NifH: Structural and Mechanistic Similarities with Proteins Involved in Diverse Biological Processes

Surobhi Lahiri, Lakshmi Pulakat and Nara Gavini
Department of Biological Sciences,
Mississippi State University, Mississippi State, MS 39762, USA

Abstract: The NifH protein is a subunit of the nitrogenase enzyme that catalyzes the reduction of atmospheric nitrogen to ammonia. This protein contains highly conserved regions including the nucleotide binding sites, metal center ligands and the Switch I and Switch II domains. A number of proteins have structural and mechanistic similarities as well as evolutionary relationships with the NifH protein, notable among them being: light independent protochlorophyllide (Pchlide) reductase (ChlL/FrxC or bChL), arsenite pump ATPase (ArsA), 2-hydroxyglutarate dehydratase Component A (CompA) involved in glutamate degradation and MinD that functions in spatial regulation of cell division. Although involved in very diverse biological processes, these proteins share an underlying common structural framework. This review mainly focuses on the structural similarities of these proteins with the NifH protein and discusses recent reports of complementation studies involving NifH and few of the proteins mentioned.

Key words: NifH, ChlL, ArsA, CompA, MinD, domain conservation, complementation

INTRODUCTION

The nitrogenase enzyme is a two-component system that consists of the iron protein (Fe-protein or NifH) and molybdenum-iron protein (MoFe-protein) working in concert to effect nitrogen reduction\[1-4\]. The 64 kDa homodimeric NifH protein has two ATP binding domains and one \[4Fe-4S\] cluster per homodimer. It supplies energy by ATP hydrolysis and transfers electrons from reduced ferredoxin or flavodoxin to the MoFe-protein\[3\]. It can bind up to two MgADP and/or MgATP molecules per dimer and is essential for coupling nucleotide hydrolysis to electron transfer for eventual nitrogen reduction\[5,6\]. It also participates in the biosynthesis and insertion of the FeMo-cofactor into the MoFe protein\[7-10\]. In the past, various proteins have been revealed to have structural and mechanistic similarities as well as evolutionary relationships with the NifH protein, notable among them being: light independent protochlorophyllide (Pchlide) reductase (ChlL/FrxC or bChL), arsenite pump ATPase (ArsA), 2-hydroxyglutarate dehydratase Component A (CompA) involved in glutamate degradation and MinD that functions in spatial regulation of cell division\[11-16\]. Although involved in diverse biological processes, these proteins have been found to bear considerable structural resemblance to the NifH protein. Mechanistically, they do play a role similar to NifH in their respective complexes, based upon the presence of ATP binding motifs in each of them\[17-21\]. Whereas ChlL, ArsA and MinD belong to the same superfamily of proteins as NifH, known as the P-loop containing nucleoside triphosphate hydrolases, the Component A of 2-hydroxyglutarate dehydratase (CompA) is a member of the Actin-like ATPase domain superfamily, as designated by the SCOP database\[22\]. An analogy between NifH and ChlL, ArsA, CompA and MinD lies in the organization of these proteins as members of their respective two component systems. In the nitrogenase complex, NifH functions as the obligate electron donor to its specific partner, the MoFe protein. This intermolecular electron transfer process requires ATP hydrolysis. In a similar manner, for the dehydratase system, CompA is the site of ATP hydrolysis and is the electron donor to the sec component, CompD\[19\]. ArsA exists in a complex with the ArsB protein forming an arsenite-antimonite \([\text{As(III)}/\text{Sb(III)}]\)-translocating ATPase. As a result of the ATP hydrolysis in ArsA, a more compact conformation of the enzyme is generated, allowing the vectorial movement of the arsenite or antimonite ions into ArsB\[13\]. It has been speculated that the MinD-MinE membrane complex is analogous to the NifH-MoFe complex and results in ATP hydrolysis, as a
from the membrane [11]; Comparison of the molecular consequence of which, MinD and MinE are released

Fig. 1: Protein sequence alignment of NifH, ChlL, ArsA1, ArsA2, MinD and CompA. The residues are colored as red, blue, magenta, green and gray based on their hydrophobicity and other properties. Regions highlighted are: the nucleotide binding domains in gray; the Switch I in yellow; the Switch II in cyan. Cysteine residues highlighted in red are the ligands for the metal centers of their respective proteins. The histidine residue highlighted in red is one of the ligands for the Sh(III) center of ArsA. It should be noted that ArsA2 sequence includes the 'linker' stretch of residues of the ArsA protein (Cr: Chlamydomonas reinhardtii, Av: Azotobacter vinelandii, Ec: Escherichia coli, Af: Acidaminococcus fermentas)

consequence of which, MinD and MinE are released from the membrane[11]; Comparison of the molecular architecture between nitrogenase and the light-independent Pchlide reductase has shown that the NifH counterpart in the light independent Pchlide reductase is ChlL. Chll is suggested to be ATP-dependent transfer of electrons from a reductant, such as ferredoxin, to the ChlB-ChlN complex via the Fe:S center[14]. Thus, in a manner similar to NifH, the proteins ArsA, ChlL, CompA and MinD also couple hydrolysis of nucleoside triphosphates to redox reactions from a metallic cluster.

