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Purification and biochemical properties of lipase from raphia palm fruit mesocarp
T. Okunwaye, J.U. Obibuzor and E.A. Okogbenin
Purification and biochemical properties of lipase from raphia palm fruit mesocarp

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Lipases hydrolyze lipids to yield fatty acids and glycerol which are used as raw materials in industries. The lipase from raphia mesocarp was extracted, purified and characterized. Assay was carried out using p-nitrophenyl palmitate as substrate. The lipase was subjected to 80% ammonium sulphate precipitation, purified on ion exchange and gel filtration chromatography. Native and subunit molecular weights were determined on a gel filtration column and 10% sodium dodecyl sulphate polyacrylamide gel electrophoresis respectively. Kinetic parameters (K\text{m} and V\text{max}), effect of temperature, pH, sensitivity to metal ions and substrate specificities of the lipase enzyme were studied. Specific activity of 17.40 µmol/min/mg and purification fold of 2.68% were obtained. Native and subunit molecular weight was 35 kDa. The K\text{m} and V\text{max} values were 0.01 mM and 20.5 µmol/min/ml respectively. The lipase enzyme was inhibited by all metallic salts used and enhanced by CaCl\text{2}. Optimal pH was 7.0 and temperature for maximal activity was 40°C. The lipase hydrolyzed coconut oil at about twice the rate of palm kernel oil and palm oil, and at a higher rate than olive oil and Raphia oil. The study reveals that raphia mesocarp is a possible source of lipase which can be of industrial use.

Key words: Lipase, raphia palm fruit, mesocarp

INTRODUCTION

Lipases are a group of enzymes that catalyses the hydrolysis (cleavage) and synthesis of various forms of lipids (Svendsen, 2000). They belong to a subclass of the esterases and have been reported to be active at the oil water interface (Ibrahim, 1996). Oilseed lipases have great potential for commercial exploitation as industrial enzymes. The demand of lipolytic enzymes has been increased due to its potential use in the various manufacturing processes of industrial goods such as food and pharmaceuticals (Boland et al., 1991; Gandhi, 1997), in the production of biodiesel and industrial detergent (Freire and Castilho, 2008) as well as in the analysis of triacylglycerols (Forgia et al., 1995), raw materials in food, cosmetics and pharmaceutical industries (Hermansyah et al., 2006). Fat splitting, with lipase as a catalyst, is advantageous compared to conventional process due to low energy consumption, low cost, high product quality, ease of purification and safer process.
The yield of plant lipases is low; sometimes it could be cheaper than other sources and does not require fermentation.

Lipase activity has been reported in several oil plant extracts which include castor bean, palm seeds, oleifera seed, sunflower seed, peanuts (Abigor et al., 2002; Henderson and Osborne, 1991; Khan et al., 1991) and palm fruit mesocarp (Abigor et al., 1985).

Raphia is an underutilized and under exploited plant that produces fruits, with improved agronomical practices; tons of the fruits can be produced for cheap source of lipase which can be used industrially. In this study we report the isolation, purification and characterization of lipase from raphia palm mesocarp to hydrolyze vegetable oils for production of fatty acid and glycerol.

MATERIALS AND METHODS

Raphia palm fruits were obtained from Mile 18 village, Edo State, Nigeria. Palm oil, coconut oil, palm kernel oil used for the hydrolysis reaction were obtained from Mega stone farms, Oba town, Ogun State, Nigeria. Raphia oil was extracted by soxhlet using n-Hexane as solvent and olive oil was bought from a supermarket in Benin, Edo State, Nigeria. All chemicals, reagents, resins, standard protein and solvents used in the experiment were of analytical grades and were obtained from Sigma chemical Company, St Louis, Mo., USA, Pharmacia Fine Chemical, Uppsala, Sweden and BDH Chemicals Limited, Poole, England.

