
The functional complexity of [NiFe] hydrogenases in sulfate reducing bacteria (genus; *Desulfovibrio* spp)

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Abstract: Sulfate-reducing bacteria are categories of bacteria and archaea that can obtain energy by oxidizing organic compounds or molecular hydrogen (H₂) while reducing sulfate (SO₄²⁻) to hydrogen sulfide (H₂S). By analysis, these organisms "respire" sulfate rather than oxygen, a form of anaerobic respiration, the oxidation of hydrogen by the primary genus of Sulfate Reducing Bacteria (*Desulfovibrio*, *Desulfovibrio desulfuricans*) is catalyzed by enzymes called Hydrogenases. Three basic types of hydrogenases have been widely isolated from the primary genus of sulfate-reducing bacteria *Desulfovibrio* which differ in their structural subunits, metal compositions, physico-chemical characteristics, amino acid sequences, immunological activities, structural gene configuration and their catalytic properties. Broadly, hydrogenases can be considered as 'iron containing hydrogenases and nickel-containing hydrogenases. The iron-sulfur-containing hydrogenase enzyme contains two ferredoxin-type (4Fe-4S) clusters and typical iron-sulfur center believed to be involved in the activation of H₂ yet it is the most sensitive domain to CO and NO₂⁻. Even though it is not featured in all species of genus *Desulfovibrio*. The nickel-(iron-sulfur)-containing hydrogenases, [NiFe] hydrogenase possess two 4Fe-4S centers and one 3Fe-xS cluster in addition to nickel and have been found in all species of *Desulfovibrio* with strong resistance to CO and NO₂⁻ so far investigated. The genes encoding the large and small subunits of a periplasmic and membrane-bound species of the [NiFe] hydrogenase have been cloned in *Escherichia coli* and sequenced, however the functional complexity of the hydrogenase system remained unexplored as a result of the metabolic diversity in *Desulfovibrio* spp. The [NiFe] hydrogenase plays an important role in the energy metabolism of *Desulfovibrio* spp. Thus, the expression of the encoded structural genes would be an excellent marker for the metabolic functionalities under specific inducible environment.

Keywords: Hydrogenases, Sulfate-Reducing Bacteria, Iron-Containing Hydrogenases, Genus: *Desulfovibrio*, Nickel-Iron-Containing-Hydrogenases, Nickel-Iron-Selenium-Containing Hydrogenases

1. Introduction

Sulfate-reducing bacteria employ different carbon sources as electron donor and channels the sulphate moiety as the respiratory substrate while other molecules are being reduced [1]. SRB especially the genus *Desulfovibrio*

ultimately require hydrogen for their metabolism [2]. Moreover, such utilization of molecular hydrogen serve as generation intermediate specially for chemiosmotic gradient via organic molecule degradation [3] and lead to production of hydrogen in sulfate-depleted media [4]. Two major models have been described to explain the energy

transduction mechanism in sulfate-reducing bacteria. The first is the so-called “hydrogen cycling” model, described by Odom and Peck [5], suggests that molecular hydrogen originated from lactate oxidation (an important intermediate metabolite) is produced within the cytoplasm and diffuses across the membrane; then, a periplasmic hydrogenase will oxidizes the diffused hydrogen while the electrons are transferred along the cytoplasmic membrane and used to reduce electron acceptors, such as sulfate, in a vectorial electron transfer process [6]. More recently, [7] proposed a generalized system that describes how does hydrogen help in different environmental condition to improves the growth of SRB species (*D. Vulgaris*). The first unified model shows that, the process of hydrogen transformation throughout in *D. vulgaris* confers an enormous important the metabolic system of the species. The second model states that the physiological function of hydrogen metabolism is to regulate the redox potential of electron carriers such as ferredoxin and/or cytochromes, in order to control the exchange of proton and vectorial electron sensation to stimulate and produce a membrane bound proton motive force [8]. Thus, the metabolic pathway designated by these two models reveals that H₂ shows an important part in the bioenergetics of sulfate-reducing bacteria; however, the specialized protein for reverse oxidation of molecular H₂ is termed Hydrogenases.

