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Physicoenzymatic Production of Monoacylglycerols Enriched with Very Long Chain Polyunsaturated Fatty Acids

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Running title: Production of PUFA monoacylglycerols

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ABSTRACT

Background: Monoacylglycerols (MAG) containing polyunsaturated fatty acids (PUFA), especially, eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA), have interesting applications. The enzymatic processing of such MAG directly from fish oils is highly interesting integrating the processing of MAG and concentration of EPA and DHA. The aim of this study was then to develop an efficient enzymatic glycerolysis system together with physical fractionation for the production of PUFA-MAG from tuna oil.

Results: Novozym 435 was eventually selected after evaluation together with the immobilized Lipase AK in tertiary alcohol based system. A further evaluation of solvent mixtures involving tertiary alcohols was made, taking the consideration of operation easiness. It turned out that a number of mixtures gave similar performances as tert-butanol (TB). Basic reaction parameters were thoroughly evaluated. In the batch reaction system with TB as solvent, the recommended conditions were: glycerol/tuna oil 4:1 mol/mol, TB/tuna oil 2:1 wt/wt, 15 wt% Novozym 435, and temperature 40 ºC. Under these conditions, the yield of MAG was up to 90% after 3 h incubation. Crude MAG from the production was fractionated to produce MAG with higher EPA and DHA content. Using acetone as solvent at 0 ºC led to ca. 50% yield of MAG but contained EPA and DHA up to 71% in comparison with ca. 30% in tuna oil.

Conclusion: Potentially practical process steps have been developed for the production of MAG containing high content of EPA and DHA from natural fish oils with high efficiency and simplicity.

Keywords: Monoacylglycerols, glycerolysis, fractionation, polyunsaturated fatty acid (PUFA), Novozym 435, tuna oil
INTRODUCTION

Monoacylglycerols (MAG) or mixtures with diacylglycerols (DAG) account for approximately 75% of the emulsifier production and have applications in different fields.\textsuperscript{1-3} In the food industry, MAG are widely used in bakery products, margarines, dairy products, and confectionary because of their emulsifying, stabilizing, and conditioning properties. They are also important in cosmetic and pharmaceutical industries as drug carriers and for consistency improvements in creams and lotions. Commercial food MAG are manufactured by chemical glycerolysis of fats and oils. High temperature (220-250 °C) and inorganic alkaline catalysts are used to accelerate the reactions. These chemical and physical processes are not suitable for heat-sensitive oils and fats because of potential deleterious effects on nutritional and biological properties.

Lipase-catalyzed glycerolysis of fats and oils at atmospheric pressure and low temperature has attracted interest in both academia and industry as a practical alternative to chemical methods in the production of commercial MAG. Several glycerolysis systems have been investigated with or without organic solvents, with immobilized or non-immobilized enzymes, and in microemulsion or other media.\textsuperscript{3-4}

Glycerolysis system with an immobilized lipase as catalyst is a three-phase system: a hydrophobic oil phase, a hydrophilic glycerol phase, and a solid enzyme phase. Because of the more hydrophilic characteristics of the enzyme, glycerol often binds to enzyme particles so that the access of oil molecules to the enzyme is difficult. The mass transfer of glycerol is also limited. Thus the reaction efficiency is usually low even though the efficiency can be improved through optimization in a narrow range. It is reported that glycerol can be immobilized on silica gel so as to overcome
these problems.\textsuperscript{1,4} The improvement is only minor, however, not to say difficulties in practical operations. Therefore, a solvent medium is actually an important solution to improve the homogeneity of the system.

A single solvent that can dissolve oil and glycerol in a homogeneous system is actually very difficult to find. The hydrocarbon solvents were generally impossible for this purpose. After evaluation, a few alcohols with more than five carbons can be considered since they contain a polar hydroxyl group and a nonpolar carbon chain. However, alcohols are naturally reaction competitors to glycerol, especially the primary alcohols. From the study of Damstrup \textit{et al.}\textsuperscript{2} and Yang \textit{et al.}\textsuperscript{5}, the use of tertiary alcohols is possible and does not involve in reactions with fatty acids, most likely due to the tertiary structure of the alcohols, which exerts strong steric hindrance to the enzyme in the system. Therefore, TB or tert-pentanol (TP) is promising for the glycerolysis system. Higher yield of MAG has been achieved with tertiary alcohols in the glycerolysis system\textsuperscript{2,5}.

