

UNIVERSITI PUTRA MALAYSIA

TRANSESTERIFICATION OF JATROPHA CURCAS L. OIL TO BIODIESEL USING Nd2O2 AND Bi2O2 -SUPPORTED CATALYSTS

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By

RABIAH NIZAH BINTI MD FAHMY

Thesis Submitted to the School Graduate Studies, Universiti Putra Malaysia, in Fulfillment of the Requirement for the Degree of Master Science

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Abstract of the thesis presented to the Senate of Universiti Putra Malaysia in fulfillment of the requirement for the Degree of Master Science

TRANSESTERIFICATION OF *JATROPHA CURCAS L*. OIL TO BIODIESEL USING Nd₂O₃ AND Bi₂O₃ -SUPPORTED CATALYSTS

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June 2015

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Biodiesel is gaining more attention since it is a renewable source of energy that can be an alternative for petroleum based diesel fuels. Other than being renewable, it is also biodegradable and non-toxic. Biodiesel is even more preferable since it can be produced easily via transesterification reaction. With various resources that can be used to produce biodiesel, Jatropha Curcas oil (JCO) is one of the feasible source since it is a non-edible oil, hence no competition over food resources. One of the major drawback of JCO is that it contains high amount of free fatty acids (FFA). In this study, catalyst with different catalyst (Nd₂O₃ and Bi₂O₃) on La₂O₃ support were compared. Bi₂O₃ catalyst shows higher catalytic activity at lower reaction conditions, hence is used for further study. Different loading of Bi₂O₃ (1,3,5,7) wt.%) using La₂O₃ as a support were prepared using wet impregnation method for simultaneous esterification and transesterification of JCO and its potential as heterogeneous catalyst was assessed. The catalysts were characterized by using Xray Diffractometer (XRD), Brunauer-Emmett-Teller (BET) surface area, Scanning Electron Microscopy (SEM) and Temperature Programmed Desorption (TPD) of CO₂ and NH₃. These catalysts were then used for transesterification reaction under different reaction conditions (methanol to oil molar ratio, amount of catalyst, reaction temperature and reaction time) to investigate the catalytic activities of the catalysts. Under optimum transesterification condition at 150°C with catalyst amount of 2 wt.%, methanol/oil molar ratio of 15:1 and reaction time of 4 h, 5BiLa catalyst gave fatty acid methyl ester (FAME) conversion of 93%. The catalytic activities were found depending on the acidity, basicity and the surface area of the catalyst used. Several tests were conducted to study the physicochemical properties of the product such as pour point, flash point, kinematic viscosity, sulphur content and cloud point of biodiesel produced. Based on the results, the synthesized biodiesel is comparable with conventional diesel in the market since it meets the international standards of biodiesel which are American Standard for testing Materials (ASTM), Europian Standard (EN) and Malaysian Standard (MS) for diesel fuel specifications.

