

**Process simulation, economic analysis and synthesis of
biodiesel from waste vegetable oil using supercritical methanol**

by

Soo Jin Lee

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Abstract

Biodiesel production using supercritical methanol received attention as an alternative method to replace the conventional alkali-catalyzed method being practiced in industry. Due to its flexibility to feedstock compared to the conventional method, the supercritical method for waste vegetable oil conversion appears to be promising in environmental and economical points of views.

Four industrial-scale biodiesel production processes were simulated using Hysys. Each process used either the conventional or the supercritical method. The first model simulated the alkali-catalyzed process using fresh vegetable oil. The second process model resembled the first one as it uses sodium hydroxide catalyst in transesterification, but pre-treatment process of waste vegetable oil was included. The third and fourth models were supercritical biodiesel production processes using waste vegetable oil. Fourth model had differences from the third one in terms of the amount of methanol being introduced to a plug flow reactor and the way of recovering methanol from reaction products, both of which can reduce energy consumption of the process. To improve the accuracy of the process simulations, properties of a model compound (triolein) of the vegetable oils were examined via thermogravimetric analysis, and the experimental data were incorporated into the simulation models. Economical aspects of the developed simulation models were then assessed using Aspen Icarus Process Evaluator.

The economic assessment revealed that supercritical processes using waste vegetable oil were competitive to the conventional process based on their better profitability indicators such as discounted payback period and net present value. The net present value prediction formulas were derived for the four processes via statistical analysis of the vegetable oil price, biodiesel selling price, by-product selling price and interest rate that were found to most strongly affect the profitability of the biodiesel production processes by sensitive analysis.

Experiments of biodiesel synthesis from waste canola oil were conducted using supercritical methanol. High methyl ester yields over 96% were achieved after 45 min of reaction time at 270°C/10 MPa with methanol to oil ratios of 1:1 and 2:1. Side reactions such as glycerol decomposition and glycerol methanolysis were confirmed by water content measurement using Karl-Fischer titration and Gas chromatography-mass spectrometry (GC-MS) analysis.

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List of Abbreviations

316SS	316 Stainless steel
ANOVA	Analysis of variance
ASME	American society of mechanical engineering
B100	100% pure biodiesel fuel
CCD	Central composite design
CS	Carbon steel
CSTR	Continuous stirred tank reactor
cwt	Centum weight (=100 pounds)
DCFROR	Discounted cash flow rate of return
DF	Degree of freedom
DPP	Discounted payback period
DSC	Differential scanning calorimeter
EAV	Equivalent annual value
EOS	Equation of state
EPA	Energy policy act
FAME	Fatty acid methyl ester
FFA	Free fatty acid
FID	Flame ionized detector
FVO	Fresh vegetable oil
G&A	General & administrative
GC	Gas chromatography
GC-MS	Gas chromatography-mass spectrometry
HETP	Height equivalent to a theoretical plate
IPE	Icarus process evaluator
IRR	Internal rate of return
KF	Karl-Fischer
L-L separator	Liquid-liquid separator
MARR	Minimum acceptable rate of return
MeOH	Methanol
MOC	Material of construction
NPV	Net present value
NRTL	Non-random two liquid model
PFR	Plug flow reactor
Prob	Probability
R&D	Research & development
RFS	Renewable fuels standards
RSM	Response surface model
SO _x	Sulphur oxide
TGA	Thermogravimetric analysis
TG-MS	Thermogravimetry and mass spectrometry

UBC	The University of British Columbia
UNIFAC	Universal functional activity coefficient
US EIA	U.S. Energy information administration
US EPA	U.S. Environmental protection agency
USDA	U.S. Department of agriculture
WEC	World energy council
WVO	Waste vegetable oil

List of Notations

A	Cross-sectional area [m ²]
A _d	Downcomer area [m ²]
A _T	Tower inside cross-sectional area [m ²]
D	Diameter [m]
D _T	Tower diameter [m]
G	Mass flow rate of vapour [kg/s]
L	Length [m]
P _c	Critical pressure [kPa]
Q	Volumetric flow rate [m ³ /s]
Re	Reynolds number
T _b	Boiling point [°C]
T _c	Critical temperature [°C]
U _f	Vapour flooding velocity [m/s]
V	Mean fluid velocity [m/s]
V _c	Critical volume [m ³ /kmol]
μ	Dynamic viscosity [kg/(m·s) or Pa·s]
ν	Kinematic viscosity [m ² /s]
ρ	Density [kg/m ³]
ρ _G	Vapour density [kg/m ³]
ρ _L	Liquid density [kg/m ³]
ω	Acentricity

Glossary

ASTM D5555-95	Standard test method for determination of free Fatty acids contained in animal, marine, and vegetable fats and oils used in fat liquors and stuffing compounds
ASTM D6751	Standard specification for biodiesel fuel blend stock (B100) for middle distillate fuels
EN14214	EU biodiesel fuel standard

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Co-authorship Statement

Chapter 2 will be revised into a manuscript co-authored with Dr. Dusko Posarac and Dr. Naoko Ellis at the University of British Columbia. As the first author, I was in charge of the thermogravimetric analysis of triolein and tripalmitin and the development of four biodiesel production process simulations incorporated with the thermogravimetric analysis result of triolein.

Chapter 3 will be revised into a manuscript co-authored with Dr. Dusko Posarac and Dr. Naoko Ellis at the University of British Columbia. My contribution to this manuscript includes assessment of the four processes from an economic perspective. I was in charge of evaluation of profitability, identification of essential elements of biodiesel costs and sensitivity analysis of net present values of the four processes.

Chapter 4 will be revised into a manuscript co-authored with Dr. Dusko Posarac and Dr. Naoko Ellis at the University of British Columbia. My contribution to this manuscript includes conducting experiments of laboratory-scale biodiesel production using a batch reactor, analyzing water content of reactants and products using a Karl Fischer volumetric titrator and analyzing a profile of fatty acid methyl esters in biodiesel products via gas chromatography.

All co-authors contributed to the identification and design of the research project and will assist in the preparation and revision of the manuscripts.

1. Introduction

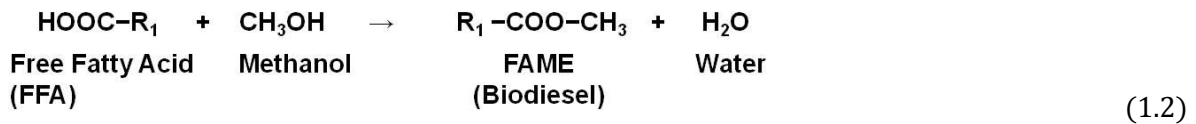
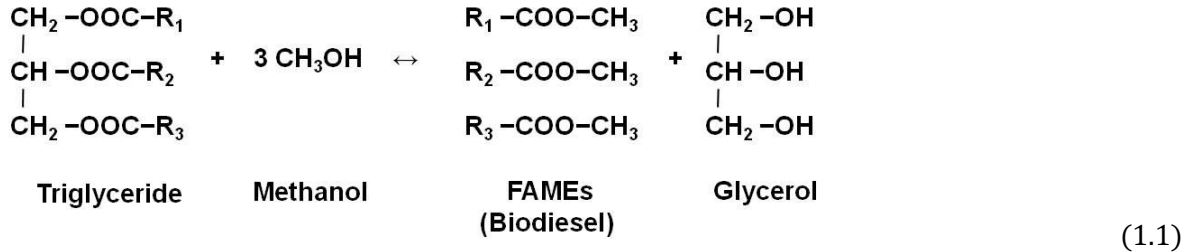
Exploitation of alternative fuels such as hydrogen, non-fossil natural gas and biofuels has been actively conducted in recent years. Skyrocketing prices of petroleum-based fuels and growing environmental concerns are boosting public attention to alternative fuels. Moreover, worldwide energy demand has steadily increased, widening the gap between the supply and demand of petroleum oil, raising concerns about the depletion of fossil fuels. According to a World Energy Council (WEC) study, if current practice is continued, worldwide energy demand in 2020 will be 50-80% higher than in 1990 (Omer, 2008).

Among the alternative transportation fuels on which research is in progress, biodiesel is one of the most promising because the infrastructure for the distribution of biodiesel is well established compared to the others. Biodiesel is distinguished from petroleum-based diesel by the fact that it is a renewable energy. It also has lower environmental impact, reducing emissions of particulate matter, carbon monoxide, total hydrocarbons and toxic aromatic and polyaromatic compounds (Lapuerta et al., 2008). Therefore, promoting innovative applications of biodiesel can contribute to the achievement of sustainable development.

Many countries have legislated the promotion of biodiesel in the forms of fuel mandates, tax incentives and funding for capital projects or fleet upgrades (Mabee, 2007). In December 2006, the Government of Canada announced that it would regulate the renewable content of diesel fuel to achieve 2% biodiesel content by 2012. In the United States, The Energy Policy Act of 2005, H.R. 6, was signed into law by President George W. Bush. This act contains renewable fuels standards (RFS) that will raise the use of biofuels to 28.4 billion litres per year by 2012, which is effectively 5% of total fuel sales (Mabee, 2007). The European Union set a policy target that mandates increasing the share of biofuels from 2% to 5.75% of the total fuel supply (based on energy content) by 2010 (Mabee, 2007). Strategic planning at the local and national levels alone, however, cannot implement a transition to sustainable energy use patterns. Only when it is combined with technological progress of biodiesel production, biodiesel can have sufficient market ability, thereby increasing its percentage of the fuel market.

Biodiesel is defined as a mixture of fatty acid alkyl esters which are commonly produced from triglycerides and alcohol through transesterification reaction shown in Equation (1.1). Methanol is the most commonly used alcohol for the reaction due to its relatively low cost and high rate of conversion. Stoichiometrically, a 3:1 molar ratio of methanol to oil is required to convert one triglyceride molecule into three methyl ester molecules. In practice, this ratio needs to be higher to provide a maximum ester yield. Transesterification consists of a sequence of three consecutive reversible reactions. The first step

is the conversion of triglycerides to diglycerides, followed by the conversion of diglycerides to monoglycerides, and finally monoglycerides to glycerol. Overall, the three fatty acids bound to the glycerol bond of triglyceride are combined to CH₃- group of methanol, yielding three fatty acid methyl esters. A catalyst is usually used in transesterification reaction to improve the reaction yield and rate. Biodiesel can also be produced from free fatty acids (FFAs) and alcohol through esterification reaction, generating water as the by-product as shown in Equation (1.2).



The type of catalysts and feedstocks used for the transesterification reaction significantly affects the unit operations involved in a biodiesel production process. Catalysts and feedstocks are also closely related to each other and affect the economic profitability of biodiesel production. Therefore, intensive research is devoted to finding pertinent feedstocks and establishing optimum reaction conditions.

1.1. Feedstocks: vegetable oils

The end cost of biodiesel mainly depends on the price of its feedstock (Canakci and Sanli, 2008). The predominant feedstocks are currently food-grade vegetable oils which mostly consist of triglycerides in the range of C12–C22. Depending on local climate and soil conditions, soybean, rapeseed, palm, canola, coconut and sunflower oils are employed in biodiesel production. The composition of the oil itself affects the properties of the biodiesel produced from the oil. For example, biodiesel from rapeseed oil, which has a high concentration of unsaturated fatty acids, has better cold

flow properties than biodiesel from palm oil, which is rich in saturated fatty acids (Gui et al., 2008). The compositions of vegetable oil are various as shown in Table 1.1.

Table 1.1. Fatty acid composition (wt%) of various vegetable oils (Source: Ramos et al (2009))

Fatty acid		Palm	Olive	Rape	Soybean	Sunflower	Peanut	Corn	Grape
Lauric	C12:0	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0
Myristic	C14:0	0.7	0.0	0.0	0.0	0.0	0.1	0.0	0.1
Palmitic	C16:0	36.7	11.6	4.9	11.3	6.2	8.0	6.5	6.9
Palmitoleic	C16:1	0.1	1.0	0.0	0.1	0.1	0.0	0.6	0.1
Stearic	C18:0	6.6	3.1	1.6	3.6	3.7	1.8	1.4	4.0
Oleic	C18:1	46.1	75.0	33.0	24.9	25.2	53.3	65.6	19.0
Linoleic	C18:2	8.6	7.8	20.4	53.0	63.1	28.4	25.2	69.1
Linolenic	C18:3	0.3	0.6	7.9	6.1	0.2	0.3	0.1	0.3
Arachidic	C20:0	0.4	0.3	0.0	0.3	0.3	0.9	0.1	0.3
Gadoleic	C20:1	0.2	0.0	9.3	0.3	0.2	2.4	0.1	0.0
Behenic	C22:0	0.1	0.1	0.0	0.0	0.7	3.0	0.0	0.0
Erucic	C22:1	0.0	0.0	23.0	0.3	0.1	0.0	0.1	0.0
Lignoceric	C24:0	0.1	0.5	0.0	0.1	0.2	1.8	0.1	0.0
Nervonic	C24:1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

However, recent research has moved toward alternative feedstocks because the use of food-grade oils has become less and less economically viable as a result of the increasing prices of crops. In addition, the use of edible oils as feedstocks for biodiesel production has been criticized for reducing destitute countries' access to food. Alternative feedstocks include waste cooking oil, yellow and brown greases, animal fats and non-edible vegetable oils such as *Jatropha* and castor oils. These potential feedstocks are expected to benefit the commercialization of biodiesel by lowering its cost and reducing the ethical issues involved in its production.

Waste vegetable oils (WVOs) are daily generated in large quantities during food preparation by frying in fast-food franchises, local restaurants, dining rooms, catering, and so on. According to Zhang et al. (2003), approximately 120,000 tonnes/yr of yellow grease are produced in Canada. Therefore, the use of this large quantity of waste oil would clearly be an environmentally friendly and economical way to produce biodiesel. However, WVOs from restaurants and food industries are known to have a large variety of qualities due to the many variables involved in the frying process, such as temperature, length of heating, and the kind of food subjected to frying. During the frying process, the physical and chemical properties of the oil change, going through various chemical reactions such as hydrolysis, polymerization and oxidation. Consequently, free fatty acids (FFAs) and

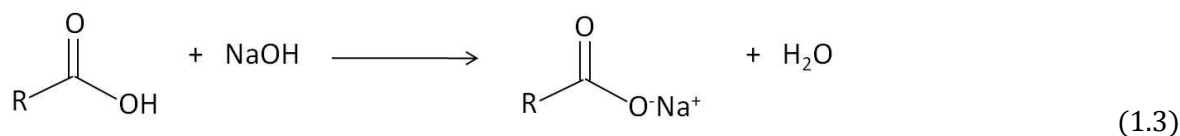
moisture contents increase due to the hydrolysis of triglycerides in the presence of food moisture and oxidation (Canakci, 2007). As an example, Tyagi and Vasishtha (1996) reported that the FFA level of fresh soybean oil changed from 0.04% to 1.51% after 70 hrs of frying at 190°C. Increases in viscosity were also reported due to polymerization, which resulted in the formation of compounds with higher molecular weight (Canakci, 2007). Therefore, using waste oils, with their variance in properties, as feedstock for biodiesel production is expected to present technical challenges.

1.2. Catalysis in transesterification

The conversion methods for producing biodiesel are divided into four categories depending on the catalysts involved in the transesterification reaction. These are the homogeneous alkali-catalyzed, homogeneous acid-catalyzed, heterogeneous catalyzed and supercritical methods.

1.2.1. Homogeneous alkali-catalyzed transesterification

Biodiesel is conventionally produced in industry through a batch process using homogeneous alkali catalysts. The most commonly used alkali catalysts are sodium hydroxide (NaOH), potassium hydroxide (KOH), and sodium methoxide (CH₃ONa). Compared to using acid catalysts, alkali-catalyzed transesterification has several advantages: a faster reaction rate, lower catalyst amount, lower reaction temperature, and a less corrosive final product (Sanli and Canakci, 2008). The process typically produces methyl esters with a high yield at a temperature of 60°C and atmospheric pressure within about 1 hr (Meher et al., 2006). However, alkali catalysts are not suitable for transesterification of feedstocks containing >0.5% FFAs (equivalent to the acid value of 1 mg KOH/g) or >0.06% water as a result of the saponification reaction shown in Equation (1.3). Soap formation partially consumes the catalyst, decreases biodiesel yield, and complicates the separation and purification steps (Vicente et al., 2004). Because of these limitations, low-cost feedstocks such as waste vegetable oils are not appropriate candidates for alkali-catalyzed transesterification unless an extra pre-treatment step is performed.



1.2.2. Homogeneous acid-catalyzed transesterification

Sulphuric acid has been the most investigated homogeneous acid catalyst. Apart from that, HCl, BF₃ and H₃PO₄ may be also used in transesterification reaction. It is reported that a homogeneous acid-catalyzed reaction is approximately 4000 times slower and achieves a lower yield than a homogeneous alkali-catalyzed reaction (Srivastava and Prasad, 2000). However, an acid-catalyzed process is generally believed to be more tolerant of high FFAs levels in the feedstock which makes it suitable for low-grade oils.

Canakci and Van Gerpen (1999) examined the conversion rate to methyl esters of soybean oil at different levels of FFAs content, using a 3% sulphuric acid catalyst. They were able to achieve 90% conversion from soybean containing 5 wt% FFAs. The level of FFAs, however, should not exceed 5% to achieve ester conversion over 90%. The authors also reported that the transesterification of soybean oil was affected by as little as 0.1 wt% water and was strongly inhibited by 5 wt% water in the oil. More recently, Kusdiana and Saka (2004) compared the effect of water content on methyl ester formation from rapeseed oil via alkali- and acid-catalyzed transesterification reactions. Approximately 70% conversion was achieved in the alkali-catalyzed reaction, whereas the conversion was only 6% in the acid-catalyzed reaction. The results from both of these studies indicate that an acid-catalyzed process is very sensitive to moisture in the feedstock oils.

1.2.3. Heterogeneous catalyzed transesterification

A heterogeneous catalyzed process is expected to decrease biodiesel production cost and reduce environmental impact by simplifying production and purification processes under mild conditions. Heterogeneous catalysts include solid base catalysts, solid acid catalysts and enzymes. Recent research is focused on exploring new and sustainable heterogeneous catalysts for transesterification reaction such as metal oxides, metal complexes, active metals loaded on supports, zeolite, resins, membranes and lipases (Kansedo et al., 2009).

Kawashima et al. (2008) investigated 13 metal oxides for transesterification reaction at 60°C with a 6:1 molar ratio of methanol to oil and 10 hr of reaction time. Catalysts with calcium such as CaZrO₃, Ca₂Fe₂O₅, CaMnO₃ and CaO-CeO₂ were found to have ester conversions of 92%, 92%, 88%, and 89%, respectively, because of their high base strength.

Research on direct use of a solid acid catalyst for biodiesel production has not been widely explored because of its limitation of a slow reaction rate and a knowledge gap in fundamental studies relating to reaction pathways of triglycerides on solid acids (Lam et al., 2010). Several studies are reported on the use of zirconium oxide (ZrO₂) as a solid acid catalyst because of its strong surface acidity. Jitputti et al. (2006) reported that methyl ester yields from palm kernel oil and crude coconut

oil were as high as 90.3% and 86.3%, respectively, using $\text{SO}_4^{2-}/\text{ZrO}_2$. More recently, titanium dioxide (TiO_2) was suggested as a solid acid catalyst. Chen et al. (2007) reported that $\text{SO}_4^{2-}/\text{TiO}_2$ with a specific surface area of $99.5 \text{ m}^2/\text{g}$ achieved a 90% methyl ester yield from cotton seed oil.

Enzymatic transesterification reactions especially those using lipase have advantages of easy recovery of product, mild reaction condition, insensitiveness to high FFA oil, and absence of by-product generation (Lam et al., 2010); however, slow reaction rate, enzyme deactivation and high costs of enzymes still remain as constraints. Currently, sources of enzymes such as *Rhizopus oryzae*, *Mucor miehei*, *Candida antarctica* and *Pseudomonas cepacia* are also under active investigation (Lam et al., 2010).

1.2.4. Supercritical transesterification

To overcome the problems associated with catalyzed processes, a supercritical process has been proposed by Saka and Kusdiana (2001). In this process, transesterification reaction proceeds under high temperature and pressure condition without use of a catalyst. Under these conditions, a single homogeneous phase of an alcohol and oil mixture is formed, which usually exists as a heterogeneous phase (two liquid phases) because of immiscibility of the polar and non-polar components. Because of better miscibility, reaction time is short. Saka and Kusdiana (2001) reported that transesterification reaction of rapeseed oil at $350^\circ\text{C}/45 \text{ MPa}$ gave a high ester yield $>95\%$ in 4 min with a methanol to oil molar ratio of 42:1. As a result of high yield and absence of a catalyst, the separating, washing, and drying processes involved in the conventional process could be eliminated in a supercritical process. The supercritical biodiesel production scheme is shown in Figure 1.1.

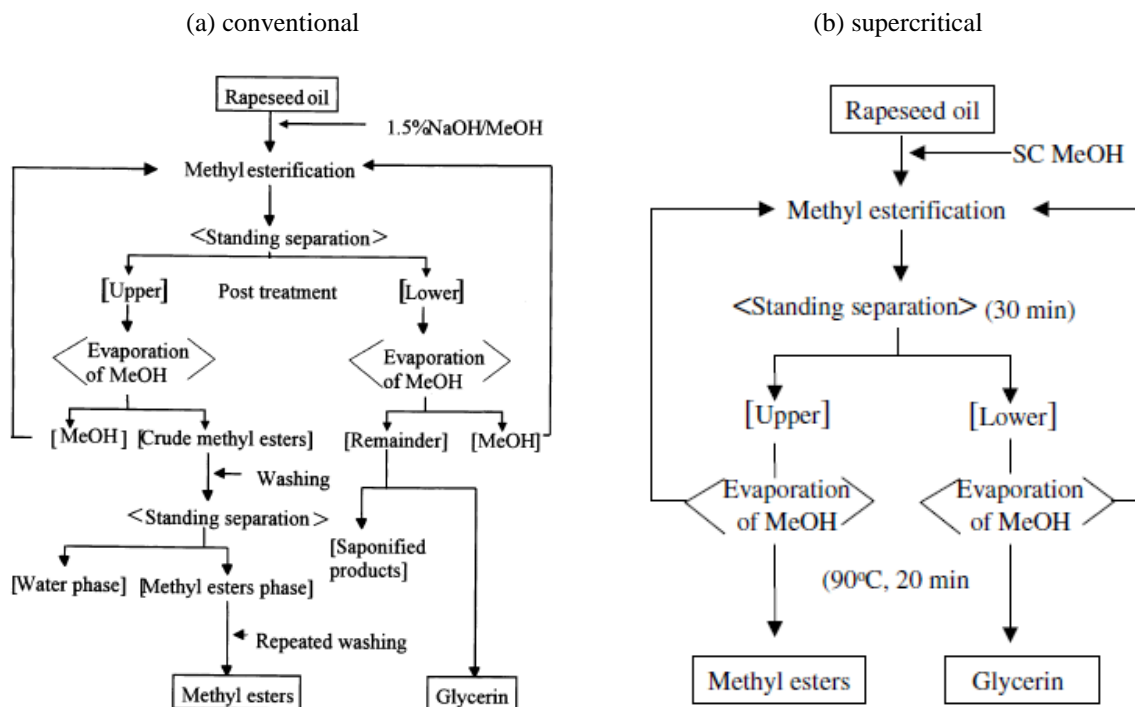


Figure 1.1. Schematic processes of biodiesel fuel production from rapeseed oil by: (a) conventional; and (b) supercritical methods (Saka and Kusdiana, 2001)

Another advantage of the supercritical transesterification reaction is that it allows relatively high content of FFAs and water in feedstocks compared with a catalyzed transesterification reaction. According to Kusdiana and Saka (2004), almost complete conversion was achieved for rapeseed oil with oleic acid content of up to 30 wt% in a supercritical reaction. The yield of methyl esters was not affected by the presence of water up to 36 wt%.

All the reported studies of supercritical transesterification reaction have shown that the reaction temperature is one of the most critical parameters for determining conversion rate. For instance, as temperature increases from 210 to 280°C, reaction rate constant increases approximately 7-fold at a pressure of 28.0 MPa and a methanol to oil ratio of 42 (Sawangkeaw et al., 2010). Varma and Madras (2007) reported the effect of temperature on the conversion of castor and linseed oils. Only 55% conversion of castor oil was obtained at 200°C after 60 min of reaction, whereas nearly complete conversion was achieved at 350°C after 40 min of reaction. Although a high temperature enhances a reaction rate, an excessively high temperature (>350°C) is not preferable because of thermal degradation of methyl esters (Imahara et al., 2008).

Stoichiometrically, a methanol-to-oil molar ratio of 3:1 is required in transesterification, and increasing the ratio to 50:1 is known to benefit the reaction as a result of increased contact area

between methanol and triglycerides (Sawangkeaw et al., 2010). For instance, a high product yield greater than 95% was obtained after only 4 min reaction time in a methanol to oil molar ratio of 42, but the yield was lower than 60% at molar ratios of 6 or less.

At present, the supercritical conversion method is in the research stage, using mostly a laboratory-scale batch or continuous reactor (up to 200 ml). To commercialize the process, there are still several challenges remaining in research and development. In particular, high temperature and pressure conditions as well as excessive amounts of methanol that result in high energy consumption in a supercritical process need to be alleviated. In addition, research on behaviour of the reaction system in large industrial plants needs to be done.

1.3. Process simulation and economic assessment

One of the best ways to conduct an economic assessment of biodiesel production is to develop a simulation model of the process. Simulating a complete biodiesel production process allows investigation of technological feasibility and limitations of the process before economic aspects are examined. Furthermore, the effect of process parameters such as operating temperatures and pressures can be studied and optimized. In this way, a reliable economic analysis of a biodiesel production process can be conducted.

Simulating industrial-scale biodiesel production processes was first attempted by Zhang et al. (2003) to assess economic profitability. Four different configurations of a biodiesel plant were developed using Hysys process simulator: a homogeneous alkali-catalyzed process using pure vegetable oil, a homogeneous alkali-catalyzed process with pre-treatment of waste vegetable oil, a homogeneous acid-catalyzed process using waste vegetable oil and a homogeneous acid-catalyzed process using hexane extraction to purify the final product. Subsequent economic assessments of the four processes showed that the acid-catalyzed process without hexane extraction was the most promising. West et al. (2008) compared homogeneous alkali-catalyzed, homogeneous acid-catalyzed, heterogeneous catalyzed (SnO) and supercritical biodiesel production processes. The results revealed that the heterogeneous process was the most economically feasible, followed by the supercritical process; however, the conversion used by West et al. (2008) in the heterogeneous process was higher than the experimental value (92.6%) provided by Abreu et al. (2005), to which the authors referred, which might have changed the economic analysis considerably. In addition, studies regarding the effects of FFAs and water contents in feedstock have not been conducted for the heterogeneous acid

catalyst, raising a question whether the technology can be practically implemented in biodiesel production using low-cost feedstocks such as waste vegetable oil.

1.4. Thesis objectives

First, a supercritical process simulation model is developed. To improve the accuracy of the simulation, a better understanding of the model compound of vegetable oil, triolein, is required. Triolein has been used as a model compound in many previous studies, but some of its chemical properties are not well known. In particular, the normal boiling point of triolein has been estimated by a few researchers, but the estimated values are not consistent with each other. Moreover, some estimated triolein boiling points are considerably different from the ones built into process modeling programs. The correct value of the normal boiling point is extremely important, since many missing properties are estimated from it. In this thesis, using thermogravimetric analysis (TGA), the normal boiling point will be measured and compared with values from the literatures and process simulators.

Second, the technological feasibility of a supercritical biodiesel production process is investigated. By replacing the normal boiling point in Hysys with the one from TGA measurements, a supercritical process simulation with improved accuracy is developed. In addition, to decrease the high energy consumption problem of a supercritical process, another supercritical process model using less energy is developed. The two supercritical processes are then compared with conventional homogenous alkali-catalyzed processes in terms of material flows, energy flows, and number and type of unit operations.

Third, the economic feasibility of supercritical processes is compared with alkali-catalyzed processes. This is done using an Aspen Icarus Process Evaluator (IPE) whose practicability has been confirmed by engineering design and construction firms. Compared with previous researches using Lang's factor method or Guthrie's bare-module concept, the economic assessment conducted in this study is expected to provide more reliable results. Moreover, intensive sensitivity analysis will be conducted on the developed process simulations.

Finally, this thesis investigates laboratory-scale biodiesel production from waste cooking oil using supercritical methods. Compared with an alkali-catalyzed method, a supercritical method is seen favourable in economical point of view because of its ability to use low-cost feedstocks. The effect of three parameters, reaction time, reaction temperatures and weight ratios of methanol to waste canola oil, will be investigated.

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2. Simulation of four continuous biodiesel production processes¹

2.1. Introduction

Biodiesel has drawn attention as an alternative energy source, especially as a substitute to petroleum-derived diesel. Feedstocks for biodiesel are mainly vegetable oils, yellow or brown greases, and animal fats. Biodiesel is produced from transesterification of triglycerides in the feedstocks. Depending on the type of feedstock, the compositions of fatty acids that compose triglycerides are varied.

One of the important parameters in the process modeling of transesterification is to select a model compound to represent the triglyceride which contains various compositions of fatty acids. Triolein has been selected as a model compound of vegetable oils in many previous studies (Zhang et al., 2003; Kasteren and Nisworo, 2007; West et al., 2008; Glisic et al., 2009; Lim et al., 2009). Triolein is composed of three oleic acids with a glycerol bond, and oleic acid is the major fatty acid in many types of vegetable oils. It composes around 40-80% of fatty acids in rapeseed, canola, olive, palm and peanut oils (Pinnarat and Savage, 2008).

Despite the significance, the chemical properties of triolein have not been thoroughly understood. The reported normal boiling point and critical properties of triolein vary significantly by different estimation methods and process modeling programs, as shown in Table 2.1. The use of comprehensive properties of the major component will lead to more accurate and reliable simulation results in process modeling.

Among the properties of triolein, the normal boiling point is of interest in predicting other properties such as a critical temperature and pressure. Goodrum and Geller (2002) applied thermogravimetric analysis (TGA) method to measure the normal boiling point of medium- and long-chain triglycerides, namely trilaurin (C12:0), trimyristin (C14:0), tripalmitin (C16:0) and tristearin (C18:0). The results were in good accordance with the Clausius-Clapeyron model and with previously published data at low pressures (Perry et al., 1949). On the other hand, Ceriani and Meirelles (2004) proposed a group contribution method to estimate vapour pressures of fatty compounds. The estimation model is a function of temperature with a few parameters to take account for various molecular groups. The parameters were obtained by regressions of more than 1300 experimental values on vapour pressures of fatty compounds (saturated and unsaturated fatty acids, fatty esters, fatty alcohols, monoglycerides and triglycerides) gathered from published papers.

¹ A version of this chapter is in preparation for submission for publication. Lee, S., Posarac, D. and Ellis, N. (2010). Simulation of four continuous biodiesel production processes.

In the present work, the normal boiling point of triolein was measured using TGA method and compared with those estimated by the group contribution method. The boiling point of triolein obtained from TGA was further applied to UNIFAC structure in a process simulator, Hysys v.2006.5, to estimate the best properties of triolein.

Table 2.1. Comparison of available properties of triolein

	Hysys	Aspen	VMGsim	Tang et al. (2006)	Glisic et al. (2007)	Weber et al. (1999)
T _b (°C)	606.8	846.85	541.7	879.9 ^a	-	-
T _c (°C)	680.9	1367	681	954.1 ^a	704.73 ^c	673.92 ^d
P _c (kPa)	360.2	470	360.2	360.2 ^a	334 ^c	468.20 ^d
V _c (m ³ /kmol)	3.09	3.09	3.09	-	-	-
ω	1.686	-	1.69	1.6862 ^b	1.978 ^c	1.686

^a Estimated by the method of Dohrn and Brunner (1991, 1994)

^b Estimated according to Han and Peng (1993)

^c Estimated by the method of Constantinou and Gani (1994, 1995)

^d Estimated by the method of Ambrose (1978) and Reid (1987)

2.2. Thermogravimetric analysis of triolein

2.2.1. Method

Triolein (Sigma grade, ≥99%) and tripalmitin (Sigma grade, ≥99%) were purchased from Sigma-Aldrich. 20μl of hermetic alumina pans and hermetic lids with a laser-drilled pin hole were purchased from TA Instruments. A triolein sample of 4.5 mg (±0.5 mg) was placed at the centre of an alumina pan and 1 mg (±0.2 mg) of alumina powder was added to assist in achieving isothermal boiling. The alumina pan was sealed with a lid using a sample encapsulation press from TA Instruments. The pan was completely sealed except a 0.050–0.100 mm diameter laser-drilled hole at the centre of the lid. The hole permits pressure balance between the sample and the environmental pressure of 1 atm. The rate of diffusion out of the pan is regarded less than the vapourization rate inside the pan due to the very small size of the hole. In the same manner, a 3.5 mg (±1 mg) of tripalmitin sample with 1 mg (±0.2 mg) of alumina powder was prepared.

TA Instruments Model TGA/DSC Q600 was used for the thermogravimetric analysis. At atmospheric pressure, a flow of 50 ml/min of nitrogen gas was used. The temperature of the furnace was programmed to ramp at 10°C/min from room temperature to 550°C and to be isothermal at 550°C

for 5 min. To ensure the consistency of the results, five and three replicate runs were conducted for triolein and tripalmitin, respectively.

Universal Analysis 2000 version 4.5a, supplied by TA Instruments, was used as software to analyze the boiling point of the samples. The boiling point was determined as an intercept of the initial isothermal base line and the tangent of the isothermal weight loss slope as shown in Figure 2.1. The tangent line was drawn in automatic mode based on the two onset points designated at the start point (20°C) of initial baseline and end point (550°C) of the secondary baseline.

2.2.2. Results and discussion

Figure 2.1 shows a weight loss curve of a sample consisted of 4.91 mg of triolein and 1.08 mg of alumina powder. The sample weight remained constant until the temperature reached at 370°C, and then rapidly decreased after 400°C. After the evaporation of triolein, a second constant line appeared at the 20% of the initial sample weight, indicating that the alumina powder added to triolein only remained in the pan. Five replicate runs all showed the same single rapid and definite weight-loss curve without any secondary transition. This fact indicates that there was not any thermal decomposition during heating since the weight curve would have had multiple transition points if the analyzed material were broken into small molecules. The average normal boiling point obtained from the replicate runs was 412.8°C with a standard deviation of 1.08°C. Except for one run where the boiling point was measured at 410.0°C, the others gave the boiling point at 413.0 or 414.0°C. Therefore, the final boiling point of triolein was determined as 412.8°C, which was the mean value of the experimental results.

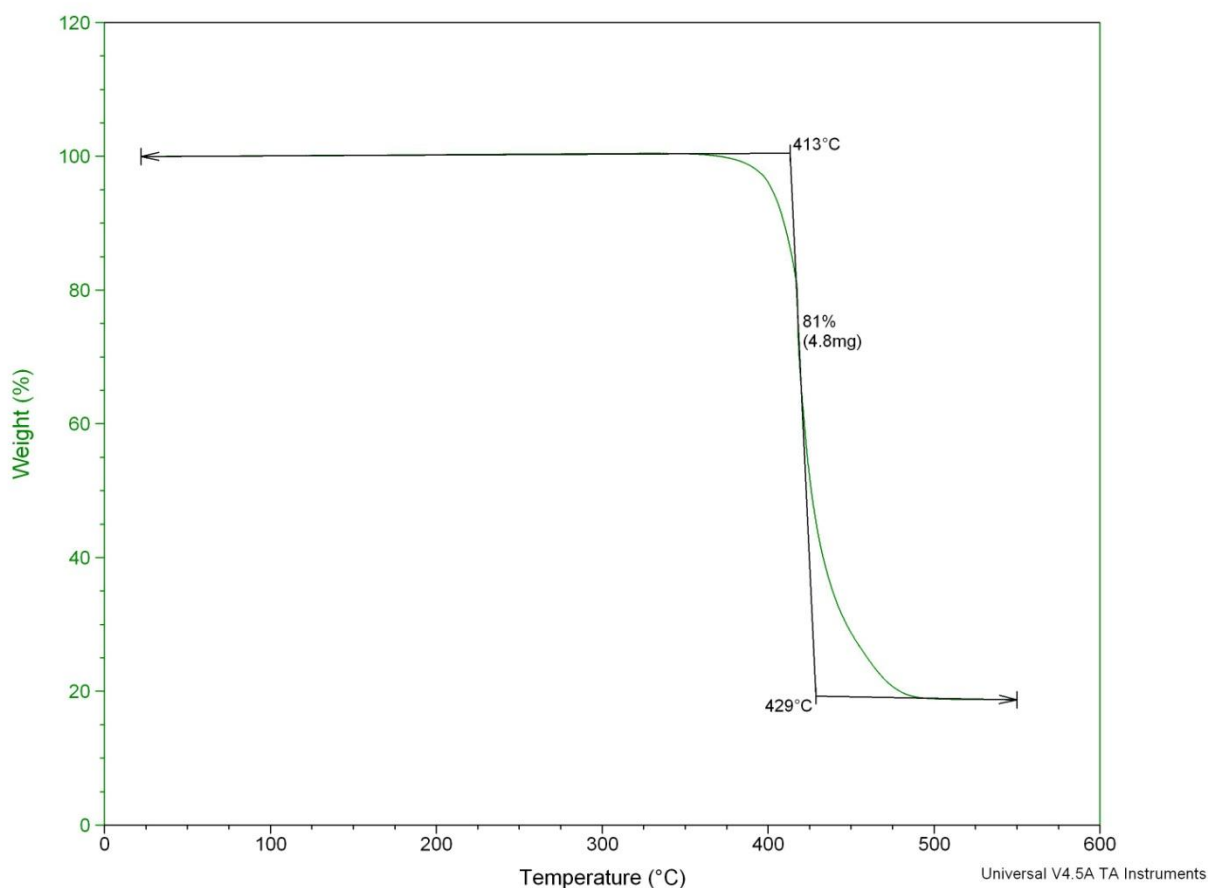


Figure 2.1. Weight loss curve of triolein (C18:1) measured in TGA

As shown in Figure 2.2, the weight loss curve of tripalmitin also had a single thermal transition point with increasing temperature. The TGA-derived boiling point of tripalmitin remained consistent in three replicate runs, measured as 409.0 or 410.0°C. The averaged boiling point of tripalmitin was 409.3°C in this study, which showed only 10°C difference from the value reported in the literature (Goodrum and Geller, 2002).

