



Effective expression of human proteins on bacterial magnetic particles in an anchor gene deletion mutant of *Magnetospirillum magneticum* AMB-1

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ABSTRACT

Biologically synthesized magnetic particles by magnetotactic bacteria (BacMPs) have promising potential in the area of functional protein display technology for various biotechnological and biomedical applications. Functional proteins fused with an anchor protein, Mms13, have been demonstrated to be an effective and stable method for the display of functional proteins on BacMPs. However, the expression of some human proteins is relatively low. Useful host strains of *Escherichia coli* have been developed for the enhanced expression of recombinant proteins using a genetic engineering approach. To improve human protein expression level on BacMPs in *Magnetospirillum magneticum* AMB-1, a mutant strain with a deleted native *mms13* gene ($\Delta mms13$ strain) was established and evaluated for effective functional protein display technology. As a result, the $\Delta mms13$ strain synthesized BacMPs with significantly improved expression of two human proteins, thyroid-stimulating hormone receptor (TSHR) and the class II major histocompatibility complex (MHC II) molecules. The $\Delta mms13$ strain could therefore be an effective strain for the display of other important human proteins on BacMPs and may be useful for further applications.

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1. Introduction

Magnetic particles are increasingly used in various applications across biomedical and environmental fields, and progressively have been incorporated as support materials in areas such as enzyme immobilization [1,2], drug delivery [3], and cell separation [4,5]. The greatest advantage of magnetic particles is the ability to manipulate them with an external magnetic field. This characteristic makes it possible to easily recover functionalized magnetic particle-bound target molecules from complex heterogeneous reaction solutions. For the effective binding of a range of targets in various reaction conditions, methodologies for magnetic particle synthesis and surface functionalization have been developed [4,6–8].

Magnetotactic bacteria synthesize nano-sized (50–100 nm), uniform, and strong ferrimagnetic bacterial magnetic particles (BacMPs). The BacMPs are covered with a lipid membrane, derived from cytoplasmic membrane through the invagination process [9,10]. Because the lipids on BacMPs are mainly comprised of phospholipids, the purified BacMPs show high dispersibility based on

the negative-charged electrostatic interaction. Previous molecular studies have documented that BacMP synthesis is regulated by a unique set of membrane proteins on the BacMPs [10–12]. Genome sequencing and identification of BacMP membrane proteins opened the door for the novel protein display technology on BacMPs to functionalize its surface using genetic engineering in *Magnetospirillum magneticum* AMB-1 (*M. magneticum* AMB-1).

A fusion protein expression system involving an anchor protein identified from the BacMP membrane and a functional protein have allowed various functionalized BacMPs to be produced thus far [13,14]. The expression plasmid, pUMG, is a stable and high copy number plasmid which has been used for the expression of fusion genes containing an anchor protein gene and target protein gene [15]. Transformation using the plasmid vector with selective integration of the promoter region and anchor protein gene allows for efficient functional protein expression onto BacMPs. Among the anchor proteins, Mms13, a major BacMP membrane protein tightly bound to core magnetite particles, has been successfully demonstrated to stably display target proteins on BacMPs [16]. In addition, highly active promoters in *M. magneticum* AMB-1 were identified and an inducible protein expression system was recently developed to avoid the toxic effects of transmembrane protein expression in bacterial cells [17]. Various expression systems for protein display on BacMPs have been developed based on the techniques of other host cells, such as *Escherichia coli* (*E. coli*). However, the amount of displayed proteins on BacMPs depends on the

Abbreviations: *M. magneticum* AMB-1, *Magnetospirillum magneticum* AMB-1; BacMP, bacterial magnetic particle; $\Delta mms13$, *mms13* deletion mutant; TSHR, thyroid-stimulating hormone receptor; MHC, major histocompatibility complex; MSGM, magnetic spirillum growth medium; GPCR, G protein-coupled receptor.

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properties of the human protein. In particular, the amount of expressed human protein or membrane protein on BacMPs is quite low. Therefore, the establishment of techniques for protein displays on BacMPs is still in demand.

Techniques for heterologous protein expression using *E. coli*, yeast, insect, and mammalian cells have been developed to enhance expression levels. The varieties of available expression plasmids, inducible expression systems, recombinant fusion partners, and mutant strains have advanced the possibilities in *E. coli*. Various mutant *E. coli* host strains have been developed by genetic engineering and widely used for many applications. For example, expression strains should be deficient in the most harmful natural proteases because one of the major problems associated with the expression of heterologous proteins in *E. coli* is the degradation of cloned gene products by host-specific proteases. In addition, *recA* negative strains and *trxB/gor* negative mutants have been used for the stabilization of target plasmids and the enhancement of cytoplasmic disulfide bond formation [18]. Thus, the genomic engineering of host strains is a useful strategy for expression of recombinant proteins.

In this study, we focused on improving the amount of displayed human protein using an anchor protein gene deletion mutant (*Δmms13*) strain, which eliminates the expression of the *mms13* gene in the genome. During expression of human proteins on BacMPs in *M. magneticum* AMB-1, the anchor protein, Mms13, appears to be expressed both from the plasmid vector as a fusion protein and from the genome as a native protein. The competitive expression of these proteins might restrict the number of functional fusion proteins on BacMPs.

