Factors Affecting the Quality of Biodiesel from Palm Fatty Acid Distillate at Palm Oil Refining Plant

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Abstract: A study on factors affecting biodiesel quality of agricultural by-products, namely palm oil derived using palm fatty acid distillate (PFAD), collected from the Oleen Palm Oil industrial refining plant. This PFAD showed free fatty acid content and a saponification value of 88.4 % and 204 mg KOH/g, respectively. An acid catalyst was successfully used to produce biodiesel in the esterification reaction, and a 97.11% conversion to biodiesel based on the European Standard EN 14214:2003 was achieved under the conditions (PFAD to methanol molar ratio 1:3.71 with 1.834 % H₂SO₄ catalyzed at 121 °C for 15 minutes). Overall, this novel process achieved highly enhanced FAME (95.82% to 97.31%) with a significantly increased reaction time (10 to 30 minutes) and catalyst requirements (1.834 % H₂SO₄).

Keywords: palm fatty acid distillate; esterification; biodiesel.

1. Introduction

Thailand is one of the productive palm oil producers in the world. In the year 2018, 15.53 million tonnes of palm oil products from Thailand were produced [1]. It is reported that about 776,750 tonnes of Palm fatty acid distillate (PFAD) were produced in Thailand during the palm oil refining process (Figure 1). PFAD containing very high free fatty acids (FFA) is a by-product of the palm oil purification process [2-4]. It is generally used in several industrials (cleaners, animal feeds, plastics). Moreover, other researchers reported in the literature biodiesel production using PFAD [4-6]. The biodiesel can be synthesized by esterifying low-quality oils containing high FFA (> 90%) palm fatty acid with alcohols, as shown in Figure 2. Therefore, there is a need for innovation. The PFAD is generally used in nonfood applications such as soap making and is also used as a power source in power plants and industrial boilers [2, 3]. To produce biodiesel from high FFA oil, esterification has been frequently used to convert the FFA content in oil to esters [6, 7]. However, excess alcohol and catalyst loading must be used in the acid-catalyzed esterification to obtain high purity and yield of biodiesel from high FFA [8-10]. In the esterification step, the generated wastewater hinders the extent of esterification, and the methanol and sulfuric acid are diluted by the generated wastewater [8-10].
PFAD samples are potential substrates for biodiesel fuel production, biochemical, soap, and oleochemical [2-4, 13]. Thailand has a great advantage in developing biodiesel production since PFAD can be used as feedstocks for biodiesel production.

This study aims at factors affecting biodiesel production quality for value-adding PFAD from the palm oil refining plant at Oleen Co., Ltd.

2. Materials and Methods

2.1. PFAD samples analysis.

The samples of PFAD were collected at the Oleen Palm Oil Plant (Samutsakorn, Thailand) and stored at room temperature (26 ± 2 ºC). The PFAD was analyzed for free fatty acid (FFA) by AOCS Ca 5a-40 [14], saponification value by AOCS Cd 3-25 [14], and fatty acid composition was determined by gas chromatography (AOCS Ce 1-62) [14].

![Figure 1. Palm Fatty Acid Distillates (A) Biodiesel PFAD (B).](image1)

![Figure 2. Esterification Reaction.](image2)

2.2. Experimental biodiesel production.

The esterification of PFAD was carried out using reactor 5 liters per bath for biodiesel production from palm fatty acid distillate with high pressure (Figure 3).

![Figure 3. Biodiesel esterification reactor 5 liters.](image3)
Additionally, the percentage conversion of FFA in the processes was compared under similar reaction conditions such as methanol to PFAD molar ratio, acid catalyst dosage, temperature, and reaction time.

