Screening, Cloning and Expression of Active Streptokinase from an Iranian Isolate of \textit{S.equisimilis} Group C in \textit{E. coli}

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\textbf{ABSTRACT}

\textbf{Introduction:} Streptokinase (SK) is a fibrinolytic protein secreted by \(\beta\)-hemolytic streptococci (\(\beta\)HS) groups A, C and G. Due to its importance as a thrombolytic drug, national screening programs in different countries for isolation of \(\beta\)HS and especially SK-producing group C (GCS) strains have been conducted. Herein, we provide data of the first screening study on \(\beta\)HS isolates in Iran for the aim of recombinant SK (rSK) production from a local strain.

\textbf{Materials and methods:} 252 streptococcal samples were collected and characterized using microbial/biochemical assays. The GCS strains were serologically confirmed. Activity of GCS supernatant cultures was determined by caseinolytic assay in comparison with the standard strain GCS9542. The SK gene of the highest producer strain was selected for production of rSK in \textit{E.coli} system. The rSKs activities were determined using chromogenic assay.

\textbf{Results:} \(\beta\)HS were detected in 75 of the collected specimens (29.4%) including groups A (25.8%), C (3.6%) and G (0.4%). Analyses by SDS-PAGE and Western blotting indicated the proper expression of 47 kDa rSK proteins in \textit{E. coli} for SK genes which were cloned from both the selected (GCS\textsuperscript{87}) and standard (GCS\textsuperscript{9542}) strains with the yields of 0.53 and 0.59 mg/ml (of the purified protein), respectively. The calculated activity for rSK\textsuperscript{87} was around 90% of rSK\textsuperscript{9542} activity (0.18x10\textsuperscript{5} IU/mg v/s 0.21x10\textsuperscript{5} IU/mg).

\textbf{Conclusion:} Results of the present study for the first time provided the possibility of producing rSK from a local and native source with comparable yields and activities similar to the standard strain.

\textbf{Introduction}

Pathologic blood clots (in the form of thrombus) can result in vascular blockage which can induce serious consequences including death (1). In a healthy haemostatic system, formation of blood clots is suppressed through conversion of zymogen plasminogen (Plg) to plasmin (the serine protease that degrades fibrin) (2). However, in pathological conditions, clinical intervention through application of plasminogen activators (also known as “thrombolytic or fibrinolytic agents”) to relieve the vein from thrombosis is required. Currently, routine thrombolytic agents in clinical applications are recombinant...
human tissue plasminogen activator (tPA), urokinase (UK) and streptokinase (SK). SK unlike UK and tPA activates Plg indirectly by complex formation and in a fibrin non-specific manner. SK exhibits significantly higher in vivo half-life compared to UK and tPA, but it’s fibrin non-specific mode of action and bacterial origin may increase the risk and side effects of thrombolytic therapies compared to the other two agents. Despite these short comings, SK gained a worldwide acceptance in developing countries due to its half-life, cost-effectiveness and shorter period of therapy. Moreover, a number of large-scale clinical trials, which have been conducted to compare the clinical efficacy of SK and tPA could not indicate a clear preference for either drug. Although most of the group A, C and G beta-hemolytic streptococci (GAS, GCS and GGS respectively) produce and secrete SK, GCS which are neither erythrogenic toxin generators nor very fastidious in growth requirements, are the preferred bacteria for SK production. Historically, GCS strain S. equisimilis H46A (ATCC 12449) was the first streptococci to be introduced as a high-yield SK secreting bacteria (for production aims) by Christensen et al. in 1945. Subsequently, Estrada et al. introduced another S. equisimilis group C (ATCC 9542) as a SK production strain in 1992. These two strains were used for SK production and served extensively as the principal source of SK gene for heterologous expression of the recombinant SK (rSK) in other hosts like E. coli and yeast.

