



**UNIVERSITI PUTRA MALAYSIA**

**STRUCTURAL STUDIES OF A CHEMICALLY MODIFIED  
THERMOSTABLE LIPASE FROM *Geobacillus* SP. STRAIN T1**

**CHEONG KOK WHYE**

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**STRUCTURAL STUDIES OF A CHEMICALLY MODIFIED  
THERMOSTABLE LIPASE FROM *Geobacillus* SP. STRAIN T1**

**By**

**CHEONG KOK WHYE**

**Thesis Submitted to the School of Graduate Studies, Universiti Putra  
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**STRUCTURAL STUDIES OF A CHEMICALLY MODIFIED  
THERMOSTABLE LIPASE FROM *Geobacillus* SP. STRAIN T1**

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**September 2007**

**Chair : Professor Abu Bakar Salleh, PhD**

**Faculty: Biotechnology and Biomolecular Sciences**

Alkylation has been successfully performed using propionaldehyde on four batches of T1 thermostable lipase (M1, M2, M3 and M4) with different degrees of modification (27% to 55%) to represent the different levels of hydrophobicity. Based on the crystal structure, T1 possessed 11 lysine residues, of which four of the lysine residues, Lys84, 102, 138 and 251 have scores between 53.7% and 95.8% exposure ratio, were totally exposed. Another four residues, Lys185, 329, 344 and 345 have a ratio between 20% and 50% (moderately exposed) and three of the lysine residues, Lys28, 207 and 229 are buried. The hydrolytic activity of the modified enzymes dropped drastically by 10 to 40-fold upon chemical modification, despite both the native and modified form showed distinctive  $\alpha$ -helical bands at 208 and 222 nm by Far Ultra-Violet Circular Dichroism (CD) spectropolarimetry. As cooperative unfolding transitions were observed, the modified lipases were distinguished from the native state, which the former



possessed a  $T_m$  in lower temperature range, 60-64 °C whilst the latter at 68 °C. Consequently, this has led us to the hypothesis of formation of a molten globule (MG)-like structure.

Subsequent analysis of both native and modified lipases by Matrix Assisted Laser Desorption/Ionization Time of Flight Mass Spectrometry (MALDI-TOF MS) study was carried out to ascertain the modifications and the location of these modifications. Four lysine residues, Lys28, 84, 207 and 329 were clearly identified from the native spectrum. As expected, Lys84 and Lys102 were clearly modified. Surprisingly, Lys185 which has a very low exposure ratio (27.5%) was also identified to be one of the modified residues. To further support the hypothesis of the formation of a molten globule, intrinsic and extrinsic fluorescence were performed. A decrease of fluorescence intensity was observed for modified lipase M1, which was modified using 0.5% of propionaldehyde. However, subsequent addition of propionaldehyde enhanced the fluorescence intensity of M2, M3 and M4, which indicated an inversion of placement for tryptophans to a more hydrophobic environment. As for extrinsic fluorescence, the alkylated lipases showed a clear enhancement of fluorescence intensity as compared to the native lipase due to the exposure of the hydrophobic interior of the enzymes.

Abstrak tesis yang dikemukakan kepada Senat Universiti Putra Malaysia sebagai memenuhi keperluan untuk ijazah Master Sains