Enzyme preparations

The Raphia fruit was allowed to ripe, the exocarp was peeled off and the mesocarp was scraped with a cleaned knife and homogenized for 5 min in 0.15M Tris buffer adjusted to pH 7.5 with KOH. The homogenate was filtered through one layer of cheese cloth and centrifuged for 30 min at 10,000 g using High Speed Refrigerated Centrifuge. It yielded a fat layer, a supernatant liquid and a pellet. The supernatant was used for the assay of lipase activity and protein concentration determination. The remaining supernatant was fractionally precipitated using ammonium sulphate.

Assay of enzyme activity

Lipase activity was determined by the colorimetric method according to Lotrakul and Dharmsthiti (1997). This method was based on the cleavage of p-nitrophenyl palmitate (p-NPP) at pH 8.0. The reaction contained 180μl of solution A (0.062g of p-NPP in 10ml of 2-propanol, sonicated for 2 min before use), 1620μl of solution B (0.4% triton X-100 and 0.1% gum Arabic in 50 mM Tris-HCl, pH 8.0) and 200μl of properly diluted enzyme sample. The product was detected at 410 nm wavelength after incubation for 15 min at 37°C. Under this condition, one unit of lipase activity (U) was defined as the amount of enzyme required to release 1μmol of p-nitrophenol (p-NP) per minute under standard assay condition.

Protein determination

Protein concentrations were determined according to Lowry's method (1951) using BSA (1mg/ml) as the standard protein, absorbance was read at 660 nm and a standard graph was plotted. The amount of protein present in the sample was estimated from the standard graph.

Enzyme purification

The crude extract from the centrifugation step was fractionally precipitated to 80% ammonium sulphate saturation (516 g/L) according to the method of Doonan (1996). After standing for 1 h in an ice bath, the suspension was centrifuged at 10,000 g for 15 min at 4°C, the supernatant discarded, and the precipitate collected and was dialyzed against Tris buffer for 4 h to remove traces of ammonium sulphate.

Ion exchange chromatography was performed using DEAE Sephadex A-50 as the resin. The resin was packed into a 1.5 x 40 cm column. 5ml of the dialyzed extract was carefully layered on the column bed surface that has been previously equilibrated with about 3 L of Tris buffer, pH 7.5, containing 1mM EDTA and was allowed to drain into the bed. The column was washed with Tris buffer to remove unbound protein, followed by elution with stepwise gradient of 1M NaCl in Tris buffer. Elution was performed at a flow rate of 30ml/h and fraction of 5ml each was collected. Protein was monitored spectrophotometrically at 280nm. The fractions were also assayed for lipase activity. The fractions having lipase activity were pooled stored in the refrigerator for further purification.

The enzyme fraction was further purified on gel filtration chromatography using Sephadex G-200 as the resin. 5ml of the DEAE-Sephadex sample was layered on the column (1.5 x 40cm). Elution was performed at a flow rate of 10ml/h and fractions of 5ml each were collected. Protein was monitored spectrophotometrically at 280nm. The fractions were also assayed for lipase activity. The fractions having lipase activity were pooled.

Determination of native molecular weight

The native molecular weight of lipase was determined by gel filtration on a column of Sephadex G-100 (1.5 x 100 cm) using the following protein markers: Blue Dextran (2,000Da), Gamma Globulin (15,000Da), Creatine phosphokinase (81,000Da), Bovine serum albumin (68,000Da), Ovalbumin (45,000Da), Alpha Chymotrypsinogen (25,700Da). Void volume (Vv) was determined using blue dextran. Apparent molecular weight was determined by semi logarithmic plot of Ve/Vo against molecular weight of the marker proteins.

Determination of subunit molecular weight

The subunit molecular weight of purified lipase extract was analyzed with sodium dodecyl sulphate polyacrylamide gel electrophoreses (SDS-PAGE) according to the method of Laemmli (1970) in 12% slab gel. About 0.1ml of purified protein was loaded onto the gel. The gel was stained with 0.1% Coomassie brilliant blue R-250.

Determination of kinetic parameters

The Michaelis-Menten constant (Km) and the maximum velocity (Vmax) were determined by measuring enzyme activity of the purified lipase at various concentrations of emulsified substrates. The kinetic parameters (Km and Vmax) were calculated by plotting 1/V against 1/[S] in Lineweaver Burck plot (1934).

