Metabolic Engineering of Cofactor $\text{F}_4\text{H}_2$ Production in *Mycobacterium smegmatis*

Ghader Bashiri$^{1,2,}$*, Aisyah M. Rehan$^{1,2}$, David R. Greenwood$^{2,3}$, James M. J. Dickson$^{1,2}$, Edward N. Baker$^{1,2}$

$^1$ Structural Biology Laboratory, School of Biological Sciences, The University of Auckland, Auckland, New Zealand, $^2$ Maurice Wilkins Centre for Molecular Biodiscovery, School of Biological Sciences, The University of Auckland, Auckland, New Zealand, $^3$ Centre for Genomics and Proteomics, School of Biological Sciences, The University of Auckland, Auckland, New Zealand

Abstract

Cofactor $\text{F}_4\text{H}_2$ is a unique electron carrier in a number of microorganisms including Archaea and Mycobacteria. It has been shown that $\text{F}_4\text{H}_2$ has a direct and important role in archaenal energy metabolism whereas the role of $\text{F}_4\text{H}_2$ in mycobacterial metabolism has only begun to be uncovered in the last few years. It has been suggested that cofactor $\text{F}_4\text{H}_2$ has a role in the pathogenesis of *M. tuberculosis*, the causative agent of tuberculosis. In the absence of a commercial source for $\text{F}_4\text{H}_2$, $\text{M. smegmatis}$ has previously been used to provide this cofactor for studies of the $\text{F}_4\text{H}_2$-dependent proteins from mycobacterial species. Three proteins have been shown to be involved in the $\text{F}_4\text{H}_2$ biosynthesis in Mycobacteria and three other proteins have been demonstrated to be involved in $\text{F}_4\text{H}_2$ metabolism. Here we report the over-expression of all of these proteins in $\text{M. smegmatis}$ and testing of their importance for $\text{F}_4\text{H}_2$ production. The results indicate that co-expression of the $\text{F}_4\text{H}_2$ biosynthetic proteins can give rise to a much higher $\text{F}_4\text{H}_2$ production level. This was achieved by designing and preparing a new T7 promoter–based co-expression shuttle vector. A combination of co-expression of the $\text{F}_4\text{H}_2$ biosynthetic proteins and fine-tuning of the culture media has enabled us to achieve $\text{F}_4\text{H}_2$ production levels of up to 10 times higher compared with the wild type *M. smegmatis* strain. The high levels of the $\text{F}_4\text{H}_2$ produced in this study provide a suitable source of this cofactor for studies of $\text{F}_4\text{H}_2$-dependent proteins from other microorganisms and for possible biotechnological applications.

Introduction

The cofactor $\text{F}_4\text{H}_2$ was first identified chemically in methanogenic Archaea in 1972 [1], although a compound with similar characteristics was previously described in Mycobacteria in the early 1960s [2,3]. Since its discovery, $\text{F}_4\text{H}_2$ and its precursor FO (so called 5-deazaflavins) have been found in a variety of microorganisms, including Archaea, bacteria and eukaryotic species (Table 1). $\text{F}_4\text{H}_2$ is named on the basis of its intense absorbance/fluorescence at 420 nm (emission 480 nm), which is redox dependent and is lost upon reduction of the cofactor. It also has unique chemical and biological characteristics; the isoalloxazine chromophore of $\text{F}_4\text{H}_2$ is structurally very similar to that of the flavins (FMN and FAD), although it is functionally similar to NAD(P)$^+$ [Figure 1]. Functionally, $\text{F}_4\text{H}_2$ is a two-electron carrier involved in hydride transfer reactions. The redox potential of $\text{F}_4\text{H}_2$ is $\text{F}_4\text{H}_2$/2$\text{F}_4\text{H}_2$ (−300 mV) is lower than those of the classical hydrogen carriers NAD(P)H/NAD(P)+2e$^-$ (−320 mV) and FADH$_2$/FAD+2e$^-$ (−219 mV) [4,5].

A key biosynthetic precursor of $\text{F}_4\text{H}_2$ is FO (7,8-didemethyl-8-hydroxy-5-deazariboflavin), comprising an isoalloxazine ring and ribitol moieties. Formation of $\text{F}_4\text{H}_2$ follows a series of biochemical reactions and is completed by the addition of a phospholactate group, and finally a poly-glutamate tail in which L-glutamate residues are linked together via γ-glutamyl bonds (Figure 2) [6,7]. The length of the poly–glutamate tail constitutes the main difference between the $\text{F}_4\text{H}_2$ cofactors from different microorganisms, the number of residues varying from 2–9. There are suggestions, however, that the type of α– or γ–glutamyl linkage in the terminal glutamate residue could also be different in some Archaeal species [6,8,9,10].

$\text{F}_4\text{H}_2$ is not commercially available and researchers working on $\text{F}_4\text{H}_2$–dependent proteins have to prepare it as required. With the discovery of new $\text{F}_4\text{H}_2$–dependent enzymes and increasing interest in $\text{F}_4\text{H}_2$–dependent reactions, especially in the case of the pathogen *Mycobacterium tuberculosis* (*Mt*), a resource with high yields of $\text{F}_4\text{H}_2$ production is required. $\text{F}_4\text{H}_2$ has been previously purified from various microorganisms, including Archaea (Methanobacterium, Methanococcus and Methanosarcina species) and Actinomycetes (Actinomadura, Actinoplanes, Streptomycetes, Rhodococcus, Nocardia and Mycobacteria species), with differing yields [11]. $\text{F}_4\text{H}_2$ purification in all cases, however, essentially follows the same principle; precipitation of cellular proteins using heat or an organic solvent, followed by separation.
of F420 from remaining cellular components based on its acidic nature [4]. In order to purify F420, a number of different chromatographic steps have been used, including ion exchange, adsorption, HPLC and gel filtration chromatography [6,9,11]. Isabelle et al. have reported thorough analyses of F420-producing microorganisms, and based on “ease of growth, fewer hazards, and lower costs” concluded that M. smegmatis is the best source for F420 production, providing there is no requirement for a particular number of glutamate residues in the F420 polyglutamate tail [11].

