

Original Research Article

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## Isolation and Screening of Fibrinolytic Enzymes producing Bacterium Strain from Soil Waste

S.M. Gopinath\* and K. Lingappa

Department of Microbiology, Gulbarga University, Gulbarga, Karnataka, India

\*Corresponding author

### ABSTRACT

#### Keywords

Fibrinolytic enzymes, isolates, fibrinolytic activity, caesinolytic and gelatinolytic activities.

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Fibrinolytic enzymes help in dissolving the fibrin clots and they have the potential use in the treatment of harmful cardiovascular diseases. High efficient fibrinolytic enzymes are produced from the fibrinolytic microbes under extreme environmental conditions. In the present work, microorganism strain that helps in the production of fibrinolytic enzymes were isolated and screened from the soil waste. Five isolates i.e., GSM-15, GSM -25, GSM -35, GSM -45 and GSM -55 were subjected to macroscopic characteristics. The isolated organism was cultured in the sterile media at 37°C for 24 hrs. The selected isolates were subjected to caseinolytic, gelatinolytic and fibrinolytic activities was done. The fibrinolytic activity of isolates GSM 25 and GSM 55 was higher than that of caesinolytic and gelatinolytic activities. The selected isolates were tested with modified media composition and at different pH values. Isolate GSM-55 showed good enzyme productivity ( $13.1 \pm 2.6$ ) U/ml at pH 8.0. Also the biochemical study indicated the isolate GSM 55 is similar to that of *Bacillus cereus* with fibrinolytic function.

### Introduction

Fibrinolytic protease is well known as a sub class of protease, it has an ability to degrade fibrin (Fujita *et al.*, 1993, 1995; Jeong *et al.*, 2001, 2004; Leonardi *et al.*, 2002; Sumi *et al.*, 1995; Wong *et al.*, 2004). Fibrin is normally formed from Fibrinogen by the action of Thrombin (EC 3.4.21.5). It is sliced by plasmin (EC 3.4.21.7) which is activated from plasminogen by tissue plasminogen activator (Voet *et al.*, 1990). In balanced condition, fibrin clots are hydrolysed by plasmin to avoid thrombosis in blood vessels. However, in unbalanced condition, as a result of pathophysiological

disorders, the clots are not hydrolysed. Deposition of fibrin in blood vessels normally increases thrombosis and other cardiovascular diseases such as high blood pressure, acute myocardial infarction, ischemic heart disease, valvular heart disease, peripheral vascular disease, arrhythmias, stroke etc. (Bode *et al.*, 1996; Yoshinori *et al.*, 2005).

Fibrinolytic agents are used to treat a wide variety of venous and arterial thromboembolic disorders; especially against acute myocardial infarction (Torrens

*et al.*, 1999). Based on working mechanism, thrombolytic agents are of two types, one is plasminogen activator which activates plasminogen into active plasmin to degrade fibrin and the other is plasmin like proteins which directly degrade fibrin (Koide *et al.*, 1982). Despite widespread uses, thrombolytic agents are expensive. They exhibit low fibrin specificity and have undesired side effects such as gastrointestinal bleeding, resistance to reperfusion and allergic reactions (Pautov *et al.*, 1990). Therefore, continuous efforts have been focused in the search of safer and less expensive thrombolytic agents from diverse sources but one must respect biodiversity. Investigation of extracellular fibrinolytic enzymes is an essential due to their wide applications in clinical, pharmaceutical, food, and bioremediation process.

## **Materials and Methods**

### **Chemicals**

All chemicals and reagents used were of analytical grade obtained from Sigma USA.

### **Determination of fibrinolytic protease activity**

The fibrinolytic protease activity was based on the hydrolytic power of the enzymes was determined by using the fibrin as substrate. Fibrinolytic protease activity was evaluated by the modified procedure of Greenberg (1957) using 2% bovine fibrin in 0.05 mM phosphate buffer (pH 8). 0.5ml fibrin solution with an equal volume of diluted enzyme solution was incubated for 10min at 37°C. The reaction was stopped by the addition of 10% cold Tri Chloroacetic Acid (TCA). The mixture was centrifuged at 3,000rpm for 10 min and to the supernatant 5ml of 0.44M Na<sub>2</sub>CO<sub>3</sub> was added, followed by the addition of 1ml of diluted Folin -

ciocalteau reagent. The mixture was incubated for 30min at 37°C, the sample develops color change, and the intensity of colour change will be measured at 660nm using Shimadzu UV-1700 (Japan) spectrophotometer against a blank reagent prepared without adding enzyme. Tyrosine is used as the reference standard.

