Beauveria bassiana Lipase expressed in Komagataella (Pichia) pastoris with potential for biodiesel catalysis

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Lipases (EC 3.1.1.3) comprise a biotechnologically important group of enzymes because they are able to catalyze both hydrolysis and synthesis reactions, depending on the amount of water in the system. One of the most interesting applications of lipase is in the biofuel industry for biodiesel production by oil and ethanol (or methanol) transesterification. Entomopathogenic fungi, which are potential source of lipases, are still poorly explored in biotechnological processes. The present work reports the heterologous expression and biochemical characterization of a novel Beauveria bassiana lipase with potential for biodiesel production. The His-tagged B. bassiana lipase A (BbLA) was produced in Komagataella pastoris in buffered methanol medium (BMM) induced with 1% methanol at 30°C. Purified BbLA was activated with 0.05% Triton X-100 and presented optimum activity at pH 6.0 and 50°C. N-glycosylation of the recombinant BbLA accounts for 31.5% of its molecular weight. Circular dichroism and molecular modeling confirmed a structure composed of α-helix and β-sheet, similar to α/β hydrolases. Immobilized BbLA was able to promote transesterification reactions in fish oil, demonstrating potential for biodiesel production. BbLA was successfully produced in K. pastoris and shows potential use for biodiesel production by the ethanolysis reaction.

Keywords: lipase, Beauveria bassiana, heterologous expression, lipase purification, ethanolysis, biodiesel

Abbreviations: BbLA, Beauveria bassiana Lipase A expressed in Komagataella pastoris; BMM, buffered methanol medium; bp, basis pair; BMMY, buffered methanol medium plus yeast; C18-Sep, Octadecyl-Sepabeads; CALB, C. antarctica lipase B; CCRD, central composite rotational design; CTAB, cetyl trimethylammonium bromide; D/d, diameter of the halo/diameter of colony; DEPC, diethylpyrocarbonate; DHA, docosahexanoic acid; EPA, eicosapentaenoic acid; EstA, Aspergillus niger esterase A; Far-UV CD, far ultraviolet Circular Dichroism; GCLI, Geotrichum candidum lipase I; GCLI, Geotrichum candidum lipase II; GS115, strain of K. pastoris used in this work; IMAC-Cu2+Cl-, immobilized metal (Cu2+Cl-) affinity chromatography; kcat, Turnover number; Kd, Michaelis constant; MM, minimal medium; PDB, protein data base; pl, isoelectric point; PNP, p-nitrophenol; PNPB, p-nitrophenyl butirate; pPIC9k, CT, vector used in this work; RCLs, R. chenensis lipases; RML, R. miehei lipase; RT-PCR, real time PCR; SDS, sodium dodecyl sulfate; SDS-PAGE, sodium dodecyl sulfate–polyacrylamide gel electrophoresis; TLL, T. lanuginosa lipase; Vmax, maximum reaction rate; YNB, yeast nitrogen base; YPM, yeast peptone medium.
Background

Microbial lipases (EC 3.1.1.3) constitute a biotechnological important enzyme group because of their versatile properties and the relative ease of large-scale production compared to animal and vegetable homologues (Singh and Mukhopadhyay, 2012). Lipases belong to the carboxyl ester hydrolase family, which present activity against carboxylic ester bonds of triacylglycerols and act on the organic-aqueous interface releasing fatty acids and glycerol (Jæger and Reetz, 1998; Bancerz and Ginalska, 2007; Lotti and Alberghina, 2007). These enzymes are able to catalyze both hydrolysis and synthesis reactions of esterification, transesterification, and interesterification, depending on the amount of water in the system (Krishna and Karanth, 2001; Gotor, 2002; Hasan et al., 2009). The different catalytic properties are very important for lipase applications, especially in the food, detergents, fine chemical synthesis, pharmaceutical, cosmetics, and biodiesel industries (Wang et al., 2002; Hasan et al., 2006; Freire and Castilho, 2008). There is therefore a strong interest in the investigation of new sources of lipases or in the enhancement of the production of existing sources.

Research on lipase applications is currently focused on the area of biofuels, since production of biodiesel is a promising substitute for diesel oil. Biodiesel can be produced from renewable resources such as vegetable oils, animal oil, and residual oil from industrial and domestic kitchens. In order to this application, lipases in aqueous-restricted environments and the presence of ethanol or methanol results in a transesterification reaction called alcoholysis, giving rise to fatty acid esters (Iso et al., 2001; Matsumoto et al., 2001; Parawira, 2009).