Our initial F420 purification trials indicated that M. smegmatis transformed to over-express the M. tuberculosis protein FGD1 (F420-dependent glucose-6-phosphate dehydrogenase 1) could produce higher levels of F420 compared with the wild type strains. This observation prompted us to thoroughly investigate the effects on F420 production of over-expression of other proteins known to be involved in F420 biosynthesis and metabolism in Mycobacteria. These include three proteins in the F420 biosynthetic pathway, viz. FbiA (Rv3261) [12], FbiB (Rv3262) [12] and FbiC (Rv1173) [13] and three other proteins which are shown to be involved in F420 metabolism: FGD1 (Rv0407) [9,14], Ddh (Rv3547) [13] and Rv0132c (author’s unpublished data).

Here we describe the development of vectors to co-express Mtb proteins in M. smegmatis. We further show that by co-expressing enzymes associated with F420 production and manipulating growth conditions, greatly increased levels of F420 can be obtained. With the growing recognition that F420 plays a crucial role in Mycobacteria and other organisms, this readily available source of the cofactor will be useful for testing its physiological and biochemical roles, and for possible applications in biotechnology.

### Materials and Methods

**Preparation of New Mycobacterial Vectors**

The pYUB1049 vector (5795 bp) is a product of ligation between the vectors pMS134 and pET28b–cmaA2 [16], resulting in a vector with a cloned gene between NdeI and BamH1 restriction sites. The pYUB1049 vector was subjected to restriction digestion using NotI (single site) and BglII (two sites) restriction sites, in order to obtain a linear vector without the multiple cloning site. The plasmid was first digested to completion with NotI (Roche Applied Science) and dephosphorylated using calf intestinal alkaline phosphatase (New England Biolabs) followed by ethanol precipitation. The NotI-cut linear pYUB1049 vector was subjected to a partial digestion with BglII (BglII isoschizomer, Fermentas) for 20 minutes and the reactions stopped using 5 μL 0.5 M EDTA. The digested vector was run on a 0.5% agarose gel and a DNA fragment corresponding to 4705 bp was excised and gel-purified.

The pET28b and pETDuet–1 vectors (Novagen) were double-digested using NotI and BglII enzymes. The resulting multiple cloning site fragments, 216 and 382 bp respectively, were ligated separately into the NotI/BglII fragment of the pYUB1049 vector using T4 DNA ligase (Roche Applied Science). Ligation mixtures were electroporated into E. coli TOP10 cells and the positive colonies were selected on low salt LB agar plates (tryptone 10 g/L, yeast extract 5 g/L, NaCl 5 g/L and agar 15 g/L, pH 8.0) containing 50 μg/mL hygromycin B. Positive clones were verified using restriction digestion and sequencing. The resulting vectors were designated as pYUB28b (4921 bp) and pYUBDuet (5087 bp), respectively.
PCR Amplification and Cloning

The open reading frames (ORFs) encoding Rv3261 (FbiA), Rv3262 (FbiB), Rv1173 (FbiC), Rv0407 (FGD1), Rv3547 (Ddn) and Rv0132c were amplified from *M. tuberculosis* H37Rv genomic DNA using *Pwo*, *Pfx* or *PrimeStar* polymerases with the primers outlined in Table 2. All constructs were cloned with either N– or C–terminal His6–tags. The amplified products for the FGD1 [17] and Rv0132c constructs were cloned using restriction/ligation cloning into the pYUB1049/pYUB28b vectors. The constructs were transformed into *E. coli* Top10 cells and plated on low salt LB agar medium supplemented with 50 μg/mL hygromycin B to select for colonies harbouring the plasmid. Positive clones were verified using colony PCR, restriction digestion and sequencing.

All other ORFs were cloned using the Gateway® cloning system into the pDESTsmg vector [18]. The Gateway® cloning system uses a nested PCR method involving two rounds of amplification in which the second round uses the product of the first round as template. Gene–specific primers are used in the first round PCR to amplify the gene of interest and generic primers are used in the second round amplification to incorporate the required recombination sites for subsequent cloning. The PCR products were cloned by recombination into pDONR221 (Invitrogen) using BP Clonase™ (Invitrogen), to generate the entry clones. The constructs were transformed into *E. coli* Top10 cells and plated on LB agar medium containing 50 μg/mL kanamycin. Positive clones were verified using BsrGI digestion and sequencing. These positive entry clones were recombined in vitro with pDESTsmg, in an LR reaction using LR Clonase™ (Invitrogen), to generate a *M. smegmatis* expression construct. Following transformation of recombinant pDESTsmg plasmids, positive clones were selected on low salt LB agar plates supplemented with 50 μg/mL hygromycin B and were verified using BsrGI digestion. The pYUBDuet vector was used to clone the F420 biosynthetic ORFs (FbiAB and FbiC) together using restriction/ligation cloning. Both FbiC and FbiAB ORFs were amplified using *PfuUltra* Fusion HS DNA polymerase (Stratagene) using the primers outlined in Table 2. FbiC was first cloned using NcoI/HindIII restriction sites and the FbiAB operon was subsequently cloned using NdeI/EcoRV restriction sites.