### **Estimation of extracellular protein**

Extracellular protein was estimated by Lowry method (Lowry *et al.*, 1951). To the 1ml of dilute protein sample (10-60 µg/ml), 5ml alkaline working solution was added and mixed well which was incubated at 37°C for 10min. To the above mixture, 0.5ml Folin Ciocalteau reagent was added, mixed well and incubated at 37°C for 30min. The absorbance was measured at 680nm in a spectrophotometer. The amount of protein present in the sample was calculated from the standard curve.

### **Screening and isolation of fibrinolytic protease producing bacteria**

To isolate the potent fibrinolytic protease producing bacteria, soil samples were collected from various places in and around Bangalore, Karnataka, India. The soil samples were selected based on the varying characteristics such as organic matter, particle size, colour of soil and geographical distribution. The samples were stored in sterile condition.

1g of the below samples was separately taken in a conical flask containing 100ml sterile water and the suspension was subjected to rotary shaker for 30 minutes. 1ml of the supernatant was serially diluted with sterile water and one ml of each of these dilutions was added to 20ml of sterile molten starch casein-agar medium maintained at 40°C.

### Samples

Sample	Source	Physical Characteristics
GUG –I	chicken centre, Shivaji Nagar, Bangalore.	brown colour sticky semi solid soil sample
GUG –II	chicken centre, Tannery Road, Bangalore	black colour sticky semi solid soil sample
GUG –III	slaughterhouse, nearby Tannery Road, Bangalore	brown colour sandy semi solid soil sample
GUG –IV	dumping yard of chicken centre near Tannery Road, Bangalore	brown colour hard soil sample containing the mixture of sand and clay
GUG –V	slaughterhouse, nearby Soldevanahalli, Bangalore	brown colour semi solid sample
GUG –VI	chicken centre, Chikkabanavara, Bangalore	brick red sticky soil sample
GUG –VII	dumping yard of chicken centre near Devasandra lake, Bangalore	red colour semi solid consisting of sand and clay sample
GUG -VIII	slaughterhouse, nearby Tin Factory, Bangalore	red colour semi solid clay sample
GUG –IX	dumping yard of chicken centre near Tin Factory, Bangalore	black colour solid consisting of sand and clay sample
GUG –X	dumping yard of fish market near Yashwantpura Railway Station, Bangalore	black colour sticky sample consisting of sand and clay

It was mixed thoroughly and plated in 10cm diameter sterile petridishes and incubated at 37°C. 75µg/ml of Flucanazole was incorporated to control the fungal contamination.

After 24h of incubation, the bacteria colonies with clear hydrolyzed zones were transferred onto starch casein agar slants.

#### Screening of isolates for proteolytic activity

##### Primary screening

The selected isolates were initially screened for their proteolytic activities i.e. caseinolytic, gelatinolytic and fibrinolytic activities.

#### Caseinolytic activity

Caseinolytic activity of the isolates was evaluated using casein-agar plate technique

(Zerdani *et al.*, 2004; Li *et al.*, 2009; Gholamhossein *et al.*, 2013). To the sterilized agar, 10% of pasteurized skimmed milk was added aseptically and the media was transferred into sterile petridish and kept aside for solidification. Then a loopful of each culture was streaked onto the medium, incubated at 37°C for 24h. The diameters of hydrolyzed zones around the colonies and the growth zones were measured. The ratio of hydrolysis zone/growth zone was calculated which gives a measure of the caseinolytic activities of the isolates.

### **Gelatinolytic activity**

20ml of sterile nutrient gelatin agar medium (McLaughlin *et al.*, 1996; Deshmukh *et al.*, 1997; Tran *et al.*, 2002; Shanmugasundaram *et al.*, 2012) was poured in sterile petridishes and inoculated with a loop full of spores from 24h old cultures and incubated at 37°C for 24h. The plates were flooded with mercuric chloride reagent (Williams *et al.*, 1971). After treating with mercuric chloride-HCl solution, the hydrolysis zone and growth zones were noted.

### **Fibrinolytic protease activity**

To check the fibrinolytic protease activity, modified fibrin plate agar (Astrup *et al.*, 1952; Kumaran *et al.*, 2011) was poured in sterile petridishes and inoculated with a loop full of spores from 24h old cultures and incubated at 37°C for 24h. The diameters of hydrolyzed zones around the colonies and the growth zones were measured. The ratio of hydrolysis zone/growth zone was calculated which gives a measure of the fibrinolytic protease activities of the isolates.

### **Design of suitable basal medium**

Fibrinolytic protease production in shake flasks was checked using different media composition (table.2). 1ml of the cell suspension from the isolates was aseptically transferred into 250ml Erlenmeyer flasks containing 50ml each of sterile medium. The contents were incubated at 37°C for 24h using rotary shaker at 70rpm. It is followed by centrifugation at 3000rpm for 10 min and the supernatant solution was tested for fibrinolytic protease activity by modified method of Greenberg 1957.

