Study of calcined eggshell as potential catalyst for biodiesel formation using used cooking oil

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Abstract: Eggshells were utilized as a cost effective catalyst to assist in biodiesel formation. Eggshells were calcined using different calcination conditions such as temperature and time. The eggshells underwent calcination under 1000 °C in which calcium carbonate was completely converted into calcium oxide under open air conditions. The calcined eggshell catalyst was characterized by X-ray diffraction (XRD). Then the eggshell derived CaO catalyst was subjected to transesterification of used cooking oil (UCO). The parametrics in the transesterification of used cooking oil such as a methanol to oil ratio, weight of catalyst and reaction temperature were evaluated. The optimum result exhibited a 4:1 methanol to oil ratio, 2wt % catalyst, and a 65 °C reaction temperature within two hours. Pure calcined CaO acted as the control of the experiment that was subject to the most optimum eggshell derived catalyst transesterification conditions. The catalytic activity for both the pure calcined CaO and the calcined eggshell derived catalyst were comparable. The biodiesel formation was identified by gas chromatography mass spectrometry (GC-MS).

Keywords: Calcination; eggshell; Calcium oxide; Transesterification; Used cooking oil.

1 Introduction

The continuous increase in crude oil prices and the depletion of petroleum reserves, has aroused researchers to develop alternative ways to replace gas-derived fuels. Biodiesel has appeared as an alternative fuel to replace gas-based fuels [1]. Costs of raw materials, usually refined vegetable oil, add to the production cost [2]. As a consequence, the ideas of used cooking oil (UCO) as raw material feedstock are generated [3]. Many scientists applied homogenous alkaline catalysts in biodiesel production due to the low cost and high availability. CaO is one of widely-used catalysts due to its abundant availability in nature, its high activity and low cost [4].

During the past few years, researchers have begun focusing on the regeneration of waste materials such as eggshells as useful substances. Eggshell waste comprises a big portion of food industrial waste as eggs are a daily diet in many countries. For instance, dried eggs, liquid eggs, and frozen eggs are the diversified form of egg products that are found in markets. An egg is considered a relatively cheap food that provides complete nutrition with protein sources (6g), unsaturated fatty acids (10%), essential minerals and vitamins such as A, D [5]. The egg industry is a leader is one of the major sectors of the world food market.

According to literature, there are many added-values derived from waste eggshells such as calcium phosphate bioceramics [6], hydroxyapatite(HA) as implanted constituents [7], and as absorption of dyes[8]. Waste eggshells consist of a majority of calcium carbonate compounds, nearly 95% of which are suitable as activated heterogeneous catalysts used in biodiesel production [9]. Calcium oxide components can be generated from eggshell waste after calcination and used in the transesterification process.

This study addresses the following objectives: one is to synthesize a potential catalyst from eggshell waste and to identify most effective calcination conditions for the calcined eggshells derived catalyst. The second objective is to determine the effect of the methanol to oil ratio, the weight of the catalyst and the reaction temperature using the most effective calcined eggshell conditions derived catalyst to form biodiesel.
2 Materials and methods

2.1 Catalyst preparation

The wasted chicken eggshells were collected by selling Char Keow Teow stall. The eggshells were peeled from their membranes to remove any interference substances and impurities such as organics materials. After that, chicken eggshells were rinsed three times with tap water. Then, the eggshells were dried in a hot air oven for 24 hours at 60°C [9].

Next, the dried eggshells were pulverized into fine powder using a blender. The powder was sieved to constant sizes such as 0.045 mm by using a buger sieve with the help of an automatic sieve shaker. Next, eggshell powders were transferred into a furnace to generate calcium oxide. The eggshells, potential catalysts were cooled down and placed in a desiccator to reduce the probability of reacting with any humidity and CO₂ in the air. The acidic particles like CO₂ and H₂O are considered poisonous to the catalyst’s active site [10]. Hence, the calcined eggshells needed to be kept in the desiccator to avoid contact with those particles. Additionally, the silica gel needed to have a blue color instead of a pink color. Blue silica gel is made up of cobalt chloride, and it will turn pink when saturated with water and cannot absorb any moisture.