Nickel-Iron hydrogenase enzyme; form a critical type of protein known to oxidize the reversible activation of molecular H₂ in both eukaryotes and some prokaryotes. The active site of the enzyme establishes a bases for chemical reaction by which the organism save and utilize energy through redox mechanism (Figure 1). This process is very important especially for deoxygenic bacteria such as sulfate-reducing bacteria of the genus *Desulfovibrio*. [9-11]. One of the practical applications of this enzyme has proven to be a renewable and more environmentally friendly energy source to such an extent that, scientist can engineer the condition to enable the [NiFe] hydrogenase to induce unilateral hydrogen production. In addition, enzymes which are analogues of [NiFe] hydrogenase can also generate hydrogen gas.

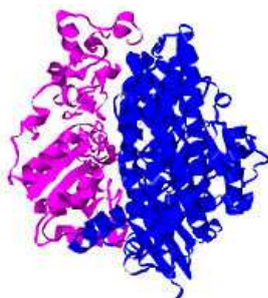


Figure 1. three dimensional structure of [NiFe] hydrogenase in *D. Vulgaris*.; the small subunit (magenta); the large subunit (blue) both are configured by Jmol (J-mol)with coordinates from 1H2A.pdb.

1.1. Structural Classification of Hydrogenase Enzymes

The enzymes are categorised into two major division based on the metals found in their active sites. The [Fe] containing hydrogenases and the [NiFe] hydrogenases including the sub-group of the [NiFeSe] hydrogenases (where a cysteinyl ligand of the nickel site is replaced by a seleno-cysteine) [12]. These enzymes have been characterized to differ in terms of their specific activity and bio-directionality [13].

Basically, the Hydrogenase enzyme are further categorised in to one of the following types depending on metal atoms they constitute in their active site. The three types are; [NiFe], [FeFe], and [Fe]. Until recently, the iron containing are known to be metal free, Thauer *et al.*, [13] in his experiment described that, metal free enzyme now called Iron containing hydrogenase are previously identified as metal free they are now called Iron containing hydrogenase enzymes [14], this include no iron-sulfur cluster and mononuclear Fe active site in contrast to [FeFe] hydrogenases. [FeFe] and [NiFe] even though contained different metals in their active site yet share some structural similarities. The similarities are;

- ✓ The entire enzyme possessed active site and a few Fe-S clusters that are found beneath the protein.
- ✓ The active site contained metallocluster with each metal; they are coordinated by cyanide (CN⁻) ligands and carbon monoxide (CO). The active site induces catalysis. [15]

The number and cellular localization of hydrogenase varies within different species of *Desulfovibrio*. For example, *D. gigas* has a single variant of [NiFe] hydrogenase while *D. Vulgaris* Hildenborough contains three types of hydrogenase, an [NiFe] hydrogenase located in the membrane, an [Fe] hydrogenase in the periplasm and an [NiFeSe] hydrogenase on the membrane adjacent to the cytoplasm [33]. The difference in their structural configuration is described in figure 2.

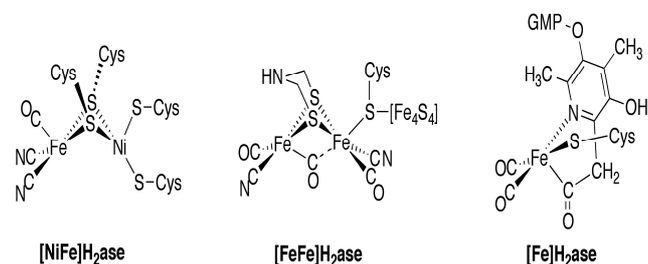


Figure 2. Structural configuration of different forms of hydrogenase found in SRB

“The sulfate-reducing bacterium *Desulfovibrio vulgaris* Hildenborough possesses four periplasmic hydrogenases that facilitate the oxidation of molecular hydrogen. These include an [Fe] hydrogenase, an [NiFeSe] hydrogenase, and two [NiFe] hydrogenases encoded by the *hyd*, *hys*, *hyn1*, and *hyn2* genes, respectively. These enzymes are present ubiquitously in both *Archaea* and *Bacteria*, including many