**KAJIAN STRUKTUR THERMOSTABIL LIPASE DARI *Geobacillus* SP. JENIS T1 YANG DIMODIFIKASI MELALUI TINDAK BALAS KIMIA**

Oleh

**CHEONG KOK WHYE**

**September 2007**

**Pengerusi: Profesor Abu Bakar Salleh, PhD**

**Fakulti : Bioteknologi dan Sains Biomolekul**

Pengalkilan telah berjaya dilaksanakan ke atas empat kelompok enzim thermostabil lipase T1 (M1, M2, M3 dan M4) dengan menggunakan propil aldehid pada beberapa peringkat modifikasi yang berbeza (27% sehingga 55%) untuk menggambarkan perbezaan hidrofobia. Berdasarkan struktur kristal, T1 mempunyai 11 lysine, yang mana empat lysine iaitu Lys84, 102, 138 dan 251 mempunyai skor di antara 53.7% dan 95.8% nisbah pendedahan, adalah dalam kategori pendedahan lengkap. Empat lysine yang lain, Lys185, 329, 344 dan 345 mempunyai skor di antara 20% dan 50% (pendedahan separa) dan tiga lysine yang terakhir, Lys28, 207 dan 229 adalah tertimbus. Sungguhpun begitu, selepas menjalani modifikasi secara kimia, aktiviti hidrolisis lipase yang telah diubahsuai didapati turun secara mendadak (10 hingga 40-kali ganda), walaupun spektrum spektropolarimetri 'Circular Dichroism' (CD) ultraungu jauh bagi kedua-dua lipase asal dan yang terubah menunjukkan struktur  $\alpha$ -helix pada 208 dan 222 nm.

Disebabkan kedua-dua lipase asal dan yang terubah menunjukkan graf transisi pembukaan lipatan yang lancar, lipase yang terubah suai hanya dapat dibezakan dengan lipase asal kerana lipase terubah suai mempunyai  $T_m$  pada julat suhu yang lebih rendah, 60-64 °C manakala lipase asal pada 68 °C. Susulan bukti ini, ia membawa kami kepada hipotesis pembentukan struktur seakan 'molten globule' (MG).

Analisis seterusnya, Spektrometri Jisim Matrik Laser Nyahserapan/Pengionan Penerbangan Masa (MALDI-TOF MS) dilakukan bagi memastikan proses modifikasi telah berlaku dan menjejaki lokasi-lokasi modifikasi tersebut. Empat lysine, Lys28, 84, 207 dan 329 dapat dikesan dalam spektrum lipase asal. Seperti yang telah diduga, Lys84 dan Lys102 telah dimodifikasi. Walau bagaimanapun, Lys185 yang mempunyai pendedahan yang sangat rendah (27.5%) juga telah dimodifikasi. Bagi menyokong hipotesis pembentukan struktur seakan MG, pendarfluor dalaman dan luaran dijalankan. Penurunan keamatan pendarfluor dapat diperhatikan bagi lipase terubah suai M1 yang diubahsuai menggunakan 0.5% propil aldehid. Walau bagaimanapun, apabila lebih banyak propil aldehid ditambah, keamatan pendarfluor lipase terubah suai M2, M3 dan M4 didapati meningkat, menunjukkan asid amino tryptofan mengubah kedudukan kepada persekitaran yang lebih hidrofobia. Bagi pendarfluor luaran pula, keamatan pendarfluor bagi lipase terubah suai adalah lebih tinggi berbanding lipase yang asal kerana bahagian dalaman enzim yang hidrofobia didedahkan.

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I certify that an Examination Committee has met on 11<sup>th</sup> September 2007 to conduct the final examination of Cheong Kok Whye on his Master of Science thesis entitled “Structural Studies of a Chemically Modified Thermostable Lipase from *Geobacillus* Sp. Strain T1” in accordance with Universiti Pertanian Malaysia (Higher Degree) Act 1980 and Universiti Pertanian Malaysia (Higher Degree) Regulations 1981. The Committee recommends that the student be awarded the Master of Science.

Members of the Examination Committee were as follows:

**Mohd. Arif Syed, PhD**

Professor  
Faculty of Biotechnology and Biomolecular Sciences  
Universiti Putra Malaysia  
(Chairman)

**Nor Aripin Shamaan, PhD**

Professor  
Faculty of Biotechnology and Biomolecular Sciences  
Universiti Putra Malaysia  
(Member)

**Irmawati Ramli, PhD**

Associate Professor  
Faculty of Science  
Universiti Putra Malaysia  
(Member)

**Noorsaadah Abd. Rahman, PhD**

Professor  
Faculty of Science  
Universiti Malaya  
(External Examiner)

---

**HASANAH MOHD GHAZALI, PhD**

Professor and Deputy Dean  
School of Graduate Studies  
Universiti Putra Malaysia

Date: 24 October 2007

This thesis was submitted to the Senate of Universiti Putra Malaysia and has been accepted as fulfilment of the requirement for the degree of Master of Science. The members of the Supervisory Committee were as follows:

