Cloning and characterization of a new site specific methyl-directed DNA endonuclease EcoBLI recognizing 5’-G(5mC)NGC-3’/3’-CGN(5mC)G-5’

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ABSTRACT

A gene coding BisI, site specific 5mC-directed DNA endonuclease recognizing DNA sequence 5’-G(5mC)NGC-3’/3’-CGN(5mC)G-5’, was recently identified in the sequenced genome of the strain-producer Bacillus subtilis T30. In this work we have undertaken a search of bisl gene homologues among the sequenced genomes of enterobacteria. DNA analysis has revealed a small group of highly homologous ORFs with unknown function including one ORF in DNA of well-known strain E.coli. This ORF WP 001276099.1 from E.coli BL21 (DE3) was amplified and cloned. An obtained recombinant strain E.coli pEcoBLI produces MD-endonuclease named EcoBLI. The new enzyme has the same substrate specificity as Bisl MD-endonuclease. Thus, ORF WP 001276099.1 from E.coli BL21 (DE3) encodes site-specific 5mC-directed DNA-endonuclease EcoBLI recognizing and cleaving DNA sequence as indicated by arrows 5’-G(5mC)NGC-3’/3’-CGN(5mC)G-5’.

INTRODUCTION

Site specific methyl-directed DNA endonucleases (MD-endonucleases) recognize and cleave DNA sequences with 5-methylcytosines (5mC), and don’t cleave unmethylated DNA. Over the past 10 years about 15 prototypes of these enzymes with different recognition sites have been described including BisI[1]. BisI recognizes and cleaves DNA sequence 5’-G(5mC)NGC-3’/3’-CGN(5mC)G-5’ as indicated by arrows. Recently a genome of Bacillus subtilis T30 - producer of BisI has been determined and a gene coding BisI has been identified[2]. BisI and other MD-endonucleases were found in bacteria of different taxonomic groups, but mainly in Microbacteriaceae and Bacillaceae families. We couldn’t detect the MD-endonuclease activity in cell lysates of Enterobacteriaceae strains, due to either an absence or extremely low enzymatic activity in bacteria of this group. In this work we have made an
attempt to find MD-endonuclease in Enterobacteriaceae strain not by biochemical methods, but based on the Enterobacteriaceae genome sequences analysis.

Nowadays, many genomes of enterobacteria are published. This allows us to find a BisI similar enzyme by a search of homologues based on the known amino acid sequence of BisI. We have used a program PSI-BLAST\textsuperscript{[3]} to carry out a homology search by method of successive iterations, at first finding firstly the most similar sequences and then the remote homologues with a low degree of similarity and identity.

Here we describe a search and cloning ecoBLI gene from bacterial genome of well-known strain E.coli BL21 (DE3) with subsequent determination of EcoBLI endonuclease substrate specificity.

**EXPERIMENTAL**

Searching for distant amino acid homologues in Enterobacteriaceae was carried out with using the PSI-BLAST program\textsuperscript{[3]}. Multiple alignment of nucleotide and amino acid sequences was carried out with using of Vector NTI Suite 7\textsuperscript{[4]}.

Amplification and cloning of the gene encoding MD-endonuclease from E.coli BL21 (DE3) strain

Amplification of BisI homologue from DNA of E.coli BL21 (DE3) laboratory strain (the Sequence Identifier in the GenBank database is ACT43858) was carried out using the following primers:

\begin{align*}
\text{Esp-1:} & \text{-} 5'\text{-} \text{CCCCCATATGAGTGCACGTGAAGCATATC} - 3' \\
\text{Esp-2:} & \text{-} 5'\text{-} \text{CGCGGATCCTTAGGGATTACACTGAC} \\
& \text{TGAAACTCTTC} - 3'.
\end{align*}

These primers contain recognition sites for FauNDI and BamHI ENases (underlined) to insert the PCR fragment into a plasmid vector.

The resulting PCR fragment was inserted into pMTL22 plasmid\textsuperscript{[5]} on FauNDI and BglII sites with T4 DNA Ligase treatment by standard cloning procedure\textsuperscript{[6]}. The resulting plasmid was called pEcoBLI. After transformation of E.coli ER2267 laboratory strain with pEcoBLI plasmid the resulting recombinant clones were seeded on Petri dishes with LB agar supplemented with ampicillin (50 µg/mL). The clones were grown overnight at 37°C and then subcultured onto separate plates with ampicillin (100 µg/mL) and again incubated overnight for further analysis. Crude lysates of obtained clones were tested for the presence of 5mC-directed DNA endonuclease activity. For further work one of the clones of E.coli pEcoBLI that gave the best results on the hydrolysis of methylated substrate plasmid was selected.

