

Discovery of Evolutionary Divergence of Biological Nitrogen Fixation and Photosynthesis: Fine Tuning of Biogenesis of the NifH and the ChlL by a Peptidyl-Prolyl *Cis/Trans* Isomerase

^{1,2}Nara Gavini, ¹Sinny Delacroix,
²Kelvin Harris Jr. and ^{1,2,3}Lakshmi Pulakat
¹Department of Biological Sciences,
Bowling Green State University, OH-40342, Bowling Green
²Department of Biological Sciences,
Mississippi State University, MS-39762, Mississippi State
³Department of Internal Medicine,
University of Missouri, MO-65212, Columbia

Abstract: Problem statement: Despite the structural and functional similarities between the nitrogenase that performs biological nitrogen fixation reaction and the Dark Protochlorophyllide Oxidoreductase (DPOR) that performs chlorophyll-biosynthesis, attempts to substitute nitrogenase-components with DPOR-components have hitherto failed. This investigation was undertaken to test if *Chlamydomonas reinhardtii* protochlorophyllide (Pchlde) reductase (ChlL) that shares some structural similarity with Nitrogenase Reductase (NifH) could complement the functions of NifH in biological nitrogen fixation of *Azotobacter vinelandii*. **Approach:** Genetic complementation studies were performed to test if the *chlL* gene and its mutants cloned under transcriptional control of *nifH* promoter (*nifHp*) in a broad-host range low copy plasmid pBG1380 could render a Nif⁺ phenotype to NifH-deficient *A. vinelandii* strains. **Results:** Expression of ChlL could render Nif⁺ phenotype to NifH-deficient *A. vinelandii* only in the absence of NifM, a nif-specific PPIase essential for biogenesis of NifH. The ChlL mutants Cys95Thr and Cys129Thr were unable to substitute for NifH. Thus, the conserved cysteine ligands of [4Fe-4S] cluster in ChlL are essential for successful substitution of NifH by ChlL. Since C-termini of NifH and ChlL demonstrated the least similarity and Pro258, a substrate for the PPIase activity of NifM, is located in the C-terminus of NifH, we posited that replacing the C-terminus of NifH with that of ChlL would render NifM-independence to NifH. The NifH-ChlL chimera could support the growth of NifH- and NifM-deficient *A. vinelandii* in nitrogen limiting conditions implying that it has acquired NifM-independence. **Conclusion/Recommendations:** Collectively, these observations suggest that NifM, an evolutionarily conserved nif-specific PPIase, could have contributed to the functional divergence of biological nitrogen fixation and photosynthesis during evolution by virtue of its ability to exert opposing effects on structurally similar substrates, ChlL and NifH.

Key words: Nitrogenase, NifH, ChlL, Dark Protochlorophyllide Oxidoreductase (DPOR), NifM, PPIase

INTRODUCTION

Functional divergence of biological nitrogen fixation and photosynthesis, the two fundamental biological processes that sustain life on earth, is still an enigma. Structural and functional similarities exist between nitrogenase that performs the biological nitrogen fixation reaction and Dark Protochlorophyllide Oxidoreductase (DPOR) that performs reduction of protochlorophyllide to

chlorophyllide during chlorophyll-biosynthesis (Brocker *et al.*, 2008; Gavini *et al.*, 2006; Georgiadis *et al.*, 1992; Sarma *et al.*, 2008; Tezcan *et al.*, 2005; Watzlich *et al.*, 2009; Yamamoto *et al.*, 2009; 2008; Yamazaki *et al.*, 2006a; 2006b; Nomata *et al.*, 2006a; 2006b). Both nitrogenase and DPOR are oxygen sensitive two-component systems. However, attempts to substitute nitrogenase components NifH or NifDK by DPOR components have not been successful thus far.

