Optimization of Phospholipase A\textsubscript{1} (PLA\textsubscript{1}) Production from a Soil Isolate \textit{Bacillus subtilis} subsp. \textit{inaquosorum} RG1 via Solid State Fermentation

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Microbial sources for phospholipase A\textsubscript{1} (PLA\textsubscript{1}) are economic and industry relevant for degumming of oils and also their product lyso-phospholipids have been widely used as emulsifying agent. Numerous PLAs have been reported, but still the few enzymes have got position in the commercial sector. Due to enormous demand of PLA\textsubscript{1} in the industrial sector, the present study was carried out to optimize PLA\textsubscript{1} production using cheaper agro-industrial waste like defatted rice bran. For this, defatted rice bran was used in solid state fermentation for the production of PLA\textsubscript{1}. One-factor at a time approach was used to obtain maximum production of 51.5 U/gm which is 2.15 folds more than un-optimized medium. The optimized medium components were (pH 7): defatted rice bran (5gm), glucose (1\% w/v), peptone (1\% w/v) and olive oil (0.5 \% v/v) with moisture content of 1:1.5 and after 48h of incubation at 37°C. This approach will provide the cleaner solution for degumming of oil as compare to the acid degumming and also help to reduce the stress of the environment by utilizing waste. This is the first report where in \textit{Bacillus subtilis} subsp. \textit{inaquosorum} was employed for PLA\textsubscript{1} production via solid state fermentation.

**Keywords**: Submerged fermentation, Solid-State Fermentation, Optimization, Classical approach, PLA\textsubscript{1}, \textit{Bacillus subtilis} subsp. \textit{inaquosorum}.

Phospholipases are one of key enzymes in oil refinery industry and play a crucial role in degumming of edible oils. Physical refining using water removes only hydratable phospholipids so there is great demand for the environment friendly technique that can also remove the non-hydratable phospholipids\textsuperscript{1} from edible oils. Various chemical processes are also used by the industries to remove phospholipids that includes acid degumming but enzymatic degumming using phospholipases is more advantageous and effective\textsuperscript{2} because these enzymes can easily convert non-hydratable phospholipids to the hydratable phospholipids and can be further eliminated by centrifugation\textsuperscript{3}.

Soil is a major reservoir for a variety of microorganisms\textsuperscript{4-6} and can be used for exploiting isolates possessing commercially important enzymes like PLA\textsubscript{1}, Lyso-phospholipids, the product formed by action of PLA\textsubscript{1} on phospholipids are also in great demand as emulsifying agent, cosmetic agent as well as drug delivery agent. Various PLA\textsubscript{1} has been purified from various mammalian systems but to obtain high titers of PLA\textsubscript{1} microbial sources are still in demand\textsuperscript{7}.
Phospholipases are basically classified into four groups depending on the site of ester bond attacked i.e. phospholipase A (PLA), phospholipase B (PLB), phospholipase C (PLC) and phospholipase D (PLD). Phospholipases A are further categorized into Phospholipases A1 (PLA1) and Phospholipases A2 (PLA2). Numerous PLA1 have been reported from a variety of microorganisms that includes PLA from Serritia liquefaciens, Fusarium sp., Aspergillus oryzae, Streptomyces. Out of which only a few enzymes are available commercially viz. PLA1 from Novozenzymes (Fusarium PLA1) and Genecor (Streptomyces PLA1).

Fermentation process is the way out for producing high volume of enzymes. Solid state fermentation is an efficient method for enzyme production with less cost and reduced risk of contamination. For solid state fermentation, agro-industrial waste can be used as a substrate. Keeping that in mind, the present study aims for production of extracellular PLA1 by using defatted rice bran as fermentation substrate.

**MATERIALS AND METHODS**

**Isolation of microorganism from soil samples**

Twenty soil samples were collected from various sites like Dhaba, Mandir, Temples and Motor -market in different regions of Chandigarh, Mohali and Shimla. One gm of soil was suspended in 10 ml followed by serial dilution. Isolates were plated on tributyrin (0.5 % v/v) agar plate and incubated at 37ºC for 24h. Isolates producing clear zone indicate lipase positive culture.

**Qualitative assay for PLA1**

Lipase positive cultures were patched on egg-yolk (5% v/v) agar plate and incubated at 37ºC for 24h. Isolates producing clear zone indicate lipase positive culture.