### **Determination of type of fibrinolytic protease produced by the isolates**

To determine the whether the enzyme

secreted by the isolate is of alkaline or acidic or neutral protease, fibrinolytic protease activity in the harvested broth was assayed by adding 0.5ml culture broth to 0.5ml of 2% fibrin solution. Further the activity was checked at different pH conditions using Citrate buffer (pH 4.0), Phosphate buffer (pH 7.0) and Carbonate buffer (pH 8.0, 10.0) by dissolving the fibrin in respective buffers (Greenberg 1957).

### **Results and Discussion**

From the ten soil samples collected 116 isolates were formed. Of the colonies formed, five isolates (Nos. 15, 25, 35, 45 and 55) were selected based on their macroscopic characteristics. They were sub cultured on starch casein agar media and subjected for secondary screening. They were designated as KLG-15, KLG -25, KLG -35, KLG -45 and KLG -55. Extent of growth and the proteolytic activities of the selected isolates were checked and the results were presented in Table.1.

Study clearly indicated that isolates KLG 25 and KLG 55 are high fibrinolytic protease producers and they were further tested with medium IV and V for better fibrinolytic protease production.

### **Design of suitable basal medium**

The composition of the medium IV and V were slightly changed in their fructose concentration and used for the production of fibrinolytic protease by isolates KLG 25 and KLG 55 to compare and design a suitable basal medium for efficient production. The results are presented in Table.4.

### **Determination of type of fibrinolytic protease produced by the isolates**

Isolates KLG -25 and KLG -55 indicated

high enzyme activity at pH 8, indicating the enzyme is an alkaline protease. Further the isolate KLG-55 showed comparatively good enzyme productivity ( $13.1 \pm 2.6$ ) U/ml.

Also, results of classification and identification clearly showed that the isolate KLG -55 is having similar characteristics as of *Bacillus cereus*.

**Table.1** Growth pattern and proteolytic activities of selected isolates Secondary Screening

Sample No.	Isolate No.	Extent of Growth (24h)			Proteolytic activity		
		CA	GA	FA	Caseinolytic activity	Gelatinolytic activity	Fibrinolytic protease activity
GUG -II	KLG 15	++	+	++	8.4	2.6	3.9
GUG -IV	KLG 25	++	++	++	7.8	5.6	6.9
GUG -V	KLG 35	++	+	++	5.3	1.9	3.1
	KLG 45	++	+	+	5.8	2.6	2.3
GUG -VI	KLG 55	++	-	++	6.8	0	8.4
CA = Casein Agar Medium GA = Gelatine Agar Medium FA = Fibrin Agar Medium Hydrolysed zone (mm) / growth zone (mm) = Ratio					++ = Good growth + = Moderate Growth - = No Growth		

Fibrinolytic protease production was checked for the five isolates in shake flasks using different media composition (table.2). The results are presented in Table.3.

**Table.2** Composition of different media for fibrinolytic protease production

Medium No.	Composition (g/100ml)
I	Glucose, 3.0; soyabean meal, 2.0; CaCl <sub>2</sub> , 0.04; MgCl <sub>2</sub> , 0.2.
II	Glucose 1.5; yeast extract, 0.5; CaCl <sub>2</sub> , 0.2.
III	Glucose, 0.1; yeast extract, 0.5; tryptone, 0.5.
IV	Fructose, 1.0; fibrin; 0.2; peptone 1.0; salt solution 5ml*.
V	Fructose, 0.5; fibrin; 0.2; peptone 1.0; salt solution 5ml*.
VI	Fructose, 1.0; fibrin, 0.2; salt solution, 5ml*.
VII	Soluble starch, 1; Casein, 0.3; KNO <sub>3</sub> 0.2; K <sub>2</sub> HPO <sub>4</sub> , 0.2; MgSO <sub>4</sub> , 7H <sub>2</sub> O; CaCO <sub>3</sub> , 0.002; FeSO <sub>4</sub> , 7H <sub>2</sub> O, 0.001.

\* Salt Solution Composition (%): KNO<sub>3</sub>, 0.2; K<sub>2</sub>HPO<sub>4</sub>, 0.2; MgSO<sub>4</sub>, 7H<sub>2</sub>O, 0.005; CaCO<sub>3</sub>, 0.002; FeSO<sub>4</sub>, 7H<sub>2</sub>O, 0.001.

