Effects of Lid 1 Mutagenesis on Lid Displacement, Catalytic Performances and Thermostability of Cold-active Pseudomonas AMS8 Lipase in Toluene

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A R T I C L E   I N F O
Article history:
Received 7 November 2018
Received in revised form 14 January 2019
Accepted 19 January 2019
Available online 25 January 2019

Keywords:
Lid engineering
Cold-active lipase
Molecular dynamics simulation
Hydrophobic contacts
Site-directed mutagenesis
Solvent accessibility
Substrate affinity
Thermostability

A B S T R A C T
Pseudomonas fluorescens AMS8 lipase lid 1 structure is rigid and holds unclear roles due to the absence of solvent-interactions. Lid 1 region was stabilized by 17 hydrogen bond linkages and displayed lower mean hydrophobicity (0.596) compared to MS38 lipase. Mutating lid 1 residues, Thr-52 and Gly-55 to aromatic hydrophobic-polar tyrosine would churned more side-chain interactions between lid 1 and water or toluene. This study revealed that T52Y leads G55Y and its recombinant towards achieving higher solvent-accessible surface area and longer half-life at 25 to 37 °C in 0.5% (v/v) toluene. T52Y also exhibited better substrate affinity with long-chain carbon substrate in aqueous medium. The affinity for pNP palmitate, laurate and caprylate increased in 0.5% (v/v) toluene in recombinant AMS8, but the affinity in similar substrates was substantially declined in lid 1 mutated lipases. Regarding enzyme efficiency, the recombinant AMS8 lipase displayed highest value of $k_\text{cat}/K_m$ in 0.5% (v/v) toluene, mainly with pNPC. In both hydrolysis reactions with 0% and 0.5% (v/v) toluene, the enzyme efficiency of G55Y was found higher than T52Y for pNPL and pNPP. At 0.5% (v/v) toluene, both mutants showed reductions in activation energy and enthalpy values as temperature increased from 25 to 35 °C, displaying better catalytic functions. Only T52Y exhibited increase in entropy values at 0.5% (v/v) toluene indicating structure stability. As a conclusion, Thr-52 and Gly-55 are important residues for lid 1 stability as their existence helps to retain the geometrical structure of alpha-helix and connecting hinge.

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1. Introduction

The structure-guided protein engineering is a strategy used to identify and regulate the molecular interactions happening around certain area of interest which involved in enzymatic functions so that it can be develop into a compelling biocatalyst for industrial applications [1]. Amid many industrial enzymes, lipases contribute to many important roles in biocatalysis process due to its extensive usages in detergents formulation, food preservations and flavorings, biodiesel productions and cosmetics active ingredients [2]. In some lipases, a mobile amphipathic structure known as lid mainly covers the catalytic active site region where the length and complexity of the structure depends entirely on the shape of enzymes. The lid is very much involved in modulating activity and substrate selectivity in lipases. Depending on the target enzymes, protein engineering on lid structures may altered important properties such as enzyme substrate specificity and local stability [3]. In addition, it is often to observe a reduction in enzyme activity whenever lid paves way to selected compounds of increasing lipophilic characteristics. Non-polar solvent such as toluene is one of the medium which supports the solubility of amphipathic molecules/compounds where interactions with lid allowing substrate access to the active site and initiation of catalysis. The intrinsic conformational changes of the enzyme initiated by lid domain are important determinants for lipase activation especially in organic solvents [4].

Mutations on lid sequence would cause in different interactions of enzyme towards the presence of organic solvents. Previously, the lid engineered Candida rugosa lipase activity and enantioselectivity was found reduced in organic solvents. Cold-active Pseudomonas fragi lipase has its chain length specificity altered while experiencing an increase in

https://doi.org/10.1016/j.csbj.2019.01.005
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thermostability, also in the presence of organic solvents [5]. Later, Jørgensen et al. described that lipase would undergo structural transition from close to open state when being exposed to high solvent polarities [4]. Normally, lid has a weaker tendency to orient itself in a preferred orientation between a polar and apolar medium. In the case of Thermomyces lanuginosus lipase E87K, a negatively-charged residues such as glutamic acid (E) that serves to stabilize the lid in a closed conformation can potentially be mutated with an opposite charge residue such as lysine (K) to allow lipase activation at high solvent polarities. This amino acid substitution on α-helix domain results in increase of energy needed to activate lid opening at low dielectric constants [4]. As the enzyme shifts to the open conformation, the hydrophobic interphase was exposed, making the enzyme even more accessible to substrate. In the meantime, lid replacement in Candida rugosa lipase has resulted in substrate-independent hydrolysis reaction of p-nitrophenol esters which responded to added detergents. The change in lid position by detergent was stimulated by the shift of reaction equilibrium of different conformation state in detergent/interfaces [6]. In cold adapted enzymes, the lid modulation depends on the conformational flexibility as part of a molecular adaptation towards consistency of interactions between lid and its surroundings [7]. Thus, it is frequently said that lid appears to display a very complex role in lipases activity, specificity and conformational stability subject to variations of amino acid sequences and its surrounding. In addition, surface accessibility is important for any residue involved with exterior interactions [8].

Previous works highlighted the importance of the lid sequence to structure, mechanism and function via protein engineering (lid swapping and site-directed mutagenesis) and computational analysis focusing on lid domain. Based on these approaches, the role of the lipase lid in substrate selectivity and activity in the presence of solvents or surfactants or deterrents can be identified. In lipases, lid was observed to have a distinct appearance in three groups. The first ones are lipases without lid, secondly, lipases with one loop or one helix lid and the last ones are lipases with two or more helical lids [9]. Cold-active Pseudomonas fluorescens AMS8 lipase shared similarity to lid sequence from lipase of Pseudomonas sp. MIS38, and thermophilic lipases from Proteus mirabilis and Geobacillus stearothermophilus [10,11,12], where these lipases clearly showed to have two helical lids [8]. Lid 1 is a short-hinged-alpha-helix lid located at residue 51–57 and is found inactive (structurally rigid) in the presence of organic solvent, whereas the second and longer alpha-helix lid 2 (148–167) is saturatedly composed with hydrophobic residues. Unlike lid 1, the role of lid 2 in interfacial activation and substrate accessibility has been associated with interactions in non-polar organic solvents.

The main objective of this work is to determine the effects of lid 1 mutation to physical and biochemical properties of cold-active Pseudomonas AMS8 lipase in hydrophobic/aqueous interphase. It is hypothesized that higher activity of lipase could be achieved in toluene if lid 1 gets shifted from 'close' to 'open' state conformation via increased in solvent-surface interactions. By making lid 1 to co-participate lid 2 in the activation of lipase, the barriers in obtaining higher lipase activity could be overcome. In current state, it was difficult to understand the catalytic behavior of cold-active lipase with two lids where the function relies heavily on lid activation and its biomechanics. In the context of current study, lid activation involves the voluntary opening and closure of alpha-helix structure in organic solvent via surface interactions to stimulate better catalysis.

