Overproduction, Purification and Refolding of codon-optimized Hepatitis B Virus X Protein Subgenotype B3 in *Escherichia coli* BL21(DE3)

Anita Artarini**, Armini Syamsidi†, Anindyajati†, Raymond R. Tjandrawinata‡, Debbie S. Retnoningrum***

1Laboratory of Pharmaceutical Biotechnology, School of Pharmacy, Institut Teknologi Bandung, Bandung, Indonesia
2Dexa Laboratories of Biomolecular Sciences, Cikarang, Indonesia

** ABSTRACT**

Hepatitis B virus (HBV) infects human and causes chronic liver infection, leading to liver cirrhosis and hepatocellular carcinoma. HBV X (Hbx) protein is known to interact with tumor suppressor protein p53 and block its translocation into the nucleus. This study outlines the overproduction of Hbx protein from HBV subgenotype B3 in *Escherichia coli* BL21(DE3), including its purification and refolding. The gene encoding Hbx was first codon-optimized and inserted into pET16b. The recombinant plasmid was then transformed into *E. coli* BL21(DE3) as an expression host. Optimization of Hbx expression was performed with variation of IPTG concentration and overproduction temperature. The results showed that Hbx protein was optimally induced by 0.075 mM IPTG and overproduction of Hbx at 17, 25, and 37°C exhibited no difference in protein level and location. The optimal refolding of Hbx was obtained using 0.1 M arginine prior to elution from Nickel column using 100 mM imidazole and 0.25 M arginine. Hbx migrates differently in SDS-PAGE reducing and non-reducing, while the melting curve pattern in TSA analysis changed after the refolding step. Essentially, this purified Hbx protein could potentially be used for interaction study with p53 and the inhibitor candidate of the protein.

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1. Introduction

Hepatitis B virus (HBV) belongs to the Hepadnaviridae family and is the causative agent of liver infection. HBV causes acute and chronic liver infection, where the chronic infection has higher risk of developing liver cirrhosis and hepatocellular carcinoma (HCC) (Chang 2007). It has been reported that two million people were infected by HBV and 650,000 died due to complication of HCC. Development of chronic infection into cirrhosis and HCC was allegedly related to the viral titer, genotype and subgenotype, as well as Hbx protein (Kew 2011; El-Serag 2012; Sunbul 2014). Infection by HBV genotype C has higher risk in developing cirrhosis and HCC, while infection with HBV genotype B may develop HCC without cirrhosis (Chemin and Zoulim 2009; Di Bisceglie 2009).

Hbx protein is the smallest HBV protein which is composed of 154 amino acids and has four disulphide bonds, between Cys7-Cys69, Cys61-Cys115, Cys78-Cys137, and Cys17-Cys143 (Sidhu et al. 2014). Hbx is known for its function in cell cycle modulation, apoptosis regulation and modulation of signal transduction. Hbx influences signal transduction of Ras, Raf, c-jun, MAPK, NFk-B, Jak-Stat, FAK, PKC, Src-dependent signalling and PI3K. Hbx also inactivates p53 by directly interacting with Hbx C-terminal region, therefore inhibiting p53 localization into nucleus (Kew 2011; Shlomai et al. 2014; Hussein 2016).

In this study, we performed overproduction, purification and refolding of Hbx protein in *Escherichia coli* BL21(DE3) for further application in p53 interaction study as well as screening for prospective inhibitors. The gene encoding for Hbx gene was first optimized according to the codon preference and %GC of *E. coli* BL21. Optimization of Hbx overproduction was performed by variations of IPTG concentration as inducer. Investigation of soluble and insoluble Hbx was performed at three
different induction temperatures. As insoluble Hbx was overproduced, purification and refolding steps were performed simultaneously according to previous study (Basu and Leong 2012) with modification. The refolded Hbx protein was characterized using reducing and non-reducing SDS-PAGE and thermal shift assay approach.

2. Materials and Methods

2.1. Bacterial Strains, Plasmids and Culture Media

*E. coli* strains, TOP 10 for cloning and BL21(DE3) for protein expression, are maintained at the laboratory of Pharmaceutical Biotechnology, School of Pharmacy, Institut Teknologi Bandung. The plasmid pET16b carrying codon-optimized Hbx gene was obtained as synthetic gene, purchased from GenScript®. The bacteria were grown at 37°C in Luria Bertani (LB) medium containing 100 μg/ml ampicillin and the protein expression was induced by isopropyl β-d-1-thiogalactopyranoside (IPTG).