The objectives here were to: (1) establish the *mms13* gene deletion mutant strain, (2) evaluate BacMP production within the *Δmms13* strain, and (3) express and evaluate production of thyroid-stimulating hormone receptor (TSHR) and class II major histocompatibility complex (MHC II) molecules on BacMPs in the *Δmms13* strain. The results reported here suggest that the novel *Δmms13* strain may be an efficient strain for the display of various human proteins on BacMPs, and a useful tool for a number of applications. This is the first report of a mutant *M. magneticum* host strain for protein display on BacMPs and it demonstrates efficient display of human proteins at the surface of BacMPs.

2. Materials and methods

2.1. Bacterial strains and culture conditions

E. coli strain S17-1 was used as the host cell for conjugation [19] and *E. coli* DH5 α was used as the host for gene cloning. Cells were cultured in LB medium containing 25 μ g/ml kanamycin or 50 μ g/ml ampicillin at 37 °C. *M. magneticum* AMB-1 (ATCC 700264) [20] was microaerobically cultured in magnetic spirillum growth medium (MSGM) at 28 °C as previously described [20,21]. Microaerobic conditions were established by purging the cultures with argon gas. *M. magneticum* AMB-1 transformants harboring each expression vector were cultured under the same conditions in medium containing 5 μ g/ml ampicillin. All other reagents were laboratory-grade, commercially available analytical reagents. Deionized distilled water was used in all procedures.

2.2. Preparation of BacMPs from *M. magneticum* AMB-1

Cultured *M. magneticum* AMB-1 cells were collected by centrifugation at 11,344 g for 10 min at 4 °C, resuspended in 10 mM phosphate buffered saline (PBS, 40 ml, pH 7.4) and disrupted by three passes through a French press cell at 1500 kg/cm² (Ohtake Works Co. Ltd., Tokyo, Japan). BacMPs were collected from the disrupted

cells using a columnar neodymium-boron (Nd-B) magnet and washed 10 times with 10 mM HEPES buffer (pH 7.4). The washed BacMPs were suspended in PBS and stored at 4 °C. The concentration of BacMPs in suspension was determined by measuring the optical density (660 nm) using a spectrophotometer (UV-2200; Shimadzu, Kyoto, Japan). A value of 1.0 corresponded to 172 μ g (dry weight) BacMPs/ml.

2.3. Construction of the *Δmms13* strain by homologous recombination

The plasmids and primers used in this study are described in Table S1. The *mms13* sequence (0.4 kb) was obtained from the NCBI (Gene ID: 3805263). The DNA fragment was amplified from *mms13* upstream (0.8 kb) and downstream (0.8 kb) sequences by PCR with primer sets M13F5-800 and M13R3-800, which contained the restriction sites for *Bam*HI. The resulting fragment was cloned into the *Ssp*I site of pUC19, and the resulting plasmid was defined as pUC19M13. After confirming the DNA sequence, the outside region of the *mms13* fragment within pUC19M13 was amplified with primers M13F3-ad and M13R5-ad. These primers had an additional 24 bp sequences for In-Fusion PCR cloning (Takara). The gentamicin-resistance gene was amplified primer set Gm5-ad and Gm3-ad. These primers also had an additional 24 bp sequences, corresponding to the additional sequences of primer set M13F3-ad and M13R5-ad (Table S1). The purified PCR product of the gentamicin-resistance gene was fused to the amplified fragment of the outside region of the *mms13* using the In-Fusion system, which generated pUC19M13updownGm^r. To construct the plasmid for homologous recombination, the sub-cloned DNA fragment M13updownGm^r in pUC19M13updownGm^r was integrated into the *Bam*HI site within pK19*mobsacB* [22], which contains *sacB* as a counter-selectable suicide marker [23], to create pK19*mobsacB*M13updownGm^r (pK19*mobsacB*M13Gm^r).

pK19*mobsacB*M13Gm^r was introduced into *M. magneticum* AMB-1 cells using *E. coli* S17-1 as the donor strain for the conjugation. The cells were incubated in MSGM solid medium containing 2.5 μ g/ml gentamicin at 28 °C as previously described [24]. Individual colonies were picked and grown in MSGM containing 2.5 μ g/ml gentamicin. The cells were then incubated on plate medium in the presence of 2.5 μ g/ml gentamicin and filtered 1% sucrose under the same conditions. Resulting colonies were analyzed by PCR and sequenced to confirm that *mms13* was replaced with the gentamicin-resistance gene (Fig. 1).

2.4. Protein profiling of BacMP membrane from *Δmms13*

Quantitation of protein on the BacMPs from the wild-type and *Δmms13* strain was performed by the Lowry method. BacMP membrane protein (1 mg) was extracted by treatment with 0.02% SDS and boiling. BSA was used as a standard for protein quantification. Extracted and purified BacMPs (3 mg) from the wild-type and *Δmms13* strain were treated by boiling in 1% SDS for 30 min to obtain the membrane proteins. BacMPs were removed by centrifugation and magnetic separation, and the supernatant (BacMP membrane protein fraction) was mixed with SDS sample buffer containing 6.25 mM Tris-HCl (pH 6.8), 5% 2-mercaptoethanol, 2% SDS, 5% sucrose, and 0.002% bromophenol blue. The membrane proteins were denatured and subjected to SDS-polyacrylamide gel electrophoresis (SDS-PAGE) in a 12.5% (wt/vol) polyacrylamide gel. The gel was stained with Coomassie brilliant blue.

2.5. Expression of *Mms13* fusion proteins in the *Δmms13* strain

Each protein expression vector was derived from pUMG (Apr; 6.4 kb) [15] and pUMGP16M13 [16], which includes the *Mms13* promoter and the coding sequence for *Mms13* in plasmid pUMG.

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