2. Materials and Methods

2.1. Prediction of Protein Stability, Solvent-accessibility and Flexibility of Lid 1 Mutant Candidates

The knowledge-based approach was used to predict the effects of site-directed mutagenesis (SDM) on protein stability and to calculate the change in thermal stability between recombinant and mutant lipases (http://marid.bioc.cam.ac.uk/sdm2/). PDB file containing the 3D coordinates of the protein atoms of AMS8 lipase was submitted to the server. Following this, a summary of analysis is presented, stressing on details of the recombinant and mutant residues, residue number and protein chain. Another list features the prediction analysis of mainchain conformation, side-chain solvent accessibility, side-chain hydrogen bonding pattern for both recombinant and mutant residues [13]. The physico-chemical features of α-helix sequence of lid 1 (51–57) and its outskirt (+ 5) residues in recombinant AMS8 lipase was explored via online server, HeliQuest (http://heliquest.ipmc.cnrs.fr/cgi-bin/ComputParams.py), This module allows specific properties such as hydrophobicity, hydrophobic moment, net charge (z) and amino acid composition for several helix types to be determined.

2.2. Lid 1 Mutant Designed, Expression, and Purification

A polar Thr-52 and neutral Gly-55 located at the center of lid 1 region were chosen as the target for mutation and substituted with tyrosine which has aromatic side chain and hydroxyl group at its benzene ring making it semihydrophobic. Mutant T52Y and G55Y lipases were obtained by site-directed mutagenesis using a previous construct recombinant pET32b–AM8S plasmid as DNA template [14,15]. Primers used for rational mutagenesis were designed via web-based automated program, QuikChange Primer Design (http://www.genomics.agilent.com/primerDesignProgram.jsp) and synthesized by Life Technology (Guangzhou, China) as shown in Table 1.

A recombinant lipase with expression vector pET32b + harboring mutation site was prepared by using QuickChange Lightning Site-Directed Mutagenesis kit (Catalog No. #210519, Agilent Technologies, USA) where a few colonies were sterilely picked and plasmid extracted for sequencing. The positive lid 1 mutant lipases were then digested with restriction enzymes, BamH1 and XhoI (NEB, USA) and re-transformed into expression host, E. coli BL21 (DE3) from Novagen®, Germany.

Subsequent expression and purification of recombinant AMS8 and its mutant lipases were carried out according to the methods previously described [15]. In brief, both recombinant AMS8 and lid 1 mutant lipases (T52Y and G55Y) were expressed as fusion protein with histidine binding domain tag for the purpose of affinity purification via Nickel-Sepharose Excel (GE, USA) followed by anion-exchange chromatography (IEC) purification by using 2 × 5 mL pre-packed column of Hitrap Q HP (GE, USA). The purified proteins were collected, pooled and concentrated using ultra-centrifugation (Amicon, Merck). The purity of expressed lipase was analysed by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS–PAGE) [16]. Protein concentrations were measured by using Bio-Rad Protein Assay (Bio-Rad Laboratories, USA) [17].

2.3. Molecular Dynamics (MD) Simulations of Recombinant and Mutated Lid 1 AMS8 Lipase

The MD simulations were performed on computed model of recombinant AMS8 and lid 1 mutant lipases using YASARA structure software (version 11.3.22, Netherlands) following previous MD settings and protocols in water or toluene [7,18] targeting specific temperature of 25 °C and 37 °C. The system was equilibrated at the steepest descent parameters, using a time step of 1 fs for intramolecular forces which calculated the intermolecular forces in every 2 simulation sub-steps. The structure was energy-minimized with AMBER03 force field using a cut off 10.486 Å and the Particle Mesh Ewald algorithm to treat long range electrostatic interactions. After the removal of conformational stress by a short steepest descent minimization via 50 steps of MD in solvent, the procedures continued by simulated annealing and converge as soon as the energy improves by -0.05 kJ/mol per atom within 200 steps. The changes of protein structure were examined using 800 saved
trjectories where each 40 simulation snapshots represents 1 ns. The MD simulation was allowed to run until 20 ns. By using final trajectories from MD simulations, the hydrogen-bond networks in recombinant AMS8 and mutant lipases, T52Y and G55Y were explored by BIOVIA Discovery Studio 2017R2 Visualiser version 17.2.0.16,349 (Dassault Systemes, 2016).

2.4. Normal Mode Analysis (NMA) and Sequence-specific Solvent Accessibilities (ProtSA) Analysis of Lid 1 Mutants

Normal mode analysis (NMA) is a fast and simple method to calculate vibrational modes and protein flexibility [19]. To get this information, the final trajectories of simulated lipase in toluene was taken and submitted to Elastic Network Model (ENM) website: http://www.sciences.univ-nantes.fr/elnemo/. ENM is an algorithm that enables quantitative description of protein behaviour starting from three dimensions (3D) view of protein assemblies to the network models showing the connecting pairs of nodes located within a distance measured under harmonic restrictions from the equilibrium structure [19]. The application offers calculations of normal or vibrational modes on all atoms by fixing it on Cα.

ProtSA calculates sequence specific protein solvent accessibilities in the unfolded ensemble by simulating the unfolded protein many times and combining the results. In the simulations, the structural model to describe the unfolded conformations representative of the unfolded protein is generated by the Flexible-Meccano algorithm [20]. The analytical software ALPHASURF is applied to calculate atom solvent accessibilities. The tool was readily available for download at: http://webapps.bifi.es/protsa/#Xbernaudo:2006.

2.5. Biochemical Properties of AMS8 Lipase and its Mutant

The hydrolysis activity of AMS8 and lid 1 mutant lipases was measured spectrophotometrically by using Kwon and Rhee method [21]. The enzyme kinetics was measured by p-nitrophenyl caprylate (pNP-C8), laurate (pNP-C12) and palmitate (pNP-C16) as substrates. Each reaction mixture consists an approximate 10 μl of 20 mM pNP solution dissolved in 2-propanol was spiked into a 100 μl of 50 mM Tris-HCl, 5 mM CaCl₂ buffer (pH 8). 5 mM CaCl₂ pH 8 was used for Dynamic Light Scattering (DLS) measurement from 20 to 70 °C. Data were analysed using Zetasizer software (v 7.11) to compare reduction in molecular size (d.nm) or volume size of proteins after experienced increase of temperature with and without toluene (0.5%, v/v).

2.7. Thermodynamic Analysis

Both recombinant AMS8 and mutant lipases were purified and assayed in substrate, pNPC at different temperatures (25–45 °C) for the determination of kcat and to plot Arrhenius graph. Activation energy (Ea) of respective lipases was calculated by identifying the slope of Arrhenius graph showing a series of data in ln kcat against 1/T (K⁻¹). or, alternatively by calculating Ea by means of formula:

$$\ln \frac{k_2}{k_1} = -\frac{Ea}{R}\left(\frac{1}{T_2} - \frac{1}{T_1}\right)$$

(1)

where, T₁ and T₂ is the temperature in Kelvin (K) at 25 °C and 35 °C. k₁ and k₂ is the kcat of lipase activity in pNPC measured at 25 °C and 35 °C, respectively.

