Purification and characterization of recombinant FAD synthetase from Neurospora crassa

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FAD Synthetase (FADS) [EC 2.7.7.2], the second enzyme in flavin cofactor biosynthetic pathway converts FMN to FAD, plays an important role in many redox reactions. Neurospora crassa FADS (NcFADS) was cloned and overexpressed in E. coli. Recombinant NcFADS was purified in high yields of ~8 mg per liter of bacterial culture using a single step glutathione sepharose affinity chromatography. SDS-PAGE and MALDI-MS revealed that NcFADS has a molecular mass of ~31 kDa. Enzyme kinetic analysis monitored by reverse phase HPLC demonstrate a specific activity and kcat of 1356 nmol/min/mg and 0.69sec⁻¹ respectively. Steady state kinetic analysis of NcFADS exhibited a Km of NcFADS for FMN is 2.7 μM and for MgATP⁻² is 88.7 μM. Isothermal titration calorimetry experiments showed that the recombinant protein binds to the substrates with apparent Km of 20.8 μM for FMN and 16.6 μM for MgATP⁻². Biophysical characterization using intrinsic fluorescence suggests that the enzyme is in folded conformation. Far-UV CD data suggest that the backbone of the enzyme is predominantly in a helical conformation. Differential scanning calorimetry data shows that the enzyme is in folded conformation. The Tm is 53 °C ± 1. This is the first report on cloning, purification and characterization of FADS from N. crassa. The specific activity of NcFADS is the highest than any of the reported FADS from any other source. The results obtained in this study is expected to pave way for intensive research aimed to understand the molecular basis for the extraordinarily high turnover rate of NcFADS.

1. Introduction

Flavoproteins play a crucial role in many biological processes [1]. Flavin mononucleotide (FMN) and Flavin adenine dinucleotide (FAD) derived from the precursor riboflavin, act as coenzymes for various flavoproteins. Therefore, flavin metabolism must be tightly regulated in the cell [2]. Most of the higher eukaryotes cannot synthesize riboflavin de novo so they depend on external dietary sources. Cells need to efficiently acquire them from absorbed food with specific transporter(s) of plasma membrane [3]. Riboflavin kinase (RFK) [ATP: riboflavin 5′phosphoryl transferase (EC2.7.1.26)] catalyzes the first step in biosynthesis of flavin cofactors by ATP-dependent phosphorylation of riboflavin to form FMN. Subsequent catalysis of ATP dependent adenylation of FMN to form FAD is carried out by FAD synthetase (FADS) or FMN adenylyl transferase (FMANAT) [ATP: FMN adenylyl transferase (EC2.7.7.2)] [4,5]. These two reactions follow a sequential bi-bi ordered mechanism [6,7].

In prokaryotes, RFK and FADS are encoded by C-terminal domain in the same gene (ribF or ribG) [8]. The gene product is a single polypeptide in the form of a bifunctional enzyme (RFK/FADS), with C-terminal showing riboflavin kinase activity and N-terminal showing adenyllyl transferase activity. RFK/FADS bifunctional enzyme was purified and characterized from certain bacteria [9–11]. Apart from the mentioned bifunctional enzyme prokaryotic monofunctional RFK’s were also reported [12,13]. In eukaryotes, both RFK and FADS are encoded by two separate genes resulting in two independent monofunctional enzymes [7,14]. Monofunctional enzymes, RFK and FADS were purified and characterized from yeast and rat tissues [4,5,15–17]. Eukaryotic RFK shares sequence similarities with the C-terminus of RFK/FADS bifunctional enzyme of prokaryotes. N-terminus of prokaryotic bifunctional RFK/FADS showed sequence similarity to the proteins of “nucleotidylyl transferase family” unlike the eukaryotic FADS that showed similarity to

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proteins of “PAPS reductase like family” [8,18].