Corresponding Author: Lakshmi Pulakat, Department of Internal Medicine, University of Missouri, MO-65212, Columbia
Tel: 573-814-6000

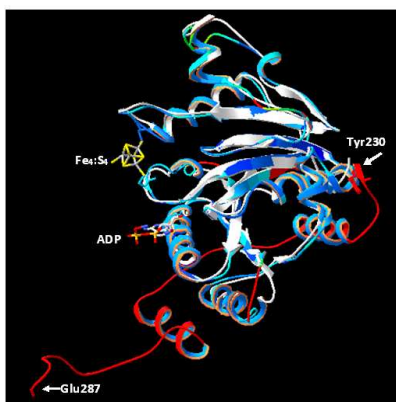


Fig. 3: The NifH monomer structure (based on 1NIP crystallographic structure of the NifH peptide from *A. vinelandii* (Georgiadis *et al.*, 1992) colored according to differences with the ChlL model. Deep View color for ChlL was white. The Deep View coloration of NifH was done according to root mean square deviation type of coloration (blue = maximum similarity; red = maximum dissimilarity) in comparison to ChlL. Bottom view shows a particularly dissimilar C-terminal region (red stretch) starting with Tyr230 and ending with Glu287 of NifH. The [4Fe-4S] cluster and the bound MgADP in NifH are indicated

Since the ChlL protein showed such high structural similarity to the NifH protein, we hypothesized that ChlL would substitute for NifH in *A. vinelandii* and that NifM may play a role in modulating the functional properties of the NifH-like proteins such as ChlL/Bchl. The purpose of this study was to determine whether ChlL could substitute for NifH in biological nitrogen fixation in presence and absence of functional NifM.

MATERIALS AND METHODS

Construction of plasmid pBG2400: An 879bp DNA fragment encoding the ORF of *chlL* flanked by EcoRV and HindIII restriction enzyme sites was generated by PCR amplification using *C. Reinhardtii* chromosomal DNA as template and cloned into the pCR2.1 TOPO vector to create plasmid pBG1382 (Suh, 2002). This fragment was then subcloned into the EcoRV and HindIII digested broad-host range expression vector pBG1380 that contained the *nifHp* (Gavini *et al.*, 2006) to generate pBG2400. Thus, the plasmid pBG2400 had the *chlL* gene under the transcriptional regulation of the *nifHp*.

Construction of ChlL mutants: PCR-mutagenesis with 'QuikChange™ Site Directed Mutagenesis Kit'

(Stratagene Products, La Jolla, CA) was used to generate these mutants according to manufacturer's protocol. Plasmid pBG2400 was used as template. To generate Cys95Thr amino acid replacement, the forward primer was: 5'-CCACCTGCCGGTGC GG GACCGGTGGTTATGTTG TAGGT-3' and reverse primer was 5'-ACCTACAACATAACCACCGGTCCCCGCACCGGC AGGTGG-3'. To generate Cys129Thr amino acid replacement, the forward primer was: 5'-GTTTTAGGTGATGTTGTTACCGGTGGCTTTGCTG CTCCA-3' and reverse primer was 5'-TGGAGCAGCAAAGCCACCGGTAACAACATCACC TAAAAC-3'.

Construction of NifH-ChlL chimera: The 873bp DNA fragment encoding *nifH* ORF was PCR amplified using pDB6 (Jacobson *et al.*, 1989b) as the template and initially cloned into the pCR2.1 TOPO vector. The EcoRV- HindIII fragment encoding *nifH* ORF was subcloned into EcoRV- HindIII digested pBG1380 to generate pBG2434 that carries *nifH* gene under the transcriptional control of *nifH* promoter (*nifHp*). The 129bp region that carries the last 42 amino acids at the C-terminus of NifH was removed via Sall digestion. Next, the DNA containing the last 55 codons of the *chlL* was PCR-amplified using a 5' primer that carries a Sall site (5'-GTTCGACAATTCTACAGTAGGAGTGTC-3') and a 3' primer with a HindIII site (5'-AAGCTTTTAAATTTAAGATAGAAATC-3'). The resultant PCR product encoding the C-terminal region of ChlL protein (55 amino acids at the C-terminal end) was cloned into the Sall-HindIII digested pBG2434 (carrying the N-terminus of *nifH*) to generate a *nifH-chlL* chimeric gene in which the DNA encoding the C-terminal region of NifH (bp745-873) was replaced by the DNA encoding C-terminal region of ChlL (bp718-882).

