Expression of membrane proteins from *Mycobacterium tuberculosis* in *Escherichia coli* as fusions with maltose binding protein

A. Korepanova, J.D. Moore, H.B. Nguyen, Y. Hua, T.A. Cross, F. Gao

Abstract

Sixteen of 22 low molecular weight integral membrane proteins from *Mycobacterium tuberculosis* with previously poor or undetectable levels of expression were expressed in *Escherichia coli* as fusions with both the maltose binding protein (MBP) and a His8-tag. Sixty-eight percent of targeted proteins were expressed in high yield (> 30 mg/L) in soluble and/or inclusion body form. Thrombin cleavage of the MBP fusion protein was successful for 10 of 13 proteins expressed as soluble proteins and for three proteins expressed only as inclusion bodies. The use of autoinduction growth media increased yields over Luria-Bertani (LB) growth media in 75% of the expressed proteins. Expressing integral membrane proteins with yields suitable for structural studies from a set of previously low and non-expressing proteins proved highly successful upon attachment of the maltose binding protein as a fusion tag.

Keywords: Membrane protein expression; Maltose binding protein; Fusion proteins

Production of membrane proteins for structural and functional studies remains a difficult task due to their highly hydrophobic nature [1,2]. During the last decade increasing efforts to express prokaryotic and eukaryotic membrane proteins have resulted in significant advancements for the expression of several bacterial transporter proteins [3–9], outer membrane proteins of bacteria [10], membrane protein complexes [11–13] and a few eukaryotic G-protein coupled receptors [14–21]. With others we have successfully expressed more than 70 membrane proteins from *M. tuberculosis* in *E. coli* using a His-tag [22]. However, a significant number of tested proteins (28%) were not expressed at detectable (Coomassie stain or Western) levels in *E. coli*. Expression of nonmembrane proteins by the Methanobacterium thermoautotrophicum structural genomics pilot project showed that almost 50% of cloned genes did not express when an N-terminal His-tag was used [23]. More importantly, 41% of our cloned membrane proteins with a His-tag that have a molecular weight less than 13.7 kDa did not express, and only 22% expressed well, such that a Coomassie stain was observed representing enough protein to proceed with structural studies. Here, we have set out to enhance the expression of small molecular weight proteins and to identify a robust protocol for doing this.

While many different vectors and tags have been used, the fusion construct with maltose binding protein (MBP) has been found to be an effective fusion partner for increased solubility and expression yields for a variety of recombinant water-soluble proteins [3,24–28]. MBP has also been tried with a few eukaryotic and prokaryotic membrane proteins [3,15,29–31]. Grisshammer and colleagues...
have achieved membrane localization and functional expression in *E. coli* for neurotensin and neurokinin-2, G-protein coupled receptors, upon fusion to the periplasmic MBP [1,14]. Expression as non-periplasmic MBP fusions in *E. coli* followed by purification and structural characterization by NMR was reported for two small membrane proteins: phospholamban and sarcolin [30].

Here, we report a successful application of the MBP fusion expression system for expression of small molecular weight integral membrane proteins from *M. tuberculosis* in *E. coli*. Some of these proteins are expressed as soluble fusion proteins obviating the need for harsh solubilizing conditions in extracting membrane proteins from inclusion bodies, but many others were expressed in high yields in inclusion bodies from which we have demonstrated isolation and cleavage of the fusion.

**Materials and methods**

**Cloning**

The DNA fragments encoding membrane proteins from *M. tuberculosis* were cloned into a modified pMALc2 plasmid [32] by PCR. Cloning was done according to standard techniques. Plasmid DNA encoding selected proteins were used as a template for PCR. All primers were purchased from IDT, Inc., USA. Pairs of gene specific primers were used to amplify DNA using standard PCR conditions and the enzyme *Pfu Turbo* DNA polymerase (Stratagene, USA), shown to have lower probability for introducing unwanted mutations. Two pairs of the restriction endonucleases were used to digest generated PCR fragments or vector DNA: *BamHI* and *HindIII* or *BamHI* and *PstI*. Sites for restriction endonucleases were introduced in the primer sequences. PCR products were gel purified after digestion with restriction enzymes and ligated into the prepared expression vector. Correct insertions into the vector were confirmed by PCR screening followed by DNA sequencing to check for in-frame insertion and lack of PCR-introduced point mutations.