The group contribution method proposed by Ceriani and Meirelles (2004) has been developed specifically for fatty compounds such as fatty esters and acylglycerols:

$$\ln P_i^{vp} = \sum_k N_k \left(A_{1k} + \frac{B_{1k}}{T^{1.5}} - C_{1k} \ln T - D_{1k} T \right) + \left[M_i \sum_k N_k \left(A_{2k} + \frac{B_{2k}}{T^{1.5}} - C_{2k} \ln T - D_{2k} T \right) \right] + Q \quad (2.1)$$

where P_i^{vp} in Pa and T in K.

The vapour pressure is estimated based on molecular structures and functional groups of the compounds of interest along with several parameters (A_{1k} , B_{1k} , C_{1k} , D_{1k} , A_{2k} , B_{2k} , C_{2k} and D_{2k}) obtained from the regression of experimental data. The correction term, Q , accounts for isomers and the effect of functional groups. Since triolein and tripalmitin belong to acylglycerols, this term is zero. The parameters for triolein and tripalmitin are shown in Table 2.2. The calculated normal boiling points of triolein and tripalmitin are shown in Table 2.3.

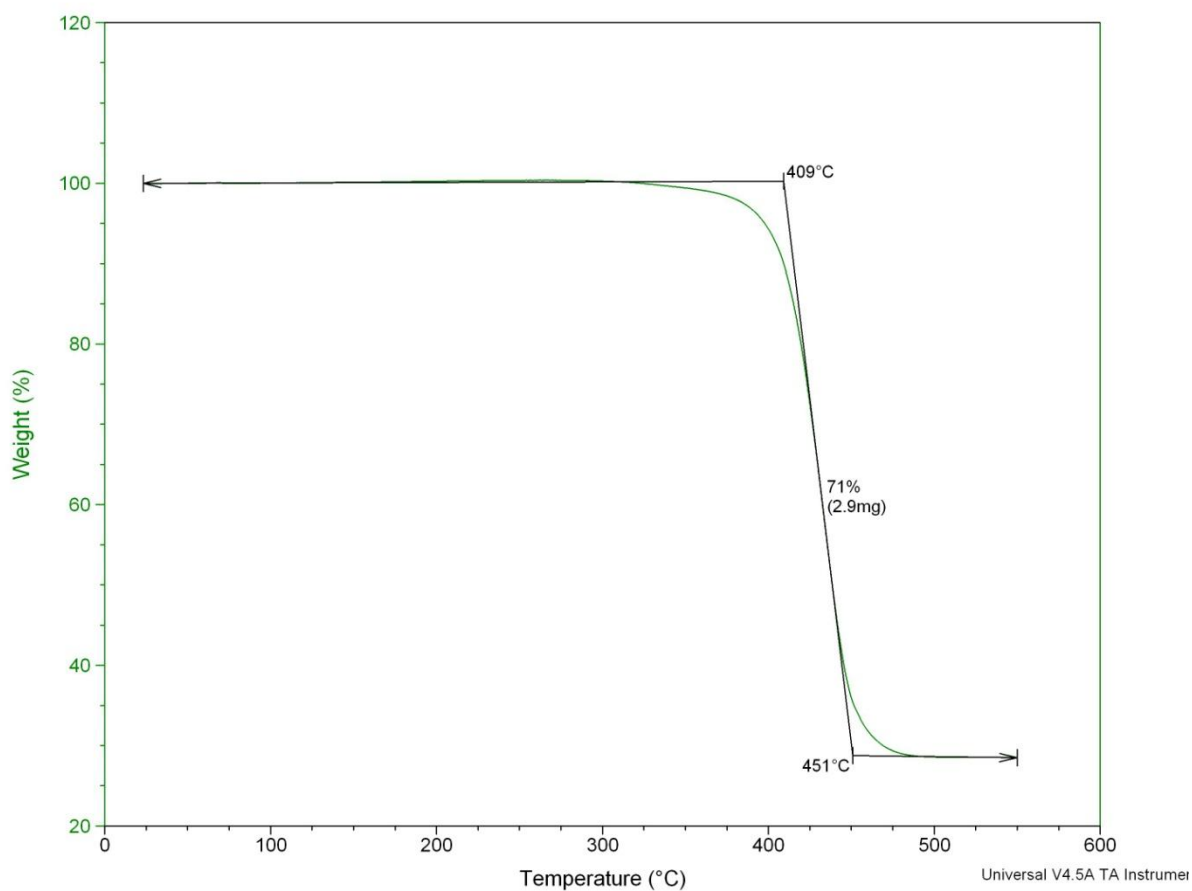


Figure 2.2. Weight loss curve of tripalmitin (C16:0) measured in TGA

Table 2.2. Parameters for Equation (2.1) (Source: Ceriani and Meirelles (2004))

Group	number of group		A_{1k}	B_{1k}	C_{1k}	D_{1k}	A_{2k}	B_{2k}	C_{2k}	D_{2k}
	triolein	tripalmitin								
CH ₃	3	3	-117.5	7232.3	-22.7939	0.0361	0.00338	-63.3963	-0.00106	0.000015
CH ₂	42	42	8.4816	-10987.8	1.4067	-0.00167	-0.00091	6.7157	0.000041	-0.00000126
CH= _{cis}	6	0	2.4317	1410.3	0.7868	-0.004	0	0	0	0
COO	3	3	7.116	49152.6	2.337	-0.00848	0.00279	10.0396	-0.00034	0.00000295
CH ₂ - CH- CH ₂	1	1	688.3	-349293	122.5	-0.1814	-0.00145	0	0	0

Table 2.3. Comparison of boiling point of tripalmitin and triolein obtained from TGA, DSC and group contribution method

Method	TGA (°C) (this study)	TGA (°C) (Goodrum and Geller, 2002)	DSC (°C) (Goodrum and Geller, 2002)	Group contribution (°C) (Ceriani and Meirelles, 2004)
Tripalmitin (C16:0)	409.3	399.08	401.88	449.6
Triolein (C18:1)	412.8	N/A	N/A	432.2

The boiling points of triolein and tripalmitin estimated by the group contribution method were approximately 30–40°C higher than the experimental results. The main reason for the differences appears to be generated from the characteristic of the group contribution method itself. Since the group contribution method was developed to be used for a broad range of fatty compounds, obtaining the boiling point of a particular compound may slightly deviate from the true value. Furthermore, the parameters in Equation (2.1) were obtained from the regression of the experimental data in literatures, which are very rare in case of the boiling point of triglycerides at atmospheric pressure. However, compared to the values listed in Table 2.1, the group contribution method estimated the boiling point much closer to the experimental results in this study.

The TGA boiling point obtained in this study was used along with the molar mass (885.4 g/mol), ideal liquid density (916 kg/m³) and UNIFAC structure to estimate critical and thermodynamic properties of triolein. Temperature dependent properties such as vapour enthalpy, vapour pressure and Gibbs free energy were simultaneously estimated. This triolein was named TGA Triolein to make distinction from the triolein built in Hysys component library, named Hysys Triolein. There were several changes observed between TGA and Hysys Triolein as shown in Table 2.4. The major differences were specific heat capacity, mass density and viscosity.

Table 2.4. Comparison of the properties of Hysys triolein, TGA triolein and canola oil

	Specific heat capacity [kJ/(kg·°C)]	Mass density [kg/m³]	Viscosity [cP]
Hysys Triolein ^a	0.131	1571	1.765
TGA Triolein ^a	1.58	909.8	33.19
Canola oil	1.67-1.97	920	57

^a Properties at 25°C and 1 atm

As shown in Table 2.4, the specific heat capacity and viscosity of TGA Triolein is 12 and 19 times, respectively, higher than Hysys Triolein. The mass density of TGA Triolein was 58% of that of Hysys Triolein. Overall, the properties of TGA Triolein were found to be much closer to the properties of real canola oil. Especially, the mass density of triolein was much improved as considering common knowledge that oil has less density than water (approximately 1000 kg/m³). These indicate that consequent improvements can be achieved in a process simulation by replacing Hysys Triolein with TGA Triolein. As an example, the difference in specific heat capacity of triolein would affect the energy consumptions in a supercritical process where the raw materials need to be heated up to high temperatures (>280°C). In addition, due to the existence of unreacted triolein, the performance of the biodiesel purification steps could also be altered.

2.3. Process simulation

The common way to produce biodiesel is via transesterification reaction using vegetable oil and alcohol. There exist several processes depending on the type of catalysts being used: homogeneous alkali-catalyzed; homogeneous acid-catalyzed; heterogeneous catalyzed; and supercritical processes. Among those, a homogeneous alkali-catalyzed process using NaOH or KOH is well understood and applied in most of industrial biodiesel plants today. On the other hand, a supercritical process is a new technique that is initially proposed by Saka and Kusdiana (2001). The process does not require any catalyst because the transesterification reaction is fast and gives a high yield of biodiesel at supercritical conditions of methanol (>8 MPa and >239.45°C) by achieving better miscibility between methanol and oil.

Each process has both advantages and disadvantages. An alkali-catalyzed process requires moderate reaction conditions (1 atm and 60°C) and is a proven process that is being used in the biodiesel industries. However, it has a limitation on utilizing low cost feedstocks such as waste vegetable oils or animal fats due to their relatively high content of free fatty acids (FFA). FFAs are known to make difficulties in purifying biodiesel product and to decrease biodiesel yield due to

saponification reaction. One of the ways to use those feedstocks is to pre-treat them through esterification using acid-catalysts. A supercritical process, on the other hand, is not affected by the presence of FFAs and water in the feedstocks (Kusdiana and Saka, 2004). A simpler biodiesel purification step is another advantage of the supercritical process. However, its high energy consumption due to high temperature and pressure conditions is a significant disadvantage. In addition, the safety concern arising from the intense operating conditions could be an issue, and further research on scaling up a laboratory-scale production unit still remains.

In this study, simulations of four biodiesel production processes were developed using Hysys. The first process, named Alkali-FVO, is an alkali-catalyzed process using fresh canola oil as the feedstock. The second process, named Alkali-WVO, is an alkali-catalyzed process with an acid-catalyzed pre-treatment step of waste canola oil. The process flowsheet and operating conditions for Alkali-FVO and Alkali-WVO processes referred to those reported by Zhang et al. (2003) and West et al. (2008). The third process, named SC-WVO, is a supercritical process using waste canola oil. The last process, named SC-LowE, is a supercritical process with slight modifications of the process flowsheet of SC-WVO process to lower energy consumption.

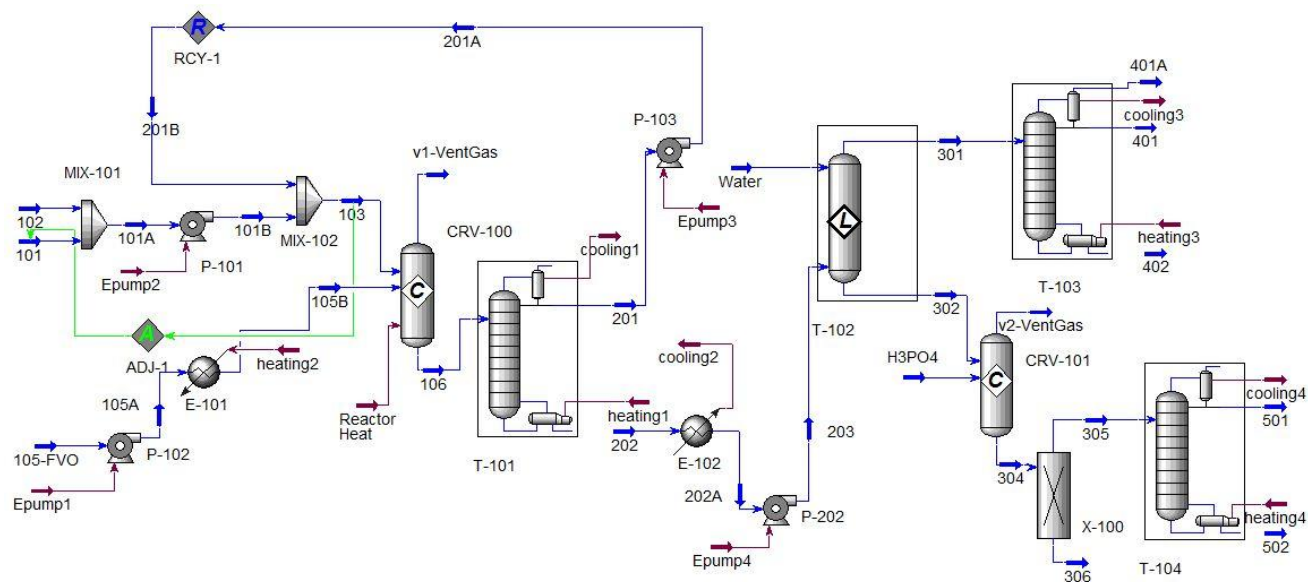
TGA Triolein ($C_{57}H_{98}O_6$) developed in Section 2.2 was used as a model compound to represent canola oil. Fresh canola oil was modeled as 100% TGA Triolein, and waste canola oil was modeled as a mixture of 94 wt% TGA Triolein and 6 wt% oleic acid ($C_{18}H_{34}O_2$). Consequently, methyl oleate ($C_{19}H_{36}O_2$) was considered as biodiesel.

Non-random two liquid (NRTL) model was selected as a property package for Alkali-FVO and Alkali-WVO processes due to the presence of polar compounds. Some interaction parameter coefficients between components were not available in Hysys library. These parameters were estimated using UNIFAC LLE model. Since an activity coefficient based model such as NRTL is not recommended to be used at pressures greater than 1000 kPa, Peng-Robinson equation of state (EOS) was used in process streams and unit operators wherever having pressures >1000 kPa.

The annual capacity of all the four processes was 40000 tonnes/year of biodiesel production based on 8000 operating hours per year.

2.3.1. Alkali-FVO process

A flow diagram and properties of main streams of Alkali-FVO process are shown in Figure 2.3.



Stream name	101	105-FVO	103	106	201	202	301	401	305	502
Pressure (kPa)	101.3	101.3	400.0	400.0	20.00	30.00	110.0	10.00	101.3	30.00
Temperature (°C)	25.00	25.00	26.69	60.00	28.20	142.8	59.39	70.97	163.1	180.0
Molar flow (gmol/hr)	17644	5929	36880	42810	17390	24880	17640	17150	6534	5743
Mass flow (kg/hr)	565.3	5250	1192	6442	574.4	5868	5284	4999	533.7	518.6
Volumetric flow (m ³ /hr)	0.7192	5.770	1.510	7.007	0.7340	48.70	6.205	5.968	4.943	0.5214
Component mass fraction										
Methanol	1.000	0.000	0.956	0.093	1.000	0.004	0.001	0.001	0.004	0.000
TGA triolein	0.000	1.000	0.000	0.041	0.000	0.045	0.050	0.002	0.000	0.000
Methyl oleate	0.000	0.000	0.000	0.778	0.000	0.854	0.948	0.996	0.000	0.000
Glycerol	0.000	0.000	0.000	0.080	0.000	0.088	0.000	0.000	0.967	0.995
NaOH	0.000	0.000	0.044	0.008	0.000	0.009	0.000	0.000	0.000	0.000
H ₃ PO ₄	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
H ₂ O	0.000	0.000	0.000	0.000	0.000	0.000	0.001	0.000	0.029	0.005
Na ₃ PO ₄	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000

Figure 2.3. Flow diagram and properties of main streams of Alkali-FVO process

Transesterification of fresh canola oil

The conditions for transesterification reaction were set at 400 kPa and 60°C. A conversion reactor, CRV-100, was used with the specified triolein conversion of 95%. 5250 kg/hr (5929 mol/hr) of fresh canola oil, 105-FVO, was pumped to 400 kPa by P-102 and heated to 60°C by E-101 before entering the reactor. Sodium hydroxide (NaOH), stream 102, was chosen as the alkali-catalyst. The amount of NaOH was 1 wt% of the canola oil and it was mixed with methanol before the reaction. The necessary amount of methanol was considered to be 6:1 molar ratio of methanol to oil and the reaction time was 1 hr (Meher et al., 2006). Since approximately 50% of the required amount of methanol was provided from the recycled stream from T-101, only 565.3 kg/hr (17640 mol/hr) of fresh methanol (stream 101) was fed to the process.

Methanol recovery

Due to the excess amount of methanol provided for the transesterification reaction, the product stream from CRV-100 contains a large amount of methanol that did not participate in the reaction. This methanol is recovered by a multi-stage distillation column and recycled back to the reactor. In the process simulation, 96% of methanol in stream 106 was recovered by using 6 theoretical stages in the distillation column. The pressures in the condenser and reboiler were set at 20.0 and 30.0 kPa, respectively, to keep the operating temperature below 200°C to avoid decompositions of methyl oleate and glycerol. As a result, the temperatures in the condenser and reboiler were 28.20 and 142.8°C, respectively.

Water washing

In the liquid-liquid extraction column, T-102, the bottom product stream from T-101 was washed with water at 110 kPa to get rid of glycerol and NaOH. The number of stages required for this separation was 4. The simulation results showed that glycerol was almost completely separated from methyl oleate using 50.0 kg/hr of water. Stream 203 also contained unreacted triolein from transesterification reaction, and 99.7% of the triolein was separated into stream 301. Since the triolein mass fraction in stream 301 was 0.0495, a distillation column, T-103, was used for the final biodiesel purification.

Neutralization and settler

Stream 302 from the liquid-liquid extraction column contains glycerol along with NaOH, which needs to be neutralized before glycerol purification. Stoichiometric amount of phosphoric acid

(H_3PO_4) was used to neutralize 52.5 kg/hr of NaOH. Water and trisodium phosphate (Na_3PO_4) were generated as reaction products. Na_3PO_4 was then separated as a precipitate in a decanter (X-100).

Biodiesel purification

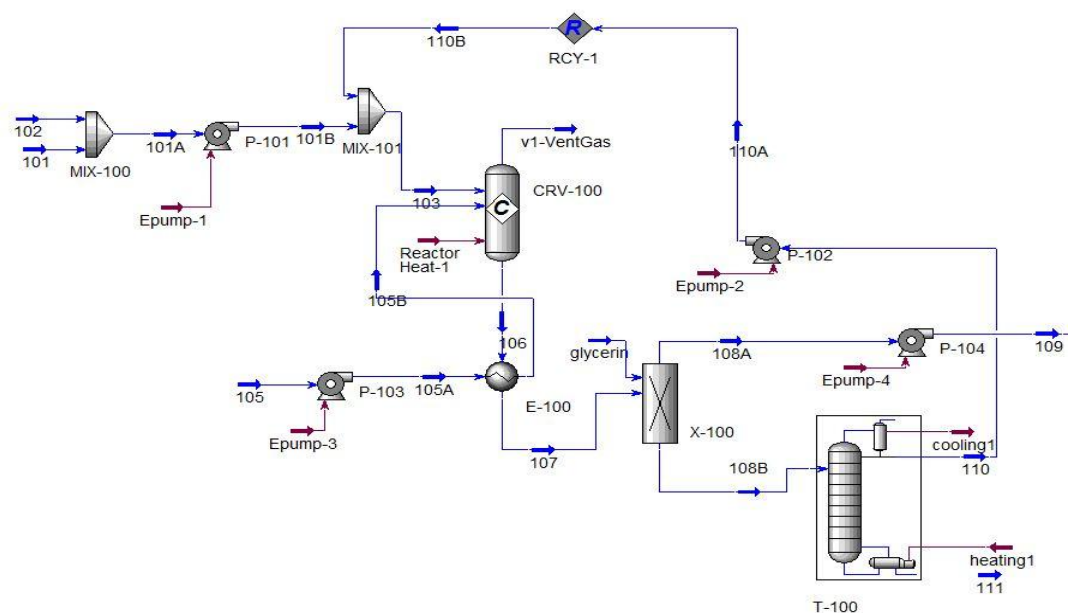
Column T-103 was designed for separating biodiesel from unreacted triolein, methanol and water. The simulated column had 6 theoretical stages in total, and the reflux ratio was 1.10. The pressures at the condenser and reboiler were 10.0 and 15.0 kPa, respectively. As a result, 96.4% of triolein was separated out in the bottom product stream. Stream 401 contained 99.6 wt% methyl oleate, 0.19 wt% triolein, 0.13 wt% methanol and 0.05 wt% water, which satisfied both European (EN14214) and American (ASTM D6751) biodiesel standards.

Glycerol purification

Glycerol was purified in the column T-102 to eliminate water and methanol contained in the feed stream. The required number of theoretical stages was 5, and the reflux ratio was 2.0. The pressures in the condenser and reboiler were set at 20.0 and 30.0 kPa, respectively, to maintain moderate operating temperature in the column to avoid decomposition of glycerol. The corresponding temperatures were 50.96 and 180.0°C in the condenser and reboiler, respectively. As a result, glycerol with high purity (99.5 wt%) was obtained as the bottom product of the distillation column.

2.3.2. Alkali-WVO process

Alkali-WVO process has additional pre-treatment steps before transesterification of waste vegetable oil with methanol. The pre-treatment process aims to decrease high FFA contents in WVO by esterification reaction in the presence of an acid catalyst. With the reduced amount of FFAs, undesirable soap formation by the reaction between FFAs and alkali catalysts is minimized. A flow diagram of the pre-treatment step is shown in Figure 2.4a.



Stream name	101	105	103	106	108A	108B	110	111
Pressure (kPa)	101.3	101.3	400.0	400.0	200.0	200.0	28.00	30.00
Temperature (°C)	25.00	25.00	34.16	70.00	46.24	46.24	35.07	68.93
Molar flow (gmol/hr)	3755	6720	40860	47580	7113	45680	36530	9150
Mass flow (kg/hr)	120.3	5275	1342	6617	5256	1841	1167	673.9
Volumetric flow (m ³ /hr)	0.1531	5.790	1.686	6.992	5.726	2.257	1.504	0.7049
Component mass fraction								
Methanol	1.000	0.000	0.958	0.189	0.002	0.672	0.997	0.110
TGA triolein	0.000	0.940	0.000	0.749	0.934	0.027	0.000	0.074
Oleic acid	0.000	0.060	0.000	0.000	0.000	0.000	0.000	0.000
Methyl oleate	0.000	0.000	0.000	0.050	0.062	0.002	0.000	0.005
Glycerol	0.000	0.000	0.000	0.000	0.001	0.258	0.000	0.705
H ₂ SO ₄	0.000	0.000	0.039	0.008	0.000	0.028	0.000	0.077
H ₂ O	0.000	0.000	0.003	0.004	0.000	0.013	0.003	0.029

Figure 2.4a. Flow diagram and properties of main streams of pre-treatment in Alkali-WVO process

Esterification of FFAs

Oleic acid, the major fatty acid in canola oil, was chosen to represent FFAs in WVO. It is reported that the amount of FFAs in WVO significantly varies depending on the source of the oil. In this study, 6 wt% FFAs in WVO was chosen (Zhang et al., 2003). The FFAs were esterified with methanol in the presence of 1 wt% sulphuric acid (based on oil) in CRV-100. The reaction was carried out at 70°C and 400 kPa at a 6:1 molar ratio of methanol to oil (Zhang et al., 2003). A complete conversion of FFAs to methyl oleate was specified in the conversion reactor, CRV-100.

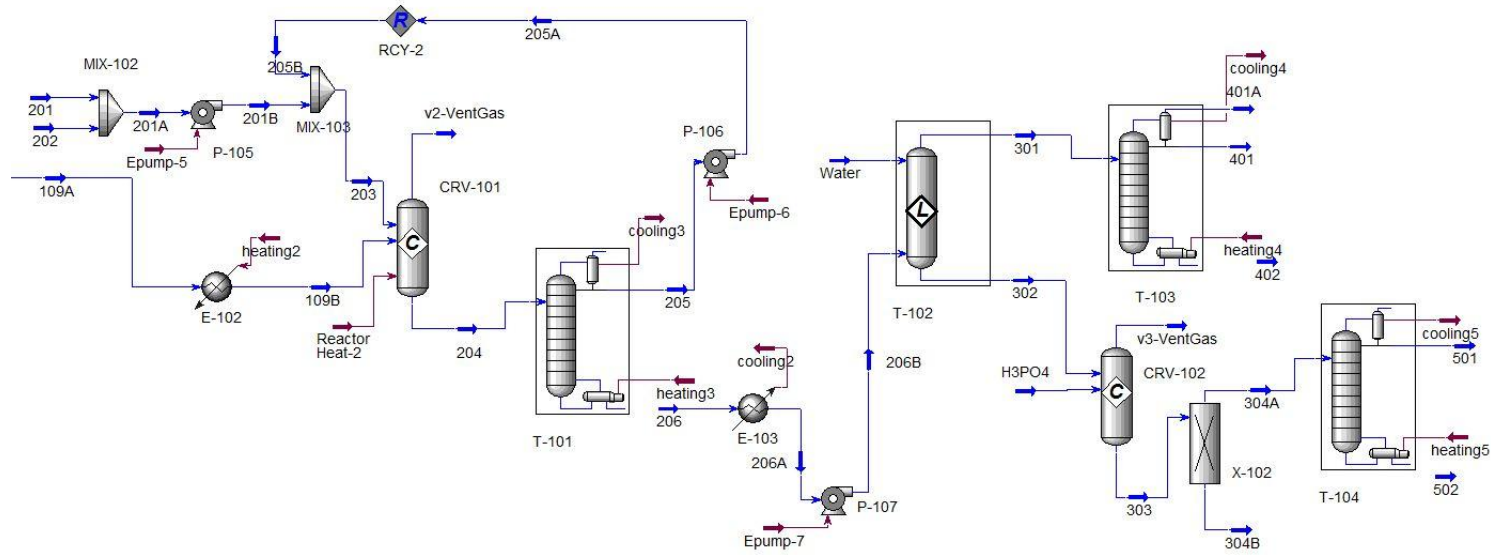
Glycerol washing

The reaction product stream consisted of triolein, methanol and sulphuric acid as well as methyl oleate and water generated from esterification reaction. A liquid-liquid extraction column was originally designed to separate an organic phase (triolein and methyl oleate) from an aqueous phase (methanol, sulphuric acid and water) by glycerol washing (Zhang et al., 2003; West et al., 2008). However, simulation results of the extraction column showed very unstable behaviour. For example, the extraction column was converged as the flow rate of glycerol varied from an extremely large amount to smaller amount. However, it was not possible to converge it using the exactly same amount of glycerol when the flow rate varied from a small amount. Therefore, a component splitter, X-100, was used instead of a liquid-liquid extraction column due to the lack of knowledge concerning the behaviour of the electrolytes such as sulphuric acid in the liquid-liquid extraction column (Zhang et al., 2003). The split fraction for all components was assumed to be 0.99. Triolein in stream 108A was then sent to CRV-101 for transesterification reaction.

Methanol recovery

A distillation column, T-100, was used to recover 94% of methanol in the feed stream using 5 theoretical stages and a reflux ratio of 5. The pressures in the condenser and reboiler were set at 28.0 and 30.0 kPa, respectively. Stream 110 containing 99.7 wt% methanol and 0.3 wt% water was then mixed with fresh methanol feed stream and sent back to the esterification reactor. The bottom stream contained 70.5 wt% glycerol along with 11.0 wt% methanol, 7.75 wt% sulphuric acid and trace amounts of water and methyl oleate. This stream was treated as a waste stream due to the presence of sulphuric acid.

After the pre-treatment of WVO, the next process flow diagram is identical to that of Alkali-FVO process. The stream table is shown below the process flowsheet in Figure 4b.



Stream name	201	109A	203	204	205	206	301	401	304A	502
Pressure (kPa)	101.3	400.0	400.0	400.0	20.00	30.00	110.0	10.00	101.3	30.00
Temperature (°C)	25.00	46.41	26.73	60.00	28.20	156.7	59.30	85.86	158.5	180.1
Molar flow (gmol/hr)	15930	7108	34130	41240	16950	24290	17560	17020	6321	5431
Mass flow (kg/hr)	510.6	5256	1104	6360	542.9	5817	5269	5002	507.3	490.6
Volumetric flow (m ³ /hr)	0.6495	5.722	1.398	6.915	0.7431	40.23	6.192	6.054	4.956	0.4933
Component mass fraction										
Methanol	1.000	0.002	0.955	0.088	1.000	0.003	0.001	0.001	0.003	0.000
TGA triolein	0.000	0.934	0.000	0.039	0.000	0.042	0.046	0.002	0.000	0.000
Methyl oleate	0.000	0.062	0.000	0.788	0.000	0.862	0.952	0.997	0.000	0.000
Glycerol	0.000	0.001	0.000	0.077	0.000	0.084	0.000	0.000	0.962	0.995
NaOH	0.000	0.000	0.045	0.008	0.000	0.009	0.000	0.000	0.000	0.000
H ₃ PO ₄	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
H ₂ O	0.000	0.000	0.000	0.000	0.000	0.000	0.001	0.000	0.035	0.005
Na ₃ PO ₄	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000

Figure 2.4b. Flow diagram and properties of main streams of Alkali-WVO process after pre-treatment

2.3.3. SC-WVO process

As stated previously, a supercritical process is not sensitive to the presence of FFAs and water in feedstocks. Therefore, waste vegetable oil was chosen as the raw material in this process. The composition of WVO was identical to the one in Alkali-WVO process. SC-WVO process basically consisted of 4 steps: heating and pressurizing of raw materials, transesterification and esterification reactions, methanol recovery and biodiesel purification. The flow diagram of SC-WVO process is shown in Figure 2.5.

Heating and pressurizing of raw materials

Methanol (stream 101) and WVO (stream 103) were pressurized to the reaction pressure (19 MPa) by P-101 and P-102, respectively. As pressurized, the temperature of the methanol stream went up to 71.9°C and the one of the WVO stream to 41.9°C. Then each pressurized stream flowed through heat exchangers (E-101 and E-102) where the heat of reaction product stream was utilized to warm up the raw materials. The reaction product stream firstly heated WVO to 293.9°C and then heated methanol to 211.0°C. The raw material streams were then mixed together in a mixer and brought to the desired temperature of 350°C by E-103.

Biodiesel production in a plug flow reactor (PFR)

In large-scale production, continuous reactors like PFRs are financially advantageous. They have a high conversion per unit volume, and do not need to be shut down and cleaned as regularly as batch reactors. Therefore, for continuous biodiesel production, a PFR was selected to carry out transesterification and esterification reactions. It is, however, noted that research on the fundamental dynamics and thermophysical properties of a mixture of the reactants and products under supercritical conditions are rare in the present stage (Chen et al., 2009).

Using a PFR in Hysys requires information about reaction kinetics such as a pre-exponential factor and activation energy of the Arrhenius equation. There are a few kinetic studies on supercritical transesterification reaction. Kusdiana and Saka (2001) reported reaction rate constant values at different temperatures based on their experimental results from rapeseed oil transesterification in a batch reactor. The Arrhenius plot of their results in supercritical methanol state gives around 47.1 kJ/mol for the activation energy and 145.87 s^{-1} for the pre-exponential factor. Varma and Madras (2007) reported that the activation energy of transesterification reaction between linseed oil and methanol was 46.5 kJ/mol and the pre-exponential factor was 119.3 s^{-1} .

Technically, a PFR together with a proper reaction kinetic data can be used in Hysys to find out the reaction conditions and reactor volumes to achieve desired reaction conversions. However, a

great care should be taken to directly use the kinetic data in literatures to simulate transesterification in a PFR. As one of the reasons, there is still a big discrepancy in the reaction conditions and the corresponding biodiesel yields among researchers. Table 2.5 shows the reaction conditions and yields as well as the type of reactor used for supercritical transesterification reaction. The optimal reaction pressures and reaction times to achieve relatively similar yields are significantly different among the researchers.

Another reason is that the thermal decomposition of FAMES at high temperatures is not taken into account in the reaction kinetics. While high temperature does increase the reaction rate, it can also cause the loss of FAMES in the products. Imahara et al. (2008) reported that FAMES become unstable and vulnerable to decomposition at high reaction temperature and pressure. Especially, poly-unsaturated FAMES like methyl linoleate (C18:2) and methyl linolenate (C18:3) were more prone to decomposition than mono-saturated and saturated FAMES. Their decomposition was pronounced at temperatures above 300°C. For example, after 20 min exposure to supercritical methanol at 350°C/43MPa, only 20% of methyl linolenate remained. Therefore, applying transesterification reaction kinetics to a PFR without considering thermal decomposition of FAMES would probably result in erroneous conversions.

In this study, the reaction conditions and biodiesel yield were obtained from Bunyakiat et al. (2006) where the reaction was carried out in a continuous method using a 5.5 m in length with 3/8-in. (9.525 mm) o.d. and 0.035-in. (0.889 mm) thickness 316SS tubing. By matching the experiment results shown in Table 2.5, 96% conversion of triolein was simulated in the PFR at 350°C and 19 MPa with 7 min of reaction time. The molar ratio of methanol to oil was 42 and the reactor was considered to be isothermal.

Table 2.5. Raw materials, reaction conditions, yields and reactor types for supercritical biodiesel production

Oil	T (°C)	P (MPa)	MeOH/oil ^a	τ (min)	B/C ^b	Yield (%)	Ref.
Sunflower	400	20	40	40	B	96	Madras et al. (2004)
Rapeseed	350	43	42	4	B	95	Kusdiana and Saka (2001)
Jatropha	320	8.4	43	4	B	100	Hawash et al. (2009)
Soybean	310	32	40	25	C	96	He et al. (2007)
Castor/ linseed	350	20	40	40	B	98	Varma and Madras (2007)
Palm	350	19	42	7	C	96	Bunyakiat et al. (2006)

^a Molar ratio

^b B-batch or C-continuous

Table 2.6. Summary of reactions involved in simulation models

	Alkali-FVO	Alkali-WVO	SC-WVO	SC-LowE
Reactions	transesterification	esterification/ transesterification	simultaneous transesterification and esterification	simultaneous transesterification and esterification
Catalyst	NaOH	H ₂ SO ₄ /NaOH	-	-
Temperature (°C)	60	70/60	350	350
Pressure (kPa)	400	400/400	19000	19000
Reaction time (min)	60	60/60	7	13.3
Methanol:oil (molar)	6:1	6:1/6:1	42:1	24:1
Conversion (%)	95	100/95	96 ^a	96 ^a
Reactor type	CSTR	CSTR/CSTR	PFR	PFR

^a For triolein. The conversion of oleic acid is 100%.

Methanol recovery

A distillation column with 4 theoretical stages was used to separate unreacted methanol. The pressures in the condenser and reboiler were at 101.5 and 106.5 kPa, respectively, and the corresponding temperatures were 65.49 and 164.1°C. Simulation result showed 99.93% recovery of methanol. The recovered methanol contained 96.1 wt% methanol and 3.9 wt% water generated from esterification reaction. It was then mixed with fresh methanol feed of 535.1 kg/hr.

Glycerol separation

After most of methanol is recovered in the distillation column, the stream was cooled down to 25°C and fed to a decanter. Two liquid phases were formed: a glycerol-rich phase and a methyl oleate-rich phase. The denser phase (glycerol-rich) consisted of 95.6 wt% glycerol, 3.96 wt% water and 0.43 wt% methanol. The separation result showed that 99.9% of glycerol in stream 109A was drained out with the denser phase in this unit.

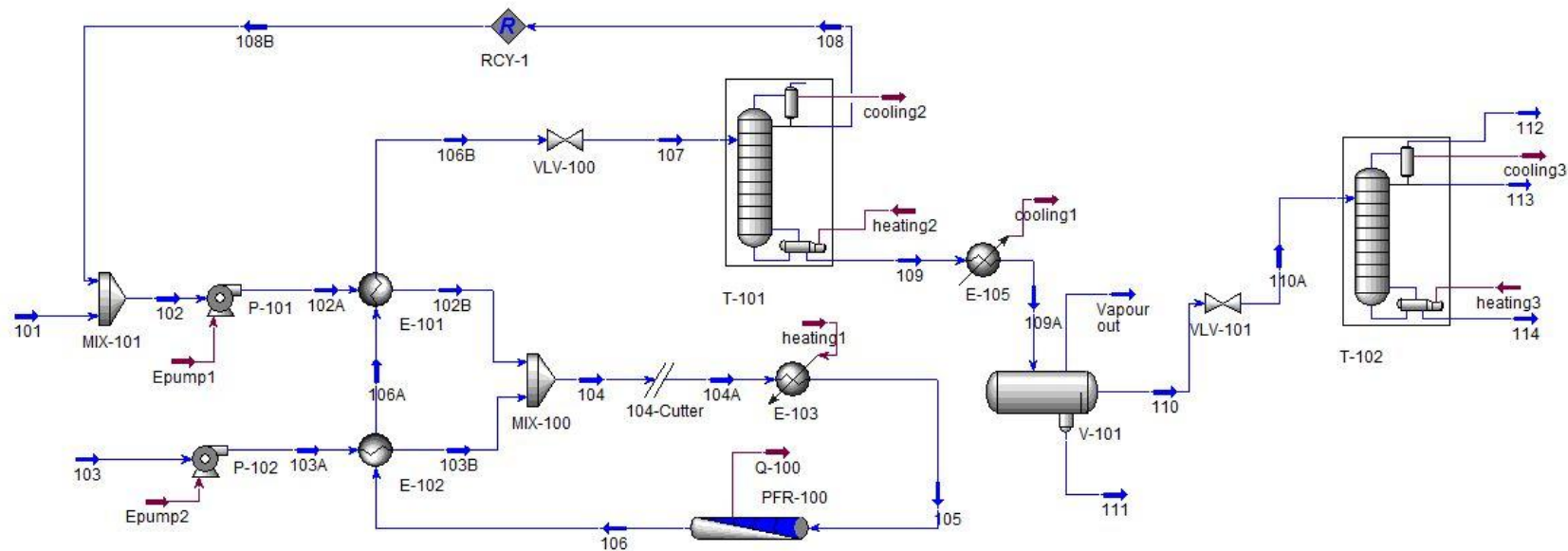
Negi (2006) reported mole fraction of glycerol in methyl oleate at different temperatures after stirring the binary system at high speed for 6 hours. A sample from the top layer was taken when two phases were clearly separated after settling. As the sample was analyzed by gas chromatography, less than 0.0025 mole fraction of glycerol was detected in the methyl oleate phase. This value was similar to the simulation result where the mole fraction of glycerol in the methyl oleate-rich phase was 0.0009. This phase was then sent to a distillation column for the final purification of biodiesel.

