Research Article

Production of Cold-Active Bacterial Lipases through Semisolid State Fermentation Using Oil Cakes

Babu Joseph,1,2 Supriya Upadhyaya,1,3 and Pramod Ramteke1,4

1 Department of Microbiology and Microbial Technology, College of Biotechnology and Allied Sciences, Allahabad Agricultural Institute-Deemed University, Allahabad-211007, India
2 College of Applied Medical Sciences, Shaqra University, Shaqra 11961, Saudi Arabia
3 Department of Microbiology, Institute of Medical Sciences, Banaras Hindu University, Varanasi 221 005, India
4 Department of Biological Sciences, School of Basic Sciences, Sam Higginbottom Institute of Agriculture, Technology and Sciences, Uttar Pradesh, Allahabad 211 007, India

Correspondence should be addressed to Babu Joseph, babuaaidu@yahoo.co.in

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Production of cold active lipase by semisolid state fermentation involves the use of agroindustrial residues. In the present study, semisolid state fermentation was carried out for the production of cold active lipase using Micrococcus roseus, isolated from soil samples of Gangotri glaciers, Western Himalayas. Among various substrate tested, groundnut oil cake (GOC) favored maximal yield of lipases at 15 ± 1°C within 48 h. Supplementation of glucose 1% (w/v) as additional carbon source and ammonium nitrate 2% (w/v) as additional nitrogen source enhanced production of lipase. Addition of triglycerides 0.5% (v/v) tends to repress the lipase production. Further mixed preparation of groundnut oil cake (GOC) along with mustard oil cake (MOC) in the ratio of 1 : 1, and its optimization resulted in improved production of cold active lipase. The enzyme exhibited maximum activity at 10–15°C and was stable at temperatures lower than 30°C. The lipase exhibited optimum activity at pH 8 and showed more than 60% stability at pH 9. Semisolid state fermentation process by utilizing agroindustrial wastes will direct to large-scale commercialization of lipase catalyzed process in cost-effective systems.

1. Introduction

The Gangotri is the second largest Himalayan glacier, situated in the Uttarkashi of Uttarakhand (India) between 30° 44’ and 30° 56’ N and 79° 4’ and 79° 15’ E, draining in northwestern direction, is an excellent source of psychrophiles. Gangotri glacier is around 30 km in length, 143 km² in area and it is the largest valley type glacier in the western Himalaya [1]. The temperature is 2–5°C in summer and subzero in winter. Psychrophiles have attracted attention as source of enzymes with potential for low temperature catalysis. Indeed a variety of cold active enzymes have been found in psychrophiles [2–4]. One such example is cold active lipases, which are largely distributed in microorganisms existing at low temperatures nearly 5°C. Lipases (triacylglycerol acylhydrolase, E. C. 3.1.1.3) are hydrolytic enzymes, which act on the carboxyl ester bonds present in acylglycerol to liberate fatty acids and glycerol. The knowledge of cold adapted lipolytic enzymes in industrial applications is increasing at a rapid and exciting rate. Cold active lipases are one of the important and widely used enzymes whose spectrum of applications has widened in many industries such as in detergent formulations, food industry, leather processing, environmental bioremediations, and fine chemical synthesis as well as in pharmaceutical industries [5]. However, their production cost limits their industrial use [6, 7]. Therefore, it is of interest to increase the productivity of fermentation processes by optimization of culture conditions. Since the raw materials employed in the culture medium contribute to total production costs, the reduction in the substrate cost would be a suitable strategy to increase the productivity of the process [8]. Conventionally, submerged fermentation is employed for production of bacterial lipase. However, emerging trends emphasize that semi-solid and solid-state fermentation systems have real potential for the judicious management of bacteria in industry.
Semisolid state fermentation (SmSSF) involves growth of microorganism on moist solid substrates in the absence of free flowing water. In SmSSF, the nature of the solid support material is an important parameter that influences the product yield and, consequently, there is a continuous search for newer and better substrates [9, 10]. Lipase production by SmSSF is suitable since it offers many advantages over submerged fermentation (SmF), which include high productivity, higher product concentration, simpler equipments and the use of low cost substrates as agro-industry waste products [11]. India is one of the world’s leading oilseeds producing country. Oil cakes have high nutritional value, as they possess high protein content (ranging from 15 to 50%). They are economically cheap, stable and dependable sources available in large quantities throughout year. The potential role of oil cake as a substrate for fermentation has been established [12]. Studies with oil cake extract showed significant results for the overproduction of lipases [13]. Moreover, the blending of different substrates shows still better performance, by providing more suitable environment for microbial growth. In the present study, the ability of Micrococcus roseus to produce extracellular cold active lipases under semi-solid state fermentation using oil cakes has been evaluated. This method may result in selecting an ideal substrate for economic production. Moreover, the maximum enzyme production by blending different substrate has been studied.