Eukaryotic FADS were cloned and over-expressed from yeasts [19], mammals [5,16,20] and plants [21]. *Saccharomyces cerevisiae* mono-functional FADS was designated as FAD1. The knockout of FAD1 gene was lethal to *S. cerevisiae* indicating the importance of FADS to various flavoproteins involved in key metabolic reactions of the cell [19]. Eukaryotic FADS is present in the cytosol or in the mitochondria. In flavoproteins involved in key metabolic reactions of the cell [19]. Amino acid sequence of putative FADS from *Table 1*. Proteins of *N. crassa* showed a high turnover number and a very low Km for its substrates [32] that might be supporting the life style of the fungi. Therefore, we hypothesize that the sequential enzyme FADS, in flavin biosynthesis pathway, may also be showing these exceptional properties to maintain the flavin content of the cell. Structural studies are the lacuna because of the non-availability of the FADS clone from *N. crassa*. The over expression of the enzyme would pave way for detailed analysis to understand the molecular intricacies responsible for its high turnover number and high specific activity compared to FADS from any other sources. In the present research, we successfully cloned, over expressed and purified the enzyme in high yields that helped in studying the kinetic and preliminary biophysical aspects of the enzyme.

2. Materials and methods

**Materials**: Taq polymerase, glutathione sepharose beads, thrombin, were purchased from Sigma-Aldrich, Saint Louis, USA. Restriction enzymes from New England Biolabs, Inc, USA. pGEX-KG vector was a gift from Dr. Junnuttala’s lab Genentech (Amersham Biosciences, United Kingdom). Qiaquick Gel extraction kit (#28704) was obtained from Qiagen, Hilden. Protease inhibitor cocktail tablets, DNA rapid ligation kit (#K1422) and first stand cDNA synthesis kit (#K1612) were from MBI-Fermentas, USA. SeeBlue-Plus2 pre-stained protein molecular weight standards were purchased from Invitrogen Life Technologies, USA. Other fine chemicals were obtained from Amersham Pharmacia Biotech, USA; Sigma-Aldrich, USA and Gibco-BRL Chemicals, USA. Components of Lucia-Bertianna media were purchased from Himedia, Mumbai. Inorganic salts and solvents used were purchased from Qualigens and Merck, India. All other chemicals used in this study was analytical grade. Glassware was purchased from Borosil, India.

Cloning of FADS from *N. crassa* (FGSC #2489): A coding sequence of FAD synthetase gene from *Neurospora* data base (BROAD MIT, *N. crassa* OR74A (NC10): NCU09223 - FAD synthetase) was identified. *N. crassa* (FGSC #2489) was grown in 1X Vogel’s supplemented medium at 28 °C for 48 h with shaking (100 rpm) in an environmental incubator. The vegetative mycelium obtained was used to isolate total RNA using TRI reagent (#T9424; Sigma, Saint Louis, USA). cDNA was synthesized from the isolated RNA using First Strand synthesis kit (#K1612, MBI-Fermentas, USA). cDNA coding for FADS of *N. crassa* was amplified using gene-specific primers; ncFADS-Forward primer 5′AGCGTGGATCCGGTATCTTATGCTTC3′ (BamHI) and ncFADS-Reverse primer 5′ACGTCGAGCTATCGATCGGACCTG3′ (Xhol). Amplified clone was cloned into pGEX-KG vector which was already pre-digested using the same restriction enzymes. Authenticity of the clone was confirmed by nucleotide sequencing.

Expression of GST-ncFADS fusion protein in *E. coli*: *E. coli* BL21 (DE3) expression host cells were transformed with pGEX-KG-ncFADS. Selection of transformed colonies was performed on Luria broth (LB) agar plates containing ampicillin (100 μg/ml). Transformed bacterial cells with the recombinant pGEX-KG-ncFADS plasmid was inoculated in 25 ml LB containing ampicillin (100 μg/ml) and grown for overnight at 37 °C with rotary shaking (250 rpm). 0.5L of LB was inoculated with overnight culture (2.5%) in the presence of ampicillin (50 μg/ml). Protein over-expression was achieved by induction with 0.6 mM IPTG when the optical density of the culture at 600 nm (A600) reached 0.6. The culture was further incubated at 37 °C for 3 h and the cells were harvested by centrifugation at 6000 rpm (Kubota 7820 RA-6 rotor). Harvested cells were subjected to sonication (3 × 10 s with 10 s pauses; amplitude 50% on Vibra-Cell VCX 750, TX, USA). Expression of ncFADS was checked by SDS-PAGE [33].

