Cloning and over-expression of Penicillin G acylase in 
*Escherichia coli* BL21

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Penicillin G acylase (PGA) is one of the most important enzymes in the pharmaceutical industry. It is utilized in the process for production of semi-synthetic penicillins. Several different penicillin acylases with various characteristics have been isolated from different bacteria and identification of bacterial isolates harboring PGA enzyme with higher industrial compatibilities is of high interest. The aim of this study is to screen for PGA producing *Escherichia coli* isolates as well as the cloning and recombinant expression of PGA for high level enzyme production. Bacteria isolated from environmental and clinical samples were identified by standard microbiological tests and then *E. coli* isolates were subjected to DNA extraction and PCR screening using primers designed on conserved region of PGA genes. The PCR product from a positive isolate were cloned and subjected to sequencing. The gene encoding for full length PGA was expressed in *E. coli* under the T7 promoter. PCR screening identified several PGA positive *E. coli*. One of the positive isolates was cloned in pGEM -T easy vector. Sequencing of the cloned gene revealed that the gene encoding Penicillin G acylase from this wild *E. coli* isolate contains an open reading frame of 2538 nucleotide encoding 846 amino acids. Analysis of the sequencing results showed that the PGA gene is highly conserved among *E. coli* strains. Recombinant expression of PGA from wild isolate in *E. coli* BL 21 resulted in a high level expression of recombinant PGA which appeared as a dense band in SDS-PAGE analysis of induced culture.

**Key words:** PGA, cloning, sequencing, recombinant expression.

**INTRODUCTION**

Traditionally, semi-synthetic penicillins were produced by chemical synthesis that needs a series of complex chemical reactions at low temperatures, using toxic compounds. These undesirable solvents need to be removed and there is a need for a rigid quality control of the downstream purification steps. In recent years, chemical synthesis has mostly been replaced by less polluting biological methods using enzymes isolated from different microorganisms (Shewale et al., 1990; Demain, 2000). These enzymatic reactions work in aqueous medium and physiological conditions without producing toxic wastes (Van Langen et al., 2001).

Penicillin G acylase (PGA) is a type II penicillin acylase that hydrolyzes Penicillin G to 6-aminopenicillanic acid (6-APA) and phenyl acetid acid (PAA) (Ohashi et al., 1989; Zietkiewicz et al., 1994). 6-APA is starting material for production of semi-synthetic penicillins (Kumar et al., 2007; Ochman et al., 1988). PGA has been found in numerous bacteria and the PGA of *Escherichia coli* has been extensively studied (Parmar et al., 2000).

Although the exact function of PGA in free-living *E. coli* have not been well understood, it is thought to act as a

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**Abbreviations:** PGA, Penicillin G acylase; 6-APA, aminopenicillanic acid; PAA, phenyl acetic acid; EMB, eosin-methylene blue; SDS-PAGE, sodium dodecyl sulfate polyacrylamide gel electrophoresis; PCR, polymerase chain reaction; IPTG, isopropyl-β-D-thiogalactopyranoside; BLAST, basic local alignment search tool; PDAB, p-dimethyl-aminobenzaldehyde.
scavenger enzyme for various natural esters and amides of PAA and its derivatives, such as hydroxyphenylacetic acid (Díaz et al., 2001). Thus, when *E. coli* encounters phenylacetylated compounds, the periplasmic PGA degrades them to PAA and this moiety then diffuses into the nucleoplasm.

The characteristics of PGA isolated from different biological and environmental sources were found to be varied in different aspects including, substrate specificity, optimum pH, temperature tolerance etc. Therefore, microorganisms have been extensively screened for isolation of novel penicillin acylases with higher compatibility with industrial deacylation requirements. In recent years, recombinant DNA technology has emerged as a potent technology for high level production of many useful proteins. In this method, desired and increased yield are achieved by proper vector and host cell selection. Production of PGA by recombinant technology could have several advantages. Firstly, recombinant strain could express higher level of recombinant protein compared to native strains. Secondly, the expression of PGA in native *E. coli* is under a complex control circuit (Wang et al., 2004), whereas in recombinant *E. coli* the expression could be under tight control. The objective of this study was to screen *E. coli* isolates from clinical and environmental samples for PGA. The gene from a positive clone was cloned and the T7 promoter and BL21 host cell were investigated for high level production of active PGA.

**MATERIALS AND METHODS**

**Bacterial strains, media, and plasmids**

The *E. coli* strains, DH5α (Invitrogen, Carlsbad, CA) and BL21 (DE3) (Novagen, USA) were used for cloning and protein expression, respectively. Strains were cultured at 37°C in Luria-Bertani (LB) broth. Ampicillin was used at a concentration of 100 μg/ml for selecting transformed bacteria. The plasmid pGEM-T Easy vector (Promega) was used for cloning of the amplified Penicillin G acylase gene (PGA) and the plasmid pET 22 b was used for protein expression.