The protein sequence comparison of NifH, ArsA, ChlL, CompA and MinD gives an overview of their similar regions (Fig. 1). However, it is difficult to detect protein relationships and the presence of different domains by direct comparisons of sequences belonging to different protein families and it is more reasonable to determine the presence or absence of domains and their family relationships by comparison of their three-dimensional structures (Fig. 2). How proteins with similar protein folding patterns perform varied functions is a prevalent question and it has been suggested that popular folding patterns have thermodynamic advantages enabling them to be stabilized by random sequences and these few advantageous folds can probably tolerate various primary structures and therefore perform different functions[23]. The structural comparison of each of the four proteins with NifH creates a general model that indicates their common ancestry at some point. The functions of NifH, ArsA and ChlL, which are broadly nitrogen fixation, arsenite resistance and photosynthesis respectively, further point out to the very first evolutionarily important processes on earth. It has been speculated that the global functions of the NifH homologs may have varied with either change in nucleotide specificity, the incorporation of new functions or the loss of others[24]. A thorough analysis of the comparative structural details of these proteins allows a better understanding of their relationship to each other. Consequently, derived from the comparison
of NifH with ChlL, ArsA, CompA and MinD, greater insight into their structure based function and phylogeny can be obtained.

**NifH: THE STRUCTURAL TEMPLATE**

The 2.9 Å crystal structure of the NifH protein from *Azotobacter vinelandii* was obtained by Georgiadis *et al.* in 1992\(^5\). It revealed that each of the subunits of the NifH dimer consisted of mixed \(\alpha\)-helix/\(\beta\)-sheet polypeptide fold, with a consensus topology of an eight-stranded \(\beta\)-sheet flanked by nine \(\alpha\)-helices. A \(\beta\)-sheet core in each monomer was formed of one short antiparallel and seven parallel \(\beta\)-strands. The binding sites critical to NifH function were determined by loops at the carboxy-terminal ends of several \(\beta\)-strands. The Nucleotide Binding Sites (NBSs) were situated in the loops following \(\beta\)1 and \(\beta\)2 and the cysteine ligands (C 97 and 132) for the [4Fe-4S] cluster were in the loops following \(\beta\)5 and \(\beta\)6\(^{26,25}\). Both cluster ligands are located near the amino terminal ends of \(\alpha\)-helices that are directed toward the cluster. As indicated by spectroscopic studies, a prominent feature of the Fe protein is the exposure of the [4Fe-4S] cluster to the solvent\(^{26}\), which is also probably one of the reasons that it is an oxygen sensitive protein. In addition to the [4Fe-4S] cluster, there are numerous Van der Waals and polar interactions in the interface beneath the cluster that help stabilize the dimer structure.

NifH displays structural similarity to other nucleotide-binding proteins including signal transduction molecules such as G-proteins and ras and energy transduction systems such as myosin\(^{27}\). It is known to be similar to the nucleotide binding proteins based on the presence of the following structural features\(^{28,29,30}\): (i) mainly parallel \(\beta\)-sheets flanked by \(\alpha\)-helices (ii) a phosphate-binding loop (P-loop) or Walker A motif\(^{31}\), containing the G-X- X- X- X-G-K-\(\gamma\) consensus sequence and (iii) two switch regions, Switch I and Switch II that interact with the \(\gamma\)-phosphate group of the bound nucleoside triphosphate. Analysis of the position of a bound ADP molecule in the X-ray structure of the Fe protein from *Azotobacter vinelandii*\(^5\) and the properties of other site-specifically altered Fe proteins\(^{8,27,32-35}\), has established the position of the nucleotide binding sites, one on each subunit, 19 Å away from the [4Fe-4S] cluster. The nucleotide-dependent switch regions are responsible for communication between the sites responsible for nucleotide binding and hydrolysis and the [4Fe-4S] cluster of the Fe protein and the docking interface that interacts with the MoFe protein upon macromolecular complex formation. In *A. vinelandii* NifH, residues 38 to 43 and 125 to 132 form the Switch I and Switch II respectively. The switch regions undergo conformational changes upon hydrolysis to the nucleoside diphosphate with consequent loss of the interaction with the terminal phosphate group. Substitutions in the nucleotide dependent switch regions of the NifH of *Azotobacter vinelandii* showed that the altered NifH proteins formed a trapped complex subsequent to a single electron transfer event\(^{36}\). The studies suggested that whereas in the structure of the native enzyme the analogous interaction between the side chains of D39 and 125 was precluded due to electrostatic repulsion, the D39N substitution allowed the formation of a hydrogen bond between the Switch I D39 and the Switch II D125; this demonstrated that the electrostatic repulsion between D39 and D125 was important for dissociation of the Fe protein: MoFe protein complex during catalysis\(^{36}\). Thus, the switch regions play a critical role in transmitting information concerning the nucleotide state to other effector molecules that bind to these regions.