2.2. Construction of codon-optimized Hbx Gene

The sequence of Hbx gene was obtained from HBV subgenotype B3 deposited in NCBI database accession number KX429626.1. The codon optimization was performed using OPTIMIZER software (http://genomes.urv.es/OPTIMIZER/) and Codon Optimization Tool (https://sg.idtdna.com/CodonOpt), based on the codon preference of *E. coli* and %GC of *E. coli*, i.e. 51.9%. For the insertion to expression vector pET16b, NdeI site was introduced at 5’ end of the Hbx gene and BamHI site was introduced at 3’ end of Hbx gene. Codon-optimized Hbx gene was then obtained as synthetic gene in pET16b. The plasmid was then characterized by migration analysis, restriction analysis using NdeI and BamHI and by nucleotide sequencing.

2.3. Overproduction, Purification and Refolding of Hbx

The Hbx protein was overproduced in *E. coli* BL21(DE3) as fusion protein with His-tag. Recombinant *E. coli* was grown at 37°C, 150 rpm and the expression of Hbx was induced at IPTG concentration of 0.025-0.2 mM. The cells were harvested at four hours after addition of IPTG. For quick check of protein expression, one ml culture was centrifuged at 4,500 g for 10 minutes and then the cell pellet was lysed using SDS-PAGE sample buffer containing 10% DTT. The sample was heated at 95°C for 10 min and then subjected to 12% SDS-PAGE analysis. After determination of optimum IPTG concentration, the expression of soluble or insoluble Hbx was investigated at three different induction temperatures, 17, 25, and 37°C. After 4 hours induction, cells were collected by centrifugation at 4,500 g for 10 minutes and then resuspended in LEW buffer (50 mM NaH₂PO₄, 300 mM NaCl, pH 8.0). Cell lysis was then performed using sonication at 15 sec pulse on and 15 sec pulse off, with 60% amplitude for 10 min. The supernatant and cell debris were then mixed with SDS-PAGE sample buffer and heated at 95°C for 10 min. Analysis of the soluble and insoluble fraction was performed using 12% SDS-PAGE.

The inclusion bodies containing Hbx protein were washed sequentially using Triton X-100 and Tris-HCl buffer, followed by solubilization in 8 M urea. The solubilized inclusion bodies were then subjected to purification and refolding. Purification of Hbx protein was performed using Ni-NTA column due to presence of His-tag in the Hbx protein. Refolding step was performed prior to elution using 2 M urea, 0.1 mM GSH, 0.01 mM GSSG with variation of arginine, 0.1 and 0.25 M. First refolding step was performed using 3 ml refolding buffer and second refolding step was performed using 10 ml refolding buffer with incubation at 4°C for 18 h. The protein was then eluted using 2 M urea, 0.25 M arginine, 0.1 mM GSH, 0.01 mM GSSG and 100 mM imidazole.

2.4. Characterization of Refolded Hbx

The refolded Hbx protein was then analyzed using reducing and non-reducing 12% SDS-PAGE. In the non-reducing SDS-PAGE, the protein was mixed with SDS-PAGE sample buffer without the presence of β-mercaptoethanol. Prior to analysis using thermal shift assay, the sample was subjected to buffer exchange using 2 M urea and 0.25 M arginine. The Hbx protein was then mixed with Sypro Orange and the melting curve profile was obtained using qPCR machine at 20-95°C.
3. Results

