Isolation, Purification and Characterization of Secondary Structure and Kinetic Study of Lipase from Indian Major Carp, Catla catla (Catla)

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Abstract

Lipases are ubiquitous enzymes which catalyze the hydrolysis of fats into fatty acids and glycerol at the water-lipid interface and reversing the reaction in non-aqueous media. Lipases occupy a place of prominence among biocatalysts owing to their novel and multifocal applications in oleochemistry, organic synthesis, detergent formulation and nutrition.

Lipase was extracted and isolated from the alimentary canal and digestive gut of Catla catla (catla). The tissue was homogenized in a ratio of 1:3 with starting buffer (0.01 M TrisHCl, pH 7.2). The crude extract thus obtained was precipitated using ammonium sulphate (20-80%). Excess salt was removed by Dialysis and the resultant dialysate (Desalted Enzyme) was subjected to DEAE-Cellulose column for ion Exchange Chromatography at a flow rate of 0.5 ml/min. Elution was carried out by a step gradient of NaCl (100-800 mM) in the starting buffer. Active fractions were pooled as Purified Fraction (PF) and were used for physical characterization of pH, temperature and effect of calcium on the enzyme activity, structural characterization, molecular characterization and for kinetic studies. The Purified Fraction (PF) showed final specific activity of 1438.72 U/mg. The optimum pH was 7.8 and the optimum temperature was found to be 20˚C. Melting Temperature (T_m) value was 42˚C and the activation energy of purified lipase was 34.82 KJ/mol/K. Thermal stability of lipase was found to be at 20˚C. Lipase activity retained up to 3 h of incubation with 10 mM and 20 mM of CaCl₂ in starting buffer. This shows that Calcium has enhancing property from the denaturation of enzyme. Michaelis-Menten constant (K_m) of lipase from Indian major carp, catla for the hydrolysis of pNPP was 6.695 mM. Turnover number (k_cat) of lipase (catla) was 0.0022 s⁻¹. Catalytic efficiency (k_cat/K_m) of lipase was 0.0033412 s⁻¹ mM⁻¹. SDS-PAGE of purified lipase (PF) revealed a homogenous single band with molecular mass of 70 kDa. Secondary structural arrangement of α helices and β strands of purified lipase results 48.51% and 9.74%, respectively using Circular Dichroism.

Keywords: Lipase; Ammonium sulfate; Purification; Characterization; Kinetic study

Introduction

The use of enzymes created opportunities for developing a green, sustainable and modern industrial chemistry due to excellent specificity, being atom economic, mild reaction conditions, energy-saving process and simplicity. In particular the use of hydrolytic enzymes such as protease, lipase and amylase in the industrial enzyme market is growing steadily.

Lipases (E.C. 3.1.1.3) can be broadly defined as enzymes that catalyze the hydrolysis of ester bonds in substrates such as vitamin esters, phospholipids, triglycerides (TGs) and cholesteryl esters [1]. Interest in lipases has been increasing in the past few years because of the actual and potential uses of this enzyme in applications like hydrolysis of fats and oils, organic synthesis, modification fats, chemical analysis and flavor enhancement in food production. The isolation and purification of lipases is reported in various models, chiefly microorganisms but there is a paucity of information about the isolation and purification from carps. Therefore, the lipases from the tissues of these aquatic vertebrates have received much less attention than those from microorganisms. Hepatopancreas as digestive organs found in the fishes remain unexplored lipolytic enzyme sources that are of potential industrial applications [2]. Purification of lipase from the Indian Major Carps may also be of benefit for the development in the field of aquaculture. Thus the present study intricacies on the isolation purification and characterization of lipase from the digestive gut of Indian Major carp, Catla catla (catla).

Materials and Methods

Culture of fish

Indian major carp Catla catla (catla) were collected from local fish market, INA, Delhi and were acclimatized at outdoor conditions for 10 days. Fish were fed with artificial diet containing 40% protein. Artificial diet was prepared by using dried fish powder, wheat flour, cod liver oil and vitamin and mineral premixes [3]. Fishes were kept in fasting condition for 48 h before sampling.

