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Crystallization of N-terminal Strep-tagged Fusion Lipase from Thermostable *Bacillus* sp. Strain 42. Mahiran Basri^a, Raja Noor Zaliha^b, Raja Abd Rahman^b, Tengku Haziya Amin Tengku Abd. Hamid^b, Abu Bakar Salleh^c. ^a*Department of Chemistry, Faculty of Science.* ^b*Department of Microbiology,* ^c*Department of Biochemistry, Faculty of Biotechnology and Biomolecular Sciences, University Putra Malaysia 43400 Serdang, Selangor, Malaysia.*

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Lipases have great potential to be used in industries due to their favourable properties such as substrate specific, enantiomerically selective, regioselective and mild reactions conditions. Lipases of microbial origin are generally more stable than lipases from animal or plant and as such they are useful source for industrial enzymes. A 1.2 kb lipase gene (AY 78735) [1], isolated from solvent stable and thermostable *Bacillus* sp. strain 42 was overexpressed using pET51b vector with *E. coli* host strain BL21(DE3) pLysS, in which the fusion lipase contains N-terminal Strep-tag II affinity tag [2]. The purified fusion lipase, at protein concentration of about 4.0 mg/mL, was induced to crystallize in 0.1 M MES buffer at pH 6.5 without the presence of salt, but in the presence of only 12% w/v PEG 20 000 as precipitant. Crystallization reactions were carried out using vapour diffusion methods at 16°C. Crystals were formed after 12 hours incubation. The crystals with size measuring around 0.04 X 0.12 mm were shown to be heavily stained with protein dyes. Lip 42 lipase is highly homologous to three crystallized lipases from thermophilic *Bacillus* sp., namely T1 lipase [3], P1 lipase [4] and L1 lipase [5]. Lip 42 protein crystals, despite having almost 97% similar homology in amino acid sequence, showed a different shape and crystallization condition. The shape of Lip 42 crystal appeared to be partly attributed to the presence of N-terminal tag.

[1]. Eltaweel, M. A., Rahman, R. N. Z., Salleh, A. B. and Basri, M., 2005, *Ann. Microbiol.*, 55:187-192. [2]. Rahman, R.N.Z., Hamid, T. H. T. A., Eltaweel, M. A., Basri, M. and Salleh, A.B., 2008, *J. Biotechnol.*, 136S : S290-S344. [3]. Leow T. C., Rahman, R. N. Z., Salleh, A. B. and Basri, M., 2007, *Cryst. Growth Des.*, 7: 2406-2410. [4]. Sinchaikul, S., Tyndall, J. D. A., Fothergill-Gilmore, L. A., Taylor, P., Phutrakul, S., Chen, S. T., Walkinshaw, M. D., 2002, *Acta Crystallogr. D.*, 58: 182-185. [5]. Jeong, S.-T., Kim, H.-K., Kim, S.-J., Pan, J.-G., Oh, T.-K. and Ryu, S.-E., 2001, *Acta Cryst. D*57, 1300-1302

Keywords : thermostable lipase; strep-tag fusion; lipase crystal

FA1-MS09-P02

Crystallization of Mutated T1 Lipase from Thermostable *Geobacillus Zalihae* Strain T1. Raja Noor Zaliha^a, Raja Abd Rahman^a, Adam Thean Chor Leow^b, Abu Bakar Salleh^c, Mahiran Basri^d. ^a*Department of Microbiology.* ^b*Department of Cell and Molecular Biology.* ^c*Department of Biochemistry,*

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A novel thermoalkaliphilic lipase producer *Geobacillus zalihae* strain T1 was isolated from palm oil mill effluent in Malaysia [1]. The mature T1 lipase was overexpressed in *Escherichia coli* harboring pGEX/T1S recombinant plasmids with lipase activity of 42 U/ml [2]. The enzyme can be crystallized up to 60°C [3]. Optimization process revealed that a balance of hydrophobic interaction, packing rate, and some flexibility was needed to obtain a good crystal solved at 1.5Å [4]. Point mutation D311E (inter-loops networking) created an additional one salt-bridge and two hydrogen bonds as compared to K344R (intra-loops networking) with additional two salt-bridges and one hydrogen bond. Denatured protein analysis revealed that mutation D311E gave higher T_m (70.59 °C) as compared to K344R (68.54 °C) and native T1 lipase (68.52 °C). The mutant D311E was able to form preliminary crystal interface with formulation 9, 13 and 21 of Crystal Screen 2 with 2 M NaCl, 30% PEG-MME-2000 and 2 M NaCl as precipitants, respectively.