Expression in *M. smegmatis*

All expression constructs were electroporated individually into the *M. smegmatis* strain mc^2^517. Preparation of electrocompetent cells and electroporation procedures were performed following published protocols [19]. Briefly, *M. smegmatis* mc^2^517 cells were grown at 37°C in 7H9/ADC/Tween80 or LB/Tween80 containing 50 μg/mL kanamycin until an OD600 ~0.7. Cells were harvested and washed three times in 10% ice–cold glycerol and finally resuspended in 10% ice–cold glycerol. Single aliquots of the resulting competent cells (40 μL) were transformed with 1 μL of DNA and a further 260 μL of 10% glycerol in 0.2 cm cuvettes. Electroporation was performed using a Bio–Rad Gene Pulser set to the following parameters: *R* = 1000 Ω, *Q* = 25 μF and *V* = 2.5 kV. Cells were immediately harvested with 1 mL 7H9/ADC/Tween80 (Difco™ and BBL™ Middlebrook) or LB/Tween80 and incubated for 5 h at 37°C with shaking. Positive

---

**Figure 1. The molecular structures of F420, the flavins and NADP⁺.** (A) Schematic representation of F420 showing different parts of the molecule, whereas (B) and (C) show the molecular structures of the flavins and NADP⁺, respectively. The atoms involved in oxidoreductive reactions are numbered in all structures.

doi:10.1371/journal.pone.0015803.g001
Metabolic Engineering of F$_{420}$ Production

**Lactate Biosynthesis**

L-lactyl-2-diphospho-5'-guanosine (LPPG)

**Tyrosine Biosynthesis**

5-amino-6-ribitylamino-2,4(1H,3H)-pyrimidinedione

**Flavin Biosynthesis**

F$_{420}^{-}$-O$_{2}$

2-phospho-L-lactate transferase (FbiA/CofD)

F$_{420}^{-}$-n

2-amino-4-hydroxyphenylpyruvate

FO synthase (FbiC/CofG and CofH)

2-phospho-L-lactate transferase (FbiA/CofD)

F$_{420}^{-}$-O$_{2}$

F$_{420}^{-}$-L-glutaryl ligase (FbiB/CofE)
Figure 2. The proposed biosynthetic pathway for cofactor F420. The FbiA, B and C are Mycobacterial proteins whereas the CofA, B, C, D, E, G and H are Archaeal proteins involved in the biosynthetic reactions. The pathway to formation of the activated phospholactate moiety (LPPG) is yet to be experimentally established in Mycobacteria. In some Archaeal species an α–linked terminal glutamate residue caps the γ–linked poly–glutamate tail, the addition of which is catalyzed by CofH.

doi:10.1371/journal.pone.0015803.g002

Table 2. Oligonucleotide primers used in the amplification of the protein coding sequences in this study.

| Construct | Primer Sequences (5′–3′) | Restriction Enzyme |
|-----------|--------------------------|--------------------|
| Rv3261 (FbiA) | Forward GCCACCGGCGCGGTTGAAAGGTCACCGTTC GTC | n.a. |
| Rv3262 (FbiB) | Forward GCCACCGGCGCGGTTGAAAGGTCACCGTTC GTC | n.a. |
| Rv1173 (FbiC–pDESTsmg) | Forward GCCACCGGCGCGGTTGAAAGGTCACCGTTC GTC | n.a. |
| FbiAB | Forward GCCACCGGCGCGGTTGAAAGGTCACCGTTC GTC | Ncol |
| Rv1173 (FbiC–pYUBDuet) | Forward GCCACCGGCGCGGTTGAAAGGTCACCGTTC GTC | EcoRV |
| Rv3547 (Ddn) | Forward GCCACCGGCGCGGTTGAAAGGTCACCGTTC GTC | HindIII |
| Rv0132c | Forward GCCACCGGCGCGGTTGAAAGGTCACCGTTC GTC | Ncol |
| Gateway Generic | Forward GCCACCGGCGCGGTTGAAAGGTCACCGTTC GTC | BarnHI |
| Gateway Generic | Reverse GCCACCGGCGCGGTTGAAAGGTCACCGTTC GTC | Ncol |

The underlined nucleotides indicate overlapping base pairs with the generic primers for the ORFs cloned using the Gateway® system. The bold underlined nucleotides show the restriction site for the appropriate enzymes, as indicated in the right–hand column.

doi:10.1371/journal.pone.0015803.t002

Transformants were selected by plating on 7H10/ADC (Difco™ and BHI™ Middlebrook) or LBT agar plates containing 50 µg/mL each of kanamycin and hygromycin B.

Protein expression was performed either in autoinduction [20], LB or 7H9/ADC media supplemented with 0.05% Tween80 and 50 µg/mL each of kanamycin and hygromycin B. A single transformed colony was selected from a 7H10/ADC plate and used to inoculate a starter culture in MDG media (25 mM Na2HPO4, 25 mM KH2PO4, 50 mM NH4Cl, 5 mM Na2SO4, 2 mM MgSO4, 0.5% D-glucose, 0.25% L-aspartate, 0.2× metal mix) [20]. The starter culture was grown for 48–72 h at 37°C and was freshly used at a dilution of 1:100 to inoculate expression cultures of ZYM–5052 autoinduction (1% tryptone, 0.5% yeast extract, 25 mM Na2HPO4, 25 mM KH2PO4, 50 mM NH4Cl, 5 mM Na2SO4, 2 mM MgSO4, 0.5% D-glucose, 0.25% L-aspartate, 0.2× metal mix), LB or 7H9/ADC. The expression cultures were grown for 4 days at 37°C for maximal expression [17], LB, MDG and 7H9/ADC cultures were induced using IPTG at a final concentration of 0.1 or 1 mM.