As for measuring Gibbs-free activation energy (ΔGi), enthalpy (ΔHi), and entropy (ΔSi) per single temperature, equations were used for calculation [24,25]. The magnitude of energy required to achieve the same position of equilibrium in both forward and reverse reactions can be explained in terms of various thermodynamic parameters such as changes of Gibbs free energy (G), enthalpy (H) and entropy (S) for both reactants and products involved in a reaction [26].

$$\Delta G_i = RTx\left[\ln \frac{k_i T}{R} - \ln k\right]$$

(2)

$$\Delta H_i = Ea - RT$$

(3)

$$\Delta S_i = (\Delta H_i - \Delta G_i)/T$$

(4)

In the above equations, kᵢ is the Boltzmann constant (1.3805 × 10⁻²³ J K⁻¹), h is the Planck constant (6.6256 × 10⁻³⁴ Js), k is the catalytic rate constant, and R is the gas constant (8.314 J mol⁻¹ K⁻¹) [24].
3. Results and Discussion

3.1. Preliminary Analysis and Outcome Predictions on Lid 1 Mutation

Lipases with special properties such as thermostable, organic solvent friendly and active at broad temperatures receive great industrial attention because of their usability under restricted reaction condition [27]. In this study, AMS8 lipase was special due to its ability to tolerate the harshness of organic solvent. Cold-active AMS8 lipase behaved as a soluble protein in water and so did other polar organic solvents. Even though the lipase is soluble, polar organic solvents had shown to cause in the destabilization of the protein structure. In addition, polar organic solvent was found not to stimulate lid activation like non-polar organic solvent did [18]. This was because the structure flexibility exhibited by lid 2 regions (148–167) enabled the lid to open and formed a tunnel for the substrate to move into the catalytic pocket. Hydrophobic core was exposed after lid was flipped to the side which attracting the non-soluble and lipid-based substrate to bind. The contributions between AMS8 lid structure and non-polar solvent interactions improves protein surface interactions as it enhances lipase catalysis in organic solvent especially toluene. AMS8 lipase possessed two lids and with molecular simulation, only lid 2 showed an observable movement upon non-polar solvent contact which was not detected at lid 1 (51–57). In MIS38 *Pseudomonas lipase*, there are two lids (lid 1: residues 140–167 and lid 2: residues 46–74) that harbour similarity in sequence homology to the currently studied AMS8 lipase. In this context, the lids in AMS8 lipase were each addressed as lid 1 for residues 51–57 and lid 2 for residues 148–167, opposite to MIS38 naming of lid. In *Pseudomonas* MIS38 lipase, both lids exhibited movement and sustain open conformation in aqueous environment [28]. The role of hydrophobic interactions in MIS38 lipase interfacial activation has been confirmed by the presence of proline which acts as a hinge residue to its lid 2 helix situated at residue 50. Meanwhile in *Aspergillus niger* lipase, prolines in the hinge domain of lid serve to stabilize the open lid conformation as a mean of substrate entry [4]. The lid 2 sequence in MIS38 lipase was “PATLVALLGG”. In this study, AMS8 lipase lid 1 sequence was differentiated by two residues, Gly-55 and Ser-60 – “PATLVGALLG”. These two amino acids substitutions would have caused lid 1 to adopt a rigid structure instead of a flexible one. In AMS8 lipase, Gly-55 has a lighter residue weight compared to Thr-55 but Ser-60 might help in establishing an intramolecular H-bonding with Gly-55 making the region highly stable and movement restricted. Ser-60 also was found to be the reason behind the lack of hydrophobic interface of lid 1 as shown in Table 2.

The movement of an α-helical lid by rotating around two hinge regions at the lipid-water interface created a hydrophobic patch surrounding the catalytic triad, resulting in activation of the lipase [29]. Instead of changing the residues of alpha-hinge Ser-60 to Gly-60 to make available on hydrophobic patch, this study focused on to improve the flexibility of the lid domain (in this case lid 1) by providing anchor (via side-chain interactions) to toluene on the alpha-helix surface of lid 1 as alternative approach for “interfacial activation” and to enforce the mobility of these H-bond saturated area by allowing the hinge to move about freely.

Our interest is to improve the hydrophobic interactions for better surface activation in non-polar organic solvent at lid 1 (51–57) by targeting residues Thr-52 and Gly-55. At pH 8, both Thr-52 and Gly-55 were found to be uncharged and to promote interactions with non-polar organic solvent, aromatic residues such as Tyr, Trp and Phe could offer better hydrophobic interactions. However, assigning an aromatic residue to a small alpha-helix lid could alter the coordination of lid in facing active site. A close check on peptide sequences showed that lid 1 was built by a successive arrangement of polar, neutral (no charge) and hydrophobic-aliphatic amino acids which do not attract any bond formations or ionic interactions between their side chains.

### Table 2

Helical domain analysis (46–63) in MIS38, AMS8 and lid 1 mutant lipases. This analysis showing the distribution of amino acid residues and the orientation of the amphipathic moments (black arrows) calculated on the basis of the hydrophobicity and side chain distribution of the amino acids in the helices.

| Name of lipase | Alpha-helix domain 46-63 | Mean hydrophobicity | Hydrophobic moment | Hydrophobic patch (residues) | Helical wheel of alpha-helix domain 46-63 | Close-up view of lid 1 and hinge | Total no. of hydrogen bonds (residues 49-59) |
|----------------|---------------------------|---------------------|---------------------|--------------------------------|--------------------------------------|--------------------------------|-----------------------------------|
| MIS38 lipase (2Z8Z) reference | GLGLPAT LVTALLG STD| 0.613  | 0.276 | A56 L49 S60 L53 |  | 13 |
| Recombinant AMS8 | GLGLPAT LVTALLG STD| 0.596  | 0.287 | A56 L49 S60 L53 |  | 17 |
| T52Y | GLGLPAT LVTALLG STD| 0.635  | 0.252 | A56 L49 S60 L53 |  | 10 |
| G55Y | GLGLPAT LVTALLG STD| 0.649  | 0.246 | A56 L49 S60 L53 |  | 8 |
and the respective solvents. Site-directed mutagenesis on lid 1 surface could facilitate various approaches associated with function prediction and stability in organic solvent. Based on helix domain analysis highlighting residues 46–63, AMS8 lipase showed a distinct distribution of hydrophobic amino acids in the helix that generates a large amphipathic moment (based on the direction pointed by arrow) supposedly making favourable hydrophobic interactions with the hydrophobic active site (Table 2).

The only difference between AMS8 lipase lid 1 and MIS38 lid 2 helix domain was the availability of hydrophobic patch that display better amphipathic moment. In this case, MIS38 lipase has better hydrophobic interaction with the active site. Both mutations on lid 1 targeting residues Thr-52 and Gly-55 did enhanced the mean hydrophobicity of AMS8 lipase after substitution with Tyrosine. Thr-52 is targeted because it is polar in characteristic and can accommodate rotamers which indicates that side chains adopt unstrained conformation. After being replaced with an aromatic amino acid such as Tyr, differences could be observed between the orientation of side chain and stalkling effects with aromatic-organic solvent such as toluene. Although tyrosine is hydrophobic, it has lower pKa due to the presence of phenolic hydroxyl (a polar side group) which makes it ionizable and highly stable on the protein surface. Mutation at Gly-55 (located between hydrophobic-aliphatic Val-54 and Ala-56) was targeted and substituted with Tyr-55 as Gly-55 did not form any important interactions (polar or non-polar) within this region. Both mutation sites especially the side-chain were predicted not to form any H-bond with other neighbouring residues and the differences in free energy strongly indicates that both mutated protein has impact on protein stability and activity as shown in Table 3. The predictions on these potential mutants were assessed through SDM, a useful tool for predicting the effects of a mutation on protein structure and function. In general, computational approaches to predict protein stability (or protein stability changes) is physics-based, knowledge-based, or a hybrid [30].