The use of lipases in biodiesel production is advantageous as compared to alkaline chemical catalysis, because the enzymatic route does not result in saponification reactions and the glycerin and biodiesel recovery is simpler with higher purity. In addition, the catalyst recovery is easy when the lipase is immobilized on inert supports, the energy cost is lower since lipases generally operate under mild temperature conditions, and as a consequence the environmental impact is significantly reduced (Yang et al., 2009; Bisen et al., 2010; Cavalcanti-Oliveira et al., 2011; Batista et al., 2013). However, alcoholysis using lipase is more expensive because of the high cost of the biological catalyst and the lower overall productivity as compared to the alkaline catalysis (Wang et al., 1988; Batista et al., 2013). The current studies on enzyme use are focused on increasing the alkaline catalysis (Wang et al., 1988; Batista et al., 2013). However, alcoholysis using lipase is more expensive because of the high cost of the biological catalyst and the lower overall productivity as compared to the alkaline catalysis (Wang et al., 1988; Batista et al., 2013). The current studies on enzyme use are focused on increasing the alkaline catalysis (Wang et al., 1988; Batista et al., 2013). However, alcoholysis using lipase is more expensive because of the high cost of the biological catalyst and the lower overall productivity as compared to the alkaline catalysis (Wang et al., 1988; Batista et al., 2013). The current studies on enzyme use are focused on increasing the alkaline catalysis (Wang et al., 1988; Batista et al., 2013). However, alcoholysis using lipase is more expensive because of the high cost of the biological catalyst and the lower overall productivity as compared to the alkaline catalysis (Wang et al., 1988; Batista et al., 2013).

Materials and Methods

Fungi Strain, Culture Medium, Total RNA Extraction and cDNA Preparation

Beauveria bassiana CFF74 was grown in Khanna medium (Khanna et al., 1995) supplemented with 1% canola oil and 0.1% peptone at 30°C, 100 rpm for 60 h. Mycelia were harvested and frozen in liquid nitrogen. Total RNA was extracted by the TRizol® (Life technologiesTM) methodology. For this purpose, 1.0 mL of Trizol was used for each 50–100 mg of mycelium. Tubes were homogenized until mycelium was completely dissolved and then, incubated for 5 min at room temperature. Samples were quickly homogenized after the addition of 200 μL of chloroform per 1 mL of Trizol used. Subsequently, samples were incubated at room temperature for 10 min and centrifuged at 12,000 g, 4°C for 15 min. The aqueous phase was transferred to another clean tube. A volume of 500 μL of isopropanol was added to the samples. After 3 h at room temperature, samples were centrifuged again at 12,000 g, 15 min, 4°C and the pellet was washed with 75% ethanol. Samples were homogenized in vortex and centrifuged at 7,000 g, 5 min, 4°C. The supernatant was discarded. After drying for 30 min at room temperature, the pellets were resuspended in 100 mL DEPC-water and incubated for 10 min at 55°C. Total RNA was quantified in NanoDrop spectrophotometer (Thermo Scientific) and analyzed in 1.5% agarose gel, in 1x TAE buffer at 80 V for 2 h. After checking the quality of the samples in the gel, mRNA was used to synthesize the complementary DNA (cDNA) using OligodT20 and Invitrogen SuperScript® III Reverse Transcriptase Kit (Life TechnologiesTM) according to the manufacturer’s specifications.

Cloning and Expression on K. pastoris

Beauveria bassiana cDNA was used as template for PCR amplification of the gene encoding B. bassiana Lipase A (BblA) using primers — BblAE, (5’-NNNNNNCTCGAGAAAGAGGCTAACCTTCTGGC GGACATTGACGG-3) and BblAR, (5’-NNNNNNGACTTCAACATCCCATATCTTCAGCAAGAG CTT-3). The underlined sequences correspond to the XhoI and EcoRI sites, respectively. The full-length BblA coding sequence was PCR amplified by an important role during the infection of insects, hydrolyzing ester bonds in the lipoproteins, fats and waxes that are abundant in the insect exoskeleton (Ali et al., 2014). Hence, the study of entomopathogenic fungi as a source of lipases for application in catalysis may yield novel enzymes with improved features over existing industrial enzymes.