**Expression constructs**

The plasmid results in the expression of the MBP fusion protein with eight histidine residues and the thrombin protease cleavage site at the N-terminus of the protein of interest to improve cleavage of MBP (Fig. 1). Importantly, two affinity tags, MBP and the His-tag, allow for a choice of the alternative purification approaches.

**Protein expression**

*Escherichia coli* BL-21(DE3) codonPlus-RP strain (Stratagene, USA) was used to express recombinant MBP fusion proteins. This strain is designed to compensate for codon usage differences between *M. tuberculosis* and *E. coli* and was successfully used in our previous expression of membrane proteins.

The *E. coli* cultures were initially grown in standard liquid media with appropriate antibiotic [33]. Briefly, 20 ml of LB growth media supplemented with 50 mg/ml ampicillin was inoculated with a 1:100 dilution of an overnight bacterial culture and incubated with agitation at 37°C to an OD$_{600}$ of $\sim$0.5. Protein expression was induced with 0.4 mM IPTG. Cells were harvested 3 h after induction by centrifugation at 5000g in an Avanti J-20 XP centrifuge (Beckman Coulter, Inc., USA). The cells were resuspended in 1 ml of water and lysed by sonication using a Sonic Dismembrator, Model 100 (Fischer Scientific, Inc.). The lysate was clarified by centrifugation for 15 min at 10,000g using a Microfuge 18 Centrifuge (Beckman Coulter, Inc.). The pellet was resuspended in 1 ml of 1% SDS and 4M urea. Insoluble and soluble fractions were checked for expression by the appearance on the SDS–PAGE gel stained with Coomassie R-250. Samples were loaded on the gel on the basis of equal volume, and the expression outcomes were assessed by visual inspection. Furthermore, the amount of expressed fusion protein was estimated for each sample during the purification step when the fusion protein was eluted from the amylose or His-tag affinity columns. The concentration was determined via the absorbance at 280 nm.

Bacterial autoinduction growth system, which utilizes lactose for the induction of protein expression has also been used for all expressed fusion proteins in an effort to increase yields [34]. Briefly, TB growth media was supplemented with 50 mg/ml ampicillin, 100 mM dibasic sodium phosphate, 50 mM monobasic potassium phosphate, 25 mM ammonium sulfate, 0.5% glycerol, 0.05% glucose, and 0.2% lactose. Twenty micro liters of prepared media were inoculated with a 1:100 dilution of an overnight bacterial culture and incubated with agitation at 37°C overnight (16h of growth). Cells were harvested by centrifugation followed by lysis and sample preparation as described above for the bacterial cells grown in LB media.

**Protein purification**

Membrane proteins from *M. tuberculosis* fused with the C-terminus of MBP were expressed in two forms: as soluble fusion proteins and as proteins directed into inclusion bodies. Dual affinity properties of the recombinant fusion protein, MBP and His-tag, allow the purification of both a
soluble recombinant protein using amylose affinity or metal affinity, as well as a protein solubilized from inclusion bodies and isolated via metal affinity chromatography.

Two hundred microliters of the LB or autoinduction media were used to grow *E. coli* cells expressing MBP fusion proteins under growth conditions identical to that described for the small expression screen. Cell debris was pelleted after lysis by centrifugation at 10,000 g. A supernatant containing soluble MBP fusion protein in a loading buffer consisting of 20 mM Tris, pH 8.0 and 200 mM NaCl was applied to an amylose resin affinity column at 4 °C. After extensive washing with a loading buffer the fusion protein was eluted from the column with 10 mM maltose followed by thrombin digest without any additional buffer change.