Western Blot Analyses

M. smegmatis cells expressing different constructs were lysed twice using a cell disruptor (Constant Systems Ltd.) and centrifuged at 16,000 g to pellet non–lysed cells and other insoluble material. Protein samples were separated on a 15% SDS–PAGE gel and transferred to polyvinylidene difluoride (PVDF) membranes using a wet transfer protocol (200mA, 3 hours) [21]. His–tagged recombinant proteins were detected using a mouse monoclonal anti–His antibody and horseradish peroxidase–conjugated anti–mouse antibody (GE Healthcare). The Luminol (ECL plus kit, GE Healthcare) chemiluminescence was detected using an LAS4000 imaging system (Fujifilm).

FO and F420 Characterization

M. smegmatis cells expressing different M. tuberculosis proteins were grown in identical conditions to late log phase or stationary phase. In all expression cultures the ZYP–5052 autoinduction media was used for F420 production experiments and the media to flask volume ratio was kept constant at 20%. In order to optimize the media for F420 production, the ZY component of ZYM–5052 media was replaced by commonly used media bases including 2× ZY, YT (0.8% tryptone, 0.5% yeast extract and 42.77 mM NaCl), TB (1.2% tryptone, 2.4% yeast extract and 0.4% glycerol), S02 (2% tryptone, 0.5% yeast extract, 8.56 mM NaCl, 2.5 mM KCl and 10 mM MgCl2) and SOC (SOB with 20 mM glycerol). Iron and sulphur supplements (ferric ammonium citrate, ferric citrate and ferrous sulphate all at 0.1 mg/mL and L–cysteine at 1 mM) were also added to the expression media as a possible requirement for the FbiC enzyme. L–glutamate and manganese chloride (1 mM final concentration) were also added to the expression media to evaluate their necessity for FbiC–mediated F420 production [22].

To ascertain the optimum growth period for F420 production, eight identical cultures of M. smegmatis cells expressing the recombinant FbiABC construct were set up. Each culture had a wild type M. smegmatis culture as a control. At 24 h intervals, one culture each of control and recombinant FbiABC–expressing M. smegmatis cells were harvested and processed to monitor the F420
production level. The procedure was carried out for eight days and the F420 production ratio for each day was calculated by dividing the F420 fluorescence from FbiABC–expressing cells by fluorescence of the wild type control.

*M. smegmatis* cells were centrifuged for 15 min at 16000×g and the resulting media were used for FO characterization. The cell pellets were washed with 25 mM sodium phosphate buffer, pH 7.0 and were subsequently resuspended in 1 mL of the same buffer per 100 mg of cells (wet weight). The cell suspensions were autoclaved at 121°C for 15 min to break the cells open and were then centrifuged for 15 min at 16000×g. Fluorescence of the media and the extract were monitored using excitation wavelength of 420 nm (405±10 nm filter) and emission wavelength of 480 nm (485±15 nm filter). All fluorescence experiments were performed using an EnVision Multilabel plate reader (Perkin Elmer) in a 96–well plate format and were carried out in triplicate.

The autoclaved cell extracts were further purified using a HiTrap QFF ion exchange column (GE Healthcare) to separate the intracellular FO from the F420. The extract was run on the column pre–equilibrated with 25 mM sodium phosphate buffer, pH 7.0 and was subsequently washed with five column volumes of buffer. Two yellow fractions were eluted at 200 and 500 mM NaCl, respectively. The purified fractions were used for mass spectrometry analysis, together with the media from the previous step. The media (1 mL) was treated with an equal volume of cold acetone to precipitate the protein and the solution was then evaporated down to <0.5 mL to drive off the acetone. A mix of water and 5% aqueous methanol with 0.1% formic acid was added to bring the final concentration of methanol to less than 1% (total volume 4 mL). All samples were then applied to a pre–equilibrated Alltech Maxi–Clean 300 mg large pore 100Å 18 SPE cartridge and washed with 4 mL 5% methanol containing 0.1% formic acid followed by 4 mL 10% methanol. Compounds were eluted with 4 mL 80% methanol containing 5 mM ammonium bicarbonate pH 8.5. Eluates were evaporated under nitrogen and redissolved in 80% methanol and 20 mM ammonium acetate ready for mass spectrometry. Samples were infused at 3 μL/min under negative electrospray conditions into an LTQ–FT mass spectrometer (Thermo Scientific). The ion intensity data were obtained using a source voltage of 2.5 kV and capillary temperature of 225°C. Ions were examined in both the ion trap and ion cyclotron resonance cells, the latter to obtain high resolution (100,000 at m/z 400) accurate mass data. This was necessary to confirm the atomic composition of the ions and help deconvolute the contribution of metal ion adducts (Na+/K+) to the levels of individual poly–glutamate species. Up to four sodium ions were adducted to produce some double charged negative ions.

**Results**

**New Mycobacterial Expression Vectors**

The pYUB1049 vector does not provide an intact multiple cloning site and does not support C–terminal His–tag expression. In order to overcome these obstacles, the pYUB1049 vector was subjected to a restriction digestion using NcoI and BlpI enzymes and a linear fragment lacking the multiple cloning site was obtained. The resulting fragment was used as a backbone that could be ligated to the intact multiple cloning site from the pET28b or pETDuet–1 vectors to produce the pYUB28b and pYUBDuet vectors, respectively. Figure 3 provides a schematic representation of the vectors with the list of unique restriction sites that can be used for cloning.

**PCR Amplification and Cloning**

Six different ORFs which are believed to be involved in F420 biosynthesis (FbiA, FbiB and FbiC) or F420 metabolism (FGD1, Ddn and Rv0132c) were amplified and cloned for expression in *M. smegmatis* as His–tagged proteins. Assuming that FbiA and FbiB ORFs are transcribed as a single operon, we investigated the possibility of cloning and co–expression of the whole F420 biosynthetic pathway (FbiAB and FbiC) in order to boost F420 production yield. The pYUBDuet co–expression vector was designed, prepared and subsequently used to clone FbiC and FbiAB ORFs, making it possible to express three different proteins from a single vector. All three proteins were expressed in their native form without His–tags.