SRB. The fully sequenced *Desulfovibrio vulgaris* Hildenborough has a total of six hydrogenases” [16]. The four enzymes including a soluble [Fe] hydrogenase (*Hyd*) are periplasmic; implies that they presumably involved in hydrogen catalysis [17], two membrane-associated [NiFe] hydrogenase isozymes (*Hyn1* and *Hyn2*) [15], and a membrane-associated [NiFeSe] hydrogenase (*Hys*) are all involved in hydrogen oxidation [18].

1.2. Structure of [NiFe] Hydrogenases

X-ray crystallography was employed for the study and the structure of [NiFe] hydrogenase enzymes was obtained from the unified results of five different sulfate-reducing bacteria viz; *D. Desulfuricans* [19], *D. Frutosovorans* [2] [20] *D. Vulgaris* Miyazaki F, [21] *D. gigas*, and *Desulfomicrobium baculatum*. [12]. As described above, the enzyme in *D. vulgaris* Miyazaki F shown in Figure 1 consist of larger subunit (blue) with a molecular mass of 62.5 kDa, and Ni-Fe active site while the smaller subunit is in magenta with a molecular mass of 28.8 kDa, and contains the Fe-S clusters at the active site.

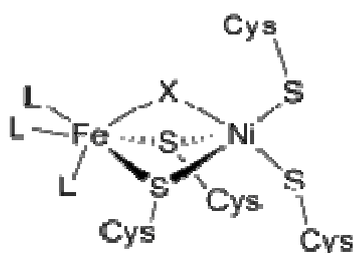


Figure 3. Structural configuration of [NiFe] hydrogenase from *D. vulgaris*

Based on the X-ray crystallography and infrared spectra, the [NiFe] enzymes active site were reported to consist of $(S-Cys)_4Ni(\mu-X)Fe(CO)(CN)_2$; the generic ligand coded X occur in form of either a sulphur, oxide, hydroperoxide, or a hydroxide in an oxidized state only (Figure 3). The nickel atom functions in redox reactions while the iron atom is consistently in a Fe (II) coordination state. [22]. However, they were identified as two carbon monoxide ($C\equiv O$) molecules and one cyanide ($C\equiv N$) molecule. [23]

1.3. Application of [NiFe] Hydrogenases

The [NiFe] Hydrogenase enzyme have numerous role to play over a wide range of biotechnology A preliminary characterization of the electron carrier system of newly isolated strain indicates the presence of a membrane bound [NiSeFe] hydrogenase, a periplasmic tetrahemic cytochrome c3, APS reductase and desulforubidin as dissimilatory bisulfite reductase [24]. The enzymes is capable of ensuring both the catalysis and consumption of

H_2 , usually this is a typical characteristics of hydrogenase family. The most important application of Hydrogenase enzymes are;

- ✓ as anode catalysts in enzyme fuel cells,
- ✓ the harnessing of the H_2 oxidation reaction to provide a source of low-potential electrons for use in reduction reactions, and
- ✓ The enzymatic photo-generation of H_2

The hydrogenase activity can provide an important resource for the cells, as an “energy valve”, when they are forced to grow in a medium, where the electron donor or acceptors are scarce. This may be relevant in environments, where bacteria can use any available metallic structures, driving to bio-corrosion processes [25][26] such as those occurring in ships hulls or heat exchangers.

1.4. Biological function of [NiFe] Hydrogenases

Scientific hypothesis suggested that, if earth were to be occupied with hydrogen, hydrogenase enzymes were potentially made to form molecular hydrogen which can be used by other microbes in a given ecosystem as a sources of energy [27]. Such microbial communities are usually bottom dwellers, thriving in a region where sources of light for photosynthesis is very minimal, thus hydrogenase enzyme evolved to generate energy for itself and other systematic biota.