The genome of B. bassiana has been partially sequenced in mid-2012 (Xiao et al., 2012), and several genes encoding proteins with putative lipase function have been annotated, suggesting that this organism might be a promising source of this enzyme. This study reports the cloning and expression of one of these putative lipase genes in Komagatella pastoris and evaluation of the potential of the heterologous lipase for biodiesel production.
heating at 95°C for 10 min, followed by 35 cycles of 95°C for 15 s, 52°C for 40 s, 72°C for 3 min and final incubation at 72°C for 7 min. The PCR product was purified and digested with XhoI and EcoRI and cloned into the expression plasmid pPIC9K (Invitrogen, Nærum, Denmark) to produce the recombinant protein fused to the α-factor secretion signal and to the 6His extension at the N- and C-terminus, respectively (Fonseca-Maldonado et al., 2013). The resulting construct (pPIC9CT-BbLA) was transformed into electrocompetent DH5α Escherichia coli, and isolated plasmid DNA was purified and the nucleotide sequence determined. The BbLA nucleotide sequence was determined by sequencing the accession number XM_008603978.1. (Xiao et al., 2012), and was deposited in the GenBank database under accession number XM_008603978.1.

The pPIC9CT-Bbla (5 μg) was linearized with MssI (Pmel) (Thermo Scientific) and used to transform competent cells of the strain K. pastoris GS115 by electroporation at 1720 V, 5.5 ms (t), using an electroporator Eporator (Eppedorf®, USA). K. pastoris GS115/pPIC9CTbbla recombinants were grown on solid Minimal Dextrose (MD) plate medium for 3 days. Ninety-six colonies were transferred to Minimal Methanol Tributyrin plate medium (MMT – 1.34% Yeast Nitrogen Base without amino acids [YNB], 1% tributyrin and 1.5% agar sterile medium added with 1% methanol when the culture medium was at 40°C) and incubated for approximately 3 days at 30°C until halos of tributyrin hydrolysis appeared.

Three colonies that produced halo were inoculated into 100 mL buffered minimal glycerol medium (1.34% YNB; 4:10−5% biotin; 100 mM phosphate buffer, pH 6.0; 1% glycerol), and cultured in an orbital shaker (200 rpm) at 30°C. After 16 h, the cells were collected and resuspended in 100 mL buffered methanol-complex medium (BMM – 1.34% YNB; 4:10−5% biotin; 100 mM phosphate buffer, pH 6.0) in an Erlenmeyer flask of 1 L, with addition of 1% methanol when the culture medium was at 40°C and incubated for approximately 3 days at 30°C until halos of tributyrin hydrolysis appeared.

Protein Purification

Culture supernatants of BbLA-G2 transformant was concentrated and buffer exchanged using Hollow Fiber Cartridge, 50000 MMWC, in QuixStand Benchtop (GE Healthcare Bio-Sciences AB, Upplands, Sweden) concentration system. Concentrated BbLA was purified by two protocols: (I) BbLA, added 500 mM NaCl, was purified by IMAC-Cu2+ resin. Resin was previously equilibrated with 500 mM NaCl in 10 mM phosphate buffer, pH 7.0. Proteins were eluted with 250 mM Imidazol and 500 mM NaCl in 10 mM phosphate buffer, pH 7.0. (II) BbLA in 10 mM phosphate buffer was purified by hydrophobic chromatography on Octyl-Sepharose resin. BbLA was eluted with increased Triton X-100 concentrations up to 1%. In both protocols, fractions containing enzyme were dialyzed in 10 mM phosphate buffer. Protein purity was checked using 12% SDS–PAGE, and protein concentration was assayed according to Bradford (1976) using bovine serum albumin as the standard.

Enzymatic Assays and Enzymatic Characterization

The catalytic activity assay, modified from Moreno-Perez et al. (2014), used a spectrophotometer with a thermostated cell and continuous magnetic stirring (500 rpm) for 2 min. Reactions were initialized by mixing 0.1 mL of the soluble BbLA or its immobilized preparations (3 suspension) were added to 2.0 mL of the pNPB solution (2.0 mM p-nitrophenyl butyrate [pNPB]) in 0.05% Triton X-100 and 25 mM sodium phosphate at pH 7.0) at 40°C. The increase in absorbance at 380 nm (ε = 4.430 M−1cm−1) resulting from pNPB release was used to calculate enzymatic activity, given as μmol of pNPB hydrolyzed per minute per mL of enzyme (U/mL) under the described conditions. Kinetic parameters were determined for pNPB hydrolysis, in the best conditions for BbLA activity. Software GraphPad Prism 6.0 was used to calculate the Vmax and Km values.

