Production and Partial Characterization of Lipase from *Pseudomonas putida*

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Abstract

The production of lipase from *Pseudomonas putida* 922 was optimized by modifying various physical parameters such as carbon source, nitrogen source, pH, salt concentration and biochemical parameters of the production medium such as temperature and incubation time of the growth medium. Oil cakes were also used as carbon source to check for an increased production of the enzyme. The bacterium was found to have a maximal growth at pH 10 with the enzyme production being highest (24 U/ml) after 48 hours at 30°C and pH 10. The optimum composition of the medium was mustard oil cake as carbon source, yeast extract or peptone as nitrogen source and 1% sodium chloride concentration. Partial characterization of the enzyme was carried out where the optimum working pH and temperature was found to be 10 and 40°C, respectively. Enzyme stability was found to lie in the pH and temperature ranges of 5-11 and 30-40°C, respectively. Partial purification of the enzyme was carried out at 80% ammonium sulphate saturation. Molecular mass of lipase was determined by SDS PAGE and found to be 45 kDa.

Keywords: Lipase; *Pseudomonas putida*; Optimization; Oil cakes; Characterization

Introduction

Lipases (triacylglycerol hydrolases) are a unique type of esterases that belong to the class 3 of enzymes and due to their specificity for hydrolyzing carboxylic acid ester bonds they have been assigned the EC number 3.1.1.3. The biological role of lipases is to carry out the hydrolysis of lipids such as triacylglycerols resulting in the production of free fatty acids along with diacylglycerols, mono-acyl glycerols or glycerol [1].

Lipases are an important class of enzymes that help micro-organisms to derive energy from triacylglycerols. Lipases can be extracellular or intracellular in nature. This nature varies from organism to organism, however, majority of the bacteria have been found to secrete extracellular lipases [2].

Lipases are found abundantly in nature and have been found to be naturally synthesized by some plants such as *Hibiscus cannabinus*, animals such as some fish and insects, fungi such as *Aspergillus sp.* and bacteria such as that produced by *Acinetobacter sp.*, *Bacillus sp.* [3] and *Pseudomonas sp.* [4].

However, lipases from microbial sources are commercially more significant since they have a lower cost of production, higher stability and a higher availability than other sources [5].

Bacterial lipases have been found to be influenced by physiochemical and nutritional factors such as temperature, pH, nitrogen and carbon sources, presence of inorganic salts, lipids and usage of agitation [6]. The major content of lipase production media is the carbon source that also acts as an inducer for lipase production [7] whereas nitrogen sources and other components of the media help in regulation of the growth of the producer organism and therefore the fermentation process [8]. Microbial lipase production is usually carried out using submerged fermentation technique [9], however solid state fermentation methods have also been found to be beneficial [10]. Lipid carbon sources have been found to be generally important for obtaining a high lipolytic enzyme yield although a few researchers have obtained substantial lipase yields without using fats or oils [4].

Lipase is one of the versatile enzymes that can be used in a number of industries such as the food industry [11]. It is also used as a biosensor for bioremediation purposes [12]. Lipase is also an effective part of detergents used in industrial and domestic laundry and dishwashing [13] while also being used in the paper industry [14].

As there is versatility in the lipase enzyme, and since the lipase from *Pseudomonas putida* has not been previously been worked upon at the production level. Therefore, the results of the present investigation will provide an insight into improving the production of lipase by aiding the development of a feasible medium and characterization of lipase.

Materials and Methods

Chemicals

All chemicals were of analytical grade and procured from Sigma chemicals (USA), Fisher Scientific (UK), Merck (Germany) and Acros Organics (Belgium).

Microorganism and culture maintenance

*Pseudomonas putida* 922 was obtained from the culture bank of IMBT, TUG, Austria and was refreshed in Lauria Bertani (LB) agar slants that were kept at 30°C for 48 hours. The slants were then placed at 4°C in a cool lab (write maker and model). The master culture was maintained in glycerol stocks at -80°C.
Preparation of inoculum

Bacterial slants that were 48 hours old were used as inoculum where loopful of bacteria were aseptically suspended in the sterilized fermentation medium.