Investigations of the dimer interface of the NifH protein have shown that in the NifH dimer, residues with >30Å of buried surface area include K 41, E 92, P 93, V 95, A 98, D 129, V 130, V 131, C 132, M 156, Y 159, K 166 and K 170. These residues, along with other residues adjacent in the sequence, mediate the subunit-subunit interactions through a series of primarily polar (hydrogen bond and salt-bridge) interactions\(^{24}\). The C-terminus of NifH has been speculated to provide an additional degree of intersubunit interaction in NifH as residues from this region wrap around the body of the opposing subunit and enhance the overall stabilization of the NifH dimer. Also, in the C-terminus region, cross-subunit salt-bridges have been known to form between K 224-E 277 and K 233- E 287\(^{24}\).

An integral part of the nitrogenase mechanism is MgATP binding to the Fe protein and hydrolysis by the Fe protein-MoFe protein complex. Through the studies of Jang *et al.*, it was confirmed that the protein interactions with the Mg\(^{2+}\) were essential for the transduction of the nucleotide hydrolysis event\(^{37}\). Analysis of the MgADP bound crystallized Fe protein revealed that S16 binds the Mg\(^{2+}\) and \(\alpha\), \(\beta\) and \(\delta\) phosphates of nucleotides and appears to form a hydrogen bond with D125 of the switch II region\(^{27,37}\). Yet another region found to be critical for the MgATP-induced conformational change was the highly conserved span of the Fe protein around A157 located at the subunit interface. This region is part of the \(\alpha\)5
helix extending from residue 151 to 176 at the subunit interface and shows substantial difference when the free Fe protein structure is compared with the structure of the Fe protein in the MgATP bound complex. Moreover, the mutation of A157 to Ser resulted in a protein that could still bind MgATP normally but was unable to undergo the MgATP-induced conformational change.[8] Further insight into MgATP binding and hydrolysis was obtained by studies based on residues K15 and D125. It was suggested that the breaking of a salt bridge between these two residues by MgATP binding was responsible for triggering a conformational change.[33,34].

The substrate reduction mechanism of the nitrogenase enzyme involves the key step of MoFe protein docking on the Fe protein. Recent studies dealing with the biochemical and structural characterization of the crosslinked complex of nitrogenase have shown the specificity of the residues involved in the transient complex formation between the Fe and MoFe proteins. It was found that only E112 from one of the two NifH subunits and K400 of the NifK subunit were crosslinked, although these residues were surrounded by numerous other charged residues that could potentially participate in this process[17]. Most recently, when the Fe and MoFe proteins were cocrystallized under the conditions of (i) no nucleotide (ii) MgADP and (iii) MgAMPPCP (an MgATP analog), the Fe protein molecules were found to be capable of occupying different interaction sites on the MoFe protein[38]; three separate docking areas were thereby identified on the surface of the MoFe protein[38]. Other studies that substituted Ala in place of F125 of the α and β subunits of the MoFe protein, separately and in combination, showed that the doubly substituted MoFe protein was unable to form a tight complex with the MgADP-AlF4− treated NifH or when using the altered L127ΔNifH, thereby suggesting that the F125 residues were involved in an early event(s) that occurred upon component protein docking and could be involved in eliciting MgATP hydrolysis[39].

Amongst the proteins that share a high level of structural-functional similarity and sequence identity/similarity with NifH Table 1 and 2, ChlL bears maximum closeness to this protein. However, the ChlL crystallographic structure is not yet available and several predictions of its structural features have been made considering its likeness to NifH[14], Table 1.

**ChlL: LIGHT-INDEPENDENT CHLOROPHYLL BIOSYNTHESIS**

The reduction of the C17-C18 double bond of the D-ring of protochlorophyllide (PChlide) to chlorophyllide (Chlide) is a major step in the chlorophyll biosynthesis pathway. The two pathways for this process involve catalysis by the enzymes NADPH : PChlide oxidoreductase (POR) and the light-independent PChlide reductase (DPOR).