**Abu Bakar Salleh, PhD**

Professor

Faculty of Biotechnology and Biomolecular Sciences

Universiti Putra Malaysia

(Chairman)

**Mahiran Basri, PhD**

Professor

Faculty of Science

Universiti Putra Malaysia

(Member)

**Raja Noor Zaliha Raja Abd. Rahman, PhD**

Professor

Faculty of Biotechnology and Biomolecular Sciences

Universiti Putra Malaysia

(Member)

**Mohd Basyaruddin Abdul Rahman, PhD**

Associate Professor

Faculty of Science

Universiti Putra Malaysia

(Member)

---

**AINI IDERIS, PhD**

Professor and Dean

School of Graduate Studies

Universiti Putra Malaysia

Date: 15 November 2007



## **DECLARATION**

I declare that the thesis is my original work except for quotations and citations which have been duly acknowledged. I also declare that it has not been previously, and is not concurrently, submitted for any other degree at Universiti Putra Malaysia or at any other institutions.

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**CHEONG KOK WHYE**

Date: 1 October 2007

## TABLE OF CONTENTS

	<b>Page</b>
<b>ABSTRACT</b>	ii
<b>ABSTRAK</b>	iv
<b>ACKNOWLEDGEMENTS</b>	vi
<b>APPROVAL</b>	viii
<b>DECLARATION</b>	x
<b>LIST OF TABLES</b>	xiii
<b>LIST OF FIGURES</b>	xiv
<b>LIST OF APPENDICES</b>	xvi
<b>LIST OF ABBREVIATIONS</b>	xvii
<b>CHAPTER</b>	
<b>1 INTRODUCTION</b>	<b>1</b>
<b>2 LITERATURE REVIEW</b>	<b>6</b>
Lipases	6
Versatile Biocatalysts	6
Thermostable Lipases	8
Analysis of T1 Lipase	9
Chemical Modification of Enzymes	15
Introduction	15
Reductive Alkylation of Amino Groups	18
Molten Globules	21
Probing Protein Structure by Modern Spectroscopy	26
Determination of Protein Conformation	26
Circular Dichroism (CD)	27
Detection and Localization of Modification Site	28
Protein Fluorescence: Utilizing Tryptophan and 8-Anilino-1-Naphthalenesulfonic Acid (ANS)	30
<b>3 MATERIALS AND METHODS</b>	<b>33</b>
Chemicals	33
Methods	36
Preparation of Culture	36
Purification of T1 Lipase	36
Sodium Dodecyl Sulphate-Polyacrylamide Gel Electrophoresis (SDS-PAGE) Analysis	37
Determination of Lipase Activity	37
Determination of Protein Concentration	38
Chemical Modification of T1 Lipase via Reductive Alkylation	38
Determination of Degree of Modification	39



Solvent Accessible Surface Area Analysis (SASA)	39
Circular Dichroism (CD) Spectroscopy and Thermal Denaturation Analysis	40
In-gel Digestion for Matrix Assisted Laser Desorption/Ionization Time of Flight Mass Spectrometry (MALDI-TOF MS) Analysis	40
Fluorescence Spectrofluorometry	43
<b>4 RESULTS AND DISCUSSION</b>	<b>44</b>
Source of T1 Lipase	44
Purification of T1 Lipase	44
Structural Studies of T1 Lipase	50
Solvent Accessibility Surface Area (SASA): Accessibility of Lysine's Side Chain	50
Alkylation Site: Reactivity of Lysine's Side Chain	52
Effect of Alkylation on Protein Conformation	59
Circular Dichroism (CD) Analysis: First Indication of a Molten Globule (MG)-like Structure	59
Thermal Unfolding	63
Determination of Alkylation Site: Matrix Assisted Laser Desorption/Ionization Time of Flight Mass Spectrometry (MALDI-TOF MS) Analysis	65
Fluorescence Spectroscopy Analysis: Shedding the Light of a Molten Globule (MG)-like puzzle	77
Formation of a Molten Globule (MG)-like Intermediate Structure	82
<b>5 CONCLUSION AND RECOMMENDATION</b>	<b>87</b>
Conclusion	87
Recommendation	88
<b>REFERENCES</b>	<b>89</b>
<b>APPENDICES</b>	<b>102</b>
<b>BIODATA OF THE AUTHOR</b>	<b>113</b>
<b>LIST OF PUBLICATIONS</b>	<b>114</b>