Fermentation technique

The bacterium was cultured using submerged fermentation and grown in sterilized medium containing 5 g of wheat bran dissolved in 50 ml of freshly prepared minimal medium containing Na$_2$HPO$_4$.2H$_2$O (6 g/L), NaCl (5 g/L), NH$_4$Cl (2 g/L), MgSO$_4$.7H$_2$O (0.1 g/L) and KH$_2$PO$_4$ (3 g/L) and incubated at 37°C at 200rpm for 48 hours in a shaking incubator.

Analytical methods

**Enzyme extraction**: After completion of incubation time, 50 ml of 0.1M phosphate buffer (pH 8) was added to the growth medium and incubated for another hour. Enzyme extraction was carried out by centrifugation of the growth medium at 5000 rpm for 5 minutes after which the supernatant was harvested and used for lipase estimation.

**Enzyme assay**: Enzyme Assay was carried out using titration method that has been previously used by Venkateshwarlu and Reddy. The reaction mixture consisted of 0.5 ml of 0.1 M phosphate buffer, 0.5 ml of enzyme extract and 1 ml of substrate (10% v/v olive oil emulsified with 10% w/v gum acacia in 0.1Ml phosphate buffer, pH 8). The experimental tube was incubated for 10 minutes at 30°C whereas no incubation time was provided for the control tube. Inhibition of the enzyme was carried out by alcohol: acetone in 1:1 ratio. Using few drops of phenolphthalein indicator, titration was carried out against 0.1 N sodium hydroxide until pink color appeared.

The enzyme was estimated by the following formula;

$$\text{Enzyme Units (U/ml/min)} = \Delta V \times N \times 1000 \times \text{Dilution factor}$$

Where, $\Delta V$ is (V2-V1) or the volume of NaOH used for experimental – the volume of NaOH used for the control.

N is the normality of NaOH used for titration.

Dilution factor is obtained due to the usage of extraction buffer for extraction

$$V\text{}(\text{sample})$$ is the amount of enzyme extract taken for the reaction mixture

T (min) is the time of incubation in minutes.

A unit (U) of enzyme is taken as the amount of enzyme (lipase) that releases 1μmol of free fatty acids per minute under specific conditions of assay.

**Protein estimation**: Protein estimation was carried out using the Bradford method. Absorbance of enzyme extract was used to calculate the amount of protein present in the enzyme extract.

The specific activity of the enzyme was calculated using the following formula

$$\text{Specific activity (U/mg of protein)} = \frac{\text{Enzyme activity (U)}}{\text{Protein content (mg)}}$$

Optimization of parameters for enhanced lipase production

The optimization was carried out by using 5 g of wheat bran as a carbon source in the medium employed for the optimization of temperature, pH, incubation time, salt concentration and nitrogen sources. Olive oil (1% (v/v)) was also used as an inducer for enhanced lipase production.

**Optimization of pH for microbial growth**: Microbial growth was optimized by inoculating the bacteria in an autoclaved medium that had pH varying from 5.0-11.0 by dissolving components of the minimal medium in the buffer of desired pH.

**Optimization of temperature**: Temperature optimization was carried out by growing the bacterium at temperatures 30, 35, 37 and 40°C in a shaking incubator at 200 rpm.

**Optimization of incubation time**: Incubation time was optimized by incubating the bacterium in wheat bran containing growth medium for 24, 48, 72 and 96 hours.

**Optimization of carbon sources**: Carbon sources were optimized by using oil cakes and carbohydrates in place of wheat bran. The oil cakes included mustard oil cake, almond oil cake, canola oil cake whereas the carbohydrates used included glucose, sucrose, fructose and maltose as 10% w/v in minimal medium.

**Optimization of nitrogen sources**: Nitrogen sources where optimized by replacing ammonium chloride with ammonium sulphate, yeast extract, malt extract, beef extract and peptone as 0.1 g/L.

**Optimization of salt concentration**: Different concentrations of sodium chloride such as 0.1, 0.5, 1.0 and 1.5 were used for identifying concentration at which high lipase production could be achieved, where 1% is the concentration present in the minimal medium used in all experiments, i.e. 5 g/L.

Production of lipase using optimized medium

The parameters used above that gave maximum lipase production were then all used together in order to provide the necessary conditions for maximal lipase production after which the enzyme was harvested and then assayed.