**Isolation of *E. coli* from samples**

In this study, *E. coli* strains isolated from water, soil and clinical specimens were screened for PGA. A total of 280 specimens were collected from water, soil and clinical specimens. Samples were transported to the laboratory and then standard microbiological tests including culture on eosin-methylene blue (EMB) agar, gram staining and biochemical tests like Oxidase, Indole, Voges-Proskauer (VP), methyl red (MR) and Citrate tests (IMVIC) (Forbes et al., 2006) were carried out for identification of isolates. Only 1 isolate was selected from each sample. The organisms were maintained on nutrient agar (Difco) slants at 4°C until they were used.

**DNA extraction and PCR screening**

Isolates were cultured in LB broth overnight at 37°C with vigorous shaking and the bacteria were pelleted by centrifugation at 9000 rpm. DNA isolation was carried out essentially as described (Montazam et al., 2009) with slight modification. Briefly, the bacterial pellet was resuspended in TE buffer (Tris 10 mM, PH 8, EDTA 1 mM) containing SDS (1%) and proteinase K (10 mg/ml) and incubated at 55°C for 2 h. After that, an equal volume of phenol-chloroform (1:1) was added, incubated at room temperature for 5 min and centrifuged for 5 min at 10000 rpm. The aqueous phase was transferred into a new tube, incubated with RNase A (10 mg/ml) for 20 min at 37°C and then DNA was precipitated by adding 2.5 volume of 100% cold ethanol. The pellet was then washed with 70% ethanol and dissolved in distilled water after drying. DNA extracted from *E. coli* isolates were subjected to PCR amplification with primer pair PGA F1, 5’- TGTGGCGGATGA TATTGTG- 3’ and PGA R1, 5’- GCGGAGTGTACGAA-3’. Primers were selected based on conserved regions of PGA genes that have already been reported. PCR amplification was carried out in a final volume of 25 μl containing 20 mM Tris-HCl (pH 8.3), 50 mM KCl, 2 mM MgCl₂, 18 mM NaCl, a 0.2 mM concentration of each 2’-deoxynucleoside triphosphate (dATP, dCTP, dTTP and dGTP), a 0.4 μM concentration of each forward and reverse primer and 1 U of Taq DNA polymerase (Fermentas, Litany). Thermocycler conditions for PCR were one cycle of 94°C for 4 min, 32 cycles of 94°C for 45 s, 50°C for 30 s and 72°C for 60 s and one cycle of 72°C for 5 min followed by a hold at 10°C. The PCR results were assessed by electrophoresis on 1% agarose gel and visualized by staining with ethidium bromide (EtBr).

**Cloning and sequencing of full length PGA gene**

After identification of PGA producing strains, PCR amplification was performed on a positive strain using primers F2, 5’-TGGCCCATGA AAAATAGAAATCGTATGATC and R1, 5’-GCTCGAGTCTCTGA GCTCGAGTCTCTGA which were complementary to 5’ and 3’ end of full length PGA gene, respectively. The PCR product was purified with a PCR purification kit (Qiagen) according to the manufacturer’s instructions and used for T-A cloning using pGEM-T easy cloning kit (Promega) which yielded plasmid clone pGEM- PGA. The reaction was transformed into an *E. coli*, DH5α competent cells and a positive clone was submitted for sequencing with M13 forward and reverse primers. Nucleotide and predicted amino acid sequences were compared to data available by the BLAST search method.

**Recombinant expression of PGA in *E. coli***

The coding region for full length PGA gene was isolated by digestion of pGEM-PCA clones with Msc I-Xho I restriction enzymes and subcloned into the Msc I-Xho I site of the pET 22b expression vector (Novagen) in frame with a carboxy-terminal six histidine tag, yielding the plasmid subclone pET22b-PCA. Sequencing with vector-based primers demonstrated that the His-tag fusion was in frame. The construct was transformed into *E. coli* BL21 strain, cultured at 37°C in LB medium containing 100 μg ml⁻¹ ampicillin to an absorbance at 600 nm (A₆₀₀) of 0.7 unit. Protein expression was induced by adding isopropyl-D-thiogalactopyranoside (IPTG) to a final concentration of 0.5 mM and analyzed by SDS-PAGE.