Purification of recombinant ncFADS: Glutathione-agarose affinity chromatography was employed for the purification of GST tagged NC-FADS. Cell pellet was resuspended in 1 × PBS (137 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 2 mM KH2PO4), pH 7.5 and subjected to sonication. The supernatant obtained after centrifugation was loaded on glutathione-agarose column (Sigma, USA) that was pre-equilibrated with 1×PBS pH 7.5 and further washed with 5 column volumes of 1×PBS pH 7.5. The bound GST-ncFADS was subjected to on-column cleavage with 1Unit of thrombin for every 200 μg of the fusion protein and incubated at 22 °C for 8 h. The eluates were collected using 1×PBS pH 7.5 followed by concentration of ncFADS using 10/30 kDa cut-off Amicon Ultra protein concentrating filters (Millipore, Bedford, MA, USA) by centrifugation at 3000 rpm (1025 + g). Homogeneity of the protein was assessed using SDS-PAGE (Table 2). ncFADS was obtained in homogeneity with a yield of 8 mg/L of culture. The authenticity of the protein sample was further verified by MALDI-TOF mass analysis. Concentration of the protein was estimated by dye-binding method according to Bradford [34] using BSA as standard.

**Measurement of ncFADS activity**: Enzyme aliquots were incubated with assay mixture (0.1 M Tris.HCl buffer pH 7.5, in the presence of 10 μM FMN, 1 mM ATP with 1 mM MgCl2) at 37 °C for 15 min, in a final volume of 0.1 mL. Reactions were terminated by the addition of formic acid (2 μL of 7 N stock). FMN and FAD were separated on Nucleosil C18 column and samples were analyzed by RP-HPLC (Gibson HPLC model 321 Pump, with online UV/VIS-156 uv–vis spectrophotometric detector and SOMA S-3370 fluorescence detector). Under test conditions 1 pmol of FAD resulted in a peak area of 20.028 units. One unit of enzyme activity was defined as that producing 1 nmol of FAD per min at 37 °C (one milliunit (mU) produces 1 pmol of FAD per min. at 37 °C). The kinetic parameters were calculated from the saturation experiments carried out at pH7.5. For FMN saturation, initial velocity (amount of FAD formed) of the catalytic reaction was determined by incubating fixed amount of ncFADS with saturating levels of MgATP2− (1 mM) and varying the

| Table 1 |
| --- |
| Amino acid sequence of putative FADS from *N. crassa*. |

![Image](image-url)
The concentration of FMN (0.1–10 μM). For MgATP $^2\text{-}\text{saturation}$, initial velocity of the reaction was determined by incubating the fixed amount of NcFADS with varying levels of MgATP (10–1000 μM) in the presence of saturating concentrations of FMN (10 μM).

**Circular Dichroism Spectroscopy:** Spectra were acquired using JASCO-720 spectropolarimeter using 0.1 mm path length quartz cell at room temperature. Far-UV spectra was measured from 200 to 250 nm averaged over 10 scans at a speed of 50 nm/min. NcFADS was diluted to final concentration of 33.3 μM in 1xPBS (pH 7.5). Necessary background corrections were made.

**Steady state fluorescence:** All fluorescence measurements were performed using Hitachi F-250 fluorescence spectrophotometer at 25 °C. The excitation wavelength was fixed at 280 nm and emission spectra were recorded from 300 to 450 nm wavelength, and the bandwidth for excitation and emission were set to 2.5 nm and 10 nm respectively. Spectrum was measured at protein concentration of 3.3 μM in 1xPBS (pH 7.5). Necessary background corrections were made.