## LIST OF TABLES

<b>Table</b>		<b>Page</b>
1	Digestion properties of various proteolytic enzymes	41
2	Purification table of T1 lipase from <i>Geobacillus</i> sp. strain T1	49
3	The accessibility and exposure of lysine residues towards solvents	51
4	Side chain reactivities	56
5	The secondary elements of native and alkylated T1 lipase with regard to their hydrolytic activity and $T_m$	61
6	Theoretical peptide mass (native T1 lipase digested with trypsin)	67

## LIST OF FIGURES

<b>Figure</b>	<b>Page</b>
1 Crystal structure of mature T1 lipase	12
2 A closer view of the closed conformation of T1 lipase	13
3 Possible alkylation sites of T1 lipase	20
4 The structure of the molten globule of cytochrome b <sub>562</sub> (left) and the structure of the cytochrome's native state (right)	23
5 Chromatography profile of GST-tagged T1 lipase on Glutathione Sepharose HP	46
6 SDS-PAGE of purification profile of T1 lipase	46
7 Chromatography profile of cleaved T1 lipase on Sephadex G-25	48
8 Chromatography profile of cleaved T1 lipase on a series of attached affinity chromatography; Glutathione Sepharose HP, Glutathione Sepharose 4FF and Benzamidine FF	48
9 The mechanism of reductive alkylation	53
10 Ionic states of amino acid side chains as a function of the pH	57
11 Far-UV CD spectra of native and alkylated T1 lipase	61
12 Thermal transition curves obtained following the change in the CD signal at 222 nm for both native and alkylated lipase	64
13 MALDI-TOF mass spectrum for a tryptic digest of native T1 lipase	66
14 MALDI-TOF mass spectrum for a tryptic digest of alkylated T1 lipase	66
15 Local milieu of Lys102, which is surrounded by Glu100, Leu101, Arg103, Gly104 and Val71	68
16 Local milieu surrounding Lys185	69
17 Lys28 being enveloped by Phe25 and Glu22, creating hindrance for propionaldehyde molecules to penetrate the vicinity	70



18	Lys207 was heavily blanketed by the electronegative carboxyl group of Asp 205 and Asp209	71
19	A heavily hydrogen bonded Lys329	72
20	Lys344 and Lys345 in a close proximity environment	72
21	Local milieu of $\epsilon$ -amino of Lys185 and Lys229 which is located 5.77 Å apart	73
22	Alike Lys84, Lys138 possessed a highly exposed $\epsilon$ -amino	75
23	Exposure of Lys251's $\epsilon$ -amino group	76
24	A highly exposed $\epsilon$ -amino group of Lys84	76
25	Intrinsic fluorescence of native and alkylated T1 lipase	78
26	Extrinsic fluorescence of native and alkylated T1 lipase in the presence of 50 $\mu$ M of ANS probe	80
27	Calibration curve for determination of oleic acid concentration colorimetrically	111
28	Calibration curve for determination of protein content by Bradford assay	112

## LIST OF APPENDICES

Appendix	Page
A Amino acid sequence of <i>Geobacillus</i> sp. T1 thermostable lipase	103
B Solvent accessibility surface area (SASA)	104
C Calibration curve for determination of oleic acid concentration colorimetrically	111
D Calibration curve for determination of protein content by Bradford assay	112