Construction of codon-optimized Hbx gene was performed based on amino acid sequence from HBV subgenotype B3. Codon optimization was performed according to the %GC and codon preference of E. coli. The required Codon Adaptation Index (CAI) is >0.7, with %GC close to %GC E. coli, 51.06%. The Hbx gene sequence was first optimized using OPTIMIZER and the result showed CAI of 0.641 and %GC of 62.1%. The sequence was further optimized manually using Codon Optimization Tool and the result showed CAI 0.705 with %GC of 50.9%, which met the requirements. The codon-optimized Hbx gene sequence showed 72.04% similarity to the original sequence (Figure 1A), while the protein sequence deduced from both sequences showed 100% similarity (Figure 1B). NdeI site and BamHI site were added to the 5' and 3' end of the sequence, respectively, to allow insertion into pET16b. Characterization of recombinant plasmid was performed using migration analysis and restriction analysis. Migration analysis showed that pET16b-Hbx has shorter migration as compared to pET16b (Figure 1C). Restriction analysis using Ncol and BamHI showed two fragments with experimental size of 5818 and 541 bp (Figure 1D). The results showed correct characters of recombinant plasmid, which were confirmed using nucleotide sequencing (data not shown).

Optimization of Hbx overproduction was performed upon various concentration of IPTG as inducer, 0.025-0.2 mM, and the total protein was analyzed using SDS-PAGE. The result showed presence of thick band at 18.4 kDa which was already induced using the lowest concentration of IPTG, which was not present in non-induced sample (Figure 2A). This size was approximately corresponding to the theoretical size of Hbx, which is 17 kDa. IPTG at 0.075 mM was then chosen for further experiments. Investigation of the solubility of the expressed Hbx was performed by fractionation of cell lysate soluble and insoluble fraction. Cell was grown to the mid-log phase and then induced with 0.075 mM IPTG for 4 hours at 17, 25, and 37°C. Cells were then lysed by sonication and the supernatant and cell debris were subjected to SDS-PAGE. The result showed that the Hbx protein was located at the cell debris at all growth and induction conditions (Figure 2B). The result conveyed that insoluble Hbx was produced and required further treatment.

The inclusion bodies containing Hbx protein were then washed and then solubilized in 8 M urea. The denatured protein was then purified using Ni-NTA column due to the presence of His-tag. Refolding steps was performed simultaneously with purification to generate the disulfide bonds present in Hbx. Initially, refolding step was performed using refolding buffer containing 2 M urea, 0.1 mM GSH, 0.01 mM GSSG, and 0.25 M arginine. According to the previous study by Basu and Leong, which used 2 M urea, 0.1 mM GSH, 0.01 mM GSSG with 0.25 M arginine as refolding buffer. Our results showed that the use of this refolding buffer resulted in elution of most Hbx protein during refolding steps. Therefore, only small amount of Hbx was present during elution (Figure 3A). Based on this result, the composition of refolding buffer was modified by reducing the concentration of arginine into 0.1 M. The result showed that only small amount of Hbx protein was washed out during refolding step, while most were obtained at elution stage with relatively high purity (Figure 3B).

Characterization of the refolded Hbx protein was performed using SDS-PAGE and thermal shift assay. Firstly, SDS-PAGE analysis was performed at reducing and non-reducing conditions. In non-reducing condition, beta-mercaptoethanol was omitted and the result may give information on the presence of disulfide bond. The result showed that after refolding the refolded Hbx protein had longer migration as compared to the denatured Hbx (Figure 4A). These results conveyed that the disulfide bonds were successfully formed during refolding stage. Secondly, thermal shift assay was employed to characterize the refolded Hbx protein. Prior to refolding, the Hbx protein showed high fluorescence intensity as compared to the refolded Hbx (Figure 4B). This result showed that after refolding, the hydrophobic residues of Hbx protein were not exposed at the protein surface. From the first derivative curve, the melting point of denatured Hbx was 57.5°C, while the melting point of refolded Hbx was 53.8°C (Figure 4C).
Figure 1. Construction of Hbx synthetic gene and characterization of recombinant plasmid. (A) Alignment of Hbx gene sequences from database and the synthetic construct, (B) alignment of Hbx protein sequences deduced from the database and the synthetic construct, (C) characterization of recombinant pET16b_Hbx using agarose gel electrophoresis, (D) restriction analysis of pET16b_Hbx using Ncol and BamHI. Circular and linear plasmids were analyzed using 1% agarose gel electrophoresis. M: 1kb DNA marker.
Figure 2. Overproduction of recombinant Hbx in *E. coli* BL21(DE3). (A) Expression of Hbx upon induction by 0.025-0.2 mM IPTG for 4 hours, (B) expression of Hbx protein in the supernatant of cell lysate and in the cell debris at 17, 25, and 37°C. Protein analyses were performed using 12% SDS-PAGE. M: protein marker, NI: non-induced, I: induced.