**Equilibrium unfolding:** Guanidinium hydrochloride (GdnCl) induced equilibrium unfolding of NcFADS was performed at a protein concentration of 3.3 μM in 1xPBS (pH7.5). The excitation wavelength was fixed at 280 nm and emission maxima was recorded based on the intrinsic tryptophan fluorescence changes with increasing GdnCl concentration, and the bandwidth for excitation and emission were set to 2.5 nm and 10 nm respectively. Necessary background corrections were made.

**Differential Scanning Calorimetry:** Calorimetric scans were performed using CSC 6300 Nano DSC III differential scanning calorimeter (Calorimetry Sciences Corporation). NcFADS with a concentration of 33.3 μM in 1xPBS (pH 7.5) was degassed under vacuum for 10min. Calorimetric scans were conducted at 1 °C/min under a pressure of 3atm. The CSC DSC software was used for the baseline subtraction and determination of the transition temperature of NcFADS.

**Isothermal Calorimetry:** Binding parameters of NcFADS to its substrates FMN and ATP was assessed by measuring the heat changes during the titration using MicroCal™ITC200 (GE Healthcare). Protein and the ligand solutions were degassed under vacuum and equilibrated at 25 °C prior to titration. The sample cell (~220 μL) contained 40–50 μM NcFADS in 1xPBS (pH 7.5). The reference cell contained MilliQ water and the substrates were in the same buffer conditions. Ligands FMN (1.5 mM), ATP (1 mM) with MgCl₂ (5 mM) were titrated with a stirring speed of 1000 rpm. A typical experiment had 30 injections with 1.2 μL each with 120sec equilibration time involved. The resulting titration curves were corrected using protein free buffer controls. $K_\text{d}$, ΔH and N were obtained from Origin ITC software supplied by Micro-Cal™ITC200 (GE Healthcare).

### Table 2

| Purification step       | Total activity (U) | Total protein (mg) | Specific activity (U/mg) | % recovery | Fold enrichment (Sp. Act) |
|-------------------------|-------------------|--------------------|--------------------------|-----------|-------------------------|
| Crude cell lysate       | 862 ± 5.36        | 10.76 ± 0.17       | 80 ± 1.45                | 100       | 1                       |
| Clarified supernatant   | 710 ± 6.35        | 6.46 ± 0.12        | 110 ± 2.51               | 81 ± 0.73 | 1.4                     |
| Thrombin cleaved NcFADS | 502 ± 9.07        | 0.37 ± 0.01        | 1356 ± 5.48              | 57 ± 1.05 | 17                      |

**Note:** The starting volume of culture for each experiment was 50 mL. Values for total activity, specific activity and % recovery have been rounded off to the nearest integer. Data are the Mean ± SE of three triplicate experiments.

### Results and discussion

**Identification and Cloning of the FADS Gene from N. crassa FGSC #2489:** A search of whole genome database of *N. crassa* (FGSC #2489, BROAD Institute, MIT) yielded a putative “FAD Synthetase” gene at locus NCU09233.1 (presently denoted as NCU09233.5) on chromosome I, coding for a hypothetical protein of 272aa (Table 1). The locus has a gene of 952 nt with three exons forming a coding region of 816 nt. This putative NcFADS is a purified from the soluble fraction to homogeneity using single step purification. NGKDKC is the conserved motif at position 71–74 for FADS but not the signal peptide and it shows single form (Fig. 1 A & B), this was a characteristic signature motif observed in all the eukaryotic FADS reported [30]. Expression of this construct resulted in the production of enzymatically active NcFADS protein in transformed *E. coli* expression cells. Total RNA (~2 μg) was isolated from *N. crassa* mycelia. The PCR amplified fragment cloned into pGEX-KG expression vector generated a recombinant plasmid pGEX-KG-NcFADS. The presence of insert was confirmed by double digestion with BamHI.