TRANS-PENGESTERAN BAGI MINYAK JATROPHA CURCAS L. KEPADA BIODIESEL MENGGUNAKAN MANGKIN Nd₂O₃ DAN Bi₂O₃ YANG DISOKONG

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Biodiesel semakin mendapat perhatian kerana ia adalah sumber tenaga yang boleh diperbaharui dan boleh menjadi alternatif kepada bahan api berasaskan petroleum diesel. Selain daripada boleh diperbaharui, ia juga bersifat biodegradasi dan tidak bertoksik. Biodiesel lebih menjadi pilihan kerana iaboleh dihasilkan dengan mudah melalui tindak balas trans-pengesteran. Daripada pelbagai sumber yang boleh digunakan untuk menghasilkan biodiesel, minyak Jatropha Curcas (JCO) adalah salah satu sumber yang boleh dilaksanakan kerana ia adalah minyak yang tidak boleh dimakan, maka tiada persaingan dengan sumber makanan. Salah satu kelemahan utama JCO ialah ia mengandungi jumlah asid lemak bebas (FFA) yang tinggi. Dalam kajian ini, pemangkin yang berbeza (Nd₂O₃ dan Bi₂O₃) kepada sokongan La₂O₃ dibandingkan. Mangkin Bi₂O₃ menunjukkan aktiviti pemangkin lebih tinggi pada keadaan tindak balas yang lebih rendah, oleh itu ia digunakan untuk kajian selanjutnya. Pelbagai muatan Bi₂O₃ (1,3,5,7wt.%) menggunakan La₂O₃ sebagai sokongan disediakan dengan menggunakan kaedah penjerapan untuk pengesteran dan trans-pengesteran serentak JCO dan potensi sebagai pemangkin heterogen dinilai. Pemangkin telah dicirikan dengan menggunakan alat pembelauan sinar-X (XRD), pengukuran luas permukaan Brunauer - Emmett -Teller (BET), pengimbas mikroskopi elektron (SEM), penyahjerapan karbon dioksida pada suhu terkawal (TPD-CO₂) dan penyahjerapan ammonia pada suhu terkawal (TPD-NH₃). Pemangkin-pemangkin ini kemudiannya digunakan untuk tindak balas trans-pengesteran pada beberapa keadaan (nisbah methanol terhadap minyak, jumlah pemangkin, suhu tindak balas dan masa tindak balas) untuk menganalisa aktiviti kesemua mangkin. Di bawah keadaan trans-pengesteran optimum pada suhu 150 °C, jumlah pemangkin 2 wt.%, nisbah methanol kepada minyak 15:1 dan masa tindak balas selama 4 jam, mangkin 5BiLa memberi penukaran asid lemak methil ester (FAME) sebanyak 93%. Aktiviti mangkin didapati bergantung kepada keasidan, kealkalian dan keluasan kawasan permukaan mangkin tersebut. Beberapa ujian telah dijalankan untuk menganalisa sifat-sifat fizikokimia biodiesel yang dihasilkan seperti takat tuang, takat kilat, kelikatan, kandungan sulfur dan takat awan. Berdasarkan keputusan tersebut, biodiesel yang

dihasilkan didapati setanding dengan diesel konvensional yang ada di pasaran memandangkan ianya dapat memenuhi piawaian biodiesel antara bangsa seperti American Standard for testing Materials (ASTM), Europian Standard (EN) dan Malaysian Standard (MS) spesifikasi untuk bahan api diesel di Malaysia.



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LIST OF ABBREVIATIONS

AES Atomic Emission Spectroscopy

ASTM American Society for Testing and Materials

BET Brunauer Emmett Teller EN European Standard FAME Fatty Acid Methyl Ester

FFA Free Fatty Acid
GC Gas Chromatography
JCO Jatropha Curcas Oil

JCPDS Joint Committee on Powder Diffraction Standards

MS Malaysian Standard
MPOB Malaysian Palm Oil Board
SEM Scanning Electron Microscopy

SV Saponification Value

TG Triglyceride

TPD Temperature Programmed Desorption

WD Wave Dispersive XRD X-Ray Diffraction

CHAPTER 1

INTRODUCTION

1.1 Energy and Renewable Energy

The world economy depends on two major energy carriers which are hydrocarbons (natural gas, gasoline, diesel fuel and heating oil) and electrical current. Although the primary energy supply for each country differs greatly, hydrocarbons are still the main source of energy. Lately, energy supply and its security have become a major issue around the world. International Energy Agency (IEA) had estimated 53 % of increase in global energy consumption by 2030, with 70 % of the growth in demand coming from developing countries. Demand for energy expected to increase over the next 24 years both in industrial countries and particularly in the developing countries like Malaysia where rapid economic growth is expected (Rahman Mohamed and Lee, 2006).

Peninsular Malaysia stands at 47 % with an energy reserved margin total of 20,493MW installed capacity. The excessive harnessing of various form of energy that made possible by the advancement of technologies resulted in unwanted byproduct such as waste and pollution. For example, fossil fuels consume and pollute water, endanger flora and fauna, generate toxic wastes and cause global warming. Due to the growth in exploration activities, Malaysia's proven oil reserves have declined in recent years and the oil production fell to 693,000 bbl/d in 2008, a 13% decrease from 2006 level. If the production rate is consistent at around 700,000 bbl/d, Malaysia's oil reserves will be exhausted in around 20 years (Mansor, 2008). There is a compelling need for energy variation and containment of the negative environmental impact on extensive use of mineral fuel. By depending mainly on oil and gas for half of a century, Malaysia started to comprehend the importance to adopt renewable energy in the energy mix and continuously reviewed its energy policy to ensure sustainable energy supply and security (Oh *et al.*, 2010).