### 3.2. Effects of Temperature to Lid 1 Mutation

Temperature-mediated protein interactions has been associated with many changes of enzymatic activities. This makes temperature, one of the crucial factors that trigger lid 2 interfacial activation in cold-active AM8 lipase other than the hydrophobic organic solvents [7,31]. It would be interesting to see the outcomes of structural changes at the lid area in response to temperature and toluene. Previous simulation of solvent-stable lipase, Lip42 in water-solvent shows the perturbation of hydrophobic clusters and structural changes due to the open-closure of the lid [32]. In this study, the conformational change after lid 1 mutation was not found in 25 °C simulation, except for the large displacement observed in recombinant AMS8 lid 2 residues (Fig. 1A). However, the change in the conformational movement was found to be linked with lid 1 when both mutants (T52Y and G55Y) were subjected to MD simulation at higher temperature, 37 °C (Fig. 1B). Recombinant AMS8 lipase appeared to have a low atomic displacement at 37 °C in which both lids turned out rigid and static. Such thermal–inducing motions are central importance to protein function and stability. A slight difference in temperature can influence the flexibility of cold-active lipase via disruption on its interfacial lid. At first, lid 1 did not assume to its role in lid activation under its normal temperature, 25 °C but after mutation, a significant difference could be observed at 37 °C in the presence of toluene.

Based on molecular dynamics analysis, lid 1 and lid 2 undergoes conformational changes with increase of temperature. Most importantly, lid 1 appeared to respond towards moderate temperature, 37 °C probably requiring higher force to stimuli its recognition to the presence of toluene. By raising the temperature to 37 °C, Gly-69 exhibited greater flexibility in T52Y and AMS8 recombinant lipase (Fig. 2A, Fig. 2B). The reason behind it is that, Gly-69 could not interact strongly with other neighbouring residues that are hydrophobic. Therefore, direct exposure of this residue to the non-polar solvents might trigger the flexibility. In terms of property, Glycine has no side chain and electrostatic charges which make the residue easily moved by surface bound water molecules via hydrophilic interactions. This highly flexible Gly-69 could also be the starting point of protein unfolding due to their large fluctuation and disorientations. The catalytic effects caused by Gly-69 flexibility can be observed through a slight increase of catalytic rate (kcat) at 35 °C. However, the activation energy required for lipase activity to occur is found higher in toluene at 25–35 °C probably due to the restricted flexibility of lid 2 when temperature starts to increase. At 37 °C, lid 2 responded by showing reductions in RMSF value. A declined lid 2 flexibility of recombinant AMS8 lipase means that overall structure of lipase became highly stable at 37 °C and required more activation energy for lipase activity.

As for T52Y, lid 1 and lid 2 were greatly affected by temperature increase (Fig. 2B). The high RMSF value of Gly-69 at 37 °C showed that T52Y capable of changing the structure of lid 1 and lid 2, while a lower RMSF values in G55Y indicated that this mutation had brought no effect towards change of lid 1 structure in particular of that site (Fig. 2C). Interestingly, Gly-69 was found located at the posterior hinge domain of lid 1 that could have influenced the subsequent shift (in RMSF) of lid 1 at 37 °C. In this study, simulation induced by temperature showed a significantly large fluctuating area on lid 1 aided by the presence of toluene. Lid 1 might have special roles and it could be based on mutual dynamics of both thermostability and solvent-recognition. Previously, activity measurements and MD simulations on psychrophilic AM88, M37 and therophilic Thermomyces lanuginosus lipase (TIL) suggest that the opening of the lid is energetically unfavorable in aqueous solutions until temperature or hydrophobic solvent triggered the interface activation [15,33,34].

Fig. 3 (A) shows that by changing the temperature from 25 °C to 37 °C, the RMSD values of recombinant lipase AMS8 declined from 7 Å to 4 Å. The reductions of RMSD values in mutant T52Y were also reported following increased in temperature. At 25 °C, T52Y showed a sharp irregular increase of RMSD between 0 and 5 ns before slowly declined to 5 Å and stabilized at 4 Å until the simulation reached 20 ns. In contrast, G55Y showed no difference in terms of overall RMSD values at both temperatures showing no significant changes on average distance, stressing on close similarity of its tertiary structure. Previously, a minimal RMSF values observed in lid 1 residues of mutant G55Y in toluene at both temperatures could be inter-relate to the lack of RMSD changes. The key indicators to describe temperature-dependence in this cold-active lipases are through protein compactness and surface area solvent accessibility (SASA). H-bond networks are crucial for protein compactness and when these interactions broke at high temperature (causing structure flexibility), both Ra and SASA started to increase [35]. In Fig. 3 (B), mutants T52Y and G55Y exhibited low in R-gyration when

### Table 3

Predicting effects of mutagenesis on lid 1 of AMS8 lipase.

| Residues substitution | Secondary structure | Solvent accessibility (%) | Side-chain forming hydrogen bond | Protein stability free energy change (ΔΔG) in Kcal/mol |
|-----------------------|---------------------|---------------------------|---------------------------------|------------------------------------------------------|
| T52Y                  | Alpha-helix         | 51.9 → 42.5 (partially accessible) | Saturated → No H-bond           | 1.98 (Stabilizing, retain or improve activity)        |
| G55Y                  | Alpha-helix         | 11.5 → 34.6 (buried to partially accessible) | No H-bond → No H-bond           | 4.78 (Highly stabilizing, negatively affect activity) |

Web server, Site Directed Mutator (SDM) [http://www-cryst.bio.cam.ac.uk/~sdm/sdm.php](http://www-cryst.bio.cam.ac.uk/~sdm/sdm.php)
temperature was set at 37 °C. However, SASA values showed no difference in both temperatures (Fig. 3 (C)). Recombinant AMS8 lipase had the opposite trend at 37 °C, showing higher R-gyration and SASA values. Heat causes T52Y and G55Y structures to stabilize and settle at the point where proteins undergo less structural changes. Water clusters breaks apart, distributed across the protein conformation but the H2O molecules should remain there. This analysis shows that mutations at lid 1 help to retain the H-bond interactions and water clusters surrounding enzyme surface.

