CLAMSHELL AND SEA SAND AS HETEROGENEOUS CATALYSTS FOR WASTE COOKING OIL BASED BIODIESEL PRODUCTION VIA TRANSESTERIFICATION REACTION

(Arsyad Ismail Sanuzi, Syuhada Mohd Tahir*, Siti Norhafiza Mohd Khazaai
Faculty of Applied Sciences, Universiti Teknologi MARA Cawangan Pahang, 26400 Bandar Tun Abdul Razak Jengka, Pahang, Malaysia
*Corresponding author: syuhadamohdtahir@pahang.uitm.edu.my
Received: 4 December 2016; Accepted: 1 December 2017)

Abstract
This study was carried out to study the effectiveness of clamshell and sea sand as heterogeneous catalyst to produce biodiesel via transesterification of waste cooking oil (WCO). WCO was first pre-treated to discard water and impurities. Clamshell and sea sand were crushed into fine powder, sieved to 500 µm particles and calcined. Next, the pre-treated WCO was used to obtain fatty acid methyl ester (FAME) via transesterification reaction. FTIR and XRD results of calcined clamshell confirmed the formation of CaO and Ca(OH)\textsubscript{2} while sea sand contains SiO\textsubscript{2} before and after calcination. 2:1 clamshell-to-sea sand ratio produced highest FAME yield. This ratio was used to study the effect of catalyst loading amount. 7 wt% catalyst produced highest FAME yield. Increasing catalyst to 10 wt% reduced the FAME yield due to the soap formation. Analysis of FAME using GC-MS showed the presence of myristic acid, palmitic acid and oleic acid. This study shows that combination of clamshell and sea sand is good potential catalyst for transesterification reaction to produce biodiesel.

Keywords: biodiesel, heterogeneous catalyst, transesterification, waste cooking oil

Abstrak
Kajian ini dijalankan untuk mengkaji keberkesanan kulit kerang dan pasir pantai sebagai mangkin heterogen untuk menghasilkan biodiesel melalui transesterifikasi minyak masak sawit terpakai (WCO). WCO dirawat terlebih dahulu untuk membuang air dan kekotoran. Kulit kerang dan pasir pantai dihancurkan menjadi serbuk halus, ditapis kepada 500 µm dan dikalsinasi. Tindakbalas transesterifikasi WCO dijalankan untuk menghasilkan metil ester (FAME). Data daripada FTIR dan XRD untuk kulit kerang terkalsinasi mengesahkan pembentukan CaO dan Ca(OH)\textsubscript{2} manakala pasir pantai pula mengandungi SiO\textsubscript{2} sebelum dan selepas kalsinasi. Nisbah 2:1 kulit kerang terhadap pasir pantai menghasilkan FAME yang paling tinggi dan nisbah ini digunakan dalam kajian seterusnya untuk melihat kesan jumlah pemangkin terhadap jumlah FAME yang dihasilkan. Mangkin sebanyak 7 wt% menghasilkan jumlah FAME yang paling tinggi. Penggunaan mangkin sebanyak 10 wt% mengurangkan jumlah FAME yang diperoleh disebabkan pembentukan sabun. Analisis FAME menggunakan GC-MS menunjukkan kehadiran asid miristik, asid palmitik dan asid oleik. Kajian ini menunjukkan campuran kulit kerang dan pasir pantai adalah mangkin yang berpotensi baik untuk tindakbalas transesterifikasi untuk menghasilkan biodiesel.

Kata kunci: biodiesel, mangkin heterogen, transesterifikasi, minyak masak sawit terpakai
Introduction

Biodiesel is an attractive alternative for conservative petroleum-based fuel and can be used directly in commercial diesel engines. Biodiesel is made of natural and renewable sources such as new and used vegetable oils and animal fats. Biodiesel made from edible vegetable oils have undesirable impact to the environment as they require a large amount of farmland [1]. Waste cooking oil (WCO) is an abundant, readily available and cheap raw material for biodiesel. The use of WCO can minimize biodiesel production costs up to 60-90% [2].