Due to the clinical importance and increasing potential of SK application, a great deal of effort has been directed towards improvement of quality and quantity of SK production. Most of these studies were focused on either optimization of production conditions for S. equisimilis H46A and 9542, or strain development using mutant strains or protein engineering using recombinant DNA technology (8-9). Sequencing studies on the SK genes and proteins from different isolates indicated that they are heterogeneous genes. In fact, the sequence identity of mature SK proteins with the same number of amino acids (414 residues), ranges from 80% to 98% (11). SK heterogeneity may reflect functional diversity of the gene products in pathogenesis, antigenic variation, solubility (12), Plg activation (13) and fibrinolytic activity (14). This implies that, alternatively, it may be possible to isolate SK protein(s) with better fibrinolytic characteristics that have clinical benefits by screening among different streptococci. In this context, a number of national screening programs to isolate SK producing beta-hemolytic streptococci (βHS) from local and regional samples have been conducted in different countries (15-17). However, to our best of knowledge, there is no prior report on screening and characterization of SK producing strains or recombinant expression of SK from Iranian isolated streptococci strains.

In the present study, an attempt was made for isolation and characterization of βHS among Iranian clinical isolates to screen for the best SK producing GCS strains and to clone and express the corresponding SK gene in E. coli, for the final aim of SK production from local and native sources.

**Materials and Methods**

**Bacterial strains and culturing conditions**

Two hundred and fifty two samples (initially assumed as streptococcal samples) were collected from patients with various non-invasive streptococcal diseases from different regions of Iran, during 2006 and 2010 (Table 1). S. pyogenes (GAS) ATCC 10403, S. dysgalactiea spp. equisimilis ATCC 9542 (GCS-9542) and S. dysgalactiea spp. equisimilis (GGS) CIP 55120 (Pasteur institute of Paris) were used as reference strains for presumptive microbial, biochemical and serological tests (Table 2). All streptococcal isolates were cultured on 5% sheep blood agar and Todd-Hewitt Broth (THB) (Difco, USA) media. The plates were incubated at 37°C overnight. Colonies surrounded by alpha or beta-haemolysis were selected for more detailed char-

**Table 1. Characteristics of the bacterial samples identified in this study**

| Streptococci group | Throat culture | Genital tract | Urine culture | Skin | Blood culture | CSF | N.D | Total |
|--------------------|---------------|---------------|---------------|------|---------------|-----|------|-------|
| GAS                | 60            | -             | 4             | 1    | -             | -   | -    | 65    |
| GCS                | 7             | 2             | -             | -    | -             | -   | -    | 9     |
| GGS                | 1             | -             | -             | -    | -             | -   | -    | 1     |
| GBS                | 51            | 20            | 27            | 3    | 1             | 3   | 4    | 109   |
| GDS                | 19            | -             | 8             | 3    | 4             | -   | 2    | 36    |
| Staphylococcus     | 8             | -             | -             | -    | -             | -   | 3    | 11    |
| Other streptococci | 16            | -             | 3             | 1    | -             | -   | 2    | 23    |
| Total              | 162           | 21            | 42            | 7    | 3             | 3   | 11   | 252   |

N.D: not determined
acterization tests (Table 2). Presumptive standard identification tests including catalase test, susceptibility to a 0.04 U bacitracin disk and SXT disk (Sulfamethoxazole 23.75µg –trimethoprim 1.25µg), CAMP test (Christie, Atkins, Munch-Petersen), PYR (Pyrolydonyl arylamidase) test, esculine hydrolysis, 6.5% NaCl tolerance and Voges-Proskauer (VP) tests were performed according to the standard protocols for determination of streptococci group A, C and G. Lancefield serotyping was performed by latex agglutination kit (Mast, UK). GCS Subspecies were further characterized by standard biochemical tests using ribose, sorbitol, lactose and trehalose fermentation (Table 2) (18). DH5α and M15 E. coli cells were cultured in Luria-Bertani (LB) medium. The pQE30 plasmid and M15 E. coli cells (Qiagen, USA) were used for cloning and expression of the SK gene and DH5α E. coli cells was used for propagation of plasmids. Kanamycin (50 µg/ml) was externally added to M15 E. coli culture.