Figure 3. Purification and refolding of Hbx protein from inclusion bodies. The solubilized inclusion bodies was purified using Ni-NTA column and refolded using refolding buffer containing 0.25 (A) and 0.1 M arginine. Protein analyses were performed using 12% SDS-PAGE. M: Protein marker, C: crude, R1: refolding step 1, R2: refolding step 2, E: elution.
4. Discussion

The Hbx protein of HBV is one factor thought to be important in development of HCC in chronic hepatitis B infection. Hbx protein is known for its activity in modulating MAPK and JAK1 signal transduction, Wnt/β-catenin, NF-kβ, and ERK/JNK transactivation, epigenetic regulation of RAR-β2, E-cadherin and p16, as well as modulation of cell cycle regulator protein, such as p21, p15, p16, p27, and Cyclin E (Ali et al. 2014). Hbx protein forms stable secondary structure with alpha-helics, beta-sheet and random coil with four disulfide bonds (Liu et al. 2009; Sidhu et al. 2014). Hbx has BH3-like motif at its carboxy terminus that is responsible for its interaction with Bcl-2 and Bcl-XL. Interaction of Hbx with Bcl-2 protein family increases cytosolic calcium concentration, apoptosis and viral DNA replication (Jiang et al. 2016). Hbx also interacts with carboxy terminus of p53, tumor suppressor protein, causing inhibition of p53 entry into nucleus. Therefore, preventing induction of proteins regulated by p53, such as induction of apoptosis (Wu et al. 2010).

To date, three-dimensional structure of Hbx has not been resolved, only BH3-like motif of Hbx in complex with Bcl-2 has been resolved. This causes difficulty in studying mechanism of interaction with other cellular protein, as well as screening for potential inhibitors.

In order to study the three-dimensional structure of Hbx that will provide essential information about its interaction with other molecules, the overproduction of Hbx is needed. To date, E. coli is known as the simplest, quickest, and most economical host. Besides, the genome has been fully revealed and a huge number of modification has been performed (Hayat et al. 2018). In this study, not only E. coli was utilized as the host of choice, but codon optimization was conducted as a strategy to increase protein yield. However, production of more complex proteins, E. coli brought about several difficulties, one of which is the issue of protein insolubility, resulting in obtaining protein as inclusion bodies rather than the soluble one. More than half of the amino acids in Hbx protein are hydrophobic. It also has four disulfide bonds in total. Such structure tends to be produced as inclusion bodies in E. coli expression system (Liu et al. 2009). Several studies did produce Hbx as soluble protein with the help of large protein fusion to improve the folding (Kay et al. 1985; Liu et al. 2009). In this study, different approach was applied; no long fusion protein was added, but a short histidine tag used for protein purification. We were focusing at
the during-production and downstream processing. Decreasing overproduction temperature to up to 17°C was expected to slow down the production, improve folding, and hence increase the yield of soluble protein, but turned out no improve was observed. Despite the abundance of target protein being produced as inclusion bodies, it required additional steps until soluble functional proteins were obtained.

The adaptation of previous study was not suitable for this system as the protein was washed from the column during refolding process done simultaneously with the purification process. Arginine as aggregation inhibitor was known to interact with tryptophan side chains; however, Hbx itself has only one tryptophan amino acid (Tsumoto et al. 2008). Lower concentration of arginine fixed the condition, resulting in soluble Hbx eluted after the refolding process. Too much arginine in the system could possibly interfere the histidine tag with Nickel column. SDS PAGE and thermal shift assay results showed that disulfide bonds were formed, and more hydrophobic residues were kept inside the tertiary structure of Hbx as the result of protein solubilization and refolding. Further, the confirmation of proper refolding should be done. There are several methods of choice available, such as Circular Dichroism to analyze the secondary structures, ELISA to confirm antibody anti-HBx recognition, and also in vitro activity assay by observing its ability to trans activate a type of cis-acting regulatory element (Kay et al. 1985; Jameel et al. 1990; Liu et al. 2009). Complete confirmation studies could then be followed by the determination of its three-dimensional structure that will be very beneficial of the study of its interaction with other molecules, especially protein.

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