Renewable energy sources (RESs) is also known as alternative energy. RESs are readily available in nature and they are primary energy resources. It is derived from those natural, mechanical, thermal and growth processes that continuously reproduced predictable quantities of energy when required. RESs use local resources that have potential to provide energy with zero or low emission of air pollutant and greenhouse gases. Renewable energy technologies manufacture marketable energy by converting natural materials into useful form of energy.

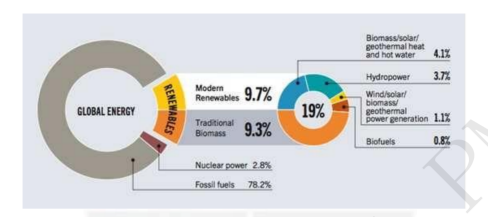


Figure 1.1 Estimated Renewable Energy Share of Global Final Energy Consumption, 2011

(Source: Mcginnet al., 2013)

By the end of 2011 (Figure 1.1), renewable energy supplied an estimated 19 % of global final energy consumption. From this total, approximately 9.3 % came from traditional biomass, geothermal heat about 4.1 %, hydropower made up about 3.7 % and an estimated 1.9 % from wind, solar, biomass, geothermal power generation and biofuels (Mcginn *et al.*, 2013).

1.2 Biofuels

Biofuel refers to liquid or gaseous fuel for transportation that are mostly produced from biomass. Biofuels mainly used in vehicles but also can be used in engine or fuel cells for electricity generation. Fuels that can be produced from biomass resources include liquid fuels such as ethanol, methanol, biodiesel and Fischer-Tropsch diesel and gaseous fuels such as methane and hydrogen. The biggest difference between biofuels and petroleum is the oxygen content. Biofuels have oxygen levels of 10 to 45 % while petroleum has almost none which makes the chemical properties of biofuels very different from petroleum (Demirbas, 2008a). Biofuels usually have very low sulphur and nitrogen levels. Some of the advantages of biofuels are easily derived from common biomass sources, carbon dioxide cycle occurs in combustion, they are environmentally friendly and biodegradable and contribute to sustainability (Kim and Dale, 2005).

With the increasing demands of energy, biofuels economy starts to grow rapidly. In Malaysia, there is biofuels policy of Malaysia which is based on Malaysia's National Biofuel Policy document. The main aim of the policy is to reduce the country's fuel import bill, further promoting the demand for palm oil, which is expected to be the primary commodity for biofuel production in Malaysia, as well as to shore up the price of palm oil especially during periods of low export demand.

1.3 Background and Potential of Biodiesel

Fatty acid methyl ester (FAME) or generally known as "biodiesel", are the best candidate for alternative diesel fuels derived from vegetable oils or animal fats. Biodiesel generally has similar physio-chemical and fuel properties as petroleum based diesel fuel. The major component of vegetable oils and animal fats are triglyceride (TG) where the 3 moles ester of fatty acids (FA) attached to one glycerol backbone. TG from vegetable oils or animal fats contains a few different FA which possesses different physical and chemical properties, making the FA profile the most important factor influencing the properties of vegetable oils or animal fats (Knothe and Gerpen, 2005).

The most common method to obtain biodiesel is through a chemical reaction known as transesterification reaction. In this reaction, TG are converted to FAME in the presence of catalyst with short chain alcohol giving glycerol as a byproduct. Vegetable oils and animal fats have high kinematic viscosity and low volatility making it unsuitable to be used as combustion fuel. Hence, vegetable oils and animal fats must undergo transesterification to reduce the viscosity of the oils (Muniyappa *et al.*, 1996).

1.4 Advantages of Biodiesel as Diesel Fuel Substitutes

Biodiesel have several distinct advantages compared to petroleum-based diesel fuel. Biodiesel is biodegradable, sustainable and environmental friendly given that it is considered to be carbon neutral. Biodiesel feedstock such as Jatropha, rape plant and palm trees absorbs carbon dioxide that released to the atmosphere when used for combustion in diesel engines (Atadashi *et al.*, 2011) as illustrated in Figure 1.2. Biodiesel also gives less exhaust emission with low level of unburned hydrocarbon, carbon monoxide and particulate matter. Compared to mineral diesel, biodiesel have lower sulphur content and no carcinogen, thus it can be called as a clean fuel. Since biodiesel is derived from renewable domestic resource, it would help to reduce the dependency on petroleum as well as preserving petroleum sources (Demirbas, 2008b).