**Table.3** Production of fibrinolytic protease (U/ml) by selected isolates in shake flask\*

Medium No.	Isolate KLG15	Isolate KLG 25	Isolate KLG 35	Isolate KLG 45	Isolate KLG 55
I	3.7	2.5	4.2	2.6	4.0
II	4.2	4.6	2.0	4.1	2.6
III	5.1	4.8	2.8	1.2	4.2
IV	1.2	7.6	1.8	1.3	8.6
V	3.8	5.7	5.0	3.5	7.0
VI	4.7	3.6	3.2	2.0	3.5
vII	1.7	2.0	2.4	1.9	2.0

\* Fibrinolytic Protease activity expressed in U/ml.

**Table.4** Production of fibrinolytic protease in modified media

Medium No.	Composition of media	Enzyme yield (U/ml)	
		KLG-25	KLG -55
I	Fructose, 1%, fibrin, 0.2%; peptone 1%; MgSO <sub>4</sub> , 0.02%, KH <sub>2</sub> PO <sub>4</sub> , 0.2%; NH <sub>4</sub> NO <sub>3</sub> , 0.5% and CaCl <sub>2</sub> 0.04%	9.4 ± 2.5	13.1 ± 3.6
II	Fructose, 0.5%; fibrin, 0.2%, peptone 1%; MgSO <sub>4</sub> , 0.02%; KH <sub>2</sub> PO <sub>4</sub> , 0.2%, NH <sub>4</sub> NO <sub>3</sub> , 0.5% and CaCl <sub>2</sub> 0.04%	8.4 ± 2.8	9.8 ± 3.2
III	Fructose 0.1%, Yeast extract 0.5%, Tryptone 0.5%	7.9 ± 2.5	8.9 ± 3.2
IV	Fructose 1%, Yeast extract 0.5%, Tryptone 0.5%	8.0 ± 2.0	6.0 ± 2.0

Maximum yield was obtained in medium IV. The composition of medium IV is Fructose, 1.0; fibrin, 0.2; peptone 1; MgSO<sub>4</sub>, 0.02%; KH<sub>2</sub>PO<sub>4</sub>, 0.2%; NH<sub>4</sub>NO<sub>3</sub>, 0.5% and CaCl<sub>2</sub> 0.04%. This was designated as the basal medium and used for further studies.

**Table.5** Fibrinolytic protease activity at different pH values

Isolate	Fibrinolytic protease activity (U/ml)			
	pH 4.0	pH 7.0	pH 8.0	pH 10
KLG - 25	1.2 ± 2.2	8.7 ± 1.4	9.2 ± 2.2	1.3 ± 2.4
KLG -55	4.8 ± 2.5	10.6 ± 1.8	13.1 ± 2.6	3.5 ± 2.0

Several effective thrombolytic agents have been identified and characterized from microorganisms, earthworms, plants, snake venoms, insects and leeches. Fibrinolytic proteases have wide applications in clinical, pharmaceutical, food, and bioremediation process. Among the various fibrinolytic proteases, bacterial extracellular fibrinolytic proteases are more attractive than those isolated from earthworms, plants, snake venoms, insects and leeches extracellular fibrinolytic proteases because of their high stability and unusual substrate specificity.

Different natural substrates collected from various places of Bangalore were screened for the isolation of potent fibrinolytic protease producing *Bacillus cereus*. A total of 116 cultures were isolated from 10 samples. All the isolates were tested for their proteolytic activity on milk casein agar medium. After preliminary studies, isolates were further screened in two different media

for their caseinolytic, gelatinolytic and fibrinolytic activities. Five promising isolates (KLG 15, KLG 25, KLG 35, KLG 45 and KLG 55), which showed good fibrinolytic activity were further screened for their enzymatic activities by shake flask method. Among these 5 isolates, isolate KLG 55 exhibited good fibrinolytic activity, while other showed poor to moderate activity. This was subjected to biochemical studies. A close study indicated that the isolate (KLG 55) was related to *Bacillus cereus* but different in some biochemical characteristics. In view of some significant differences from the reference culture, the isolate was considered to be the novel strains of *Bacillus cereus*.

Strain development for enzyme production has been an essential prerequisite for efficient production process. The main aim of strain improvement programme is to increase the product litres. The potential

productivity of the organism is controlled by its genome and therefore the genome must be modified to increase the potential yield. The process of strain improvement involves genetic modification of the culture, followed by process optimization. Therefore, the strain of *Bacillus cereus* was subjected to strain improvement program with a view to obtain increased fibrinolytic protease production.

In conclusion, the isolate KLG 55 from the soil waste indicated the effective fibrinolytic activity at pH 8, indicating the enzyme is an alkaline protease with comparatively good enzyme productivity ( $13.1 \pm 2.6$ ) U/ml. Biochemical studies clearly showed that the isolate KLG -55 is having similar characteristics as of *Bacillus cereus*. In view of some significant differences from the reference, the isolate was considered to be the novel strains of *Bacillus cereus*.

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