Partial characterization of lipase

Enzyme characterization was also carried out by determining its activity at various temperatures where the pH of the buffer remained constant, and at various pH where the temperature was kept constant. Lipase temperature and pH stability was also evaluated by keeping the enzyme for 1 hr at different temperatures and pH, after which enzyme estimation was carried out.

Partial purification of lipase

Ammonium sulphate precipitation was used in order to partially purify crude lipase so that the molecular weight of the enzyme could be determined. Solid ammonium sulphate with varying saturations (30, 40, 50, 60, 70 and 80%) was added slowly to the crude enzyme with constant stirring on a magnetic plate at 4°C and kept overnight. After that, sample was centrifuged at 12,000 rpm for 15 minutes. Supernatant and pellets were assayed for lipase activity.
Determination of molecular weight of lipase

Molecular weight of the enzyme was determined by SDS-PAGE.

Results and Discussion

Effect of pH on microbial growth

The bacterial growth was determined in the medium of different pH ranging from 5.0 to 11.0. The growth of bacteria was determined by determining absorbance at Spectrophotometer at 600 nm. Maximum absorbance obtained was 18.506 at a pH of 10 while minimum absorbance was obtained at pH 6.0 (Figure 1).

Effect of temperature on Lipase production

The temperatures used for studying the effect of temperature on lipase production included 25, 30, 35, 37, and 40°C. Lipase assay was carried out that revealed that the maximum production of lipase occurred at a temperature of 30°C and gave 24 U/mL (Figure 2).

Effect of carbon source

Numerous carbon sources such as oil cakes and carbohydrates were used for identifying the most suitable substrate for lipase production. Amongst the oil cakes mustard oil cake gave maximum lipase production (60.5 U/mL) while almond oil cake gave minimum lipase production (5.5 U/mL) (Figure 4). From among the carbohydrates the highest enzyme production was obtained by glucose and fructose (9.6 U/mL each) with maltose giving no enzyme production (Figure 5).
Effect of carbohydrates as carbon source on lipase production

From amongst the nitrogen sources used, highest production of lipase was obtained by using Peptone and Yeast Extract, each of which gave 48 U/mL. Lowest amount of lipase was produced by Ammonium Sulphate which gave 6 U/mL (Figure 6).

Effect of nitrogen sources

From amongst the nitrogen sources used, highest production of lipase was obtained by using Peptone and Yeast Extract, each of which gave 48 U/mL. Lowest amount of lipase was produced by Ammonium Sulphate which gave 6 U/mL (Figure 6).

Effect of salt concentration

Sodium chloride was used whose concentration was changed in the production medium where 1.0% is the salt concentration used in the minimal medium. Highest enzyme units were obtained when 1.0% of Sodium chloride was used (24 U/mL) while lowest enzyme units were obtained when 0.10% of salt concentration was used (6 U/mL) (Figure 7).

Production of lipase using optimized medium

Lipase production was enhanced by using optimized conditions and substances such as carbon and nitrogen sources that were meant to allow maximum growth of microorganism and lipase production as well. The results showed that the optimized medium produced 26 U/mL with a specific activity of 119.54 U/mg of protein (Figure 8).

Partial characterization of lipase

Enzyme characterization was carried out by using the enzyme extract from the optimized medium and varying conditions in order to identify the optimum temperature (Figure 9) and pH (Figure 10) for enzyme activity along with the enzyme's stability at different temperature (Figure 11) and pH (Figure 12).

The optimum working temperature of the enzyme was found to be at 40°C (56.88 U/mL) while the enzyme activity was found to decrease below and above 40°C where lipase may inactivate or achieve lower activity, respectively. The optimum working pH of the enzyme was found to be at pH 10 (26 U/mL) implying that the enzyme had the ability to work at fairly high alkaline conditions.
Thermostability of the enzyme was also assessed using different temperatures where the enzyme was found to be stable between 30 and 45°C whereas above and below this range the enzyme was unstable. Enzyme stability was also determined at different pH where the enzyme was found to be stable from pH 5 to 10.

Partial purification of lipase

Partial purification of lipase was carried out using ammonium sulphate precipitation. 5.8 fold purification was obtained with an overall yield of 32.6% when 80% of ammonium sulphate concentration was used. Dialysis was carried out to remove the remaining salts to obtain partially purified enzyme.