**Gram staining characteristic**

A loopful of overnight grown PLA positive culture in distilled water was put on a clean glass slide and a uniform smear was prepared. The smear was heat fixed by gently heating over the burner flame. After heat fixation the smear was stained with crystal violet complex for about one minute. The slide was gently washed under tap water. Next, smear was flooded with Gram’s iodine for about one minute and then slide was again washed gently with water. Decolourization of the smear was done by covering the smear with de-staining solution for 15-20 sec. Slide was again washed with water. Lastly, the smear was stained with safranin for about 40 sec followed by washing with water. Slide was blot dried and further examined under microscope (Quasmo, India) under 100X magnification.

**PLA1 production by selected isolates via submerged fermentation**

PLA positive cultures were first subjected to submerged fermentation using Nutrient Broth (NB) supplemented with olive oil (0.5% v/v). The emulsion was formed with gum- acacia (1% w/v) in mixer grinder. An inoculum was prepared in NB until O.D reached 0.8 and was added (1% v/v) to the production medium followed by incubation at 37°C in orbital shaker (Remi orbital shaker, India) with agitation speed 160 rpm for 24h. The medium was centrifuged at 10,000 rpm for 10minutes (Sigma Laborzentrifugen, Germany) and PLA1 activity assay was carried out with the supernatant.

**Quantitative assay for PLA1 activity**

The quantitation of PLA1 activity was done by using ‘Enzchek® Phospholipase A1 assay kit’ (Molecular Probes Inc. USA) that uses PLA1 dye-labeled glycerophosphoethanolamines with dye labeled acyl chain at the sn-1 position and dinitrophenyl quencher-modified head group. The standard curve for assay was prepared using lecitase provided in the kit.

Substrate-liposome mix was prepared by adding 50µl lipid-mix {30µl of 10mM of DOPC (Dioleoylphosphatidylcholine), 30µl of 10mM DOPG (Dioleoylphosphatidylglycerol) and 30µl of 2mM PLA1 substrate} slowly and steadily to the 5 ml of reaction buffer (1X) placed on a magnetic stirrer. The reaction was started by adding 50µL of substrate-liposome mix to each well of 96-well plate already containing controls and samples. The reaction was followed by incubation at 25°C for 30 minutes in dark and fluorescence was measured with excitation at 460nm and emission at 515
nm. One Unit (U) of the enzyme corresponds to one Lecitase® Ultra unit as provided in Enzchek® Phospholipase A, assay kit.

Identification by 16S rRNA gene sequencing

Two isolates with maximum PLA activity were identified using 16S rRNA gene sequencing at IMTECH, Chandigarh, India. Genomic DNA was extracted from pure cultures using HiPurATM Bacteria Genomic DNA Miniprep Purification Spin Kit (Hi-Media, India) according to the manufacturer’s instructions. The amplification of 16S rRNA gene was achieved using the following primers: 27f (5’-AGA GTT TGA TCC TGG CTC AG-3’) and 1492r (3’-ACG GCT ACC TTG TTA CGA CTT-5’). Four sequencing primers and Big-Dye Terminator Cycle Sequencing Kit (Applied Biosystems, CA, USA) were used for the sequencing of the purified 16S rRNA gene.

Sequence assembly and phylogenetic analysis

DNA sequence assembling software SEQUENCHER™ 4.10.1 (Gene Codes Corporation, MI, USA) was used to assemble and analyse the sequence data obtained. In order to find out the similar sequences, the data was used to BLAST as query in nucleotide database of NCBI (http://www.ncbi.nlm.nih.gov/). Phylogenetic tree was constructed using Clustal W software by aligning all related and acquired sequences. Neighbour-joining method was used to measure the evolutionary distances and operated in MEGA 6 and the Kimura 2 parameter models.

PLA production by solid state fermentation

The identified and selected isolate with maximum extracellular PLA activity in submerged conditions were subjected to solid state fermentation using defatted rice bran as substrate.