## LIST OF ABBREVIATIONS

ANS	8-anilino-1-naphthalenesulfonic acid
BPTI	Bovine pancreatic trypsin inhibitor
CD	Circular dichroism
DTT	Dithiothreitol
FT-IR	Fourier transform – infra red
GST	Glutathione S-transferase
IPTG	Isopropylthio- $\beta$ -D-galactoside
kDa	Kilo Dalton
LB	Luria Bertani
MALDI-TOF MS	Matrix assisted laser desorption ionization time of flight mass spectrometry
MG	Molten globule
NMR	Nuclear magnetic resonance
PBS	Phosphate buffer saline
PTH	Phenylthiohydantoin
SASA	Solvent accessible surface area
SDS PAGE	Sodium dodecyl sulphate-polyacrylamide gel electrophoresis
$T_m$	Midpoint of sigmoidal melting curves
TNBS	2,4,6-trinitrobenzenesulphonic acid



## CHAPTER 1

### INTRODUCTION

Since the booming development in enzymology, the manipulation of the enzyme structure was seen as a new way to enhance its biochemical properties, be it chemical modifications (Mine *et al.*, 2001; Basri *et al.*, 1998), protein engineering (Bornscheuer, 2002; Bornscheuer and Pohl, 2001; Harris and Craik, 1998) or by changing the reaction medium rather than the enzyme itself (Klibanov, 1989). Native enzymes are limited and not specific in its application. In contrast, modified enzymes with specific purpose are useful in certain fields. Therefore, modifications have been carried out in order to increase their activities, selectivity and stability. Chemically modified enzymes was seen as a fruitful research in Basri and co-workers' work (1998) and Ampon *et al.* (1991), as these lipases' enantioselectivity and specificity were successfully enhanced.

Protein chemists have long been interested in altering the chemical, physical and biological properties of proteins by chemically changing their structure. One of the first things discovered about proteins was how easily they were changed upon treatment with chemical reagents. Their liability to chemical reagents and reaction conditions has been a serious problem for them to be used for many purposes. The application of modern knowledge of proteins, new chemical reagents and more sophisticated analytical techniques, however, has made chemical modification of protein as one of the most useful approaches to the study many of their properties.



Numerous efforts are being made in the improvement of enzyme efficiency to exploit their environmental friendly processes more extensively. The alteration of protein surface characteristics by chemical modification is a good strategy to improve biocatalyst performance in nonaqueous conditions (Longo and Combes, 1999). Attachment of hydrophobic groups using aldehydes to the enzyme surface has been considered to increase enzyme solubility in organic solvent (Inada *et al.*, 1986). Yet, the essential of chemical modification of proteins lie beneath the opportunities for studying localization of individual amino acids in proteins, their participation in the maintenance of the native conformation (Torchilin *et al.*, 1979), their stabilization (Ryan *et al.*, 1994), conversion to molten globule structures (Dolginova *et al.*, 1992), tailoring of enzyme specificities and their structure-function relationship (Wong and Wong, 1992).

Reductive alkylation is a convenient method to convert surface exposed amino groups in proteins into their alkylamino derivatives. The functional residues are located on or near the surface of proteins, and thus are reactive to chemical reagents (Alberts *et al.*, 1983). Unlike other procedures for the modification of amino groups of proteins, reductive alkylation has little effect on the physicochemical properties of protein (Inada *et al.*, 1986). Alkyl substitution of hydrogen will increase both the bulkiness and hydrophobicity of the amino groups and reduce its ability to form hydrogen bonds (Ampon *et al.*, 1993; Means and Feeney, 1971).

One of the most interesting and well-investigated class of enzymes in this particular field is lipase or triacylglycerol hydrolases (E.C. 3.1.1.3). Lipases also

catalyze various useful reactions, for instance hydrolysis, esterification, transesterification and polyesterification reactions, and act as chiral catalysts in the production of various fine-chemicals and intermediates (Miyawaki and Nakamura, 2002; Berglund, 2001; Jääskeläinen *et al.*, 1997). The diverse functions and the enzyme specificity, both stereospecificity and regiospecificity, make lipase one of the most important biocatalyst in biotechnological applications (Shaw, 2002). The stringent requirements associated with the use of enzyme in industry reflect the need of lipases which have the desired characteristics such as thermostability and solvent stability. Thermostable lipases are expected to play a significant role in industrial processing because running bioprocess at elevated temperature lead to higher diffusion rate, increased solubility of lipids and hydrophobic substrates in water and reduced risk of contamination (Becker *et al.*, 1997).