Caseinolytic assay for streptokinase activity of bacterial culture

Activity of streptococcal supernatant cultures was determined by caseinolytic assay(19). Briefly, 50µl of overnight culture of strains in THB was added to 5ml of fresh THB and incubated at 37°C at 150 rpm. The culture supernatants were collected by centrifugation at mid-log phase (OD₆₀₀ of 0.7-0.8) and were filtered using 0.22 µm PVDF filter (Whatman, Germany). The cell-free fluid was used to fill the pre-made wells in medium containing: 5% skim milk and %1 agarose in sterile culturing plates. The same volume of human plasminogen (1 mg/ml) in a buffer containing: 150 mM NaCl and 50 mM Tris-HCl pH 7.4 was simultaneously added to the corresponding wells. Supernatant of S.equisimilis 9542 and THB were included as positive and negative controls in the corresponding wells, respectively. Plates were incubated overnight at 37°C. The clear area around the wells represented the level of SK activity of the corresponding strain.

| Table 2. Microbial and biochemical analyses undertaken for the identification of beta hemolytic streptococci in our study (adapted from ref. 18) |
|---------------------------------------------------------------|
|                  1            2         3        4         5        6        7        8        9      |
| Haemolysis       | β       | β       | β        | β/α     | β       | β       | β       | β     | β/α     |
| Growth in 6.5% NaCl | -     | a       | -        | -       | a       | +       | -       | -     | a       |
| Growth in Bile-Aesculin | -    | -       | -        | -       | -       | -       | -       | -     | +       |
| Voges-proskauer test | -    | +       | -        | -       | -       | -       | -       | -     | +       |
| Pyrrolydonylarylamidase | +    | -       | v        | v       | -       | -       | -       | -     | +       |
| Sensitive to bacitracin | -    | -       | -        | -       | -       | -       | -       | -     | -       |
| H₂O₂ production | -       | -       | -        | -       | -       | -       | -       | -     | -       |
| CAMP             | -       | +       | -        | -       | -       | -       | -       | -     | +       |
| Fermentation of Ribose | -    | +       | -        | +       | v       | +       | +       | +     | -       |
| Sorbitol         | -       | -       | v        | +       | -       | -       | -       | -     | -       |
| Lactose          | +       | v       | -        | +       | +       | v       | +       | v     | -       |
| Trehalose        | +       | -       | +        | -       | +       | +       | v       | +     | +       |
| Lancefield antigen | A     | B       | C        | C       | C       | G       | D       | f/C/A/G |

1: S. pyogenes, 2: S. agalactiae, 3: S. equi, 4: S. dysgalactiae, 5: S. zooepidemicus, 6: S. equisimilis, 7: S. spp. group G (large colony variety), 8: S. spp. group D, 9: S. anginosus.
(a) Green zone around colonies on Blood Agar, (β) Clear, colourless zone around colonies on Blood Agar, (α) some strain will grow in 4% NaCl broth, (v) variable, (+) positive result, (-) negative result.
Isolation of the streptokinase genes and plasmid construction

Genomic DNA of *S. equisimilis* (GCS-9542) and the selected GCS strain that showed the highest level of SK activity (GCS-S87) in caseinolytic assay (Figure1) was isolated by DNA extraction kit (AxyGene, USA) and used as a template for PCR-mediated isolation of SK genes. The coding region of SK gene (lacking the signal peptide sequence) was amplified by PCR using primers with inserted restriction sites for direct cloning into pQE30 vector (forward primer; *BamHI*-SK: 5'-TGGATCCATCGTCGGACCTGAGTGGCTG-3'; reverse primer; *PstI*-SKr: 5'-CGCCCGAATTCTGTTGTTAATG-3'; the sequences corresponding to restriction sites are underlined). The resulting amplified fragments were digested with *BamHI* and *PstI* and cloned into the same sites of pQE30 expression vector in tandem with the fused N-terminally 6XHis-tag and downstream of T5 promoter (Figure 2). Proper expression constructs were confirmed by restriction enzyme analysis and bidirectional sequencing. All cloning steps were performed according to standard procedures (20).