2. Materials and Methods

2.1. Bacterial Strain. Lipolytic bacterial strains were isolated from soil samples of Gangotri glacier, Western Himalayas on tributyrin agar plates [14] and observed for zone of clearance around the colonies. The organism which produced larger zone was isolated and identified as Micrococcus roseus by morphological and biochemical characteristics based on Bergey’s Manual of Determinative Bacteriology [15]. The stock culture was maintained on Nutrient broth (Hi-Media) and glycerol (50:50% v/v) at −20°C. Working cultures were prepared by two successive transfers of stock culture to nutrient broth for 24 h at 15±1°C.

2.2. Inoculum Preparation. A loopful of cells from freshly grown culture of the potential lipolytic isolate was transferred to a 250 mL conical flask containing 100 mL nutrient broth. It served as a seed culture for further use as an inoculum after 48 h of incubation at 15 ± 1°C. Five milliliters of inoculum (10⁶ cells/mL) was added to the sterilized medium for fermentation.

2.3. Semi-Solid State Fermentation. The substrates used for the production of cold active lipases were different oil cakes such as groundnut (GOC), coconut (COC), gingili (GngOC), mustard (MOC), and castor (CastOC) procured from the local market. Dry oil cake (10 g) was taken in 250 mL Erlenmeyer flask with 90 mL of double distilled water. The contents of the flask were mixed thoroughly and autoclaved at 121°C for 15 min.

2.4. Extraction of Enzyme. After fermentation, phosphate buffer (pH 8) was added (1:1) to the fermented medium and the enzyme is extracted by centrifugation at 10,000 rpm at 4°C for 10 min. The clear supernatant obtained was used as crude enzyme and stored in sterilized vials for further use.

2.5. Lipase Assay. Lipase activity was assayed using p-nitrophenyl palmitate (pNPP) as a substrate as described by Winkler and Stuckmann [16]. Briefly, substrate solution containing phosphate buffer (90 mL) with Gum Arabic (100 mg), and sodium deoxycholate (207 mg) along with 30 mg of pNPP in 10 mL of isopropanol was preincubated with the crude enzyme at 15 ± 1°C. The release of p-nitrophenol (pNP) was measured spectrophotometrically at 405 nm. One unit of lipase activity was defined as the amount of enzyme releasing 1 μmol pNP under standard assay conditions.

2.6. Optimization of Production Parameters for SmSSF Using Oil Cakes. Optimization of process parameters and manipulation of media composition are one of the most important techniques used for the overproduction of lipase to meet industrial demands. Optimization was carried out through modification of several growth parameters. The effect of an individual parameter was standardized at a time before standardizing the next parameter. The various parameters optimized for obtaining maximal lipase yield were, incubation time (24, 48, 72 and 96 h), temperature (10, 15, 25, 30 and 35°C), and pH (ranging from 5.0–10.0). The effect of lipid materials as triglycerides 0.5% v/v such as castor oil, soybean oil, olive oil, and mustard oil was determined. Further the effect of supplementation of carbon sources (1% w/v) such as glucose, sucrose, lactose, maltose, and nitrogen sources (1% w/v), that is, ammonium nitrate, potassium nitrate, peptone, beef extract, yeast extract, in the SmSSF system were also examined for production of cold active lipase. In the above process, after optimization of incubation time, oil cake showing highest lipase activity was selected as potential substrate for the SmSSF system, and further optimization of all other parameters was carried out with this substrate. For each step lipase activity was assayed to know the optimal yield.