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**Fig. 1.** (A) Cell lysates obtained before and after induction with IPTG was resolved on 12% SDS PAGE and was stained with Coomassie Blue. 1: lysate of uninduced culture; 2: lysate of induced culture showing an over-expressed band at ~56 kDa; 3: NcFADS recovered after treatment with thrombin. (B) MALDI MS Analysis of NcFADS showed a peak that corresponds to a monoisotopic mass of 30662. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)
Fig. 2. A) Resolution of FAD and FMN on Nucleosil C18 Reverse phase column using at 6.1min and 7.8min respectively; B) Standard graph for FAD (1 pmol of pure FAD = 20.028 peak area).

C) Purified NcFADS (10 ng) was incubated with FMN (10µM), ATP (1mM) and MgSO₄ (1 mM) in Tris.HCl (0.1 M, pH-7.5) at 37°C. Panel C shows pmol of FAD formed as a function of time (0-60min).

D) Purified NcFADS was incubated with FMN (10µM), ATP (1mM) and MgSO₄ (1 mM) in Tris.HCl (0.1 M, pH-7.5) at 37°C for 15min. Panel D shows rate of FAD formed as a function of enzyme concentration (2-20 ng).

E) ATP concentration dependence - Purified NcFADS (10ng) was incubated in Tris.HCl (0.1 M, pH-7.5) with FMN (10µM), ATP (10µM-1000µM) in presence of MgSO₄ (1 mM) at 37°C for 15min. Panel E shows rate of FAD synthesized expressed as pmol of FAD formed per min.

F) Lineweaver Burk plot of NcFADS for ATP dependence.

G) FMN concentration dependence - Purified NcFADS (5µg) was incubated in Tris.HCl (0.1 M, pH-7.5) with ATP (1mM), FMN (0.1µM -10µM) in presence of MgSO₄ (1 mM) at 37°C for 15min. Panel G shows rate of FAD synthesized expressed as pmol of FAD formed per min.

H) Lineweaver Burk plot of NcFADS for FMN dependence.

HPLC buffer used for resolving the peaks contain 0.1 M ammonium formate, 0.1 M Formic acid and 30%methanol.
and XhoI further ascertained by DNA sequencing. This fragment cloned in-frame with the GST coding sequence of the expression vector allowed the expression of a GST-NcFADS fusion protein from the pTac promoter (T7 and Lac promoter) of the vector. pGEX-KG-ncFADS plasmid was used to transform E. coli BL-21(DE3) cells.

**Expression of NcFADS in E. coli:** The recombinant plasmid pGEX-KG-ncFADS encoded the protein NcFADS with the GST tag at the N-terminus. SDS-PAGE of the plasmid containing E. coli BL-21 (DE3) cell cultures induced by IPTG showed an intense band at ~56 kDa corresponding to GST-NcFADS (Fig. 1A). GST tag at the N-terminus facilitated

### Table 3

Comparison of the kinetic parameters.

| Source organism | Km(μM) | Catalytic efficiency (μMsec⁻¹) | k_cat (sec⁻¹) |
|-----------------|--------|---------------------------------|---------------|
|                 | FMN    | MgATP                           | FMN           | MgATP         |
| Nc FADS         | 2.23   | 84.8                            | 0.256         | 0.0077        | 0.69          |
| Yeast           | 0.76   | 10.7                            | NA            | NA            | 0.087         |
| Human           | 0.36   | NA                              | NA            | NA            | 0.0036        |

**Fig. 3.** Structural analysis of recombinant NcFADS was determined using various biophysical techniques such as Far-UV CD, fluorescence spectroscopy, differential scanning calorimetry.

Far UV CD spectra of NcFADS (33μM) was recorded in 1xPBS (pH 7.5) at 25°C showed that the majority of the protein had helical conformation

Intrinsic fluorescence emission of 3.3μM NcFADS showed an emission maximum of 338nm indicating the folded conformation of the protein

GdnCl induced unfolding of NcFADS by steady state fluorescence showed that the protein completely unfolds at 6M GdnCl

DSC profile obtained from 33μM NcFADS in 1xPBS at pH 7.5 showed a transition from the folded to denatured state(s) at 53.3°C