Hydrogenase function to act as a “valve” in anoxic metabolism by regulating the hyper reducing equivalent in photosynthesis microbes [28]. In addition, hydrogenase were found to induces trans-membrane motive force result in to bioremediation of chlorinated compound, recovery of heavy metal contaminant from contaminated site. Examples of the microbes are pathogenic bacteria and parasite function as energy producer and virulence [27].

2. Mechanism of the [NiFe] Hydrogenase

There had been a controversy over a precise mechanism behind [NiFe] hydrogenase. The debate started in 2009 where Higuchi and his fellows examined the structure of *Desulfovibrio vulgaris* Miyazaki base on X-ray crystallography and spectroscopic data [23] they proposed that, during the excited state of catalysis, the metal (Fe) found at the active site of the enzyme remained un-static while the Ni metal ions involve in the catalysis (redox chemistry). The two redox stage of the enzymes catalysis are;

- ✓ Inactive redox states
- ✓ Active redox states and
- ✓ The CO-inhibited states (Figure 4).

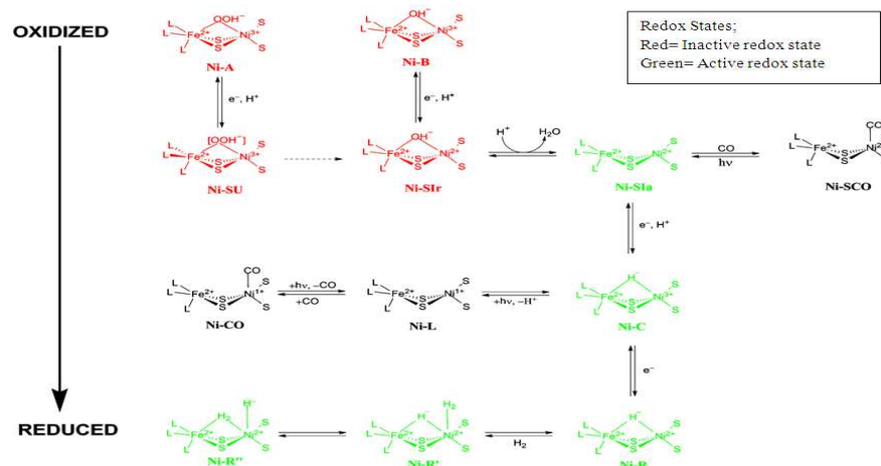


Figure 4. Redox transitions of the metal active site of [NiFe] hydrogenase [22].

(1) Resting redox state (inactive)

[NiFe] oxidizes in two forms; the unsteady state (Ni-A) and steady state Ni (Ni-B). The two stages are activated through one electron reduction. The rate conversion of Ni-B to Ni-sr occurs in second while that of Ni-su take place about one hour. The difference in the activation mode of the two stages was as a result of the differences in nature of ligand that bridge the two redox stage. Condensation reaction take place at Ni-Slr state which result to the formation of Ni-Sla state by releasing water, Hence the termed condensation reaction [29]

(2) Active Redox state;

Unlike the inactive form of [NiFe] hydrogenase enzymes, the active states constitute 3 forms viz; Ni-Sla, Ni-R and Ni-C. Ni-C is also called the light sensitive state formed through electro-reduction of Ni-Sla. Ni-C is the most sensitive form because it signifies vividly that hydrogen catalysis take place in [NiFe] hydrogenase active site [23].

(3) The CO-inhibitory state

CO, is one of the inhibitory factor that bind directly to the Ni metal ion at the active site of the enzyme changing it configuration or conformation to form Ni-Sco, Because Ni-C is photo sensor when illuminated at 100k, forms the Ni-Co state in the presence of CO. [22]

2.1. Factors Affecting the Functionality of NiFe Regulation

The understanding of the systematic hydrogenation and subsequent utilization in microbes is an ultimate goal for basic and applied research. Hydrogen metabolism is accompanied by the role of a key enzyme termed [NiFe] Hydrogenation.