2.7. Mixed Substrate Fermentation. In view of above experiments, studies were made to evaluate the mixed substrate fermentation by mixing the oil cake showing maximum lipase yield with other oil cakes in ratio 1:1. Best mixed preparation was selected; further, the above optimized ingredients were tested for mixed substrate enrichment. The enriched media were designated as media A to E.

- Media A: Combination of mixed substrate showing maximum production of cold active lipase.
- Media B: Media A + triglyceride (0.5% v/v).
- Media C: Media A + Carbon source (1% w/v).
- Media D: Media A + Nitrogen source (2% w/v).
- Media E: Media A + Carbon + Nitrogen source.
2.8. Characterization of Lipase. Activity of the crude lipase at different pH values was measured adjusting pH of the reaction mixture using (0.1 M) of following buffers: citrate buffer (pH 5.0–6.5), Tris-chloride buffer (pH 7.0–9.0), and Glycine-NaOH buffer (pH 10). The enzyme activity was assayed by method described before. To study the stability at different pH, cold active lipase was dissolved in above-mentioned buffers. These enzyme solutions were preincubated at 15°C for 1 h and relative activity was measured. To study the effect of temperature on activity of lipase, reaction mixture was incubated at different temperatures ranging from 5–50°C for 15 min and activity was determined. To study the lipase stability at different temperature, lipase was dissolved in 50 mM phosphate buffer (pH 8.0), pre-incubated at different temperatures ranging from 0 to 50°C for 1 h, rapidly relative activity was measured by the standard assay procedure.

3. Results and Discussion

Twelve lipolytic bacterial colonies were isolated on tributyrin agar medium from the five different soil samples of Gangotri glacier (data not shown). Microbiological analysis of soil samples showed that the soil of glacier region contains high bacterial count growing at 15 ± 1°C. Among these 12 lipolytic strains, isolate CAL-7 showing clear zone of maximum diameter on tributyrin agar medium at 15 ± 1°C was selected as potential strain for cold active lipase production. The isolate CAL-7 was later identified as M. roseus on the basis of biochemical tests performed (Table 1).

Selection of a suitable substrate for the production of enzyme is a primary-key factor and an extremely significant step. In the present study, the production of cold active bacterial lipase by M. roseus was tested in semi-solid state fermentation using various oil cakes as substrate. The effect of incubation time showed that there is large difference in the production of lipase with different oil cakes as substrates (Figure 1). The maximum yield of lipase (1.66 U/gds) was noted in groundnut oil cake among the various substrate tested. The suitability of GOC may be due to the high content of crude protein. A low level of lipase activity was obtained in the earlier stages of incubation which steadily reached maximum level by 48 h of incubation. Beyond 48 h, a steep loss in the production was observed which could be due to depletion of nutrients. Several workers also reported the effect of incubation time on cold active lipase production. Serratia marcescens produced lipase after 6 days of incubation [17] and Psychrobacter sp. Ant300 [18] and Pseudoalteromonas sp. wp27 [19] synthesized cold active lipase after 14 days of incubation. However, in the present study, M. roseus was able to produce cold active lipase within 48 h of incubation.