Biodiesel purification

The feed stream consisted of 96.2 wt% methyl oleate, 3.73 wt% triolein, 0.068 wt% methanol and 0.027 wt% glycerol. According to European (EN 14214) and American (ASTM D 6751) biodiesel standards, the weight percent of free glycerol is limited to ≤ 0.02 wt%. In addition, the amount of triglyceride in biodiesel is limited to ≤ 0.20 wt% in EN 14214. Therefore, further purification of the feed stream was necessary to satisfy both EN 14214 and ASTM D 6751.

Since the boiling temperatures of glycerol and methyl oleate are over 280°C , a high operating temperature was expected in the distillation column at atmospheric operating pressure. Therefore, the pressure inside T-102 was set as low as 3.0–5.0 kPa in order to avoid thermal decompositions of methyl esters. The temperatures in the condenser and reboiler were 154.8 and 264.1°C , respectively. The number of theoretical stages was 7.

Simulation result showed that the top product stream satisfied both of the biodiesel standards. The weight percents of methyl oleate, triolein and glycerol in stream 113 were 99.8%, 0.18% and 0.01%, respectively. Therefore, stream 113 was considered as the final product stream.



Stream name	101	103	104A	106	108	109	110	111	113	114
Pressure (kPa)	101.3	101.3	19000	19000	101.5	106.5	170.0	170.0	3.00	5.00
Temperature (°C)	25.00	25.00	224.3	350.0	65.49	164.1	25.00	25.00	154.8	264.1
Molar flow (gmol/hr)	16700	6577	298300	298300	274600	23660	17230	6425	16860	255.0
Mass flow (kg/hr)	535.1	5162	14250	14250	8537	5709	5204	505.3	5000	198.9
Volumetric flow (m ³ /hr)	0.6808	5.666	26.04	52.74	11.45	7.299	5.946	0.4556	6.462	0.2658
Component mass fraction										
Methanol	1.000	0.000	0.614	0.576	0.961	0.001	0.001	0.004	0.000	0.000
TGA triolein	0.000	0.940	0.341	0.014	0.000	0.034	0.037	0.000	0.002	0.932
Oleic acid	0.000	0.060	0.022	0.000	0.000	0.000	0.000	0.000	0.000	0.000
Methyl oleate	0.000	0.000	0.000	0.351	0.000	0.877	0.962	0.000	0.998	0.068
Glycerol	0.000	0.000	0.000	0.034	0.000	0.085	0.000	0.956	0.000	0.000
H ₂ O	0.000	0.000	0.023	0.025	0.039	0.003	0.000	0.040	0.000	0.000

Figure 2.5. Flow diagram and properties of main streams of SC-WVO process

2.3.4. SC-LowE process

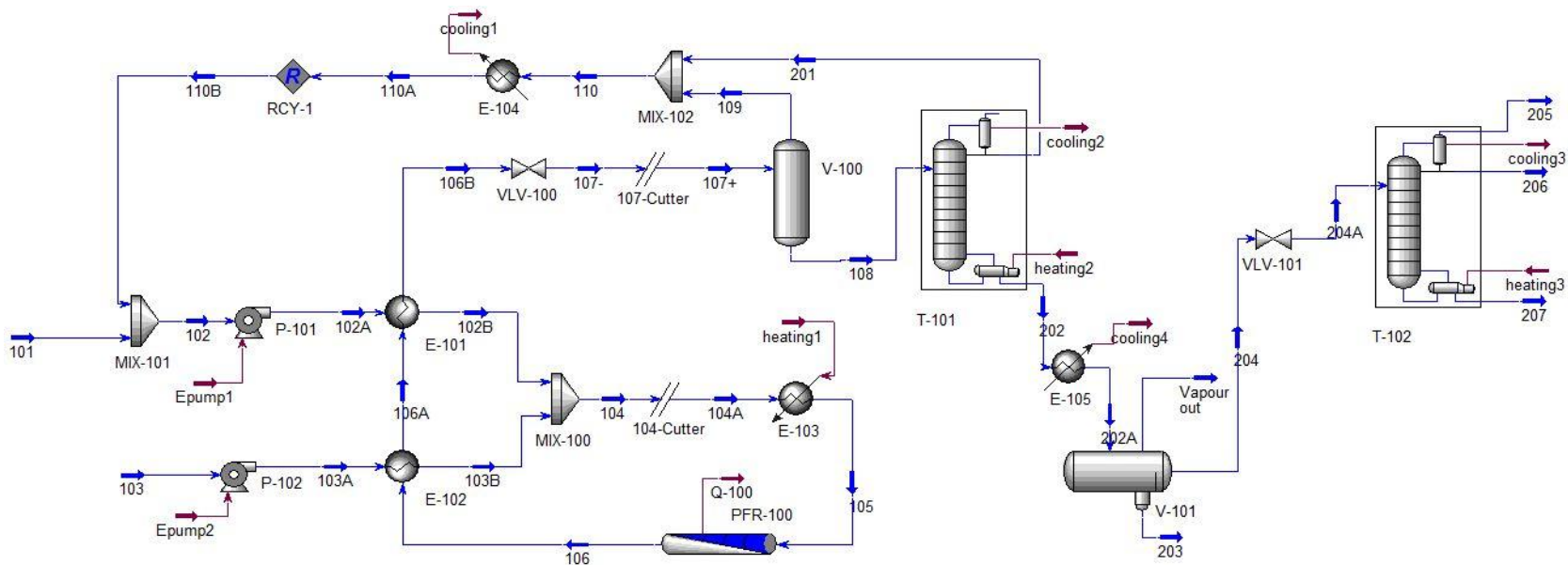
In SC-WVO process, excess amount of methanol is used to achieve better miscibility of methanol and vegetable oil. However, the large amount of methanol can result in high energy consumption in pumping and heating. For example, in Figure 2.5, the flow rate of methanol affects the energy streams of Epump1, heating 1 and heating 2. The energy consumptions are 304.5, 7501 and 24190 MJ/hr, respectively, with their sum accounting for 84.7% of the total energy consumption. In attempt to reduce the energy consumption, two features were applied in SC-LowE process: a moderate amount of methanol usage and initial methanol recovery in a flash evaporator, as shown in Figure 2.6.

Biodiesel production in PFR

Bunyakiat et al. (2006) reported that it was possible to obtain the same conversion (96%) using 24:1 methanol to oil molar ratio instead of 42:1. The corresponding reaction time increased from 420 second to 800 second to achieve the same conversion as the molar ratio decreased. Therefore, 96% conversion of triolein was simulated at 350°C and 190 MPa using 24:1 methanol to oil molar ratio as shown in Table 2.6. Using less amount methanol in the process has several advantages. It reduces the cost of raw materials as well as necessary utilities. In addition, this would result in shorter separation and evaporation steps compared to higher methanol to oil ratios, especially in a large industrial-scale plant (Bunyakiat et al., 2006). In light of the facts, using smaller molar ratio seems to be favourable despite the increased residence time.

Methanol separation in a flash evaporator

Product stream 106 consists of 41.3 wt% methanol, 48.0 wt% methyl oleate, 5 wt% glycerol and trace amounts of unreacted triolein and water. Since the boiling point of methanol is far lower than the rest of the components, a significant amount of methanol can be separated out as a vapour in a flash evaporator (V-100) by adjusting the pressure or temperature. VLV-100 was used to depressurize stream 106B from 19 to 0.2 MPa so that most of methanol evaporates whereas the other components remain mostly in the liquid phase. Using this procedure, 89.6% of methanol in the feed stream was separated into stream 109.



Stream name	101	103	104A	107+	108	109	201	202	203	204	206	207
Pressure (kPa)	101.3	101.3	226.6	200.0	200.0	200.0	101.5	106.5	170.0	170.0	3.000	5.000
Temperature (°C)	25.00	25.00	19000	96.50	96.50	96.50	68.02	162.8	25.00	25.00	198.6	265.0
Molar flow (gmol/hr)	16000	6577	181300	181300	41960	139300	18250	23716	6586	17130	16850	250.0
Mass flow (kg/hr)	512.7	5162	10430	10433	6232	4201	524.1	5708	507.6	5201	5000	197.3
Volumetric flow (m³/hr)	0.6522	5.691	14.54	2148	7.504	7.299	0.6813	7.296	0.4641	5.946	6.766	0.2639
Component mass fraction												
Methanol	1.000	0.000	0.465	0.413	0.072	0.919	0.851	0.000	0.000	0.000	0.000	0.000
TGA triolein	0.000	0.940	0.465	0.019	0.031	0.000	0.037	0.034	0.000	0.037	0.002	0.939
Oleic acid	0.000	0.060	0.030	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
Methyl oleate	0.000	0.000	0.000	0.480	0.803	0.000	0.000	0.877	0.000	0.962	0.998	0.061
Glycerol	0.000	0.000	0.000	0.046	0.078	0.000	0.000	0.085	0.953	0.000	0.000	0.000
H₂O	0.000	0.000	0.040	0.042	0.016	0.081	0.149	0.004	0.047	0.000	0.000	0.000

Figure 2.6. Flow diagram and properties of main streams of SC-LowE process

2.3.5. Energy consumption

Energy consumptions of the four process simulations are analyzed for the comparison of energy intensity of the processes as well as each of the unit operations. Table 2.7 shows the energy consumptions of the simulation models at the annual capacity of 40,000 tonnes biodiesel production.

Table 2.7. Energy consumption of the four process simulation models (unit: kW)

	Alkali-FVO	Alkali-WVO	SC-WVO	SC-LowE
Pumps	1.043	1.688	124.23	87.73
Heaters	83.45	34.05	2084	1429
Distillation columns (Reboiler)				
Methanol recovery	664	2279 (T-100) 873.8 (T-101)	6718	792.7
Biodiesel purification	2069	2033	1917	1618
Glycerol purification	31.92	36.78	-	-
Total	2349	5258	10843	3927

As a result, Alkali-FVO had the lowest energy consumption among the simulation models. The biodiesel purifying distillation column was the most energy intensive process in the model, accounting for 88% of the total energy usage. Alkali-WVO consumed more energy than Alkali-FVO because of the high energy consumption of the methanol distillation columns. The first methanol distillation column, T-100, is located in the pre-treatment step, and it consumes 2279 kW which is nearly half of the total energy consumption. This is much larger compared to the energy usage in T-101 which separates methanol from the transesterification reaction products. The reason for the prominent difference in the energy consumption of the two distillation columns is derived from the methanol content in the column feed streams. Stream 108B in the pre-treatment step contains 1237 kg/hr of methanol in the total flow rate of 1841 kg/hr, whereas stream 204 has 560 kg/hr of methanol out of 6360 kg/hr of the stream flow rates. The difference in the methanol content is the consequence of the methanol usages in the reactors. The methanol is fed at 6:1 methanol to oil molar ratio to both the esterification (CRV-100) and transesterification reactors (CRV-101). However, in the pre-treatment process, the amount of oleic acid for esterification reaction is very small and only 1 mol of methanol is consumed by 1 mol of oleic acid. This explains the high methanol content in stream 108B. On the other hand, in the transesterification reaction, 3 mols of methanol are used by 1 mol of triolein, and it results in lower amount of unused methanol in stream 204.

In comparison with alkali-catalyzed processes, the two supercritical processes had a large energy requirement for pumping and heating of raw materials to achieve the harsh reaction conditions.

Despite this fact, the energy consumption in SC-LowE process was the second lowest among the simulation models. Comparing the two supercritical models, SC-LowE indicated much lower energy consumption than SC-WVO. In SC-WVO process, the methanol distillation column was the most energy intensive process, accounting for 62% of the total energy consumption. Other than the fact that a distillation column itself is a very energy consuming operation, because of the high molar ratio of methanol to oil in this process, the methanol column used a significant amount of energy. On the other hand, the methanol distillation column in SC-LowE consumed 792.7 kW, which was only 11.8% of the energy usage in the methanol column in SC-WVO. This is because, as mentioned before, a large portion of methanol was separated in the flash evaporator. Another reason is the lower methanol to oil molar ratio used in this process. The reduced flow rate because of the lower methanol usage also decreased the energy consumption in pumps and heaters. Taking these into account, SC-LowE would benefit from not only energy saving but also from the reduced sizes of the methanol distillation column, pumps and heaters.

2.4. Conclusion

Thermogravimetric method was applied to determine the normal boiling point of triolein. From the five TGA measurements, the average boiling point was defined as 412.8°C. The boiling point of tripalmitin found in this study was consistent with the previously published literature. A new component, “TGA triolein”, was created in Hysys using the boiling point value and UNIFAC molecular structure. Its properties were compared with the triolein properties provided by the component library in Hysys. The results showed significant improvements in accuracy of triolein properties when TGA boiling point was applied.

Four simulation models were developed using TGA triolein as one of the components. Each process had certain advantages and disadvantages. For example, the conventional alkali-catalyzed process used fresh vegetable oil, which made it less favourable process in terms of economical and ethical points of views. However, it used much less energy and less methanol compared to alkali-catalyzed process using waste vegetable oil. Alkali-WVO process had moderate energy usage and used low cost feedstock. However, the process flowsheet was the most complicated among the process models. On the other hand, both supercritical processes used waste vegetable oils, and the processes were less complex due to the absence of a catalyst. To decrease energy consumption involved with a supercritical process, SC-LowE model was developed. The model featured a 24:1 methanol to oil

molar ratio and a flash evaporator to primarily recover methanol. These two features led to significant decrease in energy usage in the supercritical process.

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3. Economic assessment of biodiesel production processes²

3.1. Introduction

Over the past few decades, the world's energy consumption has rapidly risen due to increasing population and industrialized economy. To meet the increasing energy demand, the world has relied mostly on fossil fuels. One of the biggest problems of fossil fuels, however, is that they are not renewable. It is anticipated that the global conventional oil peak is 5-10 years away from now (Bentley, 2002). Another problem is that the use of fossil fuels has caused environmentally negative effects such as global warming. In addition, the production of fossil fuel is affected by political and economic issues because of the locations of reservoirs and the finite quantity of oil. These drawbacks have driven people to look for alternative energy resources.

Biodiesel has many advantages over conventional petroleum-derived diesel fuel. It is biodegradable, thus more environmentally friendly, and renewable. Compared to the combustion of petroleum diesel, the combustion of biodiesel emits less CO, SO_x, particulate matter and unburned hydrocarbons. According to Yee et al. (2009), utilization of palm biodiesel would generate an energy yield ratio (output energy/input energy) of as high as 3.53, meanwhile absorbing higher amounts of CO₂ from atmosphere than the amount of CO₂ emission from production and consumption. Biodiesel has a relatively high flash point (150°C), making it less volatile and therefore safer to transport and handle than petroleum diesel. Furthermore, it can be used directly in diesel engines with little modifications and provides additional lubrication that can reduce engine wear and extend engine life.

Biodiesel is commonly produced from food-grade vegetable oils using an alkali-catalyzed transesterification process. However, using food-grade oils is becoming economically not feasible due to the increasing prices of crops. The prices of vegetable oils have been consistently rising over the last few years as shown in Figure 3.1. As a result, low cost feedstocks such as waste vegetable oils (WVOs) from restaurants and animal fats are receiving attentions as potential feedstocks for biodiesel production.

² A version of this chapter is in preparation for submission for publication. Lee, S., Posarac, D. and Ellis, N. (2010). Economic assessment of biodiesel production processes.

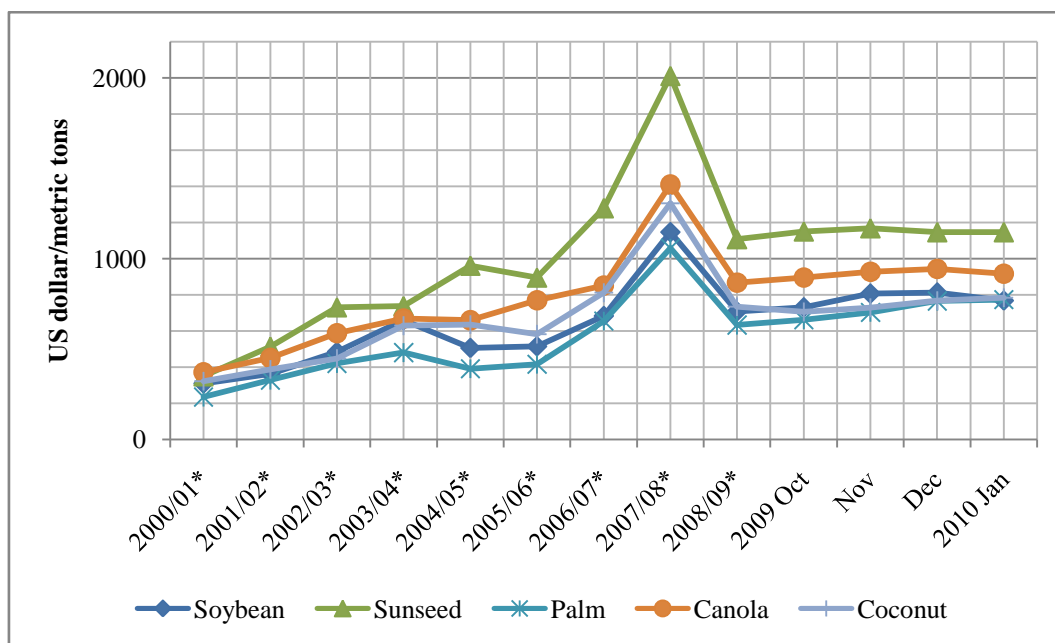
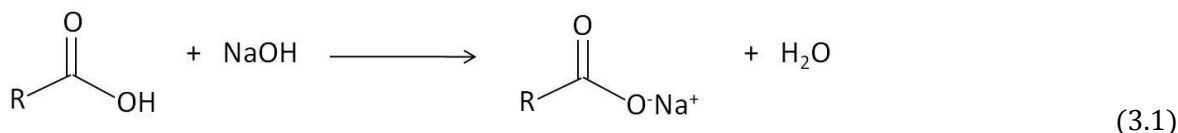


Figure 3.1. The prices of vegetable oils from Oct 2000 to Jan 2010 (Source: USDA (2010a))
 * Average from October to the following September.

The properties of WVOs are related to the compositions of the fresh vegetable oils (FVOs) and various types of processes where it has been generated. In food frying, vegetable oils are used at high temperatures ($>190^{\circ}\text{C}$) which cause various chemical reactions such as hydrolysis, polymerization and oxidation during conversion. The free fatty acids (FFAs) content was found to increase due to the hydrolysis of triglycerides in the presence of food moisture and oxidation (Canakci, 2007). For example, the FFA level of fresh soybean oil changed from 0.04% to 1.51% after 70 hrs of frying at 190°C (Tyagi and Vasishtha, 1996). The increases in viscosity were also reported due to polymerization, which resulted in the formation of higher molecular weight compounds (Canakci, 2007). In the same paper, it was reported that the levels of FFAs and water content of the waste restaurant oils and animal fats varied widely, being as high as 18% and 41.8%, respectively. Therefore, a biodiesel production process must be able to tolerate a wide range of feedstock properties in order to make use of waste vegetable oils.

Unfortunately, the feedstocks with high FFAs and water contents are known to give lower yields in the conventional alkali-catalyzed transesterification reaction due to the saponification reaction. As shown in Equation (3.1), the soap formation partially consumes the catalysts, decreasing biodiesel yields. It also complicates the separation and purification steps. Therefore, pre-treatment is necessary for the low cost feedstocks to eliminate FFAs and water before starting transesterification reaction.



Supercritical methanol transesterification is one of the ways to utilize those feedstocks. Kusdiana and Saka (2004) reported that almost complete conversion was achieved using supercritical methanol and vegetable oil containing FFAs up to 30 wt%. In comparison with the supercritical method, vegetable oil with 20 wt% FFAs resulted in 35% conversion in the alkali-catalyzed method. In addition, the yield of methyl esters from the supercritical reaction was not affected by 30 vol. % water contents in the feedstock.

There have been studies on economic feasibility of biodiesel production processes, especially regarding the types of feedstocks and the types of catalysts in transesterification reaction. Bender (1999) reviewed 12 economic feasibility studies for the projected biodiesel production costs involving several feedstocks and operational scales. The author reported that the production costs ranged from US\$0.30/litre for soybean-based biodiesel to US\$0.69/litre for rapeseed-based biodiesel. The rough projected costs of biodiesel from vegetable oil and waste grease were estimated in the range of US\$0.54–0.62/litre and US\$0.34–0.42/litre, respectively, indicating the great advantage of using waste grease.

Zhang et al. (2003) presented an economic comparison of the four continuous processes using acid and alkali catalysts in FVOs and WVOs. Based on the after-tax rate of returns, it was determined that the alkali treatment worked best for the fresh oil, but the acid-catalyzed process, even with much longer process time, worked best for the WVO. The alkali transesterification process for WVO was not cost effective, primarily because esterification pre-treatment steps were required to reduce the FFA contents. A supercritical process, however, was not investigated in their study.

Lim et al. (2009) conducted the economic analysis of processes for supercritical biodiesel production from rapeseed oil and compared the profitability to the alkali-catalyzed process. The results showed that the total manufacturing cost of the supercritical process was lower than the alkali-catalyzed process due to the absence of catalyst and higher by-product (glycerol) credit. Despite the higher total capital cost of the supercritical process, its lower manufacturing cost resulted in shorter payout time than the alkali-catalyzed process. The faster payout time of the supercritical processes even without using waste vegetable oil feedstocks indicates the advantage of a supercritical process.

Further investigation on the supercritical biodiesel production from WVO was reported by Kasteren and Nisworo (2007). In this study, propane was used as a co-solvent to alleviate harsh

supercritical reaction conditions. Their reaction conditions were 280°C/12.8 MPa with 24:1 methanol to oil molar ratio. As a result, the required selling prices for biodiesel were predicted to be US\$ 0.17/litre, US\$0.24/litre and US\$0.52/litre at the plant capacities of 125,000, 80,000 and 8,000 tonnes biodiesel/yr, respectively. This fact indicates the plant capacity plays a significant role in determining the profitability of the process.

This thesis deals with four kinds of continuous biodiesel production processes using fresh and waste vegetable oils as the feedstocks have been investigated: alkali-catalyzed using FVO (Alkali-FVO), alkali-catalyzed with pre-treatment of WVO (Alkali-WVO), supercritical using WVO (SC-WVO) and supercritical using WVO with reduced energy consumption (SC-LowE) based on 40,000 tonnes of biodiesel production per year.

To our knowledge, no economic analysis of a biodiesel plant has been done using a software program designed for process evaluation. Previous studies on economic estimations were based on the overall factor method of Lang or bare-module concept introduced by Guthrie (Zhang et al., 2003; Kasteren and Nisworo, 2007; You et al., 2008; West et al., 2008; Lim et al., 2009; Santana et al., 2010). On the other hand, Aspen Icarus software systems have been field-tested for more than 30 years on commercial plants and are in use worldwide by owner companies and engineering design and construction firms (Seider et al., 2004).

The most accurate approach in estimating equipment costs would be using quotations directly from vendors. However, particularly in the early stages of the cost estimation procedure, it is often hard to obtain sufficient information or detailed quotes. The advantage of using Icarus Process Evaluator (IPE) is that it provides the necessary specifications for detailed design, estimation and economic data. The detailed design allows detailed modifications of the process equipments, which is not possible in Lang factor or bare-module method. In addition, more accurate estimation of a total capital investment can be achieved by the provided database of the field-tested industrial standard design as well as cost modeling used by project evaluators.

Therefore, AspenTech's IPE was used in this study to provide improved accuracy of economic analysis of a biodiesel plant by applying the accumulated data in the software systems. The scheme of the work for the economic analysis in this study is as follows:

- Loading Hysys simulation results into IPE
- Setting general specifications including location and starting date of a plant, process description and complexity, plant type (e.g., grass roots), currency, necessary utilities, designating raw material and product streams
- Relating each unit in simulation to a specific type of process equipments. For example, a heater in Hysys simulation model is specified as a floating head heat exchanger in IPE.

- Setting general investment parameters including the prices of raw materials, products and utilities, tax rates, economic life of project, salvage value, depreciation method, operating hours per period and labour costs.
- Evaluation of total capital investment, total manufacturing cost and project profitability
- Identification of factors greatly contributing to the costs for biodiesel and sensitivity analysis of profitability to the factors.

3.2. General specifications for economic analysis in IPE

3.2.1. General specifications for capital costs

The general specifications defined here influence the design and cost estimate by defining system defaults and environmental variables.

First of all, a project is to be defined as one of the five categories in Process Description: new and unproven process, new process, redesigned process, licensed process and proven process. Depending on which category a project belongs to, the contingency percent is adjusted accordingly. For example, the new and unproven process makes the contingency value high compared to the proven process. Process Description also drives the design allowances for all equipment whose material cost is system-generated. Therefore, a higher design allowance is made for a new and unproven process compared to a proven one. A proven process was chosen for Alkali-FVO and Alkali-WVO and a new process was chosen for SC-WVO and SC-LowE.

Process Complexity option is used to adjust contingency percent along with Process Description. It has five sub-categories: highly complex, slightly complex, typical, somewhat simple and simple. Highly complex implies high temperature/pressure and also more controllers and instrumentation. Therefore, highly complexity was chosen for the supercritical processes and typical was chosen for the alkali-catalyzed processes.

In Project Location, the geographical location of the project site is defined and it influences various location dependent costs such as freight (domestic and ocean), taxes, wage rates and workforce productivities. For example, the domestic and ocean freights and taxes are specified as the different percentages of materials costs depending on the locations as shown in Table 3.1. All the four processes were defined to be located in North America, with the currency used in this study is in U.S. dollar.

Table 3.1. The freights (domestic and ocean) and taxes as the percentages of material costs in different locations (units: %)

	North America	South America	Central America	Europe	Asia	Africa
Domestic Freight	4	5	5	1	1	4
Ocean Freight	0	8	5	0	0	8
Taxes	6.25	4.00	4.00	0.00	6.00	4.00

A new chemical processing plant could be a grass-roots plant, with no other chemical plants nearby, or could be an addition to an existing integrated complex. For a grass-roots plant, it is required to construct auxiliary facilities including utilities, waste treatment and road facilities as well as other new facilities such as a cafeteria and a maintenance shop (Seider et al., 2004). In the integrated complex, the auxiliary facilities can be shared among the various plants in the complex (Seider et al., 2004). The electrical power distribution and process control systems can also be affected by the type of the plant. IPE allows five options to choose a Project Type: grass roots/clear field, plant addition-adjacent to existing plant, plant additions-inside existing plant, plant addition-suppressed in infrastructure and plant modification/revamp. In this study, a grass-roots plant was chosen for all four processes.

Contingencies account for uncertainty in the estimate and the possibility of not accounting for all of the costs involved (Seider et al., 2004). It is expressed as the percentage of fixed capital cost and influenced by the combination of Process Description, Process Complexity and Project Type. With processes for which a company has considerable experience, the cost of contingency is much lower than when the processes are new and just discovered by a research group (Seider et al., 2004). With the specifications defined earlier, the contingency was set at 18.0% for the alkali-catalyzed processes and 32.5% for the supercritical processes. A summary of general specifications for capital cost estimation is shown in Table 3.2.

Table 3.2. The general specifications for capital costs of the four biodiesel production processes

	Alkali-FVO	Alkali-WVO	SC-WVO	SC-LowE
Process Description	proven process	proven process	new process	new process
Process Complexity	typical	typical	highly complex	highly complex
Process Control			digital	
Project Location			north America	
Currency			U.S. dollar	
Project Type			grass roots	
Contingency (%)	18.0	18.0	32.5	32.5
Estimated Start date of Basic Engineering			2009/01/01	
Soil condition around site			soft clay	
Pressure vessel design code			ASME ^a	

^a American Society of Mechanical Engineers, Section VIII, Division 1

3.2.2. Utility, raw material and product specifications

The list of the utilities used in this study is shown in Table 3.3. Steam is a typical heat source to provide necessary energy to heat up process streams. For large chemical plants, steam is often generated at the highest pressure level and then let down to the lower pressures through turbines or valves (Turton, 2003). The highest-pressure steam is often generated at 44.3 barg (4531 kPa) and superheated to 400°C to produce more efficient power production in turbines. The superheated stream is then desuperheated to produce high-pressure saturated steam (41 barg, 254°C) before being used in heat transfer equipments (Turton, 2003). One of the reasons that saturated steam is preferred over superheated steam is that the specific heat capacity for superheated steam varies considerably according to pressure and temperature. This makes it hard to estimate a required heat transfer surface area. Another reason is the amount of heat given up by the superheated steam as it cools to saturation temperature is relatively small in comparison to its enthalpy of evaporation. This implies lower rates of heat transfer whilst the steam is superheated. In addition, in a heat exchanger, using superheated steam can lead to the formation of a dry wall zone close to the tube sheet and the dry wall area can quickly become fouled.

In a supercritical process, the temperature of high-pressure saturated steam is not sufficient to heat up the reactant streams to reaction temperature (350°C). Therefore, a hot circulating heat transfer fluid along with fuels such as coal and natural gas are needed for the high-temperature heating in a furnace or fired heater. Some researchers (Lim et al., 2009), however, reported only high-pressure

steam as a heating source in their supercritical processes, implying the undesirable usage of superheated steam in their research.

In this study, hot temperature heating oil was used as the utility to provide heat in the process wherever the operating temperatures were greater than 254°C. The specific heat of the heating oil was 2.1059 kJ/(kg·K) and the inlet and exit temperatures were set at 385.0°C and 357.2°C, respectively, as specified in IPE. The cost for the hot circulating heat transfer fluids like the heating oil mostly comes from the fuel that is burned to heat the circulating heat transfer fluid (Turton, 2003). According to Turton (2003), typical efficiencies for these heaters range from 60% to 82% based on lower heating value of the fuel, or it can be as high as 90% with air preheating economizers. In this study, 90% efficiency was assumed with natural gas as the fuel source. The utility cost for the circulating heating oil was calculated by Equation (3.2) based on the natural gas price of \$6.00/GJ (Turton, 2003). The prices of utilities are listed in Table 3.3.

Utility cost for the circulating heating oil

$$\begin{aligned}
 &= \frac{\text{Natural gas price}}{\text{heater efficiency}} (\text{specific heat of heating oil})(\Delta T \text{ of heating oil in the heater}) \\
 &= \frac{\$6.00}{\text{GJ}} \frac{1}{0.9} \left(2.1059 \frac{\text{kJ}}{\text{kg} \cdot \text{K}} \right) (385 - 357.2 \text{ K}) \frac{1 \text{ GJ}}{10^6 \text{ KJ}} \\
 &= \$0.0004/\text{kg of heating oil used in the heater} \tag{3.2}
 \end{aligned}$$

Table 3.3. The prices of utilities used in this study

Utility	Price
Chilled water ^a	\$1/tonne
Cooling water ^a	\$0.013/tonne
Circulating heating oil	\$0.4/tonne
High-pressure steam (4201 kPa) ^b	\$16.64/tonne
Medium-pressure steam (1135 kPa) ^b	\$13.71/tonne
Low-pressure steam (690 kPa) ^b	\$12.68/tonne

^a From Seider et al. (2004)

^b From Turton (2003)

The price of fresh canola oil was obtained from a report released by USDA Foreign Agricultural Service (USDA, 2010a). According to the report, the canola oil price was \$916/tonnes in January 2010, followed by soybean oil price (\$769/tonne).

The prices of yellow grease and biodiesel were obtained from a national weekly summary report released by USDA Agricultural Marketing Service (USDA, 2010b). In the report, the price of yellow grease was recorded at \$23.50-24.50/cwt (100 lb) in Central U.S. in April 2010, which is equivalent to \$530/tonne. During the same period, biodiesel (B100) price was \$3.27-3.40/gallon in Iowa, Illinois and Ohio, which is around \$0.99/kg with the assumed density of 870.5 kg/m³.

The information of the prices of methanol and glycerine were obtained from the sample reports released by ICIS Pricing. According to the reports, methanol price in US Gulf was \$260.37-277.06/tonne in October 2009 (McGinn, 2009) and the prices of pharmaceutical grade and crude glycerine were in the range of \$1212.54-1433.00/tonne and \$750-970/tonne, respectively, during the same period (Lefebvre, 2009). In this study, the averages of the reported prices were taken for methanol, pharmaceutical grade and crude glycerine, which were \$268.71/tonne, \$1322.77/tonne and \$860/tonne, respectively. The prices of other chemicals are shown in Table 3.4.

Table 3.4. The prices of raw materials and products used in this study

Name	Specification	Price (\$/tonne)
Methanol	-	268.71
Fresh vegetable oil	canola oil	916
Waste vegetable oil	-	530
Glycerol	pharmaceutical grade	1322.77
	vegetable glycerine	860
Sodium hydroxide ^a	-	200
Sulfuric acid ^a	-	60
Phosphoric acid ^a	-	340
Biodiesel (B100)	qualified to meet EN 14214 and ASTM D 6751	990

^a From West et al. (2008)

3.3. Mapping and sizing

In this section, the specific types and the material of construction (MOC) of process equipments were defined and sized according to principles outlined in the literatures (Luyben, 2002; Seider et al., 2004; Chohey, 2004).

Pumps

A centrifuge pump is one of the most widely used pumps in chemical plants. It can be used with head from 15.24 to 975.4 m and kinematic viscosity less than 100 centistokes (Seider et al., 2004). Therefore, it was selected for the pumps in the alkali-catalyzed processes. In the supercritical processes, however, external rotary gear pumps were used due to the relatively high pressure requirement (19 MPa). Driver powers for these pumps were calculated with 75% efficiency in Hysys simulation. These values were used in IPE to calculate the cost of pumps including their motor drivers.

Heat exchangers

A wide variety of heat exchangers is available for heating and cooling system. Shell-and-tube heat exchangers are usually considered first because their area to volume ratio is very high. Depending on the design, a shell-and-tube exchanger is divided into floating head, fixed head, U-tube and kettle vapourizer. In the four simulation models, fixed and floating head heat exchangers were used as the basic heat exchangers wherever the required heat transfer areas were larger than 2 m². The heat exchangers with heat transfer areas smaller than 2 m² were considered as double-pipe heat exchangers.

Sizing heat exchangers was done using Icarus Heat Exchanger Sizing Expert. For the calculations, the information of the process streams such as inlet and exit temperature and the components in the streams were used. Based on the primary fluid component, the stream is classified into several categories and the following design parameters are estimated: latent heats, fouling resistance, specific heat capacity of the fluid, liquid film resistance, overall heat transfer coefficient and therefore heat transfer area. In addition, the detailed estimations of the number of tubes, tube length, tube diameter, and other internal components of the heat exchangers were performed simultaneously. The details of heat exchangers are shown in Tables 3.6 through 3.9.

Reactors

The CSTR reactors in the alkali-catalyzed processes were specified as agitated tanks. Based on the residence time, which is 1 hr for both of the transesterification and esterification reactions, the reactor sizes were determined. The length to diameter ratio was assumed to be 3:1.

For the PFRs in the supercritical processes, the reactor type used in the studies on continuous biodiesel production was referred to. Most of the researchers used stainless steel tubes in their laboratory-scale experiments (Bunyakiat et al., 2006; Silva et al., 2007; He et al., 2007; Vieitez et al., 2008; Marulanda et al., 2010). It was considered appropriate to use a pressure vessel to represent a continuous reactor in the industrial-scale process.

Seider et al. (2004) reported that at high flow rates, i.e., at high Reynolds numbers, in a long tubular reactor, the PFR model is generally valid because turbulent flow may approximate plug flow without appreciable axial mass and heat transfer. On the other hand, PFR model is not valid in laminar flow at Reynolds numbers below 2100 because of the parabolic velocity profile. Therefore, the dimension of the pressure vessel was specified to have Reynolds number greater than 8000 in order to achieve turbulent flow. Reynolds number of a fluid in a pipe or tube is defined as Equation (3.3)

$$Re = \frac{\rho VD}{\mu} = \frac{VD}{\nu} = \frac{QD}{\nu A} \quad (3.3)$$

Where ρ = density of the fluid (kg/m^3), V = mean fluid velocity (m/s), D = hydraulic diameter of the pipe (m), μ = dynamic viscosity ($\text{kg}/(\text{m}\cdot\text{s})$ or Pa·s), ν = kinematic viscosity (m^2/s), Q = volumetric flow rate (m^3/s) and A = cross-sectional area (m^2).

For a circular tube or duct, the hydraulic diameter is same as the geometrical diameter. Equation (3.3) can be re-expressed as Equation (3.4) to calculate a diameter with which Reynolds number is greater than 8000.

$$Re = \frac{4\rho Q}{\pi\mu D} > 8000 \quad (3.4)$$

The required parameters for Equation (3.4) such as a volumetric flow rate, density and viscosity of the fluid were obtained from Hysys simulation data.

The residence time and reaction conversion were obtained from experimental data in Bunyakiat et al. (2006). According to the study, the residence times of 7 and 13.3 min are required to obtain 96% conversion to methyl esters using 42:1 and 24:1 methanol to oil molar ratios, respectively. For the inlet flow rate of $53.24 \text{ m}^3/\text{hr}$ to the PFR in SC-WVO, the necessary volume of the reactor was calculated to be 6.21 m^3 . In SC-LowE process where 13.3 min of residence time was used, the required reactor volume was calculated to be 7.23 m^3 . The dimensions of the reactors are shown in Tables 3.8 and 3.9.

Distillation columns

The first step in sizing the distillation column was to determine the column type between a tray and a packed tower. It was determined based on an initial diameter estimated from the F-Factor

method (Luyben, 2002). If the diameter of a column was estimated to be larger than 0.9 m, the column was specified as a tray column. The final column diameter, D_T , was calculated from the Equation (3.5)

$$D_T = \left[\frac{4G}{(0.8 U_f)\pi \left(1 - \frac{A_d}{A_T}\right) \rho_G} \right]^{0.5} \quad (3.5)$$

where G = mass flow rate of vapour (kg/hr), U_f = vapour flooding velocity (m/s), A_d = downcomer area (m^2), A_T = tower inside cross-sectional area (m^2) and ρ_G = vapour density (kg/m^3).

The vapour flooding velocity, U_f , was computed from an empirical capacity parameter, C , based on a force balance on a suspended liquid droplet:

$$U_f = C \left(\frac{\rho_L - \rho_G}{\rho_G} \right)^{0.5} \quad (3.6)$$

where ρ_L = liquid density (kg/m^3).