2.2.1. Effect of methanol for biodiesel production from PFAD.

Biodiesel was produced from the PFAD from the palm oil refining plant at Oleen Co., Ltd., single-step esterification method. In this, esterification process the PFAD was mixed with methanol in the ratio of 1:1.24, 1:2.47, and 1:3.71 (molar ratio). 98% concentrated sulfuric acid (1% v/v PFAD) was added as a catalyst to enhance the esterification reaction to the mixture. The mixture of PFAD, methanol, and the catalyst was allowed to react in the reactor at 121 °C for 30 minutes under high pressure of 15 bar approximately. The mixture obtained consisted of two layers; the upper layer of methanol and sulfuric acid and the lower layer consisting of the esterified oil. With the help of a separating funnel, the esterified oil was separated from the methanol and other impurities. Finally, sodium sulfate was added to methyl ester to remove water vapors and then filtered with a filter paper. The reaction yield was analyzed for FFA by AOCS Ca 5a-40 [14], and the percent conversion of esterification was calculated. Percent conversion was measured by total free fatty acid before esterification and after esterification.

\[
\text{Conversion (\%)} = \left(\frac{(a-b)}{a}\right) \times 100
\]

When;

\begin{align*}
a & = \text{total free fatty acid before esterification} \\
b & = \text{total free fatty acid after esterification}
\end{align*}

The reaction runs and FFA analysis were done in triplicate, and the results are expressed as mean value and standard deviation (SD). Results were analyzed statistically using SPSS software version 21.0. Data were tested by analysis of variance ANOVA and evaluate significant difference by LSD at the P=0.05 level.

2.2.2. Effect of acid catalyst dosage for biodiesel production from PFAD.

Applying the procedure detailed in section 2.2.1, the product containing the highest percentage of conversion was prepared again under the corresponding suitable conditions. Other batches of biodiesel were prepared under the same conditions as above (maximum percent conversion) and differing loads of acid catalyst dosage (0.611, 1.223, 1.834, and 2.445 %). The reaction yield was analyzed for FFA by AOCS Ca 5a-40 [14], and the percent conversion of esterification was calculated. Results were analyzed statistically using SPSS software version 21.0. Data were tested by analysis of variance ANOVA and evaluation of significant difference by LSD at the P=0.05 level.

2.2.3. Effect of temperature and time for biodiesel production from PFAD.

By applying the procedure detailed in sections 2.2.1. and 2.2.2, the product containing the highest percentage of conversion was prepared again under the corresponding suitable conditions. Other batches of biodiesel were prepared under the same conditions as above (maximum percent conversion) and differing loads of temperature (70, 100, 121, and 130 °C). The reaction yield was analyzed for FFA by AOCS Ca 5a-40 [14], and the percent conversion of esterification was calculated. Results were analyzed statistically using SPSS software
version 21.0. Data were tested by analysis of variance ANOVA and evaluation of significant difference by LSD at the $P = 0.05$ level.

Then the highest percent of conversion was prepared again under the corresponding suitable conditions and differing loads of time (10, 15, 20, 25, and 30 minutes). The reaction yield was analyzed for FFA by AOCS Ca 5a-40 [14] and the calculated percent conversion of esterification. Each prepared biodiesel was then placed in a closed Eppendorf tube and stored for one week in refrigeration (below $10 \pm 2 \, ^\circ\mathrm{C}$) before measuring the fatty acid methyl ester (FAME) content by a TLC-FID analyzer. Results were analyzed statistically using SPSS software version 21.0. Data were tested by analysis of variance ANOVA and evaluation of significant difference by LSD at the $P = 0.05$ level.

2.4. Determination of FAME.

Fifty microliters of the sample after esterification were mixed with 50 $\mu$L of chloroform, and the percentage of FAME was analyzed by a TLC-FID analyzer (IATROSCAMTM MK-5, Iatron Laboratories Inc., Tokyo, Japan).

3. Results and Discussion

3.1. Composition of PFAD.

PFAD and crude palm oil (CPO) were analyzed for FFA and saponification. Acidity and saponification value was $88.5 \pm 0.1\%$ and $204.2 \pm 1.1 \, \text{mg \, KOH/g \, oil}$, and $4.6 \pm 0.2\%$ and $200.5 \pm 1.1 \, \text{mg \, KOH/g \, oil}$ for PFAD and CPO, respectively (Data not show). Fatty acid compositions of samples are given in Table 1. PFAD and CPO are confirmed as sources of saturated and monounsaturated fatty acids.