The light-independent PChlide reductase plays an important role in the ability of anoxygenic photosynthetic bacteria, cyanobacteria, nonvascular plants, ferns and gymnosperms to form chlorophyll in

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**Table 1: Structural-functional features of NifH, ChlL, MinD, ArsA and CompA**

|                  | NifH | ChlL | MinD | ArsA | CompA |
|------------------|------|------|------|------|-------|
| Localization     | Cytoplasmic | Chloroplast | Peripheral membrane | Extrinsic membrane protein | Cytoplasm |
| O2 sensitivity   | Sensitive | Sensitive | Insensitive | Insensitive | Sensitive |
| Molecular weight | –64 kDa | –64 kDa (?) | –35 kDa (monomer) | –63 kDa | –54 kDa |
| Metal center     | 4Fe4S | 4Fe4S (?) | -- | ShIII/AsIII | 4Fe4S |
| Folding pattern  | 3 layers: α/β/α parallel or mixed | 3 layers: α/β/α parallel or mixed | 3 layers: α/β/α parallel or mixed | 3 layers: α/β/α parallel or mixed | 3 layers: α/β/α mixed-beta-sheets of 5 strands, strand 2 is antiparallel to others |
|                  | beta-sheets of variable sizes | beta-sheets of variable sizes (?) | beta-sheets of variable sizes | beta-sheets of variable sizes | |
| Dimerization     | Homodimer | Homodimer (?) | ATP dependent dimerization | Pseudodimer formed by connection of ArsA 1 and ArsA 2 domains through a short linker | Homodimer |
| Metal Center     | C97, C132 | C95, C129 | -- | H148 (A1), S420 (A2) | C127, C166 |
| Ligands          | 10 GKGGIGKSTTT 20 | 7 GKGGIGKSTTS 17 | 9 GKGGVGTKTSS 19 | 15 GKGGIGKSSS 20 (A1) | 7 GIDVG 11 |
|                  | 343 KGGVGKTMTA 344 (A2) | 343 KGGVGKTMTA 344 (A2) | 343 KGGVGKTMTA 344 (A2) | 343 KGGVGKTMTA 344 (A2) | 343 KGGVGKTMTA 344 (A2) |
| Nucleotide binding domain | 10 GKKIADST 45 | 37 DPKHDST 43 | 43 DFDIHLRNL 51 | 45 DPAAG 49 (I-A1) | 104 GQGD 107 |
|                  | 125 DVLGDVV 131 | 122 DVLGDVV 128 | 120 DSPAG 124 | 365 DPAAG 369 (II-A1) | 127 CAAGTG 132 |
Table 2: Percentage identity and percentage similarity scores (shown in parenthesis) between proteins having structural similarity to NifH

|        | NifH | ChlL | MinD | ArsA1 | ArsA2 | CompA |
|--------|------|------|------|-------|-------|-------|
| NifH   | 100% (100%) | 32% (63.8%) | 15.3% (52%) | 17.1% (53.7%) | 15% (48.2%) | 11% (49.6%) |
| ChlL   | 32% (63.8%) | 100% (100%) | 17.7% (56.6%) | 14.3% (47.4%) | 15.6% (47.4%) | 14.7% (49.1%) |
| MinD   | 15.3% (52%) | 17.7% (56.6%) | 100% (100%) | 17.7% (50%) | 18.1% (51.5%) | 11.5% (50.7%) |
| ArsA1  | 17.1% (53.7%) | 14.3% (47.4%) | 17.7% (50%) | 100% (100%) | 25% (61%) | 10% (46.6%) |
| ArsA2  | 15% (48.2%) | 15.6% (47.4%) | 18.1% (51.5%) | 25% (61%) | 100% (100%) | 12.2% (49.6%) |
| CompA  | 11% (49.6%) | 14.7% (49.1%) | 11.5% (50.7%) | 10% (46.6%) | 12.2% (49.6%) | 100% (100%) |

The scores were calculated using the ClustalW alignment program from the San Diego Supercomputer Center (SDSC) server. Identical residues within a pair of aligned proteins were used to compute the percentage identity; identical as well as similar residues were used to determine the percentage similarity.

The accessory gene products required for the mobilization of sulfur and iron for Fe-S cluster formation and for the activity and stabilization of NifH include NifS, NifU and NifM[47,48]. An inspection of the Chlamydomonas genome database for the equivalents of these accessory proteins did reveal their matches, leading to further evidence of a Fe-S cluster in ChlL[46].