The evaluation of the effect of surfactants on BbLA activity was made with 0.05% Triton X-100 was replaced by 0.05% Tween 20, Tween 80, Tergitol 40, CTAB or SDS. A control without surfactant was performed. Temperature and pH effects were evaluated by a 2° Rotational Central Composite Design. The pH (and encoded) values tested were 4.59 (−1.41), 5.0 (−1), 6.0 (0), 7.0 (+1) and 7.41 (+1.41), and temperature (and encoded) values were 35.9°C (−1.41), 40°C (−1), 50°C (0), 60°C (+1) and 64.1 °C (+1.41). Enzyme activity was evaluated discontinuously, where 50 μL of the reaction was added to 50 μL of supersaturated sodium tetraborate, after 1 and 2 min reaction. pNPB hydrolysis was measured at 410 nm (ε = 3.420 M−1cm−1) in a SpectraMax 96-well plate Spectrophotometer (Molecular Devices). Three central point repetitions were added to verify the analysis reproducibility. A confidence level of 95% was considered in the analysis of the variables effect. A polynomial equation was constructed for response surface analysis, and the Student’s t-test was used to verify the statistical significance of the regression coefficients. Statistical software Statistica v.12.0 was used to analyze the experimental data.

Deglycosylation of Recombinant Protein by Endoglycosidase H

Deglycosylation of purified recombinant BbLA was performed by Endo Hf (BioLabs, New England), with modifications to the manufacturer’s protocol. The deglycosylation reaction comprised 45 μL purified BbLA (80 μg protein) added to 5 μL 10X denaturation buffer, mixed and heated at 100°C for 10 min. Thereafter, 10 μL G5 reaction buffer, 0.5 μL of Endo Hf and 39.5 μL of MilliQ H2O were added. The reaction was incubated at 37°C for 24 h. Deglycosylation was evaluated using 12% SDS–PAGE.

CD Spectroscopy

Circular dichroism spectra between 190 and 250 nm (far-UV CD) were measured using a JASCO 810 spectropolarimeter (JASCO,
Tokyo, Japan) at a protein concentration of 0.04 mg/mL in 0.1 cm path-length quartz cuvettes in 10 mM phosphate buffer pH 7.0. The mean of four accumulated spectra was measured, from which the spectrum of a buffer blank was subtracted.

Three-Dimensional Model of the BbLA

The secondary sequence was inferred using the PDBSum platform. Homology modeling method (Deane and Blundell, 2003) was carried out using the I-Tasser (Zhang, 2008). Based on the X-ray structure of EstA (PDB ID: 1UKC), which was selected as template because an X-ray structure of the more homologue lipase from B. bassiana was not available. The primary sequence of lipases (BbLA and EstA) were aligned with highest scoring lipase (B. bassiana ARSEF 2860) obtained by BLAST using MUSCLE 3.6 (Edgar, 2004), to recognize the catalytic triad. The model quality structure was evaluated through PROCHECK program (Laskowski et al., 1993). The figures were generated using PyMOL.

BbLA Immobilization on Duolite A568 and Sepabeads-C18 Resins

The BbLA crude extract was dialyzed and equilibrated in 10 mM phosphate buffer, pH 7.0. A sample of 5 mg protein BbLA solution was added to 1 g of support Duolite A568 (derivative named BbLA-Duolite) or Sepabeads-C18 (derivative named as BbLA-C18). The immobilization mixture was maintained at pH 7.0, overnight, at 25°C. The activity of the immobilized enzyme was measured by the pNPB assay.

Enzymatic Synthesis of Ethyl Esters of Omega-3 Fatty Acids

Ethyl ester synthesis by BbLA was performed as described by Moreno-Perez et al. (2014). First, 0.26 g of dried immobilized lipase was added to the substrate solution containing 0.623 mmol of sardine oil and 6.2 mmol of ethanol dissolved in 4.11 mL cyclohexane. Second, 0.26 g of dry molecular sieves was added to the reaction mixture for the BbLA reaction in the absence of water. The final concentration of sardine oil in the solution was 125 mM, and the reaction was carried out at 25°C. Aliquots of 100 µL were withdrawn every 24 h, and diluted in 300 µL of HPLC mobile phase. BbLA-derivatives used in this experiment were dried using acetone solutions in water. The final concentration of acetonitrile in the solution was 125 mM, and the reaction was carried out at 25°C. Aliquots of 100 µL were withdrawn every 24 h, and diluted in 300 µL of HPLC mobile phase. BbLA-derivatives used in this experiment were dried using acetone solutions in water with increasing concentrations of acetonitrile up to 100%. Product formation was monitored by HPLC, where the control consisted of the same quantities of reagents, but without the presence of catalyst.