Biodiesel can be produced by two methods: acid esterification and alkaline transesterification. The common catalysts for acid esterification are H$_2$SO$_4$ and Fe$_2$(SO$_4$)$_3$. Unfortunately, this method produced water along with the desired ester and inhibits transesterification of glyceride. Alkaline transesterification is more preferred method because it is 4000 times faster than acid esterification method. Two types of catalyst in alkaline transesterification are homogeneous and heterogeneous catalysts. Homogeneous catalyst (NaOH and KOH) is effective and affordable but highly sensitive to water and free fatty acid content and require high energy consumption for purification [2]. Researchers have found various types of heterogeneous catalysts such as ion exchange resins, sulfated oxides, transition metal oxide, boron group, alkaline earth metal oxide, mixed metal oxide, alkali metal oxides, waste material, carbon based and enzyme based [2]. Heterogeneous catalysts are environmentally friendly, economical and have high chance for heat integration [3].

CaO, an alkaline earth metal oxides is an ionic crystal and is an established heterogeneous catalyst. The Lewis acidity of the metal cation due to its modest electronegativity makes it a good heterogeneous catalyst [4]. The conjugated oxygen anions in the crystal structure posses strong basic property. In this study, clamshell was chosen as the renewable resource for CaO. Clamshell has high content of calcium carbonate (CaCO$_3$), when calcined at proper temperature, will be converted into CaO. In this study, CaO from clamshell will be combined with SiO$_2$ from seasand in order to increase the catalytic effectiveness.

Materials and Methods

The clamshells were cleaned and dried in oven at 105 °C for 24 hours and crushed into a fine powder. Next, the powder was sieved through 500 µm. The powder obtained was then calcined in a furnace at 900 °C for 3.5 hours [5]. The sea sand was rinsed, dried, screened to extract larger particles and sieved through 500 µm. Then, sea sand was calcined at 800 °C for 2 hours [6]. The catalysts were characterized by using FTIR and XRD.

WCO sample was collected from a restaurant at Bandar Pusat Jengka, Pahang. WCO was first filtered to remove suspended solid particles and heated at 80 °C. Then, the oil was washed with water (approximately 10% of the oil volume) and centrifuged at 600 rpm for 10 minutes to remove water soluble salt impurities. Lastly, the organic phase collected and kept under vacuum at 80 °C for 30 minutes to remove water [6].

Transesterification process carried out in three-necked round bottom flask equipped with reflux, thermometer and magnetic stirrer. WCO and methanol with a molar ratio of 1:12 were mixed with catalyst and refluxed at 60 °C for 6 hours at 600 rpm. The solid catalyst was separated by filtration afterwards. The resulting liquid was placed in a separatory funnel and left for 24 hours at room temperature, until two layers were formed. The top layer was FAME in methanol and the bottom layer was glycerol. The methanol layer was collected and evaporated in a rotary evaporator to obtain the biodiesel (FAME). Table 1 shows the catalysts composition used in this study to observe the effect of catalyst ratio for 5 wt% weight of catalyst over weight of WCO. The optimum ratio with highest yield of biodiesel was used to further study on the effect of loading amount of the catalyst at 7 and 10 wt%. All temperature and time taken for transesterification reaction were kept constant.

FTIR spectra were recorded using Spectrum 100, Perkin Elmer, in the range of 450-4000 cm$^{-1}$ wavenumbers. The catalysts were characterized before and after calcined using XRD model D-5000 Siemen. The data was collected at diffraction angle 20 from 3° to 80° at the rate of 0.04 s$^{-1}$. The FAME was analyzed with gas chromatography-mass spectrometry (GC-MS) using Agilent 5977A MSD Mass Spectrometer/Data System at initial temperature of 125 °C for 4 min 15 °C per minute temperature increment until reaching final temperature at 275 °C [7].
Table 1. Catalysts composition used for 5 wt% weight of catalyst over weight of WCO

<table>
<thead>
<tr>
<th>Weight of Catalyst Over Weight of WCO (%)</th>
<th>Clamshell-to-Sea Sand Ratio</th>
<th>Clamshell (g)</th>
<th>Sea Sand (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1:0</td>
<td>0.50</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>0:1</td>
<td>0</td>
<td>0.50</td>
</tr>
<tr>
<td>5</td>
<td>1:1</td>
<td>0.25</td>
<td>0.25</td>
</tr>
<tr>
<td></td>
<td>1:2</td>
<td>0.17</td>
<td>0.33</td>
</tr>
<tr>
<td></td>
<td>2:1</td>
<td>0.33</td>
<td>0.17</td>
</tr>
</tbody>
</table>