Hysys simulation data were used in Equations (3.5) and (3.6). The necessary parameter and factors were obtained from Seider et al. (2004). The default type of trays in a tower was a sieve tray. If Hysys reported weeping had occurred, valve trays were selected instead. The height of a tray tower was calculated from Equation (3.7).

$$\begin{aligned} &\text{Height of a distillation column (m)} \\ &= 1.25 \times \text{the number of actual stages} \times \text{tray spacing (m)} \end{aligned} \quad (3.7)$$

A distillation column with a calculated diameter <0.9 m was specified as a packed column. The final diameter of a packed column was determined using modified Sherwood correlation (Chopey, 2004). The procedure to calculate the diameter and height of a packed column is as follows:

1. Between random and structured packings, a random packing was chosen due to its relatively low cost.

2. Once the arrangement of packing has been determined, a type and size of packing was chosen. Among the various packing types, metal Pall rings, which can withstand high temperatures, were chosen in this study.
3. The diameter of the packed distillation column was determined by modified Sherwood correlation described in Chopey (2004). The correlation is based on liquid and vapour flow rates through the column and packing configuration represented as packing factor, F , in y-axis. The pressure drop was assumed to be 0.50 inches of water per foot for a moderate-pressure distillation.
4. The height of the column was determined using Equation (3.8).

Height of a packed column (m)

$$= 1.25 \times \text{number of theoretical plate} \times \text{HETP (Height Equivalent to a Theoretical Plate, m)} \quad (3.8)$$

The value of HETP for a packed column was chosen depending on the packing sizes as shown in Table 3.5.

Table 3.5. HETP values for a packed column (Chopey, 2004)

Nominal packing size, for slotted rings or Intalox saddles	HETP
1 in	1.5 ft (0.46 m)
1.5 in	2.2 ft (0.67 m)
2 in	3.0 ft (0.91 m)

Gravity separators

The gravity separators in the four processes were specified as horizontal liquid-liquid separators. The volumes were determined based on 1 hr residence time with the separators half full. The aspect ratio of L/D was 3.

Liquid-liquid extraction column

Liquid-liquid extractors are often a neglected part of process plants because they are not well understood and are generally a small part of the overall process scheme (Chopey, 2004). Commercially important extractors can be broadly classified into mixer-settlers, centrifugal devices, static column contactors and agitated column contactors (Chopey, 2004). The difference between a static and

agitated column is that an agitated column provides a better degree of mixing of the immiscible liquids and is used for an extraction column requiring a large number of stages. In this study, a static column type was selected because of the moderate number of required stages as estimated in Hysys simulation. Due to a lack of information to thoroughly size a liquid-liquid extraction column, a rather simple method of sizing an extraction column was used as described in Seider et al. (2004).

Table 3.6. Equipment summary for Alkali-FVO process

Pumps	P-101	P-102	P-103	P-202
Type	centrifuge	centrifuge	centrifuge	centrifuge
Efficiency	75%	75%	75%	75%
MOC	316SS	CS	CS	CS
Flow (kg/hr)	617.8	5250	574.4	5868
Press in (kPa)	101.3	101.3	20.0	30.0
Press out (kPa)	400.0	400.0	400.0	120.0
Driver power (kW)	0.0858	0.6384	0.1033	0.2154
Heat Exchangers	E-101	E-102		
Type	double pipe	fixed head		
Area (m ²)	1.0	5.97		
MOC	CS	CS		
Duty (kW)	83.45	307.8		
Shell in (°C/kPa)	25.3/400	142.8/30		
Tube in (°C/kPa)	164/690	23.8/345		
Vessels/Reactors	CRV-100	CRV-101	X-100	
Type	agitated reactor	agitated reactor	L-L separator	
Orientation	vertical	vertical	horizontal	
MOC	316SS	316SS	316SS	
T (°C)	60.0	161.2	161.2	
P (kPa)	400.0	101.3	101.3	
Residence time (min)	60	40	60	
Diameter (m)	1.847	0.720	0.802	
Height/Length(m)	5.541	2.160	2.406	
Tower	T-101	T-102	T-103	T-104
Application	distillation	extraction	distillation	distillation
# of actual stages	9	6	9	8
Type of trays / Packing materials	valve	sieve	valve	1 in. Pall rings
MOC	CS	CS	CS	CS
Diameter (m)	0.90	0.602	2.440	0.3048
Height (m)	6.858	9.144	6.858	4.572

Table 3.7. Equipment summary for Alkali-WVO process

Pumps	P-101	P-102	P-103	P-104	P-105	P-106	P-107
Type	centrifuge	centrifuge	centrifuge	centrifuge	centrifuge	centrifuge	centrifuge
Efficiency	75%	75%	75%	75%	75%	75%	75%
MOC	316SS	CS	CS	CS	SS316	CS	CS
Flow (kg/hr)	173.1	1167	5275	5256	560.6	542.9	5817
Press in (kPa)	101.3	28.0	101.3	200.0	101.3	20.0	30.0
Press out (kPa)	400.0	400.0	400.0	400.0	400.0	400.0	120.0
Driver power (kW)	0.0199	0.2073	0.6405	0.4242	0.0778	0.1046	0.2141
Heat Exchangers	E-101	E-102	E-103				
Type	float head	double pipe	fixed head				
Area (m ²)	35.18	1.0	6.47				
MOC	316SS	CS	CS				
Duty (kW)	849.58	34.05	358.11				
Shell in (°C/kPa)	25.3/400	46.4/400	156/30				
Tube in (°C/kPa)	70/400	164/690	23.8/345				
Vessels/Reactors	CRV-100	CRV-101	CRV-102	X-102			
Type	agitated reactor	agitated reactor	agitated reactor	L-L separator			
Orientation	vertical	vertical	vertical	horizontal			
MOC	316SS	316SS	316SS	316SS			
T (°C)	70.0	60.0	157.1	157.1			
P (kPa)	400.0	400.0	101.3	101.3			
Residence time (min)	60	60	40	60			
Diameter (m)	1.863	1.830	0.712	0.904			
Height/length (m)	5.589	5.490	2.136	2.712			
Tower	T-100	T-101	T-102	T-103	T-104	X-100	
Application	distillation	distillation	extraction	distillation	distillation	extraction	
# of actual stages	8	9	6	9	8	6	
Type of trays / packing materials	valve	valve	sieve	valve	1 in. Pall ring	sieve	
MOC	316SS	CS	CS	CS	CS	CS	
Diameter (m)	1.524	0.9144	0.613	2.286	0.3048	0.646	
Height (m)	6.096	6.858	9.144	6.858	4.572	9.144	

Table 3.8. Equipment summary for SC-WVO process

Pumps	P-101	P-102		
Type	gear	gear		
Efficiency	75%	75%		
MOC	316SS	316SS		
Flow (kg/hr)	9084	5162		
Press in (kPa)	101.3	101.3		
Press out (kPa)	19000	19000		
Driver power (kW)	84.57	39.66		
Heat Exchangers	E-101	E-102	E-103	E-104
Type	float head	float head	float head	float head
Area (m ²)	40.69	14.78	92.64	37.88
MOC	316SS	316SS	316SS	CS
Duty (kW)	1608.2	765.14	2083.6	489.8
Shell in (°C/kPa)	71.9/19000	41.9/19000	385/2533	164.1/106.5
Tube in (°C/kPa)	304/19000	350/19000	224/19000	23.8/345
Vessels/Reactors	PFR-100	V-101		
Type	continuous pressure vessel	L-L separator		
Orientation	horizontal	horizontal		
MOC	316SS	CS		
T (°C)	350	25		
P (kPa)	19000	170		
Residence time (min)	7	60		
Diameter (m)	0.8	1.76		
Height/length (m)	12.36	5.28		
Tower	T-101	T-102		
Application	distillation	distillation		
# of actual stages	6	10		
Type of trays / Packing materials	sieve	valve		
MOC	CS	CS		
Diameter (m)	2.0	3.1		
Height (m)	4.572	7.620		

Table 3.9. Equipment summary for SC-LowE process

Pumps	P-101	P-102			
Type	gear	gear			
Efficiency	75%	75%			
MOC	316SS	316SS			
Flow (kg/hr)	5433	5162			
Press in (kPa)	101.3	101.3			
Press out (kPa)	19000	19000			
Driver power (kW)	49.55	39.66			
Heat Exchangers	E-101	E-102	E-103	E-104	E-105
Type	float head	float head	float head	fixed head	float head
Area (m ²)	24.67	11.53	65.54	53.99	38.04
MOC	316SS	316SS	316SS	CS	CS
Duty (kW)	992.2	707.3	1460.7	1533.6	514.7
Shell in (°C/kPa)	64.76/19000	41.94/19000	385/2533	69.23/101.5	170.8/106.5
Tube in (°C/kPa)	287.7/19000	350.0/19000	226.5/19000	35.00/344.98	35.00/344.98
Vessels/Reactors	PFR-100	V-100	V-101		
Type	continuous pressure vessel	knock-out	L-L separator		
Orientation	horizontal	vertical	horizontal		
MOC	316SS	CS	CS		
T (°C)	350	96.18	25.0		
P (kPa)	19000	200.0	170.0		
Residence time (min)	13.3	10	60		
Diameter (m)	0.8	1.22	1.76		
Height/length (m)	14.38	3.81	5.28		
Tower	T-101	T-102			
Application	distillation	distillation			
# of actual stages	6	10			
Type of trays / Packing materials	2 in. Pall rings	valve			
MOC	CS	CS			
Diameter (m)	0.765	2.90			
Height (m)	6.858	7.62			

3.4. Investment parameters

Investment parameters to evaluate the profitability of a project were specified by including a project life, salvage value, tax rate, depreciation method, desired rate of return, escalation rates, working capital percentage, operating cost parameters, and facility operation parameters.

A project life is defined as a specific length of time over which the profitability of different projects is to be compared (Turton, 2003). The lives typically used for this purpose are 10, 12 and 15 years (Turton, 2003). In this study, a ten-year analysis period was chosen. Accordingly, all the equipment in each process was assumed to equally have ten-year of useful lives for the sake of simplicity. Salvage value is an estimated value of fixed capital investment at the end of the project life. It was conservatively specified to be zero. Taxation is generally very complex and frequently changes over time. The corporate income tax in US varies from 15% to 39% depending on the net taxable income. In addition, state, city and other local taxes are also charged. For most large corporations, the overall taxation rate is often in the range of 40% to 50% (Turton, 2003). In this study, the tax rate was specified to be 40%. As a depreciation method, a straight line method most commonly used was selected. The escalation rates for project capital, products, raw materials, operating and maintenance labour and utilities were set at the default values of IPE as shown in Table 3.10. Working capital, which indicates funds for the operating costs required for the early operation of the plant, was set at 15% of fixed capital cost.

Operating cost parameters include operating charges, costs for plant overhead and General and Administrative (G&A). Operating supplies and laboratory charges cover the cost of the miscellaneous items required in order to run the plant and the cost of analyzing products. They were specified as 10% of the maintenance cost, and 16.3% of the operating labour cost. Plant overhead indicates expenses which are not directly related to plant operation such as the costs of providing the following services: cafeteria, employment and personnel, fire protection, inspection and safety, first aid and medical, janitorial, automotive and other transportation, and recreation (Seider et al., 2004). It was estimated as 50% of operating labour and maintenance costs. G&A expenses represent the general and administrative costs such as administrative salaries/expenses, R&D, product distribution, financing and selling costs. It was specified as 15% of subtotal operating costs, as shown in Table 3.10.

Facility type, operating mode, the length of start-up period and operating hours per year can be specified in facility operation parameters category, which are shown in Table 3.10.

Table 3.10. Investment analysis parameter for the four biodiesel production processes

Name	Value	Units
Economic life of project	10	year
Tax rate	40	%/year
Desired rate of return / Interest rate	20	%/year
Salvage value	0	% of initial capital cost
Depreciation method	straight line	
Escalation Parameters		
Project capital escalation	5	%/year
Products escalation	5	%/year
Raw material escalation	3.5	%/year
Operating and maintenance labour escalation	3	%/year
Utilities escalation	3	%/year
Project Capital Parameters		
Working capital percentage	15	% of fixed capital cost
Operating Costs Parameters		
Operating supplies	10	% of maintenance/year
Laboratory charges	16.3	% operating labour/year
Maintenance	6.0	% of fixed capital cost
Plant overhead	50	% of operating labour and maintenance costs/year
G&A expenses	15	% of subtotal operating costs/year
Facility Operation Parameters		
Facility type	chemical processing facility	
Operating mode	continuous processing – 24 hours	
Length of start-up period	20	weeks
Operating hours per period	8000	hours/year
Process fluids	liquids and gases	

3.5. Project evaluation

3.5.1. Total capital investment

Table 3.11 shows the total capital costs for the four biodiesel production processes. The cost of the transesterification reactor was higher than any other equipment in Alkali-FVO, SC-WVO and SC-LowE. In Alkali-WVO, the methanol distillation column in the pre-treatment had the highest cost. Overall, the purchased equipment cost of SC-WVO and SC-LowE were higher than the other processes. Especially, the costs of a transesterification reactor, pumps and heat exchangers of the two supercritical processes were significantly higher than Alkali-FVO process: about 3 times higher in reactor and pump costs; and 25 times higher in heat exchanger costs. The reasons for the high costs of this equipment were mainly the high operating pressure and temperature. For the reactors, the material of construction was 316SS in all of the four processes. However, the high operating temperature and pressure in SC-WVO and SC-LowE increased the thickness of the reactors, and thereby their cost. Operating conditions also contributed to the increases in the sizes of the pumps and heat exchangers in SC-WVO and SC-LowE processes.

The total direct costs of Alkali-FVO, Alkali-WVO, SC-WVO and SC-Low E processes were estimated as \$3.594, \$5.336, \$5.592 and \$5.425 million, respectively. Despite the use of an extra unit (flash evaporator), SC-LowE process had a slightly lower total direct cost compared to SC-WVO due to the reduced expenditure for the methanol recovery column, pumps and heat exchangers. The reduction was brought by smaller sizes of the equipment for processing less amount of methanol. The largest number of pieces of equipment in Alkali-WVO process generated the highest costs for equipment setting, civil, steel, instrumentation, electrical, insulation and paint.

The total capital investment for Alkali-FVO was estimated as \$11.1 million, which was the lowest among the four processes. Alkali-WVO had the highest total capital investment estimated as \$15.6 million. As previously stated, the largest number of equipment units in this process caused the highest indirect costs except for the contingency fees. Because the contingent fees were rated high for the new unproven processes like the supercritical processes, the fees were estimated as \$3.20 and \$3.17 million for SC-WVO and SC-LowE processes, respectively. The fees for Alkali-FVO and Alkali-WVO processes were \$1.48 and \$2.08 million, respectively. The total capital investment for SC-WVO was \$15.0 million, followed by \$14.8 million of SC-LowE.

Table 3.11. Total capital investment for biodiesel production processes at the capacity of 40,000 tonnes of biodiesel production/yr (unit: \$ millions)

Type	Description	Alkali-FVO	Alkali-WVO	SC-WVO	SC-LowE
Direct items					
Reactor	Esterification	-	0.255	-	-
	Transesterification	0.252	0.249	0.662	0.726
	Neutralization	0.080	0.080	-	-
Column	Methanol recovery ^a	0.093	0.256 ^b /0.099	0.211	0.108
	Biodiesel purification ^a	0.186	0.182	0.241	0.227
	Glycerol purification ^a	0.065	0.047	-	-
	Water washing	0.034	0.034	-	-
Other	Glycerol washing	-	0.073	-	-
	Pumps	0.017	0.031	0.063	0.052
	Heat exchangers	0.020	0.059	0.651	0.557
	L-L separator	0.025	0.027	0.026	0.026
	Flash evaporator	-	-	-	0.021
Total purchased equipment		0.772	1.39	1.85	1.72
Equipment setting		0.016	0.023	0.021	0.023
Piping		0.690	1.12	1.82	1.77
Civil		0.133	0.187	0.123	0.126
Steel		0.054	0.087	0.034	0.037
Instrumentation		1.21	1.68	0.968	0.991
Electrical		0.497	0.529	0.492	0.489
Insulation		0.195	0.283	0.255	0.246
Paint		0.028	0.035	0.023	0.021
Total direct cost		3.594	5.336	5.592	5.425
Others^c		4.050	5.454	3.627	3.703
G&A Overheads		0.150	0.219	0.215	0.209
Contract Fee		0.402	0.526	0.411	0.412
Contingencies		1.48	2.08	3.20	3.17
Fixed capital cost		9.67	13.6	13.0	12.9
Working capital		1.45	2.04	1.96	1.94
Total capital investment		11.1	15.6	15.0	14.8

^a Includes the costs for a distillation column, condenser, distillate accumulator, reflux pump and reboiler

^b Indicates a pre-treatment process

^c This item is the total of the following costs: design, engineering, and procurement costs, material charges (freight and taxes) and construction field indirect costs (fringe benefits, burdens, consumables/small tolls, insurance, equipment rental, field services, field office construction supervision, and plant start-up)

3.5.2. Total manufacturing cost

The results of the total production costs of the four processes are shown in Table 3.12. The total production cost for biodiesel in Alkali-FVO was calculated to be \$50.9 million, which was the highest among the four processes. The lowest total production cost was \$25.9 million of SC-LowE process. Figure 3.2 shows the contributions of sub-items to the total production costs for biodiesel. The oil feed cost was found to be the major contributor to the total biodiesel production cost in all the processes. As shown in Figure 3.2, it comprises 84.0%, 63.9%, 71.3% and 75.4% of the total manufacturing costs of Alkali-FVO, Alkali-WVO, SC-WVO and SC-LowE processes, respectively. Due to the usage of fresh vegetable oil feedstock, the percentage of oil feedstock was especially high in Alkali-FVO process. However, the other processes using waste vegetable oil decreased the oil feed costs by approximately 42% in comparison with Alkali-FVO.

The total manufacturing cost of Alkali-WVO process was \$35.0 million, which was the second highest among the four processes. Compared to Alkali-FVO, the cost for raw materials except for the oil feed was higher because of the usage of larger amount of methanol, sulphuric acid catalyst and glycerol for washing solution in the liquid-liquid extraction column in the pre-treatment process. The utility, maintenance and indirect manufacturing costs of Alkali-WVO were also slightly higher than Alkali-FVO due to the complexity of the process. However, using waste vegetable oil feedstock saved \$10.8 million per year in the total manufacturing cost of Alkali-WVO.

Both of SC-WVO and SC-LowE processes showed lower total manufacturing costs than the alkali-catalyzed process. Because the reaction conversion in the supercritical processes was 96%, which was slightly higher than the conversion in the alkali-catalyzed processes (95%), the oil feed cost was decreased by \$0.48 million per year in comparison with Alkali-WVO process. The costs for methanol feed in the supercritical processes were similar to those in the alkali-catalyzed processes despite the use of high molar ratio of methanol to oil in the reactions. As sufficient amount of methanol required for the reaction was supplied by the recycled methanol stream from the methanol distillation column, it resulted in similar amounts of fresh methanol feed stream in every process.

The costs of utilities in the supercritical processes were higher than the alkali-catalyzed process due to the harsh reaction conditions. The larger amount of methanol to be evaporated in the distillation columns also increased the utility costs in the supercritical processes. The usage of electricity mostly came from the agitated reactors in Alkali-FVO and Alkali-WVO processes and from the pumps in SC-WVO and SC-LowE processes. Comparing SC-LowE with SC-WVO, SC-LowE saved approximately \$1.16 million per year in the cost of steam (1135 kPa) simply because of the application of the flash evaporator prior to the methanol distillation column. The saving on the steam cost was much larger than the relatively small cost for purchasing and installing the flash evaporator.

The glycerol credits in Alkali-FVO and Alkali-WVO processes were \$5.49 and \$5.19 million, respectively, which were higher than those in the supercritical processes due to the high purity of the glycerol. The difference in the market prices of glycerol depending on its purity and the amount of the crude glycerol to be purified are important indicators to determine on the construction of a glycerol purifying distillation column. For example, if the crude glycerol stream in Alkali-FVO process was not purified and sold at the lower price of \$860/tonnes, the glycerol credit would have been decreased by \$1.98 million per year. In other words, by constructing the glycerol purifying distillation column at the expense of \$0.065 million, the increase of \$1.98 million per year in glycerol credit was achieved. In the same manner, Alkali-WVO process could also increase its profit by constructing the glycerol purifying column.

The revenue from the sales of biodiesel was \$39.6 million per year in each process on the assumption that all the biodiesel products were sold in the market. Compared to the total manufacturing costs, the revenue was large enough to make profits in Alkali-WVO, SC-WVO and SC-LowE processes. However, the total manufacturing cost was \$6.2 million dollars higher than the earnings from the biodiesel sales in Alkali-FVO process, indicating that the process was in deficit. To make Alkali-FVO process profitable, it seems inevitable to raise the biodiesel selling price and/or decrease the raw material costs.

Table 3.12. Total manufacturing cost, glycerol credit and revenues from biodiesel sales in the four processes at the capacity of 40,000 tonnes of biodiesel production/yr (unit: \$ millions)

Process	Alkali-FVO	Alkali-WVO	SC-WVO	SC-LowE
Direct manufacturing cost				
Oil feed	38.46	22.36	21.88	21.88
Methanol	1.215	1.356	1.15	1.24
Catalysts	0.201	0.215	-	-
Water washing	0.000	0.000	-	-
Glycerol washing	-	3.296	-	-
Steam (4201 kPa)	-	-	-	-
Steam (1135 kPa)	0.006	0.007	1.328	0.168
Steam (690 kPa)	0.132	0.563	-	-
Circulating heating oil	0.407	0.400	0.787	0.616
Cooling water	0.019	0.019	0.081	0.030
Chilled water	0.234	1.745	-	-
Electricity	0.066	0.099	0.115	0.089
Operating labour	0.800	0.800	0.600	0.600
Supervision	0.280	0.280	0.280	0.280
Maintenance	0.580	0.817	0.783	0.775
Operating supplies	0.058	0.082	0.078	0.078
Laboratory charges	0.143	0.176	0.143	0.143
Subtotal	42.40	32.22	27.23	25.90
Indirect manufacturing cost				
Overhead	0.730	0.948	0.831	0.828
Depreciation	0.822	1.350	1.201	1.096
Property Taxes	0.224	0.315	0.302	0.299
Insurance	0.097	0.136	0.130	0.129
Subtotal	1.873	2.749	2.464	2.352
General and Administrative costs	6.641	5.245	4.454	4.238
Total production cost	50.9	40.2	34.15	32.49
Glycerol credit	5.49	5.19	3.47	3.45
Total manufacturing cost	45.8	35.0	30.7	29.0
Revenue from biodiesel	39.6	39.6	39.6	39.6

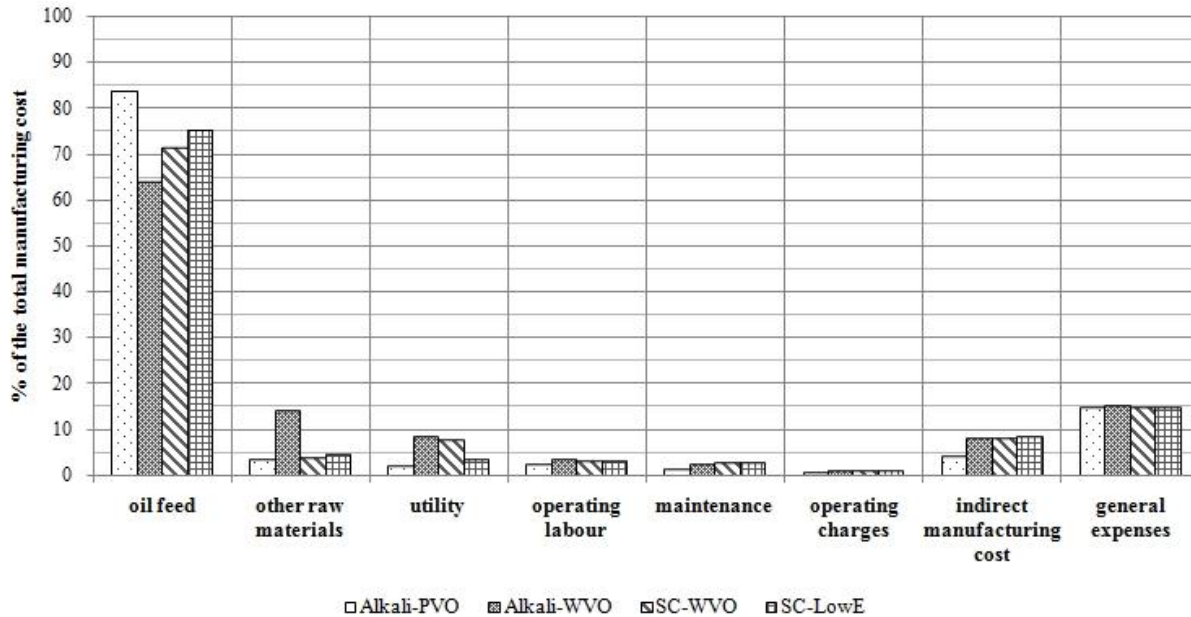


Figure 3.2. The proportions of subitems in the total production cost for biodiesel of the four processes

3.5.3. Profitability analysis

In the process of making an investment decision, the profit anticipated from an investment must be judged relative to some profitability standards. A profitability standard is a quantitative measure of profit with respect to the capital investment required to generate that profit. Several methods are used for project evaluation, and the methods can be divided into discounted techniques and non-discounted techniques. Non-discounted cash flow techniques do not consider the time value of money and therefore are not suitable for final project evaluation. The methods include payback period and return on investment. Discounted cash flow techniques are more rigorous profitability measures which involve consideration of the time value of money and estimates of the cash flows throughout the life of the process. The methods include the discounted payback period (DPP), net present value (NPV), equivalent annual value (EAV) and discounted cash flow rate of return (DCFROR).

Discounted payback period is the time required, after start-up, to recover the fixed capital investment as all cash flows discounted back to time zero. It corresponds to the time when the cumulative present value crosses over from negative to positive hitting zero. The project with the shortest time is the more favourable. However, it may lead to an economically incorrect decision, as it does not reflect cash flows after payback. The project with the shortest DPP might not produce the highest return at the end of the project life. Therefore, it should be only used to provide initial screening or supplemental information in conjunction with an analysis performed using different methods.

Net present value is defined as the sum of all cash inflows and outflows as they are discounted to the present worth by the given interest rate. A positive NPV indicates that a project is acceptable and the higher the NPV, the better the potential project. To calculate NPV, an appropriate interest rate (or discount rate) needs to be defined first. This internal interest rate usually represents the minimum acceptable rate of return (MARR) that must be earned for a project to be accepted. It is normally adjusted to account for the uncertainties and risks associated with the project. In this study, the internal interest rate was set at 20% as shown in Table 3.10.

Discounted cash flow rate of return also known as internal rate of return (IRR) is defined as the interest rate at which all the cash flows are discounted in order to bring the net present value to exactly zero. It represents the highest after-tax interest at which the project can just break even (Turton 2003). If the IRR is greater than the internal interest rate, the project is regarded to be profitable.

Among the various ways to measure profitability, DPP, NPV and DCFROR were the interests of this study and the results of the four processes are shown in Table 3.13 including the break-even price of biodiesel. The break-even price was defined as the price at which the revenue from selling biodiesel product is the same as total manufacturing cost of each process in Table 3.12 (Zhang et al., 2003). Since Table 3.12 shows before-tax values, taxes are not applied to the break-even prices.

Table 3.13. The profitability measures and the break-even price of biodiesel

	Alkali-FVO	Alkali-WVO	SC-WVO	SC-LowE
Discounted payback period (years)	-	9.2	5.0	4.2
Net present value (\$ millions)	-26.697	1.901	16.451	21.086
Discounted cash flow rate of return (%)	-	22.4	41.7	49.6
Break-even price of biodiesel (\$/kg)	1.15	0.875	0.767	0.726
(\$/litre)^a	1.00	0.762	0.668	0.632

^a Density of 870.5 kg/m³ was applied.

Except for Alkali-FVO, the NPV of each process appeared to be positive. Both of the supercritical processes showed much higher NPV than alkali-catalyzed processes due to the lower total manufacturing costs. The DPP of SC-WVO and SC-LowE were 5.0 and 4.2 years, respectively, which were about 4 years shorter than Alkali-WVO. As previously stated, for Alkali-FVO process, the margin from biodiesel and glycerol sales was not high enough to calculate the DPP and DCFROR. The DCFROR were 22.4%, 41.7% and 49.6% for Alkali-WVO, SC-WVO and SC-LowE processes, respectively. Since these values were higher than the MARR which was set at 20% and the NPVs were greater than zero, all of the three processes were considered economically profitable. Overall, SC-

LowE was evaluated as the most favourable process, followed by SC-WVO and Alkali-WVO in order. Alkali-FVO process was considered to be an unacceptable project.

The break-even price of biodiesel was approximately \$1.00/litre for Alkali-FVO. It decreased by 23.8% in Alkali-WVO process. This result was different from Zhang et al. (2003) where the break-even price of Alkali-WVO (\$884/tonne) was estimated to be higher than Alkali-FVO (\$857/tonne). This was due to the total manufacturing cost of Alkali-WVO being estimated to be higher than the one of Alkali-FVO in the results of Zhang et al. (2003). In their study, the oil feed cost comprised 61.2% of the total manufacturing cost, whereas it was 84% in Alkali-FVO process of this study. Therefore, replacing the fresh vegetable oil with waste vegetable oil would have a greater effect in this study, thus significantly reducing the total manufacturing cost of Alkali-WVO. The higher proportion of the oil feed cost in this study compared to Zhang et al. (2003) appears to be derived from the larger capacity of the plant. As the capacity of the plant increases, the indirect manufacturing cost tends to increase relatively slower than raw material costs, resulting in a greater proportion of the raw material costs in the total manufacturing cost.

Figure 3.3 shows the U.S. diesel retail prices including taxes from January 1995 to April 2010 (US EIA, 2010). The break-even prices of the four processes are similar to or slightly higher than the diesel retail prices during 2009-2010. It should be noted that a plant profit and taxes were excluded and therefore the break-even price underestimates the real commercial price of biodiesel. Thus, to make biodiesel competitive to petroleum diesel in the fuel market, sufficient government support such as in a form of subsidy is required.

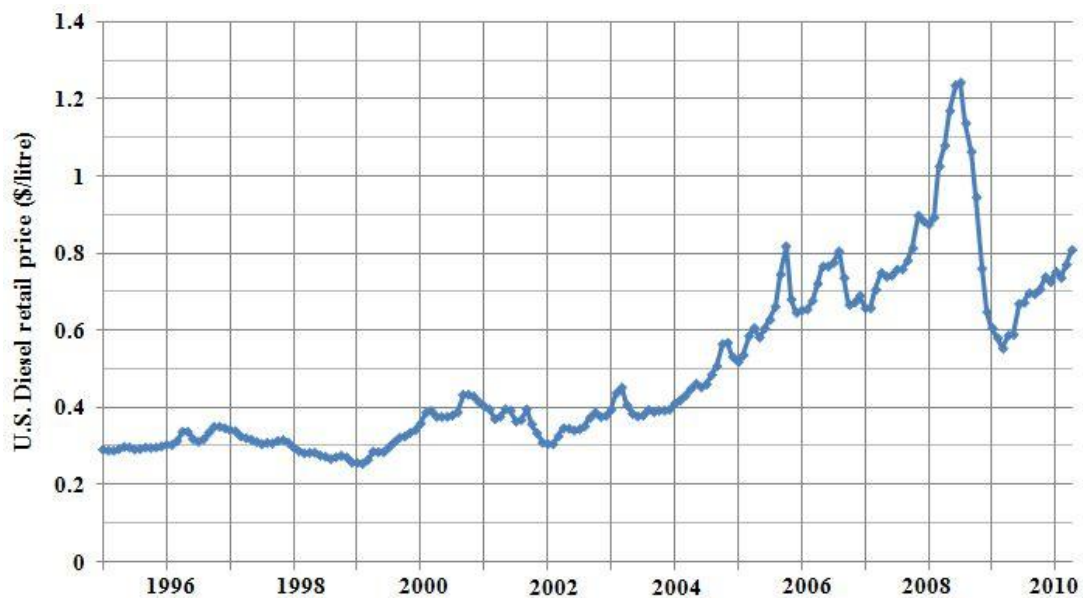


Figure 3.3. The diesel retail prices in United States from January 1995 to April 2010 (Source: US EIA (2010))

3.6. Sensitivity analysis on net present value

3.6.1. Single parameter analysis

Sensitivity analysis examines the effect of positive and negative changes in project parameters on the criteria of profitability such as NPV and DCFROR. The parameters generally examined are raw material prices, selling prices of products, levels of capacity, project lives, interest rates, capital cost and components, operating cost and components and salvage values.

Sensitivity analyses for the four biodiesel production processes were conducted to investigate the sensitivity of the NPV to changes in a variety of parameters. Six parameters were investigated: price of oil feed; glycerol credit; biodiesel selling price; capacity of a plant; interest rate; and the life of a plant. The prices of oil feed, glycerol and biodiesel were varied by ± 10 , 20, 30 and 40% from the original values as shown in Table 3.4. The capacities of the plant investigated were 8,000 and 160,000 tonnes of biodiesel production per year. The interest rate varied from 10% to 40%. Finally, the expected life of the plant was prolonged to 15 and 20 years.

The sensitivity analysis results of the oil prices are shown in Figure 3.4. The NPVs of the four processes were significantly changed by the prices of oil feed. The results showed that every 10% increase in the oil price approximately decreased the NPV by \$15, \$6.8, \$6.4 and \$6.4 million in Alkali-FVO, Alkali-WVO, SC-WVO and SC-LowE, respectively. It should be noted that the absolute value of the percentage of the oil price in Figure 3.4 is higher in Alkali-FVO than the other processes due to the high price of the fresh vegetable oil. Alkali-FVO process was, however, more sensitive to the oil price than any other processes even when the same absolute values of the changes in oil price were considered. The slopes of the trend lines of each process were evaluated and shown in Figure 3.4. The steeper the slope, the more sensitive the process is. The sensitivity to the oil price slightly increased in order of Alkali-WVO, SC-WVO and SC-LowE.

Figure 3.5 shows the effect of changes in glycerol price on the NPV of each process. The results indicated that an increase of \$0.92 and \$0.89 million in NPV of SC-WVO and SC-LowE, respectively, could be caused by an increase of \$0.086/kg in the price of crude glycerol. An increase of \$0.132/kg in the price of pharmaceutical grade glycerol resulted in an increase of \$2.1 and \$1.31 million in the NPV of Alkali-FVO and Alkali-WVO, respectively. For Alkali-WVO process, the NPV turned to negative value as the glycerol price was decreased by 15%. Overall, the influence of glycerol credit is much less than the influence of the prices of oil feed.

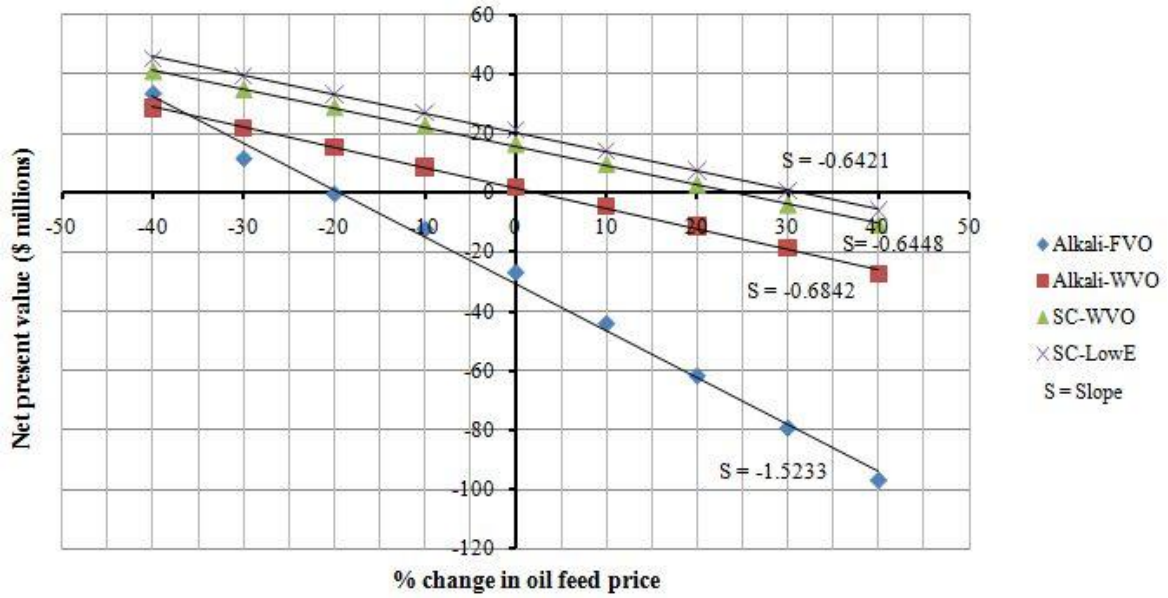


Figure 3.4. Sensitivities of the NPVs of the four processes to the price of oil feeds

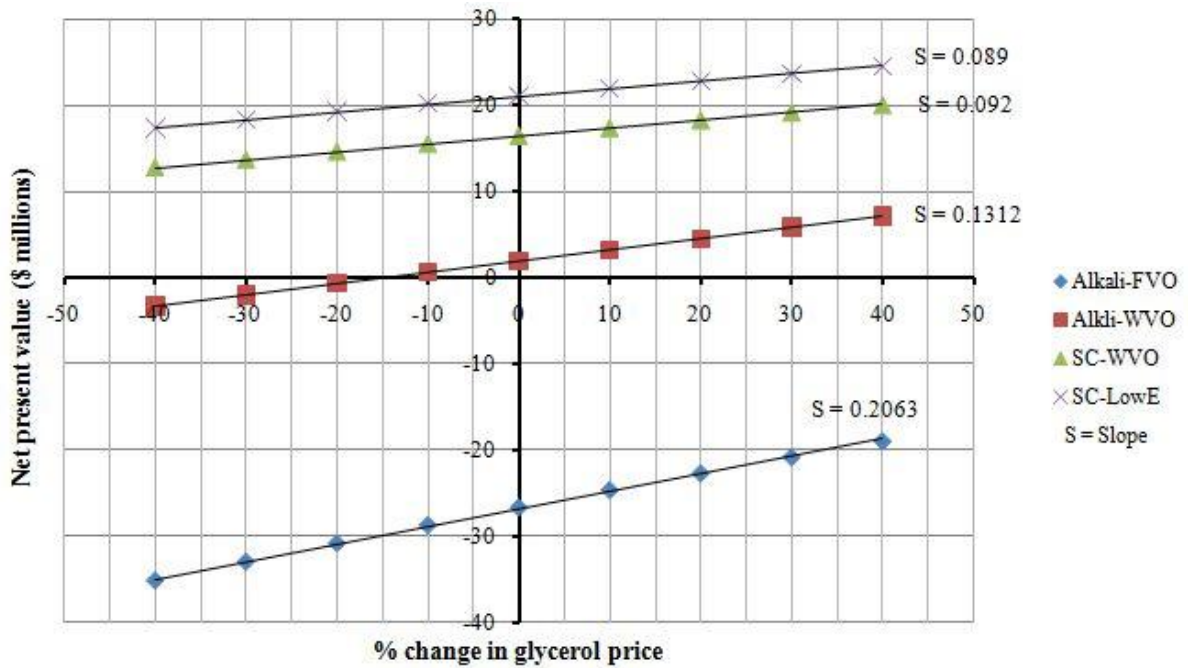


Figure 3.5. Sensitivities of the NPVs of the four processes to the price of glycerol by-product

The sensitivity analysis results of biodiesel selling price are shown in Figure 3.6. The $\pm 10\%$ changes in the price of biodiesel ($\$0.099/\text{kg}$) change the NPVs of Alkali-FVO, Alkali-WVO, SC-WVO and SC-LowE by $\pm \$13.7$, $\$11.1$, $\$10.5$ and $\$10.3$ million, respectively. The results also indicate that the NPV of Alkali-WVO, SC-WVO and SC-LowE processes are more strongly influenced by biodiesel selling prices than the oil feed prices. For Alkali-FVO process, the effect of the oil price was more significant than the biodiesel price. Consequently, the oil feed price appears to play the most important role in the profitability of Alkali-FVO.

