SYNTHESIS OF ALUMINA-CaO-KI CATALYST FOR THE PRODUCTION OF BIODIESEL FROM RUBBER SEED OIL

(Sintesis Mangkin Alumina-CaO-KI untuk Penghasilan Biodiesel daripada Minyak Biji Getah)

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Abstract

In this research, biodiesel was produced by transesterification of rubber seed oil with methanol catalyzed by three types of alumina-supported catalyst. The catalysts were Al2O3-CaO, Al2O3-KI, and Al2O3-CaO-KI. The catalysts were prepared by heating a mixture of alumina and the corresponding salt in a furnace at 700 °C. Transesterification reactions were carried out at 65 °C using a mixture of rubber seed oil: methanol of mass ratio 1:9 with different catalyst loadings between 0 and 3.5%. The optimum catalyst loading was at 2.0% for all types of catalyst and Al2O3-CaO-KI gave the highest yield (91.6%) followed by Al2O3-KI (90.7%) and Al2O3-CaO (63.5%). The catalyst Al2O3-CaO-KI also gave high biodiesel yields over a wider range of catalyst loadings compared to the other two catalysts. Reactions were also carried out at different temperatures (25, 40 and 65 °C). It was observed that the yield increased drastically with increasing reaction temperature. At all temperatures the Al2O3-CaO-KI catalyst gave the highest yield. Therefore, the study showed that among these three catalysts the most productive was Al2O3-CaO-KI, (100:30:35).

Keywords: alumina-supported catalyst, biodiesel, rubber seed oil, transesterification

Abstrak

Dalam kajian ini, biodiesel dihasilkan melalui tindak balas transesterifikasi minyak biji getah dengan metanol menggunakan tiga jenis mangkin tersokong alumina. Mangkin-mangkin tersebut adalah Al2O3-CaO, Al2O3-KI dan Al2O3-CaO-KI. Mangkin-mangkin itu disediakan dengan pemanasan campuran alumina dan garam yang berkaitan di dalam relau pada suhu 700 °C. Tindak balas transesterifikasi dijalankan pada suhu 65 °C dengan menggunakan campuran minyak biji getah: metanol dengan nisbah jisim 1:9 dengan beban mangkin yang berbeza di antara 0 hingga 3.5%. Beban mangkin yang optimum adalah 2.0% untuk semua jenis mangkin dan mangkin Al2O3-CaO-KI memberikan peratus hasil yang paling tinggi (91.6%) diikuti oleh Al2O3-KI (90.7%) dan Al2O3-CaO (63.5%). Mangkin Al2O3-CaO-KI juga memberikan peratus hasil biodiesel yang tinggi pada julat beban mangkin yang lebih besar berbanding dengan dua mangkin yang lain. Tindak balas juga dijalankan pada suhu-suhu yang berbeza (25, 40 dan 65 °C). Didapati bahawa peratus hasil meningkat secara mendadak dengan peningkatan suhu tindak balas. Pada sebarang suhu beban mangkin Al2O3-CaO-KI memberikan peratus hasil yang paling tinggi. Oleh itu, kajian ini menunjukkan bahawa di antara ketiga-tiga jenis mangkin ini Al2O3-CaO-KI, (100:30:35) adalah yang paling produktif.

Kata kunci: mangkin tersokong alumina, biodiesel, minyak biji getah, transesterifikasi
Introduction