**Protein expression in *E. coli***

*E. coli* M15 cells, which carry multiple copies of pREP4 plasmid that tightly regulate recombinant protein expression (21) were used as an expression host for pQE30 plasmids according to the manufacturer’s protocol (Qiagen, USA). Briefly, after transformation of bacterial cells with the recombinant plasmids pSK9542 and pSK87 using the standard CaCl$_2$ method (20), expression of the target fusion protein was induced at OD$_{600}$ of 0.5–0.6 by isopropyl-β-D-thio-galactoside (IPTG) to a final concentration of 0.5 mM. Cells were harvested by centrifugation after 6 hours of incubation at 37°C and stored at -20°C for purification steps.

**Purification and refolding of expressed SK**

The His6-SK fusion proteins were purified by nickel affinity chromatography under denaturing conditions according to the manufacturer’s protocol (Qiagen, USA). The pellet was dissolved in denaturing binding buffer (8 M Urea, 100 mM NaH$_2$PO$_4$, 100 mM Tris-Cl pH 8.0) and supernatant of the solubilized suspension after centrifugation was loaded onto Ni–NTA agarose resin. After washing steps, the fusion proteins were eluted with the elution buffer (8 M Urea, 100 mM NaH$_2$PO$_4$, 100 mM Tris-Cl pH 4.5). The eluted proteins were refolded via dialysis in refolding buffer (20mM Tris-HCl pH7.4 and 10% Glycerol). Polyethylene Glycol (PEG) 20000 was used for protein concentration according to standard procedures (22). The protein concentration was determined by standard Bradford assay and optical density at 280nm (OD$_{280}$).

**SDS-PAGE and Western blot analyses of rSK**

SDS-polyacrylamide gel electrophoresis was performed for protein expression assay. For western blotting, proteins were transferred to nitrocellulose membrane and the membrane was blocked by 5% BSA. Mouse anti-penta His monoclonal antibody (Qiagen, USA) was used as the primary antibody and goat anti-mouse IgG conjugated to HRP (Horse Radish peroxidase) (Qiagen, USA) as the secondary (tracking) antibody. The bound antibodies were detected using 3, 3-Diaminobenzidine (DAB) (Qiagen, USA).

**Chromogenic assay of purified rSK activity**

SK activity was determined by chromogenic substrate as previously described (23). Purified rSK (5nM) was added to a microtiter plate containing 0.2 mM of chromogenic substrate S-2251 (H-D-valyl-L-leucyl-L-lysine-p-nitroanilide dihydrochloride; Sigma, USA) and 200 mM of human plasminogen (Sigma, USA) at 37°C in a total volume of 100 µl of assay buffer (50 mM Tris-HCl, 150 mM NaCl, pH 7.4). Hydrolysis of S-2251 was measured at 405 nm every 5 min for 60 min in a microplate reader (BioHIT, UK). The protein activity was calculated using standard activity curve of streptase® (CSL, Behring, Germany).
Results

Screening of β-haemolytic streptococci (βHS) and isolation of SK producing GCS

A total of 252 samples (Table 1) were examined by microbiological and biochemical assays (Table 2). βHS including group A, C and G were found in 75 out of 250 streptococci specimens (29.4%). Group A was the dominant Lancefield serogroup found in 65 out of 250 streptococci specimens (25.8%) followed by GCS (9 out of 250; 3.6%) and GGS (1 out of 250; 0.4%), respectively. Throat culture was the common source of GAS (60 out of 250 isolates), followed by urine culture (4 out of 250 isolates) and soft tissue (1 out of 250 isolates). GCS were less common and were totally isolated from respiratory tract (7 out of 250 isolates) and genital tract (2 out of 250 isolates). More detailed fermentation analyses on isolated GCS (Table 2) could determine eight *S. dysgalactiae* subsp. *equisimilis* strains and one strain of *S. dysgalactiae* subsp. *dysgalactiae* (Table 3; sample codes are based on in house coding). Most of GCS isolates showed low to moderate SK activity in caseinolysis assay, except for GCS-S87 that showed predominant SK activity compared to the reference strain (GCS-9542) and thus was selected for further cloning studies (Figure 1).