As mentioned earlier, biodiesel possesses similar physico-chemical properties as diesel fuel so it can be used directly without further modification of diesel engine. There are two ways to use biodiesel in engines. It can be used as 100 % biodiesel fuel or it can be blended with diesel fuel since biodiesel is completely miscible with petroleum diesel fuel. A pure (100 %) biodiesel is known as B100 or "neat" fuel. Biodiesel blends are referred as BXX where XX indicates the amount of biodiesel blends in the fuel (i.e. a B10 blend is 10 % biodiesel and 90 % petroleum based diesel). Biodiesel fuels generally have excellent lubricity which helps to prolong engine life and reduce the need for maintenance (Demirbas, 2010). Table 1.1 summarized the advantages of biodiesel as diesel fuel substitute.



Figure 1.2 Biodiesel Production Cycle (Source:Mann, 2011)

Table 1.1 Advantages of Biodiesel as Diesel Fuel Substitutes

	Sustainability
Economic impacts	Fuel diversity
Economic impacts	Agricultural development
	Reducing the dependency of crude petroleum
	Greenhouse gases reduction
Environmental impacts	Biodegradable
Environmental impacts	Carbon sequestration
	Lower sulphur content
	Domestic targets
Energy accurance	Supply reliability
Energy assurance	Ready availability
	Renewability

1.5 Global Biodiesel Market Demands

The energy demand especially in transportation sector increasing each day due to the rapid urbanization worldwide. This increasing need of energy contributes to the development of the renewable energy making biodiesel as one of the fastest growing industries.

The global biodiesel production from 2000 to 2012 shown in Figure 1.3 and in table 1.2. Global production of biodiesel reached over 22.5 billion litres in 2012. 41 % of the production was from Europe, with major contribution from Germany which produced 2.7 billion liters in 2012. United States held the top spot for world's leading biodiesel producer with 3.6 billion liters followed by Argentina (2.8 billion liters), Brazil and Germany (2.7 billion liters) and France (1.9 billion liters) in 2012 (Mcginn *et al.*, 2013).



Figure 1.3 Global Biodiesel Production 2000-2012

(Source: Mcginn et al., 2013)

Table 1.2Global Biodiesel Production of Top 14 Countries (Mcginnet al., 2013)

Country	Biodiesel (billion litres)
United States	3.6
Brazil	2.7
Germany	2.7
Argentina	2.8
France	1.9
China	0.2
Canada	0.1
Thailand	0.9
Indonesia	1.5
Spain	0.5
Belgium	0.4
Netherlands	0.5
Colombia	0.3
Austria	0.4

World production and consumption of biodiesel grows rapidly during 2007 to 2012. The high percentages of biodiesel production from European Union (EU) countries mainly comes from the European government supports in terms of production incentives including tax incentives and loan guarantees. Furthermore, the new biofuel policy such as tax exemption, mandates and incentives for biodiesel introduced in countries such as United States, Europe, Brazil and Asia had encouraged the production and consumption of biodiesel to the fullest (Oh *et al.*, 2010).

1.6 Progress of Biodieselin Malaysia

The first biodiesel program in Malaysia was initiated by Malaysian Palm Oil Board (MPOB) in 1982 using palm oil as feedstock. Malaysian government funded the program in order to bring palm biofuel to local and international markets. MPOB collaborated with local oil company Petronas and constructed the first pilot plant two years later. This plant successfully produces 3000 tonnes biodiesel annually. In August 2006, the first commercial biodiesel plant started its operation. Malaysia had announced its national biodiesel policy in 2005 resulting in the government approving around 92 licenses for biodiesel projects to produce 10.2 million tonnes per year (Lopez and Laan, 2008). MPOB reported that Malaysia has exported 47,790 and 95,010 tonnes of biodiesel in 2006 and 2007 (MPOB, 2008).

Since biodiesel production in Malaysia mainly related to palm oil, the government determined to develop palm oil industries by promoting palm biodiesel. In 2011, Petronas, Shell, BHP, ExxonMobil and Chevron have been allocated with a start up fund worth RM 1 million each by MPOB to set up infrastructure for B5 biodiesel blending facilities. By June 2011, B5 biodiesel sold at six petrol stations located in Putrajaya (Times, 2011). Due to the unstable price of palm oil, Jatropha has become another alternative as biodiesel feedstock. MPOB carried out some performance test on Jatropha biodiesel alongside with Malaysian Rubber Board and The National Tobacco Board. Malaysian Rubber Board engaged in the seed breeding of Jatropha while The National Tobacco Board is responsible to gauge the suitability of cultivating Jatropha on bris soils in the northern part of Malaysia (Cottrell and Hoh, 2011).