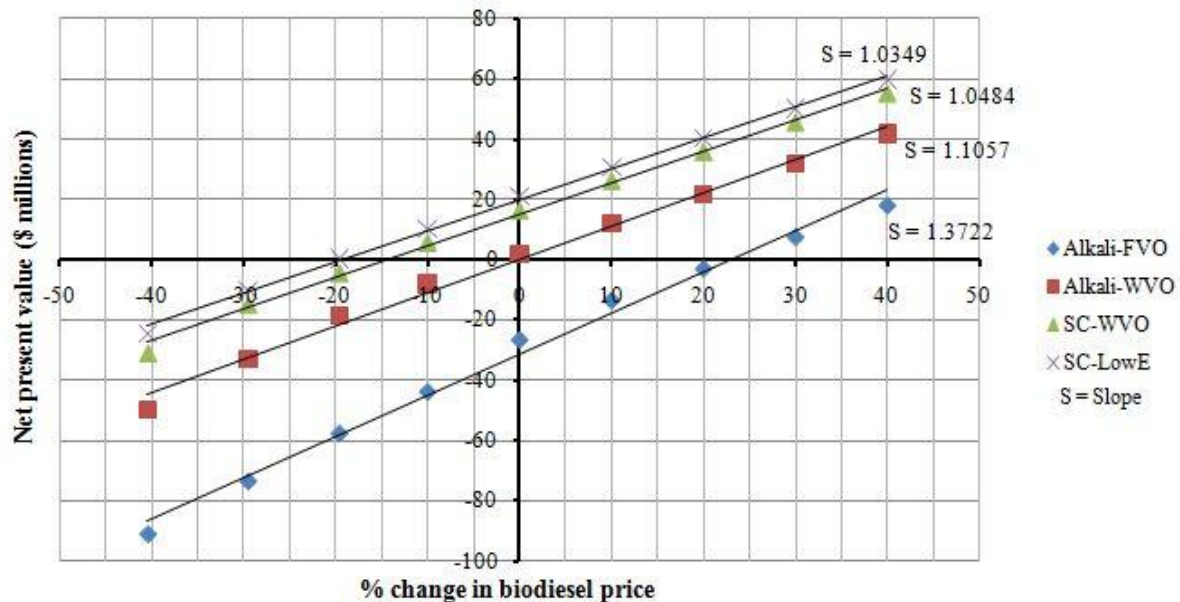


Figure 3.6. Sensitivities of the NPVs of the four processes to the price of biodiesel product

The sensitivity analysis of the interest rate is shown in Figure 3.7. The results showed that the NPVs of the four processes were very sensitive to the interest rate used for discounting cash flows. Since cash flows are discounted to time 0, the cash flows closer to the end of the project are heavily discounted. Therefore, as the interest rate increases, the revenues from biodiesel sales near the end of the project are heavily discounted, whereas the capital investment at time 0 remains constant. As a result, the NPVs of Alkali-WVO, SC-WVO and SC-LowE decreased as the interest rate increases. On the other hand, the deficit financing in Alkali-PVO over the life of the project became heavily discounted as the interest rate increases and it resulted in the increases in the NPVs.

Figure 3.8 shows the influence of the economic lives of the four processes on their NPVs. The results showed that the longer economic lives of each process increased the NPVs due to the longer

period of time to gain revenues from the biodiesel sales. As the economic life was prolonged from 10 to 20 years, the NPVs were increased by \$3.51, \$10.6, \$13.7 and \$14.6 million, respectively, in Alkali-FVO, Alkali-WVO, SC-WVO and SC-LowE processes.

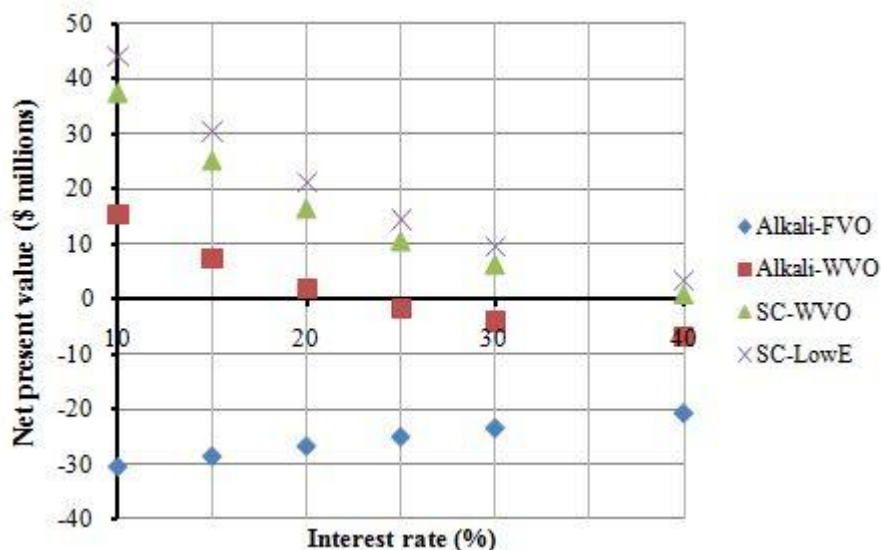


Figure 3.7. Sensitivities of the NPVs of the four processes to interest rates

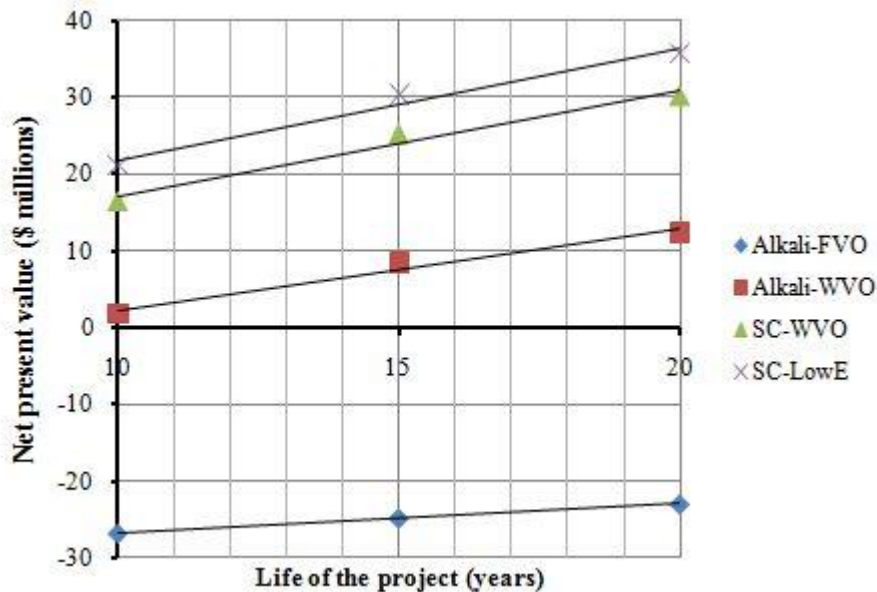


Figure 3.8. Sensitivities of the NPVs of the four processes to the life of projects

A capacity of a plant is another important factor that varies profitability of a project. Due to the growing concerns about preserving environment, many countries around the world have encouraged to use eco-friendly energy sources such as biodiesel. As an example, in June 2008, the

Canadian federal government announced plans to use of biodiesel and ethanol in Canada’s energy supply. According to the plans, the renewable fuels standard would require 5% renewable content in gasoline by 2010 and 2% renewable content in diesel and home heating oil by 2010. The mandated renewable content in diesel is approximately equivalent to 435,000 tonnes per year based on 25×10^6 m³ of annual Canadian diesel fuel market. In the United States, the US Environmental Protection Agency (EPA) released proposed revisions to the National Renewable Fuel Standard program which was established under the Energy Policy Act of 2005 (EPA, 2009). The revised statutory requirements establish new specific volume standards for cellulosic biofuel, biomass-based diesel, advanced biofuel, and total renewable fuel that must be used in transportation fuel each year as shown in Table 3.14.

Table 3.14. Renewable fuel volume requirements for renewable fuel standard (RFS2) from 2009 to 2015 for the U.S. (Source: EPA (2009)) (units: 10⁶ m³)

Year	Biomass-based diesel	Cellulosic biofuel	Total advanced biofuel	Total renewable fuel
2009	1.9	-	2.3	42.0
2010	2.5	0.38	3.6	49.0
2011	3.0	0.95	5.1	52.8
2012	3.8	1.9	7.6	58.0
2013	3.8	3.8	10.4	62.6
2014	3.8	6.6	14.2	68.7
2015	3.8	11	21	78

Based on the expected increase in biodiesel demand, the biodiesel production capacities of the four processes were expanded from 40,000 to 160,000 tonnes per year. In addition, a smaller scale of 8000 tonnes per year was investigated, which appears most frequently in literatures (Zhang et al., 2003; Kasteren and Nisworo, 2007; West et al., 2008; You et al., 2008; Lim et al., 2009). The two capacities, 8,000 and 160,000 tonnes per year, were firstly simulated in Hysys for the four different processes, namely Alkali-FVO, Alkali-WVO, SC-WVO and SC-LowE. Then each simulation was evaluated in IPE in the manner as described in from Sections 3.2 through 3.4. The details of the total capital investment and total manufacturing cost are shown in Tables A.1 and A.2 in Appendix A. The total capital investments, net present values and break-even prices of biodiesel are shown in Figure 3.9.

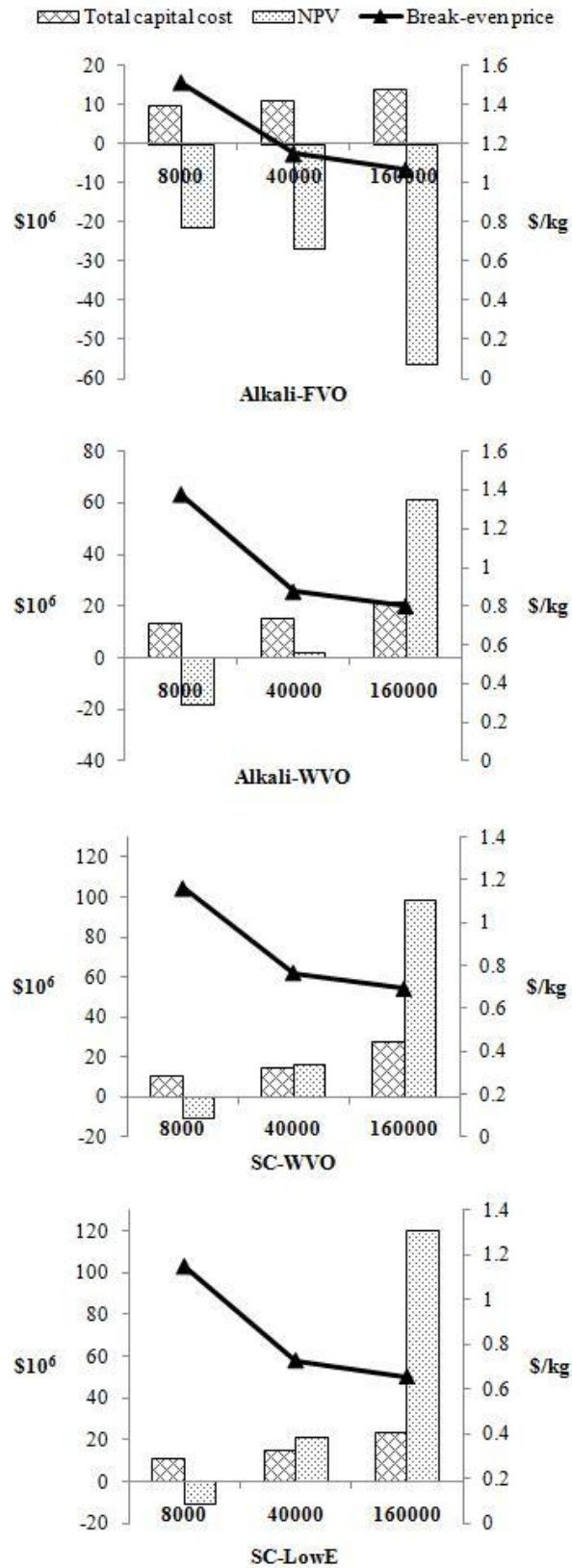


Figure 3.9. Comparison of total capital cost, NPV and break-even price of each process (x-axis: capacity in tonnes biodiesel/yr)

As the capacity diminished to 8,000 tonnes biodiesel/yr, the NPVs of all the four processes became negative values, indicating the production capacity was too small to make enough revenue to recover total capital investment. At the capacity of 160,000 tonnes biodiesel/yr, every process showed high NPVs except for Alkali-FVO process. Due to high prices of the raw materials compared to biodiesel price in Alkali-FVO, the total manufacturing costs for biodiesel were higher than the revenue from biodiesel sales at every capacity, resulting in a lower NPV in Alkali-FVO process as the capacity was expanded. It is noticeable in Figure 3.9 that total capital costs are slow to increase as the production capacity increases, implying a larger production capacity augments profitability of a biodiesel production process because of its relatively small amount of capital costs, conforming to the trend reported by Kasteren and Nisworo (2007). However, compared with the results in Kasteren and Nisworo (2007), the capital costs for the capacity of 8,000 tonnes/yr of this study are thought to be overestimated. The reason for the highly estimated total capital costs appeared to be derived from the costs for instrumentation and indirect costs. As seen in Table A.1, for the four types of processes, the purchased equipment costs increased 3.9–7.7 times as the capacity increased 20 folds (from 8,000 to 160,000 tonnes/yr); whereas, the other costs such as equipment setting, piping, civil, instrumentation, electrical, insulation, others, contract fee, and contingencies increased less than double. Furthermore, these costs were given a great deal of weight on the total capital investments at the capacity of 8,000 tonnes/yr. In IPE, the costs for these items can be manipulated by changing material and man-hour indexes for process equipment and installation bulks. Therefore, for the small capacity, the indexes for the items seem to have to be properly adjusted in the stage of specifying general specifications.

3.6.2. Multiple parameters

In Section 3.6.1, the effects of a single parameter on the profitability of the biodiesel production processes were studied. In reality, however, most of the parameters continuously change over time in an irregular way and the actual values of these parameters are occasionally unknown at the time of evaluation. In addition, the parameters are more likely to vary simultaneously. Nevertheless, there are not sufficient research findings on the effect of multiple parameters on the profitability of biodiesel production processes. Therefore, the objective of this section was to obtain further statistical analysis data using the design of experiments to investigate the effects of simultaneous changes in the parameters. The investigated processes were Alkali-FVO, Alkali-WVO, SC-WVO and SC-LowE at the capacity of 40,000 tonnes biodiesel/year.

The design of experiment selected was Response Surface Method (RSM) coupled with Central Composite Design (CCD) using JMP 8 software (SAS Institute Inc.). The parameters investigated were the prices of oil feed, biodiesel and glycerol and the interest rate because the four parameters were

found to have the most significant effects on the NPVs in Section 3.6.1. The response was the NPV of each process. Table 3.15 shows the four parameters at the different levels used in the design of experiments. After experimental tables were made for each of the processes, the NPV was calculated using IPE. The calculated NPVs of Alkali-FVO are shown in Table A.3 in Appendix A. The results of the Analysis of Variance (ANOVA) of Alkali-FVO process are shown Table 3.16.

Table 3.15. The four parameters in central composite design to investigate NPVs of Alkali-FVO, Alkali-WVO, SC-WVO and SC-LowE processes

Variables	Specification	Unit	Level				
			a	-	0	+	A
Oil feed price	FVO	\$/kg	0.1832	0.5496	0.916	1.2824	1.6488
	WVO		0.106	0.318	0.53	0.742	0.954
Biodiesel price	-	\$/kg	0.198	0.594	0.99	1.386	1.782
Glycerol price	pharmaceutical	\$/kg	0.2646	0.7938	1.323	1.8522	2.3814
	crude		0.172	0.516	0.86	1.204	1.548
Interest rate	-	%	0	10	20	30	40

Table 3.16. Analysis of variance and effect tests for each parameter of Alkali-FVO process

Analysis of Variance					
Source	Degree of Freedom	Sum of Squares	Mean Square	F ratio	Prob >F
Model	14	207532.33	14823.7	233.3583	< 0.0001
Effect Tests					
Term	Estimate	Sum of Squares	F ratio	Prob >F	
Biodiesel price	57.9212	80516.909	1267.514	< 0.0001*	
FVO price	-63.9471	98141.507	1544.964	< 0.0001*	
Glycerol price	8.4131	1698.719	26.7416	< 0.0001*	
Interest rate	7.3551	1298.334	20.4386	0.0002*	
Biodiesel price × FVO price	12.7501	2601.051	40.9463	< 0.0001*	
Biodiesel price × Glycerol price	-1.0057	16.185	0.2548	0.6190	
FVO price × Glycerol price	1.0315	17.024	0.2680	0.6101	
Biodiesel price × Interest rate	-24.8585	9887.120	155.6451	< 0.0001*	
FVO price × Interest rate	26.0840	10886.001	171.3697	< 0.0001*	
Glycerol price × Interest rate	-3.5249	198.796	3.1295	0.0914	
Biodiesel price × Biodiesel price	-5.6084	1006.531	15.8450	0.0007*	
FVO price × FVO price	-6.2336	1243.467	19.5749	0.0002*	
Glycerol price × Glycerol price	-0.6793	14.765	0.2324	0.6347	
Interest rate × Interest rate	-0.4300	5.917	0.0932	0.7632	

The predicted formula of NPV was derived from the linear combination of the parameters and the values of their corresponding estimates. The prediction formula of the NPV can be expressed with the fourteen terms in Table 3.16 plus an intercept. The formula can be expressed in a simpler form after eliminating insignificant parameters. Looking for a F -ratio greater than 2 in absolute value or using the 0.05 significance level ($\text{Prob} > F$ is lower than 0.05) is a common rule of thumb for judging the significance of the parameter. After eliminating the insignificant terms, the NPV was finally expressed in coded factors as shown in Equation (3.9).

$$NPV = -26.697 + 57.9212 A - 63.9471 B + 8.4131 C + 7.3551 D + 12.7501 AB - 24.8585 AD + 26.084 BD - 5.6084 A^2 - 6.2336 B^2 \quad (3.9)$$

where $A = (\text{biodiesel price}(\$/\text{kg}) - 0.99)/0.396$, $B = (\text{FVO price}(\$/\text{kg}) - 0.916)/0.3664$, $C = (\text{glycerol price}(\$/\text{kg}) - 1.323)/0.5292$, and $D = (\text{interest rate}(\%) - 20)/10$.

A, B, C and D are expressed in the form of $(X_i - X_{i,0})/\Delta X_i$ where $X_{i,0}$ is the value at level 0 and ΔX_i is a difference value of adjacent levels in Table 3.15. The ranges for biodiesel price, FVO price, pharmaceutical-grade glycerol price, and interest rate are: $0.198 \leq \text{biodiesel price}(\$/\text{kg}) \leq 1.782$, $0.1832 \leq \text{FVO price}(\$/\text{kg}) \leq 1.6488$, $0.2646 \leq \text{pharmaceutical glycerol price}(\$/\text{kg}) \leq 2.3814$, and $0 \leq \text{interest rate}(\%) \leq 40$, respectively.

Figure 3.10 shows the comparison of actual NPV values and those calculated by Equation (3.9). The solid and dotted lines represent the line of perfect fit and the 0.05 significance curve, respectively. As a point stands close to the line of perfect fit, the error between the actual and predicted values becomes zero. The data points in Figure 3.10 are contiguous to the line of perfect fit. The calculated R^2 was 0.99. Therefore, it is expected that the formula can be used as relatively simple means to predict profitability of Alkali-FVO process from any of the combinations of the oil feed price, glycerol price, biodiesel selling price and interest rate within the ranges in Table 3.15.

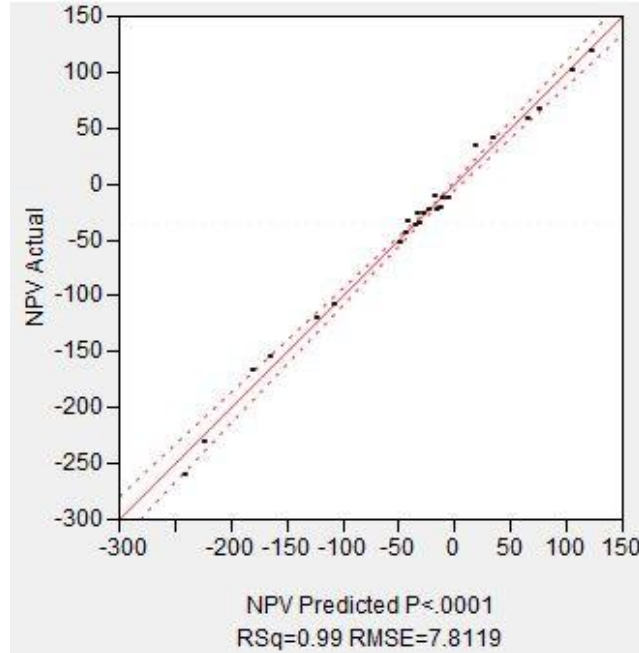


Figure 3.10. A comparative plot of NPV values calculated by the prediction formula to the actual values (- - - : 0.05 significance curve)

The NPV correlations of Alkali-WVO, SC-WVO and SC-LowE processes were developed in the same manner. For Alkali-WVO process,

$$\begin{aligned}
 NPV = & 1.6884 + 50.8435 A - 34.1184 B + 6.3613 C - 8.8331 D + 7.3361 AB - 21.776 AD \\
 & + 14.3841 BD - 4.7346 A^2 - 2.4894 B^2 + 4.1646 D^2
 \end{aligned} \quad (3.10)$$

For SC-WVO process,

$$\begin{aligned}
 NPV = & 14.8393 + 49.9542 A - 31.8746 B + 4.2861 C' - 15.7146 D + 6.5495 AB - 20.99 AD \\
 & + 13.3526 BD - 4.5113 A^2 + 6.1192 D^2
 \end{aligned} \quad (3.11)$$

For SC-LowE process,

$$\begin{aligned}
 NPV = & 19.587 + 49.4227 A - 31.3785 B + 4.2480 C' - 17.4959 D + 6.0322 AB - 20.7246 AD \\
 & + 13.0959 BD - 4.3212 A^2 + 6.5714 D^2
 \end{aligned} \quad (3.12)$$

where $A = (\text{biodiesel price } (\$/\text{kg}) - 0.99)/0.396$, $B = (\text{WVO price } (\$/\text{kg}) - 0.53)/0.212$, $C = (\text{pharmaceutical-grade glycerol price } (\$/\text{kg}) - 1.323)/0.5292$, $C' = (\text{crude glycerol price } (\$/\text{kg}) - 0.86)/0.344$, and $D = (\text{interest rate } (\%) - 20)/10$.

The ranges for biodiesel price, WVO price, pharmaceutical-grade glycerol price, crude glycerol price, and interest rate are: $0.198 \leq \text{biodiesel price (\$/kg)} \leq 1.782$, $0.106 \leq \text{WVO price (\$/kg)} \leq 0.954$, $0.2646 \leq \text{pharmaceutical glycerol price (\$/kg)} \leq 2.3814$, $0.172 \leq \text{crude glycerol price (\$/kg)} \leq 1.548$ and $0 \leq \text{interest rate (\%)} \leq 40$, respectively.

3.7. Conclusion

The economic assessment of the four types of biodiesel production plants: Alkali-FVO, Alkali-WVO, SC-WVO and SC-LowE, was performed, based on the 40,000 tonnes biodiesel production per year capacity. The assessment was done using Aspen Icarus software system which has been successfully field-tested on worldwide commercial chemical plants, but has not been used yet in studies on economic analysis of biodiesel plants.

The results showed that the most significant variable affecting the production cost of biodiesel was the oil feed. It comprised 64-84% of the biodiesel manufacturing costs. The total capital cost of Alkali-WVO process was \$4.5 million higher than Alkali-FVO process due to the pre-treatment process of waste vegetable oil, but the total production cost was around \$10 million lower than Alkali-FVO due to the low cost of the feedstock oil. The supercritical processes, SC-WVO and SC-LowE, had similar values of total capital cost compared to Alkali-WVO. However, they had lower manufacturing costs due to the absence of catalysts and the consequent simpler purification processes. The estimated DPP, NPV and DCFROR indicated that SC-LowE was the most financially promising process, followed by SC-WVO, Alkali-WVO and Alkali-FVO in order. Therefore, further research in developing a pilot-scale continuous supercritical process is expected to be necessary.

Among the parameters investigated in the sensitivity analyses, biodiesel selling price, prices of feedstock oil and by-product glycerol and interest rate were found to be the most significant variables affecting the economic viability of the four processes. The four parameters were further statistically analyzed based on the design of experiment approach using Response Surface Method coupled with Central Composite Design. From the results, prediction formulas of the NPV were derived for each of the four processes. The formulas can be used for various combinations of the four significant parameter values to predict economic feasibility.

Acknowledgments

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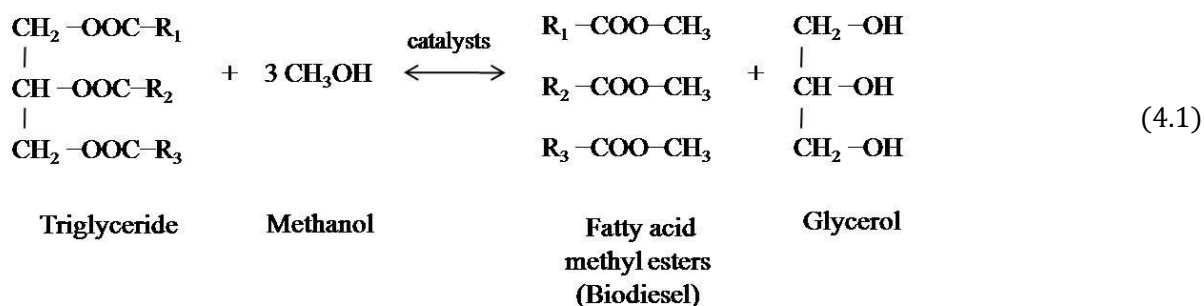
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4. Biodiesel production from used canola oil using supercritical methanol³

4.1. Background

Biodiesel is a mixture of fatty acid alkyl esters, which are generally produced through transesterification using feedstocks containing triglycerides. Equation (4.1) shows the transesterification reaction between triglyceride and methanol that produces fatty acid methyl esters (FAMEs) and glycerol. The conventional method for biodiesel production is transesterification of fresh vegetable oil in the presence of homogeneous base catalysts such as sodium hydroxide and potassium hydroxide in order to achieve reasonable reaction rates and yields. However, it is well known that the conventional base catalytic processes are inefficient when using low-cost feedstocks such as waste vegetable oils because of a large amount of free fatty acids (FFAs) and water present. These impurities are known to consume the alkali catalysts via saponification reaction, thereby decreasing biodiesel yields. Moreover, a mixture of soaps, biodiesel and un-reacted compounds causes emulsification during a washing process which makes it difficult to obtain biodiesel of high quality (Marulanda et al., 2010). The inability to use low-cost feedstocks has recently challenged biodiesel production through the conventional method due to constantly increasing prices of fresh vegetable oils.



Researchers have endeavoured to overcome the weaknesses of the homogeneous catalyzed processes by focusing on developing appropriate heterogeneous solid catalysts. Heterogeneous reactions have been reported to be able to tolerate high FFAs contents in feedstock oils. However, the solid catalysts have low reactivity rates due to mass transfer limitation between liquid and solid phases of reactants and catalysts (Tan et al., 2010).

³ A version of this chapter is in preparation for submission for publication. Lee, S., Posarac, D. and Ellis, N. (2010). Biodiesel production from used canola oil using supercritical methanol.

In a recent development, it is reported that the low-cost feedstock might be successfully used in the transesterification reaction using methanol under its supercritical conditions. Saka and Kusdiana (2001) reported that over 95% conversion to methyl esters was achieved in 240 seconds using rapeseed oil and supercritical methanol. It was also reported that the dielectric constant of liquid methanol which tends to decrease in the supercritical state increased the solubility of oil in methanol, resulting in a single phase methanol/oil system. This process also has an advantage in purification of the product mixture due to the absence of a catalyst.

In later studies, effects of water and FFAs in feedstocks on supercritical biodiesel production have been studied (Kusdiana and Saka, 2004). It was reported that complete conversions were always achieved with the water contents up to 36 wt% in the rapeseed oil. In addition, as treated by supercritical methanol, the FFAs in the oil were simultaneously converted to methyl esters by esterification reaction. Since this process can utilize low-cost feedstocks, research on supercritical biodiesel production is actively in progress. Table 4.1 summarizes the reported experimental data of biodiesel production using supercritical methanol.

As shown in Table 4.1, supercritical reaction has been investigated in a wide range of reaction conditions ($T = 250\text{--}450^\circ\text{C}$; $P = 9.6\text{--}43\text{ MPa}$; residence time = 2–40 min). The consequent biodiesel yields range from 66% to 100%. In the early stage of research on supercritical transesterification reaction, high temperature ($>350^\circ\text{C}$) and pressure ($>20\text{ MPa}$) conditions were used to obtain desirable methyl ester yields in short time. However, more recently, it has been proposed to use relatively moderate reaction conditions due to thermal degradation of FAMEs at high temperature. Varma and Madras (2007) reported thermal decomposition of FAMEs at the reaction temperatures above 350°C .

Table 4.1. Summary of reported experiment data of biodiesel production using supercritical methanol

Oil (co-solvent)	T (°C)	P (MPa)	MeOH/oil ratio (mol/mol)	Residence time (min)	B/C^a	Yield (%)	Reference
Waste oil	287	N/A	41	30	B	99.6	Demirbas (2009a)
Linseed	250	N/A	41	8	B	98	Demirbas (2009b)
Soybean	100–320 ^b	32	40	25	C	96	He et al. (2007)
Rapeseed	350	43	42	4	B	95	Saka and Kusdiana (2001)
Sunflower	400	20	40	40	B	96	Madras et al. (2004)
Canola	420–450	40	11–45	4	C	~100	Iijima et al. (2004)
Coconut	350	19	42	6.67	C	95	Bunyakiat et al. (2006)
Palm	350	19	42	6.67	C	96	
Castor/linseed	350	20	40	40	B	~100	Varma and Madras (2007)
Chicken fat	400	41.1	6	6	B	88	Marulanda et al. (2010)
Palm oil	372	15–25	40	16	B	81.5	Tan Gui et al. (2010)
Fodder radish oil	317	10–18	39	27	B	97.0	Valle et al. (2010)
Soybean (C₃H₈/MeOH=0.05)^c	280	12.8	24	10	B	98	Cao et al. (2005)
Soybean (C₃H₈/MeOH=0.05)^c	288	9.6	65.8	10	B	99	Hegel et al. (2007)
Soybean (CO₂/MeOH=0.1)^c	280	14.3	24	10	B	98	Han et al. (2005)
Soybean (CO₂/MeOH=0.1)^c	350–425	10–25	3–6	2–3	C	~100	Anitescu et al. (2008)
Palm oil (heptane/MeOH=0.2)^c	280	15	30	20	B	66	Tan et al. (2010)
without heptane	360	22	30	20	B	80	

^a B-batch or C-continuous

^b Gradual heating

^c Molar ratio of co-solvent to methanol

Imahara et al. (2008) reported that reaction temperature in a supercritical methanol process should be lower than 300°C, preferably 270°C as reaction pressures are >8.09 MPa to maximize the yield. Furthermore, poly-unsaturated fatty acid methyl esters such as methyl linoleate (C18:2) and

methyl linolenate (C18:3) were more susceptible to thermal decomposition compared to mono-unsaturated (methyl oleate, C18:1) and saturated methyl esters (methyl stearate, C18:0). The thermal behaviour of soybean biodiesel decomposition was investigated with thermogravimetry (TG) and mass spectrometry to the thermogravimetry (TG-MS) instruments by Chien et al. (2009). These authors reported that smaller esters, CO₂, aldehydes and many hydrocarbon fragments can be found as the expected by-products from biodiesel decomposition.

To avoid the thermal degradation of FAMEs, addition of the third component (“co-solvent”) such as propane (Cao et al., 2005; Hegel et al., 2007), CO₂ (Han et al., 2005; Anitescu et al., 2008) and heptane (Tan et al., 2010) has been proposed. As can be seen in Table 4.1, the presence of a third component appeared to decrease reaction temperatures and pressures to achieve the similar biodiesel yields from reactions without any additional component. These authors claimed that the co-solvent increased mutual solubility between triglyceride and methanol, thereby aiding transesterification reaction. Imahara et al. (2009), however, reported that the improvement in reaction rate due to the third component addition was merely the effect of the increased reaction pressure.

Table 4.1 also shows that most of the research studies were conducted with food-grade vegetable oils despite the emphasized favourable properties of the supercritical process with low-quality feedstocks. Although scarce, the research on supercritical transesterification with waste oil showed very encouraging biodiesel yields (Demirbas, 2009a; Marulanda et al., 2010). Results of economic analyses performed on industrial-scale supercritical transesterification processes were also encouraging compared to the conventional biodiesel production processes (Anitescu et al., 2008).

To decrease the production cost of biodiesel with a supercritical method compared to the conventional alkali-catalyzed processes, low-cost feedstocks and reasonable yields must be obtained. Accordingly, the main objective of this research is to conduct a supercritical transesterification study with waste canola oil under relatively moderate temperature and pressure conditions to avoid thermal degradation of FAMEs. Any other additional components other than raw materials were not used. The effects of process variables such as reaction time, reaction temperature and weight ratio of methanol to oil were assessed. The interactions of the variables were also studied via analysis of variance (ANOVA). Based on the ANOVA results, an empirical mathematical model correlating a FAME yield from the waste canola oil to the investigated variables was developed. Product analyses indicated the by-product glycerol obtained from transesterification reaction went through side reactions such as decomposition and etherification.

4.2. Experimental

4.2.1. Materials and apparatus

Waste canola oil was provided by Place Vanier residence on the University of British Columbia (UBC) Vancouver campus, Canada. The waste oil was brownish in colour compared to the fresh one provided from the same place, as shown in Figure 4.1, but transparent in appearance as any solid impurities were filtered. Methanol (99.9%) was purchased from Fisher Scientific. For the gas chromatography (GC) analysis of product samples, *n*-heptane (purity $\geq 99.0\%$) was purchased from Fluka and used as solvent. An analytical grade kit with various FAMES including methyl myristate (C14:0), methyl palmitate (C16:0), methyl stearate (C18:0), methyl oleate (C18:1), methyl linoleate (C18:2), methyl linolenate (C18:3), methyl arachidate (C20:0) and methyl behenate (C22:0) for with GC standards was purchased from Supelco. Methyl palmitoleate (C16:1) and methyl cis-11-eicosenoate (C20:1) (99% purity) were purchased from Sigma-Aldrich. Methyl heptadecanoate ($\geq 99.0\%$, Sigma) was used as the internal standard.

Supercritical reaction was carried out in a batch-type autoclave reactor of 100 ml (Autoclave Engineers) made of Stainless Steel 316. A temperature and pressure controller (Autoclave Engineers) was used in order to carry out reactions in isothermal conditions. A material for the other parts of experimental settings such as tubing, valves and a cylinder were made of Stainless Steel 316.



Figure 4.1. Fresh canola oil (left) and waste canola oil (right) from UBC Place Vanier residence

4.2.2. Procedures

Figure 4.2 shows a schematic of the experimental setup for supercritical biodiesel production. To avoid any reaction that would occur between methanol and waste canola oil during heating to reaction temperature, they were charged separately into the autoclave reactor and the oil container, respectively. The waste canola oil of 8.5 g (± 0.5 g) was initially injected into the oil container made of Stainless Steel 316 using a pipette through Valve 2. Once the oil was loaded, Valve 2 was closed. Then, the waste oil was preheated to around 110°C by a heating tape wound on the container before it was injected into the reactor. Meanwhile, a necessary amount of methanol (depending on the desired weight ratio of methanol to oil) was placed into the autoclave reactor vessel and the vessel was closed. Reaction temperature was set by the reactor controller so that the heating jacket on the reactor vessel started heating the vessel to the reaction temperature. The pressure inside the reactor was monitored with a pressure gauge attached to the reactor. The reactor temperature was measured by a thermometer inserted in the reactor and monitored in real time on the controller display.