RESULTS AND DISCUSSION

ChlL can substitute for NifH in biological nitrogen fixation reaction only in the absence of NifM: Two NifH-deficient *A. vinelandii* strains, one NifM-positive (*nifM*⁺ *A. vinelandii* DJ54 (Gavini *et al.*, 1994; Robinson *et al.*, 1987) and one NifM-negative (*nifM*::*kan* *A. vinelandii* BG98 (Gavini *et al.*, 2006) respectively, were used to test the ability of the ChlL to substitute for the NifH in nitrogen fixation reaction by *A. vinelandii*. Both strains were transformed with pBG2400 that carries the *C. reinhardtii chlL* gene cloned under the transcriptional regulation of the *nifHp* and the ability of the transformants to grow under nitrogen limiting conditions was assessed as follows.

Table 1: Replacing the cysteine ligands Cys95 and Cys129 with Thr inhibits the ability of the ChIL to substitute for the NifH in NifH-deficient NifM-minus *A. vinelandii* BG98

Strain	Plasmid	Codon/amino acid for position 95 of ChIL	Codon/ amino acid for position 129 of ChIL	Growth on BN ⁺ medium with Chloramphenicol	Growth on BN ⁻ medium
<i>A. vinelandii</i> DJ54 ($\Delta nifH$) NifM ⁺	pBG2400	TGT/Cys	TGT/Cys	+++	---
<i>A. vinelandii</i> BG98 <i>nifM</i> ⁻ NifH-deficient	pBG2400	TGT/Cys	TGT/Cys	+++	+++
<i>A. vinelandii</i> BG98 <i>nifM</i> ⁻ NifH-deficient	pBG2406	ACC/Thr	TGT/Cys	+++	---
<i>A. vinelandii</i> BG98 <i>nifM</i> ⁻ NifH-deficient	pBG2428	TGT/Cys	ACC/Thr	+++	---
<i>A. vinelandii</i> DJ54 ($\Delta nifH$) NifM ⁺	pBG2406	ACC/Thr	TGT/Cys	+++	---
<i>A. vinelandii</i> DJ54 ($\Delta nifH$) NifM ⁺	pBG2428	TGT/Cys	ACC/Thr	+++	---

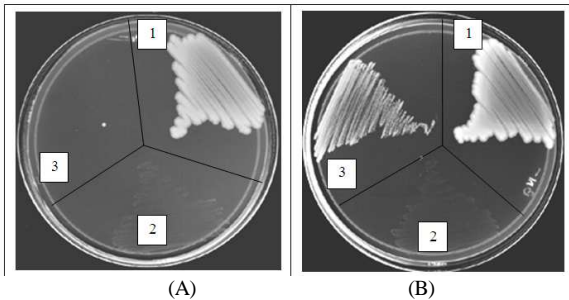


Fig. 4: Comparative growth of NifH-deficient *A. vinelandii* strains carrying plasmids with or without *chlL* gene on Burk's nitrogen free (BN⁻) agar plate is shown. A) *nifM*⁺ $\Delta nifH$ *A. vinelandii* DJ54 carrying pBG1380 that does not harbor *chlL* (marked 2) or pBG2400 that harbors *chlL* (marked 3); B) *nifM*⁻ (*nifM* gene interrupted by insertion of *Kan*^R) *A. vinelandii* BG98 that expresses a non-functional NifH and harboring pBG1380 (marked 2) or pBG2400 (marked 3). Wild type *A. vinelandii* was used as control (marked as 1) on both plates. The pBG1380 is a parental plasmid that harbors the *nifH* promoter^[2]. The pBG2400 carries the *chlL* under the transcriptional control of the *nifHp* of pBG1380, therefore, the *chlL* is expressed under nitrogen limiting conditions. The presence of pBG2400 allowed the growth of NifH-deficient *nifM*⁻:*kan* *A. vinelandii* BG98, on BN⁻ agar plates (B 3), while it did not support the growth of NifH-deficient *nifM*⁺ *A. vinelandii* DJ54 on BN⁻ agar (A 3). Transformations were repeated at least six times

Since the plasmid carries a chloramphenicol resistance (*Cm*^R) marker, the transformants were originally selected on Burk's Nitrogen plus (BN⁺) (Strandberg and Wilson, 1968) medium supplemented with 72 $\mu\text{g mL}^{-1}$ of chloramphenicol. Then, the ability of the *chlL* gene to render Nif⁺ phenotype to *A. vinelandii* strains DJ54 and BG98 was monitored by growing the transformants on Burk's nitrogen free (BN⁻) media containing molybdenum

(Strandberg and Wilson, 1968) that imits expression of alternate nitrogenases (Betancourt *et al.*, 2008).