Omega-3 PUFA have received much attention in recent years because of the health benefits they offer, including reduced risk of coronary disease, prevention of certain cancers, and improved immune function.\textsuperscript{6,7} Omega-3 PUFA-containing MAG are interesting for many potential uses or applications in food, drug, or cosmetic production. Their MAG forms may offer new possibilities in different applications. We have intended to synthesize MAG from fish oil with higher content of omega-3 PUFA through alcoholysis of fish oil with 1,3-specific lipases.\textsuperscript{8,9} The reaction strategy was mainly to produce 2-MAG since usually more omega-3 PUFA is located at 2-position of fish oil.\textsuperscript{6,8}
One possible approach to obtain a dedicated fraction with different melting points is the application of fractionation. There are several approaches available to fractionate fats and oils including dry fractionation (without solvent), wet fractionation (with solvent), and super-critical fluid fractionation.\textsuperscript{10,11} Using these fractionation processes, lipid fractions with different nutritive properties can be produced since the melting behavior of lipids is strongly related to the number of double bonds, meaning PUFA fractions and their derivatives can have very different melting properties from the rest fractions in the mixture.

Therefore, in this study, we designed the production of MAG rich in omega-3 PUFA (EPA and DHA) into a two-step operation. In the first step, an efficient glycerolysis system should be set up for the enzymatic production of MAG from fish oil. The system with tertiary alcohols was considered for the above consideration, as high yields of MAG can be expected in the system after optimization.\textsuperscript{2,5} Under such a possibility, the second step was targeted to fractionate the MAG containing PUFA. Normally, such MAG have much lower melting points than MAG containing saturated or monounsaturated fatty acids. Therefore, a physical fractionation system was also studied to isolate the omega-3 PUFA containing MAG from other MAG.

**MATERIALS**

Crude tuna oil from Skipjack tuna head, with water content of 4.4\% and free fatty acid content of 0.36\%, was provided by Chotiwat Industrial Co. Ltd. (Hat Yai, Thailand). The oil was prepared from crude tuna oil by a conventional pressing method. The refined oil was achieved through degumming, neutralization, bleaching, and deodorizing. The major fatty acid compositions of the refined oil (wt\%) was as following: C14:0, C16:0, C18:0, C18:1, C18:2, C20:5 and C22:6 (4.2, 30.6, 9.3, 17.3, 2.6, 6.7 and 29.0, respectively). The glycerol was analytical grade with 0.2\% water.
The properties of TB are boiling point 83 °C, melting point 25 °C, relative density (water=1) 0.8, octanol/water partition coefficient (log $P_{o/w}$) 0.4, and with colorless appearance. Commercially immobilized lipase, Novozym 435, from *Candida antarctica* lipase B, was obtained from Novozymes (Bagsvaerd, Denmark) and *Pseudomonas fluorescens* lipase (Lipase AK) was a gift from Amano Pharmaceutical Co. Ltd (Nagoya, Japan). Accurel EP-100, a microporous polypropylene powder (particle size < 400 μm), was a gift from Akzo Nobel Membrana (Obernburg, Germany). All other chemicals and solvents used were of reagent grade or analytical grade.

**METHODS**

**Preparation of the immobilized lipase**

Accurel EP-100 (10 g) was added to 100 mL of buffer (pH 7) containing 100 U/mL Lipase AK and the mixture was stirred with a magnetic bar at 100 rpm for 30 min. Afterward, 100 ml of 0.1M phosphate buffer (pH 7) was added and the suspension was filtered through a Buchner funnel by vacuum. The immobilized enzyme (IM-AK) was washed with 100 mL of 0.1M phosphate buffer to remove the unbound enzyme.