Fusion proteins expressed in inclusion body form were reconstituted from the pellet obtained after lysis and 10,000 g centrifugation in a buffer containing 20 mM Tris, pH 7.9, 5 mM imidazole, 500 mM NaCl, and 6 M urea for at least 2 h at 4 °C. A sample was applied to the Chelating Sepharose Fast Flow column charged with Ni²⁺ and equilibrated with an identical buffer. After extensive washing with 20 mM imidazole, the fusion protein was eluted from the column with 300 mM imidazole. The isolated protein was dialyzed extensively to decrease the concentration of imidazole in the sample prior to thrombin digest while maintaining the urea concentration at 1 M to prevent aggregation and precipitation of a fusion protein.

**Thrombin digest**

Thrombin from human plasma (catalog #T6884) was purchased from Sigma, (USA). Thrombin digest of MBP fusion proteins was selectively optimized for the amount of thrombin added per mg of fusion protein and the duration of the cleavage reaction. The reactions were supplemented with additives such as 0.01% SDS, 1 M urea, and 10% glycerol in order to facilitate the cleavage. The choice of additives was based on the thrombin digest recommendations developed by Novagen, USA. The completion of the thrombin digest was verified via the correct shifts in the positions of the protein bands on a polyacrylamide gel.

**Results**

**Target selection and cloning**

Twenty-two integral membrane proteins from *M. tuberculosis* were cloned into a modified pMALc2 plasmid for expression as fusions to the C-terminus of non-secreted MBP (Table 1). Previously, these proteins did not yield any detectable expression as fusions with short N-terminal Histags [22] except for Rv0463, Rv0882, Rv0961, Rv1305, and Rv2076c which showed low level expression identified via western blots. Proteins selected for MBP fusions had a molecular weight range between 7.1 and 13.7 kDa and a number of putative transmembrane helices that varied

<table>
<thead>
<tr>
<th>Membrane protein from <em>M. tuberculosis</em></th>
<th>Fusion protein (MW, kDa)</th>
<th>Membrane protein (MW, kDa)</th>
<th>#TM</th>
<th>LB media growth</th>
<th>Autoinduction media growth</th>
<th>Thrombin digest (%)</th>
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“o” denotes no observed expression, and “+” though “+++++” represents increasing expression level, where “+” corresponds to an expression yield between 10 and 30 mg of fusion protein per L of the bacterial culture, “+++” corresponds to yields between 40 and 60 mg, “+++++” corresponds to fusion protein yields between 70 and 90 mg, and “++++++” corresponds to yields of fusion protein exceeding 100 mg. The amount of protein was determined by absorbance at 280 nm after eluting the sample from an amylose or metal affinity column. Abbreviations are as follows: #TM, the number of putative transmembrane helices; MW, molecular weight in kDa; n/a, not applicable; n/t, not tested.
between one and four. These proteins were selected for expression trials because they are attractive targets for structural characterization by NMR.

Expression

Sixteen of 22 (72%) *M. tuberculosis* membrane proteins were expressed as MBP fusions in *E. coli*. Results of the expression testing are shown in Table 1. We consider the expression of the MBP fusion protein in soluble form preferable because soluble proteins can be easily purified in one step using affinity chromatography and the thrombin digest can be done under non-denaturing conditions.

Significant variations in the amount of expressed protein were observed in the different cellular compartments. In considering cultures in both LB and autoinduction media, 13 fusion constructs out of 16 (81%) were expressed as soluble proteins and all 16 were expressed in inclusion bodies. The highest levels of expression in soluble form were achieved for Rv0476, Rv1440, and Rv1824 with yields up to 120 mg of fusion protein per L of bacterial culture as estimated by measuring the absorbance at 280 nm after purifying the fusion proteins via an amylose affinity column. Most of the tested proteins showed moderate expression yields in the soluble form, between 10 and 30 mg of fusion protein per L of bacterial culture (e.g., Rv2876, Fig. 2, panel b). For most of these fusion constructs the amount of expressed protein was substantially greater in the insoluble fraction than in the soluble fraction. Thus, the Rv1824 fusion expressed in LB media has three fold (based on fraction) than in inclusion bodies (Fig. 2, panel a). For three proteins (Rv3155, Rv3346c, and Rv3355c) expression was detected only in the insoluble fraction (Fig. 2, panel c).