It was found that pBG2400 could not render a Nif⁺ phenotype to the *A. vinelandii* DJ54 (Fig. 4a), but it rendered Nif⁺ phenotype to the *A. vinelandii* BG98 (Fig. 4b).

Since the difference between the two strains is that DJ54 has an intact *nifM*, whereas BG98 has a disrupted *nifM*, we concluded that the ChIL could restore nitrogenase activity in the absence of functional NifH- but only if NifM was also absent. In summary, the ChIL can replace the NifH-function in biological nitrogen fixation by *A. vinelandii*. However, the NifM, the accessory PPIase essential for biogenesis of functional NifH, has a negative effect on the compensatory ability of ChIL. This interpretation is consistent with the observations that (a) the *chlL* gene could not complement the $\Delta nifH$ of *nifM*⁺ *A. vinelandii* DJ54 and (b) the BchL protein (similar in structure and function to the ChIL) isolated from *nifM*⁺ *A. vinelandii* was unable to substitute for the NifH protein in nitrogenase assay (Sarma *et al.*, 2008).

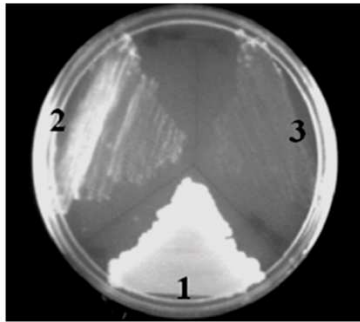
Cys95 and Cys129 of the ChIL are required for substitution of NifH by ChIL in biological nitrogen fixation reaction:

The cysteine ligands Cys97 and Cys132 of the NifH peptide are conserved in the ChIL peptide (Cys95 and Cys129 of the ChIL peptide respectively; Fig. 1). Replacing these conserved cysteines of NifH impairs its function (Howard *et al.*, 1989). We posited that the Cys95 and Cys129 of the ChIL have a similar role in the ability of the ChIL to participate in nitrogen fixation reaction. Therefore we generated ChIL mutants Cys95Thr and Cys129Thr. In both mutants a TGT to ACC conversion was made that resulted in the codon change TGT (Cys) to ACC (Thr) and created a new *PinA1* restriction enzyme site that facilitated identification of these mutants. Locations of the mutations were confirmed by nucleotide sequencing. As shown in Table 1, neither of the mutant *chlL* genes was able to support the growth of *nifM*⁻ NifH-deficient *A. vinelandii* BG98 on BN⁻ medium.

```

P00459 [1-290]nifHA.v.  MAMRQCAYIGGGIGKSTTTQLVAALAEKMKVMIKVCDDPKADSTLILHSAQNTIME 60
NifH-ChlL              MAMRQCAYIGGGIGKSTTTQLVAALAEKMKVMIKVCDDPKADSTLILHSAQNTIME 60
Q00469 [1-293]chlL    ---MLAVYGGIGKSTTTQNLVAALAEKMKVMIKVCDDPKADSTLILHSAQNTIME 57
                        * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * *
P00459 [1-290]nifHA.v.  MAEAEQVDELDLVDLIRAGYGGVVCVSEGGELRQVGCAGRGVITAINFLREBGAEDDL 120
NifH-ChlL              MAEAEQVDELDLVDLIRAGYGGVVCVSEGGELRQVGCAGRGVITAINFLREBGAEDDL 120
Q00469 [1-293]chlL    LSSKDYHYEDLWPEVDIYGGYGGVVCVSEGGELRQVGCAGRGVITAINFLREBGAEDDL 116
                        : : : : : * * * * * : * * * * * : * * * * * : * * * * * : * * * * *
P00459 [1-290]nifHA.v.  DFVFDVLDVVCVGGFAMFIRENKAQEIYVCSGEMMAYANNISKGIYVYANGSGVRL 180
NifH-ChlL              DFVFDVLDVVCVGGFAMFIRENKAQEIYVCSGEMMAYANNISKGIYVYANGSGVRL 180
Q00469 [1-293]chlL    DVILFDVLDVVCVGGFAMFIRENKAQEIYVCSGEMMAYANNISKGIYVYANGSGVRL 174
                        * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * *
P00459 [1-290]nifHA.v.  GGLICNSRNTDEDELLIALANLGTQMIHEVFDNVVQAEIRHMTVIEYDPAKQADE 240
NifH-ChlL              GGLICNSRNTDEDELLIALANLGTQMIHEVFDNVVQAEIRHMTVIEYDPAKQADE 240
Q00469 [1-293]chlL    AGLIGN-RTSKR--DLIDYVEACMPVLEVLIEEIRISRVKGLTFEMSNINMTSA 231
                        : * * * * * : * * * * * : * * * * * : * * * * * : * * * * *
P00459 [1-290]nifHA.v.  YRALARKVNNLLVINPFIHDELELLMEFG-IMEVEDESIVGTAREV----- 290
NifH-ChlL              YRALARKVNNLLVINPFIHDELELLMEFG-IMEVEDESIVGTAREV----- 290
Q00469 [1-293]chlL    H--MDGSGNSGVTSVGETSEYIYCNLYMADQLTSEGVVLELAEKELFLSDP 289
                        * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * *
P00459 [1-290]nifHA.v.  -----
NifH-ChlL              YLR 304
Q00469 [1-293]chlL    YLR 293
    
```