When the lipase yield in semi-solid state fermentation system was compared with that of submerged fermentation system the lipase activity was reported to show 1.6-fold

### Table 1: Morphological and biochemical characteristics shown by the potential isolate CAL-7.

| Tests                        | Results               |
|------------------------------|-----------------------|
| Colony morphology           | Round/circular        |
| Configuration               |                       |
| Margin                      | Entire                |
| Elevation                   | Convex                |
| Surface                     | Smooth/glistening     |
| Density                     | Opaque                |
| Pigments                    | Rosy red              |
| Grams reaction              | Positive              |
| Shape                       | Cocci                 |
| Size                        | Small                 |
| Arrangement                 | Single or in pairs    |
| Spore                       | Absent                |
| Biochemical tests           |                       |
| Catalase                    | +                     |
| Oxidase                     | –                     |
| Indole                      | –                     |
| Methyl red                  | +                     |
| Voges Prausker              | –                     |
| Citrate utilization         | –                     |
| Urease                      | –                     |
| Nitrate reduction           | +                     |
| Gelatin hydrolysis          | –                     |
| Casein hydrolysis           | –                     |
| Starch hydrolysis           | –                     |
| Cween 80 hydrolysis         | –                     |
| Hydrogen sulphide production| –                     |
| Motility test               | –/w                   |
| Sugar fermentation          |                       |
| Glucose                     | A                     |
| Xylose                      | A                     |
| Sucrose                     | A                     |
| Fructose                    | A                     |
| Maltose                     | A                     |
| Galactose                   | A                     |
| Mannitol                    | A                     |

+: Positive, –: Negative, A: Acid, w: weak.
increase in SmSSF system (data not shown). Similar result was reported by [20], which pointed out the superiority of solid state fermentation over traditional suspension culture. Therefore the SmSSF showing higher lipase yield was found to be an appropriate system for the overproduction of cold active lipase for industrial benefits.

*Micrococcus roseus* was found to be a psychrophilic microorganism having an optimum temperature for growth at 15 ± 1°C with maximum lipase yield. On increasing the temperature to 35°C, cold active lipase production was tremendously decreased (Figure 2). The absence of lipase activity in the culture supernatant above 35°C is likely due to the thermolability of enzyme. The cold adapted microorganisms tend to have good growth at low temperature and production of cold active lipase is considered to be temperature dependent and thermolabile [21].

In the present case the isolate was found to be growing and producing the lipases over wide range of pH values (Figure 3). The maximum production of lipase was obtained at pH 8.0 (3.66 U/gds). An alkaline stable lipase is used for the enzymatic hydrolysis of tributyrin [22]. The cold active lipase produced owing to its alkaline nature seems to be of considerable importance in industrial processes such as leather processing, sewage treatment, and detergent formulations.

The supplementation of different triglycerides, that is, castor oil, soybean oil, olive oil, and mustard oil at 0.5% concentration was not desirable as it suppressed the production of lipase in SmSSF medium (Table 2). This might be because the substrate itself would have provided the necessary lipid source. It might be due to the fact that higher oil content resulted in the formation of a biphasic system which prevented not only the accessibility of water to the microorganism but also prevented the oxygen transfer and nutrient assimilation by the microorganism from the substrate. Butter oil, corn oil, or olive oil inhibited production of lipase by *Penicillium roqueforti* [23]. Further addition of inducers can be avoided which offers economic benefits for industrial scale production of cold active lipases. The impact of additional carbon sources at 1% (w/v) was studied (Table 2). It was found that, glucose when used as an additional carbon source induced maximum production of cold active lipase (4.66 U/gds). However, sucrose, lactose and maltose were found to be notably repressing the lipase production. Glucose is an easily available cheap carbon source which is easily utilized by the bacteria, hence it can be efficiently used for improved production of cold active lipases.

The effect of supplementation of inorganic and organic nitrogen sources in SmSSF system is shown in Table 2. Ammonium nitrate and peptone (1% w/v) when used as additional nitrogen sources increased the production of cold active lipase from *M. roseus* upto 5.33 U/gds in each case. However, supplementation of potassium nitrate, beef extract, and yeast extract do not have any significant role in increasing the production of cold active lipase. Similar to above findings, few reports are available on peptone as the best carbon source [24] and increase in production of lipase when ammonium nitrate was supplied as inorganic nitrogen source to the organism [25].