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Keywords: lipase; rational design; protein crystallization

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Hydrophilic Polymers as Efficient Protein Surface Shielding Agents in Protein Crystallization. Jindřich Hašek^a, Tereza Skálová^a, Jarmila Dušková^a, Jan Dohnálek^a, Petr Kolenko^a, Tomáš Koval^a, Andrea Štěpánková^a. ^a*Department of molecular structure, Institute of Macromolecular Chemistry AV CR, Praha 6, Czech Republic.*

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The paper introduces a new concept of “protein surface shielding agents” into protein crystallization. This concept follows from “analytical crystallography”, i.e. analysis of processes leading to formation of crystals and the analysis of intermolecular contacts observed in the crystalline state “crystal contact areas” (CCA) [1]. It shows that large molecules have many various modes of mutual adhesion but only some of these “adhesion modes” (AM) are suitable for compact stacking of macromolecules into the crystal lattice. It shows also that different adhesion modes lead to crystals of different diffraction quality, and to different space groups, and some adhesion modes are

not compatible with any stable crystalline state. A helpful term in description of the controlled crystallization process is “*dominating adhesion mode*” (*DAM*), i.e. the most probable adhesion realized between two macromolecules under the given conditions, usually but not necessary, the highest affinity adhesion mode. Preferences of adhesion modes can be regulated e.g. by a selective mutation of protein surface but also much more easily by changing pH, buffer composition, anions, additives, precipitants etc. Any chemicals adhering temporarily to the specific places on the protein surface and blocking efficiently unwanted adhesion modes between proteins are called “*protein surface shielding agents*” (*PSSA*). Because of complex effects of components used in the crystallization solution, many different low-efficiency PSSA's have already been intuitively used long time in crystallization screens under the concepts of additives, precipitants, anions. In the case that DAM leads to formation of a single crystal we receive a unique *group of crystal contact areas* (*GCCA*) related directly to the specific crystal properties (space group, water contents, diffraction quality, twinning, etc.). It is evident that well diffracting crystals are formed only when a single adhesion mode dominates but a reverse statement need not be true. Because of the possible strain induced by deposition of macromolecules in the growing molecular cluster, the contact areas realized in the crystal are not necessarily identical with the contact areas in solution. However, the contact areas of the dominating adhesion mode should be very similar in both states. Hydrophilic polymers are generally very efficient PSSAs, because they protect a much larger area on protein surface than their low-molecular weight equivalents upon binding to protein surface. Thus the polymers have generally better propensity to eliminate unwanted adhesion modes between protein molecules forming a crystal.

Practical part: We made an experimental scan of an efficiency of a number of water soluble polymers and co-polymers as novel *protein surface shielding agents* in the crystallization process. Sixteen novel water soluble polymers were tested for their efficiency in protein crystallization. Six of them were selected as precipitants to two new polymer crystallization screens (POLYA, POLYB) /2/. The screens were independently tested in six laboratories on 30 different globular proteins. Supported by GA AV IAA500500701.

[1] Hašek J., *Zeitschrift für Kristallogr.*, **2006**, 23, 613.

Keywords: polymers; co-polymers; protein crystallization; protein surface shielding agents; molecular adhesion

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Expression, Purification and Crystallization of Human Bile Acid-coA: Amino Acid N-Acyltransferase (BAAT). Laura Civiero^a, Stefano Capaldi^a, Massimiliano Perduca^a, Hugo Luis Monaco^a. ^a*Biocrystallography Lab, Department of Biotechnology, Verona, Italy.*
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Bile acid-coenzyme A:Amino Acid N-Acyltransferase

(BAAT) is the sole enzyme responsible for conjugation of primary and secondary bile acids to taurine and glycine [1].

The human cDNA BAAT sequence was cloned into the pET15 vector and its expression was tested in several *E.coli* strains. Because of the toxicity associated with the overexpression of BAAT in several strains, BL21 (DE3) C41 was selected to express the recombinant protein. Only with this specific strain, a reasonable amount of the protein was obtained in the soluble fraction.

The purification steps involved Nickel-sepharose chromatography followed by reverse IMAC after histidine-tag cleavage. Size-exclusion and hydrophobic interaction chromatography steps were also included to increase the purity of the sample. After these steps the protein shows as a single band in both SDS and native PAGE, The purified enzyme was assessed for activity and was found to be able to catalyze the conjugation of taurine to CoA-fatty acids [2].

The presence of non physiological inter- and/or intramolecular disulfide bounds between the three cysteine residues present in the protein sequence as a source of sample microheterogeneity was also examined. Three cysteine mutants (C235A, C372A and C373A, and the triple mutant without cysteines) were constructed and tested following the same protocols used for the wild type protein and no significant differences in the behaviour of the four species were found.

Crystallization trials with the apo- and holo-wild type enzyme, as well as with the three mutants, are in progress but up to now only microcrystals not adequate for X-ray diffraction studies have been obtained.

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Novel Crystallisation Plate Imaging System Using ANS-Based Fluorescence at the EMBL Hamburg High-Throughput Crystallisation Facility. David Watts^a, Victor Lamzin^a, Jochen Muller- Dieckmann^a, Matthew Groves^a. ^a*EMBL, c/o DESY, Building 25a, Notkestrasse 85, Hamburg, 22603 Germany.*

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We have previously demonstrated that the addition of low concentrations (μM - nM) of a nonspecific fluorescent dye (1,8-ANS) does not significantly hinder crystal growth in a set of trial proteins [1]. The resulting fluorescence images collected on a high-resolution fluorescence microscope show that these protein crystals are strongly contrasted against other common crystallisation drop phenomena, such as precipitate and phase separation. However, the experiments were performed on a fluorescence microscope and no specialised equipment currently exists for the visualisation of ANS-based fluorescence (excited at 365 nm). Here we present a novel and cost-effective device for visualisation