3.3. Normal Mode Analysis (NMA) and Sequence-Specific Solvent Accessibilities (ProtSA) of Lid 1 Mutants

Normal mode analysis (NMA) is used to investigate collective motions and large-scale conformational transitions in proteins [36,37]. Apart from localized motions (detected by MD simulation), large-scale motions (detected by NMA) also play important role in the functional motions of an enzyme. According to matrix that compares 3 out of 5 modes of computed NMA analysis, recombinant AMS8 lipase has greater local flexibility at residues 140–170 specifically located at lid 2 regions (Fig. 4 A). In T52Y, 2 out of 5 modes were found to show greater CO distance at residues 153–190 whereas in G55Y, 4 out of 5 modes were found to have largest fluctuated CO distance starting from residues 48–59 and 162–172 (Fig. 4B and 4C). Gly-159 and Leu-163 exhibited largest increase in CO distance by referring mutated Tyr-52 as a reference point. Leu-162 and Pro-168 were greatly distanced from Tyr-55 CO backbone. These highly flexible residues were mostly found at lid 1 help to retain the H-bond interactions and water clusters surrounding enzyme surface.

There were two pre-indications that could define the overall role of lid 1 of AMS8 lipase after mutations. First, lid 1 did not participate in the process of interfacial activation. There was very little flexibility shown by this region in organic solvent and with increase of temperature [7,18]. This means, lid 1 would have existed as an imitation of a smaller lid which looks quite similar to lid 2 in Pseudomonas sp. MIS38 [10,38]. Secondly, the distance and coordination of active site residues were found to be greatly affected by lid 2 movements in the presence of toluene. The effects of mutations at residue Thr-52 and Gly-55 of AMS8 lipase showed a slight movement at lid 1 region and minimal disruption on lid 2 activation as the changes in folding by toluene causing the two lids to move against each other. Mutation that targets Thr-
52 and Gly-55 on lid 1 confirmed the importance of this region to temperature stability subjected to biophysical changes. Altered optimal temperature, melting temperature, half-life and toluene stability was appropriate to describe the dependency of AMS8 lipase on the rigidity of lid 1. Molecular dynamics simulations at 37 °C showed that by mutating Thr-52 and Gly-55, the short lid 1 of AMS8 lipase may experience larger atomic movements. These movements were known to cause in regional flexibility of lid 1 without sacrificing much of the protein stability and was able to increase the solvent accessible surface area (ASA).

Since the native state of protein can be recognized by losing the solvent accessible surface area (ASA), it was highly anticipated that the structural changes exhibited by these mutants at this temperature was contributed by protein folding [39]. One way to provide a realistic comparison of folding-unfolding state between mutants is through ProtSA, a web application that calculates sequence-specific solvent-accessibilities and the influence of point mutations on the protein stability [40]. Mutant T52Y and G55Y shared a number of 17 residues (which includes active site, Asp-255) identified to be within 45–90% of unfolded SASA whereas majority of residues were located within 90–110% of unfolded SASA. Mutant T52Y has a higher mean unfolded residue indicated by higher value in SASA as compared to G55Y (Fig. S1 and Table S1). Based on this output, ProtSA depicts that both mutated region of lid 1 appeared to gain a large percentage of solvent exposure upon mutation and was expected to destabilize the folded conformation of lid 1 when interacted with hydrophobic organic solvent. Hence, this data explained the actual indications of atomic movement involved residues from lid 1 and its surrounding during the molecular dynamics simulation in toluene.

3.4. Kinetic Outputs of Lid 1 Mutations Based on Small-, Medium- and Long-Substrate Length

Activation of lipase involves the displacement of the lid to expose the active site, suggesting that the dynamics of the lid could be of mechanistic and kinetic importance [41]. For recombinant AMS8 lipase, the positive changes in $K_m$ had not resulted in any significant improvements to the $k_{cat}$ and $k_{cat}/K_m$ (Table 4). The $k_{cat}/K_m$ in AMS8 lipase was meagrely raised in 0.5% (v/v) toluene in the presence of all three pNP substrates. G55Y was found to share similar preference with recombinant AMS8 in hydrolysing shorter pNP esters. Both recombinant and mutant G55Y lipases scored high $k_{cat}$ and $k_{cat}/K_m$ value following the low $K_m$ observed for pNPC hydrolysis in 0.5% (v/v) toluene as shown in Table 4. Unlike these two lipases, mutant T52Y favours a longer chain carbon ester (pNPP) without the presence of toluene and shorter pNPC in 0.5% (v/v) toluene. The reason behind this change was probably due to the lowest total number of hydrogen bonds attained.
by mutant T52Y in water (9H-bonds) compared to recombinant AMS8 (12H-bonds) and G55Y (11H-bonds). The loss of hydrogen bond in T52Y makes the region slightly flexible and allows longer chain substrate to bind. The numbers of H-bond at lid 1 (highlighting residues 49–59) were all reduced in the presence of toluene as observed in the final simulation trajectories. This might be one of the reason in recombinant AMS8 to have improvements in $K_m$ values at 0.5% (v/v) toluene but not in 25 °C and 37 °C.

Albeit the improvement of mutant T52Y on substrate (pNPP) selectivity, the overall lipase activity and stability in 0.5% (v/v) toluene remained low in comparison to the recombinant. This suggests that mutation on lid 1 brings minor alteration in chain-length specificity and major changes towards molecular adaptations to organic solvent stability. Other similar cases involved substitutions of lid residues includes cold-adapted _Pseudomonas fragi_ lipase (PFL). This study targets T137 and T138 located at lid domain to valine and asparagine, leading to successfully modified lipase chain-length preference profile by increasing the relative activity towards C-8 substrates [42]. A thermostable _Rhizopus chinensis_ lipase (RCL) showed that by introducing a disulfide bond in the hinge of lid domain between F95C and F214C, the mutant RCLCYS had improved the $K_m$ value of pNPP at 1.5 fold higher. However, RCLCYS preferred pNPA based on $k_{cat}$/$K_m$ [29]. This mutation however did not convene to organic solvent adaptation properties.