1.7 Problem Statements

Biodiesel production industry grown exponentially in many countries as alternative fuels. A lot of studies had been done with growing interest in biodiesel production. Despite the fact that biodiesel offers several advantages over fossil fuels, the major concern about biodiesel are the cost and economic issues. Homogeneous catalysts are typically used for transesterification reaction. Although homogeneous catalyst produces high yield of biodiesel at lower reaction conditions, it is difficult to separate the catalyst from the product and needs purification of the product that leads to environmental pollution. Hence, heterogeneous catalysts are used in this research to overcome the weakness of homogeneous catalyst since it is more efficient and environmental friendly.

Conventionally, vegetable oils such as canola, soybean and corn are normally used as feedstocks for biodiesel production. However, nowadays, we are facing "food vs. fuel" dilemma regarding the risk of diverting corps for biofuel production which harms the food supply on global scale. Non-edible oils meet these requirements because they are inedible and can be grown in waste land with low fertilizer. In this

research, *Jatropha Curcas* oil (JCO) which is a non-edible oil are selected as a feedstock to replace the food corpses. Jatropha is reported as one of the best candidate for future biodiesel production (Lim and Teong, 2010).

Unfortunately, crude JCO contains high amount of free fatty acid (FFA). An acid and base catalyzed two step method usually used for oils with high FFA's. In this method, the oil is pretreated with an acidic catalyst before transesterification performed by using a basic catalyst. This reaction requires several reactions, washing and separation stages which is not economic. To overcome the disadvantageous of the two step process for biodiesel from oils containing high FFA, a new class of heterogeneous catalysts with both acid-base properties is developed. To improve these catalysts, a bi-functional acid-base heterogeneous catalyst is developed. Lanthanum oxide (La₂O₃)found to contain moderate Lewis acid sites (La³⁺) and base sites (O²⁻, OH). The role of metal dopant in lanthanum oxide catalyst was extendedly revised. In this research, the influence of two different metal substitutions and influence of different weight percent of Bi³⁺ substituted on lanthanum oxide catalyst were investigated towards Jatropha conversion. Furthermore, the parameter biodiesel optimization transesterification reaction such as methanol/oil molar ratio, catalyst loading, reaction time and reaction temperature were studied to get high biodiesel conversion.

1.8 Objectives

The objectives of this research are:

- To synthesize Nd₂O₃-La₂O₃ and Bi₂O₃-La₂O₃catalysts via impregnation method.
- 2. To characterize the catalysts using several methods:
 - (a) X-Ray diffraction (XRD) Analysis
 - (b) Brunauer-Emmett-Teller (BET) surface area analysis
 - (c) Temperature programmed desorption of ammonia (TPD-NH₃)
 - (d) Temperature programmed desorption of carbon dioxide (TPD-CO₂)
 - (e) Scanning electron microscopy (SEM) and energy dispersive X-Ray (EDX)
 - (f) Inductively coupled plasma-atomic emission spectrometer (ICP-AES)
- 3. To produce biodiesel from crude JCO in the presence of Nd₂O₃-La₂O₃ and Bi₂O₃-La₂O₃ catalysts and investigate the catalytic performance of the catalysts towards biodiesel production.

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APPENDIX

A. BET Surface Area Analysis

Data collected was evaluated by using BET adsorption isotherm Equation 3:

$$\frac{P}{V(P_0 - P)} = \frac{1}{V_m C} + \frac{(C - 1)}{V_m C} (\frac{P}{P_0})$$
 (3)

Where:

P = equilibrium partial pressure of adsorbate gas in equilibrium with the surface at -196 °C