---

**Figure 3. A schematic representation of the vectors designed in this study.** The unique restriction sites in the multiple cloning sites of each vector are indicated. doi:10.1371/journal.pone.0015803.g003
Expression of Proteins in *M. smegmatis*

The six F420 biosynthetic or metabolic ORFs cloned into pYUB1049/pYUB208/pDESTsmg vectors were expressed in *M. smegmatis* as individual proteins. Each of these proteins were cloned with either N- or C-terminal His-tags, making it possible to detect the protein expression using monoclonal anti-His antibodies. The western blotting experiments indicated that all proteins were expressed in *M. smegmatis* cells, as shown by appearance of correctly-sized bands for the appropriate proteins (data not shown).

The expression of proteins from the pYUBDuet vector could not be detected using western blotting, as they did not contain any tags; however, their successful expression could be inferred from FO and F420 production as discussed later.

**Cofactor F420 Production**

Individual *M. smegmatis* cultures harbouring six different constructs (FbiA, FbiB, FbiC, FGD1, Ddhn and Rv0132c) were grown in order to find out the over-expression effect of these targets on F420 production. Three different media were initially used to express the proteins; LBt with IPTG induction, MDG with no or low induction using IPTG, and ZYM-5052 autoinduction media. Based on growth rate and cell mass, ZYM-5052 media was selected as the best media and was used to continue F420 production experiments. The fluorescence signals of the expression media and the cell extracts were monitored at 420 nm, enabling the detection of both FO and F420. It has been reported that FO comprises 1–7% of the total intracellular deazaflavin in Mycobacteria [8]; we used fluorescence at 420 nm to evaluate the F420 contents of the cellular extracts without taking into account the small portion of the fluorescence signal coming from FO.

The experimental results indicate that FGD1 over-expression increases F420 production by almost two-fold compared to the wild type strain (Figure 4, A). Cells expressing other Mtb proteins did not show a significant increase in F420 yield, however. Cells expressing the FbiC construct (pDEST-FbiC) showed a strong blue-green colour in the media. This is presumably due to the presence of fluorescent FO in the media which diffuses out of the cells as FO does not have any charge on the molecule to cause retention inside the cell (Figure 4, A) [23]. Mass spectrometry confirmed that FO was indeed responsible for the distinct fluorescence of the media (m/z 362.09870 [M+H]+; C14H15N3O7 requires 362.09862). This observation could be explained by over-expression of the FbiC protein leading to higher FO synthesis. Because the cells could not convert the over-produced FO to F420, the excess was presumably lost from the cells, either by diffusion or by active export.

This observation provided the motivation for us to co-express the FbiAB operon together with FbiC, hoping that over-expressed FbiA and FbiB proteins would be able to convert the synthesised FO into F420 inside the cells. The pYUBDuet vector was used to clone FbiABC ORFs together; FbiC was first cloned, resulting in the pYUBDuet-FbiC construct, after which FbiAB was introduced to obtain pYUBDuet-FbiABC. Both these constructs were used to investigate the effect on FO/F420 production (Figure 4, B). Cells expressing FbiC alone (pYUBDuet-FbiC) consistently showed more than 10-fold higher FO levels in the expression media compared to the wild type strains. It is an interesting observation that FO production by the pYUBDuet-FbiC construct is much higher (>50%) than by the pDESTsmg-FbiC construct, with the former expressing FbiC as the native protein whereas the latter has an N-terminal His-tag. In contrast, F420 production from pYUBDuet-FbiC was not significantly elevated compared to wild type. By expressing the FbiAB operon together with FbiC (pYUBDuet-FbiABC), however, F420 production was consistently more than five times higher inside the cells (Figure 4, B). These results clearly indicate that the cells express functional recombinant proteins resulting in much higher intracellular F420 levels.

*M. smegmatis* cells expressing the pYUBDuet-FbiABC construct were then used to find out the optimum time period for F420 production. The F420 production was monitored for eight days using ZYM-5052 media and the F420 production ratio was calculated and plotted versus the day of culture. The results indicated that the F420 levels were the highest on day four of the culture, after which the levels gradually decreased. Based on this result, the best time to harvest the cells for F420 purification is 4–5 days after setting up the expression culture (Figure 4, C).

Subsequently, a set of experiments was performed to find out the best media formulation to grow the cells for F420 production using an autoinduction protocol. ZY produced the highest F420 yield among ZY, YT, TB, SOB and SOC media. Bioinformatic analysis has indicated that FbiC is a protein with possible Fe-S clusters. In addition the reaction catalyzed by an archaeal homologue of FbiB requires L-glutamate and manganese chloride [22]. The expression media were therefore also supplemented with iron/sulphur and L-glutamate/manganese additives. The results indicated that supplementation of the expression media with either of these additives does indeed increase the F420 production yield (Figure 4, D). Surprisingly, cultures with an L-glutamate/manganese supplement did not have extra FO in the media, implying that the cells could convert all the produced FO into F420 inside the cells (Figure 4, D). It seems that the limiting factor in producing F420 from over-produced FO was the supply of the required L-glutamate/manganese.

The FO/F420 produced by the cells expressing the FbiABC construct was purified and analysed using mass spectrometry. The results show two predominant fractions; a 200 mM NaCl fraction mainly composed of FO and a 500 mM NaCl fraction of exclusively F420 with more than 95% being F420-6 and F420-7 species (Figure 5). This result is in line with the previously published results of F420 extracted from the wild type *M. smegmatis* cells having the major species of F420-5 to F420-7 [9,11], implying that the over-expression of the FbiABC construct does not change the F420 production profile.

