An Optimized Medium for Screening of L-Asparaginase production by *Escherichia coli*

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**Abstract:** Purified L-asparaginase II from *Escherichia coli* has been supplied and employed in the acute leukemia and other malignant neoplasms chemotherapy. L-asparaginase II gene (*ansB*) in *E. coli* is under regulation and certain conditions is needed for expression of this gene. In this investigation, the various concentrations of modified M9 medium ingredients and various carbon source were tested to optimize the medium for expression and identification of L-asparaginase in *E. coli*. Finally a semi-quantitative plate assay for L-asparaginase producing *Escherichia coli* is reported.

**Key words:** L-asparaginase, *E. coli*, medium, *ansB*

**INTRODUCTION**

In 1953 Kidd found that guinea pig serum inhibited a number of transplantable lymphomas in mice and rats as well as certain spontaneous and radiation-induced leukemias in mice. Broome has presented some evidence that the antitumor principle in guinea pig serum is L-asparaginase [1]. Deamidation of L-asparagine by extracts of *E. coli* was first reported in 1957 [2]. Later, Mashburn and Wriston observed that L-asparaginase (L-asparagine amidohydrolase, Enzyme Commission 3.5.1.1) purified from cell extract of *E. coli* has an antitumor activity similar to that of guinea pig serum [3]. Although other microorganisms such as *Aerobacter*, *Bacillus*, *Erwinia*, *Pseudomonas*, *Serratia*, *Xanthomonas*, *Photobacterium* [4], *Streptomyces* [5], *Proteus* [3], *Vibrio* [6] and *Aspergillus* [7] have a potential for asparaginase production, purified enzyme from *E. coli* has been supplied and employed in the clinical application for acute leukemia and other malignant neoplasms in human. L-asparaginase belongs to an amidase group that hydrolyses the amide bond in L-asparagine to aspartic acid and ammonia. Unlike the normal cells, neoplastic cells cannot synthesize L-asparagine due the absence of L-asparaginase synthetase. Therefore, they obtain the required asparagine from circulating pools. For this reason, intravenous injection of free enzyme results in selective neoplastic cell death, directly by the depletion of circulating asparagine levels or indirectly from some other metabolite of the asparaginase reaction [7-9].

*E. coli* has two isozymes of L-asparaginase [10], L-asparaginase I (*AnsA*) which is found in the cytoplasm and has a low affinity for L-asparagine (EC1; Km = 3.5 mM) and L-asparaginase II (*AnsB*) that is a periplasmic enzyme used in the treatment of acute lymphoblastic leukemia and in contrast, it is a high-affinity enzyme (EC2; Km = 10 pM). *AnsA* is thought to be constitutively produced but *ansB* gene expression is changed by aeration, carbon source and variation of available amino acids [1,2,11]. In this investigation, an optimized medium for semi-quantitative plate assay and screening of L-asparaginase producing *Escherichia coli* is reported based on modified M9 medium [12]. In this medium, L-asparagine is used as the sole nitrogen source. The production of L-asparaginase by *E. coli* leads to ammonia formation and increase in pH of the medium. A pH indicator (phenol red) makes medium pink around the colonies producing L-asparaginase.

**MATERIALS AND METHODS**

**Isolation and identification of *E. coli*:** Environmental samples were collected from waste water and the Khoshk River, Shiraz, Iran (in February 2008). Clinical
Optimization of medium: The various concentrations of Na₂HPO₄·2H₂O, KH₂PO₄, NaCl, L-asparagine, MgSO₄·7H₂O, CaCl₂·2H₂O, carbon sources and phenol red were tested as ingredients of medium; one factor was changed while others were constant (Table 1). The pH of the media was adjusted at 7.0.

Stock solution of the phenol red was prepared (2.5% in ethanol, pH 7.0), filtered by 0.2 µm cellulose filter and added to autoclaved media. E. coli isolates were inoculated on the prepared plates and incubated at 37°C for 48h. The Modified M9 medium was used as blank. After incubation, L-asparaginase activity were reported as (-), trace or (+) based upon intensity of produced pink color and pink zone diameter.

RESULTS

Studies with different concentrations of ingredients revealed that the most appropriate concentrations in 1 liter of medium are KH₂PO₄ (0.75 g), NaCl (0.5 g), L-asparagine (10 g), Maltose (1 g), agar (17 g) and phenol red (0.05 g). On optimized medium with above concentrations among the 130 isolates of E. coli, 35 isolates produced trace pink zone and 8 isolates produced (+) zone (Fig. 1), while only 4 trace zones were produced on modified M9 medium (Table 2).

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FIG. 1: Pink zone production by L-asparaginase producing E. coli on optimized medium

| Isolation source | No. | Total isolates | Pink zone intensity on the modified M9 medium | Pink zone intensity on the optimized medium |
|------------------|-----|----------------|---------------------------------------------|---------------------------------------------|
| River            | 47  | 44             | 3   | 0   | 24 | 19 | 4 |
| Water            | 35  | 34             | 1   | 0   | 21 | 11 | 3 |
| Stool            | 33  | 33             | 0   | 0   | 30 | 3  | 0 |


discussion

The ansB gene of E. coli is regulated by catabolite activator protein (CAP; also called cyclic AMP receptor protein or CRP)[1,2,11]. The low level of intracellular glucose results in a high level of cAMP, which means high cAMP-CAP complexes. cAMP-CAP binds to a specific site on the DNA that is located adjacent to the promoter for the genes regulated by CAP. The -10 and -35 motifs in the promoter sequence for such genes are not a perfect match to the consensus sequence for σ⁷⁰ promoters. cAMP-CAP bound to the CAP binding site increases the binding of σ⁷⁰ RNA polymerase to the promoter and increases gene expression[13]. Therefore, glucose is an inhibitor for ansB gene. We changed the carbon source of medium from glucose, to disaccharide for reduction in the level of intracellular glucose and more AnsB production; maltose was the best. Also, in contrast to modified M9 medium, Na₂HPO₄ has been eliminated to reduce bufferic properties of medium and little ammonia identification. The optimum concentration of phenol red in the medium is 0.05 mg L⁻¹ which keeps the colour yellow for a better identification of colour change to pink for trace amounts of L-asparaginase. High concentrations of phenol red giving the colour of medium dark red make difficult to identify the pink zone. The observed results encourage us to continue further screening for L-asparaginase production in the other microorganisms (Bacteria and Fungi) on our optimized medium.
CONCLUSION

The optimized medium is a sensitive medium for L-asparaginase detection and trace amounts of produced enzyme by E. coli. This medium can be used as proper medium for other L-asparaginase producing microorganisms.

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