The excessive use of fossil fuels has stirred environmental concern due to their non-renewable nature. Therefore, as an alternative biodiesel is becoming a popular choice of renewable fuel. Biodiesels are fatty acid methyl esters that can be produced from fats or oils from plants or animals. There are two chemical reactions which can be used to produce biodiesel, namely esterification, and transesterification. The more commonly applied reaction is transesterification since it involves only a single step reaction between oil and methanol. Transesterification reaction can be carried out with or without the presence of a catalyst [1]. However, the use of a catalyst can reduce the production time by speeding up the reaction. There are three types of catalysts which are acid, base and enzyme. Acids catalysed transesterification is a slow reaction, therefore is not favourable [2]. On the other hand, enzymes are too costly. The cost of enzyme-catalyzed production of biodiesel is about twice that of the alkali-catalyzed process [3]. Therefore, this research focuses on the application of a base catalyst to produce biodiesel from rubber seed oil. Base catalysts can be in the form of a homogeneous catalyst which is soluble in the reaction medium or heterogeneous catalyst which are insoluble. However, the homogeneous base catalyst is sensitive to water and free fatty acid content in the feedstock where it will cause the formation of soap. This will cause difficulty in the purification of the biodiesel because soap will lead to the formation of oil-water emulsion [4].

Currently, the use of heterogeneous catalysts is gaining attention among researchers since they are reusable and therefore could save the production cost. These catalysts are in the form of metal oxides such as alkaline earth metal oxides, MnO and TiO [5, 6]. Even though these catalysts are reusable, but some studies have shown that they are subjected to leachate where part of the solid dissolves in the reaction medium and thus limiting the reusability [7]. An approach to overcome this problem is to bind the catalyst to a solid support. One of the materials that have been studied extensively as a support is an alumina.

As to evaluate the catalytic activity of alumina supported, they were used KI, CaO, and CaO-KI to transesterify palm oil into biodiesel. The alumina supported-KI showed the highest conversion [8-10]. However, since palm oil is edible oil, then there is an issue of competition with the food supply. Moreover, there was reported, 925 million people were facing with malnutrition [11]. As such, in this research, we are utilizing rubber seed oil, which is a non-edible oil to produce biodiesel. The aim of this research is to compare the catalytic activity of alumina-CaO, alumina-KI, and alumina-CaO-KI in the transesterification of rubber seed oil with methanol.

Materials and Methods

Rubber seeds were collected from local rubber plantations. Aluminium oxide powder, potassium iodide and calcium acetate were purchased from R&M Chemicals. Methanol (99%) was from HmbG Chemicals. The chromatographic standards used were methyl heptadecanoate as internal standard and the FAME standard mixture GLC-10. Both were from Sigma-Aldrich. GLC-10 consists of methyl palmitate, methyl stearate, methyl oleate, methyl linoleate and methyl linolenate.

Rubber seeds were crushed, and the kernel was separated from the shell. The kernel was dried in the oven at 60 °C overnight and then milled by using a household blender. Rubber seed oil was extracted from the kernel by a Soxhlet extractor using petroleum ether for 6 hours. The solvent was evaporated in a rotary evaporator. The oil was then kept in a refrigerator.

Three types of catalysts were prepared. They are alumina-CaO-KI, alumina-CaO, and alumina-KI. The alumina-CaO-KI catalyst was prepared as follows. A 33.8 g of calcium acetate and 14 g of potassium iodide were dissolved in 100 mL water in a beaker. Then 40 g of alumina was added to the mixture. The mixture was stirred for 4 hours at room temperature. Then it was dried in an oven at 100 °C overnight. The dried mixture was then calcined at 700 °C for 5 hours. The mass ratio of alumina: CaO: KI in the catalyst is 100: 30: 35. The mass of CaO produced from calcium acetate was calculated based on TGA data. The alumina-CaO was prepared with the same method but without adding potassium iodide. Alumina-KI was also prepared with the method but without adding calcium acetate.

The Brunauer-Emmett-Teller (BET) surface area analysis of porous properties of the catalysts was determined by Micromeritis ASAP 2000. Adsorption N₂ gas within 6 hours at 250 °C (de-gassed). Investigation of the chemical
composition and morphology of the catalysts done by using FESEM-EDX (JSM 7800F). While, X-Ray Diffraction analysis (XRD) was achieved by using Rigaku with Cu Kα X-ray as a source, 20 range from 2° to 80° with step sizes of 0.02°, at a scanning speed of 1° min⁻¹.