(A)



(B)

Fig. 5: (A) Comparison of the amino acid sequence of the NifH-ChlL chimera with the NifH and the ChlL. Conserved proline residues are marked by the open box. The C-terminal region of the NifH-ChlL chimera that corresponds to that of the ChlL peptide is highlighted in yellow. (B) Growth of the NifH-deficient *nifM* *A. vinelandii* strain BG98 carrying the parental plasmid pBG1380 (marked 2) and pBG1380-derivative expressing the *nifH-chlL* chimeric gene (marked 3) on BN⁻ medium is shown. Wild type *A. vinelandii* (marked 1) served as control. Thus, the NifH-ChlL chimera could support growth of *nifM* *A. vinelandii* strain BG98 in nitrogen limiting conditions indicating that replacement of the C-terminal region of the NifH with that of the ChlL resulted in partial relief from NifM-dependence. Experiments were repeated at least six times

These observations suggested that the Cys95 and Cys129 of the ChlL could play roles analogous to that of Cys97 and Cys132 of NifH in stabilizing the [4Fe-4S] cluster of ChlL.

A *nifH-chlL* chimera could render Nif⁺ phenotype to *A. vinelandii* BG98: As shown in Fig. 1 and 3, the C-termini of the NifH and the ChlL are highly dissimilar. Therefore, the fact that NifH is not functional in the absence of NifM while ChlL is not functional in the

presence of NifM might be traced to this region. We have shown previously that the Pro258 located in the C-terminus of the NifH is one of the substrates for the PPIase activity of NifM (Gavini *et al.*, 2006).

Because the C-terminus of the ChlL is dissimilar to that of the NifH and the ChlL could substitute the NifH in the absence of the NifM in nitrogen fixation, it is conceivable that the C-terminus of the ChlL would render NifM-independence to the NifH. To test this idea, we analyzed the effect of replacing the C-terminal region of NifH with that of ChlL. The DNA encoding the C-terminal region of NifH (bp745-873) was replaced by the DNA encoding C-terminal region of ChlL (bp718-882) to construct the *nifH-chlL* chimeric gene. Therefore, the resulting NifH-ChlL chimera did not contain Pro258 of the NifH. Amino acid sequence of the NifH-ChlL chimera is shown in Fig. 5a. *A. vinelandii* BG98 transformants expressing the *nifH-chlL* chimeric gene were capable of growing on BN⁻ medium (Fig. 5b). Thus, the *nifH-chlL* chimera could render partial NifM-independence to *A. vinelandii* BG98.