Sampling of tissue

Two fishes were anesthetized with MS 222 (Tricaine methanesulfonate) before sacrifice. Individual fish was dissected; digestive system and associated glands (hepatopancreas) were removed from the body, cleaned, weighted and immediately frozen at -20°C till further use.

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Preparation of crude extract

Digestive systems and hepatopancreas collected from two fishes were pooled; total weight of tissue was recorded and homogenized in sample buffer (10 mM Tris-HCl and pH 7.2) in the ratio of 1:3. Homogenized solution was passed through pretreated cheese cloth for the separation of excess fats. The cheese cloth was treated by keeping it in 1% EDTA solution for 12 h. The suspension was then centrifuged for 30 min at 13000 rpm at 4°C. The floating fat phase was removed and the solution was filtered through Buchner funnel (Borosilicate grade-2) under vacuum (ROCKER-300, TARSONS, Mumbai, India), volume of the solution was noted and thus formed solution is called crude extract.

Estimation of protein

Protein concentration was measured by the method of Bradford [4]. Bovine serum albumin was used as a standard.

Assay of lipase activity

Lipase was assayed according to the method of Winkler and Stuckman [5]. The basis of this assay is the colorimetric estimation of para-Nitrophenol (pNP) released as a result of enzymatic hydrolysis of para-Nitrophenyl palmitate (pNPP). The substrate was prepared by dissolving pNPP (30 mg) in 10 ml of isopropanol mixed with 90 ml of 0.05 M Sorensen’s phosphate buffer (pH 8.0) containing 207 mg of sodium deoxycholate and 100 mg of gum arabic. Aliquot (1.2 ml) of the freshly prepared substrate was pre-warmed at 37°C and incubated with the enzyme extract of 50 µl for 15 min, at 37°C. Then the absorbance was recorded at 410 nm against enzyme free control. One enzyme unit is defined as 1 nmol of p-Nitrophenol enzymatically released from the substrate ml⁻¹ min⁻¹. The extinction coefficient of para-Nitrophenol is 15, 000 cm² mg⁻¹.

Purification of crude sample

Crude extract is subjected to ammonium sulfate fractionation [6]. Saturated ammonium sulfate (20%) was slowly added to the crude extracts with constant stirring for 2 h after final addition of ammonium sulfate. Then centrifuged at 13,000 rpm at 4°C for 30 min and the supernatant was brought to 80% saturation by further addition of ammonium sulfate. Then the sample was centrifuged at 13000 rpm at 4°C for 30 min and the precipitate was collected; re suspended in the sample buffer in 1:1 ratio. The solution was pipetted into the cuvette and the baseline was scanned from 200-260 nm using CD machine (Applied PhotoPhysics, Chirascan™ CD Spectrometer). Few lyophilized aliquots were used for molecular characterization.
Results and Discussions

Purification of enzyme

Lipase was extracted and concentrated by using ammonium sulfate precipitation in the cut off range 20-80%. Initial specific activity was found to be 406.1 units in the crude extract (Table 1). Resultant precipitate was subjected to dialysis to remove excess salts. Dialysate specific activity was found to be 408.3 with recovery of 42.2%. Dialysate was subjected to purification by using ion exchange chromatography on DEAE-Cellulose. Bound fraction was eluted by using step gradient of sodium chloride in the range of 100-800 mM. An active peak in fraction numbers 48-55 and minor peaks were found in the elution profile of DEAE-Cellulose column (Figure 1).

Active fractions were pooled as Purified Fraction (PF), the final specific activity of purified lipase was found to be 1438.72 units with recovery of 8.01% and purification fold was 3.54. The purified fraction was further processed for physical, kinetic and molecular characterization. Nayak et al. [13] used SEPHADEX G100 Chromatography to purify intestinal lipase from Labeorohita and obtained 3.6 fold purify with a specific activity of 38.4 U/mg. Gorgun and Akpinar [14] purified lipase from the liver of carp Cyprinus carpio by combined PEG-6000 on Q Sepharose on Sepharacryl S200-HR on Phenyl Sepharose CL-4B. 75.5 fold was purified and 90.38 U/mg specific activity was obtained. Purification fold of 34.5 was obtained with 6.21 U/mg specific activity when lipase was purified from the dorsal part of Cirrhinus reba using gel filtration on Sephadex G-50 (F-1) and DEAE Cellulose [15].