**Results and Discussion**

Figure 1 shows the FTIR spectrum and the absorption peaks of uncalcined and calcined clamshell, respectively. Uncalcined clamshell showed three main absorption peaks belonging to three vibration modes for $\text{CO}_2^-$ molecule; asymmetric stretch, out-of plane bend and in-plane bend. During calcination, $\text{CaCO}_3$ will decompose into CaO and $\text{CO}_2$, as shown in equation 1 [8]. Exposure of CaO to atmospheric air cause the formation of $\text{Ca(OH)}_2$ as shown in equation 2, hence a sharp OH bond stretching band observed at 3640 cm$^{-1}$ [5]. The presence of CaO and OH absorption peaks prove the complete decomposition of $\text{CaCO}_3$ at the chosen calcination temperature.

\[
\text{CaCO}_3 \rightarrow \text{CaO} + \text{CO}_2 \quad (1)
\]

\[
\text{CaO} + \text{H}_2\text{O} \rightarrow \text{Ca(OH)}_2 \quad (2)
\]

FTIR spectrum of uncalcined and calcined sea sand in Figure 2 shows an almost similar absorption peaks, means calcination did not change the chemical structure of sea sand and calcination was not necessary. The four major absorption peaks observed were classified as quartz or silicon dioxide, SiO$_2$. The peaks at 778.84 cm$^{-1}$ and 777.95 cm$^{-1}$ refers to Si-O symmetrical stretching vibration while 1086.93 cm$^{-1}$ and 1084.92 cm$^{-1}$ refers to Si-O asymmetrical stretching [9].
Figure 2. The FTIR spectra of sea sand; a) uncalcined and b) calcined

Figure 3 represents the XRD diffractograms for uncalcined and calcined clamshell. The result revealed that the composition of uncalcined clamshell dominantly consists of CaCO$_3$ (aragonite). The diffraction peaks for CaCO$_3$ can be observed at 2\(\theta\) = 26.34\(^{\circ}\), 33.24\(^{\circ}\) and 45.93\(^{\circ}\). After calcination at 900 \(^{\circ}\)C, CaCO$_3$ completely transformed into CaO by releasing CO$_2$. The peaks of CaO (lime syn) appeared at 2\(\theta\) = 32.38\(^{\circ}\), 37.52\(^{\circ}\), and 53.98\(^{\circ}\) with (111), (200) and (220) orientations. In addition, Ca(OH)$_2$ (portlandite) also observed at 2\(\theta\) = 18.16\(^{\circ}\) and 34.26\(^{\circ}\) due to hydration of CaO by atmospheric moisture [10]. These XRD result proves the calcination temperature of clamshell, 900 \(^{\circ}\)C for 3.5 hours was suitable to completely decompose CaCO$_3$ into CaO.
Figure 4 represents the XRD patterns for uncalcined and calcined sea sand. The result revealed that the composition of uncalcined sea sand dominantly consists of SiO₂. These intensity diffraction peaks can be observed at 2θ = 20.89° and 26.84°. Meanwhile, the peaks of SiO₂ for calcined sea sand were observed at 2θ = 20.94° and 26.70°. SiO₂ was the major constituent due to high quartz content rather than any mineral in the sea sand. The JCPDS standard pattern number 46-1045 values for SiO₂ (quartz) are 2θ = 20.82° and 26.62°.

![XRD diffractograms of uncalcined sea sand and calcined sea sand (■:SiO₂)](image)

According to GC-MS result, each samples of FAME produced contain palmitin. Their peak ranges between 10.00 – 12.00 minutes. Uzun et al. [11] stated that five properties of WCO-based FAME is myristic acid, palmitic acid, oleic acid, linolenic acid, and linoleic acid. However, the column for GC-MS instrument used in this study only capable to detect three of them; myristic acid, palmitic acid and oleic acid. Figure 5 shows the GC-MS spectra of FAME obtained and detailed in Table 2.
Arsyad Ismail et al: CLAMSHELL AND SEA SAND AS HETEROGENEOUS CATALYSTS FOR WASTE COOKING OIL BASED BIODIESEL PRODUCTION VIA TRANSESTERIFICATION REACTION