Inspite of many genetic and biochemical investigations of DPOR, the lack of a crystallized ChlL protein has deterred a detailed structural insight into the protein. As a step further to elucidate the structural aspects of the ChlL protein, we superimposed the protein sequence derived ChlL onto the NifH template (PDB ID: 1NIP), using the Swiss PDB (Deep View) protein modeling software (Fig. 3). The resultant ChlL protein model generated showed a marked resemblance to the NifH overall architecture and folds. As shown in Fig. 4, upon superimposition of a ChlL monomer on the
NifH monomer structure colored according to differences with the ChlL model. The superimposition of ChlL on NifH indicates very few regions of dissimilarity (shown in red). A root mean square deviation type of coloration of the NifH protein (in comparison to ChlL) shows a particularly dissimilar 'tail' region spanning residues 230-289 of NifH. Each dissimilar region is indicated by the residues it is comprised of. The [4Fe-4S] cluster and the bound MgADP in NifH are indicated.

NifH monomer, there was a wide overall structural alignment, however, the following residues in NifH appeared non-aligned: A1-R3, A53, D117, N142, K143, R187-R191, Y230-G283. Particularly, the tail region of NifH, comprised of residues 230-283 (Fig. 4), differed extensively, since no such region of ChlL could be structurally aligned to it. Such absence of a tail region in ChlL could explain their phylogenetic divergence and possibly account for their functional differences.

Besides ChlL, CompA is another oxygen sensitive protein that has structural similarity to NifH and is a homodimer containing a [4Fe-4S] cluster Table 1. CompA shares ~50% protein sequence similarity with NifH Table 2.

**CompA: GLUTAMATE DEGRADATION**

The hydroxyglutarate pathway, which involves the syn-elimination of water from (R)-2-hydroxyglutaryl-CoA is key to glutamate degradation in *Acidaminococcus fermentas* [49,50]. This important step in the process of glutamate degradation is catalyzed by the enzyme 2-hydroxyglutaryl-CoA dehydratase. This enzyme is the product of the hgdCAB genes and consists of the protein components, component A (CompA) and component B (CompD). CompA is an extremely oxygen-sensitive activator or initiator whereas CompD is moderately oxygen-sensitive and is the actual dehydratase. For activity of the enzyme, both protein components, Mg$^{2+}$, ATP and a strong reducing agent are required. ATP is bound and hydrolyzed by CompA whilst the substrate binds to CompD [51].

The X-ray crystallized structure of recombinant CompA protein from *A. fermentas* was determined at 3 Å resolution by Locher et al. in 2001 [19]. The CompA protein resembles an L shape, with its two arms being roughly similar in length. It has eight α-helices and ten β-strands that form two flat sheets [19]. These two sheets meet at an angle of 60° and the second sheet positions at the corner of the L and therefore is at the center of the protein. Helices 2 and 8, which are almost perpendicular to each other, fill up the large gap between the two β-sheets [19]. The ADP is bound between the two edges of the closely situated β-sheets. The β-phosphate of the ADP is positioned ~19 Å from the [4Fe-4S] cluster that lies at the dimeric interface of CompA [19]. The nucleotide binding pocket in CompA is formed of residues in strands 1 and 6, containing the Gly7-X-Asp-X-Gly11 and the Val100-X-Asp-X-Gly104 motif (Table 1), as found in sugar kinases [52].

Though NifH and CompA are from unrelated phylogenetic families, several structural resemblances exist between them. Like NifH, CompA is a homodimeric enzyme (54 kDa), with one ADP bound to each subunit and a single [4Fe-4S] cluster at its dimeric interface. The [4Fe-4S] cluster in both, CompA and NifH, are solvent exposed (~20 Å$^2$) instead of buried, as observed in other [4Fe-4S] cluster containing proteins [19,53]. Excluding the cluster, the interface between CompA monomers accounts for only ~800 Å$^2$ buried area [19]. From predictions based on protein sequence comparisons and subsequent structural analysis of the X-ray crystallized CompA, the [4Fe-4S] cluster was found to be coordinated by two cysteine residues, C 127 and 166, provided by each monomer [19]. The coordination of the Cys ligands with the [4Fe-4S] cluster in NifH and CompA is shown in Fig. 5A and 5B respectively. Two helices, each from one subunit, pointing with their N-termini towards the [4Fe-4S] cluster and forming a helix-cluster-helix angle of 105°, is a remarkable feature of CompA [54]. Such similar architecture has also been noted in NifH, revealing a helix-cluster-helix angle of 150° [55]. Further, as observed in the NifH complex with MoFe protein in the
presence of ADP-AlF_4^−, the CompA also probably opens to an angle of 180° upon ATP binding. This open structure then resembles an archer shooting arrows, wherein the helix-cluster-helix with two bound ADP could be superimposed on the string of the archer's bow, hence leading to the term archerase for CompA[56].