 P_o = saturated pressure of adsorbates at corresponding temperature

V =volume of gas adsorbed at STP

 V_m = volume of monolayer gas adsorbed at STP on the catalyst

surface

C = BET constant related to the enthalpy of adsorption

BET equation can be rearranged into linear form to:

$$y = mx + c$$

Where:

$$y = \frac{P}{V(P_0 - P)}$$

$$m = \frac{(C-1)}{V_m C}$$

$$x = \frac{P}{P_0}$$

$$c = \frac{1}{V_m C}$$

Hence, a graph of $\frac{P}{V(P_0-P)}$ versus $\frac{P}{P(P_0-P)}$ be plotted and the surface area of the catalyst can be calculated by using $\frac{P(P_0-P)}{V_mC}$ as the value of m and $\frac{1}{V_mC}$ as the value of c (y-intercept)

B. Determination of Saponification Value& Molecular weight

The saponification value was determined by Equation 4:

$$SV = \frac{56.1 \text{ N } (V_b - V_c)}{W} (4)$$

Where:

SV is the saponification value in mg/g

Vb is the volume (mL) of HCl solution used for blank;

Vs is the volume (mL) of the solution used for determination of the sample;

N is the normality of HCl;

W is the weight (g) of the test portion;

56.1 is the molecular weight of KOH

Average molecular weight (MW) of triglyceride (TG) of oil can be estimated by Equation 5:

AverageMWofTG =
$$\frac{56.1 \times 1000 \text{ mg} \times 3}{\text{SV}} (5)$$

Where:

56.1 is molecular weight of KOH

The saponification value for crude JCO used in this study:

Saponification Value =
$$\frac{56.1 (0.5 \text{ N}) (21.70 - 7.65 \text{ mL})}{2.004 \text{ g}}$$
$$= 188.64 \text{ mg/g}$$

Average molecular weight of crude JCO:

Average MW of TG =
$$\frac{56.1 \times 1000 \text{mg} \times 3}{188.64 \text{ mg/g}}$$

= 892.19 g/mol

C. Determination of Acid Value& %FFA

The acid value was determined by Equation 6:

$$AV = \frac{(V_b - V_c) N 56.1}{W}$$
 (6)

Where:

ΑV is the acid value in mg/g

Vb is the volume (mL) of KOH solution used for blank;

Vs is the volume (mL) of the solution used for determination of the sample;

N is the normality of KOH;

W is the weight (g) of the test portion; 56.1 is the molecular weight of KOH

FFA content can be estimated by Equation 7:

$$\frac{\%FFA}{28.2} = \frac{(V_b - V_c) N}{W} (7)$$

Where:

Vh is the volume (mL) of KOH solution used for blank;

is the volume (mL) of the solution used for determination of the sample; Vs

N is the normality of KOH;

W is the weight (g) of the test portion

28.2 is molecular weight of oleic acid divided by 10

By combining Equation 6 and 7, % FFA can be calculated as in Equation 8: $\% FFA = \frac{AV}{1.99}(8)$

$$%FFA = \frac{AV}{1.99}(8)$$

The acid value for crude JCO used in this study:

Acid Value =
$$\frac{(8.10 - 7.50 \text{ mL})(0.1 \text{ N})56.1}{0.2011 \text{ g}}$$
$$= 16.73 \text{ mg/g}$$

The %FFA for crude JCO used in this study:

$$\%FFA = \frac{16.73}{1.99}$$
$$= 8.40\%$$

D. Determination of FAME conversion

The conversions of the jatropha oil into biodiesel were calculated from the weight of the glycerol obtained using these equations Equation 9 and 10as in below:

$$Conversion(\%) = \frac{\text{moleofesperimentalgsyceros}}{\text{moleoftheoreticalgsyceros}} \times 100\%$$
 (9)

$$Conversion(\%) = \frac{\frac{maccofglycerol}{molecularweightofglycerol}}{\frac{maccoftryglyceride}{molecularweightoftriglyceride}} \times 100\%$$
(10)

E. Optimization conditions for La₂O₃ support (page31)

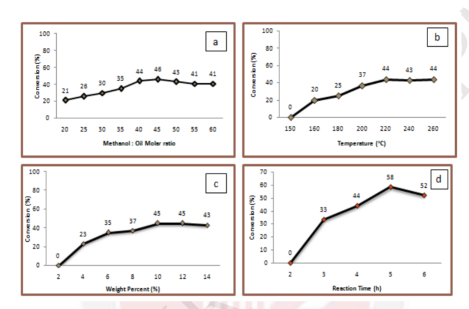


Figure A1. (a) Effect of Me:Oil molar ratio, (b) Effect of reaction temperature, (c) Effect of catalyst amount and (d) Effect of reaction time