Figure 5. GC-MS spectrum of FAME using 7 wt% loading amount with 2:1 clamshell-to-sea sand ratio

Table 2. The scan range of FAME properties 2:1 clamshell-to-sea sand ratio

<table>
<thead>
<tr>
<th>Properties</th>
<th>Scan Range (minutes)</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>5 wt%</td>
<td>7 wt%</td>
<td>10 wt%</td>
<td></td>
</tr>
<tr>
<td>Palmitic acid</td>
<td>7.774</td>
<td>7.774</td>
<td>7.775</td>
<td></td>
</tr>
<tr>
<td>Stearic acid</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Lauric acid</td>
<td>4.713</td>
<td>4.713</td>
<td>4.714</td>
<td></td>
</tr>
<tr>
<td>Palmitoleic acid</td>
<td>7.597</td>
<td>7.597</td>
<td>7.592</td>
<td></td>
</tr>
<tr>
<td>Ascorbic acid</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Glycidyl palmitate</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Glycidyl oleate</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Cyclopropanebutanoic acid</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
</tbody>
</table>

To observe the effect of clamshell-to-sea sand ratio on FAME production, five ratios were chosen: 1:0, 0:1, 1:1, 1:2, and 2:1. Loading amount used was fixed at 5 wt%. The result shown in Figure 6. It was found that the highest percentage FAME yield obtained was 71.1% with clamshell-to-sea sand ratio of 2:1. This result shows that the combination of two catalysts increase the percentage yield significantly and the greater amount of clamshell in the catalyst ratio gave higher biodiesel yield. The presence of CaO and Ca(OH)₂ in the calcined clamshell was believed as the factor. O²⁻ in CaO and OH⁻ in Ca(OH)₂ are the adsorptive sites that will extract H⁺ from methanol to form methoxide anions that will then react with the fatty acid in triglyceride molecules to form FAME.
The clamshell-to-sea sand ratio of 2:1 was used to further study the effect of catalyst loading on FAME production. As shown in Figure 7, 7 wt% loading amount of the catalysts produced highest FAME yield of 75.3%. Increasing the loading amount from 5 to 7 wt% increase the percentage yield but further increase to 10 wt% reduced the yield. High basicity in CaO and Ca(OH)\(_2\) helps in breaking the bond of free fatty acid chain with glycerol in producing FAME. Insufficient catalyst dosage will reduce the breaking process results in incomplete transesterification and reduce the FAME yield. FAME yield increased with the increase in the catalyst loading amount since more triglyceride molecules will be broken. However, excess alkaline catalyst loading could results in more triglyceride participation leading to soap formation and thus decrease biodiesel yield especially in a longer retention time [12].

Figure 6. Effect of clamshell-to-sea sand ratio on FAME yield using 5 wt% catalyst loading

Figure 7. Effect of loading amount of catalyst on FAME yield using 2:1 clamshell-to-sea sand ratio
Conclusion
Waste cooking oil-based biodiesel (FAME) was produced via transesterification reaction. The effect of ratio and loading amount of combination of alkaline heterogeneous catalyst; clamshell as source of CaO and sea sand as source of SiO$_2$ was studied. FTIR and XRD results showed the calcination of clamshell at 900 °C for 3.5 hours were suitable to completely decompose CaCO$_3$ into CaO. Presence Ca(OH)$_2$ also observed as the result of reaction of CaO with atmospheric moisture. Combination of two catalysts was found to increase the percentage yield significantly and the greater amount of clamshell in the catalysts ratio gave higher FAME yield. The optimum ratio for clamshell-to-sea sand was 2:1 and this ratio was used to study the effect of loading amount of the catalysts in FAME percentage yield. Increasing the loading amount from 5 to 7 wt% increase the percentage yield but further increase to 10 wt% loading amount reduced the yield due to the soap formation especially in a longer retention time. 7 wt% loading amount of the catalysts produced highest FAME yield of 75.3%. In conclusion, combination of clamshell and sea sand possesses good potential as heterogeneous catalyst in WCO-based FAME production.

Acknowledgements
The authors would like to thank Universiti Teknologi MARA and Ministry of Higher Education, Malaysia for research grant no RAGS/1/2014/SG01/UITM/3.

References