The Switch II region in NifH includes the Cys132 residue that coordinates the [4Fe-4S] cluster and consequently links the nucleotide and the cluster in NifH[55]. The Switch II region interacts with the γ-

phosphate of the nucleotide and upon hydrolysis changes its conformation to the nucleoside diphosphate state. Thus, in NifH, the regulation of electron transfer by protein conformational change is directed by the nucleotide hydrolysis. In CompA, a structural analogue equivalent to the NifH Switch II region has been mapped to the residues 104 to 107 and 127 to 132, positioned between the ADP and [4Fe-4S] cluster; these segments connect the nucleotide-binding site to the cluster and are most likely involved in coupling ATP hydrolysis to the changes in the cluster-binding region[19].

Based on the evolutionary distances, CompA appears to have diverged from NifH at a much earlier time than ChlL, MinD and ArsA (Fig. 6). MinD and ArsA are both oxygen insensitive and bear highly similar structural folds to NifH (Table 1). Within this group of NifH similar proteins (Table 1), MinD and ArsA are marked by the absence of the [4Fe-4S] cluster. The ArsA protein could be considered to be a dimer of two NifH like subunits ArsA1 and ArsA2; however only a monomeric form of MinD has been available for study by researchers, eventhough there is evidence for ATP dependent dimerization of MinD[12].

MinD: SPATIAL REGULATOR OF CELL DIVISION

The peripheral membrane ATPase MinD, along with MinC and MinE, is a component of the Min system responsible for correct placement of the division site in E. coli cells[58]. MinD helps in blocking unnecessary septation events at the poles by rapidly migrating from one cell pole to the other. The binding
of MinE to MinD induces hydrolysis of ATP and the release of MinD into the cytoplasm\[59\]. The ATP binding cycle induced by MinE results in the rapid movement of MinD from one cell pole to the opposite cell pole, forming alternating broad polar zones\[60\]. It was recently reported that MinC and MinD, which were previously thought to be diffusely associated with the membrane at the poles, were in reality organized into extended spirals that coiled around the cell\[61\]. Although it has been shown that MinD dimerizes in the presence of ATP, only its monomeric form with ADP, AMPPCP or without a nucleotide has been crystallized till now\[18,62,63\]. These crystallized MinD were obtained from three different archaeal species, *Pyrococcus horikoshii* OT3, *Pyrococcus furiosus*, *Archaeoglobus fulgidus* based on the relative ease of their crystallization. The MinD molecule from *P. horikoshii* was found to consist of a β-sheet comprised of 7 parallel and 1 antiparallel strands and 11 peripheral α-helices\[63\]. Though NifH and MinD share only ~16% identical residues (Table 2), the overall folding topology of the two proteins are clearly related to each other\[63\].

Since the dimeric form of crystallized MinD was unavailable, Lutkenhaus and Sundaramoorthy used the NifH structure as a template for modeling a dimeric MinD\[11\]. We attempted a similar structural alignment for this review, using the crystallized MinD (PDB ID: 1HYQ) and NifH (PDB ID: 1NIP) structures as shown in Fig. 7. The superimposition of two MinD monomers (green and blue) on a NifH dimer template (yellow) revealed that other than certain loops at the surface of the two proteins, an additional loop and an α-helix present near the C-terminus in NifH, the superimposition yielded a good fit, similar to the observation of Lutkenhaus and Sundaramoorthy \[11\]. Likewise, Hayashi et al. found that NifH and MinD exhibited similar architectural regions in their ATPase and GTPase domains (β7, β6, β1, β5, β2, β4, β3, the P-loop, α1, α7, α8) yielding 1.2 Å r.m.s.d. value for 81 C α atoms\[68\]. Both NifH and MinD contain the deviant Walker motif A [X-K-G-G-X-X-K-(T/S)]\[21\]. It was however observed that the C97 and C132 ligands of the [4Fe-4S] cluster of NifH were not conserved in *P. horikoshii* MinD. Upon protein sequence comparison of NifH and MinD from *A. fulgidus*, a distinct presence of the Fer4_NifH conserved domain, typically found in [4Fe-4S] iron-sulfur cluster binding proteins and NifH/frxC protein families, was detected in MinD\[64\], spanning residues 133-240 of MinD that correspond to residues 144-251 of NifH.