F. TPD- CO₂ and TPD-NH₃graphs for preliminary study (pg32)

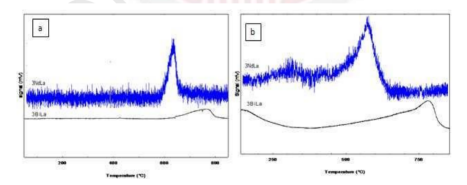


Figure A2 (a) TPD-CO $_2$ profiles for 3Bila and 3NdLa catalysts, (b) TPD-NH $_3$ profiles for 3BiLa and 3NdLa catalysts

BIODATA OF STUDENT

Born in Port Dickson, Negeri Sembilan on 18th August 1988, Rabiah Nizah attended her primary school at Sekolah Kebangsaan Port Dickson from 1995 to 2000. From year 2001-2003, she went to secondary school at Sekolah Menengah Kebangsaan Tinggi Port Dickson and had her PMR. Then in 2004-2005, she went to Sekolah Menengah Sains Muar located Muar, Johor and did her SPM there. After that, she continues her study at Johor Matriculation College, Tangkak, Johor in Physical Science for two semesters. In 2007, she was accepted to further her study in Universiti Putra Malaysia (UPM) and she managed to complete her Bachelors degree majoring in Petroleum Chemistry in 2010. Next, she pursued her second degree in Master of Science, majoring in Catalysis at Catalysis Science and Technology Research Centre, Faculty of Science, UPM under Professor Taufiq Yap Yun Hin's supervision. During her research period, she gained experiences in operating various instruments such as x-ray diffraction (XRD), BET surface area, temperature desorption programmed of carbon dioxide (TPD-CO₂), gas chromatography (GC), scanning electron microscopy (SEM) and so on. Her research was supported financially by Ministry of Education.

LIST OF PUBLICATIONS

Published papers:

- 1. Rabiah Nizah, M.F., Taufiq-Yap, Y.H. and Hussein, M.Z. (2013) Production of Biodiesel From Non-edible *Jatropha Curcas* Oil via Transesterification Using Nd₂O₃-La₂O₃ catalyst. *Advance Materials Research* 620: 335-339.
- 2. Rabiah Nizah, M.F., Taufiq-Yap, Y.H., Umer Rashid, Teo, S.H., Shajaratun Nur, Z.A. and Aminul, I. (2014). Production of biodiesel from non-edible *Jatropha Curcas* oil via Bi₂O₃-La₂O₃ catalyst. *Energy Conversion and Management*. 88: 1257-1262.

List of seminars/ conferences/ workshops attended:

- 1. The 7th International Conference of Chemical Engineering on Science and Applications (ChESA), Banda Aceh, Indonesia. (Oral presenter).
- 2. International Conference on X-Rays And Related Techniques in Research & Industry, Universiti Sains Malaysia, 2012. (Oral Presenter).
- 3. 19th IUPAC International Conference on Chemical Research Applied to World Needs (CHEMRAWN XIX), Putra World Trade Center (PWTC), 2011. (Participant).
- 4. Response Surface Methodolgy (RSM) Workshop, Institute of Bioscience, Universiti Putra Malaysia, 2011. (Participant).
- 5. X-Ray Fluorescence Workshop, Universiti Kebangsaan Malaysia, 2011. (Participant).
- 6. 2nd Seminar on Catalysis Science and Technology (2nd CAST), Universiti Putra Malaysia. (Participant)
- 7. Public Lecture: Energy Issues and Renewable Energy from Biomass/Wastes by Professor Dr. Kuniyuki Kitagawa, Universiti Putra Malaysia, 2010. (Participant)
- 8. PutraCAT Public Lectures: Hydrogen for Sustainable Energy by Professor Dr. Ichiro Naruse and Professor Dr. Yukihiko Matsumura, Universiti Putra Malaysia, 2013. (Participant).

- 9. PutraCAT Public Lecture: Rational Catalyst Design A case study of thermo neutral reforming catalyst development by Dr. Shakeel Ahmed, Universiti Putra Malaysia, 2012. (Participant)
- 10. PutraCAT Public Lecture: Renewable Energy from Waste Biomass by Assoc. Prof. Dr. Larence M. Pratt, Universiti Putra Malaysia, 2012. (Participant)