Cloning, expression, purification and refolding of the rSK

Using the SKf and SKr primers and genomic DNA of *S. equisimilis* GCS-9542 and *S. equisimilis* GCS-S87 as template, PCR reactions resulted in a single band of the expected length (1242bp) of SK gene for both strains (Figure 3). Cloning steps for insertion of SK gene in pQE30 vector is illustrated (Figure 2). Restriction enzyme analyses of the recombinant vector harboring SK gene (pSK87) (Figure 3) and nucleotide sequence analyses (not shown) confirmed the accuracy of cloning procedures.

Expression of pSKC plasmids produced rSK with a predicted molecular mass of about 47 kDa, harboring a 6XHis domain which was appended to the N-terminus of the native molecule. Crystal structure of streptokinase shows that the N-terminal of the enzyme is unfolded (24), thus, addition of polyhistidine at the NH₂-terminus of the enzyme was expected to have little or no effect on the catalytic activity. Analysis of protein profile of un-induced and induced cell lysates on SDS-PAGE proved expression of SK by both strains (Figure 4A). Analysis of the purified rSK further indicated two major bands on the gel (Figure 4A). The upper band which was the most prominent protein corresponded to the full length SK (47 kDa) and the lower band (around 44 kDa) might be related to the digested form of SK as previously suggested (25). Accordingly, the eluted (purified) recombinant proteins were identified by the presence of the same two bands in western blot analysis (Figure 4B). Evaluation of the expression efficiencies for the rSK proteins by concentration measurements at OD₂₈₀ and Bradford assay indicated yields of 0.53 mg/ml (rSK9542) and 0.59 mg/ml (rSKS87) for the purified proteins.

Table 1. Distribution of identified GCS subspecies in different samples

| No. | Sample No. | culture Source | Disease                  | Sub species of GCS |
|-----|------------|----------------|-------------------------|-------------------|
| 1   | S-04       | Human throat    | Streptococcal pharyngitis | *S. equisimilis*   |
| 2   | S-05       | Human throat    | Streptococcal pharyngitis | *S. equisimilis*   |
| 3   | S-08       | Human vagina    | Puerperal fever          | *S. equisimilis*   |
| 4   | S-87       | Human throat    | Streptococcal pharyngitis | *S. equisimilis*   |
| 5   | S-91       | Human throat    | Acute tonsillitis        | *S. dysgalactiae*  |
| 6   | S-31       | Human vagina    | Puerperal fever          | *S. equisimilis*   |
| 7   | K-17       | Human throat    | Acute tonsillitis        | *S. equisimilis*   |
| 8   | K-19       | Human throat    | Acute tonsillitis        | *S. equisimilis*   |
| 9   | K-34       | Human throat    | Acute tonsillitis        | *S. equisimilis*   |

Figure 3. Restriction Enzyme Analysis of Recombinant pQE-S87
Lane 1: DNA Marker 1kb, lane 2: pQE87 digested by *BamHI*-PstI (3461 and 242bp fragments corresponding to vector and PCR fragments respectively, Lane 3: digestion of pQE30 by *BamHI* (346bp); Lane 4: PCR product of SKC-S87 (242bp). The corresponding bands were indicated by arrows.
Biological activity assay of streptokinase by chromogenic method

The chromogenic assay in the absence of fibrin is known as an approved an internationally standard assay for streptokinase activity (Third International Standard for streptokinase; National Institute of Biological Standard and Controls, NIBSC, 2004UK) (26). Employing this method and chromogenic substrate S-2251, a standard curve based on definite activity of Streptase® was plotted (Data not shown). Subsequently, the biological activity (which represents the activity of SK in International Units per ml of total volume; IU/ml) and specific activity (which refers to the activity of rSK per mg of total protein; IU/mg) were calculated based on the plotted standard curve. The calculated values for the biological activities were 11200 IU/ml (rSK9542) and 10720 IU/ml (rSKS87) and for specific activities were $0.21 \times 10^5$ IU/mg (rSK9542) and $0.18 \times 10^5$ IU/mg (rSKS87).