The water content of the immobilized enzyme was adjusted by different methods\(^\text{12}\). The first method was vacuum drying in a desiccator at room temperature for 12 h ($a_w=0.389$). The second method was acetone washing and evaporation ($a_w=0.019$). And the last method was to equilibrate the enzyme over saturated LiCl solution in a desiccator at 25 °C for 16 h ($a_w=0.113$). The water activity ($a_w$) of the prepared enzymes was measured with an Aqualab Water Activity Meter (Decagon Devices, Inc., Washington, USA) at room temperature, as shown in the parenthesis.
Enzymatic glycerolysis of tuna oil

The mixture of 10 g of tuna oil, required amount of glycerol and TB was incubated in a capped 25-mL flask at the designed conditions on a 400 rpm shaker. The reaction was initiated by the addition of lipases. At selected intervals, 0.25 mL of reaction mixture was withdrawn and the lipase was removed by filtration and the solvent was removed by vacuum. All samples were stored at -20 °C before analysis.

Experimental repeatability for batch reactions was conducted through three experiments under the following condition: temperature 45 °C, glycerol/tuna oil molar ratio 4.5:1.0, TB/tuna oil 2.2:1.0 (w/w), 15 wt% lipase (based on oil and glycerol), and no additional water.

Fractionation of PUFA-MAG from reaction mixture of glycerolysis of tuna oil

The product collected after reaction under the optimal conditions was subjected to solvent removal under vacuum and was named as crude MAG. The crude MAG in 0.1 g was dissolved in 30 mL of different solvents or mixtures. Acetone and hexane are commonly cited in the literature and used industrially. Therefore, these two solvents were selected together with one mixture between the two solvent in 50/50 (v/v). The fractionation was conducted under different temperatures. Based on melting points of different MAG fractions, the 10, 4, and 0 °C were selected for evaluation. The fractionation was conducted in a selected solvent and temperature for 3 h. Afterwards the samples were centrifuged at the same temperature for 30 min at 10000 rpm. The supernatant was removed and the solid was washed several times with the same solvent cooled to the same temperature. The liquid parts were collected together. The solvent was removed form both solids and liquids by a vacuum evaporator. The two fractions were then weighed and used for further analysis.
Analysis of acylglycerols by TLC-FID

The components of oil phase were analyzed with a thin-layer chromatography with flame ionization detector (TLC/FID) (IATROSCAN MK5, Iatron Laboratories Inc., Tokyo, Japan) for the content of TAG, 1,2(2,3)-DAG, 1,3-DAG, MAG and free fatty acids (FFA).\textsuperscript{13} The samples diluted in chloroform/methanol (2:1 v/v) were spotted onto the chromarod and developed for 35 min in a mixture of benzene/chloroform/acetic acid (50:20:0.7, v/v/v). After developing and drying, the rods were subjected to scanning with FID. Standards were used to identify the peaks. The peaks areas were normalized and used for evaluation of reactions. Triplicate analysis was conducted and the averages were used.

Analysis of fatty acids compositions

The fatty acid compositions of acylglycerol species were determined by converting into fatty acids methyl esters followed by GC analysis. After evaporating excess solvent of the sample, the mixture was applied to normal silica gel TLC-plate and developed in benzene/chloroform/acetic acid (50:20:0.7, v/v/v). After drying, the MAG band was scraped off and methylated with 0.5%NaOH in methanol (1000 $\mu$L), for 10 min at 60 °C. The methyl esters were extracted with $n$-hexane (300 $\mu$L) for 1 min. The $n$-hexane layer was washed with 200 $\mu$L distilled water and dried over anhydrous sodium sulfate. Analysis was carried out with a Perkin-Elmer Autosystem XL-GC gas chromatograph (Perkin-Elmer Corporation, Norwalk, CT) on a FFFAP column (PERMABOND-FFFAP DF-0.25, 25m×0.25mm i.d., MACHEREY-NAGEL, Germany). The carrier gas was helium at a flow rate of 0.5 mL/min (15 psi) and operated in a split ratio of 50:1. The temperature was started from 150 °C for 0.50 min and increased at the rate of 4 °C/min to 170 °C, followed with the rate of 5 °C/min to 195 °C, and further with the rate of 10 °C/min to and 215 °C and held there for 14
min. Injector and detector temperatures were 250 °C. Response factors were determined using a standard mixture of fatty acid methyl esters. Duplicate analyses were carried out for all samples. The relative standard deviation was less than 4.1% for all results more than 10% and less than 6.6% for results less than 10%.