The limitation of enzymatic catalysis in toluene could also be due to the implications of H-bond because the H-bonding process continuously competes with the presence of bulk water. This mechanistic insight suggested indiscriminate strengthening of H-bonds correlates poorly with experimental binding affinity [43]. Hence, an increase in H-bonds of lid 1 could interfere the binding of substrate due to the intensifying stability. Earlier in a closed-conformation of recombinant AMS8 lipase, there were a total of 17H-bond interactions found between residues 49 to 59 as shown in Table 2. After 20 ns simulation in toluene, the lipase exhibited reductions in the total number of H-bonds at similar area at 47% (Fig. 5A) When temperature was increased from 25 to 37 °C, the number of H-bonds lowered to 8 (53% reductions) where new H-bond interactions Ala-51-Leu-49, Leu-57-Val-54, Leu-58-Gly-55 and Ala-56-Leu-58 were formed (Fig. 5B). Other interactions were found preserved with increased in temperature. Based on this structures, lid 1 appeared rigid before simulations in toluene because of the interactions of H-bond between Thr-52 and Pro-68. The H-bond between Thr-52 and Pro-68 are found to be intact in lipase MIS38 and AMS8. Unlike in T52Y and G55Y lipase structures, this H-bond disappeared and left the coil structure to freely circumnavigate the lid domain. In T52Y, the hydrogen-bond interactions were reduced by 40% after underwent 20 ns of simulations in toluene at 25 °C. A few new H-bond interactions that appeared after 20 ns of simulation includes Gly-55-Tyr-52 and Gly-59-Ala-56 (Fig. 5C). However, the H-bond networks changed at 37 °C when new bonds connecting Ala-51-Leu-49, Ala-56-Tyr-52, Gly-59-Gly-55 and Gly-59-Gly-55 were formed (Fig. 5D). This results indicated that mutation at Tyr-52 had a direct influence on the formation of H-bond at lid 1 through decline in its number from 10 to 6 at 25 °C. At 37 °C, the number of H-bonds were increased to strengthen alpha-helix structure. Both recombinant AMS8 and T52Y lipases exhibited higher flexibility at this area (Fig. 2A and B) due to the reorientation of the coil part which occurred in residues Gln-64 to Ile-70. Mutant G55Y exhibited growing numbers of hydrogen bonds between the residues 49 to 59 from 25 °C to 37 °C in toluene. In G55Y lipase, Tyr-55 formed hydrogen bonds with residues, Leu-58 and Gly-59 at 25 °C (Fig. 5E) whereas two more H-bonds formed with Thr-52 and Ser-60 at 37 °C which contributed to increase in rigidity of lid 1 structure (Fig. 5F). The coil and turns which creates the boundary of lid 1 in G55Y was pointing downwards showing lack of structure differences when being simulated at 37 °C. Compared to all mutant lipases, lid 1 from recombinant AMS8 lipase had the highest number of H-bonds that affect the degree of flexibility. By introducing hydrogen bond via tyrosine (a classic hydrogen bond donor) at residue 52 and 55 and subsequently interrupting other H-bonds at its area of vicinity, lid 2 flexibility comes to constraint in toluene. In this study, both alpha-helix lids from cold-active AMS8 lipase ensued a “see-saw” approach, showing its response to balance structural stability and flexibility. Based on this concept, there would hardly be a dual movement on two lids happened at the same time.

3.5. Biochemical and Biophysical Changes Following Lid 1 Mutations

In most lipases, reduced activity in organic solvent (in this case, toluene) was not always caused by protein denaturation due to organic solvent effects. It is more likely to happen because of the difference in half-life of enzyme in organic solvent [44]. The activating effects shown by recombinant AMS8 lipase in 0.5% (v/v) toluene verified that this cold-loving enzyme resist the presence of small amount of toluene which improves the $K_m$ of pNP carbon-chain 8, 12 and 16 hydrolysis. The fact that toluene enhanced substrate binding defined the possibility on structural adaptation towards the presence of organic solvent during hydrolysis reactions. Because of toluene, this substrate finds its way to
efficient binding with catalytic sites improving $k_{cat}$ and $k_{cat}/k_{m}$ of recombinant AMS8 lipase. Recombinant AMS8 lipase was found to be highly stable and active at optimal temperature of 45 °C as shown in Table 5.

Other mutants featured on lid area such as T52Y shared optimal temperature with G55Y via significant reduction of 10 °C temperature. Half-life of T52Y was longer at 25 °C (> 10 h) without toluene but shorter in 0.5% (v/v) toluene (8 h). When temperature was increased to 37 °C, the half-life of mutant T52Y declined but remained active at half its activity up to 9 h without toluene and 5 h with 0.5% (v/v) toluene. The low substrate affinity (higher $K_m$) of T52Y in 0.5% (v/v) toluene in pNP-palmitate could share similar trends to its affinity in a long-chain fatty acid substrate, olive oil at 25 °C because of its longer half-life. The half-life trend at 0.5% (v/v) toluene however was shorter in G55Y. At 25 °C, mutant G55Y achieved longer-half-life in 0.5% (v/v) toluene but shorter in toluene-free hydrolysis reaction. As temperature increased, G55Y half-life dropped at 1.5 h in which toluene did not set to improve its half-life at 37 °C similar to the rest of lipases. Apparently, thermal stability of T52Y and G55Y was very much declined by the reducing half-life in 0.5% toluene as compared to parent lipase. In other study, lid mutants of cold-adapted *Pseudomonas fragi* lipase (PFL) exhibited thermal stability following the time-dependent loss of activity at 29 °C, a temperature where subtle changes in stability can be observed [42]. Parent lipase (PFL) on tricaprylin was found reduced by 50% after 4 h incubation, whereas, under the same conditions, mutant T137 V still retained 90% of its activity and mutant T138 N remained active at 70%.

Given a range of toluene concentrations (0–10%, v/v), most lipases exhibited tolerance at 1–3% (v/v) toluene (Fig. S2). Because all lipases expressed high stability and activity at low concentration of toluene, a decline on conformational stability would be expected at any concentration of toluene exceeding 3% (v/v). This might be due to the relatively high viscosity of the solvents, which hindered efficient interaction between the enzymes and substrates [44]. Based on widely reported scientific evidence, toluene is considered highly toxic to most microorganisms and enzymes secreted by them even though being exposed at lowest toluene concentration of 0.1% (v/v) [45].

Aggregation and precipitation (Fig. S3) of protein mostly occurred due to the exposure of protein in incompatible solvent [46]. In all lipases studied, denaturation precedes aggregation in toluene showing that aggregate protein may also exhibited partially denatured state. Without toluene, the aggregation points appeared before denaturation. Both circumstances placed recombinant AMS8 and lid 1 mutant lipases at different state of folding/unfolding. Through this, toluene was depicted as a driving-force in supporting the progression of protein folding and minimizing structure instability when it comes to high temperature. For lid 1 mutants, the outcome of melting temperature suggested different verdicts on the progression of protein denaturation in the presence of toluene. Mutant G55Y showed to exhibit melting point ($T_m$) of 60.01 °C whereas T52Y $T_m$ was slightly lower at 57.38 °C. In toluene, both

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**Table 4**

Michaelis Menten kinetics measuring $K_m$ and $k_{cat}$ of mutants T52Y and G55Y. Three types of pNP substrates were used in kinetic study at 25 °C, stated as A, B and C, where A signifies pNP palmitate (C-16), B for pNP laurate (C-12) and C for pNP caprylate (C-8).

| WT AMS8 | T52Y | G55Y |
|---------|------|------|
|         | 0% toluene | 0.5% toluene | 0% toluene | 0.5% toluene | 0% toluene | 0.5% toluene |
| $K_m$ (mM) | | | | | | |
| A: 2.59 ± 2.7 | A: 0.79 ± 0.1 | A: 32.47 ± 5.1 | A: 0.79 ± 0.1 | A: 45.92 ± 6.1 | A: 0.79 ± 0.1 | A: 7.11 ± 6.4 |
| B: 1.85 ± 2.9 | B: 0.69 ± 0.3 | B: 48.17 ± 0.0 | B: 0.69 ± 0.3 | B: 45.45 ± 9.3 | B: 0.69 ± 0.3 | B: 15.58 ± 19.3 |
| C: 0.73 ± 2.0 | C: 0.63 ± 0.7 | C: 32.47 ± 5.1 | C: 0.63 ± 0.7 | C: 45.45 ± 9.3 | C: 0.63 ± 0.7 | C: 7.19 ± 4.9 |

| $k_{cat}$ (s$^{-1}$) | $V_{max}$/[ET] | $k_{cat}/K$ (mM$^{-1}$ s$^{-1}$) |
|---------------------|----------------|-------------------------------|
| A: 2.16 × 10$^{-7}$ | A: 5.02 × 10$^{-6}$ | A: 6.92 × 10$^{-6}$ |
| B: 1.86 × 10$^{-7}$ | B: 5.13 × 10$^{-6}$ | B: 7.44 × 10$^{-6}$ |
| C: 2.87 × 10$^{-7}$ | C: 1.97 × 10$^{-5}$ | C: 1.56 × 10$^{-4}$ |

| A: 8.37 × 10$^{-8}$ | A: 6.92 × 10$^{-6}$ |
| B: 2.09 × 10$^{-7}$ | B: 3.17 × 10$^{-7}$ |
| C: 3.69 × 10$^{-8}$ | C: 1.08 × 10$^{-6}$ |