The constitution of Ni and Fe in the enzyme make it metalloenzyme, like other group of metallo-protein, Hydrogenase enzyme is very sensitive to some factors such as high temperature, oxygen inactivation, CO, CN and other environmental factors in order to obtained high functional efficiency of the enzyme . Favourable condition

must be provided for smooth biotechnological application.

2.2. Expression of the [NiFe] Hydrogenase Gene in Hydrogenase in *Desulfovibrio spp*

An analytical research was carried out to study the cellular function of [NiFe] hydrogenase by comparing gene (*hyd* and *hyn1*) and novel artificially constructed mutant gene (*hys* and *hyn-1-hyd*) [30]. The two distinct growth rates were compared to the wild type growing in a medium where lactate and hydrogen is used as a source of carbon for desulfurization. The expression of gene was studied using microarray hybridization and real time Polymerase Chain Reaction using mRNA. Such mRNA was extracted from cell reported to grow in three different culture condition viz; 5% hydrogen as a sources of electron donor , 50% hydrogen and mM lactate [31].

The results reveals that; the strain growing in 50% H₂ and lactate as a source of electron donor containing enzymes lacking Fe at the active site were adversely subjected to rudimentary growth. It was further reported that, the cells that suffered the most are the once growing in small amount of hydrogen as low as 5% H₂, hence lacking [NiFeSe] at the active site. Thus the impairment order for the expression of *hyd* gene follows; lactate <50% hydrogen < 5% while the expression of *hys* gene follows the reverse case. It is thus suggested that, there is a strong correlation between the intracellular hydrogen concentration and the growth of strain with lactate and 50% hydrogen where as grow in low amount of intracellular hydrogen is associated with growth in 50% hydrogen concentration. This induces low activity and high affinity of hydrogenase enzyme containing [NiFeSe], perhaps, it is clearly understood that the concentration of hydrogenase gene, although the distinct influence is yet to be known.

2.3. Analysis of Distinct Genes Function

Caffrey and his co-workers had described certainly the expressivity of multiple gene arising from hydrogen and lactate metabolism [30]. It was pointed out that, majority of the expression stimulation were induced in a cell growing

in a medium rendered with hydrogen of 50%, 5%, vol/vol or lactate as source of electron donor. Up regulation of the expression of *hyn* gene and down regulation of the of the *hyd* gene were as a results of changes from lactate to hydrogen. Such comparison had establishes a relationship that the gene of hydrogenase enzymes are regulated by several environmental signals. The effectors include the availability of Fe^{2+} [32], Ni^{2+} [33], Se^{4+} [18], and electron acceptors such as oxygen [31].

The expression of Nickel-Iron hydrogenase gene in *Desulfovibrio spp*; Hydrogen metabolism is a central bioenergetic pathway in *Desulfovibrio spp*. that involves hydrogen consumption and production under both sulfate-reducing and fermentative conditions. Different classes of hydrogenases, viz., [NiFe], [NiFeSe], and [Fe] hydrogenases, have been identified within the genus [34].

Their genes are not uniformly found among *Desulfovibrio strains* [35]. Complexity of the hydrogen metabolism and the hydrogenase system in *Desulfovibrio spp*. makes it difficult for the gene to be expressed. The results of the study finally indicate that the transcription of the [NiFe] hydrogenase gene is correlated with bacterial growth and therefore with the metabolic activity, i.e., sulfate reduction rate, in various *Desulfovibrio spp*. Thus, the detection of the [NiFe] hydrogenase mRNA appears to be a suitable tool with which to determine metabolically active sample in environmental population. The [NiFe] hydrogenase plays an a crucial role in energy metabolism of *Desulfovibrio spp*. [36]. Therefore, its expression should be an excellent marker for the metabolic activity of those bacteria in the environment

2.4. Expression of the [NiFe] Hydrogenase Gene in Pure Cultures (Bioreactor System)

According to the experiment of [10], the identification of [NiFe] hydrogenase gene fragments and mRNA in *Desulfovibrio spp*. were investigated and described below in anaerobic methanogenic bioreactor. In parallel, DNA and RNA were reported to be isolated and used as templates for PCR and RT-PCR, respectively. The PCR products were analyzed by DGGE to visualize the individual members of the *Desulfovibrio* populations. As previously demonstrated by [37] experiments.