Transesterification reaction was carried out in a three-neck flask equipped with a thermometer, a condenser, and a magnetic stirrer. 10 g of rubber seed oil was mixed with 90 g methanol and various amounts of the catalyst. The amount of catalyst used were between 0 and 3.5% (weight percent to total reactants). The reactions were carried out at 65 °C and stirred at 600 rpm for 5 hours. After the reaction was completed, the catalyst was removed by filtration, and the reaction mixture was poured into a separatory funnel. Two layers formed where the lower layer is glycerol, and the upper layer is biodiesel (FAME) solution in methanol. The glycerol layer was discarded, and biodiesel was obtained from the upper layer by evaporation with a rotary evaporator. Reactions were also carried out at 25 °C and 40 °C using 2% catalyst loading to study the effect of temperature on the biodiesel yield.

Quantification of biodiesel was done by Agilent 7890B gas chromatographer equipped with 5977A MSD and Zebron ZB-Wax column (30 m x 0.25 mm x 0.25 μm). The injector and MS transfer-line temperatures were set at 240 °C and 280 °C, respectively. The column temperature was held at 110 °C for 2 minutes and then programmed to 240 °C at 6 °C/min. The methyl esters were quantified by an internal standard method using methyl heptadecanoate as internal standard and compared against GLC-10 FAME standards (Supelco).

**Results and Discussion**

The BET surface area, pore volume and pore size of different catalysts from the previous and currently studies are reported in Table 1. Referring the table, the surface area of the catalysts is ranging from 4.9283 – 48.8130 m²/g. The literature data of Al₂O₃ act as a catalyst showed ranging 120 – 265 m²/g of the surface area. The decreasing in surface area was considering the sequence of catalyst displacement on the metal oxide, Al₂O₃. During the impregnation and calcination method, the reduction of BET surface area also occurred and reflecting the pore volume and pore size. The phenomena were apparently due to the insertion of CaO or KI into pores and surface of Al₂O₃. An identical measurement has been discussed by several researchers [9, 10, 12-14]. Referring to this range of pore size (2 to 50 nm = mesopore), the average pore size of the catalysts ranged from 10.9 to 18.6 nm categorize as mesoporous type. Meanwhile, the pore diameter less than 2 nm is microporous and on top of 50 nm is macroporous.

### Table 1. BET characterization of the catalysts.

<table>
<thead>
<tr>
<th>Samples</th>
<th>Asri et al. [10]</th>
<th>Zabeti et al. [9]</th>
<th>Present Study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface area (m²/g)</td>
<td>83.77</td>
<td>54.551</td>
<td>83.74</td>
</tr>
<tr>
<td>Pore volume (cm³/g)</td>
<td>0.135</td>
<td>0.083</td>
<td>0.223</td>
</tr>
<tr>
<td>Pore size (x 10⁻⁸m)</td>
<td>3.421</td>
<td>3.821</td>
<td>63.37</td>
</tr>
</tbody>
</table>

The crystal morphology exposes through the micrograph of the catalyst. The CaO and KI on the surface of amorphous Al₂O₃ create porous materials. The impregnation and calcination method, made spherical crystals on the surface become obvious needle-shaped spicules of Al₂O₃ with a porous apparent in all of the catalysts. As can be seen in Figure 1, no distinct different between KI and KI/CaO samples thus propose highly dispersed on the surface of support catalyst. Also, notified the sustained structure of Al₂O₃ after impregnation with another compound was important. Almost similar observation reported by [8]. The composition of catalysts is displayed in EDX analysis. The result showed the existence of Al (29.55%), O (18.69%) and Ca (7.04%) for Al₂O₃/CaO. Meanwhile for Al₂O₃/KI, confirms the composition of O (23.77%), I (23.48%), Al (13.48%) and K (9.25%). Also, the EDX
The XRD profile of the catalysts is presented in Figure 2. The result demonstrated the characteristic peaks at 2θ ranging from 20° to 70°. It showed large reflection at 29.66°, 25.62° and 25.46°, respectively and confirmed to be Al₂O₃. This is recommended that the structure of Al₂O₃ undefined although loading with other compound. Meanwhile, small reflection of diffraction peaks from KI and CaO suggesting it was well dispersed on the surface of the support catalyst [15].