Effect of temperature on enzyme activity

A thermostated, water-jacketed reaction chamber was employed in determining the temperature dependence of lipase activity. The
optimum temperature was determined by measuring the lipase activity at different temperatures of 30, 40, 50, 60, and 70°C after 1 h incubation at pH 7.5 at standard conditions. The values of lipase activities were plotted against temperature.

Thermostability of lipase

Thermal stability was determined by incubating the enzyme at different temperatures ranging from 30 to 70°C for various length of time (30, 60, 90, 120, and 150 minutes) at pH 7.5. Lipase activity was measured and expressed as a percentage of the lipase activity.

Effect of pH on enzyme activity

The effect of pH on the activity of purified enzyme was studied by measuring the enzyme activity at various pH range of 3.5 to 10.0 for 1 h at room temperature. The lipase activity was assayed under standard conditions.

Enzyme inhibition

The following salts at the concentration of 5–20mM were tested for their possible inhibitory effects on the activity of the purified lipase enzyme: zinc chloride, manganese chloride, calcium chloride, potassium chloride and EDTA at room temperature for 1h and the lipase activities were determined.

Effects of substrates (hydrolysis reactions)

Palm oil, raphia oil, oliveoil, palm kernel oil and coconut oil were used as lipase substrates. The degree of lipolysis was determined by the assay method described previously above.

RESULTS

The elution profile of the crude extract from ion – exchange chromatography column of DEAE Sephadex A-50 is presented in Figure 1. The enzyme was not bound to the DEAE-Sephadex A-50 as it came out in the flow through fractions.No lipase activity was recovered in the fractions obtained by gradient elution. The crude protein was adsorbed on the DEAE- Sephadex A-50 column and this was eluted with a stepwise NaCl gradient(Figure 1), single peak was obtained. The active fractions from the peak was pooled together and used for further purification gelfiltrationchromatography. Single peak was also obtained, active fractions were pooled and used to assay the enzyme. A summary of the purification procedures is presented in Table 1. The partially purified enzyme from the mesocarp of raphia had a specific activity of 17.40 µmol/min/mg, a purification fold of 12.68 and a percentage yield of 5.79.

Five millilitre (5ml) of the dialyzed extract was carefully layered on the DEAE Sephadex A-50 column bed (1.5 x 40 cm) surface that has been previously equilibrated with about 3 L of Tris buffer, pH 7.5, containing 1mM EDTA and was allowed to drain into the bed. The column was washed with Tris buffer to remove unbound protein, followed by elution with 1M NaCl in Tris buffer. Elution was performed at a flow rate of 30ml/h and fraction of 5ml each was collected. Protein was monitored spectrophotometrically at 280 nm. The fractions were also assayed for lipase activity. The fractions 7 to 21 were pooled for further purification.

Five millilitre (5 ml) of the DEAE-Sephadex sample was
Table 1. Summary of purification of lipase from raphia palm mesocarp.

<table>
<thead>
<tr>
<th>Step</th>
<th>Volume (ml)</th>
<th>LPA (µmol/min/ml)</th>
<th>LPC (mg/ml)</th>
<th>TLA (µmol/min/ml)</th>
<th>TLP (mg/ml)</th>
<th>SP (µmol/min/mg)</th>
<th>Yield</th>
<th>Fold</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude extract</td>
<td>100.00</td>
<td>19.30</td>
<td>14.06</td>
<td>1930.00</td>
<td>1406.00</td>
<td>1.37</td>
<td>100.00</td>
<td>1.00</td>
</tr>
<tr>
<td>(NH₄)₂SO₄ ppt</td>
<td>8.00</td>
<td>23.05</td>
<td>16.17</td>
<td>184.40</td>
<td>129.36</td>
<td>1.42</td>
<td>9.55</td>
<td>1.03</td>
</tr>
<tr>
<td>Ion Exchange</td>
<td>5.70</td>
<td>31.15</td>
<td>3.18</td>
<td>177.55</td>
<td>18.12</td>
<td>9.79</td>
<td>9.19</td>
<td>7.13</td>
</tr>
<tr>
<td>Gel filtration</td>
<td>3.40</td>
<td>32.90</td>
<td>1.89</td>
<td>111.86</td>
<td>6.42</td>
<td>17.40</td>
<td>5.79</td>
<td>12.68</td>
</tr>
</tbody>
</table>