The analyzed fatty acid composition of the oils used in this study (Table 1) is consistent with published values. PFAD and CPO contain high percentages of palmitic (16:0) and oleic (18:1) acids as major components. PFAD and CPO show a higher proportion of saturated fatty acids (56.9 and 48.0 respectively) compared to unsaturated fatty acids. But CPO contains less palmitic acid (41.49 \%) than PFAD (49.46\%). PFAD contains 35.0 \% of monounsaturated fatty acids, including 33.95 \% of oleic (18:1) acid. The percentage of oleic acid in CPO is higher than in PFAD, and the same applies to polyunsaturated fatty acids. Subsequently, the percentage of trans fatty acid in PFAD is higher than in CPO, probably because of the catalysis of isomerization by the activated bleaching clay or high temperature due to physical refining [2-4].

| Fatty acid            | PFAD (%) | CPO (%) |
|-----------------------|----------|---------|
| Saturated fatty acid  |          |         |
| Lauric (C12:0)        | 0.05     | 0.33    |
| Myristic (C14:0)      | 1.29     | 0.99    |
| Palmitic (C16:0)      | 49.46    | 41.49   |
| Stearic (C18:0)       | 5.17     | 4.49    |
| Arachidic (C20:0)     | 0.45     | 0.40    |
| Behenic (C22:0)       | 0.22     | 0.09    |
| Other                 | 0.23     | 0.19    |
| Monounsaturated fatty acid | 35.0   | 41.7     |
| Palmitoleic (C16:1 n-7c) | 0.14 | 0.17 |
| Elaidic (C18:1 n-9t)  | 0.67     | 0.11    |
| Oleic (C18:1 n-9c)    | 33.95    | 40.18   |
| Eicosenoic (C20:1 n-9c) | 0.14 | 0.14 |
| Other                 | 0.10     | 1.08    |
Table 2. Effect of PFAD to methanol molar ratio on esterification with 1.834 % H₂SO₄ catalyzed at 121 °C for 30 minutes.

| PFAD : Methanol (molar ratio) | FFA (%)        | Conversion (%) |
|-------------------------------|----------------|---------------|
| 1 : 1.24                      | 13.34±0.02ab   | 84.61abc      |
| 1 : 2.47                      | 2.97±0.01bc    | 90.02bc       |
| 1 : 3.71                      | 2.16±0.01c     | 97.56a        |

* a Values within the same column having the same or without superscript are not significantly different (p>0.05); Data is written as mean value and standard deviation.

3.2.2. Effect of acid-catalyzed.

The product containing the highest percentage of conversion was prepared again under the corresponding suitable conditions. The addition of methanol to the reaction mixture was investigated using a molar ratio of PFAD to methanol 1:3.71, and differing loads of acid-catalyzed 0.611-2.445 % (Table 3). When the acid-catalyzed was 1.223 %, the FAME conversion was higher than 90 %, and when the acid-catalyzed was 1.834 %, it was 97.56 %. This might be due to a diffusion limitation of methanol in the acid-catalyzed esterification on the PFAD. When the temperature was 121 °C, the FAME conversion reached 97.56% in a reaction time of 30 minutes. The ester content could be reached more than 97% compared to the EN 14103 standard (96.5 % min).

Table 3. Effect of acid-catalyzed on esterification of PFAD to methanol molar ratio 1:3.71 at 121 °C for 30 minutes.

| H₂SO₄ (%) | FFA (%)        | Conversion (%) |
|-----------|----------------|---------------|
| 0.611     | 11.22±0.01a    | 87.12bc       |
| 1.223     | 6.67±0.01bc    | 90.02bc       |
| 1.834     | 2.16±0.01c     | 97.56a        |
| 2.445     | 2.15±0.01c     | 97.25a        |

* a Values within the same column having the same or without superscript are not significantly different (p>0.05); Data is written as mean value and standard deviation.
3.2.3. Effect of reaction temperature.