Discussion

In the present study, most of the clinical samples were isolated from the throat of human beings (Table 1). The reason behind this strategy was the fact that the respiratory tract is being used by most βHS as a port of entry into the body of their hosts (27). Prior epidemiological studies indicated that GCS is not the dominant group in human streptococcal diseases (28). Accordingly, among all streptococci isolates in this study, only 9 strains of GCS were identified (Tables 1 and 3). Among isolated GCS, S.equisimilis was the predominant subspecies identified (8 out of 9) and only one S. dysgalactiae strain could be detected in our samples (Table 3). These data are in accordance with previous studies reporting S.equisimilis as the major subspecies to be screened among GCS (29). Finally, GCS group had the lowest frequency among screened βHS in our study (only 1 out of 250). This finding is in accordance with previous reports which designated GCS as a minor human pathogen (30). Among GCS isolated S.equisimilis strains, only GCS-9542 showed similar activities to reference strain (GCS-9542) in radial caseinolytic assay (Figure 1) and all the other 8 isolated GCS strains indicated less activity. Due to the presence of the same genetic regulating factor for both hemolytic and SK activity (31), the quantity of β-hemolytic activity on blood agar has been considered as an important characteristic for screening of SK producing streptococci in most of the previous studies (8, 15, 17). However, in the present study both hemolytic and caseinolytic activities were the criteria considered for selection of the SK producing GCS. Accordingly, the GCS-S87 (Figure 1 and Table 3) was selected for isolation of SK gene and production of rSK. For both expressed proteins (SK of GCS-9542 and GCS-S87 strains) SDS-PAGE and western blotting results (Figure 4B and C) indicated the appearance of both a 44 kDa and full length protein (47kDa). These results are in accordance with prior observations which had suggested that C-terminal region of SK is vulnerable to proteolytic attack in all examined expression systems (32). This phenomenon however does not affect plasminogen activation potential of rSK as previously demonstrated (25). Measurement of pure and refolded rSK9542 and rSKS87 proteins indicated yields of 0.53 mg/ml and 0.59 mg/ml, respectively. The design of expression strategy depends on process economics and end-use of the purified protein. In the case of SK, expression of inclusion bodies is shown to be useful for obtaining large amounts of protein, provided that refolding is sufficient and recovery of protein is high (35). However,
by employing a different expression system and optimization of medium, purification and refolding process, literature has reported a wide range of SK expression (15-65% of total protein) and recovery (purity 100% and recovery 68%) (33).

SK specific activity assay of rSKS87 presented about 88.1% of the activity of SK9542 which is a recognized activity for a new strain. In a very recent study in Egypt, it was reported that the clot lytic activities of both recombinant form of SK expressed from a locally isolated strain and commercial SK were almost similar (15). In the present study, however, analysis of rSK activities was performed by chromogenic method in the absence of fibrin according to the European Pharmacopeia (EP) (26). National Institute for Biological Standard and Control (NIBSC) has recommended that the analysis for SK activity is to be carried out under two different conditions (i.e. both in the presence and absence of fibrin). It has been noted that rSK from different sources are sensitive to the assay format (34). For instance, the presence of fibrin may change the result of the analysis of SK activity compared with the data obtained from chromogenic assay based on the EP recommendations for analysis of SK activity (34). Therefore determination of activity of SKS87 in the presence of fibrin and comparison with SK9542 and a commercial SK might be considered in future studies. In addition the GCS-S87 strain in our study might be subjected to different mutation methods for the aim of strain improvement as previously suggested for improving SK production (14, 22).