The analysis of enzymatic activity in E.coli recombinant clones with pEcoBLI was performed by adding the lysate of bacteria into 20 µL of reaction mixture containing 1 µg of pFsp4HI3 plasmid DNA\textsuperscript{[7]} previously linearized with Dril restriction enzyme (pFsp4HI3/Dril) in SE-buffer «B» (10 mM Tris-HCl, (pH 7.6 at 25°C), 10 mM MgCl\textsubscript{2}, 1 mM DTT) and incubating the reaction mixture for 1 hour at 37°C. Bacterial lysate was obtained by suspending ~ 0.5-1 µL of biomass grown on a Petri dish in Lysis Buffer (10 mM Tris-HCl, pH 8.5, 0.1 mg/mL Lysozyme, 0.5 M NaCl, 1 mM EDTA, 1 g/L Triton X-100). The products of digestion were determined by electrophoresis in 10 g/L agarose gel.

**The strain producer growth**

The recombinant E.coli strain producer transformed with pEcoBLI was grown in the flasks containing 200 mL of LB broth (10 g/L Tryptone, 5 g/L Yeast extract, 5 g/L NaCl, pH 7.5) with 100 µg/mL ampicillin and cultured in a shaker at 30°C and stirring at 150 rev/min to later log phase, then induced with 0.5 mM IPTG for 4 hours. The cells were precipitated by centrifugation at 5,000 rpm for 30 min at 4°C in the centrifuge J2-21 (“Beckman”, USA), rotor JA-10. The cells pellet was stored at -20°C.

**Enzyme purification**

EcoBLI enzyme preparation was purified by one chromatographic step on Heparin-agarose (“Sigma”, USA). All purification steps were carried out at 0-4°C. Frozen cell paste was thawed with 50 mL of Buffer A (10 mM Tris-HCl (pH 7.5), 0.1 mM EDTA, 7 mM 2-mercaptoethanol) containing 1 mM Phenylmethylsulfonyl fluoride (PMSF), 0.1 mg/mL Lysozyme and disrupted by sonication on the ultra-
sound desintegrator Soniprep 150 (MSE, UK) using five 1 min impulses with 1 min intervals. Cell debris was removed by centrifugation at 15,000 rpm for 30 min in the centrifuge J2-21 (“Beckman”, USA), rotor JA-20. The supernatant was loaded onto Heparin-agarose column (20 mL) equilibrated with Buffer A with 0.2 M NaCl and washed with two volumes of Buffer A with 0.2 M NaCl. Adsorbed material was eluted with 500 mL of linear gradient (0.2-1 M NaCl) in Buffer A. The fractions containing a peak of EcoBII activity were collected and dialyzed against 300 mL of Buffer A with 550 g/L Glycerol for 16 hours. The purified enzyme preparation was stored at -20°C. Target enzyme activity assay in chromatographic fractions was performed by addition of 2 µL aliquots from the fractions to 20 µL of reaction mixture containing pFsp4HI3/DrlI in SE-buffer «B» for 1 hour at 37°C. The products of digestion were determined by electrophoresis in 10 g/L agarose gel.

Sequencing

Determination of the nucleotide sequence was carried out by Sanger-method on the automated sequencer ABI 3130xI Genetic Analyzer (“Applied Biosystems”, USA) according to manufacturer’s recommendations.

Enzymes preparations, DNA, deoxynucleoside triphosphates and synthetic oligonucleotides, as well as molecular weight markers “1 kb DNA Ladder” used in the work, were from “SibEnzyme Ltd.” (Russia). For dilution of enzyme preparations the SE Buffer «B100» (10 mM Tris-HCl, (pH 7.6 at 25°C), 50 mM KCl, 0.1 mM EDTA, 200 µg/mL BSA, 1 mM DTT, 500 g/L Glycerol) was used.

RESULTS AND DISCUSSION

BisI MD-endonuclease distant homologues search in databases of enterobacteria protein sequences

A search of the amino acid sequences homologs of MD-endonuclease BisI (Genbank AJW87312)[2] among Enterobacteriaceae putative proteins was made with PSI-BLAST program. After two rounds of iterations fifty amino acid sequences with unknown functions from enterobacteria...
ria strains displaying a homology (~30-50% similarity and ~10-12% identity) to the BisI protein sequence were found. Among 50 amino acid homologues we have revealed four very similar hypothetical proteins 143-144 amino acid residues in length belonging to *Escherichia* (ACT43858 and AKN48098), *Cronobacter* (CCJ93299) and *Klebsiella* (KEG36084). The amino acid sequences alignment of BisI MD-endonuclease and a group of four homologous putative proteins is presented in Figure 1.