Physical characterization

Optimum pH: Effect of pH on lipase activity of the digestive tissue of Catla catla was measured in series of buffer range for pH 2.0 to 10.2 using pNPP as substrate at 37°C. Optimum pH in the study was found to be 7.8 (Figure 2). In the acidic pH at 2.5, the lipase activity was 10% and then it gradually increased up to 75% at pH 4.0. Thereafter activity was gradually decreased; activity was declined by 25% and was 50% at pH 7.0. In the pro alkaline condition, activity increased to maximum (100%) at pH 7.8 and thereafter it decreased finally to 30% at pH 10.0. Triglyceride lipase pH of Gadus morhua was 8.25 [16] using Olive oil as substrate and pH of purified lipase of Cyprinus carpio was 8.0 using the substrate pNPB [14]. These results were in a strong correlation with those of present study. However under acidic conditions, the charge distribution and conformation were changed and enzymes could not bind to substrate properly [17].

Optimum temperature: Effect of temperature on the hydrolysis of pNPP of lipase is shown in Figure 3. Optimum temperature was found to be at 20°C. Activity was found to be 60% at 10°C and then it increase by 40% with maximum activity (100%) at 20°C. Therefore for every

![Figure 1](image1.png)

Table 1: Summary of Catla catla lipase purification. The basis of this assay is the colorimetric estimation of para-Nitrophenol (pNP) released as a result of enzymatic hydrolysis of para-Nitrophenyl palmitate (pNPP). One enzyme unit is defined as 1 nmol of p-nitrophenol enzymatically released from the substrate ml⁻¹ min⁻¹. The extinction coefficient of para-nitrophenol is 15000 cm⁻² mg⁻¹.
rise in the temperature, lipase activity gradually decreased with final activity of 10% at 80°C. In the present study lipase showed optimum activity at 20°C.

Optimum temperature was found to be 15°C for *O. mykiss* liver lipase [18], 37°C in *Cyprinus carpio* [14], 25°C to 30°C for *Gadus morhua* digestive lipase [16] and 35°C for *C. reba* dorsal part lipase [15].

**Melting temperature:** Melting temperature of purified lipase of *Catla catla* was evaluated by using the relative activity of lipase at various temperatures by using pNPP as the substrate. Melting temperature (T_m) of catla lipase was 42°C (Figure 4) as exactly 50% of the activity lost at this particular temperature.

**Activation energy:** Arrhenius Activation energy of purified lipase from *Catla catla* was determined by using the I_r, relative activity. The activation energy was calculated as 34.82 KJ/mol/K (Figure 5). The activation energy of cyanobacterial lipase from *Spirulina platensis* was calculated by using Arrhenius equation and it was found as 146.34 kJ/mol [19].

**Temperature stability:** A 60.2% lipase activity was recorded at 30°C with incubation period of 30 min (Figure 6) and was stable after 60 min of incubation retaining 55% activity. Similar trend was observed at 40°C with incubation period of 30 min and thereafter it decreased to 10% after 60 minutes of incubation showing less stability. Half of the activity was found at 30°C after 30 minutes of incubation and then it slightly decreased to 25% final activity after 60 min of incubation.

Residual activity of lipase of *Catla catla* was stable at temperature 20°C up to 60 minutes of incubation whereas residual activity of lipase gradually declined at 30°C and 40°C. The enzyme activity was clarified to be stable within the temperature range of 30-60°C for *C. reba* dorsal part lipase [15].
Kinetic study

Kinetic constants $K_m$ and $K_{cat}$ for lipase from digestive gut or hepatopancreas of *Catla catla* were calculated using Michaelis-Menten and Lineweaver-Burk plot (Figures 7a and 7b) and were shown in Table 2. $K_m$ and $K_{cat}$ for the pure lipase of *Catla catla* were found to be 6.695 mM and 0.0022 s$^{-1}$, respectively. Catalytic efficiency of lipase was 0.0003412 s$^{-1}$ mM$^{-1}$.