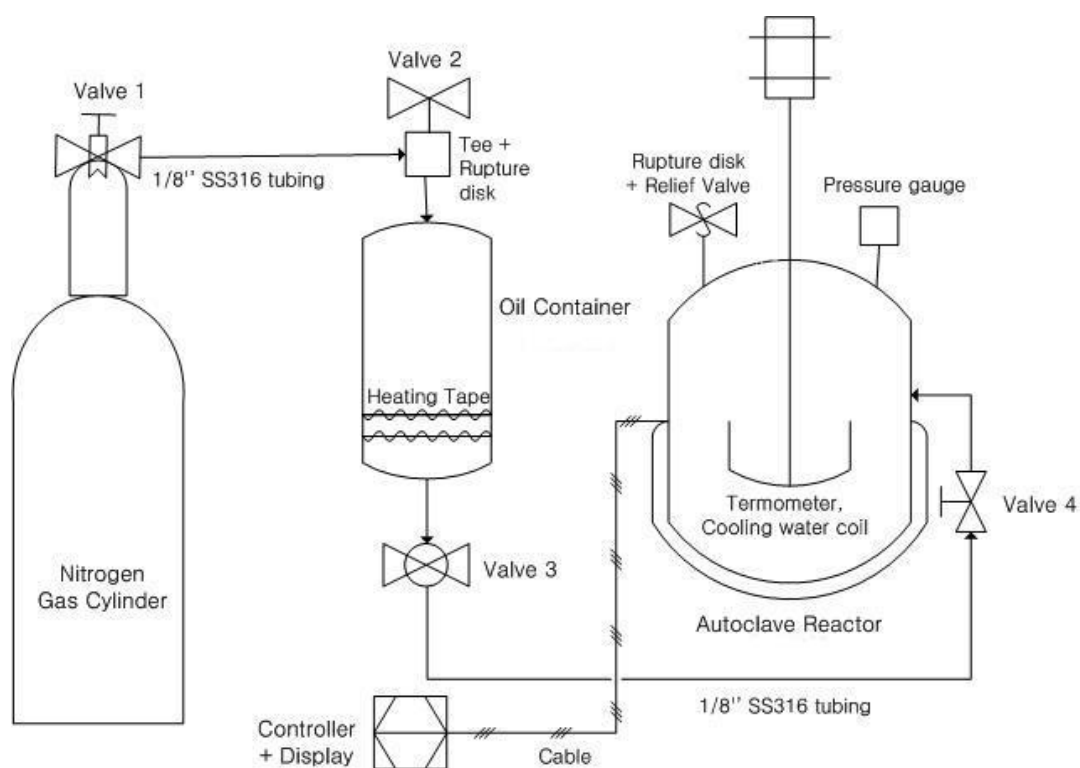


Figure 4.2. The experimental setup for supercritical biodiesel production

The pressure inside the autoclave reactor was monitored as both the waste oil and methanol reached the desired temperatures. It was 3–4 MPa lower than the nitrogen gas pressure discharged from the gas cylinder, and the pressure difference was used to flush the waste oil into the autoclave reactor. By opening Valve 1 on the gas cylinder, both nitrogen gas and the preheated waste oil flowed

to Valve 4. Then, the nitrogen gas and oil were injected into the autoclave as Valve 4 opened. The reactor temperature dropped around 10°C from the original temperature with the injection of waste oil, but increased back within a minute due to relatively small amount of the oil. Meanwhile, the autoclave reactor was pressurized with nitrogen until the reaction pressure of 10 MPa was achieved, and then Valve 4 was closed. Reaction time was measured from the time when the oil and nitrogen were injected. The reactor temperature was kept at $\pm 4^\circ\text{C}$ during the reaction by the controller, while chilled water immediately kicked in through the coil to quench the reaction after the desired reaction time. As a result, the temperature inside the reactor dropped below 200°C within 30 seconds. Consequently the pressure also dropped. Once the reactor temperature dropped to 20°C, the remaining pressures inside the oil container and autoclave were released by opening Valve 2 and Relief valve. Then, the reactor vessel was opened and the product was analyzed by gas chromatography.

4.2.3. Analytical procedure

The acid number (mg KOH/g) of the waste oil was determined by titration with potassium hydroxide solution (0.1 wt% KOH in distilled water). 10 ml of 2-propanol was added to 1 g of the waste canola oil in a 250 ml flask. A few drops of phenophtalein indicator were added into the flask and it was titrated with the KOH solution until a distinct and sharp end point was noted. The solution in the flask was kept stirred using a magnetic stirrer during titration. The procedure was repeated three times and the averaged value was determined as the final acid number of the waste canola oil. Since the waste canola oil contained no free acids other than fatty acids, the acid number was converted to percent free fatty acids by a suitable factor (1.99) as described in ASTM D5555-95.

The water content for the waste oil was assessed by means of Karl Fischer (KF) volumetric titration using 794 Basic Titrino apparatus from Metrohm shown in Figure 4.3. The KF volumetric reagent was purchased from Fisher Scientific. Extra dry methanol (water <50 ppm) purchased from Acros Organics was used as the solvent. Before analyzing the waste oil sample, the titer value of the KF reagent (the amount of water titrated by 1 ml of KF reagent) was checked using distilled water as recommended by Metrohm. Triplicate runs were performed on the water content analysis for all samples. The details of the procedure and titration reports are included in Appendix B.



Figure 4.3. Karl Fisher volumetric titration apparatus for measurement of water content

The composition and quantity of methyl esters in the product samples were determined using gas chromatography (HP 5890 Series II) equipped with a flame ionized detector (FID). A Varian WCOT fused silica capillary column (CP8723) of $60\text{ m} \times 0.25\text{ mm} \times 0.25\text{ }\mu\text{m}$ film thickness was used with helium as carrier gas. The temperature program started at 180°C (held for 5 min) and ramped to 220°C at $5^\circ\text{C}/\text{min}$. The temperature was held at 220°C for 15 min and ramped to 250°C at $5^\circ\text{C}/\text{min}$. The holding time at the final temperature (250°C) was 5 min. The temperatures of the injector and detector were set at 290°C and 300°C , respectively. A standard sample was prepared in heptane with a known composition of methyl esters, and $1\text{ }\mu\text{l}$ of the prepared solution was injected into the column.

Various methyl esters in the product samples were identified by comparing the retention times of each FAME to the ones obtained from the standard sample as shown in Figures 4.4 and 4.5. The methyl esters were then quantified using the ratio of the areas under the peaks of each methyl ester to the internal standard and the known concentration of internal standard. The fatty acid profile of biodiesel produced from the waste canola oil is compared with literatures in Table 4.2.

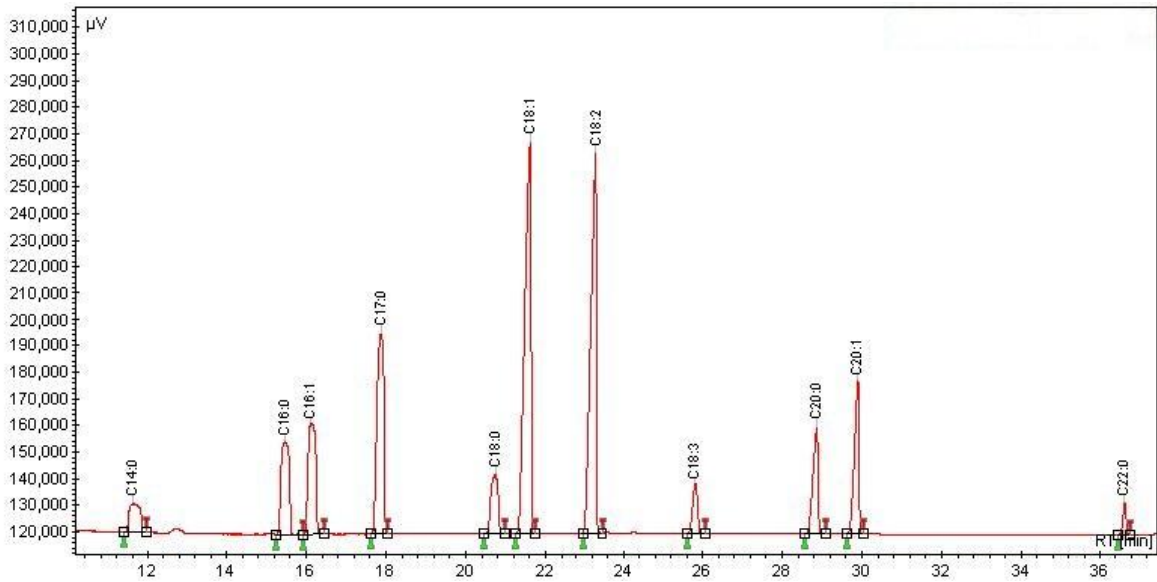


Figure 4.4. Gas chromatogram of the standard sample for peak identification of methyl esters

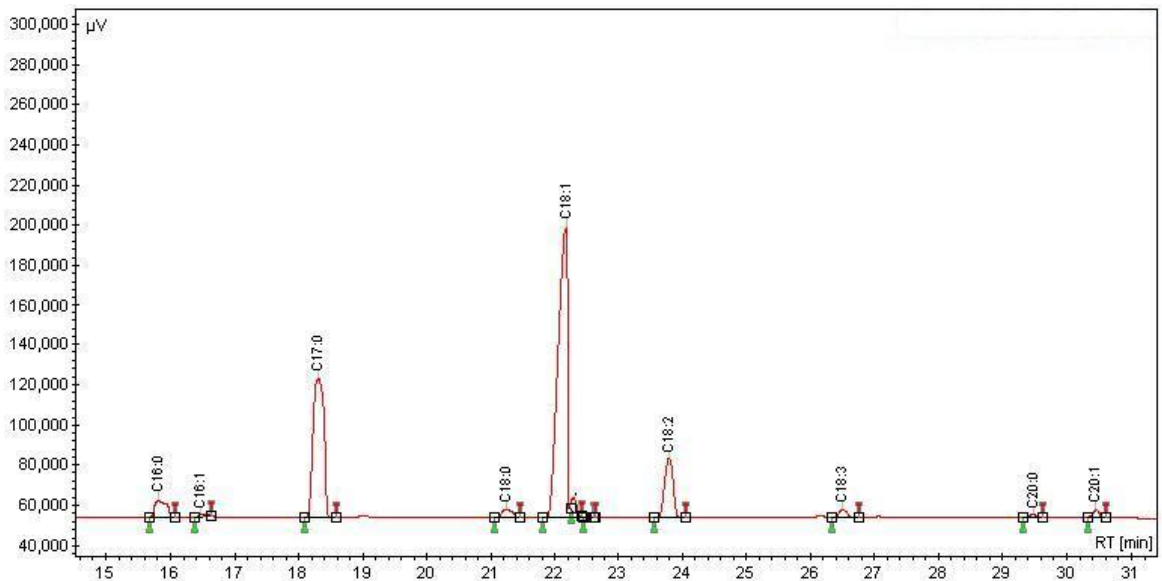


Figure 4.5. Gas chromatogram of a product sample from the waste canola oil

Table 4.2. Profile of fatty acid methyl esters in biodiesel from different feedstocks

Feedstock for biodiesel	Fatty acid methyl ester composition in biodiesel (wt%)								
	C16:0	C16:1	C18:0	C18:1	C18:2	C18:3	C20:0	C20:1	C22:0
Corn oil ^a	11.8	0.0	2.0	24.8	61.3	0.0	0.0	0.0	0.0
Safflower seed oil ^a	7.3	0.0	1.9	13.6	77.2	0.0	0.0	0.0	0.0
Soybean oil ^a	13.9	0.0	2.1	23.2	56.2	4.3	0.0	0.0	0.0
Jatropha oil ^a	14.9	0.0	9.5	40.5	34.7	0.3	0.0	0.0	0.0
Rapeseed oil ^a	3.5	0.0	0.9	64.1	22.3	8.2	0.0	0.0	0.0
Chicken fat ^b	24.6	5.8	8.4	46.1	2.4	0.0	0.0	0.0	0.0
Waste canola oil ^c	3.2	0.0	0.3	84.3	10.0	0.8	0.5	0.8	0.2

^a From Gopinath et al. (2009)

^b From Marulanda et al. (2010)

^c From this study (run 19)

The yield of FAME was calculated by Equation (4.2).

$$\text{Yield (\%)} = \frac{\text{Total weight of methyl esters produced}}{\text{Total weight of oil used for reaction}} \times 100 (\%) \quad (4.2)$$

4.2.4. Statistical analysis

The process parameters used in this study are: the reaction time; reaction temperature; and weight ratio of methanol to waste oil. The effect of the process parameters on the yield of methyl esters was investigated by using customized design of experiments available in JMP8 (SAS institute Inc.). Table 4.3 shows the experimental conditions along with the yields of methyl esters. The experimental order was done randomly. Upon completion of all the experimental runs, the obtained responses (the yield of FAMEs) were then fitted in a quadratic model using regression analysis.

Table 4.3. Experimental conditions and the yields of methyl esters from the waste canola oil

Run	Time (min)	Temperature (°C)	MeOH/oil (w/w)	Yield (%)
1	15	240	1	1.5
2	15	240	1.5	3.2
3	15	240	2	3.8
4	15	255	1	2.2
5	15	270	1	8.9
6	15	270	2	13.6
7	30	240	1	1.5
8	30	240	2	12.3
9*	30	255	1.5	11.4
10*	30	255	1.5	13.6
11*	30	255	1.5	12.7
12	30	255	2	18.5
13	30	270	1.5	60.9
14	45	240	1	4.4
15	45	240	1.5	16.5
16	45	240	2	24.5
17	45	255	1.5	62.8
18	45	270	1	96.4
19	45	270	2	101.6

* Triplicate runs for reproducibility

4.3. Results and discussion

4.3.1. FFA and water contents of waste canola oil

The FFA and water contents of the waste canola are shown in Table 4.4. The FFA content was 4.0 wt% and the water content was less than 0.01 wt%. Other than Karl Fischer titration, thermogravimetric analysis (TGA) of the oil was performed to check volatile components as well as water present in the oil. As shown in Figure 4.6, the weight loss while the oil was heated at 110°C for an hour was 0.009%, indicating the oil contained a negligible amount of those components. Canakci (2007) reported that the FFA levels in waste oils can vary from 0.7% to 41.8%, and moisture contents from 0.01% to 55.38% as analyzing restaurant grease and animal fat samples from 13 different locations. The FFA and water contents of the waste canola oil in this study were within those ranges, but were relatively less contaminated compared to those having FFA and water contents greater than 10%.

Table 4.4. Comparison of the yield of methyl esters from used frying oil, waste palm oil and waste canola oil (Source: Kusdiana and Saka (2004))

Vegetable oil	FFA content (wt%)	Water content (wt%)	Yield of methyl esters (%)	
			Alkali-catalyzed	Supercritical
Used frying oil	5.6	0.2	94.1 ^a	96.9 ^b
Waste palm oil	>20.0	>61.0	No reaction ^a	95.8 ^b
Waste canola oil (this study)	4.0	<0.01	N/A	101.6 ^{c,d}

^a Reaction conducted at 60°C/1 atm for 1 hr using 1.5 wt% sodium hydroxide (Kusdiana and Saka, 2004)

^b Reaction conducted at 350°C/43 MPa for 4 min (Kusdiana and Saka, 2004)

^c The yield from run 19 of this study

^d Sample calculations of yield are shown in Appendix C

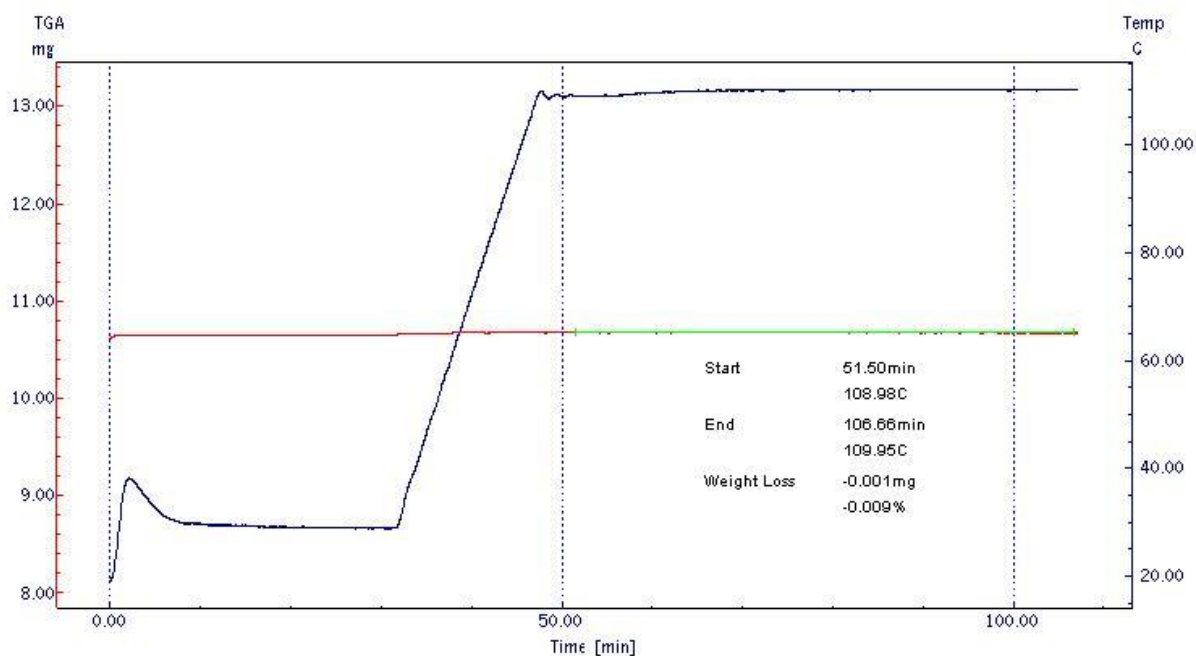


Figure 4.6. Thermogravimetric analysis of waste canola oil used in the supercritical experiments

The FFA and water contents of the waste canola oil were compared to those of used frying oil reported in Kusdiana and Saka (2004). As shown in Table 4.4, the FFA concentrations of the waste canola oil and used frying oil were similar to each other, whereas the water content was relatively low in the waste canola oil. The yield from this study was greater than 96.9% of the used frying oil supposedly because of the longer reaction time. Table 4.4 also compares the yields of methyl esters from alkali-catalyzed and supercritical reactions. The results showed that supercritical reaction had a higher yield than the alkali-catalyzed reaction. The difference of the yields became greater as the waste

palm oil was used as the feedstock. With FFA >20.0 wt% and water >61.0 wt% in the oil, the supercritical reaction still had a yield as high as 95.8% whereas no reaction occurred in the alkali-catalyzed reaction.

4.3.2. Supercritical transesterification of waste canola oil

Table 4.3 shows that the FAME yield is very low at the temperature of 240°C. In the experiments 14–16, despite the reaction time was as long as 45 min, the yields were low, ranging from 4.4% to 24.5%. The result can be explained by the critical temperature of a triglyceride–methanol mixture. The critical temperature of pure methanol is 239.45°C, and the methanol exists in the supercritical state above the temperature. As triglycerides are added to the methanol, the critical temperature required to achieve a single phase state of the mixture increases. Hegel et al. (2007) reported that the lower methanol/oil ratio increased the temperature required to bring the system into a supercritical phase. In this sense, the mixture of the waste oil and methanol will need to be at the temperature greater than 239.45°C to be in the supercritical single phase. Therefore, the experiments conducted at 240°C would have not reached the state, resulting in the low yields.

The high yields over 96% were obtained from the 45 min reactions at 270°C/10 MPa (runs 18 and 19). The result from run 18 was comparable to the value reported by Demirbas (2009a) in Table 4.1 where 99.6% of yield was obtained after 30 min of reaction at 287°C.

The reaction time was also found to be an important factor to the FAME yield. When the reaction time was as short as 15 min (runs 1–6), the FAME yields were in the range of 1.5–13.6%. Although the temperature and weight ratio were at the highest levels of this study, the yield was only 13.6% in run 6. Comparing with the reported values in Saka and Kusdiana (2001), Marulanda et al. (2010) and Tan et al. (2010) where 95%, 88% and 81.5% yields were obtained within 4–16 min, the 13.6% yield was very low. The reason was, as stated previously, much more moderate reaction conditions of this study compared to the literatures as shown in Table 4.1. To achieve a reasonable yield in the mild reaction conditions, therefore, it is unavoidable to trade off an increase in the reaction time. Consequently, if the moderate reaction conditions are applied, it will bring a raise in the expense for a larger size of reactor in a continuous supercritical process.

The yields from the replicate runs 9, 10 and 11 ranged from 11.4% to 13.6% with an average value of 12.6%. The standard deviation was calculated to be 0.903%, indicating the experimental error was small.

The experimental results shown in Table 4.3 were analyzed using analysis of variance (ANOVA) to develop a regression model for the FAME yield. The results of ANOVA are summarized in Table 4.5.

Table 4.5. The results of ANOVA for the supercritical biodiesel production experiments

Analysis of Variance					
Source	DF	Sum of Squares	Mean Square	F ratio	Prob >F
Model	9	17337.750	1926.42	37.9156	<0.0001*
Error	9	457.272	50.81		
Total	18	17795.023			
Effect Tests					
Source	DF	Sum of Squares	Mean Square	F ratio	Prob >F
Time	1	7369.392	7369.392	145.044	<0.0001*
Temperature	1	6939.654	6939.654	136.586	<0.0001*
Weight ratio	1	126.206	126.206	2.484	0.1495
Time × Temperature	1	3488.731	3488.892	68.668	<0.0001*
Time × Weight ratio	1	74.406	74.406	1.464	0.2570
Temperature × Weight ratio	1	26.264	26.264	0.517	0.4904
Time × Time	1	172.045	172.045	3.386	0.0989
Temperature × Temperature	1	312.723	312.723	6.155	0.0349*
Weight ratio × Weight ratio	1	16.222	16.222	0.319	0.5858

The *F* ratio in Table 4.5 represents the ratio of the mean square for the model divided by the mean square for error. The ANOVA results showed that the probability if the null hypothesis is true, that is the chance that a larger *F* ratio would occur due to noises in the experiment was less than 0.0001 (= 0.01%). Subsequently, the significant parameters to the FAME yield were identified from the effect tests. Among the single parameters, reaction time and temperature were found to be significant as their values of Prob >*F* are less than 0.0001. ‘Time × Temperature’ parameter also showed Prob >*F* value less than 0.0001, indicating there is a considerable interaction effect between the reaction time and reaction temperature on the FAME yield. On the other hand, among the 9 parameters analyzed, the Prob >*F* values of ‘Weight ratio × Weight ratio’, ‘Temperature × Weight ratio’ and ‘Time × Weight ratio’ were the highest as of 0.5858, 0.4904 and 0.2570, respectively, indicating they are less significant to the FAME yield. The parameters were involved with “Weight ratio” in common. The methanol/oil weight ratios of 1.0, 1.5 and 2.0 in this study are approximately equivalent to 28, 41 and 55 molar ratios, respectively, if triolein is regarded as triglyceride of the waste oil. As shown in Figure 4.7, the effect of increasing amount of methanol in supercritical reactions is great in lower molar ratios, but diminished in high molar ratios (>35). Thus, in this study, the effect of weight ratio on the FAME yield was assessed relatively less significant than reaction temperature and reaction time.

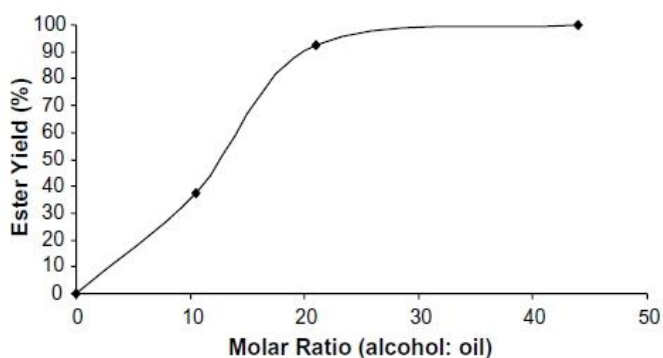


Figure 4.7. Effect of molar ratio on methyl ester yield from supercritical transesterification of Jatropha oil (Hawash et al., 2009)

A second order quadratic equation to predict a FAME yield was developed, therefore, excluding the three insignificant parameters. The prediction formula is shown in Equation (4.4).

$$\text{Yield} = 17.76 + 25.64 A + 23.73 B + 3.41 C + 19.00 AB + 7.29 A^2 + 8.94 B^2 \quad (4.4)$$

where $A = (\text{Time (min)} - 30)/15$, $B = (\text{Temperature (}^\circ\text{C)} - 255)/15$ and $C = (\text{Weight ratio} - 1.5)/0.5$.

A, B and C are expressed in the form of $(X_i - X_{i,0})/\Delta X_i$. Since the time, temperature and weight ratio varied at three different levels in Table 4.3, $X_{i,0}$ and ΔX_i represent the average value of the three levels and the difference value between adjacent levels, respectively. The ranges for time, temperature, and weight ratio are: $15 \leq \text{time (min)} \leq 45$, $240 \leq \text{temperature (}^\circ\text{C)} \leq 270$, and $1 \leq \text{weight ratio} \leq 2$, respectively.

The coefficients in Equation (4.4) indicate the relative significance of the corresponding terms to the yield. Therefore, according to the Equation (4.4), the reaction time has the highest effect on the FAME yield among the variables, followed by reaction temperature and the interaction of the time and temperature in order. The FAME yields predicted by the formula developed in Equation (4.4) were then compared to the actual yields as shown in Figure 4.8. The R^2 value of the fit was 0.9680 which was close to unity.

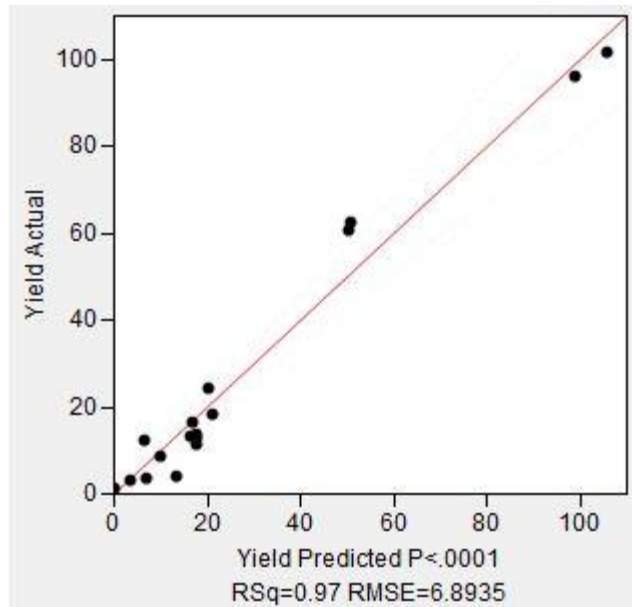


Figure 4.8. A plot for comparison of the actual yields and predicted yields

4.3.3. Interaction effects on the FAME yield

In Figure 4.9, the effects of reaction temperature and time on the FAME yield at a fixed methanol/oil weight ratio of 2 are shown as two-dimensional (top) and three-dimensional plots (bottom). According to the ANOVA results in Table 4.5, this interaction term was found to significantly affect the FAME yield. As seen in Figure 4.9, at 270°C, the yield considerably increased as increasing reaction time. However, the effect of reaction time was less notable at lower reaction temperature. Therefore, to achieve a reasonable transesterification reaction rate, a reaction temperature should be >240°C. The response surface plot in Figure 4.9 showed that the maximum yield could be obtained at 270°C with a reaction time of 45 min.

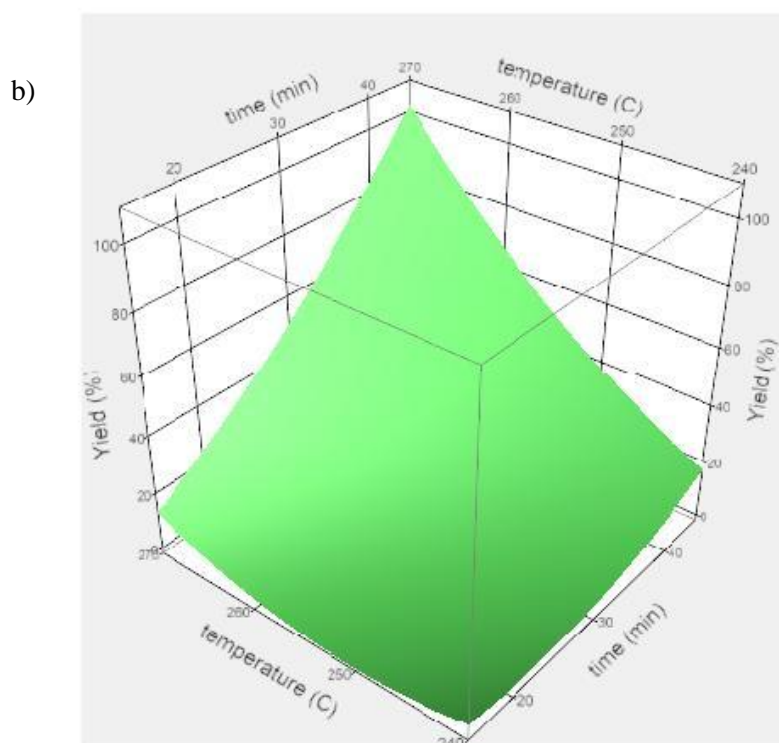
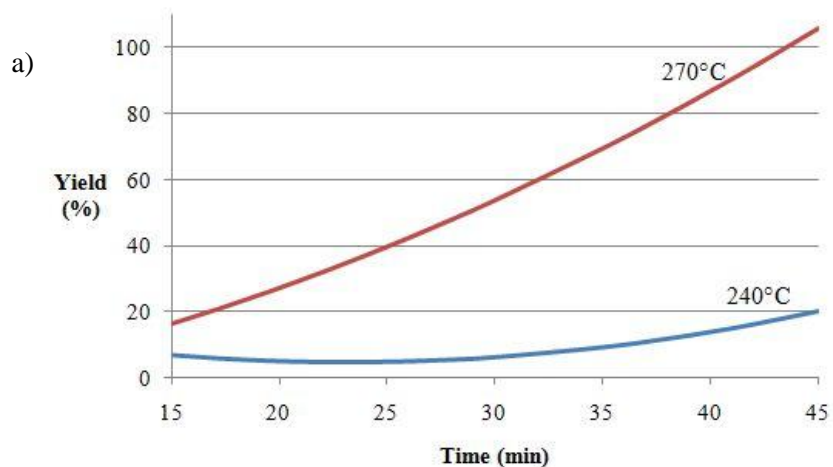


Figure 4.9. a) Two-dimensional and b) response surface plots of FAME yield against reaction time and temperature as predicted by Equation (4.4)

In Figure 4.10, the effects of reaction time and weight ratio on the yield of FAME at fixed reaction temperature of 270°C are shown as a two-dimensional plot. Figure 4.11 shows the effects of reaction temperature and weight ratio on the yield with the reaction time kept constant at 45 min. In the two figures, it is shown that doubling the weight ratio from 1 to 2 resulted in only a minor increase in the FAME yields. Both of the two graphs showed similar trends for the FAME yield, indicating the

interactions of ‘time × weight ratio’ and ‘temperature × weight ratio’ are insignificant. It corresponded to the ANOVA results presented in Table 4.5 where the Prob >F values of the two interaction parameters are 0.2570 and 0.4904, respectively.

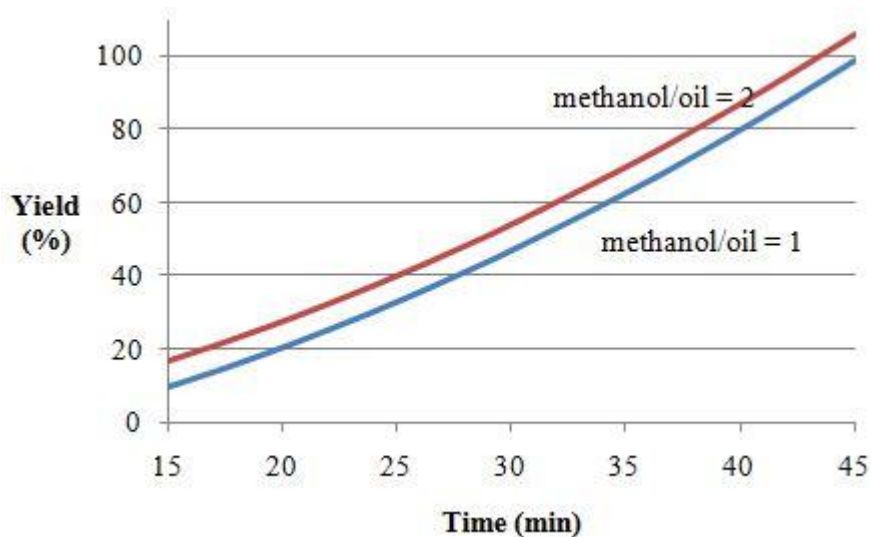


Figure 4.10. FAME yields against reaction time and weight ratio predicted by Equation (4.4)

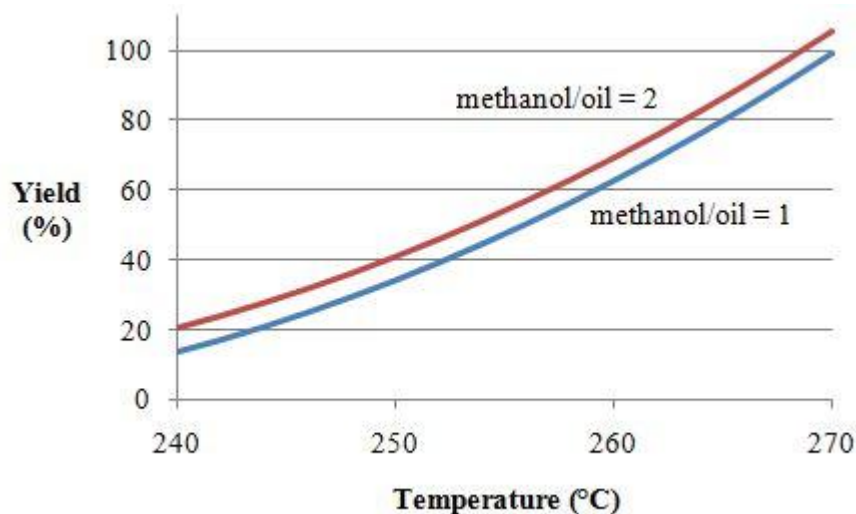


Figure 4.11. FAME yields against reaction temperature and weight ratio predicted by Equation (4.4)

4.3.4. Glycerol reactions

A noteworthy feature was found in the products from runs 18 and 19. Figure 4.12 shows a single yellowish liquid phase formed in the product from run 18. The single liquid phase products contrasted with those from the other experiments where two liquid phases were observed with a bright

yellowish layer at the top and dark yellowish layer at the bottom. Both run18 and run19 were conducted at the highest temperature (270°C) and for the longest reaction time (45 min) of this study. Thus, it was postulated that some kinds of side reactions had occurred under the supercritical methanol conditions and relatively small amounts of methanol and glycerol left at the end of the process remaining in solution with biodiesel.

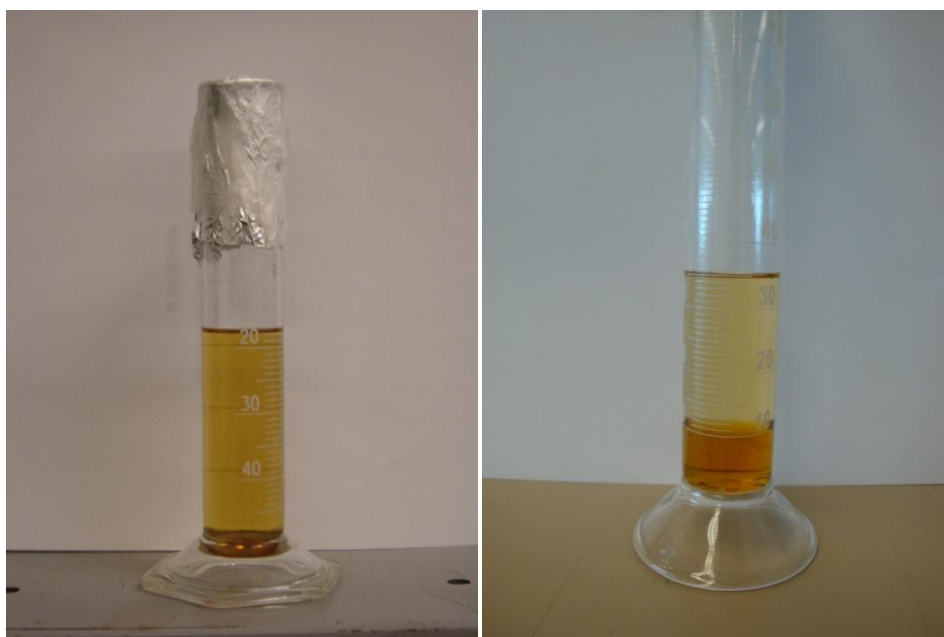
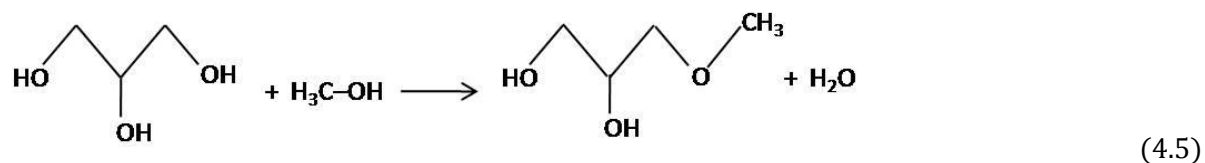


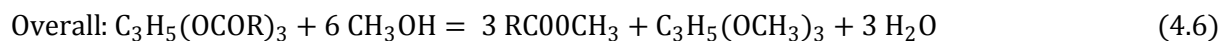
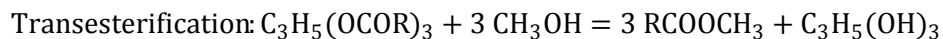
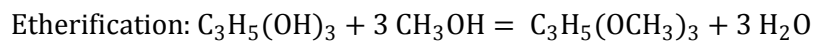
Figure 4.12. The comparison of products with a single phase (left, run18) and with two phases (right, run17)

Recently, Marulanda et al. (2010) reported that a product with a single yellowish liquid phase was obtained from supercritical transesterification of chicken fat. As a reason for the disappearance of glycerol/methanol phase, it was claimed that glycerol was thermally decomposed or reacted with excess methanol according to the following type of reaction in Equation (4.5).