The lipase used in the present study, from *Geobacillus* sp. strain T1 was isolated together with 28 other putative lipase producers from palm oil mill effluent in Malaysia by Leow and co-workers in 2004 (Leow *et al.*, 2004). T1 was selected for further studies as T1 has the highest lipase production rate, 0.15 Uml<sup>-1</sup>. Later, it was identified as *Geobacillus* sp. strain T1, which has a maximum activity at 70 °C and pH optimum of 9.0. It was cloned, over-expressed and purified to homogeneity before the protein was used for chemical modification purposes (Rahman *et al.*, 2005). The fact this is a thermostable enzyme added new perspective to this study as this is the first studies carried out on a thermostable enzyme. The crystal structure of this 43 kDa protein has been resolved at 1.8 Å resolutions (Matsumura *et al.*, 2007).

Since the late 1960s, the molecular mechanisms of thermostability of enzymes from thermophilic microorganisms have attracted much interest from scientists from the fundamental point of view and from engineers engaging in biotechnology from the applied point of view (Mozhaev and Martinek, 1984; Mozhaev *et al.*, 1988a). With the rapid development of chemical reagents and methods, chemical modification coupled with spectroscopy has become a handy tool with the growing needs for identifying and modifying the functional amino acid residues in proteins.

Over the past 20 years, there have been major developments in the application of spectroscopic methods to systems of biological interest. Physical scientists, biological scientists and engineers have an impressive array of powerful and elegant tools for gathering qualitative and quantitative information about composition and structure of matter. With the increase in interdisciplinary research in recent years, the need for accurate and sensitive methods for the analysis of biomolecules has been increasingly important for both the chemist and biologist. The three-dimensional structure of a protein is governed by its primary structure (sequence of amino acids) and its environment. Changes in either of these can have important effects on its properties. Modifying any of its amino acid residues necessarily changes the primary structure of a protein. Thus, sophisticated analytical methods such as circular dichroism spectropolarimetry (CD) (Choi *et al.*, 2005; Kelly *et al.*, 2005; Hosseinkhani *et al.*, 2004; Greenfield, 1999), matrix-assisted laser desorption/ionization time of flight mass spectrometry (MALDI-TOF MS) (Wa *et al.*, 2006; Henzel *et al.*, 2003) and

fluorescence spectroscopy (Mossavarali *et al.*, 2006; Hosseinkhani *et al.*, 2004) have become standard tools to probe these changes.

Thus, the principal objective of this thesis is to probe the effect of modification on lipase via reductive alkylation by means of spectroscopy methods. Hence, the research was undertaken with the following specific objectives:

- To modify T1 lipase using propionaldehyde through reductive alkylation with different degree of modification.
- To investigate the effect of modification on T1 lipase's structure by using CD, MALDI-TOF MS and fluorescence spectroscopy.
- To evaluate the extent of modification on T1 lipase's structure using both spectroscopy data and structural analysis.



## CHAPTER 2

### LITERATURE REVIEW

#### Lipases

##### **Versatile Biocatalysts**

The demand for industrial enzymes is ever increasing owing to their applications in a wide variety of processes. Enzyme-mediated reactions are attractive alternatives to tedious and expensive chemical methods. Enzymes find great use in a large number of fields such as food, dairy, pharmaceutical, detergent, textile, fine chemicals and cosmetic industries (van Beilen and Zhi Li, 2002). With the realization of the biocatalytic potential of lipases in both aqueous and nonaqueous media in the last one and a half decades, industry has shifted towards utilizing this enzyme for a variety of reactions of immense importance. Enzymes are used quite extensively as industrial catalysts as they offer the following advantages in comparison with chemical catalysts (Humphrey and Lee, 1983):

1. They are specific, thus minimizing the undesirable side reactions and by-products.
2. They are relatively cheap when used in crude form.
3. They are effective catalysts for chemical conversion, offering mild conditions for reaction; low temperatures and pressures.
4. They are relatively non-toxic and thus, acceptable for applications in food processes and medicinal purposes.
5. They are effective within a wide range of substrate concentrations.