Statistical analysis

The SPSS program analysis was used for data analysis. Analysis of variance and t-test were used to evaluate the significance and difference of data. Values were considered significant at $P < 0.05$ level.

RESULTS AND DISCUSSION

Selection of lipases for glycerolysis

Enzyme characteristics can have determinant functions for the product development and process development. In recent progress of enzymatic production of MAG in solvent systems, Novozym 435 was recommended for the tertiary solvent system. Kaewthong and H-Kittikun, however, concluded from a solvent screening that IM-AK showed good activity in the system used. To find an appropriate catalyst for the aimed MAG processing, the two immobilized lipases concluded from the above studies were selected for further evaluation, i.e. Novozym 435 from Candida antarctica B lipase (nonspecific) with the hydrolytic activity of 9.8 U/mg immobilized enzyme and IM-AK from Pseudomonas fluorescens lipase (1,3-specific lipase) with the hydrolytic activity of 0.97 U/mg immobilized enzyme. In the TB and hexane media, experiments were conducted at the ratio of organic solvent to tuna oil 2.2:1.0 (w/w), immobilized enzyme 100 U/g (based on total substrates), 4.5:1.0 (mol/mol) glycerol/tuna oil, 45 °C, and reaction time of 8 h. The results showed that the reaction by both enzymes in hexane was slow with very low TAG conversion (less than 20%), while in TB much more MAG were formed (data not shown). In TB,
Novozym 435 showed the highest activity with 90% yield of MAG, while IM-AK gave 70% conversion but with 21% FFA (Fig. 1). High FFA content is a problem for industrial applications. It obviously came from the higher water content in the immobilized lipase. We therefore studied the pretreatment of the IM-AK to see if the performance could be further improved because it could be a cheaper alternative to Novozym 435. Therefore, the IM-AK was pre-treated to reduce the water content and the following different water activities (a_w) were obtained as 0.369, 0.113, and 0.019. As seen from Fig. 2, once water content was down, the activity of the immobilized enzyme was decreased as well, meaning the enzyme was water dependent. This implies that the lipase needs higher amount of water to maintain the activity, but such a high amount of water will consequently lead to the stronger hydrolysis reaction so as to form higher amount of FFA. This behavior makes difficult for the use of IM-AK in such reactions where polar solvents are used to exert stronger water partitioning from enzymes. Therefore we conclude that IM-AK is not quite suitable for the reaction system even though quite good reaction conversion can be obtained in high water content situations (Fig. 1). As widely demonstrated and also proved in this study, Novozym 435 has less water dependence and its catalytic activity did not drop even in very polar systems with ethanol.\textsuperscript{2,5,16} After all, Novozym 435 was selected for further process studies, even though it is a costly commercial lipase. With its low water requirement, the process can have high benefit, in which a very low FFA content can be obtained in the products. This is a very important issue for industrial applications since higher FFA content will lead to the loss of oils as well as difficulty in processing.
Evaluations of solvent mixtures for glycerolysis of tuna oil