The ArsA protein that is the catalytic subunit of the As(III)/Sb(III)-translocating ArsAB complex, has long been speculated to be a distant relative of NifH\[13\]. The ArsB subunit serves as a membrane anchor for ArsA and forms the anion-translocating sector of the ArsAB pump\[65\]. The crystallized structure of the ArsA protein\[20\] has revealed many structural details that establish its similarity to the NifH protein. ArsA is a 583-residue polypeptide that has homologous N-terminal (A1) and C-terminal (A2) halves, indicating an evolutionary gene duplication and fusion\[66\]. This 63 kDa ATPase contains two consensus nucleotide-binding motifs, one each in the A1 and A2 halves, indicating an evolutionary gene duplication and fusion\[66\]. The A1 and A2 halves are held together by a short linker peptide\[67\]. ArsA, like NifH exists in a more open conformation in the presence of ADP and contains two NBSs. The two NBSs are each filled with MgADP and lie at the interface between A1 and A2. In each NBS, Mg2+ is octahedrally coordinated by the β-phosphate of ADP, several water molecules and a threonine hydroxyl. When compared to D39 and D129 residues of the NifH protein that play a role in the MgATP hydrolysis, D45 from the ArsA1 NBS directly coordinates with Mg2+ whereas D364 of the ArsA2 NBS coordinates indirectly to it via a water molecule\[20,35,68\].

ArsA is a homodimer in its catalytically active form\[69\] and dimerization is favored by the formation of a three-coordinate complex among three specific

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**Fig. 7:** Comparisons between NifH and MinD. The MinD monomers (blue and green) have been superimposed on a NifH dimer (yellow). The [4Fe-4S] cluster of NifH is shown as a spacefilled body (magenta)

**ArsA: OXYANION TRANSLOCATING ATPASE SUBUNIT**

The ArsA protein that is the catalytic subunit of the As(III)/Sb(III)-translocating ArsAB complex, has long been speculated to be a distant relative of NifH\[13\]. The ArsB subunit serves as a membrane anchor for ArsA and forms the anion-translocating sector of the ArsAB pump\[65\]. The crystallized structure of the ArsA protein\[20\] has revealed many structural details that establish its similarity to the NifH protein. ArsA is a 583-residue polypeptide that has homologous N-terminal (A1) and C-terminal (A2) halves, indicating an evolutionary gene duplication and fusion\[66\]. This 63 kDa ATPase contains two consensus nucleotide-binding motifs, one each in the A1 and A2 halves, indicating an evolutionary gene duplication and fusion\[66\]. The A1 and A2 halves are held together by a short linker peptide\[67\]. ArsA, like NifH exists in a more open conformation in the presence of ADP and contains two NBSs. The two NBSs are each filled with MgADP and lie at the interface between A1 and A2. In each NBS, Mg2+ is octahedrally coordinated by the β-phosphate of ADP, several water molecules and a threonine hydroxyl. When compared to D39 and D129 residues of the NifH protein that play a role in the MgATP hydrolysis, D45 from the ArsA1 NBS directly coordinates with Mg2+ whereas D364 of the ArsA2 NBS coordinates indirectly to it via a water molecule\[20,35,68\].

ArsA is a homodimer in its catalytically active form\[69\] and dimerization is favored by the formation of a three-coordinate complex among three specific
cysteine thiolates (Cys-113, Cys-172 and Cys-442) and the effector, Sb(III) or As(III) \cite{70,71}. Each Sb(III) is coordinated by one A1 residue and one A2 residue. These residues include His148 (A1) and Ser420 (A2); Cys113 (A1) and Cys422 (A2); Cys172 (A1) and His453 (A2). The oxygen insensitivity of ArsA in comparison to NifH could most reasonably be assigned to the difference in the metal center of both these proteins.