CONCLUSION

Our results show that the NifM, a NifH-specific PPIase that is essential for biogenesis of the NifH protein, has a role in disabling structurally similar ChlL from participating in the biological nitrogen fixation reaction. Significance of PPIase-substrate interactions are particularly highlighted in many pathological conditions. For example, overexpression of human Pin1 is implicated in the formation of Lewy bodies in Parkinson's Disease, while the same protein has a beneficial effect in Alzheimer's disease, since it regulates amyloid precursor protein processing and amyloid beta production (Pastorino *et al.*, 2006; Ryo *et al.*, 2006). Similarly, Macrophage Infectivity Potentiators (MIPs) are PPIases expressed by bacterial pathogens, however, they interact with host-cell proteins and alter their functions to establish infection (Kohler *et al.*, 2003). These examples show that the molecular interactions between PPIases and proteins that share structural similarity to their natural substrates result in pathogenesis. The example that has emerged from this study is that of a PPIase which could have contributed to the functional divergence of two fundamental biological processes (nitrogen fixation and photosynthesis) during evolution. This is because this PPIase prototype (NifM) would render functionality to one substrate (NifH) and hinder the function of the other structurally similar substrate (ChlL) so that

nitrogen fixation is favored under conditions that lead to NifM expression (such as nitrogen limitation). These findings represent a unique example of an accessory protein playing a vital part in the evolutionary divergence of biological processes.

The observation that ChlL mutants Cys 95Thr and Cys129Thr were unable to substitute for NifH further extends the structure-function similarity of the NifH and ChlL related to their mechanistic involvement in nitrogen fixation. Although the structure of ChlL is not yet solved, these observations strengthen the similarities in the role of Cys ligands of the (4Fe-4S) cluster of ChlL in electron transfer by the ChlL to that of the NifH. On the other hand, our studies also highlight the dissimilarity of the C-termini of the NifH and the ChlL. The C-terminus of the NifH is involved in the NifM-dependence of the NifH due to the presence of Pro258 (Gavini *et al.*, 2002). In contrast, the C-terminus of the ChlL could render NifM-independence to the NifH as shown by the functional NifH-ChlL chimera (Fig. 5). It is conceivable that a protein similar to the NifH-ChlL chimera could have served as a common ancestor for the NifH and the ChlL before the functional divergence of biological nitrogen fixation and photosynthesis during evolution.

ACKNOWLEDGMENT

We thank members of Gavini/Pulakat laboratories for valuable comments and technical help. Funding for this research was provided by NSF grant # MCB-1041718 to LP.

REFERENCES

- Betancourt, D.A., T.M. Loveless, J.W. Brown and P.E. Bishop, 2008. Characterization of diazotrophs containing Mo-independent nitrogenases, isolated from diverse natural environments. *Appl. Environ. Microbiol.*, 74: 3471-3480. PMID: 18378646
