Sequence alignment of DvH-FD2 and DvH-FDH1
5 (Figure S3), ICP-MS quantification of metals in DvH-FDH2 (Table S3), Comparison of
6 electronic spectra of DvH-FDH2 and DvH-FDH1 (Figure S4), Simulation of Fe/S centers of
7 formate-reduced DvH-Fdh2 prepared under aerobic conditions (Figure S5), EPR
8 parameters of DvH-FDH2 (Table S4), Steady-state kinetic parameters of SRB-FDHs
9 compiled from the literature (Table S5), Minimal catalytic model used to assess BV kinetics
10 (Scheme S1), Ground truthing published enzyme kinetics data of Dd-FDH3 (Figure S6) and
11 DvH-FDH1 (Figure S7), Steady state kinetics of DvH-FDH2 (Table S6), 2 mM BV reduction
12 by DvH-FDH2 (Figure S8), 20 mM BV reduction by DvH-FDH2 (Figure S9), GO and catalase
13 do not affect the kinetics of BV reduction by DvH-FDH2 (Figure S10), Optimizing DCPIP
14 and PES for aerobic enzyme kinetics (Figure S11), Reduction of PES/DCPIP by DvH-FDH2
15 in air (Figure S12), Anaerobic reduction of PES/DCPIP by DvH-FDH2 (Figure S13), Effect of
16 azide on the reduction of PES/DCPIP (Figure S14), Effect of nitrate on the reduction of
17 PES/DCPIP (Figure S15), ^{13}C NMR spectra of ^{13}C -formate + DvH-FDH2 (pH 7.5) (Figure
18 S16), ^{13}C NMR spectra of ^{13}C -formate + DvH-FDH2 + PES (pH 7.5) (Figure S17), ^{13}C NMR
19 spectra of ^{13}C -formate + DvH-FDH2 (pH 6) (Figure S18), ^{13}C NMR spectra of ^{13}C -formate
20 + DvH-FDH2 + PES (pH 6) (Figure S19), ^{13}C NMR spectra of ^{13}C - NaHCO_3 (pH 6) (Figure
21 S20), ^{13}C NMR spectra of ^{13}C -formate (pH 6) (Figure S21), ^{13}C NMR spectra of ^{13}C -formate
22 (pH 7.5) (Figure S22), ^1H NMR spectra of ^{13}C -formate (pH 6) (Figure S23), DvH-FDH2 lacks
23 catalase activity (Figure S24), Assessing the effect of variables on amplex red assay (Figure
24 S25), Evaluating how variables impact the CBA assay (Figure S26), O_2 uptake by DvH-
25 FDH2 is not impacted by SOD addition (Figure S27), Acetylated cytochrome c is not

1 reduced during aerobic FDH2 catalysis (Figure S28), Initial rates of native cytochrome c
2 reduction is unaffected under aerobic and anaerobic conditions (Figure S29), Spectra of
3 native cytochrome c reduced by FDH2 (Figure S30), Lack of effect of SOD and catalase on
4 the aerobic reduction of equine cytochrome c by DvH-FDH2 (Figure S31), Confidence
5 metrics of AlphaFold2.1 prediction (Figure S32), Variation in backbone RMSD (Figures S33
6 and S34), Residue-residue distance map (Figure S35), Active site of DvH-FDH2 (Figure
7 S36), MOYLS4 medium recipe (Protocol S1), DvH growth in 10 L carboy (Figure S37), BV
8 assay workflow (Figure S38), Aerobic PES/DCPIP assay workflow (Figure S39), and
9 Anaerobic PES/DCPIP assay workflow (Figure S40), and TOC Graphic (OutFoxing Oxygen
10 via Redox Bifurcation)

11 This information is available free of charge on the ACS Publications website.

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OutFOXing Oxygen via Redox Bifurcation