GdnSCN induced unfolding of NcFADS by steady state fluorescence showed that the protein completely unfolds at 3.5M GdnSCN

| Source organism | Km(μM) | Catalytic efficiency (μMsec⁻¹) | k_cat (sec⁻¹) |
|-----------------|--------|---------------------------------|---------------|
|                 | FMN    | MgATP                           | FMN           | MgATP         |
| Nc FADS         | 2.23   | 84.8                            | 0.256         | 0.0077        | 0.69          |
| Yeast           | 0.76   | 10.7                            | NA            | NA            | 0.087         |
| Human           | 0.36   | NA                              | NA            | NA            | 0.0036        |
the purification of recombinant NcFADS using the glutathione agarose affinity chromatography. This protein was purified to homogeneity by a single-step purification procedure. The purified NcFADS migrated as a single intense band of ~31 kDa on SDS-PAGE under reduced conditions (Fig. 1A). MALDI-MS analysis of the purified NcFADS showed a molecular mass of 30.6 kDa (Fig. 1B) that was also in agreement with the theoretical mass of 30.7 kDa. The yield of the protein was ~8 mg per liter of the culture. Such high yields of over-expressed protein would enable us to carry out the structural studies of enzyme.

**Functional characterization of NcFADS:** NcFADS activity was tested using a standard RP-HPLC based enzyme assay. Under standard assay conditions, the rate of production of FAD increased linearly with time up to 60 min (Fig. 2C) and protein concentration (Fig. 2D) up to 20 μg. The purified NcFADS showed a specific activity of 1356 U/mg with 17-fold purification (Table: 2). The steady state kinetic parameters of NcFADS were determined using Lineweaver Burk plot of 1/v (initial turnover number) versus 1/[FMN] or 1/[ATP]. FMN adenylation reaction was shown to follow a sequential ordered bi-bi mechanism in eukaryotes [6, 7]. NcFADS showed a typical hyperbolic saturation curve for both the substrates (Fig. 2E, G). The K_M for FMN, calculated from Lineweaver Burk plots (Fig. 2H) was 2.23 ± 0.2 μM and for MgATP^2^- was found to be 84.8 ± 2 μM (Fig. 2F). NcFADS showed a k_cat value of 0.69 sec^-1. The catalytic efficiency for FMN and MgATP^2^- obtained for NcFADS were 0.256 μM s^-1 and 0.0077 μM s^-1 respectively. As shown in the Table 3, FADS from other eukaryotic sources like yeast showed the K_M values for FMN and ATP as 0.76 μM and 10.7 μM respectively and k_cat value as 0.087 sec^-1 [8]. In humans K_M value for FMN reported was 0.36 μM and the k_cat value was 0.0036 sec^-1 [28]. K_M values for NcFADS were in comparison with other FADS reported as they were all in μM range but the k_cat of NcFADS was ~8-fold high compared to yeast and ~200 times higher than human enzymes. Catalytic efficiency of NcFADS for FMN is 25-fold higher to the human and 2-fold higher to C. glabrata FADS. Earlier reports also indicate the high turnover number of RFC from N. crassa [32]. In our present study NcFADS also showed a very high turnover number compared to other eukaryotic FADS (Table 3). The high turn-over numbers and the catalytic efficiencies of both the enzymes RFC and FADS from N. crassa indicate the requisite of the flavins in this organism that also supports its lifestyle.

**Biophysical characterization of NcFADS:** Structural analysis of recombinant NcFADS was determined using various biophysical techniques such as Far-UV CD, fluorescence spectroscopy, differential scanning calorimetry. Far-UV CD provides valuable information about the gross secondary structure of proteins. Spectrum of NcFADS showed a double minimum at 208 nm and 222 nm (Fig. 3A) indicates that majority of the secondary structure in NcFADS is α-helix. According to SCOP classification eukaryotic FADS belongs to α/β family of proteins. Further confirmation of the secondary structure was done by submitting the NcFADS absorbance values to the K2D3 software [35]. The resulting percentage of α-helix and β-sheet is ~65 and 2 respectively. The secondary structural information from both the reported crystal structures of FADS from C. glabrata (3FWK) and S. cerevisiae (2WSI) [8, 30] showed 40% of the helical content and <5% of total β-content. NcFADS shows ~70% sequence similarity with the reported PDB structures, thus the NcFADS experimental data is also in agreement with the SCOP classification.