**Discussion**

The cofactor F420 has an important role in the metabolism of Archaea and has been the subject of numerous studies over the years since its identification. It is now clear that this importance applies also to Mycobacteria, for which there is growing evidence that F420 plays a key role in defence against oxidative and nitrosative stress [5,24]. Consistent with this, the number of identified F420-dependent enzymes from Mycobacteria is growing, with nine new examples recently described [25]. A recent partial phylogenetic profiling study has proposed that there are at least 28 separate F420-dependent enzymes in *M. tuberculosis*, suggesting that F420 has a pivotal role in redox reactions of this pathogenic mycobacterium [26]. Few of these enzymes have been characterised, however, and research into their functions, and the role of F420, are handicapped by the fact that there is no commercial source for this cofactor, which can only be obtained in relatively low yield from the wild type *M. smegmatis* strain. A major aim of this study was to increase the F420 production yield in *M. smegmatis* by cloning and expression of the genes involved in F420 production and metabolism.

**Mycobacterial Expression Vectors**

The pYUB1049 plasmid is a T7 promoter-based vector for which expression can be induced by IPTG or autoinduction. This vector has previously been used as a shuttle vector for cloning of
Mycobacterial genes into \textit{E. coli} and subsequent expression of proteins in \textit{M. smegmatis} \cite{17,27}. The pYUB1049 vector has been also converted to a Gateway\textsuperscript{\textregistered} cloning system compatible vector, pDESTsmg \cite{18}. In this study, two different vectors were designed and prepared from the parental pYUB1049 vector; the pYUB28b vector is used for restriction/ligation cloning of single genes with the capability of expressing N– and C–terminal His–tags, whereas the pYUBDuet vector is a co–expression vector for simultaneous expression of two genes in a Mycobacterial host. Our experimental results demonstrate the application of T7–promoter based co–expression vectors in \textit{M. smegmatis} that could also be useful in other contexts. Although there have been previous reports of co–expression systems for Mycobacteria \cite{28,29,30,31,32,33}, to the best of our knowledge, this is the first T7–promoter–based vectors which can be routinely used for expression of a wide range of ORFs in a Mycobacterial host.

**F420 Production**

FbiC is annotated as FO synthase \cite{34}, catalysing the transfer of the hydroxybenzyl group from 4-hydroxyphenylpyruvate (a tyrosine precursor) to 5-amino-6-ribitylamino-2,4(1H,3H)-pyrimidinedione (an intermediate in flavin biosynthesis) to form FO \cite{23}. FO is the first intermediate with a complete deazaflavin chromophore in the F420 biosynthesis pathway \cite{13,23} providing the rationale for believing that this reaction might be the rate limiting step in F420 biosynthesis. FbiA and FbiB are believed to be involved in this reaction.

**Figure 4. FO/F_{420} production by \textit{M. smegmatis} cells expressing different recombinant proteins.** (A) comparative FO/F_{420} production between wild type strain and six different recombinant strains expressing proteins involved in F_{420} biosynthesis or metabolism. (B) FO/F_{420} production using co–expression vector compared to the wild type strain. (C) F_{420} production ratio by \textit{M. smegmatis} cells expressing FbiABC construct over wild type strains for eight days. (D) Effect of iron/sulphur and L–glutamate/manganese additives on the F_{420} production (FO is shown as black and F_{420} as grey). In panels (A), (B) and (D) the error bars are derived from experiments carried out in triplicate.

doi:10.1371/journal.pone.0015803.g004
to be involved in production of F_{420} from the precursor FO molecule; FbiA in generating F_{420}-0 from FO and FbiB in adding glutamate residues to F_{420}-0 to produce F_{420} with a poly-glutamate tail of variable length ([Figure 2] [12]). In all Mycobacterial species with the genome sequences completed to date ([as of October 2010]) FbiA is located immediately upstream of FbiB (www.TBdb.org). A detailed analysis indicates that the start site for the FbiB ORF overlaps with the last four base pairs of the FbiA ORF, though in a different reading frame, implying that they might be transcribed as a single operon for expression. In fact, it has been shown in M. bovis that these two ORFs are transcribed together as a single mRNA species [12]. This genetic arrangement made it possible to co-express the FbiAB operon and FbiC gene together, using the pYUBDuet vector, in order to increase F_{420} yield. Based on our results, the optimum condition to produce F_{420} by M. smegmatis cells expressing recombinant FbiABC is a culture with autoinduction media using ZY base over 4–5 days supplemented with iron, sulphur, L-glutamate and manganese. Using these optimal conditions, the F_{420} production yield was up to 10–times higher compared with the wild type strains.

The main limiting factor in F_{420} production, based on our results on over-expression of the three enzymes FbiA, FbiB and FbiC from the F_{420} biosynthetic pathway, appears to be the availability of the FbiB reaction substrate/cofactor. It does not seem that the FbiC reaction is the limiting step of the pathway even when the media are not supplemented with L-glutamate/manganese; excess FO was always present in high quantities in the media indicating that the two over-expressed FbiA and FbiB proteins are still not capable of converting all FO to F_{420}. An alternative possibility is that FbiA and FbiB need other accessory protein(s) in order to perform the conversion more efficiently; in fact another ORF in M. smegmatis (MSMEG_2392) has been shown, by transposon mutagenesis studies, to be involved in F_{420} biosynthesis from FO [35]. Biochemical studies need to be performed using purified enzymes in order to study the kinetics in detail and determine the rate limiting step of the pathway.