Vol = Volume (ml); LPA = lipase activity (µmol/min/ml); LPC = lipase protein concentration (mg/ml); TLA = total lipase activity (µmol/min/ml); TLP = total lipase protein (mg/ml); SPA = specific activity (µmol/min/mg). Yield = percentage fold = purification fold.

Figure 2. Elution profile of Lipase from raphia palm fruits, (gel filtration on Sephadex G-200).

Figure 3. Standard molecular weight curve.

ml each were collected. Protein was monitored spectrophotometrically at 280nm (Figure 2). The fractions were also assayed for lipase activity. The fractions having lipase activity were pooled and used for analysis.

Molecular weights determination

The calibration curve on Sephadex G-100 (1.5 x 100 cm) for the determination of native molecular weight is shown in Figure 3. The native molecular weight of lipase from raphia mesocarp was 35 kDa.

The picture and calibration curve for the SDS-polyacrylamide gel electrophoresis on 12% slab gels of lipase from Raphia mesocarp is shown in Figure 4. The molecular weight of lipase was estimated to be 35 kDa.

The Lineweaver Burk plot indicating the ($K_m$ and $V_{max}$) of lipase from raphia mesocarp is shown in Figure 5.
Figure 5. Lineweaver Burk plot of lipase activity (mesocarp of raphia palm.) \( K_m = 0.01 \text{ mM} \), \( V_{\text{max}} = 20.5 \text{ µmol/min/ml} \).

Figure 6. Effect of temperature on purified lipase from Raphia palm mesocarp.

The results on effect of temperature, thermostability, effect of pH on lipase from Raphia mesocarp are shown in Figures 6, 7 and 8 respectively. Figure 9 shows the hydrolytic activity of the lipase on some vegetable oils and the effect of salt of metal ions on lipase activity are seen in Figure 10.

**DISCUSSION**

The precipitation of the enzyme was carried out using ammonium sulphate since it was highly soluble in water, cheap and had no deleterious effect on structure of protein. The specific activity obtained from the purification of lipase from Raphia mesocarp after further purification using Sephadex G 200 and DEAE Sephadex A-50 resins was 17.40, yield of 5.79% and purification fold of 12.68. Lipase from *Propionibacterium acidpropionic* Sephadex G-50 has a 33-fold increase in specific activity (Sarada and Joseph, 1992).

The purified lipase was monomeric and had a subunit and native molecular weight of 35 kDa obtained on gel filtration which is similar to the molecular weight of *Aspergillus niger* lipase 35 kDa on Sephadex G-100 gel filtration chromatography and SDS-PAGE (Sugihara et al., 1988), and also monomeric lipase of *Pseudomonas fragi* and *Pseudomonas fluorescens* with molecular weights of 33 kDa.

In many cases, lipases appear to obey Michaelis—
Menten kinetics (Guit et al., 1991). Michaelis–Menten kinetics are characterized by two parameters, $K_m$ and $V_{max}$. The latter is the maximum rate of reaction and $K_m$ is a measure of the affinity of an enzyme for a particular substrate. The kinetic parameters ($K_m$ and $V_{max}$) of lipase obtained by Lineweaver Burk plot when $p$-nitrophenyl palmitate was used as substrate were 0.01mM and 20.5$\mu$mol/min/mg for the mesocarp of raphia. For a *Pseudomonas aeruginosa* lipase, Pancreac'h and Baratti (1996) reported $V_{max}$ value of 30 mmol/min when pNPP was used as substrate. The $K_m$ reported in this study is an indication that the enzyme from the germinated palm seeds has high affinity for the substrate and could catalyse the hydrolysis reaction for production of glycerol and fatty acid very fast.