HPLC Analysis

Reactants and products were analyzed using a reverse-phase column (Ultrabase-C8, 150 mm × 4.6 mm, 5 µm) by RP-HPLC (Spectra Physic SP 100 coupled with a UV detector Spectra Physic SP 8450). Products were eluted at a flow rate of 1.5 mL/min with acetonitrile/water/CH3COOH (80:20:0.1 v:v), pH 3. The UV detection was performed at 215 nm. Product yields were calculated from the pure peak areas of known quantities of EPA ethyl ester [retention time (RT) of 9 min] and docosahexaenoic acid (DHA) ethyl ester (RT of 12 min).

Reproducibility of the Results

All data are the mean of at least three independent experiments.

Results and Discussion

Cloning and Expression of Beauveria bassiana Lipase A in Komagataella pastoris

A putative triacylglycerol lipase gene (GenBank access number EJP62207.1) was chosen from several genes described as lipase or lipase-like in the B. bassiana genome. The gene, named BbLA (Beauveria bassiana lipase A) is 1,641 bp long and encodes a protein with 546 amino acid residues, including a 19-amino acid signal peptide. BbLA has a theoretical molecular weight and pi of 59.58 kDa and 5.22, respectively. The BbLA cDNA without the signal peptide sequence was obtained by RT-PCR from B. bassiana cDNA with primers BblAF and BblAR. BbLA was cloned into the expression vector pPIC9k_CT in-frame with the yeast α-factor secretion signal. The resulting plasmid was sequenced and named pPIC9CT-BbLA. BbLA is translated into a 554 amino acids protein (including a His tag at the C-terminus), which has the catalytic triad and the common consensus sequence (G-X-S-X-G) for lipases (Figure 1).

Linearized pPIC9CT-BbLA was used to transform K. pastoris GS115. Ninety-six transformants were screened for the presence of hydrolysis halos in MM solid medium containing methanol and tributyrin (Figure 2). Three colonies that showed a higher D/d ratio (D/d) were selected. These clones were called BbLA F5, BbLA G2 and BbLA H1.

After 120 h induction with 1% methanol in liquid medium, the culture supernatants from the three clones showed similar lipase activity (Figure 3A). Transformant BbLA G2 was chosen for further studies because it showed the highest ratio D/d (Figure 2). BbLA showed better lipase expression in BMM medium after 120 h of methanol induction, demonstrating that buffering the culture medium and long induction times favor protein expression (Figures 3B,C).

Purification and Characterization of BbLA

Beauveria bassiana lipase A was purified by two protocols. The first protocol employed IMAC-Cu2+, which exploited the addition of a His-tag at the N-terminal of BbLA. The second protocol involved hydrophobic interaction chromatography using Octyl-sepharose at low ionic strength, which exploited the fact that BbLA is a lipase and therefore has a strongly hydrophobic substrate binding region. Both these protocols resulted in purified BbLA (Figure 4). IMAC-Cu2+ showed a slightly higher purification factor, while the Octyl-sepharose chromatography showed higher recovery (Table 1). Therefore,
Beauveria bassiana lipase A was activated by Triton X-100, Tween 20, Tween 80, Tergitol and CTAB, but was completely inactivated by SDS. The best condition for BbLA activation was with 0.05% Triton X-100 (Supplementary Figure S1).

A CCRD was used to analyze the optimal temperature and pH for BbLA reaction. For BbLA activity, the two variables showed a linear effect and temperature showed a quadratic effect (Figure 5A). A reduced second order model for BbLA activity is described by equation 1.

\[
\text{Activity (U/mL)} = 15.96 - 2.43(T) - 5.09(T)^2 - 6.56(pH)^2
\]

where, T and pH are encoded temperature and pH values, respectively.

Table 2 shows ANOVA results from reduced models generated. The determination coefficient was 0.9112. \( F_{\text{calculated}} \) value was higher than \( F_{\text{listed}} \) (Table 2). Therefore, this model was considered predictive and statistically significant at 95% confidence level. This result was analyzed by response surface methodology, and generated a 3D surface graph (Figure 5B). These results led to the estimation of ideal temperature and pH reaction conditions for BbLA, which were 50°C and pH 6.0, corresponding to the central points assayed.

The \( K_M, V_{max}, k_{cat} \) and \( K_{cat}/K_M \) of BbLA recombinant against \( p\text{NPB} \), pH 6.0 at 50°C, were 0.5546 μM,
Recombinant BbLA expression in *K. pastoris*. (A) Methanol induction of BbLA colonies F5, G2, and H1; (B) BbLA expression in different culture medium of the BbLA-G2 colony; (C) SDS–PAGE electrophoresis of BbLA expressed by BbLA-G2 colony over increasing time intervals on methanol induction. The control was *K. pastoris* transformed with empty vector. M, molecular weight marker; C, control. Cultures were incubated at 30°C, with 200 rpm. The arrow indicates the purified lipase.