The product containing the highest conversion percentage was prepared again under the corresponding suitable conditions. The addition of methanol to the reaction mixture was investigated using a molar ratio of PFAD to methanol 1:3.71 and differing loads of temperature 70-130 °C (Table 4). When the temperature was 70 °C, the FAME conversion was lower than 90 %, and when the temperature was 121 °C, the FAME conversion reached 97.56 % in a reaction time of 30 minutes. The ester content could be reached more than 97 % in comparison with the EN 14103 standard. However, in all temperature cases, the conversion of PFAD to FAME increased and was higher than 85 %. In addition, according to the kinetic theory, when the temperature is increased, the pressure also increases. As the particle gains kinetic energy, the mass transfer rate between the oil-methanol-catalyst phases increases, hence, producing FAME yield in a shorter time [16-20].

Table 4. Effect of temperature on esterification of PFAD to methanol molar ratio 1:3.71 with 1.834 % H₂SO₄ catalyzed for 30 minutes.

| Temperature (°C) | FFA (%)    | Conversion (%) |
|-----------------|-----------|---------------|
| 70              | 9.62±0.01a| 89.12b        |
| 100             | 6.17±0.01b| 93.02b        |
| 121             | 2.16±0.01c| 97.56a        |
| 130             | 2.34±0.01c| 97.35a        |

* a Values within the same column having the same or without superscript are not significantly different (p>0.05); Data is written as mean value and standard deviation.

3.2.4. Effect of reaction time.

To investigate the effect of the reaction time on the esterification reaction rate, a series of experiments has been performed by varying the reaction time from 10 to 30 minutes, as shown in Table 5. The FAME yield increased non-significantly after 15 minutes and remained constant when the reaction time was further increased to 30 minutes. Once the reaction system reached the desired temperature, the reaction was rapid, and the FAME yield immediately reached 97.11–97.31 %. This proves the potential of the acid catalyst under high temperature and pressure to shorten the time of the esterification reaction.

Table 5. Effect of time on esterification of PFAD to methanol molar ratio 1:3.71 with 1.834 % H₂SO₄ catalyzed at 121 °C for 10-30 minutes.

| Time (minutes) | FFA (%)    | Conversion (%) | FAME (%) |
|----------------|-----------|---------------|----------|
| 10             | 3.26±0.02a| 96.31b        | 95.82a   |
| 15             | 2.34±0.01b| 97.35a        | 97.11a   |
| 20             | 2.26±0.01b| 97.44a        | 97.23a   |
| 25             | 2.20±0.01c| 97.51a        | 97.32a   |
| 30             | 2.16±0.01c| 97.56a        | 97.31a   |

* a Values within the same column having the same or without superscript are not significantly different (p>0.05); Data is written as mean value and standard deviation.

In Thailand and Europe, biodiesel quality is assessed under the provisions and the requirements of quality standard EN 14214. Quality control was performed for the biodiesel produced under the optimum process conditions specified above (i.e., the molar ratio of PFAD to methanol 1:3.71 and obtained about 97.11% FAME yield at 121 °C, with 1.834 % H₂SO₄ catalyzed and 15 minutes reaction time), based on the European Standard EN 14214:2003. Density at 15 °C, acidity number, methyl ester content, the content of monoglycerides, diglycerides, triglycerides, total and free glycerol were determined and measured. To further
scale up biodiesel production from PFAD, all quality characteristics specified by the European standard EN 14214 should be determined.

4. Conclusions

Palm fatty acid distillate makes this by-product a suitable starting feedstock for biodiesel manufacture. The highest FAME yield produced was 97.11\% in the presence of 1.834 \% H\textsubscript{2}SO\textsubscript{4} catalyst loading, 1:3.71 PFAD:methanol molar ratio, at 121 \degree C within 15 minutes. In conclusion, the acid catalyst with high temperature and pressure showed potential to enhance the esterification reaction rate of PFAD with low biodiesel production costs, high FAME yields, and short reaction times.

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Conflicts of Interest

The authors declare that they have no conflict of interest. The funders had no role in the study's design, in the collection, analyses, or interpretation of data, in the writing of the manuscript, or in the decision to publish the results.

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