**Biologic assay for PGA production by recombinant Bacteria**

*E. coli* BL21 containing pET 22b-PCA construct was cultured in media I (1% Tripton, 0.3% Beef extract and 0.15% Phenyl acetic acid) to an absorbance of 0.8 and induced by addition of IPTG (1 mM). After two hours, the cells were collected by centrifugation and PGA production was estimated by the p-dimethyl-aminobenzaldehyde (PDAB) method (Shewale et al., 1987). Briefly, the cells
were resuspended in 0.1 M phosphate buffer, pH 7.5 and incubated for half of an hour with 5 mg of penicillin G. The reaction was stopped by the addition of 3 mL of a (1:1) mixture of 0.05 M NaOH - 20% acetic acid and centrifuged. Production of 6APA was estimated by incubation of supernatant with pDAB. One unit of enzyme activity was defined as the amount of enzyme required to produce 1 μmol 6-APA per minute at 37°C.

RESULTS

PCR screening for PGA producing E. coli isolates

In this study, screening of 280 specimens (environmental and clinical specimens) for E. coli resulted in isolation of 180 E. coli bacteria. PCR screening of E. coli isolates for PGA production resulted in identification of 5 positive strains.

Amplification, cloning and sequencing of PGA

PCR amplification of E. coli genomic DNA with primers designed on conserved region of PGA gene resulted in a PCR band of 815 bp. Sequencing of the PCR products confirmed their identity as a part of PGA gene. The nucleotide sequence was deposited in NCBI Gene Bank under accession number HM011571. Amplification of full length PGA produced a PCR product of about 2500 bp (Figure 1). The PCR product was ligated to the pGEM-Teasy vector and cloning was verified by PCR and restriction digestion (Figure 2). Sequencing of cloned gene revealed that the gene consisted of 2538 bp encoding a protein of 846 amino acids. Multiple alignment of predicted amino acid sequence with PGA sequences in data base revealed a homology ranging from 90 - 94%. This shows that PGA sequence is highly conserved among E. coli strains.

Recombinant expression of PGA in E. coli

The insert containing the full length PGA was subcloned into the pET 22 b expression vector (Novagen) in frame with a C-terminal His-tag fusion for affinity purification (Figure 3). Induction with IPTG of E. coli BL21 transformed with expression cassette resulted in a high expression of recombinant protein as appeared in SDS-PAGE analysis of lysate of induced bacteria. Figure 4 shows Commassie blue-stained SDS-PAGE gels of E. coli culture before and after induction along with the bacteria without expression construct.

Biological assay of recombinant PGA expression

To compare the PGA activity of recombinant E. coli BL21 carrying the PGA gene on pET 22 b expression vector
with wild *E. coli* isolate, the bacteria were cultured in media I and induced by addition of different concentration of IPTG. Culture was continued for high density (Lee, 1996), harvested, incubated with penicillin G and assayed for 6APA production. The results showed that induction with 1 mM IPTG can result in a high level of PGA activity by *E. coli* BL21 that were equivalent to 150 unit per gram (wet weight) of recombinant bacteria. This activity was about 300% more than that exhibited by wild type *E. coli*.

**DISCUSSION**

Biological screening methods have been used for isolation and identification of bacteria producing PGA but this method is labor and time consuming. In this project, the PCR technique was utilized for identification of penicillin G acylase producing *E. coli* strains from clinical and environmental samples. We found 5 PGA producing *E. coli* strains among 280 screened isolated. The results of our study showed that this procedure is a simple, rapid and efficient method for identification of PGA producing bacterial strains. This result was in line with previous report on application of PCR technique for identification of 7ACA positive bacterial strains (Luo et al., 2005).

The gene from one positive isolates was cloned in the pGEM-Teasy vector and used for DNA sequencing. Sequence analysis showed the gene composed of 2538 nucleotide encoding 846 amino acids. BLAST analysis of obtained sequence showed that the gene contains 98% homology to previously reported PGA gene from *E. coli* strains. Homology with other reported PGA genes were 96% to *Achromobacter xylosoxidans* (Cai et al., 2004) and 81% homology to *Kluyvera citrophila* (Guisán et al., 1993). The results of our study showed that the PGA gene is highly conserved among *E. coli* strains. The high conservation of PGA gene in *E. coli* isolates may indicate important benefit of this enzyme for survival of these bacteria in different environmental condition. Assay for penicillin G acylase production by *E. coli* BL21 expressing recombinant PGA showed high level of enzyme activity that was 3 times more than PGA activity of wild type *E. coli* strain. These results are consistent with previous reports indicating the advantages of recombinant DNA technology for production of PGA enzyme (Olsson et al., 1985; Garcia et al., 1986; Polderman-Tijmes et al., 2002).

**Conclusion**

This study showed the suitability and benefit of PCR technique for rapid detection of Penicillin G acylase in wild *E. coli* isolates from different origins. The results of our study could expand the application of PCR as a cheap and rapid method for screening of isolates with pharmaceutical and biological important characteristics. The results of this project also showed that recombinant technology could be used for high level production of PGA.
REFERENCES