Sodium dodecyl sulfate–polyacrylamide gel electrophoresis electrophoresis of BbLA purification. (A) Purification in IMAC – Cu²⁺. (B) Purification on Octyl-sepharose. M, molecular weight marker; CD, crude extract of BbLA; S, IMAC supernatant; C, concentrated BbLA. The arrows indicate the band corresponding to BbLA. See Methods section for further experimental details.

85.67 μmols.min⁻¹.mg⁻¹, 7.139 × 10⁴.s⁻¹, and 1.29 × 10¹¹.M⁻¹.s⁻¹, respectively (Table 3). Hill coefficient was higher than 1.0, which suggested that BbLA can be an allosteric enzyme. Allosteric lipase modulation is not common, but it is described as possible and biologically relevant (Kohler and Wunsch, 2007).

**BbLA Deglycosylation**

*Beauveria bassiana* lipase A presents no potential site for O-glycosylation, but includes six potential N-glycosylation sites. Purified BbLA showed a diffuse band in SDS–PAGE centered around 78 kDa, which after treatment with EndoHf, an endoglucanase that cleaves N-glycosylated oligosaccharides,
shifts to 60 kDa (Figure 6). Assuming that these forms represent the glycosylated and deglycosylated forms, respectively, then the saccharide content of the glycosylated BbLA is approximately 30%.

Molecular Modeling

The far-UV circular dichroism spectrum of purified BbLA at pH 7.0 presents a maximum at 194 nm and a minimum at 207 and 220 nm, which are spectral features typical of proteins rich in α-helix (Figure 7A). Nevertheless, the spectral feature generated is not characteristic of proteins exclusively formed by α-helices.

Aspergillus niger esterase (EstA, PDB code Q6ED33) (Bourne et al., 2004), shares 55.56% identity with BbLA and was used as a template for structural modeling of the BbLA on the I-TASSER platform. The secondary structure topology is showed in Figure 7B for the sequence length of 554 amino acids, where 61 residues are found in beta strands (forming β-sheets), 153 in α-helices, 14 in 3-10 helix and 326 in other secondary structure motifs such as beta hairpins, beta bulges, beta turns, and gamma turns. The model of the enzyme is a monomer presenting a α/β fold comprised of four central stranded β sheet flanked by 22 α-helices. The superposition between BbLA and EstA structures (Figure 8A) generates an RMSD score of 0.37. The catalytic triad is formed by Ser-Glu-His (Figure 8B).

**BbLA Immobilization and Application of BbLA-Derivatives in Fish Oil Ethanolysis**

*Beauveria bassiana* lipase A was immobilized on Duolite and C18-Sep by reversible anionic and hydrophobic interaction, respectively. Derivatives generated from this immobilization were named BbLA-Duolite and BbLA-C18, and had 8.65 and 12.35 U/g of support, respectively. Duolite and C18-Sep immobilized 83 and 98% of the BbLA present in the crude extract. BbLA-Duolite and BbLA-C18 were dried and added to a reaction medium free of water, in order to favor the transesterification reaction.

Eicosapentaenoic and DHA acids are abundant in sardine oil, and fatty acid ester formation after 24 h reaction revealed BbLA-C18 produced 7.35-fold more EPA-ester and 13.61-fold more DHA-ester than produced by BbLA-Duolite after 48 h reaction. However, BbLA-Duolite was more selective than BbLA-C18 (Table 4).

![Figure 5](image-url) | Figure 5: Experimental design for assessing pH and temperature effects on BbLA activity. (A) Pareto plot for CCDR. (B) Surface response for pH and temperature variables. The analysis was performed by Statistica 12.0 software.

![Figure 6](image-url) | Figure 6: Western blotting of purified BbLA showing a shift to 60 kDa. Assuming that these forms represent the glycosylated and deglycosylated forms, respectively, then the saccharide content of the glycosylated BbLA is approximately 30%.

![Figure 7A](image-url) | Figure 7A: Spectral features typical of proteins rich in α-helix. Nevertheless, the spectral feature generated is not characteristic of proteins exclusively formed by α-helices.