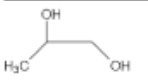
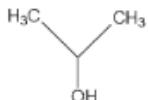
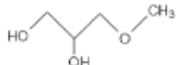
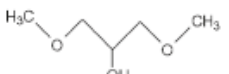
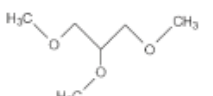
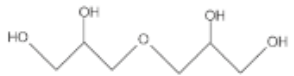
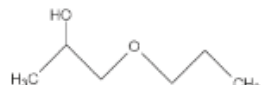
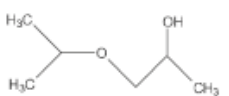
The glycerol methanolysis (etherification) reactions can produce methyl, dimethyl and trimethyl glycerol ethers, which can be directly used as a fuel additive in the biodiesel fuel, positively affecting the viscosity and the pour point (Noureddini et al., 1998; Aimaretti et al., 2009; Marulanda et

al., 2010). As coupling a complete etherification reaction with the transesterification reaction, the following global reaction equation can be written (Marulanda et al., 2010).



Equation (4.6) shows that water present in products can be related to the disappearance of glycerol. The authors also proposed several organic compounds as glycerol decomposition products and identified them by gas chromatography–mass spectroscopy (GC-MS). The identified decomposition products of glycerol are shown in Table 4.6.

Table 4.6. Identified glycerol reaction products by GC-MS (Marulanda et al., 2010)

glycerol decomposition products	name
	1,2-propanediol
	2-propanol
	1,2-propanediol, 3 methoxy
	2-propanol, 1,3 dimethoxy
	propane, 1,2,3 trimethoxy
	diglycerol
	2-propanol, 1 propoxy
	2-propanol 1(1-methylethoxy)

Side reactions of glycerol and methanol were also reported by Aimaretti et al. (2009). It was reported that, as reacting refined soy oil with supercritical methanol for 30–90 min reaction times, free glycerol contents continuously decreased from 0.054% to 0.031%, whereas water contents in the

reacting mixture increased. The authors postulated that glycerol could be decomposed into smaller molecules such as acrolein, acetaldehyde and acetic acid, generating water as a product. Another option was etherification with methanol to produce glycerol ethers as previously shown in Equation (4.5).

The presence of the glycerol reactions proposed by Aimaretti et al. (2009) and Marunlanda et al. (2010) can be proven by measuring water contents in reactants and products of supercritical methanol process. Since both thermal degradation and etherification reaction of glycerol generate water as a product, an increase in water content will be observed if those reactions have occurred. To verify this, waste canola oil and dry methanol were prepared and their water contents were analyzed by Karl-Fischer titration. As reacting them under the same conditions with those of run 19, a single-phase product was obtained. The product was kept perfectly sealed and let settled for a day. Then, the water content of the product was measured in the same way. Details of the titration results are shown in Appendix B. The summarized results are shown in Table 4.7.

Comparing the reactants (waste oil and methanol) with the product, water contents had increased from being negligible to 0.23 g after the reaction. Other than glycerol reactions, another source of water was esterification of FFAs (4 wt%) in waste oil. Calculation of the amount of water from the esterification reaction is shown in Equation (D.1) in Appendix D. The calculation shows that only 0.017 g water can be produced in maximum if all the FFAs (assumed as oleic acid) were converted into methyl esters. Therefore, it was confirmed that the water appearance in the product was mostly contributed to the glycerol reactions.

Table 4.7. Water contents in the reactants and a product of supercritical reaction

Materials	Water content (g)
Waste oil	0.0002 ^a
Dry methanol	0.0013 ^b
Biodiesel solution	0.23 ^c

^a Standard deviation: 0.0001 g

^b Standard deviation: 0.0002 g

^c Standard deviation: 0.012 g

GC-MS of the biodiesel product sample was performed to identify components in the product. Details of the GC-MS analysis chromatogram are presented in Appendix E. Results in Figure 4.13 and Table 4.8 show that 3-methoxy-1,2-propanediol, 1,3-dimethoxy-2-propanol and diglycerol in the sample were identified. These components were the glycerol reaction products shown in Table 4.6

previously. Glycerol in the sample was also identified, indicating the by-product glycerol was not entirely consumed by the glycerol reactions. Methanol in the product sample was also identified by comparing its retention time (0.79 min) with the one of pure methanol. Quantification of the glycerol and methanol was not performed in this study.

Several chromatographic peaks observed during the beginning 5 min of GC-MS analysis verified a variety of chemical components present in the product samples. It indicated unknown side reactions occurred during the supercritical methanol reaction. As a candidate for the side reactions, Aimaretti et al. (2009) proposed that reaction of glycerol and supercritical methanol can result in formic acid ethyl ester by water loss and subsequent internal molecular rearrangement. A possibility of losing one oxygen atom and one water molecule by reaction between glycerol and supercritical methanol was also proposed by the authors. Those components can go through further chemical reactions particularly under high temperature conditions. It appears that chemical components generated from the unknown reactions are dissolved in the biodiesel solution of this study, also aiding un-reacted methanol dissolved in the single phase.

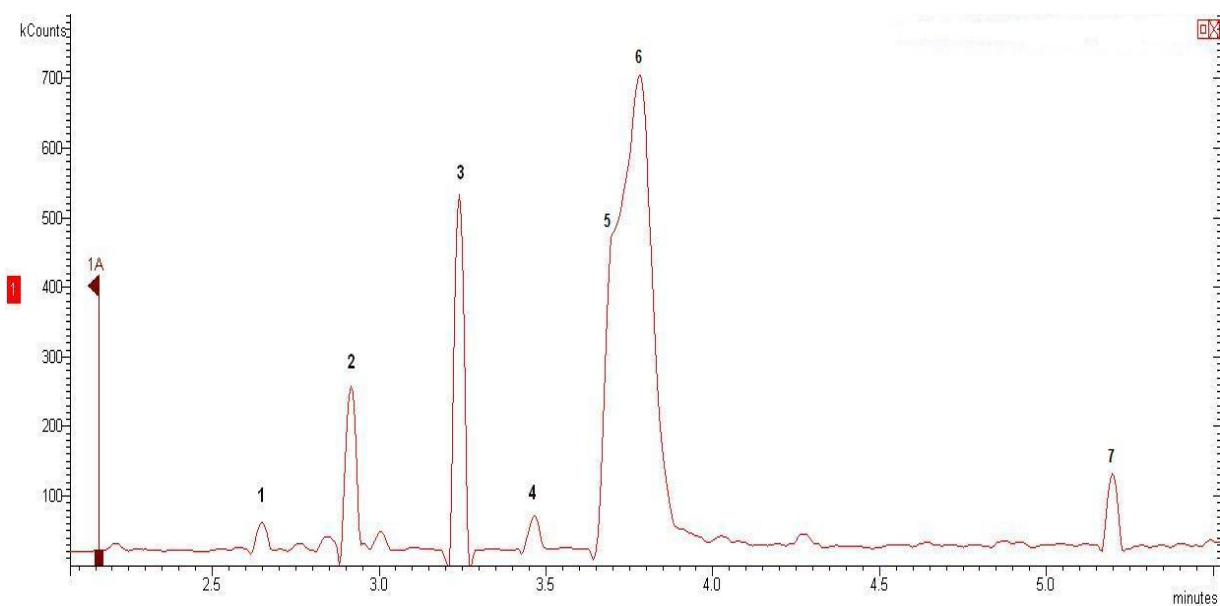


Figure 4.13. Gas chromatography–mass spectroscopy (GC-MS) chromatogram of the product from supercritical reaction at 270°C/10 MPa for 45 min

Table 4.8. Peak identification of the components in Figure 4.13

Peak	Name
1	1,3-dimethoxy-2-propanol
2	3-methoxy-1,2-propanediol
3	hexanoic methyl ester
4	1,2,4 butanetriol
5	glycerol
6	diglycerol
7	octanoic methyl ester

4.4. Conclusion

Biodiesel was produced from waste canola oil through supercritical methanol process. Results obtained through the study suggested the potential of supercritical method in biodiesel production as the fatty acid methyl ester yields close to 100% were obtained after 45 min of the reactions at 270°C/10 MPa. The advantage of the method was to perform the reactions under relatively moderate conditions, which could avoid thermal degradation of the methyl esters. Through the analysis of variance of the experimental results, reaction time and temperature were found to be significant to the yields and their interaction effects were also identified.

Single phase products were produced from 45 min reactions at 270°C/10 MPa. The increase in water contents after the supercritical reaction confirmed that glycerol reactions such as glycerol decomposition and glycerol methanolysis occurred during the supercritical reaction. Products from the glycerol reactions were identified by GC-MS analysis. Observation of several chromatographic peaks indicated other side chemical reactions also occurred during the 45 min supercritical reaction, resulting in the single phase product. The candidates for those side reactions were not investigated in this study, leaving behind more studies on analysis of the components in the reaction product.

Acknowledgments

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5. General discussion, conclusions and recommendations

5.1. General discussion

Chapter 2 presented simulated models of four different biodiesel production processes. While developing the simulated models, the normal boiling point of triolein was found to be inconsistent depending on its source. For example, Hysys reported its boiling point as 606.8°C, but another process simulator, Aspen, reported it as 846.85°C. As previously explained, the normal boiling point is of importance since it is used, along with the molecular structure, to predict the other remaining properties of the compound. The thermogravimetric measurement of the boiling points of triglycerides, proposed by Goodrum and Geller (2002), was therefore very useful because it provided results rapidly and with relatively simple preparation. In addition, the thermogravimetric measurements were proven by the same authors to be valid for long-chain triglycerides such as tristearin (C18:0) whose carbon chain length is the same as the triolein (C18:1) of our interest. Through thermogravimetric measurement, this thesis found the normal boiling point of triolein to be 412.8°C, slightly higher than the normal boiling point of tristearin (403.12°C) reported by Goodrum and Geller (2002). Technically, the normal boiling point of an unsaturated compound is lower than for a saturated one, which suggests that the experimental result for the normal boiling point of triolein was measured higher than was expected. The normal boiling point of triolein could be also measured with a differential scanning calorimeter (DSC). This measurement was omitted in this thesis. Providing the result from the DSC along with the result of thermogravimetric measurement would provide a more solid clue as to the normal boiling point of triolein.

The heat capacity of methanol also varies between Hysys and the literatures. It is 3.604 kJ/(kg·°C) at room temperature (25°C) in Hysys, which is 1.124 kJ/(kg·°C) higher than the value of 2.48 kJ/(kg·°C) from Filatov and Afanas'ev (1992). Applying the heat capacity from the literature to the Hysys simulation would result in lower utility costs in the models of the biodiesel production process. The savings on utility costs would be particularly large in the supercritical processes because the energy consumption involved with heaters and distillation columns for methanol are greater in this process than for alkali-catalyzed processes. For example, in the SC-WVO process, the different value for heat capacity results in an energy consumption difference of $(1.124 \text{ kJ/kg}^{-1} \cdot \text{°C}^{-1}) \cdot (8749.5 \text{ kg/hr}) \cdot (8000 \text{ hr/yr}) \cdot (350.0\text{°C} - 224.3\text{°C}) \cdot (1 \text{ GJ}/10^6 \text{ kJ}) = 9889.5 \text{ GJ/yr}$ in the heater (E-103) for pre-heating methanol to the reaction temperature. However, the order of profitability for the four biodiesel processes will not be altered by replacing the Hysys heat capacity with the value from the literature because the utility costs are only a small part of the total manufacturing costs.

The general layout of Alkali-FVO and Alkali-WVO processes in Chapter 2 referred to the process flow diagrams described in Zhang et al. (2003a) and West et al. (2008). The glycerol washing column in the Alkali-WVO process was represented as a liquid-liquid extraction column in Hysys, but the simulation results were not fully understood as a result of the abnormal behaviour of the simulated liquid-liquid extraction column. As stated in Chapter 2, the simulated extraction column had some problems in converging with the flow streams containing electrolytes, such as sulphuric acid. As the flow rate of the glycerol stream decreased from extremely large to a certain, smaller amount, the extraction column converged. However, it did not converge if the flow rate increased from a small amount to the same amount at which it had previously converged. Zhang et al. (2003a) also commented on the lack of knowledge concerning the behaviour of electrolytes in the liquid-liquid extraction column and used two component splitters after the column to practically separate glycerol, sulphuric acid and methanol from the rest of the components.

The amount of glycerol required for the washing process was not completely certain. Because of the abnormal behaviour of the extraction column, the amount of glycerol required could not be determined from the simulation results. Zhang et al. (2003a) used the glycerol washing process in their simulation model, but they did not report how they determined the amount of glycerol required for the washing process. Thus, in this thesis, the flow rate of glycerol in simulations was determined based on the assumption that it is approximately proportional to the capacity of the biodiesel plant being simulated. In Chapter 3, the cost of the glycerol washing process was found to be approximately 10% of the direct manufacturing cost, becoming the second largest cost factor after the cost of the waste oil feedstock. Therefore, more accurate data for the required amount of glycerol would provide more accurate washing process costs in Alkali-WVO process.

Chapter 3 focused on an economic assessment of the four biodiesel production process models developed in Chapter 2. The economic assessment conducted in this thesis is different than previous studies (Zhang et al., 2003b; Kasteren and Nisworo, 2007; West et al., 2008; You et al., 2008; Lim et al., 2009; Santana et al., 2010) because this thesis used, for the first time, a software package designed specifically for economic process evaluations (Icarus Process Evaluator—IPE).

Alteration of the general specifications discussed in Section 3.2 can be simply made in the early stages of the economic analysis. Alterations of this type were not included in this thesis. For example, the biodiesel plant could be re-located in different regions in the world, allowing an investigation into the effects of regional factors such as freight, taxes, wage rates, and workforce productivity on the profitability of a biodiesel plant.

An economic assessment of the four biodiesel production processes revealed the significance of the oil feedstock cost, which accounted for 63.9% to 84.0% of the total production cost. Except for

the Alkali-FVO process, which used expensive fresh vegetable oil, the other processes showed positive net present values due to the use of waste vegetable oil. The results indicate that the utilization of low-cost feedstocks would be a desirable feature for biodiesel production to become competitive.

Another important consideration was whether a reaction conversion rate lower than Bunyakiat et al. (2006) would result in negative profitability of the supercritical processes. The sensitivity analysis of the SC-WVO process showed that the net present value of the process would become negative if the biodiesel selling price decreased by around 15%. If the decrease in revenue from biodiesel sale resulted from a decrease in the net yield of biodiesel in the process (i.e., a decrease in the reaction conversion) instead of from a reduction in the biodiesel selling price, negative NPV would result from approximately a 15% reduction in conversion. Moreover, if we consider the increase in the cost of the purification processes as a result of the lowered conversion, a reduction in the reaction conversion of less than 15% would make the profitability of the SC-WVO process negative. Unfortunately, research on the continuous production of biodiesel in a supercritical methanol process is scarce. Consequently, the validity of the economic assessment of the supercritical processes in this study would be more clearly understood once more research is conducted in the future.

The biodiesel selling price used for the economic assessment was \$0.99/kg, equivalent to \$0.861/litre (given the biodiesel density of 870 kg/m³). The price was researched on April 16th, 2010. During the same period, the average petroleum diesel price in Canada was \$0.746/litre. In the urban cities such as Vancouver and Toronto, the prices were \$0.761/litre and \$0.772/litre, respectively, whereas the price was even higher in remote areas, such as Whitehorse (\$0.882/litre). In the same manner, geographically related factors, such as freight costs, will affect the biodiesel selling price.

Chapter 4 of this thesis describes the experimental work on biodiesel synthesis from waste canola oil via a supercritical methanol process. This experiment investigated the effects of reaction time, reaction temperature, and weight ratios of methanol to oil on the yield of fatty acid methyl esters (FAMEs), and a yield prediction formula for FAMEs was developed. Originally, the supercritical reactions were planned to be carried out at a higher pressure (17 MPa) with a broader temperature range (240–315°C) than the investigated conditions reported in Chapter 4. Unfortunately, the reaction conditions were limited by the allowable operating conditions of the autoclave o-ring, which would be destroyed at high temperature and pressure (>11 MPa). Supercritical reactions lasting 45 min at 270°C/10 MPa resulted in 96.4% and 101.6% of FAME mass yields using methanol:oil ratios of 1:1 and 2:1, respectively. The product from these reaction conditions appeared to be in a

single yellowish liquid phase, which did not separate into two phases after being kept at 19°C for 2 weeks.

The single phase product was thought to be a result of the decomposition of the glycerol from the transesterification reaction, as well as its partial reaction with methanol. Aimaretti et al. (2009) reported that reactions conducted at the conditions of 280°C and 6.25–7.55 MPa produced practically no glycerol by-product. The authors reported that glycerol was transformed into products of smaller molecular size and water, and the water reacted with the triglycerides of the reacting mixture to form free fatty acids, which are then converted into methyl esters at longer reaction times (60–120 min). However, as the products in this thesis were kept in a cold room (4°C) for two days, they separated into two phases, like the products from runs 1–17, indicating that by-product glycerol was present in the single phase. Glycerol presence was also confirmed by gas chromatography-mass spectrometry (GC-MS), which detected a peak associated with glycerol. The presence of a single, liquid phase raised questions concerning the adequacy of the residence time (1 hr) in the 3-phase separator used in previous simulation models (SC-WVO and SC-LowE). In light of the fact that the reaction time was relatively short (7–14 min) and since no single-phase product was reported by Bunyakiat et al. (2006), the residence time used in the simulation models was considered to be adequate.

5.2. Conclusions

This thesis begins with the measurement of the normal boiling point of triolein through the thermogravimetric method. The boiling point was used for later process simulations and an economic analysis of a supercritical biodiesel production process. Based on the thermogravimetric result, the normal boiling point of triolein was found to be 412.8°C, which was lower than the value reported by Hysys, Aspen and VMGsim programs used by previous studies of the process simulation of biodiesel production. Significant improvements in the specific heat capacity, mass density and viscosity were observed as the normal boiling point found by the thermogravimetric analysis was applied to triolein in Hysys.

Using the triolein with the improved properties, simulation models of four different kinds of biodiesel production processes were developed: a homogeneous alkali-catalyzed process using fresh vegetable oil as the feedstock (Alkali-FVO); a homogeneous catalyzed process using waste vegetable oil as the feedstock (Alkali-WVO); a supercritical process using waste vegetable oil as the feedstock (SC-WVO); and a supercritical process with slight modifications to the SC-WVO process in an attempt to lower energy consumption. The process flowsheets and energy and material balances were presented in Chapter 2. A comparison of the energy consumption of the four processes showed that the supercritical process (SC-WVO) consumed 4.6 times more energy than the Alkali-FVO process as a result of the large amount of methanol evaporated in the distillation column and the reaction conditions of high temperature and pressure.

The economic assessment, conducted in Chapter 3, showed that the advantageous use of the low-cost feedstock in the supercritical process compensated for the high utility cost when compared to the conventional alkali-catalyzed process. The costs for the feedstock oil comprised 84.0%, 63.9%, 71.3% and 75.4% of the total production costs of biodiesel using Alkali-FVO, Alkali-WVO, SC-WVO and SC-LowE processes, respectively, considerably affecting the processes' economics.

Based on the economic analysis, SC-LowE is the most promising process among those investigated. The discounted payback period, net present value, discounted cash flow rate of return and break-even price of biodiesel were 4.2 years, \$21.086 million, 49.6% and \$0.726/kg, respectively. The net present values of the SC-WVO, Alkali-WVO and Alkali-FVO processes were \$16.451 million, \$1.901 million and -\$21.697 million, respectively. These results indicate the potential of the supercritical processes compared to the conventional alkali-catalyzed process.

Sensitivity analysis of the six parameters (biodiesel selling price, feedstock oil price, by-product glycerol price, interest rate, life of the biodiesel plants and capacity) was conducted in order to systematically examine the effect of positive and negative changes in project parameters on

profitability. From these results, biodiesel selling price, feedstock oil price, by-product glycerol price and interest rate were found to be the significant factors contributing to the net present values of the four processes. These were further studied through the Response Surface Method (RSM) coupled with central composite design (CCD) using JMP software. Through the analysis of variance (ANOVA) of the results, prediction formulas of the NPV for each of the four processes were developed.

The significance of the usage of low-cost feedstock and the potential of a supercritical process resulting from the economic assessment brought about the idea to conduct experiments on synthesizing biodiesel from waste canola oil using supercritical methanol. Supercritical reactions with the waste canola oil and methanol gave the FAME yields close to 100% at the reaction conditions of 270°C/10 MPa after 45 min of reaction time. Due to the relatively moderate reaction conditions, the reaction time necessary to achieve a high FAME yield was longer than the time reported in literatures (Saka and Kusdiana, 2001; Madras et al., 2004; Demirbas, 2009; Valle et al., 2010). Experiments showed that the conversion of the waste oil increased significantly with increasing reaction time and reaction temperature. The effect of methanol to oil weight ratio on the FAME yield was relatively less significant. The interaction between reaction time and reaction temperature was identified by the ANOVA of the experimental results.

5.3. Recommendations

For the future research, several recommendations are proposed based on the work conducted for this thesis.

- Experiments should be performed to verify the quantity of glycerol used for the washing process in the liquid-liquid extraction column in Alkali-WVO process.
- For the comparison with the properties of triolein estimated by Hysys, it is recommended to apply the normal boiling point of triolein determined by the thermogravimetric analysis to other process simulation programs.
- Since the reaction conversion (96%) in the supercritical process simulation models referred to the experimental data reported by Bunyakiat et al. (2006) where coconut and palm oils were used instead of waste vegetable oil as the feedstocks, it is recommended to perform experiments to verify the reaction conditions and conversions in SC-WVO and SC-LowE processes using waste vegetable oil.

- It is also recommended to conduct a kinetic study of thermal decomposition of methyl esters, which can be combined with transesterification and esterification reaction kinetics. The data would enhance the simulation of a plug flow reactor in SC-WVO and SC-LowE processes, and could be usefully applied as scaling up the processes.
- The simulated waste vegetable oil in Alkali-WVO, SC-WVO and SC-LowE processes contains 5 wt% free fatty acid, but in practice waste vegetable oil contains diverse ranges of free fatty acids and water as reported by Canakci (2007). Therefore, the feedstock oil to be used in a simulation model could be expanded to include those with FFAs contents greater than 5 wt% and water content.
- The reason why the products from run 18 and run 19 in Chapter 4 formed a single phase and took relatively long time to be separated into two phases was not fully investigated in this thesis. Experiments need to be performed to find out in which reaction conditions single phase products are generated by varying the key parameters such as reaction time, reaction temperature, reaction pressure and weight ratio of methanol to oil. Verifying components in the products could subsequently be done using GC-MS and compared to those in other products having two phases from the beginning.
- Around 99% of by-product glycerol was removed from the biodiesel in the simulated 3-phase separators in the supercritical process models (SC-WVO and SC-LowE). Performing experiments with a mixture of the similar composition with that of the feed stream to the 3-phase separator are recommended to verify the simulation results.

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Appendix A: Economic assessments for the biodiesel plant capacities of 8,000 and 160,000 tonnes/yr biodiesel

Table A.1. Total capital cost for the four processes (unit: \$ millions)

Process		Alkali-FVO		Alkali-WVO		SC-WVO		SC-LowE	
Capacity (tonnes/year)		8,000	160,000	8,000	160,000	8,000	160,000	8,000	160,000
Direct items									
Reactor	Esterification	-	-	0.091	0.657	-	-	-	-
	Transesterification	0.109	0.642	0.13	0.624	0.113	1.934	0.335	1.695
	Neutralization	0.05	0.106	0.026	0.105	-	-	-	-
Column	Methanol recovery	0.067	0.179	0.162 ^a /0.129	0.552 ^a /0.174	0.104	0.619	0.068	0.256
	Biodiesel purification	0.11	0.597	0.097	0.844	0.153	0.88	0.124	0.578
	Glycerol purification	0.048	0.053	0.021	0.079	-	-	-	-
	Water washing	0.021	0.050	0.024	0.046	-	-	-	-
	Glycerol washing	-	-	0.09	0.156	-	-	-	-
Other	Pumps	0.017	0.019	0.031	0.034	0.023	0.141	0.023	0.141
	Heat exchangers	0.005	0.03	0.037	0.125	0.293	1.76	0.321	1.744
	L-L separator	0.009	0.031	0.006	0.031	0.015	0.041	0.015	0.041
	Flash evaporator	-	-	-	-	-	-	0.02	0.097
Total purchased equipment		0.437	1.71	0.844	3.43	0.701	5.37	0.908	4.55
Equipment setting		0.014	0.027	0.025	0.042	0.014	0.046	0.016	0.04
Piping		0.542	0.82	0.929	1.59	0.905	4.13	0.885	3.159
Civil		0.106	0.19	0.155	0.298	0.099	0.235	0.105	0.229
Steel		0.041	0.089	0.066	0.146	0.021	0.056	0.03	0.077
Instrumentation		1.07	1.28	1.55	1.8	0.866	1.202	0.881	1.005
Electrical		0.494	0.664	0.515	0.706	0.471	0.681	0.482	0.682
Insulation		0.153	0.273	0.226	0.429	0.173	0.452	0.169	0.356
Paint		0.019	0.031	0.033	0.045	0.015	0.036	0.015	0.028
Total direct cost		2.879	5.086	4.344	8.487	3.286	12.21	3.481	10.13
Others		3.74	4.48	5.124	6.4	3.187	5.226	3.26	4.73
G and A Overheads		0.12	0.202	0.183	0.331	0.134	0.447	0.14	0.372
Contract Fee		0.368	0.46	0.483	0.644	0.335	0.64	0.343	0.567
Contingencies		1.28	1.84	1.824	2.855	2.256	6.02	2.348	5.133
Fixed capital cost		8.39	12.1	11.96	18.72	9.2	24.54	9.57	20.93
Working capital		1.26	1.81	1.8	2.81	1.38	3.68	1.44	3.14
Total capital investment		9.65	13.9	13.8	21.53	10.58	28.22	11.01	24.07

^a Indicates a pre-treatment process

**Table A.2. Total manufacturing cost, glycerol credit and revenues from biodiesel sales of the four processes
(unit: \$ millions)**

Process	Alkali-FVO		Alkali-WVO		SC-WVO		SC-LowE	
	8,000	160,000	8,000	160,000	8,000	160,000	8,000	160,000
Direct manufacturing cost								
Oil feed	7.69	153.4	4.58	88.86	4.448	87.83	4.448	87.83
Methanol	0.246	4.673	0.282	5.247	0.236	4.446	0.236	4.658
Catalysts	0.065	0.542	0.043	3.471	-	-	-	-
Water washing	0.000	0.000	0.000	0.000	-	-	-	-
Glycerol washing	-	-	0.757	13.06	-	-	-	-
Steam (4201 kPa)	0.005	1.388	0.004	1.297	0.449	6.087	0.106	2.115
Steam (1135 kPa)	-	-	-	-	-	-	-	-
Steam (690 kPa)	0.024	0.059	0.112	1.621	-	-	-	-
Circulating heating oil	0.082	1.599	0.084	2.354	0.073	2.661	0.227	2.461
Cooling water	0.004	0.083	0.007	0.111	0.015	0.276	0.012	0.153
Chilled water	0.046	1.45	0.067	6.984	-	-	-	-
Electricity	0.044	0.118	0.054	0.223	0.048	0.348	0.047	0.318
Operating labour	0.6	0.8	0.8	0.8	0.6	0.6	0.6	0.6
Supervision	0.28	0.28	0.28	0.28	0.28	0.28	0.28	0.28
Maintenance	0.503	0.724	0.717	1.123	0.552	1.473	0.574	1.256
Operating supplies	0.05	0.072	0.072	0.112	0.055	0.147	0.057	0.126
Laboratory charges	0.143	0.176	0.176	0.176	0.143	0.143	0.143	0.143
subtotal	9.785	165.4	1.592	125.7	6.898	104.3	6.731	99.94
Indirect manufacturing cost								
Overhead	0.692	0.902	0.899	1.101	0.716	1.176	0.727	1.068
Depreciation	0.72	1.016	1.195	1.822	0.791	2.429	0.823	1.746
Property Taxes	0.194	0.279	0.277	0.433	0.213	0.568	0.222	0.485
Insurance	0.084	0.121	0.12	0.187	0.092	0.245	0.096	0.209
Subtotal	1.69	2.318	2.491	3.543	1.812	4.418	1.867	3.508
General expenses	1.721	25.16	1.579	19.39	1.306	16.31	1.29	15.52
Total production cost	13.2	192.9	12.105	148.7	10.02	125	9.888	119
Glycerol credit	1.094	21.7	1.061	20.5	0.723	14.3	0.719	14.3
Total manufacturing cost	12.1	171.2	11.0	128.2	9.3	110.7	9.2	104.7
Revenue from biodiesel	7.92	158.5	7.976	158.1	7.922	158.3	7.922	158.3

Table A.3. Experimental tables of central composite design and the responses (NPV) of Alkali-FVO process

Pattern	Parameters				NPV (\$ millions)
	Biodiesel price (\$/kg)	FVO price (\$/kg)	Glycerol price (\$/kg)	Interest rate (%)	
0000	0.99	0.916	1.323	20	-26.697
00A0	0.99	0.916	2.3814	20	-12.393
+++−	1.386	1.2824	1.8522	10	-26.197
0000	0.99	0.916	1.323	20	-26.697
0a00	0.99	0.1832	1.323	20	66.416
−+−−	0.594	1.2824	0.7938	10	-260.209
−++−	0.594	1.2824	1.8522	10	-231.314
00a0	0.99	0.916	0.2646	20	-44.123
0000	0.99	0.916	1.323	20	-26.697
A000	1.782	0.916	1.323	20	58.72
++++	1.386	1.2824	0.7938	30	-34.454
+−−+	1.386	0.5496	0.7938	30	34.368
a000	0.198	0.916	1.323	20	-154.669
+−++	1.386	0.5496	1.8522	30	41.479
0000	0.99	0.916	1.323	20	-26.697
−−−+	0.594	0.5496	0.7938	30	-23.873
0A00	0.99	1.6488	1.323	20	-167.367
+−−−	1.386	0.5496	0.7938	10	101.267
0000	0.99	0.916	1.323	20	-26.697
++−−	1.386	1.2824	0.7938	10	-51.844
000a	0.99	0.916	1.323	0	-33.86
0000	0.99	0.916	1.323	20	-26.697
−+++	0.594	1.2824	0.7938	30	-119.929
000A	0.99	0.916	1.323	40	-20.662
+−+−	1.386	0.5496	1.8522	10	118.604
0000	0.99	0.916	1.323	20	-26.697
−++++	0.594	1.2824	1.8522	30	-108.078
−−+−	0.594	0.5496	1.8522	10	-10.817
0000	0.99	0.916	1.323	20	-26.697
0000	0.99	0.916	1.323	20	-26.697
++++	1.386	1.2824	1.8522	30	-23.368
0000	0.99	0.916	1.323	20	-26.697
−−−−	0.594	0.5496	0.7938	10	-36.364
−−++	0.594	0.5496	1.8522	30	-12.893
0000	0.99	0.916	1.323	20	-26.697
0000	0.99	0.916	1.323	20	-26.697

Appendix B: Water content measurement by Karl Fischer volumetric titration

The water content of the waste canola oil and product from run 19 in Chapter 4 were determined using an automatic Karl Fischer titration apparatus (794 Basic Titrino from Metrohm). The procedure is as follows:

- Approximately 40 ml extra dry methanol and a magnetic stir pill are placed in the titration cell.
- The cell is placed on the titration stand and the magnetic stirrer is switched on.
- Load a titration method named “KFSample” and press the “Start” button.
- Titrate the methanol to dryness until a constant drift of approximately 10–20 $\mu\text{l}/\text{min}$ is achieved. This step is called conditioning.
- Place a syringe filled with the oil sample on the balance and tare.
- Press “Start” on the titrator. The display will change, prompting to add the sample and enter the sample size.
- Remove the syringe from the balance and inject approximately 0.5 g of sample into the cell. The sample should fall directly into the methanol and not touch the walls of the cell, the electrode or anything else.
- After the sample has been dispensed, place the syringe back on the balance to obtain an accurate differential sample weight. Enter the sample weight using a keypad and press “Enter”
- This procedure is repeated at least three times to ensure accuracy.

The water content in percentage is calculated via the following formula

$$\text{Water content (\%)} = \frac{\text{EP1} \times \text{C39} \times \text{C01}}{\text{C00}} \quad (\text{B. 1})$$

where EP = end point (ml), C39 = titer value (mg of water titrated by ml of KF reagent), C01 = 0.1 (conversion factor to percentage) and C00 = sample weight (g).

Each of the reported water contents in Chapter 4 is an average of three replicate measurements. The printed reports of all the measurement are included on the next pages.

1) Water contents in the waste canola oil in Tables 4.4 and 4.7

```

NuNu 794.0010
date 2010-07-16 time 15:31 6
U(init) 272 mV SET IpolKFSample
smpl size 0.8718 g
EP1 0.008 ml 18 mV
RS1 0.004 %
-----

```

```

*fm
794 Titrimo NuNu 794.0010
date 2010-07-16 time 15:31 6
SET Ipol KFSample
>calculations
RS1=EP1*C39*C01/C00*3;%
C00= 0.8718
C01= 0.1
C39= 4.1498
-----

```

```

*sf
794 Titrimo NuNu 794.0010
date 2010-07-16 time 15:31 6
method id 1/C21 id 2/C22 id 3/C23 mean t/s n
-----

```

```

794 Titrimo NuNu 794.0010
date 2010-07-16 time 19:28 7
U(init) 178 mV SET IpolKFSample
smpl size 0.6013 g
EP1 0.000 ml 250 mV
RS1 0.000 %
-----

```

```

*fm
794 Titrimo NuNu 794.0010
date 2010-07-16 time 19:28 7
SET Ipol KFSample
>calculations
RS1=EP1*C39*C01/C00*3;%
C00= 0.6013
C01= 0.1
C39= 4.1498
-----

```

```

*sf
794 Titrimo NuNu 794.0010
date 2010-07-16 time 19:28 7
method id 1/C21 id 2/C22 id 3/C23 mean t/s n
-----

```

```

794 Titrimo NuNu 794.0010
date 2010-07-16 time 21:25 8
U(init) 272 mV SET IpolKFSample
smpl size 0.5242 g
EP1 0.006 ml 16 mV
RS1 0.005 %
-----

```

```

*fm
794 Titrimo NuNu 794.0010
date 2010-07-16 time 21:25 8
SET Ipol KFSample
>calculations
RS1=EP1*C39*C01/C00*3;%
C00= 0.5242
C01= 0.1
C39= 4.1498
-----

```

```

*sf
794 Titrimo NuNu 794.0010
date 2010-07-16 time 21:25 8
method id 1/C21 id 2/C22 id 3/C23 mean t/s n
-----

```

2) Water contents in the methanol in Table 4.7

```

794 Titrimo      NuNu  794.0010
date 2010-07-19  time 11:36      2
U(init)         294 mV SET IpolKFSample
smpl size      0.4615 g
EP1            0.010 ml          185 mV
RS1            0.009 %
-----

```

```

*fm
794 Titrimo      NuNu  794.0010
date 2010-07-19  time 11:36      2
SET Ipol        KFSample
>calculations
RS1=EP1*C39*CO1/CO0:3:%
CO0=            0.4615
CO1=            0.1
C39=            4.1498
-----

```

```

*sf
794 Titrimo      NuNu  794.0010
date 2010-07-19  time 11:36      2
method id 1/C21 id 2/C22 id 3/C23      mean      t/s      n
-----

```

```

794 Titrimo      NuNu  794.0010
date 2010-07-19  time 14:40      3
U(init)         278 mV SET IpolKFSample
smpl size      0.2660 g
EP1            0.008 ml          22 mV
RS1            0.012 %
-----

```

```

*fm
794 Titrimo      NuNu  794.0010
date 2010-07-19  time 14:40      3
SET Ipol        KFSample
>calculations
RS1=EP1*C39*CO1/CO0:3:%
CO0=            0.2660
CO1=            0.1
C39=            4.1498
-----

```

```

*sf
794 Titrimo      NuNu  794.0010
date 2010-07-19  time 14:40      3
method id 1/C21 id 2/C22 id 3/C23      mean      t/s      n
-----

```

```

794 Titrimo      NuNu  794.0010
date 2010-07-19  time 16:54      4
U(init)         287 mV SET IpolKFSample
smpl size      0.6521 g
EP1            0.012 ml          72 mV
RS1            0.008 %
manual stop
-----

```

```

*fm
794 Titrimo      NuNu  794.0010
date 2010-07-19  time 16:54      4
SET Ipol        KFSample
>calculations
RS1=EP1*C39*CO1/CO0:3:%
CO0=            0.6521
CO1=            0.1
C39=            4.1498
-----

```

```

*sf
794 Titrimo      NuNu  794.0010
date 2010-07-19  time 16:54      4
method id 1/C21 id 2/C22 id 3/C23      mean      t/s      n
-----

```

3) Water contents in the biodiesel solution in Table 4.7

```

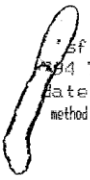
*fr
794 Titrimo      NuNu  794.0010
date 2010-07-20  time 11:45      2
U(init)         288 mV SET IpolKFSample
smpl size      0.6500 g
EP1            1.880 ml           239 mV
RS1            1.200 %
-----

```

```

*fm
794 Titrimo      NuNu  794.0010
date 2010-07-20  time 11:45      2
SET Ipol        KFSample
>calculations
RS1=EP1*C39*CO1/CO0;3;%
CO0=            0.6500
CO1=            0.1
C39=            4.1498
-----

```



```

*sf
794 Titrimo      NuNu  794.0010
date 2010-07-20  time 11:45      2
method id 1/C21 id 2/C22 id 3/C23  mean  t/s  n
-----

```

```

*fr
794 Titrimo      NuNu  794.0010
date 2010-07-20  time 14:39      3
U(init)         290 mV SET IpolKFSample
smpl size      0.7224 g
EP1            1.862 ml           248 mV
RS1            1.070 %
manual stop
-----

```

```

*fm
794 Titrimo      NuNu  794.0010
date 2010-07-20  time 14:39      3
SET Ipol        KFSample
>calculations
RS1=EP1*C39*CO1/CO0;3;%
CO0=            0.7224
CO1=            0.1
C39=            4.1498
-----

```

```

*sf
794 Titrimo      NuNu  794.0010
date 2010-07-20  time 14:39      3
method id 1/C21 id 2/C22 id 3/C23  mean  t/s  n
-----

```

```

*fr
794 Titrimo      NuNu  794.0010
date 2010-07-20  time 18:28      4
U(init)         290 mV SET IpolKFSample
smpl size      0.6273 g
EP1            1.806 ml           244 mV
RS1            1.195 %
-----

```

```

*fm
794 Titrimo      NuNu  794.0010
date 2010-07-20  time 18:28      4
SET Ipol        KFSample
>calculations
RS1=EP1*C39*CO1/CO0;3;%
CO0=            0.6273
CO1=            0.1
C39=            4.1498
-----

```

```

*sf
794 Titrimo      NuNu  794.0010
date 2010-07-20  time 18:28      4
method id 1/C21 id 2/C22 id 3/C23  mean  t/s  n
-----

```

Appendix C: Sample calculations of FAME yields

The yields of FAME were calculated by Equation (C.1) in this thesis. Accordingly, the yield of 101.6% in Table 4.4 was calculated as shown in Equation (C.2). The concentrations of methyl esters were measured by gas chromatography analysis.

$$\text{Yield (\%)} = \frac{\text{Total weight of methyl esters produced}}{\text{Total weight of oil used for reaction}} \times 100 (\%) \quad (\text{C.1})$$

$$\begin{aligned} \text{Yield} &= \frac{\text{Concentrations of methyl esters } \left(\frac{\text{g}}{\text{ml}}\right) \times \text{Product volume (ml)}}{\text{Total weight of oil used for reaction (g)}} \times 100 (\%) \\ &= \frac{0.267 \times 32.0}{8.486} \times 100 (\%) = 101.6\% \end{aligned} \quad (\text{C.2})$$

The yield over 100% in the experiment 19 appears to be attributed to the FFAs contained in the feedstock and the nature of the transesterification and esterification reactions. Simple calculations of the yields from transesterification and esterification reactions are shown in Equation (C.3) and (C.4), respectively. Methyl oleate was used as an example because it is the dominant component in the biodiesel produced in this study. Assuming 100% conversion of transesterification, 3 mols of methyl ester are produced from 1 mol of triolein. Because the mass of 3 mols of methyl oleate are slightly heavier than the mass of 1 mol of triolein, the yield calculated based on the definition in Equation (C.1) becomes 100.46%. In the same manner, the yield from esterification of oleic acid is calculated to be 104.97% as complete conversion of oleic acid was assumed. Thus, the theoretical FAME yield for the waste oil containing 96 wt% triolein and 4 wt% oleic acid is 100.64%, assuming 100% conversion of both transesterification and esterification reactions. The 101.6% yield in run 19 is 0.95% higher than the theoretical yield, indicating almost complete conversion of fatty compounds in the waste canola oil was achieved.

$$\begin{aligned} \text{triolein 1 mol (885.4 g)} &\rightarrow \text{methyl oleate 3 mols (889.4 g),} \\ \therefore \text{Yield} &= \left(\frac{889.4}{885.4}\right) \times 100(\%) = 100.46\% \end{aligned} \quad (\text{C.3})$$

$$\begin{aligned} \text{oleic acid 1 mol (282.46 g)} &\rightarrow \text{methyl oleate 1 mol (296.49 g)} \\ \therefore \text{Yield} &= \left(\frac{296.49}{282.46}\right) \times 100(\%) = 104.97\% \end{aligned} \quad (\text{C.4})$$

$$\begin{aligned} \text{Waste oil (triolein 96 wt\% + oleic acid 4 wt\%)} &100.00 \text{ g} \rightarrow \text{methyl oleate } 100.64 \text{ g} \\ \therefore \text{Yield} &= \left(\frac{100.64}{100.00}\right) \times 100(\%) = 100.64\% \end{aligned} \quad (\text{C.5})$$

Appendix D: Calculations of the amount of water produced from esterification reaction of FFA

Esterification: $\text{FFA} + \text{CH}_3\text{OH} \rightarrow \text{Methyl ester} + \text{H}_2\text{O}$

\therefore Water produced (g)

$$\begin{aligned} &= \text{FFA reacted (mol)} \times \text{water molar mass} \left(\frac{\text{g}}{\text{mol}} \right) \\ &= \frac{\text{weight fraction of FFA} \cdot \text{weight of waste oil used (g)}}{\text{oleic acid (FFA) molar mass (g/mol)}} \times \text{water molar mass} \left(\frac{\text{g}}{\text{mol}} \right) \\ &= \frac{0.04 \cdot 6.538}{282.46} \times 18 = 0.017 \end{aligned} \quad (\text{D. 1})$$

Appendix E: Gas chromatography-mass spectrometry (GC-MS)

A Varian Factor Four™ capillary VF-5ms column of 30 m × 0.25 mm × 0.25 μm film thickness was used with helium as carrier gas. The temperature program started at 60°C (held for 2 min) and ramped to 270°C at 20°C/min. The holding time at the final temperature (270°C) was 2 min. The temperature of the injector was set at 270°C. A 1 μl of the biodiesel sample in Table 4.7 was injected into the column. Details of peak identifications of the components in Table 4.8 are shown on the next pages. The chromatographic peaks of methyl esters with a carbon chain longer than C14 appeared in the retention times >10.0 min.