The nucleotide sequences of corresponding genes have been extracted from Genbank database and compared. A comparison of DNA structure of these genes reveals 93-99% identity of the sequences. Surprisingly, *Escherichia* (ACT43858) turned out to be a well known laboratory strain *Escherichia coli* BL21 (DE3) used in many laboratories for genetic engineering experiments. In our experiments we couldn't isolate any MD-endonuclease from biomass of *Escherichia coli* BL21 (DE3) cells. We have cloned open reading frame (ORF) WP_001276099 from *Escherichia coli* BL21 (DE3) to determine this ORF function. Structure of Esp-1 and Esp-2 primers complemented to 5’ and 3’ ends of the ORF WP_001276099.1 is provided in the methods.

**Amplification and gene cloning of EcoBLI MD-endonuclease**

The target ORF (WP_001276099.1) was successfully amplified from *Escherichia coli* BL21 (DE3) DNA. After elution from agarose gel, the amplified fragment 432 bp in length was treated with FauNDI and BamHI ENases and ligated with pMTL22 vector, previously digested with FauNDI and BglII. After transformation of *E. coli* ER2267 cells with a ligation mixture a recombinant *E. coli* strain was obtained. Determination of MD-endonuclease activity in the cell lysate of this strain was obtained. Determination of MD-endonuclease activity in the cell lysate of this strain was carried out with pFsp4HI3 plasmid DNA carrying methylated sites 5’-G(5mC)NGC-3’/3’-CGN(5mC)G-5’ [7], previously linearized with DrI (pFsp4HI3/DrI). DrI ENase cleaves the plasmid at the unique site 5’-GACNNNNNGTC-3’. The result of this experiment is presented in Figure 2. Unlike the original *E. coli* BL21 (DE3) the recombinant strain produced MD-endonuclease called EcoBLI. A recombinant strain was named *E. coli* pEcoBLI. From the above described experiments we can conclude that ORF WP_001276099.1 is coding MD-endonuclease EcoBLI but this ORF is silencing in original *E. coli* BL21 (DE3) strain.

**Determination of EcoBLI substrate specificity**

To determine a substrate specificity of the new enzyme we have made the treatment of pFsp4HI3/ DrI DNA with EcoBLI and control BisI (0.5 units of activity in μL, u.a./μL) enzyme preparations. This plasmid comprises the gene coding Fsp4HI DNA Methyltransferase (MTase) resulting in the first cytosines occur to be methylated in all sequences 5’-GCNGC-3’ on both strands [8]. pFsp4HI3 also contains three hypermethylated DNA regions due to the
presence of two or three overlapped Fsp4HI MTase recognition sites. The resulting data of this DNA cleavage by 2\textsuperscript{\text{L}} of EcoBLI and BisI are shown in Figure 3. This figure shows that 2\textsuperscript{\text{L}} of EcoBLI cleaves pFsp4HI3/DraI (lane 2) forming DNA fragments of the same lengths that BisI does (lane 3). Thus EcoBLI as well as BisI recognizes and cleaves the sequence 5'\textsuperscript{-}GCNGC-3' when containing at least two 5-methylcytosines and has specific activity.

Figure 4 demonstrates the results of comparative digestion of pFsp4HI3/DraI DNA with diluted EcoBLI and BisI enzyme preparations. The substrate DNA was incubated with aliquots of EcoBLI and BisI consistently diluted in 2 times (from 2 up to 1/64 \text{ \mu L}) with the Buffer “B100”.

As can be seen from this figure the results of pFsp4HI3/DraI digestion with EcoBLI and BisI are identical when enzyme’s quantities are maximal (2 \text{ \mu L} of the enzyme with concentration 0.5 u.a./\text{ \mu L}), that is EcoBLI can digest BisI recognition site containing two 5-methylcytosines: 5’-G(5mC)NGC-3’/3’-CGN(5mC)G-5’). But at the same time the pictures of hydrolysis with diluted enzyme preparations have certain differences. BisI proved to have an equal efficiency of hydrolysis its recognition site containing two, three or four 5-methylcytosines\cite{1} whereas EcoBLI has the definite preferences.