One of the factors that affect the activity of enzymes is temperature and is often used to characterize enzyme. In this study, the optimum temperature of lipase from Raphia mesocarp was 40°C. Activity was rapidly lost at temperatures greater than 50°C. This is similar to the report of Abigoret al. (2003) on lipase from germinating seeds of *Jatropha curcas L.* Itikhar et al. (2010c) reported that the optimum temperature for lipase from *Rhizopus oligosporus* was 40°C. The relative activity of the partial purified cocoa lipase has an optimum temperature range of 30-40°C (Ratna, 2008).

The thermal stability experiment showed that lipase from this plant source was stable at 40°C for 60 min, retaining about 95% of its activity. The lipase enzyme was observed to be sensitive to thermal activation; this
could be attributed to thermal distortion of its structure.

According to Sharon et al. (1998), the changes in pH affect the protein structure and the enzyme activity. The maximum activity of lipase from raphia mesocarp was observed to be at pH 7 which dropped onwards, this is similar to results of lipase from germinating *Jatropha curcas* L. seed whose pH optima was near neutrality (Abigor et al., 2003), *Pentaclethra* lipase which was also near neutrality (Enujiugha et al., 2004) and partial purified cocoa lipase which had an optimum pH of 7.0 (Ratna, 2008).

Metal ions have been reported to stimulate lipase-catalyzed hydrolysis of oil by removing the fatty acids from the oil-water interface and allowing lipase to act freely on oil molecules (Ohnishi et al., 1994). From this study of the effects of metallic salts on lipase activity from raphia mesocarp, all the metal salts used inhibited the lipase activity at differing levels of inhibition. The highest level of inhibition was observed with Zn\(^{2+}\) while the least inhibition was observed with Mn\(^{2+}\). Other salts used in this experiment included the activity of the lipase enzyme. Monnet et al. (2012) reported that calcium ion was found to enhance lipase activity, while Mn\(^{2+}\), Na\(^{+}\), K\(^{+}\), Fe\(^{3+}\), Fe\(^{2+}\) and Zn\(^{2+}\) strongly inhibited the same. Sharon et al. (1998) reported a lipase of *P. aeruginosa* KKA-5 that retained its activity in presence of Ca\(^{2+}\) and Mg\(^{2+}\) but was slightly inhibited by Mn\(^{2+}\), Cd\(^{2+}\), and Cu\(^{2+}\). Rahman et al. (2005) stated that metal ions will bind to the enzyme and change the enzyme’s conformation to achieve better stability and hence greater activity. EDTA was also observed to inhibit the activity of the lipase and this is in agreement with the report of Monnet et al. (2012) that lipase from dormant ripe and unripe *Terminalia catappa* Linn. were strongly inhibited by EDTA, thus suggesting that the lipases are metalloenzymes. According to Enujiugha et al. (2004), the inhibition could be attributed to its chelating process of the system and thereby disrupting the formation of the enzyme substrate complex. This invariably affects the formation of the end product (Enujiugha et al., 2004).

The result of the lipolysis activity shows that the enzyme hydrolyses coconut oil at a higher rate than other oils used. Palm oil and palm kernel oil were hydrolysed at comparable rates; also olive oil and raphia oil were hydrolysed at comparable rates. The result shows that the lipase hydrolyses medium chain triacylglycerol at a faster rate than long chain triacylglycerol. This is in contrast to the findings of Abigor et al. (2003), which reported that lipase from germinating *J. curcas* L. seeds hydrolyzed long-chain TAG than medium chain triacylglycerol.

In conclusion, apart from the fact that this work established the presence of lipase from Raphia mesocarp, it also revealed that this enzyme from these sources had some physico-chemical properties that are similar to what have been obtained from other plant and microbial sources. Some of these properties could make the enzyme to find extensive applications not only in chemical, pharmaceutical, food, or leather industries but also in biodegradation of some materials.

Conflict of interest

Authors declare no conflict of interest.

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