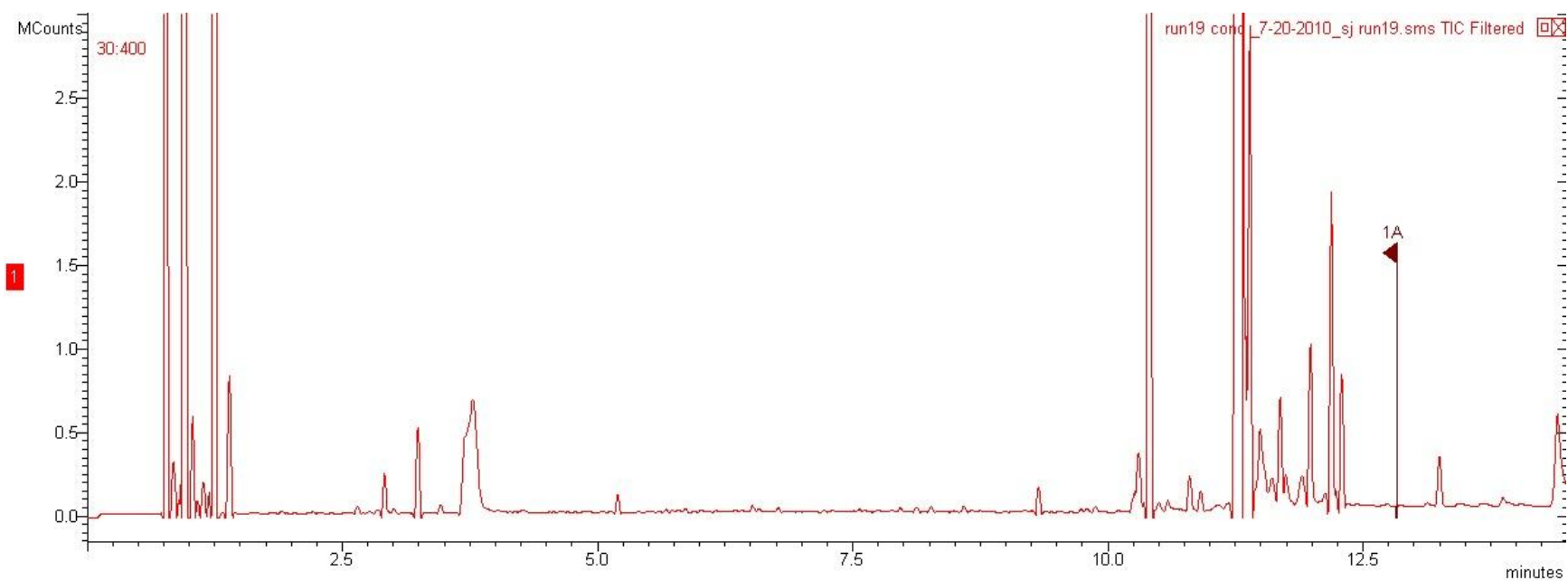


Figure E.1. GC-MS chromatogram of the single phase biodiesel product in Table 4.7

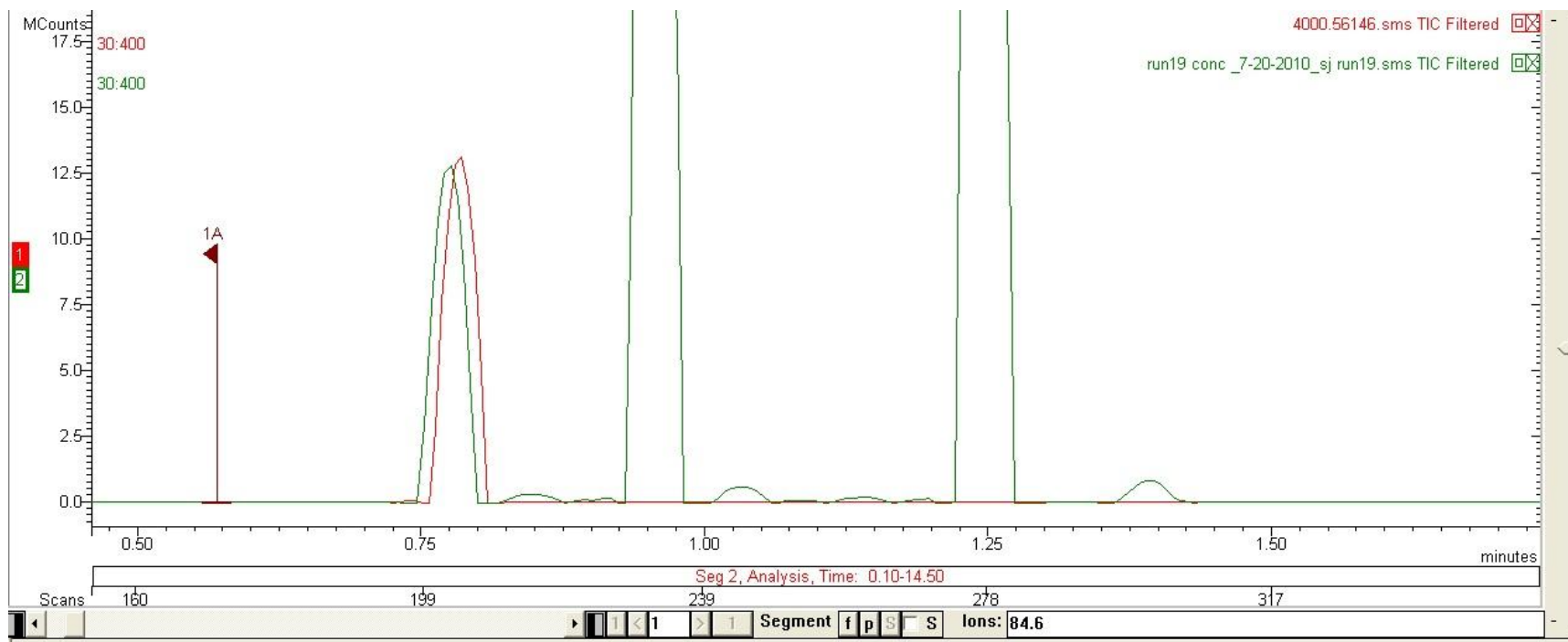


Figure E.2. Identification of methanol in the biodiesel sample (green) by comparing retention time with pure methanol (red)

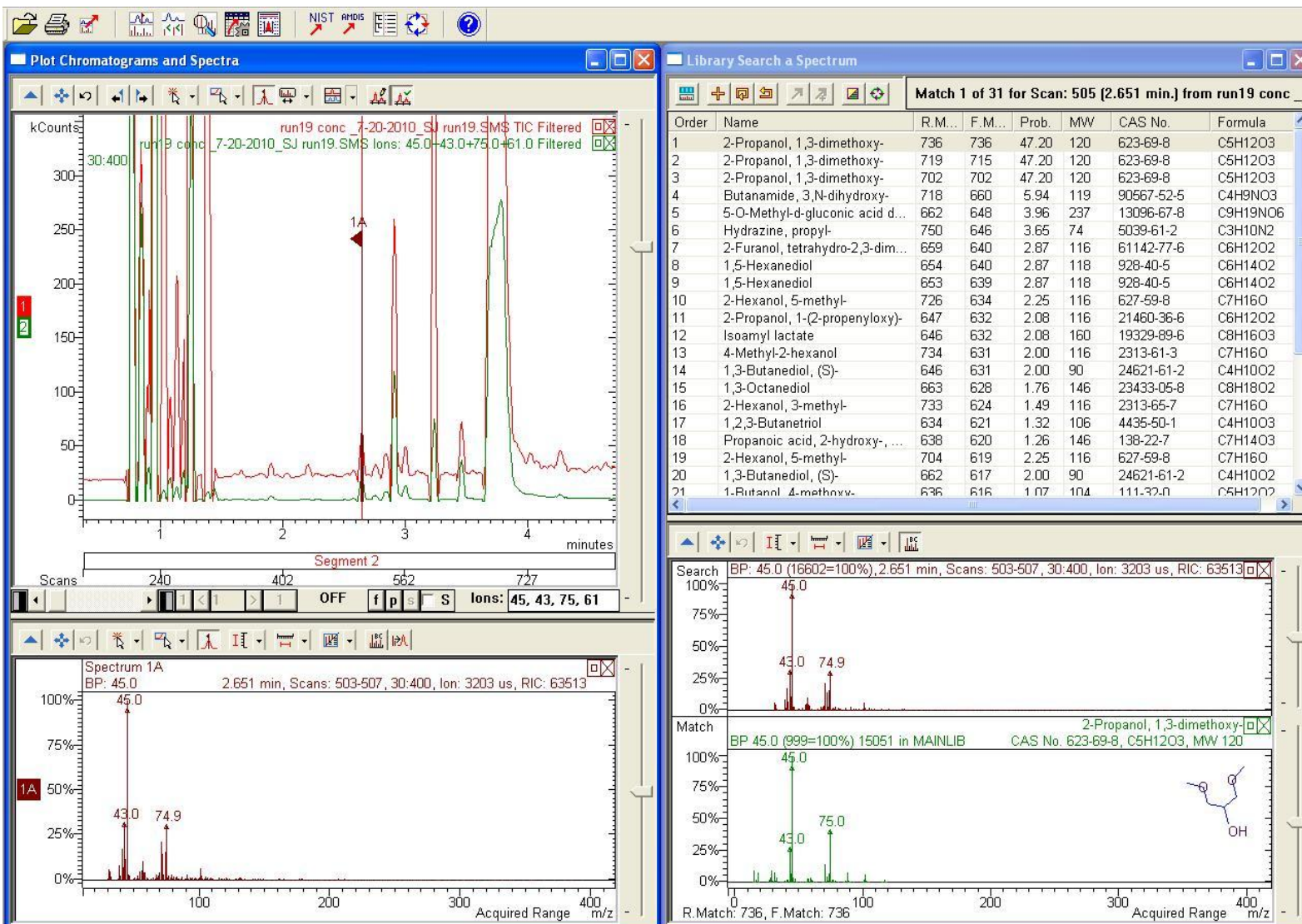


Figure E.3. Identification of 1,3-dimethoxy-2-propanol in the biodiesel sample by GC-MS

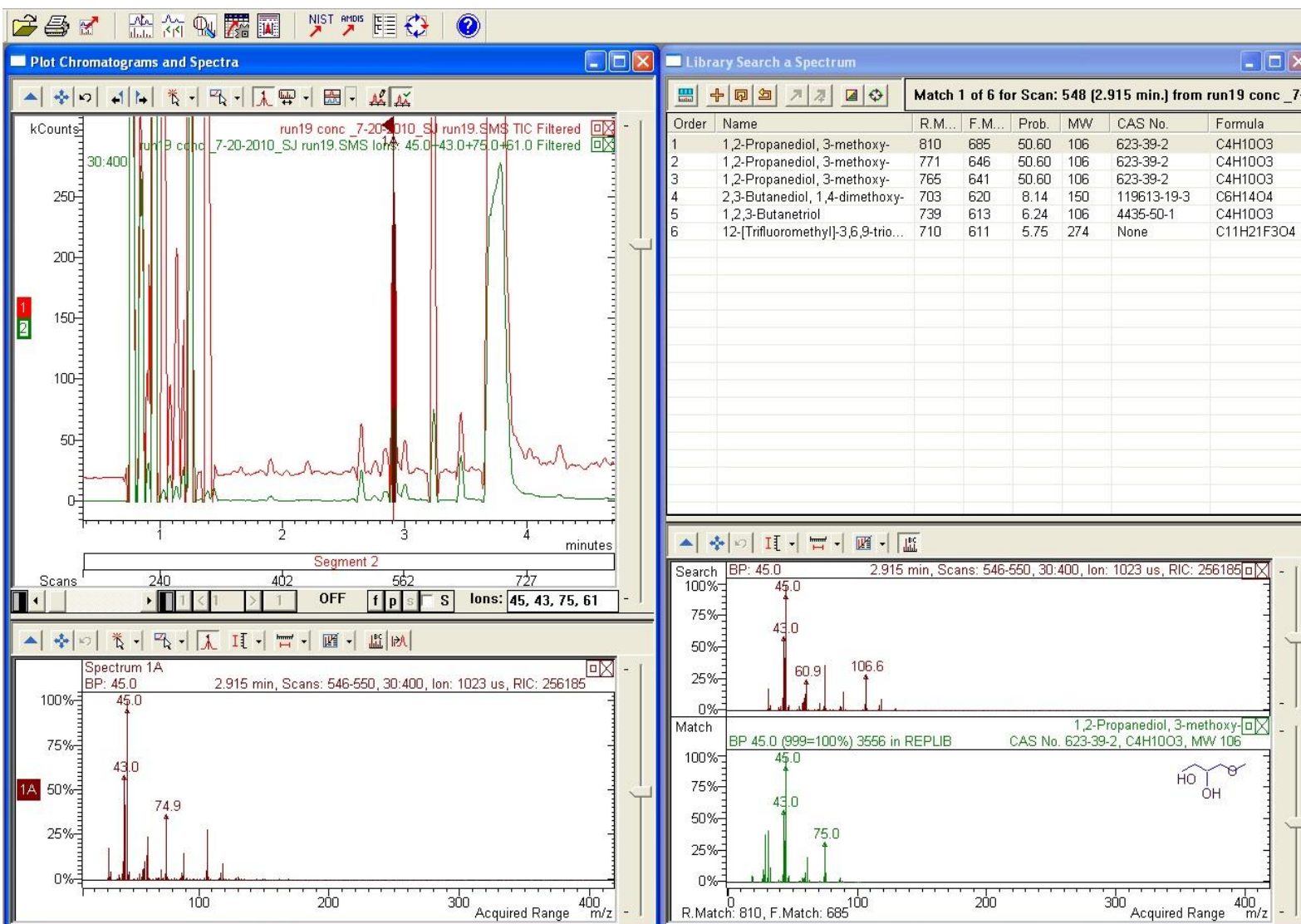


Figure E.4. Identification of 3-dimethoxy-1,2-propanediol in the biodiesel sample by GC-MS

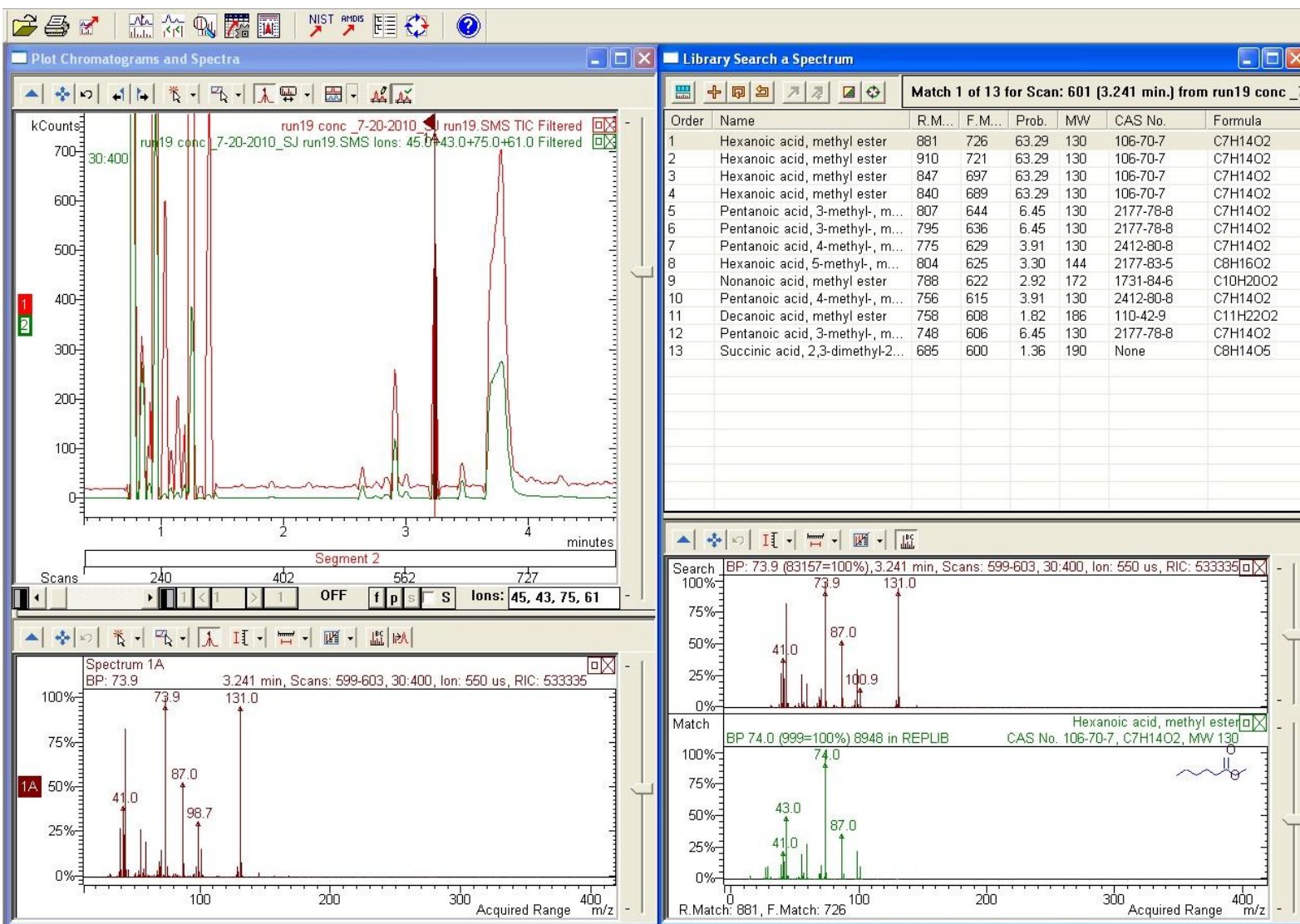


Figure E.5. Identification of hexanoic methyl ester in the biodiesel sample by GC-MS

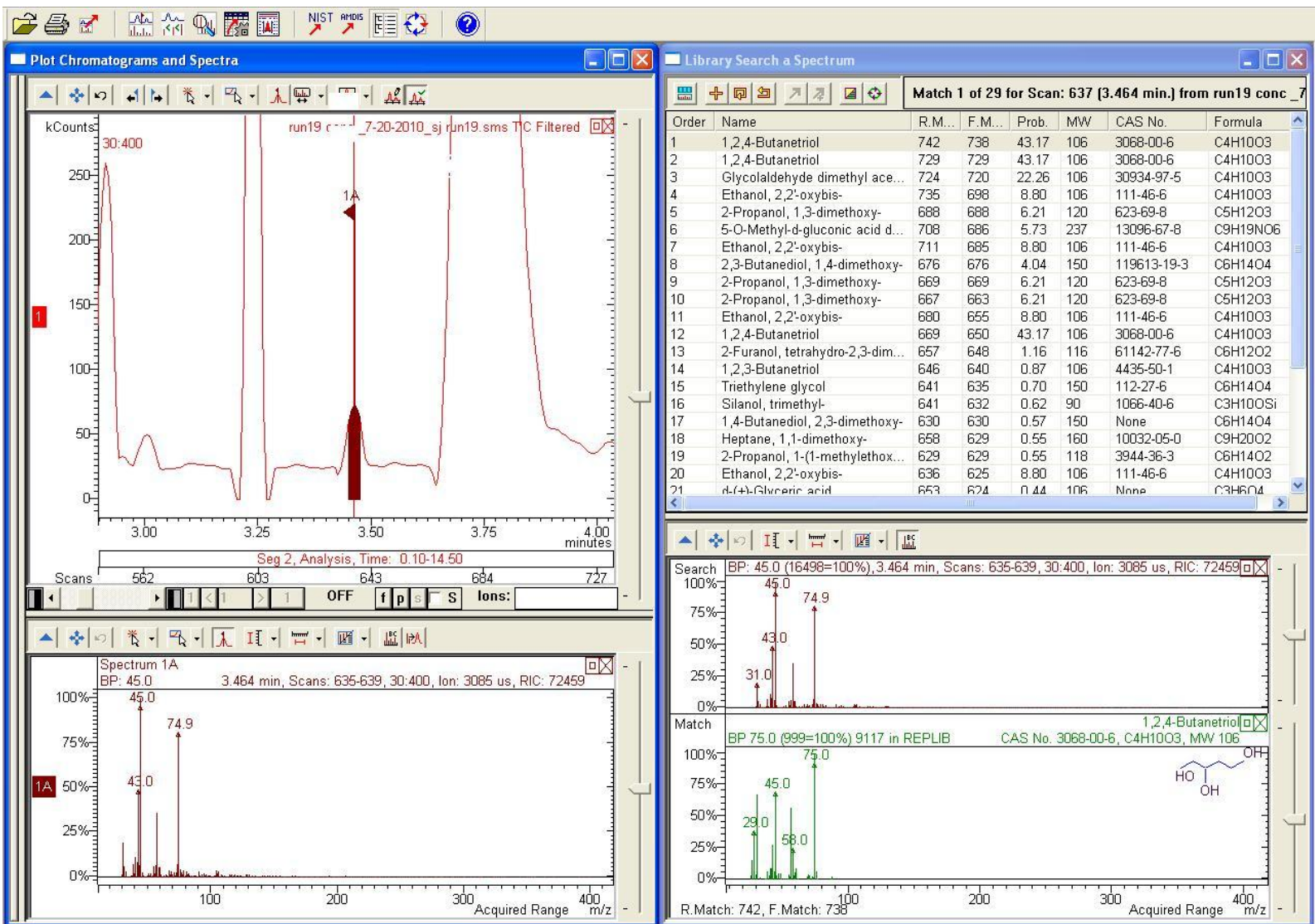


Figure E.6. Identification of 1,2,4-butanetriol in the biodiesel sample by GC-MS

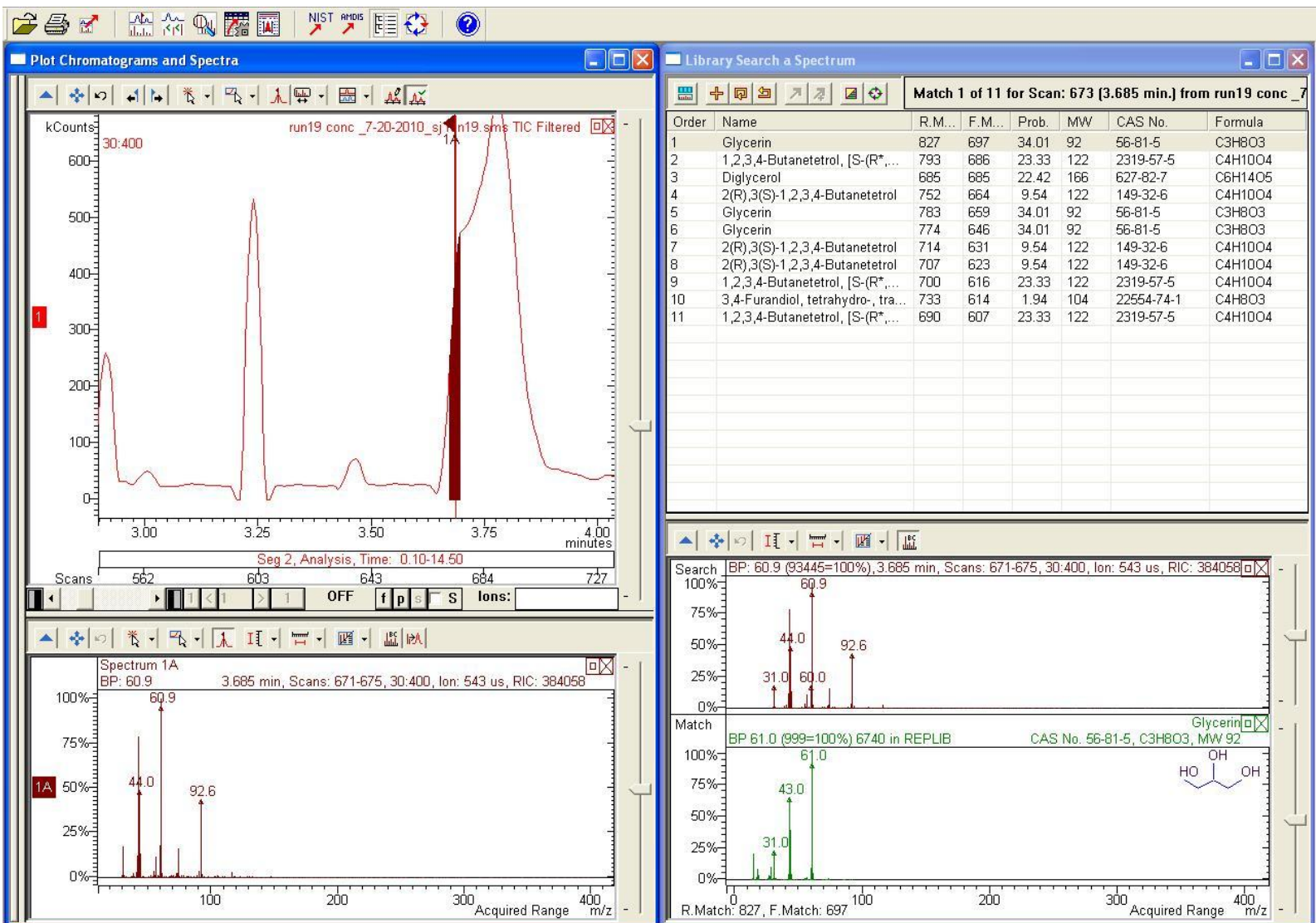


Figure E.7. Identification of glycerol in the biodiesel sample by GC-MS

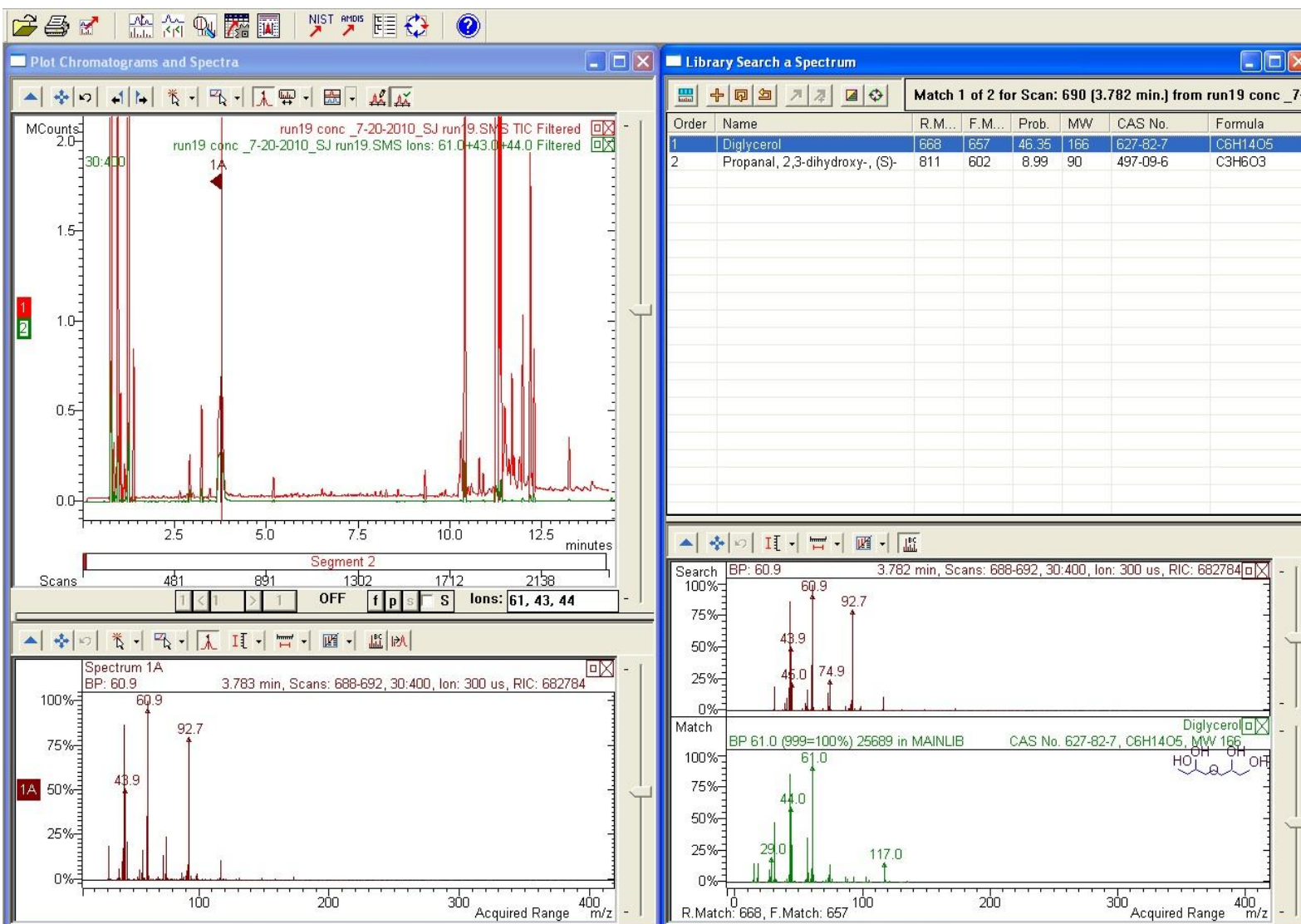


Figure E.8. Identification of diglycerol in the biodiesel sample by GC-MS

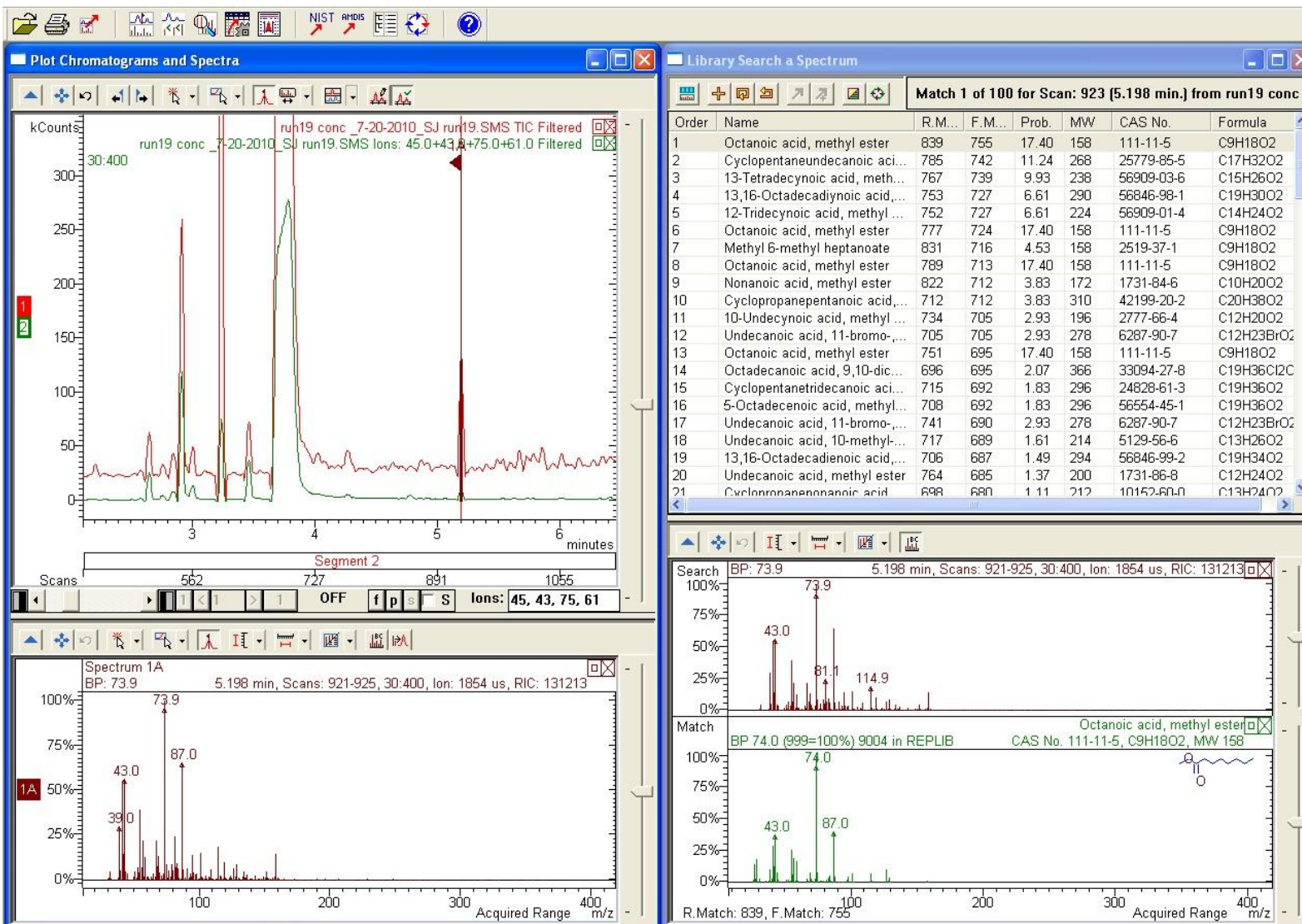


Figure E.9. Identification octanoic methyl ester in the biodiesel sample by GC-MS