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**Fig. 4.** Comparing one from five modes of distance fluctuation map of recombinant AMS8 (A), mutants T52Y (B) and G55Y (C) between all pairs of Cα atoms at the end simulation trajectories in toluene (25 °C). Distance increases of flexible blocks appeared in blue whereas decreases in distance of flexible regions are spotted in red. This matrix displays the maximum distance fluctuations attributed by the strongest 10% of the residue pair distance changes. Every pixel corresponds to a single residue. Grey lines are drawn for every 10 residues- and yellow lines for every 100 residues (counting from the upper left corner). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Fig. 5. Hydrogen bond interactions at Thr-52 and Gly-55 in recombinant AMS8 (A, B), T52Y (C, D) and G55Y (E, F) in toluene. Lid 1 structures (highlighting residues 46–70) showed the structure at 25 °C (A, C and E) and 37 °C (B, D and F) simulation in toluene. The helix structure of lid 1 was presented in red colored ribbon. The pointing arrow indicates the direction in hinge movements coordinated by turns (shown in green ribbon) and coil structure (grey ribbon). H-bonds are shown in green-lines connecting residues at 49–59, revealing the total numbers of H-bond interactions within this area. In the list of H-bond interactions, the mutated residues (with H-bond) are highlighted in red font and new H-bond interactions (which exclusively appeared in either 25 or 37 °C) are differentiated by green font. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
mutants $T_m$ dropped to 46.5 °C showing their higher tendency of protein denaturation. A decline in denaturation point of T52Y and G55Y mutants at 2% (v/v) toluene illustrates the weakening of hydrophobic interactions among buried residues. As a consequence of this situation where more buried peptide groups and side chains are exposed to solvent, the tightly packed non-polar side chain and peptide groups are getting reduced in number. Losing the intramolecular hydrogen bonds required for buried residues exposes the packing of close-packed spheres and groups in the interior of a protein to protein unfolding or denaturation [47]. The denatured states of cold-active protein can be understood from the increased formation of randomly coiled structure. Thus, it is clear that two important conditions must be met by lipase to co-exist in a solvent, in which firstly, the native state of protein must be favored and second, protein must be active and soluble in its optimum concentration of solvent.

3.6. Thermodynamic Profiles of Lid 1 Mutants in Toluene

The difference of energy states for converting reactant into product can be determined by using $\Delta G$, $\Delta H$ and $\Delta S$. Gibbs free energy describes the capacity of a thermodynamic system to do maximum or reversible work at a constant temperature and pressure. It is said that negative $\Delta G$ is obtained when the system reaches an equilibrium state at constant pressure and temperature, but a fast drop in $\Delta G$ will also indicates a sign of protein denaturation [48]. In this study, all lipases generally exhibited decline in $\Delta G$ after temperature was switched from 25 °C to higher temperature (35 °C) in the presence of 0.5% (v/v) toluene (Table 6). Through the negative changes of free energy, these lid 1 mutants and parent AMS8 lipases were thermodynamically driven and act as biocatalyst under a spontaneous reaction to form product [49] as $\Delta G$ can tell the difference of energy between reactants and products. Compared to recombinant AMS8, all mutants exhibited lower $\Delta G$ values in toluene and without 0.5% (v/v) toluene at both temperatures. This finding suggested that lid 1 mutants could display higher catalytic rate in response to the rise in temperature regardless in which system. Subsequently, $\Delta G$ values can be use in determining the stability of protein-substrate complex in toluene based on the enthalpy-entropy balance. The cold-adapted enzymes invariably show a more negative entropy and a lower enthalpy of activation than their mesophilic orthologs [50].

Enthalpy is a measure of total energy of a thermodynamic system in which higher enthalpy shows lower catalytic rates at respective temperature [48]. Most of lipases exhibited decline in $\Delta H$ when measured from 25 °C to 35 °C with and without 0.5% (v/v) toluene. Low $\Delta H$ value in mutant G55Y indicates that this mutant has higher catalytic rates for $p$NPC hydrolysis in toluene. Since $k_{cat}$ was measured based on its reaction to smaller-chain substrate ($p$NPC) in toluene, this result confirmed that the enthalpy changes not only caused by differences in temperature but also due to its association with substrate specificity. In reaction without toluene, recombinant AMS8 was found to exhibit higher catalytic rate due to low $\Delta H$. In this study, recombinant AMS8 and T52Y lipases were found to have catalytic inefficiencies when used for catalysis in toluene and this was shown via the decline in $k_{cat}$. Nonetheless, the activity improved when the lipases were used in toluene-free reaction.

Entropy is a measure of distributed heat energy over the thermodynamic system. Heat always flows from regions of higher temperature to regions of lower temperature with its positive and negative signs indicating the overall increase and decrease in degree of the freedom of the system, respectively [48]. The increase in negative entropy will or might lead to further instability of protein creating a softer structure. This corresponds to the entropic effects commonly found in enzymes that are adapted by evolution to work at low temperatures [50]. Mutant T52Y was found to have a slight increase of $\Delta S$ value from 25 °C to 35 °C in 0.5% (v/v) toluene indicating lower degree of structure disorder whereas the other lipases did not show changes in structure disorder at these temperatures. On this basis, mutant T52Y was found to be
Biophysics and biochemical properties of mutants T52Y and G55Y.

Table 5

| 1. Changes of secondary structure following increase in temperature 20–30 °C | Recombinant AMS8 | TS2Y | G55Y |
|---|---|---|---|
| 2. Optimal temperature (°C) | 45 (± 0.04) | 35 (± 0.01) | 35 (± 0.06) |
| 3. Half-life at 25 °C (min) a) without toluene | 300 (5 h) | 300 (5 h) | 300 (5 h) |
| b) with 0.5% (v/v) toluene | 480 (8 h) | 480 (8 h) | 480 (8 h) |
| 4. Half-life at 37 °C (min) a) without toluene | 300 (5 h) | 300 (5 h) | 300 (5 h) |
| b) with 0.5% (v/v) toluene | 480 (8 h) | 480 (8 h) | 480 (8 h) |
| 5. Stability in toluene (v/v, %) at 25 °C (range of concentration 0–10 v/v, %) | 1 | 3 | 3 |
| 6. Aggregation point, Tagg (°C) by DLS a) Without toluene | 35 | 40 | 50 |
| b) 0.5% (v/v) toluene | 62.5 | 50 | 50 |
| 7. Denaturation point, Tdn (°C) by CD a) Without toluene | 45.8 | 57.38 | 60.01 |
| b) 2% (v/v) toluene | 53.5 | 46.5 | 46.5 |

Notes: Differences in toluene concentration measured for Tagg and Tdn in DLS and CD are based on the allowed detection limit in respective equipment. Ideal concentration of non-polar solvent in CD was identified at 0.5–5% (v/v) and 0.5–1% (v/v) in dynamic light scatterings (DLS) when performed under gradually increased temperature 20–80 °C. All assays were performed in 3-replicates (± standard deviation).