![Figure 7B](image-url) | Figure 7B: Secondary structure topology for the sequence length of 554 amino acids, where 61 residues are found in beta strands (forming β-sheets), 153 in α-helices, 14 in 3-10 helix and 326 in other secondary structure motifs such as beta hairpins, beta bulges, beta turns, and gamma turns.

![Figure 8A](image-url) | Figure 8A: The superposition between BbLA and EstA structures generates an RMSD score of 0.37. The catalytic triad is formed by Ser-Glu-His.

![Figure 8B](image-url) | Figure 8B: The superposition between BbLA and EstA structures generates an RMSD score of 0.37. The catalytic triad is formed by Ser-Glu-His.
Discussion

BbLA is a B. bassiana putative triacylglycerol lipase gene. This gene encodes a protein with 546 amino acid residues, 59.58 kDa and pl 5.22. BbLA protein has the catalytic triad and the common consensus sequence (G-X-S-X-G) for lipases. This sequence was the only consensus sequence identified in studies on non-redundant sequences of lipases deposited in the public data banks. The serine residue present in this pentapeptide is the nucleophilic amino acid present in the catalytic triad (Jaeger et al., 1994, 1999; Schrag and Cygler, 1997; Lotti and Alberghina, 2007).

BbLA F5, BbLA G2 and BbLA H1 clones presented similar lipase activity. BbLA G2 showed better lipase expression in BMM medium after 120 h of methanol induction. In comparison, Thermomyces lanuginosus lipase shows improved expression when grown in BMMY medium (similar to BMM, but including yeast extract) pH 7.0, with 1.2% methanol added every 24 h, and 144 h induction (Fang et al., 2014). These results corroborate those obtained in this work and demonstrate the importance of controlling the pH and induction over relatively long periods.

Beauveria bassiana lipase A was purified by affinity chromatography (IMAC-Cu²⁺) and hydrophobic interaction chromatography (Octyl-sepharose). The BbLA recovery after IMAC purification (39.13%) was superior to that observed for GCLI (13%) and GCLII (14%) for IMAC purification of Geotrichum candidum lipase expressed in Saccharomyces cerevisiae (Bertolini et al., 1995). However, it was lower than the recovery obtained for Bacillus licheniformis lipase (64%) also fused to a His-tag and expressed in Escherichia coli (Madan and Mishra, 2009). BbLA purified with Octyl-Sepharose had 75.58% recovery. Under low ionic strength conditions, only proteins with large hydrophobic sites, such as lipases, adsorb to hydrophobic resins (Palomo et al., 2002, 2006; Mateo et al., 2007). Indeed, hydrophobic interaction chromatography is often used as one of the steps for lipase purification, such as lipases produced by Pseudomonas aeruginosa (Bertolini et al., 1995) and Staphylococcus warneri (Volpato et al., 2011).

Surfactants activation is common to lipases. Triton X-100, Tween 20, Tween 80, Tergitol and CTAB activated BbLA. While, SDS inactivated it. These results are similar to those previously reported in the literature, such as those obtained for Thermosyntropha lipolytica lipases A and B, which are activated by sodium cholate, CTAB, Triton X-100, Tween 20, Tween 80 and SDS (Salameh and Wiegel, 2010). The lipase from Thermomyces lanuginosus was also activated by all the surfactants used, including SDS (Mogensen et al., 2005). On the other hand, Penicillium chrysogenum lipase was inhibited by almost all detergents used, including SDS, Triton X-100 and Tween 20, the only exception being cholic acid (Bancerz et al., 2005).

Central composite rotational design and response surface showed that the best temperature and pH reaction were 50°C and pH 6.0, respectively. Similarly to the results presented in this work, the optimum conditions obtained for Penicillium verrucosum lipase activity (Kempka et al., 2008) were 44°C and pH 7.0 and 40°C and pH 5.0 for Penicillium sp. lipase (Wolski et al., 2009).

Beauveria bassiana lipase A expressed in K. pastoris is approximately 30% N-glycosylated. The glycosylation of BbLA was higher than that observed for the Y. lipolytica Lip2, expressed in K. pastoris, which has 12% of its molecular weight corresponding to glycosylation (Wang et al., 2012). However, not all proteins expressed by K. pastoris are glycosylated. For example, two RCLs expressed in this system showed no evidence for glycosylation after treatment with EndoH (Yu et al., 2009).

Beauveria bassiana lipase A far-UV circular dichroism spectrum presented features typical of proteins rich in α-helix. But BbLA is not exclusively formed by α-helix, because the secondary structure topology showed that 61 amino acids residues would be in the β-sheet configuration. A similar structure is observed for other lipase enzymes from mammalian and bacterial origin, where the number of α-helices and β-sheet differ from one species to another (Messaoudi et al., 2011).