As demonstrated in a few recent publications,\textsuperscript{2,5} tertiary alcohols are suitable solvents for the efficient glycerolysis system with very short reaction time but high MAG yields. However, TB is solid in room temperature (melting point 25-26°C) so as making the process operation difficult while TP is much more expensive (2-3 fold higher than TB). Therefore, a mixture could be a better choice for practical and cost-effective processes. Therefore the mixtures of the two solvents as well as the mixtures with hexane were evaluated for the reaction system to offer possibilities for different selections. The glycerolysis reaction was carried out in such solvent mixtures and their results are shown in Fig. 3. Tertiary alcohols and their mixtures generally gave higher yields of MAG, even though there were slightly differences between each other. The mixtures of tertiary alcohols with low amount of hexane (20%) also gave reasonably good result, but higher amount of hexane led to lower yields of MAG. Yields of MAG 90-95% were occurred in mixtures of TB/hexane (down to 20% v/v hexane) and TB/TP in various ratios (20:80, 50:50, and 80:20 v/v). This offers a variety of possibilities of solvent selection in practical uses. As seen from the studies of the different mixtures, particularly with tertiary alcohols, the reaction behavior is very similar each other in terms of reaction conversion and enzyme activity (Fig. 3 and data not shown). In practical uses, different decisions can be made depending on the easiness of the process and cost of the solvents as well as other subjective considerations. To simplify the study for the fractionation part, TB was selected for the following experiments since more information has been accumulated in large scale operations concerning TB evaporation procedures and its safety approval from the authority.
Evaluations of other parameters on the MAG yield

Various parameters for the reaction systems have been already evaluated in the early studies. Due to the use of tuna oil where contains high content of DHA and EPA in this study, we had concerns whether the reaction will be seriously affected since the early work used the linoleic acid dominated sunflower oil as materials. Therefore, we still made the evaluation of various parameters. After all, the effects of parameters with the use of tuna oil were very similar to those previous studies (data not shown). Therefore only a general summary is given below. The effects of enzyme loading, amount of solvent, substrate ratio and temperature on the glycerolysis of tuna oil were performed. In the batch reactions, 15 wt% Novozym 435 based on total substrates (glycerol and oil) gave the maximum reaction performance and was used for further reactions. The weight ratio of TB to tuna oil of 2.0:1.0-2.5:1.0 showed high MAG production with no significant difference. Therefore, the weight ratio of TB to tuna oil of 2.0:1.0 was selected. For the effect of glycerol amount, the result showed that the molar substrate ratio of glycerol and oil of 4.0:1.0-4.5:1.0 had no significant difference on MAG production. Therefore, in this study, 4.0:1.0 (mol/mol) glycerol/tuna oil was decided. For the effect of temperature, glycerolysis of tuna oil was carried out at 30 to 50 °C and the results showed that the temperature of 40-50 °C showed high MAG production with no significant difference. Therefore, the temperature of 40 °C was selected for the production since lower temperature was recommended with respect to the product quality.

MAG production under optimal conditions

The recommended conditions for MAG production were finalized as using TB as the medium, the molar ratio of glycerol to tuna oil of 4.0:1.0, the weight ratio of TB to tuna oil with 2.0:1.0, using 15 wt% Novozym 435 (based on glycerol and tuna oil),
and no additional water. The temperature was controlled at 40 °C. Under these conditions, the yield of MAG of 90.8 wt% was obtained after 3 h incubation and the remained TAG was only 5.5 wt%. A time course under such production conditions is also conducted (Fig. 4).

The major fatty acid compositions of the MAG fraction after separating by thin layer chromatography were determined by gas chromatography as follows: C14:0, C16:0, C18:1, C18:2, C20:5 and C22:6 (3.5, 29.7, 8.6, 17.1, 3.6, 6.3 and 30.5 wt%, respectively). The fatty acid compositions had no significant difference from that of the original tuna oil. The result indicated that the reaction gave little fatty acid selectivity for the formation of MAG. This is a reasonable conclusion since the reaction was an interesterification process where positional and fatty acid selectivity of the lipase will place no difference for the product formation.

Fractionation of the reaction mixture from glycerolysis of tuna oil

Temperature fractionation of fats or oils or their derivatives can be regarded as a thermo-mechanical separation process and has been widely used in industry. Individual species (for example TAG or MAG) for a given material are selectively crystallized from the liquid phase at different temperatures. During cooling of the liquid oil or melted material, the species with the highest melting point preferentially crystallized, resulting in solid phase within the system. For natural fats and oils, they are mostly complex mixtures of individual TAG that can contain from one to three different fatty acyl residues on their glycerol backbone. Because of this there is the large variation in the melting points of the TAG species, which complicates the fractionation process. For MAG product, single fatty acid residue is attached to glycerol backbone, the melting point profile is largely dependent on the fatty acids attached. Therefore, a simple separation of MAG with different fatty acids having
different unsaturation is theoretically possible. In particular, EPA and DHA have 5-6
double bonds, the melting points of their MAG will be largely different from rest of
fatty acids in tuna oil. For this reason, a fractionation system with solvent used (so-
called wet fractionation) was studied.