![Figure 2: Cold active lipase production at different temperatures using groundnut oil cake as substrate.](image2)

![Figure 3: Cold active lipase production at different pH using groundnut oil cake as substrate.](image3)

**Table 2: Effect of triglycerides, carbon source, and nitrogen sources in cold active lipase production.**

| Additional sources | Lipase activity (U/gds) |
|-------------------|------------------------|
| Control           | 3.66                   |
| Castor oil        | 3.00                   |
| Triglycerides (0.5% v/v) |  |
| Soybean oil      | 1.33                   |
| Olive oil        | 1.33                   |
| Mustard oil      | 0.66                   |
| Carbon source (1% w/v) |  |
| Glucose          | 4.66                   |
| Sucrose          | 3.33                   |
| Lactose          | 2.00                   |
| Maltose          | 2.00                   |
| Nitrogen source (1% w/v) |  |
| NH₄NO₃           | 5.33                   |
| KNO₃             | 2.66                   |
| Peptone          | 5.33                   |
| Beef extract     | 3.33                   |
| Yeast extract    | 3.33                   |
yield was obtained in mixed substrate preparation from *Candida rugosa* [26]. The supplementation of GOC along with MOC (1:1), that is, media A with different nutrient sources individually such as triglycerides, glucose as carbon source, ammonium nitrate as nitrogen source as well as with the combination of glucose and ammonium nitrate, which were designated as media B, C, D, and E were tested and results obtained are given in Figure 5. The increased production of cold active lipase was obtained with glucose 5.00 U/gds, ammonium nitrate 5.33 U/gds, as well as with the combination of glucose and ammonium nitrate 6.66 U/gds. Production of cold active lipases from *M. roseus* was reported to show 150% increased production after optimization of mixed preparation. Similarly, a higher lipase production by *Aspergillus niger* on optimization of mixed substrate in ratio of 1:1 [27] was obtained in fungal fermentation process.

The optimum reaction temperature for lipase was 15°C when p-nitro phenyl palmitate (p-NPP) was used as substrate. Reaction mixture when incubated at 40°C showed less activity. *Micrococcus roseus* lipase was stable up to 30°C but rapidly inactivated at higher temperature above 40°C when incubated for 1 h (Figure 6). The activity of the cold enzyme presents an apparent optimal activity around 35°C and retains about 20% of its activity at 0°C, whereas the activity of the mesophilic lipases is close to zero at temperatures below 20°C and increases at temperatures above 60°C [28]. The increased catalytic activity at low temperatures and decreased thermostability of psychrophilic enzymes suggest that there is a relationship between stability and activity to maintain the activity at low temperature. This marked liability of *M. roseus* lipase together with its high catalytic efficiency near 15°C clearly denotes that it is cold active enzyme. Maximum activity and stability of cold active lipase was obtained at pH 8.0. The enzyme was stable under a pH range of 7.0–9.0 with slightly more in alkaline conditions (Figure 7). Lipases showing high stability and activity over a wide range of pH and activity under nonconventional conditions are of great interest. The major commercial application for alkaline stable lipases is the use in laundry and household detergents. *M. roseus* lipase was stable (>60%) over a pH range of 7.0–9.0; therefore, it has great potential for application in detergent industry as an additive used for cold washing.

4. Conclusion

Agricultural waste utilization for industrial development is one of the developing areas in modern industrial biotechnology. As shown in present research, the cold active bacterial lipases could be economically produced by *M. roseus* under semi-solid state fermentation using cheaply...
available groundnut oil cake as potent substrate. However, blending of groundnut oil cake along with mustard oil cake (1:1) under optimized condition showed better yield of cold active lipase. Groundnut oil cake and mustard oil cake along with glucose and ammonium nitrate was found to be a promising substrate for industrial production of cold active lipase in SmSSF by providing all the nutrients for the anchorage of cells. Further, it is reported that production of cold active lipase through SmSSF might not only reduce the cost of production but also results in manyfold increase in the lipase yield when compared with submerged fermentation. The study thus conducted has tried to expand the horizon of our knowledge on the cold active bacterial lipases which represents an extremely versatile group of bacterial extracellular enzymes that are capable of performing a variety of important reactions, thereby presenting a fascinating field for future research.

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