ArsA is twice the size of NifH, yet since it consists of two similar domains connected by a short linker, each ArsA monomer could actually be considered to be a pseudodimer, as regarded herein. Information obtained on ArsA (PDB ID: 1IHU) and NifH (PDB ID: 1G5P) protein families from the SCOP database \cite{22} and their comparison based on structural modeling made it evident that the core \(\beta\)-sheet pattern in both proteins, made up of at least 7 parallel \(\beta\)-strands, could fit with minimum root square deviation upon superimposition (Fig. 8A). It was also observed that the Cys97 ligands for the [4Fe-4S] cluster from the two NifH monomers could be aligned to the Cys113 and Cys422 residues of ArsA1 and ArsA2 respectively (Fig. 8B). Moreover, the Cys132 ligands for the [4Fe-4S] cluster of NifH corresponded to the His 148 (A1) and His453 (A2) of the antimony site. The superimposition of the ArsA1 or ArsA2 with a NifH monomer revealed a clear coincidence of the metalloid center of ArsA and the [4Fe-4S] cluster of NifH. Furthermore, the D\(^{142/447}\)TAPTGH\(^{148/453}\) signature sequence of ArsA1 and ArsA2\cite{20} has an exact counterpart in NifH and is believed to correspond to the Switch II region of G-proteins\cite{37}. The comparison of the residues of ArsA1 and ArsA2 corresponding to the region of NifH involved in complex formation with MoFe (NifDK) protein yielded a similar patch of acidic residues. E112 of NifH and K400 of the NifDK protein which specifically crosslink during this process are found within a pocket of other close acidic residues and basic residues respectively\cite{17}. E112 in NifH is surrounded by multiple carboxylic acids, including E68, 71, 73, 110, 111, 112, 116 and D69\cite{17}. Although E112 is highly specific for the crosslinking, it is not a conserved residue among the NifH protein sequences. However, the acidic patch surrounding it is well conserved. The ArsA2 region comprising of residues from L398 to G439, when compared with the region G94 to F135 of NifH, revealed eight glutamate residues, two of which had exact counterparts in NifH and E112 of NifH corresponded to D417 of ArsA2 (Fig. 9A). The buried interface between the A1 and A2 domains of ArsA was found to be relatively small (<10%), compared with the total surface of the enzyme\cite{20}, rendering ArsA as a hollow protein with a large central cavity (Fig. 9B). The hinge region between the ArsA1 ArsA2 polypeptides is formed by a 25-residue linker peptide in the region of 289-313\cite{67}. The structural comparison of the linker region of ArsA by superimposition on NifH indicated the absence of such a stretch in NifH.

Fig. 8: Structural comparisons between ArsA and NifH. (a): Superimposition of ArsA (cyan) on NifH (yellow) shows near coincidence of their metal clusters and a good fit of the \(\beta\)-strand structures. [4Fe-4S] cluster of NifH shown spacefilled in yellow and Sb(III)/As(III) metal center is spacefilled in red (b): A comparison of the metal centers and ligands of NifH and ArsA. Both protein structures are represented as blue ribbons and the metal centers are shown in magenta (spacefill). C97 and 132 for NifH and C113 and C172 for ArsA2 are indicated by arrows. Also C422 implicated as a ligand from ArsA1 is indicated.
Fig. 9: (a): Protein sequence alignment of NifH and ArsA2 in the region spanning of and corresponding to important residues in NifH required for complex formation with the MoFe protein (Gly 94 to Phe 135 in NifH and Leu 398 to Gly 439 in ArsA). An acidic patch can be observed in ArsA, indicated by the presence of eight glutamate residues (highlighted in black). Glu 112 of NifH corresponds to Asp 417 of ArsA2. Asp 417 could be also considered as Asp 99 when ArsA2 is taken as a single unit. (b): The ArsA is a hollow protein with a central cavity as seen in this view. The protein is completely spacefilled and α-helices are in red, β-strands are in orange, coils in gray and part of the metal center seen is in yellow.

Strong evidence for the functional conservation of essential structural domains in NifH and ArsA was recently obtained by substitution of the ArsA2 half of ArsA by NifH, in E. coli[7]. The positive complementation suggested that these proteins were either derived from a common ancestral protein or that one of these was the actual precursor protein from which the other has evolved by divergence. The study also emphasized the importance of the conservation of the group of motifs present in both proteins that allowed NifH to substitute for the function of ArsA2.

EVOLUTIONARY RELATIONSHIPS

The structural similarities between NifH and the proteins ArsA, MinD, CompA and ChlL led to the question that whether all these proteins have diverged from a single closely related ancestor or from one of the proteins amongst this group. Computation of the phylogenetic distances between NifH and these proteins suggested that ChlL and MinD were most closely related to NifH whereas ArsA and CompA were relatively more distant from NifH.

In order to visualize the relationships between these structurally related proteins, a phylogram based on their amino acid sequence alignment was generated (Fig. 6). Due to the functional diversity of the proteins, direct evolutionary interpretations should be avoided but as shown in Fig. 6, there exist three broad divisions that phylogenetically classify the polypeptides NifH, ArsA1, ArsA2, ChlL, MinD and CompA. The phylogenetic tree suggests that NifH and ChlL share maximum similarities and have probably diverged most recently compared to others. It also suggests that both NifH and ChlL could have evolved from a MinD ancestor and that ArsA1, ArsA2 and CompA, though structurally similar to NifH, may have diverged farther away during evolution.

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