Temperature is a critical issue for fractionation. Theoretically, MAG with
C16:0 to C18:0 have a melting point (mp) in the range between 69-75 °C under pure
lipid phase. The C18:1 based MAG has a mp around 24 °C and C18:2 based MAG
around 9 °C. No information for the mp of EPA and DHA based MAG, but a melting
point much lower can be expected. Once solvent applied, the melting behavior is
completely different from pure lipid phase. Both solvent and concentration in the
solvent can have effect on the crystallization temperatures. Based on application of
wet fractionation in industry as well as literature, three temperatures (0, 4, 10 °C)
were selected for this study.

Solvent is another issue. Acetone and hexane have been commonly applied in
industry and many previous studies. Considering the higher polarity of the material,
the polarity of solvent may have effects on the fractionation process. Therefore, both
solvents were selected for further evaluation including their mixtures.

The effects of solvent and temperature on yield and fatty acid compositions
were evaluated. Table 1 shows that percentage of EPA and DHA were higher in liquid
fraction than solid fraction. The yield of liquid fraction was decreasing in general with
the decreasing of temperature. Consequently, the EPA and DHA content in the MAG
of the liquid fraction was increasing. The effect of solvent mixing was not very
significant. There was a tendency that better fractionation was obtained in the acetone
system than in the hexane system. Yang et al. found that the percentage of saturated
fatty acid of stearin decreased with increasing solvent polarity, and percentage of EPA
and DHA increased with solvent polarity and fractionation temperature. Lee and Foglia\textsuperscript{11} also reported that fractionation with acetone at low temperature was effective for enriching the monounsaturated fatty acid of chicken fat in the liquid fraction. Yokochi \textit{et al.}\textsuperscript{18} reported that the winterization process with acetone at -20 °C showed higher separation efficiency for tri-unsaturated TAG into liquid fraction than the other solvents.

In general, C16:0 and C18:0 were dramatically reduced in the liquid fraction and increased in the solid fraction.\textsuperscript{11,12} The oleic acid was also changing but not highly consistent. The crystallization of oleic acid based MAG may need further lower temperature. As commonly known, the yield of the liquid fraction will be reduced with the decreasing of the temperature. The loss of the liquid fraction which is trapped by the solids, will also increase. With the present set-up, a liquid fraction with around 50% yield of MAG contained 70% EPA and DHA at 0 °C using acetone was obtained.

\textbf{CONCLUSION}

Physicoenzymatic production of MAG containing PUFA especially EPA and DHA was investigated. A few solvent mixtures were suitable for production of MAG by using Novozym 435 as a catalyst in glycerolysis of tuna oil. A few reaction parameters have been evaluated including solvent amount, substrate ratio, enzyme load, and temperature. The yield of MAG up to 90.8% could be achieved with suitable conditions. The temperature fractionation under different solvents was evaluated in order to produce a fraction with higher content of EPA and DHA. Temperature was a critical parameter for effective fractionation. A liquid fraction under 0 °C fractionation could be obtained with around 70% EPA and DHA and in a yield of
MAG around 50%. A possibility of enriching the EPA and DHA into MAG has been built.