As was mentioned above pFsp4HI3 plasmid comprises two sequences 5’-GCNGCNGC-3’ and one sequence 5’-GCNGCNGCNGC-3’ which contain two and three overlapped Fsp4HI MTase recognition sites respectively\cite{8}. So as a result of Fsp4HI methylation the sequence 5’-GCNGCNGCNGCNGC-3’/3’-CGNCGNCGCNGC-5’ comprises two 5’-GCNGCNGCNGCNGC-5’ sites with three 5-methylcytosines and the sequence 5’-GCNGCNGCNGCNGCNGC-3’/3’-CGNCGNCGNCGCNGC-5’ comprises one central 5’-GCNGCNGCNGCNGCNGC-5’ site with four 5-methylcytosines and two sites with three 5-methylcytosines. The analysis of the pFsp4HI3 pri-
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mary structure conducted by using the Vector NTI Suite 7 showed that the resulting cleavage pFsp4HI3/Drl at only three hypermethylated sites leads to appearance of ~3000 ~ 490 (double fragment) and ~340 bp DNA fragments. As it is seen from Figure 4 these fragments were well visualized using 1-1/8 μL of EcoBLI (lanes 2-5, left) as opposed to BisI. In the case of BisI these fragments are visualized worse – only with using 1/2-1/4 μL of the enzyme (lanes 3-4, right). These fragments disappear completely in the case of DNA substrate treatment with 2 μL of EcoBLI or 2 and 1 μL of BisI.

These data suggest that unlike BisI new MD-endonuclease EcoBLI cleaves the variants of the sequence 5’-GCNGC-3’/3’-CGNCG-5’ with three or four 5-methylcytosines better if compared with the case of methylation only two cytosines.

As expected EcoBLI like BisI didn’t cut DNA substrates methylated with following MTases: HpaII (5’-C(5mC)GG-3’), MspI (5’-(5mC)CGG-3’) and Mml.Hgal+M2.Hgal 5’-GA(5mC)GC-3’/5’-G(5mC)GTC-3’) (data not shown).

Therefore a substrate specificity of the new enzyme EcoBLI from E.coli BL21 (DE3) is similar to one of MD-endonuclease BisI from B.subtilis T30.

Determination of DNA position hydrolysis for EcoBLI.

Determination of DNA position hydrolysis for EcoBLI was carried out by comparing of EcoBLI, Pkr and BisI digestion products of oligonucleotide duplex D1/D2, formed from synthesized oligonucleotides D1 and D2 (estimated EcoBLI recognition sequence is underlined):

D1: 5’-GAGTTTAG(5mC)GG(5mC)TATCGATCC-3’
D2: 5’-GGATCGATAG(5mC)CG(5mC)TAAACTC-3’.

Figure 5 presents the autoradiograph of the labeled duplex *D1*/D2 digested with the enzymes after the electrophoresis in 200 g/L polyacrylamide gel with 7M urea.

Figure 5 shows that DNA fragments formed by Pkr and EcoBLI digestion (lanes 2 and 3, respectively) of D1*/D2 duplex have different electrophoretic mobilities suggesting that these enzymes have different hydrolysis positions relatively to the recognition site. At the same time the electrophoretic mobilities of DNA fragments formed by BisI and EcoBLI are identical (lanes 3 and 4, respectively). Thus, EcoBLI and BisI have the same hydrolysis positions with respect to the recognition sequence 5’-GCNGC-3’.

CONCLUSIONS

A search of bisI gene homologues among the sequenced genomes of enterobacteria has been made and ORF WP 001276099.1 from a well-known strain E.coli BL21 (DE3) was amplified and cloned. An obtained recombinant strain E.coli pEcoBLI produces MD-endonuclease named EcoBLI with the same substrate specificity as BisI MD-endonuclease. EcoBLI has a recognition sequence 5’-G(5mC)^NGC-3’/3’-CG(5mC)G-5’ and cleaves it before the central nucleotide “N” in both strands of DNA forming single-nucleotide 5’-protruding ends. We identified ORF WP 001276099.1 from the strain E.coli BL21 (DE3) as a silenced gene coding MD-endonuclease.

We believe that EcoBLI as well as BisI MD-endonucleases can be used in epigenetics for detec-
tion and analysis of methylation pattern in mammalian DNA, which usually contains a considerable number of 5-methylcytosine bases. Such type of DNA methylation is known to play a significant role in the regulation of genes expression[9].

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DECLARATION

The manuscript is original and is not published or communicated for publication elsewhere either in part or full

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