The use of autoinduction media resulted in an increase in the expression yields for 12 of the tested proteins. While inclusion body expression increased for 11 of 12, the expression in soluble form increased for just five proteins: Rv0460, Rv0514, Rv1440, Rv1824, and Rv2876. Rv1824 fusion protein showed the most dramatic improvement with the use of autoinduction media as the amount of soluble protein was tripled (Fig. 3). Importantly, Rv0460 and Rv0514 MBP fusions were only expressed in soluble form by the autoinduction media. While autoinduction improved the expression for a number of proteins, there were several proteins (Rv0476, Rv0882, Rv2144c, and Rv3346c) for which expression was not improved, and two (Rv1305 and Rv2128) for which expression in the soluble form was only achieved in LB media.

Thrombin digest of MBP fusion proteins

Our attempts to use a thrombin cleavage site were successful (>30% cleavage) for 13 of the 16 (81%) expressed proteins (Table 1). However, fusion proteins showed significant variation in their ability to be cleaved with the protease (Table 1). The 13 soluble fusion proteins were purified using amylose affinity chromatography prior to thrombin digest. The concentration of fusion protein was assessed via protein absorbance at 280 nm and an appropriate amount of thrombin was added to the samples while still in the elution buffer. In most cases, 3–10 U of thrombin were used to cleave 1 mg of fusion protein at room temperature overnight. The additives such as 0.01% SDS or 1 M urea as well as 10% glycerol were selectively added to the reaction mixtures to optimize the cleavage. An example of cleavage optimization efforts for Rv1824 fusion protein is presented in Fig. 4. A low concentration of SDS was found to be most effective in facilitating the cleavage of fusion proteins. Nine of the 13 (69%) fusion proteins expressed in soluble form were completely cleaved.

If fusion proteins were expressed in soluble form, thrombin digest was performed only on the soluble fraction. Three fusion proteins, Rv3155, Rv3346c, and Rv3355c, were not expressed in soluble form and were isolated from inclusion bodies using a His-tag affinity column and digested with thrombin in the presence of 1 M urea. It required more protease to cleave the protein from inclusion bodies than protein expressed in soluble form; at least 15 U of thrombin were added for each mg of fusion protein. A further increase in the thrombin concentration did not improve the fraction cleaved. We estimate that only 30–50% of the fusion protein was cleaved with the protease.

Discussion

A desirable expression system should produce a large amount of stable protein and permit efficient protein purification for subsequent structural or functional studies. Fusion protein approaches are popular and MBP fusions have been extensively utilized for water-soluble protein expression. However, relatively few transmembrane proteins and interfacial membrane proteins have been tested for expression as fusions with the large MBP [1,14,15,29,30]. Fusions of hydrophobic membrane proteins with the hydrophilic MBP domain have been shown to reduce toxicity and improve stability for bacterio-opsin and bacteriorhodopsin [35,36]. This study reports on the expression of multiple membrane proteins using MBP as a fusion partner. Sixteen membrane proteins from *M. tuberculosis* were expressed in *E. coli* here, only three of which had been expressed previously and these had poor yields. The achieved levels of expression with MBP fusions make functional and structural analyses feasible even though only a small fraction of the microgram per L of culture is the desired protein product. Variation in expression yield is likely to be influenced by the size of the polypeptide, its amino acid content and three-dimensional conformation. It has been suggested that MBP has chaperon-like features and facilitates correct protein folding [24,37,38]. Thus, MBP may provide a protein scaffold that prevents the membrane protein portion of the fusion from aggregating, while MBP maintains its own structural integrity verified by its binding to an amylose column. Final separation of the membrane protein and its MBP partner after thrombin digest requires
the addition of a detergent to create a membrane mimetic environment suitable for the membrane protein [39].

Comparing the expression outcomes in LB media and in the autoinduction media shows that there is no universal solution for protein expression of these low molecular weight membrane proteins. While it was rare that expression was observed in one media and not the other, the yields of both the soluble and insoluble forms of the fusion protein did vary substantially warranting an effort to “fine tune” the expression conditions.