[NiFe] hydrogenase mRNA was detected by RT-PCR in various cell of *Desulfovibrio spp* growing exponentially in a medium where hydrogen or lactate is used as an electron donor in the presence of sulphate. However, no growth of *D. desulfuricans* DSM 1926 has been reported with hydrogen as an energy source, as was also demonstrated by Devereux *et al.* [38]. Thus, mRNA was not analyzed. In addition, *D. vulgaris* DSM 644, *D. desulfuricans* DSM 1926, and *D. baculatus* DSM 2555 were grown in lactate-sulfate medium. Growth, sulfide production, and the expression of the [NiFe] hydrogenase gene were monitored until the cultures reached the stationary phase. They described that [NiFe] hydrogenase mRNA was detectable in exponentially growing cells at 5 to 30 h after inoculation,

as determined by RT-PCR amplification of a ca. 2440-bp

According to the results of [36], DGGE analysis of amplified [NiFe] hydrogenase gene fragments allows for the differentiation of the various *Desulfovibrio* strains due to their different migration behaviour caused by sequence variations within their PCR products. [NiFe] hydrogenase gene fragments were amplified from DNA isolated from a sample taken directly out of the reactor. Positive results were also reported to be found for DNA obtained from subsamples that were incubated with hydrogen to induce gene expression, respectively. The corresponding mRNA was detectable in only the bioreactor subsample which was incubated with hydrogen. Agarose gel electrophoresis of the PCR products revealed that the sizes of all amplified fragments were as expected for the [NiFe] hydrogenase of *Desulfovibrio spp*. DGGE analysis of the amplified gene fragments showed several bands with identical electrophoresis patterns. However, their results showed that the [NiFe] hydrogenase gene is present in all *Desulfovibrio* strains tested so far in *Desulfovibrio spp*. Which has a single hydrogenase gene, the [NiFe] hydrogenase is responsible for all the aspects of hydrogen metabolism [10]

Thus the Members of *Desulfovibrio spp*. containing different numbers of hydrogenase genes, including the following: *D. vulgaris* DSM 644, which possesses at least three different hydrogenase genes; *D. desulfuricans* DSM 1926, containing two hydrogenase genes; and *D. gigas*, which contains only the [NiFe] hydrogenase gene [35] have been analysed and the transcription of the [NiFe] hydrogenase gene were functionally demonstrated in all strains growing in hydrogen-sulfate and lactate-sulfate media.

3. Conclusion

The versatility of metabolism experience by sulfate-reducing bacteria of the genus *Desulfovibrio* has established that Hydrogenases are the key enzymes involved in the hydrogen production/consumption [39]. The three most widely researched species of SRB (genus *Desulfovibrio*) usually show different behaviour in all traits in terms of hydrogen evolution. Differences in the type, localization, specific activity and number of hydrogenase, is responsible for the metabolic performances of the enzymes towards electron donors and acceptors. In fact *D. vulgaris* Hildenborough shows an enhanced hydrogen yield, which is due to the higher specific activity of the [Fe] hydrogenase as compared with [NiFe] and [NiFeSe] hydrogenases in *D. gigas* and *D. desulfuricans*, respectively [10]. It is clear however that all the reducing power of the metabolic chain is driven for hydrogen production vis -avis indicating the existence of hydrogen cycling in SRB [1]. The ability of *Desulfovibrio* species to handle hydrogen in a reversible mode shows that these micro-organisms have a high capacity to quickly acclimatize to their system of metabolism despite the variation of environmental changes. Hydrogenase activity

can provide an important resource for cells when they grow in environments containing scarce electron donors or acceptors. Random and site directed mutagenesis can be done to produce other hydrogenases of high stable function for use in different biotechnological applications. However this SRB require to be treated with biological curiosity as a diversification tool to investigate other factors that could trigger [NiFe] hydrogenase efficiency in SRB.

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