Production of biodiesel by transesterification of rubber seed oil refluxed with methanol was carried out by using three alumina-supported catalysts. Biodiesel yield was observed by varying the catalyst loading between 0 to 3.5 %. As shown in Figure 3, the yield increased with increasing catalyst loading from 0 to 2% loading for all of the
catalysts. The highest yield of biodiesel with alumina CaO-KI, alumina-KI and alumina-CaO were 91.6%, 90.7% and 63.5% respectively. It was found that the optimal catalyst loading was at 2.0% but decreasing yield of biodiesel was observed with further increasing amount of catalyst. As expected, alumina-CaO-KI gave the highest yield of biodiesel probably due to the combination of factors; the high basicity of alumina-KI catalyst [16] and higher activity in the mild reaction condition of CaO catalyst [17]. Similarly, high biodiesel yield (95%) was also observed from palm oil using same catalyst [10].

![Graph showing effect of catalyst loading on biodiesel yield](image1)

**Figure 3.** Effect of the catalyst loading on the biodiesel yield

![Graph showing effect of temperature on biodiesel yield](image2)

**Figure 4.** Effect of temperature on the biodiesel yield

The decline of biodiesel yield when the catalyst loading exceeded 2.0% was due to the mixing effectiveness of the reaction mixture. Beyond the optimum catalyst loading, there is a mass transfer limitation of the reactants. Previous studies by other researchers also showed the same trend [18, 19].

Calcium oxide is a popular heterogeneous catalyst because of its availability. However, the activity of this catalyst will decline after multiple uses due to leachate into the reaction medium. An approach to overcome this problem is to bind the calcium oxide onto a catalyst support such as alumina [9]. Other than calcium oxide, some potassium salts supported on alumina also showed high catalytic activity. Previously, it was discovered that alumina-supported potassium iodide with 35% KI loading gave the highest catalytic activity among several alumina-supported potassium salts [8]. It was also observed that when CaO was combined with this catalyst the optimum loading of
CaO was 30% as used in our study [10]. In this study, we tried to investigate whether a combination of CaO and KI supported onto alumina gives a higher catalytic activity compared to when any of them are used separately. Amazingly, the alumina-CaO-KI catalyst was capable of producing higher biodiesel yields compared to alumina-CaO or alumina-KI.

According to Figure 4, the yield increased drastically when the temperature was raised from 25°C to 65°C. At 25°C the yield for all catalysts was only between 4.7% and 14.8%, whereas at 65°C it was between 63.5% and 91.6%. At all temperatures, Alumina-CaO-KI gave the highest yield. The study was done only up to a maximum reaction temperature of 65°C because it is the boiling temperature of methanol and thus, the highest temperature that can be achieved by refluxing. Moreover, to increase the temperature beyond that, special equipment is required. Previously, a study using alumina supported CaO as a catalyst also showed a drastic increase in biodiesel yield between 55 °C and 65 °C [9]. Similarly, the same trend was also observed using a catalyst made of a mixture of calcium oxide and manganese oxide [20].

**Conclusion**

Biodiesel was successfully produced from rubber seed oil by transesterification with methanol using alumina-supported CaO-KI as the catalyst. In this research, we have studied the activity of three types of alumina-supported catalysts, namely alumina-supported CaO, alumina-supported KI and alumina-supported CaO-KI. It was proven that a combination of CaO and KI supported onto alumina had a higher catalytic activity than any of the salts supported separately onto alumina. The highest biodiesel yield achieved was 91.6% using alumina-CaO-KI (100:30:35) with a 2.0% catalyst loading.

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