Expression results suggest that proteins are more frequently expressed in soluble form as fusions with MBP when they have fewer transmembrane helices and hence fewer hydrophobic residues. In analyzing the combined results from LB and autoinduction media nine out of 12 proteins (75%) with one or two transmembrane helices are expressed in soluble form while four out of 10 proteins (40%) with three or four transmembrane helices are expressed in soluble form. Even if the pool of proteins is restricted to just those that are expressed, the percentages still favor those with a small number of helices by 83–60%. There is no such trend for proteins expressed in inclusion bodies. Similarly, for proteins with only a His-tag there was no significant expression correlation with the number of transmembrane helices over a larger data set [22]. For soluble protein expression the data also suggests a correlation
with molecular weight—10 out of 13 proteins (70%) having a molecular weight less than 10.0 kDa are expressed in the soluble fraction in LB and/or autoinduction media while six out of 12 proteins (50%) having a molecular weight greater than 10.0 kDa are expressed in the soluble fraction. This correlation is likely to result from a correlation between molecular weight and the number of transmembrane helices which exists for such small membrane proteins and consequently the observed molecular weight correlation supports the transmembrane data analysis described above.

Separation of a target protein from the fusion partner may present a major challenge [37]. Incomplete enzymatic digestion with a protease is often due to inaccessibility of the protease site. Attempts to use a thrombin cleavage site for structural studies, although there have been successful structural efforts even for membrane proteins at much lower expression levels. Out of 22 proteins that were not expressed at such levels with only a His-tag, five were expressed here at levels greater than 30 mg/L as an MBP fusion in solution. Disappointingly, only two of these were successfully cleaved, Rv1824 and Rv2876. However, 14 of these 22 proteins were expressed at levels greater than 30 mg/L as an MBP fusion in inclusion bodies and all these high expression proteins for which thrombin cleavage was attempted (3) were substantially cleaved suggesting that an increased effort with the inclusion body fraction is warranted. Considerable success has been achieved with thrombin cleavage by using a modest detergent concentration and consequently the results suggest that this approach is a good choice for a first effort cleavage strategy, but others should be tried if thrombin fails. Conditions for detergent solubilization may need to be screened to optimize the percent cleavage. Here, only a low percentage of SDS has been tested.

We have shown in the present study that multiple small integral membrane proteins from Mycobacterium tuberculosis are successfully expressed as fusions to the C-terminus of non-secreted MBP in E. coli. Most of the target proteins did not show any expression while fused with only an N-terminal His-tag in our previous study [22]. Expression of small molecular weight proteins was a clear weakness in the N-terminal His-tag approach with only 22% of the cloned proteins expressing well enough for structural studies. Here, the MBP fusion strategy represents an excellent second approach for this class of proteins, as the percent of well-expressed membrane proteins having a molecular weight <13.7 kDa is increased to 63%. As the hydrophobic portion of the fusion protein increases one might expect that proteins with a larger number of helices might not produce much protein in the soluble form and, in fact, there is some evidence for this. The results show that MBP fusions are an effective vector choice for low molecular weight membrane proteins and that the choice of media can significantly affect expression and expression yield. While LB media is a good choice for demonstrating expression, the yields of both soluble and insoluble forms were most frequently increased with autoinduction media.

Acknowledgments

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References


![Fig. 3. The results of small scale protein expression screening using 12% Tricine gels (as in Fig. 2) for Rv1824 MBP fusion protein in LB and autoinduction (AI) media. The lanes are as follows: MW, molecular weight marker; NI, noninduced cells from LB media; Sol, soluble fraction; IB, inclusion bodies fraction.](image)

![Fig. 4. Polyacrylamide gel results of a thrombin digest of Rv1824 MBP fusion protein using gels as described in Fig. 2. Thrombin digest results in two protein bands on the gel: 12.7 kDa for Rv1824 and 42.5 kDa for MBP with the addition of the His-tag. The lanes are as follows: NC, not cleaved; MW, molecular weight marker; SDS, 0.01% SDS added to reaction mixture; Urea, 0.8M urea added to reaction mixture; U, units of thrombin per mg fusion protein added to reaction mixture. In all cases the reaction mixture was incubated at room temperature for 14 h.](image)


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