In summary, to our best of knowledge, we reported the results of the first national screening of βHS isolated from local and regional samples of Iran for the first aim of isolating a native SK producing GCS. βHS were found in 75 out of 252 specimens (29.4%) including groups A (25.8%), C (3.6%) and G (0.4%). Cloning of SK genes from the selected GSC strain (GCS87; which showed the highest SK activity in caseinolytic assay) and the standard GCS9542 strain in E.coli M15 resulted in the proper expression of 47 kDa SK proteins after IPTG induction. One-step purification by Ni-NTA affinity chromatography followed by refolding steps provided yields of 0.53 mg/ml and 0.59 mg/ml for rSK 9542 and rSKS87, respectively. Using the chromogenic assay via application of S-2251 substrate, comparable biological activities of: 11200 IU/ml and 10720 IU/ml and specific activities of: 0.21x10^5 IU/mg and 0.18x10^5 IU/mg were obtained for SK9542 and SKS87, respectively. rSK from a local source with comparable yields and activities to the standard strain was expressed. Optimization of expression condition, purification and refolding process will aid to improve the yields of product for future production plans.

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References

1. Annmol Kumar AK, KK Palicharya KKP, K. Serha Ram KSR, KBS Sambhasiva Rao KRSSR. Evolutionary trend of thrombolytics. Int J Bio Sci Bio Technol 2010; 2:53-68.
2. Banerjee A, Chisti Y, Banerjee UC. Streptokinase—a clinically useful thrombolytic agent. Biotechnol Adv 2004; 22:287-307.
3. Kunamneni A, Abdelghani TTA, Ellaiya P. Streptokinase—the drug of choice for thrombolytic therapy, J Thromb Thrombolysis, 2007; 23:9-23.
4. Baruah DB, Dash RN, Chaudhari M, Kadam S. Plasminogen activators: A comparison. Vasc Pharmacol 2006; 44:9-9.
5. Della G, Miocardico SNL. GISSI-2: a factorial randomised trial of alteplase versus streptokinase and heparin versus no heparin among 22,490 patients with acute myocardial infarction. Lancet 1990; 336:65-71.
6. Ridker PM, O’Donnell C, Marder VJ, Hennekens CH. Large-scale trials of thrombolytic therapy for acute myocardial infarction: GISSI-2, ISIS-3, and GUSTO-I. Ann Intern Med 1993; 119:530.
7. Siki N, Baruda A. A history of streptokinase use in acute myocardial infarction. Texas Heart Institute J 2007; 34:338-337.
8. Abdelghani TTA, Kunamneni A, Ellaiya P. Isolation and mutagenesis of streptokinase producing bacteria, Am J Immunol 2005; 3:129-249.
9. Araki R, Roohvand F, Noruzian D, sardari S, Aghasadeghi MR, Khanamad H, et al. A comparative study on the activity and antigenicity of truncated and full-length forms of streptokinase. Polish J Microbiol 2011; 60:243-252.
10. Malie H. Polymorphism of the streptokinase gene: implications for the pathogenesis of post-streptococcal glomerulonephritis. Zentralblatt fur Bakteriologie. Int J Med Microbiol 1993; 278:246-253.
11. Malie H, Steiner K, Gase K, Mechold U, Ellinger T. The streptokinase gene: allelic variation, genomic environment and expression control. Dev Biol Stand 1995; 85:183.
12. Pupo E, Baghbaderani BA, Lugo V, Fernandez J, Paez R, Torrens I. Two streptokinase genes are expressed with different solubility in Escherichia coli W3110. Biotechnol lett 1999; 21:1119-1123.
13. Keramati M, Roohvand F, Esfamejad Z, Nikbin VS, Aslani MM. PCR/RFLP-based allelic variants of streptokinase and their plasminogen activation potencies. FEMS Microbiol Lett 2012; 335:79-85.
14. Towdros W, Norgren M, Kronwall G. Streptokinase activity among group A streptococci in relation to streptokinase genotype, plasminogen binding, and disease manifestations. Microbial Pathogenesis 1995; 18:53-65.
15. Al Sohaimy S, Aleem E, Hafez EE, Esmaiil SS, El-Saadani M, Moneim NA. Expression of recombinant Streptokinase from local Egyptian streptococcus sp. SalMarEg. Afr J Biotechnol 2011; 10:5900-5911.
16. Freed E. Production and partial purification of streptokinase by streptococcus sp. Int J Students Project (IJSPP) Biotechnol 2010; 11:39.
17. Doss HM, Manohar M, Singh NA, Mohanasrinivasan V, Devi CS. Studies on isolation, screening and strain improvement of streptokinase producing hemolytic streptococci. World J Sci Technol 2011; 1:74-77.
18. Garrity GM, Brenner DJ, Krieg NR, Bergey’s manual of systematic bacteriology. Springer: 2005.
19. Kim DM, Lee SJ, Kim IC, Kim ST, Byun SM. Asp41-His48 region of streptokinase is important in binding to a substrate plasminogen. Thrombosis Res 2000; 99:93-98.
20. Sambrook J, Russell DW. The condensed protocols from molecular cloning: a laboratory manual: Cold Spring Harbor Laboratory Press: 1989.