Our previous crystal structures of M. tuberculosis FGD1 [9], together with other F_{420}-containing crystal structures from different Archaeal species [36,37,38,39], have indicated that the F_{420} poly-glutamate tail is not required for reaction catalysis; the poly-glutamate tail is extended into the solvent and it seems that this is a conserved feature of the enzymes that use F_{420} in oxidoreduction reactions. We propose that, therefore, the high yields of F_{420} from M. smegmatis strains expressing the recombinant FbiABC proteins, regardless of the number of glutamate residues in the poly-glutamate tail, identify this as a valuable source of F_{420} that might be used with enzymes purified from other microorganisms. Furthermore, the high yield of FO/F_{420} also opens a door for possible biotechnological applications.

Depositions

The nucleotide sequences for pYUB23b and pYUBDuet vectors have been deposited in the National Centre for Biotechnology Information (NCBI) under GenBank HQ247814 and HQ247815 accession numbers, respectively. The vectors are available upon request.

Acknowledgments

The pYUB1049 vector and M. smegmatis mc²517 were kindly provided by Professor W. R. Jacobs, Albert Einstein College of Medicine.

Author Contributions

Conceived and designed the experiments: GB. Performed the experiments: GB AMR DRG. Analyzed the data: GB DRG. Contributed reagents/materials/analysis tools: JMJD. Wrote the paper: GB JMJD ENB. Acquisition of funding/supervision: ENB.

References

1. Cheeseeman P, Toms-Wood A, Wolfe RS (1972) Isolation and Properties of a Fluorescent Compound, Factor 420, from Methanobacterium Strain M.S.H. J Bacteriol 112: 527–531.
2. Cousins FB (1960) The prosthetic group of a chromoprotein from mycobacteria. Biochim Biophys Acta 40: 532–534.
3. Sutton WB (1964) Properties of a new TPN-like electron transport component from Mycobacterium phlei. Biochem Biophys Res Commun 15: 414–419.
4. DiMarco AA, Bobik TA, Wolfe RS (1990) Unusual coenzymes of methanogenic bacteria. Annu Rev Biochem 59: 355–394.
5. Purwantini E, Mukhopadhyay B (2009) Conversion of NO2 to NO by reduced coenzyme F420 protects mycobacteria from nitrosative damage. Proc Natl Acad Sci U A 106: 6333–6338.
6. Eirich LD, Vogels GD, Wolfe RS (1976) Distribution of coenzyme F420 and properties of its hydrolytic fragments. J Bacteriol 40: 29–27.
7. Bair TB, Isabelle DW, Daniels L (2001) Structures of coenzyme F(420) in Mycobacterium species. Arch Microbiol 170: 37–43.
8. Bashiri G, Square CJ, Moreland NJ, Baker EN (2008) Crystal structures of F420-dependent glucose-6-phosphate dehydrogenase FGDI involved in the activation of the anti-tuberculous drug candidate PA-824 reveal the basis of coenzyme and substrate binding. J Biol Chem 283: 17331–17341.
9. Graupner M, White RH (2003) Methanococcus jannaschii Coenzyme F420 Analogos Contain a Terminal α-Linked Glutamate. J Bacteriol 185: 662–4665.
10. Isabelle D, Simpson DR, Daniel L (2002) Large-scale production of coenzyme F420-5,6 by using Mycobacterium smegmatis. Appl Environ Microbiol 68: 5750–5755.
11. Choi KP, Bair TB, Baie YM, Daniels L (2001) Use of transposon Tn5367 mutagenesis and a nitroimidazopyran-based selection system to demonstrate a requirement for fbiA and fbiB in coenzyme F(420) biosynthesis by Mycobacterium tuberculosis. J Bacteriol 183: 7058–7066.
12. Choi K-P, Kendrick N, Daniels L (2002) Demonstration that fbiC is required by Mycobacterium bovis BCG for coenzyme F420 and FO biosynthesis. J Bacteriol 184: 2420–2429.
13. Purwantini E, Daniels L (1996) Purification of a novel coenzyme F420-dependent glucose-6-phosphate dehydrogenase from Mycobacterium smegmatis. J Bacteriol 178: 2981–2986.
14. Singh R, Maannatha U, Bodoff HI, Ha YH, Niyomrattanakit P, et al. (2008) PA-824 kills nonreplicating Mycobacterium tuberculosis by intracellular NO release. Science 322: 1392–1395.
15. Huang CC, Smith CV, Glickman MS, Jacobs WR, Jr., Sacchettini JC (2002) Crystal structures of mycoid acid cyclopropane synthases from Mycobacterium tuberculosis. J Biol Chem 277: 11559–11569.
18. Goldstone RM, Moreland NJ, Bashiri G, Baker EN, Shain L, Jaffe J (2008) A new Gateway vector and expression protocol for fast and efficient recombinant protein expression in Mycobacterium smegmatis. Protein Expr Purif 57: 81–87.

19. Cirillo JD, Weisbrod TR, William R, Jacobs J (1993) Efficient electrotransformation of proteins from polycarylamide gels to nitrocellulose sheets: procedure and some applications. Proc Natl Acad Sci U S A 70: 4396–4394.

20. Studier FW (2005) Protein production by auto-induction in high-density shaking cultures. Protein Expr Purif 41: 207–234.

21. Towbin H, Staehelin T, Gordon J (1979) Electrophoretic transfer of proteins from polyacrylamide gels to nitrocellulose sheets: procedure and some applications. Proc Natl Acad Sci U S A 76: 4350–4354.

22. Noeck B, Evdokimova E, Proudfoot M, Kudritka M, Grochowski LL, et al. (2007) Structure of an amide bond forming F420:gamma-glutamyl ligase from Archaeoglobus fulgidus – a member of a new family of non-ribosomal peptide synthetases. J Mol Biol 372: 456–469.

23. Graham DE, Xia H, White RH (2003) Identification of the 7,8-didemethyllum-8-hydroxy-5-deazabovallan synthase required for coenzyme F420 biosynthesis. Arch Microbiol 180: 455–464.