Using different substrates, it appears that lipases purified from different tissues of fish have exhibited different $K_m$ and $V_{max}$ values. $K_m$ and $V_{max}$ values for *Cyprinus carpio* [14] liver lipase were 0.17 mM and 2.6 μmol/ml·dk using p-NPB as substrate respectively. Harmon et al. [20] found the $K_m$ and $V_{max}$ values for *O. mykiss* liver lipase. Another study focused on liver lipase of *O. mykiss* determined $K_m$ and $V_{max}$ values as 0.12 mM and 0.40 U/mg, respectively, using p-Nitrophenyl acetate (p-NPA) as substrate [21]. Aryee et al. [22] indicated that medium and long chain p-Nitrophenyl substrates were useful substrates for the lipase purified from the viscera of *M. cephalus*. Using p-Nitrophenyl palmitate (p-NPP) as a substrate, they calculated $K_m$ and $V_{max}$ as 0.22 mM and 20 μmin$^{-1}$ mg$^{-1}$.

The catalytic efficiency ($k_{cat}/K_m$) of lipase was found to be 1.5×10$^6$ M$^{-1}$ s$^{-1}$. These results showed that $K_m$ value of the lipase (p-NPP as substrate) from *S. platensis* was appreciably lower than $K_m$ values of lipase from other sources such as *A. niger* F044 (7.37 mM) [23], *Bacillus stearothermophilus* MC 7 (0.33 mM) [24] and *Bacillus sp.* J33 (2.5 mM) [25].

Effect of calcium

The Effect of Calcium chloride on the stability of lipase from *Catla catla* was examined at pH 7.2 and 37°C (Figure 8). Lipase activity increased by 8.2 times in 2 mM Calcium chloride till 2 hours and
thereafter it decreased at the end of 4 hours of incubation. Similarly, lipase activity increased by 6.2 times in 10 mM Calcium chloride till 2 hours and thereafter decreased to initial value at the end of 4 hours of incubation. Its maximum activity increased 8.5 times in 20 mM calcium chloride till 1 hour and then it gradually decreased to the initial value at the end of 4 hours of incubation. Incubation of purified trypsin with 2 mM CaCl₂ showed an increase in the activity up to 8 h. The enzyme activity was 11.7% higher after 4 and 8 h of incubation compared to the initial [26].

SDS-PAGE

The SDS-PAGE of purified lipase is shown in Figure 9. A commercial product of low molecular weight marker was (Bio rad) was loaded for comparison. In the crude extract there were three bands with molecular weight 89, 68 and 32 kDa. Two protein bands with molecular weight 30 and 67 kDa were observed in dialysate fraction. Protein bands were purified and reduced to a single homogenate band with 70 kDa molecular weight in ion exchange chromatography, DEAE-Cellulose. Other studies found the molecular weight as 74 kDa for liver lipase of C. carpio [14], 87 kDa for lipase from the dorsal part of Cirrhinus reba [15], 40-43 kDa for O. mykiss liver lipase [20], 43 kDa for S. aurita digestive lipase [27]. As can be seen from these results, we may conclude that lipases have different molecular weight depending on the tissue and the fish species under investigation.

Circular dichroism

The data was analyzed using K2D2 software. The CD spectra (Figure 10) obtained in the UV range 200-260 nm suggest that the secondary
The structural arrangement of lipase contains 48.51% of α helices, 9.74% β-sheets and 41.75% random conformations.

The CD spectra of secondary structure analysis of thermostable lipase from *Bacillus sp.* showed that the lipase structure contained 38.6% α-helices, 2.2% β-strands, 23.6% turns and 35.6% random conformations [28].

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Author’s Contribution

Experimental set up and data has been collected by Neelima Boora and Prasidhi Tyagi. Paper was compiled and written by Kameshwar Sharma.

References