ACKNOWLEDGEMENTS

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REFERENCES


Table 1. Effect of solvents ratios\(^a\) and temperatures on the major fatty acid compositions of monoacylglycerol fractions.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Solvent mixture (hexane /acetone, v/v)</th>
<th>Yield of liquid and solid fractions (wt%(^b))</th>
<th>Major fatty acid content (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>C14:0</td>
</tr>
<tr>
<td>10</td>
<td>100/0</td>
<td>L 84.9</td>
<td>2.9</td>
</tr>
<tr>
<td></td>
<td></td>
<td>S 15.1</td>
<td>6.6</td>
</tr>
<tr>
<td></td>
<td>50/50</td>
<td>L 69.8</td>
<td>3.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>S 30.2</td>
<td>2.7</td>
</tr>
<tr>
<td></td>
<td>0/100</td>
<td>L 63.1</td>
<td>3.9</td>
</tr>
<tr>
<td></td>
<td></td>
<td>S 36.9</td>
<td>1.6</td>
</tr>
<tr>
<td>4</td>
<td>100/0</td>
<td>L 67.9</td>
<td>2.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>S 32.1</td>
<td>6.7</td>
</tr>
<tr>
<td></td>
<td>50/50</td>
<td>L 71.6</td>
<td>2.2</td>
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<td></td>
<td></td>
<td>S 28.4</td>
<td>6.7</td>
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<td></td>
<td>0/100</td>
<td>L 65.3</td>
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<td></td>
<td>S 44.7</td>
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<td>0</td>
<td>100/0</td>
<td>L 52.8</td>
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<td></td>
<td>S 47.2</td>
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<td>L 49.4</td>
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<tr>
<td></td>
<td></td>
<td>S 52.5</td>
<td>5.3</td>
</tr>
</tbody>
</table>

\(^a\)MAG/solvent mixture, 1:30 (v/v): fractionation conditions: 3 h at designated temperature after rapid cooling from room temperature; fraction temperature were 10, 4 and 0 °C.

\(^b\)wt% recovery of liquid and solid fractions at the same temperature. L-liquid fraction; S-solid fraction.
FIGURE CAPTIONS:

Fig. 1. Glycerolysis time courses of tuna oil in tert-butanol. Reaction conditions: temperature 45 °C, glycerol/tuna oil molar ratio 4.5:1.0, tert-butanol/tuna oil 2.2:1.0 (wt/wt), 15 wt% lipases (based on total substrates), and no additional water. Abbreviations: TAG (triacylglycerol), DAG (diacylglycerol), MAG (monoacylglycerol), FFA (free fatty acid), AK (Immobile lipase AK), 435 (Novozym 435)

Fig. 2. Glycerolysis time courses of tuna oil in tert-butanol by immobilized lipase AK (a). dry with acetone (b) adjusted water content with $a_w = 0.113$ by saturated salt (LiCl) (c). dry in vacuum. Reaction conditions: temperature 45 °C, glycerol/oil molar ratio 4.5:1.0, tert-butanol/oil 2.2:1.0 (w/w), 15 wt% Immobilized Lipase AK (based total substrates) and no additional water. See Fig. 1 for abbreviations.

Fig. 3. Effects of solvent mixtures on glycerolysis of tuna oil. Reaction conditions: temperature 45 °C, glycerol/tuna oil molar ratio 4.5:1.0, reaction time 3 h, 15 wt% Novozym 435 (based on total substrates) and no additional water.

Fig. 4. Time course of glycerolysis by Novozym 435 in TB. The reaction mixture contained the mole ratio of glycerol to tuna oil with 4.0:1.0, the weight ratio of TB to tuna oil with 2.0:1.0, reaction time 3 h, 15 wt% Novozym 435 (based on glycerol and tuna oil) and no additional water.
Fig. 1

Lipid profile (wt%) vs. time (min)

- TAG-435
- TAG-AK
- FFA-435
- FFA-AK
- DAG-435
- DAG-AK
- MAG-435
- MAG-AK

Time (min) from 0 to 240
Fig. 2

![Bar chart](image)

**Drying method for immobilized Lipase AK**

- **Vacuum (aw=0.369)**
- **LiCl solution (aw=0.113)**
- **Acetone (aw=0.019)**

Lipid Profile (wt%)

- TAG
- FFA
- DAG
- MAG
Fig. 3

Mixture (%v/v) vs. Monoacylglycerol content (Wt%)
Fig. 4

Lipid profile (wt %)

- TAG-435
- DAG-435
- MAG-435
- FFA-435

Time (min)