- Brocker, M.J., D. Watzlich, F. Uliczka, M. Saggi and F. Lenzian *et al.*, 2008. Substrate recognition of nitrogenase-like dark operative protochlorophyllide oxidoreductase from *Prochlorococcus marinus*. *J. Biol. Chem.*, 283: 29873-29881. PMID: 18693243
- Chen, L., N. Gavini, H. Tsuruta, D. Eliezer and B.K. Burgess *et al.*, 1994. MgATP-induced conformational changes in the iron protein from *Azotobacter vinelandii*, as studied by small-angle x-ray scattering. *J. Biol. Chem.*, 269: 3290-3294. PMID: 8106367
- Christiansen, J., D.R. Dean and L.C. Seefeldt, 2001. Mechanistic features of Mo-containing Nitrogenase. *Annu. Rev. Plant. Physiology. Plant. Mol. Biol.*, 52: 269-295. PMID: 11337399
- Curatti, L., J.A. Hernandez, R.Y. Igarashi, B. Soboh and D. Zhao *et al.*, 2007. *In vitro* synthesis of the iron-molybdenum cofactor of nitrogenase from iron, sulfur, molybdenum and homocitrate using purified proteins. *Proc. Natl. Acad. Sci. U S A.*, 104: 17626-17631. PMID: 17978192
- Edlich, F. and G. Fischer, 2006. Pharmacological targeting of catalyzed protein folding: the example of peptide bond cis/trans isomerases. *Handb. Exp. Pharmacol.*, 172: 359-404. PMID: 16610367
- Finan, T.M., 2002. Nitrogen Fixation: Global Perspectives. 1st Edn., CABI, Oxford, ISBN: 0851995918, pp: 553.
- Gavini, N. and B.K. Burgess, 1992. FeMo cofactor synthesis by a *nifH* mutant with altered MgATP reactivity. *J. Biol. Chem.*, 267: 21179-21186. PMID: 1400428
- Gavini, N., L. Ma, G. Watt and B.K. Burgess, 1994. Purification and characterization of a FeMo cofactor-deficient MoFe protein. *Biochemistry.*, 33: 11842-11849. PMID: 7918402
- Gavini, N., N.S. Tungtur and L. Pulakat, 2006. Peptidyl-prolyl cis/trans isomerase-independent functional NifH mutant of *Azotobacter vinelandii*. *J. Bacteriol.*, 188: 6020-6025. PMID: 16885471
- Georgiadis, M.M., H. Komiva, P. Chakrabarti, D. Woo and J.J. Kornuc *et al.*, 1992. Crystallographic structure of the nitrogenase iron protein from *Azotobacter vinelandii*. *Science*, 257: 1653-1659. PMID: 1529353
- Howard, J.B. and D.C. Rees, 1996. Structural basis of biological nitrogen fixation. *Chem. Rev.*, 96: 2965-982. PMID: 11848848
- Howard, J.B., R. Davis, B. Moldenhauer, V. L. Cash and D. Dean *et al.*, 1989. Fe: S cluster ligands are the only cysteines required for nitrogenase Fe-protein activities. *J. Biol. Chem.*, 264: 11270-11274. PMID: 2500438
- Howard, K.S., P.A. McLean, F.B. Hansen, P.V. Lemley and K.S. Koblan *et al.*, 1986. *Klebsiella pneumoniae nifM* gene product is required for stabilization and activation of nitrogenase iron protein in *Escherichia coli*. *J. Biol. Chem.*, 261: 772-778. PMID: 3001082
- Jacobson, M.R., K.E. Brigle, L.T. Bennett, R.A. Setterquist and M.S. Wilson *et al.*, 1989a. Physical and genetic map of the major *nif* gene cluster from *Azotobacter vinelandii*. *J. Bacteriol.*, 171: 1017-1027. PMID: 2644218
- Jacobson, M.R., V.L. Cash, M.C. Weiss, N.F. Laird and W. E. Newton *et al.*, 1989b. Biochemical and genetic analysis of the *nifUSVWZM* cluster from *Azotobacter vinelandii*. *Mol. Gen. Genet.*, 219: 49-57. PMID: 2615765