Thermodynamically stable in the presence of toluene. As for enthalpy-entropy equilibrium in toluene, TS2Y exhibited high enthalpy with increasing entropy. This result was an indication of a functionally compromised (reducing rate of activity) but stable lipase. In a normal system (a reaction without toluene), G55Y showed the highest enthalpy of activation, ΔH a sign where more energy is required for the products to form (substrate turnover). The high enthalpy value was followed with increase in entropy, ΔS that further intensify protein stability. As a consequence of these analyses, it was found that temperature is a good precursor to determine the stability of AMS8 lipase and lid 1 mutants as well as the extent of their activity. In this study, toluene facilitates stability and activity of lid 1 mutant TS2Y through its improvement from precursor to determine the stability of AMS8 lipase and lid 1 mutants.

Table 6

Thermodynamics parameters of recombinant AMS8 and lid 1 mutant lipases, TS2Y and G55Y with and without 0.5% (v/v) toluene. All thermodynamic values were derived based on the independent calculations of kcat on temperatures (25° and 35 °C) by using eqs. 2, 3 and 4.

| Temperature (°C) | Gibbs Free Energy of Activation Energy, ΔG (kJ/mol) | Enthalpy, ΔH (kJ/mol) | Entropy, ΔS (kJ/mol) |
|---|---|---|---|
| | 0.5% (v/v) Toluene | No toluene | 0.5% (v/v) Toluene | No toluene | 0.5% (v/v) Toluene | No toluene |
| 25 | | | | |
| WT | −113.98 | −119.62 | 101.74 | −144.21 | 0.72 | −0.08 |
| TS2Y | −114.63 | −120.72 | 312.23 | −39.31 | 1.43 | 0.27 |
| G55Y | −121.19 | −118.26 | −50.20 | 112.63 | 0.24 | 0.77 |
| 35 | | | | |
| WT | −121.22 | −118.78 | 101.65 | −144.29 | 0.72 | −0.08 |
| TS2Y | −131.56 | −121.58 | −39.40 | 112.55 | 1.53 | 0.27 |
| G55Y | −123.55 | −126.00 | −50.29 | 112.55 | 0.24 | 0.77 |
Table 7
Estimated activation energy in kJ/mol derived from aforementioned Eq. 1. Formula used to calculate was slope = − Ea/R where R is a constant equal to 8.314 J/molK−1 using temperature 25 °C to 35 °C.

| Recombinant | 0.5% (v/v) toluene | Without toluene |
|-------------|-------------------|-----------------|
| T52Y        | 104               | −142            |
| G55Y        | 314               | −36.834         |
|             | −47               | 115.11          |

difficult to bind to the active site which results in a drop of reaction constant (kcat).

4. Conclusion

The effects of mutations at residue Thr-52 and Gly-55 of AMS8 lipase showed a slight movement at lid 1 region and minimal disruption on lid 2 activation. Mutation that targets T52Y and G55Y on lid 1 confirmed on the importance of this region to temperature stability according to the results of biophysical analysis and molecular dynamics simulations. Analysis on the altered optimal temperature, melting temperature, half-life and toluene stability clearly described the dependency of AMS8 lipase structure stability on the rigidity of lid 1. As an added advantage to these work, mutant T52Y showed preference to the long carbon-chain substrate or, pNP-16 via Michaelis-Menten constant (Km) which seems to support hydrolysis without toluene at 25 °C. This mutant however continued to fancy hydrolysis of smaller-chain pNP (pNPC) in 0.5% (v/v) toluene. In 0.5% (v/v) toluene, catalytic efficiency of G55Y is higher than T52Y in 2 out of 3 substrates tested. Mutants T52Y and G55Y unfold faster in 2% toluene than without solvent. Analysis on protein denaturation profiles was found not to be correlated with increase in toluene concentrations. Both enthalpy and entropy exhibited decline in value indicated high catalytic activity and higher degree of structure loss when toluene was added. All lipases exhibited decline in Gibbs free energy and ΔH‡ when measured from 25 °C to 35 °C in 0.5% (v/v) toluene. There was least change occurred to ΔS‡ between these temperatures. Therefore, pNPC hydrolysis in mutant lid 1 lipases showed to have higher catalytic rate in toluene between 25 and 35 °C. Instead of temperature, the structure loss was very much dependent on the toluene concentration. The enzyme will become a highly relevant industrial biocatalyst if it is able to utilize broad catalysis on the toluene concentration. The enzyme will become a highly relevant industrial biocatalyst if it is able to utilize broad catalysis on the toluene concentration.

Author Contributions

All authors reviewed the manuscript. N.Y. and M.S.M.A designed the experiments. N.Y. conducted all experiments. N.Y. analysed all data, performed data curation and wrote and revised the manuscript. A.T.C.L contributed to the software for molecular dynamics simulation (YASARA version 11.3.22). N.H.A.K., A.B·S., M.S.M.A were the Principal Investigators of Enzyme and Microbial Technology Research Centre whom contributed equally in research discussion, instrumentation, purchasing of reagents and chemicals. All authors read and approved the final manuscript.

Funding

This research was funded by Putra Graduate Initiative (IPS), Universiti Putra Malaysia with grant number 9536300. N.Y. was a recipient of Graduate Research Fellowship (GRF) scholarship awarded by School of Graduate Studies, Universiti Putra Malaysia.

Conflicts of Interest

There was no conflict of interest. The funder had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision to publish the results.

Appendix A. Supplementary data

Supplementary materials: Fig. S1: The relation (in percentage) between the unfolded SASA of the individual residues of T52Y (A) and G55Y (B) and that of the average unfolded SASA for that residue type for a test set of 19 other proteins, Fig. S2: Toluene stability analysis based on toluene concentration (0, 1, 3, 5, 7, 10%), Fig. S3: Aggregation point (or tendency to oligomerize/aggregate) found in purified protein of recombinant AMS8 (A, B), T52Y (C, D), and G55Y (E, F) with and without 0.5% (v/v) toluene, Fig. S4: Arrhenius plot with calculated ln kcat and 1/T (K−1) to estimate the value of activation energy (Ea) of the system, Table S1: Mean and total unfolded residue of SASA obtained at Thr-52 and Gly-55 from recombinant (WT) AMS8, Tyr-52 from mutant (Mut T52Y) and Tyr-55 from mutant (Mut G55Y). Supplementary data associated with this article can be found in the online version, at doi:https://doi.org/10.1016/j.csbj.2019.01.005.

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