Table 1. Gram Character, Hydrolysis Capacity

| S.No | Isolate number | Gram Staining             | HC     |
|------|----------------|----------------------------|--------|
| 1    | S2-1           | Isolated, thin, Gram positive rods | 0.11   |
| 2    | S3-1           | Isolated, Gram positive rods | 1.55   |
| 3    | S3-2           | Isolated, Gram positive rods | 1.37   |
| 4    | S6-3           | Very small, Gram negative rods | 1.36   |
| 5    | S7-2           | Small, Gram positive rods | 1.60   |
| 6    | S7-5           | Very small, isolated Gram negative rods | 1.72   |
| 7    | S7-6           | Isolated, Gram negative rods | 1.25   |
| 8    | S8-1           | Gram negative rods | 1.50   |
| 9    | S8-3           | Gram positive rods in chains | 3.00   |
| 10   | S9-1           | Isolated, Gram positive thick rods | 1.40   |
| 11   | S9-2           | Small, isolated, Gram negative rods | 1.25   |
| 12   | S10-2          | Isolated, Gram positive rods | 1.44   |
| 13   | S10-7          | Isolated, Gram positive rods | 1.20   |
| 14   | S11-4          | Very small, Gram negative rods | 1.18   |
| 15   | S12-2          | Small, Gram positive rods | 1.75   |
| 16   | S12-3          | Gram positive rods in chains | 1.44   |
| 17   | S13-1          | Gram negative rods | 1.37   |
| 18   | S14-1          | Gram negative, coco-bacilli, | 1.90   |
| 19   | S14-2          | Gram positive cocci | 1.28   |
| 20   | S14-3          | Very small, Gram negative rods | 1.44   |
| 21   | S15-1          | Gram negative cocci | 1.87   |
| 22   | S15-3          | Small, thin, Gram negative rods | 1.76   |
| 23   | S16-3          | Gram negative rods | 1.50   |
| 24   | S16-4          | Very small, Gram positive cocci | 1.75   |
| 25   | S16-5          | Gram negative cocci | 1.62   |
| 26   | S17-4          | Gram negative cocci | 1.05   |
| 27   | S17-5          | Gram positive cocci | 1.60   |
| 28   | S18-1          | Gram negative thin small rods | 1.50   |
| 29   | S20-2          | Gram positive rods, isolated | 1.36   |
| 30   | S20-3          | Gram positive rods, in chains | 1.38   |
| 31   | S20-4          | Gram positive rods, in chains | 1.50   |
The olive oil emulsion (0.5% v/v) was formed in mixer grinder using 1% (w/v) gum acacia as emulsifying agent. An inoculum (1.5ml) was obtained in NB until O.D_{600} reaches 0.8 and added to autoclaved defatted rice bran (5gm) with 6ml of olive oil emulsion maintaining surface to moisture content ratio of 1:1.5. The flasks were incubated at 37°C under static conditions for 48h. The content of the flask were suspended in 100ml of distilled water and kept at shaking 160rpm for 1h. Then centrifugation was carried out at 10,000rpm for 10minutes and supernatant was used for PLA₁ activity assay.

Classical approach for optimization in solid state fermentation

Different parameters like incubation time, incubation temperature, moisture conditions, oil sources, nitrogen and carbon sources were taken in account to optimize the culture conditions for maximum production of PLA₁ by solid state fermentation.

**Incubation time**

For optimization of incubation time, medium was prepared consisting of 5g defatted rice bran with 6 ml olive oil emulsion (0.5%, pH 7). The stirring of olive oil with water was carried out in a mixer grinder with 1% (w/v) gum acacia to prepare olive oil emulsion. The medium was sterilized and inoculum (1.5 ml) was added and incubated at 37°C for 24 to 120h.

**Incubation temperature**

The optimum temperature was determined by incubating the same medium at different temperatures (25-45°C) for 48h.

**Moisture content**

The moisture content in the medium was optimized by incubating 5gm defatted rice bran and inoculums (1.5 ml) with following substrate to moisture ratio (rice bran : olive emulsion + inoculums): 1:1, 1:1.5; 1:2; 1:2.5; 1:3; 1:3.5 (pH 7) and medium was incubated at 37°C, for 48h.

**Oil source**

Different oil sources (0.5 % v/v) were used as inducer for PLA₁ production like sunflower oil, olive oil, soy-bean and groundnut oil. The moisture content of the medium was maintained at 1:1.5 by using 6ml of oil emulsion with 1.5ml of inoculums in 5gm defatted rice bran and incubated at 37°C for 48h.

**Carbon source**

The carbon source in the medium was optimized by taking different carbon sources (1% w/v) like glucose, sucrose, fructose, xylose, maltose in the same medium and incubated at 37°C for 48h.

**Nitrogen source**

The nitrogen source in the medium was optimized using 1% (w/v) of malt extract, yeast extract, peptone, urea and soybean meal in the medium and was incubated at 37°C for 48h under static conditions.