Intrinsic tryptophan fluorescence would be an excellent probe to measure the tertiary structure of proteins. Changes in the fluorescence emission of tryptophans in proteins have been widely reported as a consequence of conformational transitions, subunit association, ligand interactions or denaturation. NcFADS has five tryptophan residues at position 87,101,194,206 and 212 of the primary amino-acid sequence (Table 1). NcFADS in its native state showed an emission maxima around 338 nm indicate that the tryptophan residues are buried in the hydrophobic environment of the protein. Guanidinium chloride induced equilibrium unfolding of NcFADS is monitored by the changes in the tryptophan fluorescence, showed that NcFADS completely unfolds at 6M GdnCl. Concentration of the denaturant at which 50% of the molecules remain in the denatured state is estimated to be 3.2 M. The free energy of unfolding of NcFADS is estimated to be 748.8calmol^-1. These studies indicate that NcFADS is in stable native conformation.

This denaturation was not successful with GdnCl so we later used Guanidinium thiocyante induced equilibrium unfolding of NcFADS is monitored by the changes in the tryptophan fluorescence shows that NcFADS

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**A.**

![Image](image1.png)

**B.**

![Image](image2.png)

**Fig. 4.** Isothermal colorimetric profile of NcFADS: Upper panel represents the raw thermograms and the lower panel represents the binding isotherms with integrated heats for NcFADS titrations. Both the titrations were performed in 1xPBS (pH7.5). A. NcFADS (50 μM) in the cell binds to 1 mM ATP with 5 mM MgCl_2 present in the syringe. B. NcFADS (40 μM) in the cell interacts with 1.5 mM FMN present in the syringe.
completely unfolds at 6 M GdmSCN. Concentration of the denaturant at which 50% of the molecules remain in the denatured state is estimated to be 1.59 M. The free energy of unfolding of NeFADS is estimated to be 1.4kCal/mole (Fig. 3E).

Thermodynamic stability of NeFADS was measured by subjecting the enzyme to denaturation with gradual increase in temperature. Differential scanning calorimetry (DSC), directly measures the enthalpic changes, accompanied by the folding and unfolding of proteins which are most commonly exothermic in nature. It directly measures the Tm of a protein indicating the temperature at which half the protein population was in equilibrium with the native and denatured state(s). Tm of NeFADS was found to be ~53.3 °C. The heat capacity (Cp) observed at this temperature was 4.99 kcal/mol K. The ∆H and ∆S obtained were ~222.43 kcal/mol and 0.68 kcal/K mol respectively, showing that NeFADS is a fairly stable enzyme.

IFT Analysis for the Interaction of NeFADS with FMN and ATP: Isothermal titration calorimetry is the direct measure of the binding affinity of a protein with its ligand and also a tool for studying the thermodynamic properties of the biological molecules. In the present study ITC was used to measure the binding affinities of both the substrates FMN and MgATP2− to NeFADS. Isothermograms obtained were hyperbolic. Isothermograms represent the binding of ligands as well as the heat changes occurring during binding. A single binding site was observed for both the substrates. Kd values for FMN was 20.8 μM and for MgATP16.6 μM respectively (Fig. 4). The lower k2 value of MgATP2− compared to FMN is consistent with the MgATP to be 1.59 M. The free energy of unfolding of NcFADS is estimated to be which 50% of the molecules remain in the denatured state is estimated in equilibrium with the native and denatured state(s). Tm of NeFADS was found to be ~53.3 °C. The heat capacity (Cp) observed at this temperature was 4.99 kcal/mol K. The ∆H and ∆S obtained were ~222.43 kcal/mol and 0.68 kcal/K mol respectively, showing that NeFADS is a fairly stable enzyme.

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