Keramati M et al
Recombinant Streptokinase from Iranian Streptococci
Iran J Basic Med Sci; Vol. 16, No. 4, Apr 2013

626
21. Crowe J, Henco K. The QIAexpressionist. DIAGEN GmbH, Hilden, Germany: 1992; p. 2.
22. Ingham KC. Precipitation of proteins with polyethylene glycol. Methods Enzymol 1990; 182:301-306.
23. Wohl RC, Summaria I, Robbins K. Kinetics of activation of human plasminogen by different activator species at pH 7.4 and 37 degrees C. J Biologic Chem 1980; 255:2005-2011.
24. Wu XC, Ye R, Duan Y, Wong SL. Engineering of plasmin-resistant forms of streptokinase and their production in Bacillus subtilis: streptokinase with longer functional half-life. App Environ Microbiol 1998; 64:824-830.
25. Pimienta E, Ayala JC, Rodríguez C, Ramos A, Van Mellaert I, Vällin C, et al. Recombinant production of Streptococcus equisimilis streptokinase by Streptomyces lividans. Microb Cell Fact 2007; 6:20-31.
26. Sands D, Whitton C, Longstaff C. International collaborative study to establish the 3rd International Standard for Streptokinase. J Thromb Haemost 2004; 2:340-345.
27. Winn WC, Koneman EW. Koneman’s color atlas and textbook of diagnostic microbiology. Lippincott Williams & Wilkins; 2006.
28. Hughes JM, Wilson ME, Brandt CM, Spellerberg B. Human infections due to Streptococcus dysgalactiae subspecies equisimilis. Clin Infect Dis 2009; 49:766-772.
29. Barnham M, Kerby J, Chandler R, Millar M. Group C streptococci in human infection: a study of 308 isolates with clinical correlations. Epidemiol Infect 1989; 102:379-390.
30. Davies MR, McMillan DJ, Sriprakash KS, Chhatwal GS. Distribution of group A streptococcal virulence genes in group C and G streptococci. Int Cong Series 2006; 1289:184-187.
31. Steiner K, Malke H. Dual control of streptokinase and streptolysin S production by the covRS and fasCAX two-component regulators in Streptococcus dysgalactiae subsp. equisimilis. Infect Immun. 2002; 70:3627-3636.
32. Kim MR, Choeng YH, Chi WJ, Kang DK, Hong SK. Heterologous production of streptokinase in secretory form in streptomyces lividans and in nonsecretory form in escherichia coli. Microbiol Biotechnol 2010; 20:112-117.
33. Balagurunathan B, Ramchandra NS, Jayaraman G. Enhancement of stability of recombinant streptokinase by intracellular expression and single step purification by hydrophobic interaction chromatography. Biochem Engineering Journal. 2008;39(1):84-90.
34. Longstaff C, Thelwell C, Whitton C. The poor quality of streptokinase products in use in developing countries. J Thromb Haemost 2005; 3:1092-1094.