**RESULTS AND DISCUSSION**

**Isolation of cultures from soil samples**

In total, 82 lipase positive isolates were screened on the basis of degradation of tributyrin.
in lipase plate assay. Out of 82, only 31 isolates were found to have PLA activity on the basis of formation clear zone on egg-yolk agar plate. The Gram staining and HC on egg-yolk agar plate of the selected 31 isolates is displayed in Table 1.

**PLA production by submerged fermentation**

The PLA positive isolates were subjected to submerged fermentation. The supernatant activity of 31 isolates (U/ml) is displayed in Table 2.

**Identification of bacteria by 16S rRNA sequencing**

Two isolates with maximum PLA activity S8-3 and S14-1 were selected for identification. The 16S rRNA gene sequence of the S8-3 displayed 99% identity with *Bacillus subtilis* subsp. *inaquosorum* KCTC 13429 and S14-1 strain showed 99.78% identity with *Acinetobacter radioresistens* DSM 6976T. The S14-1 was not used for further studies due to its pathogenic nature. Nucleotide sequence obtained after sequencing of S8-3 was submitted in GENBANK under Accession no. KY088040 with strain name RG1. The Phylogenetic tree of both isolates was constructed by Neighbor joining method (Fig. 1) on the basis of sequence analysis.

### Optimization of the cultural conditions

*Bacillus subtilis* subsp. *inaquosorum* RG1 was subjected to solid state fermentation for PLA production. The optimization was carried out by classical approach by taking different factors into account including incubation temperature, incubation time, moisture content, different oils, different nitrogen and carbon sources (Figure 2 and Figure 3). Maximum PLA activity of 51.55 U/gm was obtained after the incubation of 48h at 37°C with moisture content of 1:1.5 using glucose and peptone as carbon and nitrogen source respectively with 0.5% olive oil as an inducer.

PLA production (23.2 U/gm) was observed after incubation of 48h and after that there is a decrease in PLA, that may be due to decrease in viable number of cells\(^{23}\) and depletion of nutrient availability\(^{24}\).

Substrate to moisture ratio of 1:1.5 was found to be more suitable for PLA production. The moisture content ratio plays a crucial role in the growth of microorganism ultimately affecting enzyme productions by solid state fermentation\(^{25}\).

Temperature is a crucial factor for optimum enzymatic activity and overall metabolism\(^{26}\). In our study, optimum PLA production was observed at
37°C. Our results are in accordance with previous reports wherein 37°C was the best temperature for maximum PLA₁ production in *Pseudomonas gessardii* and *Trichosporon sp.*²⁷.

In addition to physical factors, medium composition also affects the enzyme production as it play an important role in overall growth of microorganisms. Various carbon and nitrogen source and different oils have different effects on growth of microorganisms and finally influenced enzyme production. Thus, these parameters need to be

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**Fig. 2.** Effect of incubation conditions on PLA₁ production by *Bacillus subtilis* subsp. *inaquosorum* (a) incubation time (24h-120h) (b) incubation temperature (25-45°C) (c) moisture content (1:1-1:3.5) via solid state fermentation

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**Fig. 3.** Effect of medium composition on PLA₁ production by *Bacillus subtilis* subsp. *inaquosorum* (a) oil variation (b) carbon sources (d) nitrogen sources

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optimized for better production yields of the desired enzyme.

In our study, 0.5% of olive oil is observed to be the best inducer for PLA production. Similar results were observed for lipase production from Bacillus sp.28,29. Among the carbon and nitrogen sources used, glucose and peptone were found to be the best carbon and nitrogen source respectively. In contrast, xylose and ammonium sulphate were found as good carbon and nitrogen source for PLA in Serriatia sp.11.

The use of PLA has been restricted owing to its low stability and less availability. In addition, crystallographic and structural data for most of the PLAs, have not been obtained yet. So there is an increasing trend to isolate new microorganisms for PLA production which can provide a platform to further improve these microbes to make them competent enough to be exploited at industrial scale.

CONCLUSION

In the present study, a PLA producing microorganism was isolated from soil and identified as Bacillus subtilis subsp. inaquosorum. Maximum PLA production of 51.55 U/gm was observed which is 2.15 folds more than un-optimized medium via solid state fermentation. This is the first report wherein PLA was produced using one factor at a time approach. Moreover, the present study also provided a clean approach to successfully produce high titers of enzyme and will reduce burden of the environment by utilizing waste as substrate.

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