24. Darwin KH, Ehrt S, Gutierrez-Ramos J-C, Weich N, Nathan CF (2003) The evolution of the methylenetetrahydromethanopterin dehydrogenase. Biochemistry 42: 10098–10105.

25. Afuhammer SW, Warkentin E, Ermler U, Hagemeier CH, Thauer RK, et al. (2004) Biochemical and genetic characterization of two families of F420H2-dependent reductases from Mycobacteria that catalyze allantoin degradation. Mol Microbiol 52: 761–757.

26. Selengut JD, Haft DH (2010) Unexpected Abundance of Coenzyme F420-dependent enzymes in the Genomes of Mycobacterium tuberculosis and other Actinobacteria. J Bacteriol 192: 5758–5768.

27. Rohsen J, McKenzie JL, Curson R, Cook GM, Arcus VL (2009) The vanBPC operon from Mycobacterium smegmatis is an autoregulated toxin-antitoxin module that controls growth via inhibition of translation. J Mol Biol 390: 353–367.

28. Chang Y, Mead D, Hoddva H, Brum JF, Wolf PG (2009) One-plasmid tunable coexpression for mycobacterial protein-protein interaction studies. Protein Sci 18: 2316–2325.

29. George KM, Yuan Y, Sherman DR, Barry CE, 3rd (1995) The biosynthesis of cyclopropane mycolic acids in Mycobacterium tuberculosis. Identification and functional analysis of CAMAS-2. J Biochem 270: 27292–27298.

30. Hazlitt K, Widdel F, Wolfe RS (1989) Expression of secondary alcohol dehydrogenase in methanogenic bacteria and purification of the F420-dependent enzyme involved in cell carbon synthesis of Methanobacterium thermoautotrophicum. J Bacteriol 171: 604–613.

31. Johnson EF, Mukhopadhyay B (2005) A new type of sulfite reductase, a novel coenzyme F420-dependent enzyme, from the methanarchaeon Methanocaldococcus jannaschii. J Biol Chem 280: 38776–38786.

32. Jacobson FS, Daniels L, Fox JA, Walsh CT, Orme-Johnson WH (1982) Purification and properties of N5, N10-methylene-F420 and related biological assay of an unusual flavenoid cofactor involved in lincomycin biosynthesis. J Antibiot (Tokyo) 47: 472–474.

33. Jacobson FS, Daniels L, Fox JA, Walsh CT, Orme-Johnson WH (1982) Purification and properties of an N8-hydroxy-5-deazafavolin-reducing enzyme from Methanobacterium thermoautotrophicum. J Biol Chem 257: 3385–3388.

34. Deppenmeier U, Blaut M, Mahlmann A, Gotschalk G (1998) Membrane-bound F420H2-dependent heterodisulfide reductase in methanogenic bacteria. Science 282: 1963–1966.

35. Zeikus JG, Fuchs G, Kenealy W, Thauer RK (1983) Coenzyme F420: heterodisulfide oxidoreductase, a proton-translocating redox system in methanogenic bacteria. Proc Natl Acad Sci U S A 87: 9449–9453.

36. Last JF, Linder D, Stetter KO, Thauer RK (1994) F420H2: quinone oxidoreductase from Archaeoglobus fulgidus. Characterization of a membrane-bound multisubunit complex containing FAD and iron-sulfur clusters. Eur J Biochem 225: 503–511.

37. Szegf, Bryant MP, Wolfe RS (1975) Factor 420-dependent pyridine nucleotide-linked hydrogenase system of Methanobacterium ruminantium. J Bacteriol 121: 104–192.

38. Szegf, Bryant MP, Wolfe RS (1975) Factor 420-dependent pyridine nucleotide-linked formate metabolism of Methanobacterium ruminantium. J Bacteriol 121: 192–196.

39. Zekiu JG, Fuchs G, Kenealy W, Thauer RK (1977) Oxidoreductases involved in cell carbon synthesis of Methanobacterium thermoautotrophicum. J Bacteriol 132: 604–613.

40. Fuchs G, Stupperich E (1982) Autotrophic CO2 fixation pathway in Methanobacterium thermoautotrophicum. Zentralbl Bakteriol Hyg Abt 1 Orig C 3: 277–288.

41. Hartzell PL, Zelius G, Eschante-Senemger JM, Donnelly MI (1983) Coenzyme F420 dependence of the methylene-tetrahydromethanopterin dehydrogenase of Methanobacterium thermoautotrophicum. Biochim Biophys Res Commun 133: 884–890.

42. Ma K, Thauer RK (1990) Purification and properties of N3, N10-methylene-tetrahydromethanopterin reductase from Methanobacterium thermoautotrophicum (strain Marburg). Eur J Biochem 191: 187–193.

43. Widdel F, Wolfe RS (1989) Expression of secondary alcohol dehydrogenase in methanogenic bacteria and purification of the F420-specific enzyme involved in cell carbon synthesis of Methanobacterium thermoautotrophicum. J Bacteriol 171: 185–191.

44. Vermeij P, Vanke E, Keltjens JT, Van der Drift C (1994) Purification and characterization of coenzyme F430 synthetase from Methanobacterium thermoautotrophicum (strain delta H). Eur J Biochem 226: 185–191.

45. Coats JH, Li GP, Kuo MS, Yurek DA (1989) Discovery, production, and biological assay of an unusual flavonoid cofactor involved in lincomycin biosynthesis. J Antibiot (Tokyo) 47: 472–474.

46. McCormack JRD, Morton GO (1982) Identity of co-factoric factor 1 of Streptomyces aureofaciens and fragment FO from coenzyme F420 of Methanobacterium sp. J Am Chem Soc 104: 4014–4015.

47. Bieleski RL, Waxdal M (1968) Oxidoreductase with and without its substrates bound. EMBO J 20: 6561–6569.