Aspergillus niger esterase A (Bourne et al., 2004) was used as a template for structural modeling of the BbLA. Esterases (EC 3.1.1.1) and lipases (EC 3.1.1.3) can be differentiated on the basis of their substrate specificity and by comparison of sequence-related structural features. In contrast to esterases, lipases display a significant difference in the distribution of
hydrophobic amino acid residues in the vicinity of their active sites (Chahinian and Sarda, 2009). The catalytic triad formed by Ser-Glu-His is the same as that in the *Geotrichum candidum* lipase, where glutamic acid replacing the usual aspartate (Schrag et al., 1991).

The three-dimensional model of the BbLA has proportionally more α-helix than β-sheet, but the two secondary structures are present. BbLA circular dichroism and molecular model are consistent with structures of lipases and esterases, because showed similar structure to α/β hydrolase. Despite the validation of structure through Ramachandran plot have presented less than 90% of residues in most favored regions. The predicted model is consistent with other lipases structures from eukaryotic and prokaryotic organisms.
Beauveria bassiana lipase A molecular modeling. (A) Superposition of structures from BbLA (blue) and EstA (green). The figure was made by I-TASSER (Zhang, 2008) using a TM-align structural alignment program. (B) 3D model of BbLA showing the catalytic triad (Ser, Glu, and His). The ribbon is colored with a rainbow spectrum from N-terminus (blue) to C-terminus (red).

Table 3 | Beauveria bassiana lipase A kinetic parameters obtained using pNPB as substrate in 25 mM phosphate buffer, pH 6.0, after incubation at 50°C, for 2 min.

|     | \( V_{\text{max}} \) (\( \mu \text{mol}.\text{min}^{-1}.\text{mg}^{-1} \)) | \( K_M \) (\( \mu \text{M} \)) | \( K_{\text{cat}} \) (s\(^{-1} \)) | \( K_{\text{cat}}/K_M \) (M\(^{-1}\).s\(^{-1} \)) | \( n \) |
|-----|-------------------------------------------------|-----------------|-----------------|-----------------|-----|
|     | 85.67                                           | 0.5546          | 7.139 \( \times \) 10\(^4 \) | 1.29 \( \times \) 10\(^{11} \) | 1.46 |

GraphPad Prism 6.0 was used to calculate \( V_{\text{max}} \), \( K_M \) and \( n \). \( n \), Hill coefficient.

Table 4 | Sardine oil Ethanolysis catalyzed by Duolite-BbLA and BbLA-C18 derivatives in the presence of cyclohexane.

| Derivative        | Selectivity (EPA/DHA) | Ethanolysis (%) |
|-------------------|-----------------------|-----------------|
|                   |                      | EPA            | DHA            |
| BbLA-Duolite\(^a\) | 5.57                 | 0.66           | 0.18           |
| BbLA-C18\(^b\)    | 2.98                 | 4.85           | 2.45           |

The reaction occurred at 25°C, 100 rpm. \(^a\)48 h of reaction. \(^b\)24 h of reaction.

Selectivity were 29 and 12, respectively (Moreno-Perez et al., 2014). It is noteworthy that the TLL also showed differences in selectivity when immobilized in the same supports used for BbLA.

Biodiesel production by BbLA-C18 may be improved through the study of several factors, such as increasing the amount of enzyme on the support, changing the organic solvent used, altering the temperature and the efficiency of transesterification on other oils. The present results demonstrate that the search for new enzyme sources together with the modulation of the activity by enzyme immobilization are valuable strategies for the development of biotechnological routes for biofuels production.

**Conclusion**

A B. bassiana Lipase A was expressed as a heterologous protein in K. pastoris, and the biochemical characteristics of the BbLA show potential for biodiesel production. The differential immobilization, by anionic and hydrophobic interaction, can generate products with different proportions of EPA and DHA esters, at mild temperatures, reducing the energy costs for biodiesel production.

**Author Contributions**

This work is part of the doctorate thesis of ACV and she is the main author. AC, RF-M, RW, and FT had collaboration in part of the experiments with molecular biology and fungal expression (enzyme assays optimum temperatures and heterologous protein expression). RW performed the circular dichroism. CC performed the molecular modeling BbLA and its analysis. BP, GF-L, FF and MGP had collaboration in the immobilization and application of enzyme. JJ designed some experiments; MLTMP contributed with the experimental design and the final manuscript.
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Supplementary Material

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2015.01083
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