Determination of molecular weight of lipase

Partially purified enzyme was treated with SDS-PAGE in order to determine its molecular weight. The figure below shows (a) M represents marker of ACT-IDMW-24, (b) L1 represents lipase which is precipitated by ammonium sulphate at 80% saturation, (c) L2 represents crude enzyme. The molecular mass of lipase was found to be 45 kDa.
Discussion

Varying pH of the growth medium indicated that P. putida grows well in an alkaline medium and its growth decreases in strongly alkaline and acidic medium. Somewhat similar results were obtained by Patil and Chaudhri [1,5] who showed that P. aeruginosa grew best in pH range of 8-10.

By changing the incubation temperature of the growth medium, the optimum temperature for lipase production was identified as 30°C. Increasing or decreasing the temperature causes a decrease in enzyme units showing that P. putida is a mesophilic bacterium 30°C. Similar results have been shown by Padhiar et al. [15] for P. aeruginosa and Kiran et al. [16] who found 30°C as the optimum temperature for enzyme production from Pseudomonas sp.

Increasing the incubation time showed that by 24 hours lipase production had been initiated and at 48 hours the bacterium had obtained logarithmic growth phase. After 48 hours the enzyme units tend to decrease suggesting that enzyme may have been either degraded or become non-functional due to decrease in lipidic substrate or increase in acidity after consumption of medium. Such a trend has also been reported by Chigusa et al. [17] with P. aeruginosa EF2 and Kathiravan et al. [18].

When different carbon sources such as carbohydrates and oil cakes were used, results suggest that carbohydrates were not able to stimulate lipase production as oil cakes were able to do so. Maximum enzyme production with mustard oil cake suggests that mustard oil cake is a better inducer for lipase than coconut oil cake and other carbon sources used. Such results have also been reported by Joseph et al. [18] when they used oil cakes as substrates for lipase production. While with carbohydrates Kathiravan et al. [18] found glucose and fructose giving similar results while maltose inhibited lipase production. They are also similar to results by Pogaku et al. [19].

Results using different nitrogen sources show that yeast extract and peptone were better for lipase production than other nitrogen sources. Yeast extract and peptone gave similar results when used by Sirsisha et al. [20], Mobarak-Qamsari et al. [21] and Lau et al. [22].

When different salt conditions were used, the result suggests that high salt concentration tend to inhibit lipase production while low salt concentration does not aid in lipase production. Such results were also obtained by Sangeetha et al. [23] who used salt concentration to obtain high lipase yield.

The optimized medium was found to produce lesser enzyme units than other media used for optimization because of the pH of the optimized medium that was at 10 which was found to be better for growth of P. putida but not optimal for lipase production. The conditions of pH may not have been suitable for maximal lipase production. Similar reports were also given by Sekhon et al. [24] who found that good bacterial growth occurred at pH 4-11 whereas lipase production was maximum at pH 7 and decreased above or below this pH.

The results of optimum working temperature for lipase are confirmed by studies done by Gilbert et al. [25] showed that lipase activity decreased after 45°C while Zaliha et al. [26] found lipase activity to be optimum above 40°C. The results for varying optimum pH coincide with various studies carried out by Mobarak-Qamsari et al. [21] which reveal similar results that the enzyme has the ability to work at pH 9 and 10. Thermostability results are similar to such studies carried out on lipase by Ranjitha et al. [27] where the enzyme was stable at 30°C while pH stability studies also agree with studies carried out by Annamalai et al. [28] who also observed that the lipase from Bacillus licheniformis was stable at pH 9.

The molecular mass of lipase obtained from Pseudomonas putida was determined by utilizing SDS-PAGE. The results revealed that molecular mass of lipase was 45 kDa. Less number of bands was observed in partially purified sample compared to the bands of crude enzyme, which proved that lipase partial purification was exceptional. Marker used during procedure was ACT-IDMW-24 (Figure13). Longshaw et al. [29] identified a lipase gene in Staphylococcus epidermidis having molecular size of 45 kDa.

Conclusion

In the present work lipase from Pseudomonas putida has been found to give substantial production of lipase by submerged fermentation using mustard oil cake and yeast extract or peptone in production medium and differs from other lipases that it has a good stability at a wide temperature. It also is significant that it gives good activity in alkaline pH and high temperature therefore making it suitable to be used as a component in laundry detergents etc.

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