- Kohler, R., J. Fanghanel, B. Konig, E. Lüneberg and M. Frosch *et al.*, 2003. Biochemical and functional analysis of the Mip protein: Influence of the N-terminal half and of peptidyl/prolyl isomerase activity on the virulence of *Legionella pneumophila*. *Infect. Immun.*, 71: 4389-4397. PMID: 12874317
- Lei, S., L. Pulakat and N. Gavini, 1999. Regulated expression of the *nifM* of *Azotobacter vinelandii* in response to molybdenum and vanadium supplements in Burk's nitrogen-free growth medium. *Biochem. Biophys. Res. Commun.*, 264: 186-190. PMID: 10527862
- Lei, S.H., L. Pulakat, K.C. Parker and N. Gavini, 1998. Genetic analysis on the NifW by utilizing the yeast two-hybrid system revealed that the NifW of *Azotobacter vinelandii* interacts with the NifZ to form higher-order complexes. *Biochem. Biophys. Res. Commun.*, 244: 498-504. PMID: 9514861
- Nomata, J., M. Kitashima, K. Inoue and Y. Fujita, 2006a. Nitrogenase Fe protein-like Fe-S cluster is conserved in L-protein (BchL) of dark-operative protochlorophyllide reductase from *Rhodobacter capsulatus*. *FEBS. Lett.*, 580: 6151-6154. PMID: 17064695
- Nomata, J., T. Mizoguchi, H. Tamiaki and Y. Fujita, 2006b. A second nitrogenase-like enzyme for bacteriochlorophyll biosynthesis: reconstitution of chlorophyllide a reductase with purified X-protein (BchX) and YZ-protein (BchY-BchZ) from *Rhodobacter capsulatus*. *J. Biol. Chem.*, 281: 15021-15028. PMID: 16571720
- Pastorino, L., A. Sun, P.J. Lu, X.Z. Zhou and M. Balastik *et al.*, 2006. The prolyl isomerase Pin1 regulates amyloid precursor protein processing and amyloid-beta production. *Nature*, 44: 528-534. PMID: 16554819
- Peters, J.W. and R.K. Szilagy, 2006. Exploring new frontiers of nitrogenase structure and mechanism. *Curr. Opin. Chem. Biol.*, 2: 101-108. PMID: 16510305
- Petrova, N., L. Gigova and P. Venkov, 2002. Dimerization of *Rhizobium meliloti* NifH protein in *Saccharomyces cerevisiae* cells requires simultaneous expression of NifM protein. *Int. J. Biochem. Cell. Biol.*, 34: 33-42. PMID: 11733183
- Rahfeld, J.U., K.P. Rucknagel, B. Schelbert, B. Ludwig and J. Hacker *et al.*, 1994. Confirmation of the existence of a third family among peptidyl-prolyl cis/trans isomerases. Amino acid sequence and recombinant production of parvulin. *FEBS. Lett.*, 352: 180-184. PMID: 7925971
- Robinson, A.C., D.R. Dean and B.K. Burgess, 1987. Iron-molybdenum cofactor biosynthesis in *Azotobacter vinelandii* requires the iron protein of nitrogenase. *J. Biol. Chem.*, 262: 14327-14332. PMID: 3477546
- Rubio, L.M. and P.W. Ludden, 2005. Maturation of nitrogenase: A biochemical puzzle. *J. Bacteriol.*, 187: 405-414. PMID: 15629911
- Ryo, A., T. Togo, T. Nakai, A. Hirai and M. Nishi *et al.*, 2006. Prolyl-isomerase Pin1 accumulates in lewy bodies of parkinson disease and facilitates formation of alpha-synuclein inclusions. *J. Biol. Chem.*, 281: 4117-4125. PMID: 16365047
- Sarma, R., B.M. Barney, T.L. Hamilton, A. Jones and L.C. Seefeldt *et al.*, 2008. Crystal structure of the L protein of *Rhodobacter sphaeroides* light-independent protochlorophyllide reductase with MgADP bound: A homologue of the nitrogenase Fe protein. *Biochemistry*, 47: 13004-13015. PMID: 19006326
- Strandberg, G.W. and P.W. Wilson, 1968. Formation of the nitrogen-fixing enzyme system in *Azotobacter vinelandii*. *Can. J. Microbiol.*, 14: 25-31. PMID: 5644401
- Suh, M.H., 2002. Engineering a simplified functional nitrogenase by gene fusion and DNA shuffling. Doctoral Dissertation, Bowling Green State University.
- Tezcan, F.A., J.T. Kaiser, D. Mustafi, M.Y. Walton and J.B. Howard *et al.*, 2005. Nitrogenase complexes: Multiple docking sites for a nucleotide switch protein. *Science*, 309: 1377-1380. PMID: 16123301
- Watzlich, D., M.J. Brocker, F. Uliczka, M. Ribbe and S. Virus *et al.*, 2009. Chimeric nitrogenase-like enzymes of (bacterio)chlorophyll biosynthesis. *J. Biol. Chem.*, 284: 15530-15540. PMID: 19336405
- Yamamoto, H., J. Nomata and Y. Fujita, 2008. Functional expression of nitrogenase-like protochlorophyllide reductase from *Rhodobacter capsulatus* in *Escherichia coli*. *Photochem. Photobiol. Sci.*, 7: 1238-1242. PMID: 18846289
- Yamamoto, H., S. Kurumiya, R. Ohashi, Y. Fujita, 2009. Oxygen sensitivity of a nitrogenase-like protochlorophyllide reductase from the cyanobacterium *Leptolyngbya boryana*. *Plant. Cell. Physiol.*, 50: 1663-1673. PMID: 19643808

- Yamazaki, S., J. Nomata and Y. Fujita, 2006a. Differential operation of dual protochlorophyllide reductases for chlorophyll biosynthesis in response to environmental oxygen levels in the cyanobacterium *Leptolyngbya boryana*. *Plant. Physiol.*, 142: 911-922. PMID: 17028153
- Yamazaki, S., J. Nomata and Y. Fujita, 2006b. Nitrogenase Fe protein-like Fe-S cluster is conserved in L-protein (BchL) of dark-operative protochlorophyllide reductase from *Rhodobacter capsulatus*. *FEBS. Lett.*, 580: 6151-6154. PMID: 17064695