Different conditions were applied to synthesize eggshells as potential catalysts. For instance, eggshells calcination using a muffle furnace under 800 °C for 4 hours (Condition 1), eggshells calcination using a tube furnace under 1000 °C for two hours with nitrogen gas flow (Condition 2), and eggshells calcination using a tube furnace under 1000 °C for two hours without nitrogen gas flow (Condition 3).

Pure calcium oxide acted as a experimental control and was subjected to the same calcination conditions.

2.2 Characterization of catalyst using X-Ray Diffraction (XRD)

In order to prove the potential of eggshells as catalysts, XRD analysis was conducted in the uncalcined or calcined chicken eggshell powder. Apart from that, pure calcium oxides either uncalcined or calcined were observed in XRD.

With the 2θ range from 10° to 90°, the XRD powder diffraction analysis of the samples were carried out using Cu Kα radiation with a 2°/min scanning speed and a 0.04° step size[9].

The components found in the respective calcination methods were identified. The diffraction pattern was determined by referring to the International Centre for Diffraction Data (ICDD) using the powder diffraction file PDF-2 Release 2014 as a reference database.

2.3 Pre-treatment of used cooking oil

Used cooking oil was collected from the cafeteria of the Jeli Campus at the University of Malaysia, Kelatan. First, dispersed particles and any residues in the oil were filtered using cheese cloth. Then, centrifugation was employed to discard any impurities or fat sediment. Then the oil was filtered using filter paper to get a better quality oil. Next, oil was heated at 110°C for 50 minutes to evaporate any trace amount of moisture in a 2L beaker with magnetic stirrer [11].

2.4 FFA identification

Total quantities of FFA in the oil samples were identified using the titration method [12] before any reactions were carried out. First, the amount of 0.5 grams of potassium hydroxide (KOH) was weighed and dissolved into 500 mL of distilled water in a 1 L beaker. The potassium hydroxide (KOH) was then transferred into a burette.

1 mL of used cooking oil was dissolved into 10 mL of isopropyl alcohol in a 50 mL beaker. The beaker consisted of used cooking oil. The isopropyl alcohol was heated in a water bath for 15 minutes at 60 °C. Color changes were observed ranging from a dark brown to a plain yellowish solution as the used cooking oil slowly dissolved into isopropyl alcohol.

The dissolved solution of used cooking oil together with isopropyl alcohol was then transferred into a 500 mL Erlenmeyer flask. Then two drops of phenolphthalein were added. Potassium hydroxide (KOH) was added slowly from the burette into the alcohol-oil mix solution, and the solution mixture was swirled gently. The light pink color appeared which lasted for 10 seconds revealing the endpoint measurement and the final volume of potassium hydroxide (KOH) which was then recorded. The experiment was repeated three times to collect the average titration value.

The calculation formula for the FFA% from a titration value is shown in Equation 1

\[
\text{FFA\%} = \frac{(t - b) \times N \times 28.2}{v} \quad (1)
\]
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$t$ indicates amount (mL) of titration sample

$b$ indicates amount (mL) of the blank sample

$N$ indicates normality (N) of the titration sample

$v$ indicates quantity (g) of the oil sample

2.5 Synthesis of biodiesel through transesterification

The reaction took place using a 250 mL Erlenmeyer flask, a magnetic stirrer, a reflux condenser, a hot plate, a thermometer, and a 1 L beaker which acted as a water bath. First, the calcined eggshell catalyst was mixed with methanol for one hour to produce calcium methoxide. 50 mL of used cooking oil was then transferred accordingly into a new Erlenmeyer flask. The calcium methoxide solution which had stirred for one hour was poured into the oil. The content was refluxed under magnetic stirring for two hours in a closed vessel to avoid the evaporation of methanol[13]. The reaction duration was fixed at 4 hours due to the longer reaction time, there will be hindrance to the catalyst hence suppress the biodiesel yield [14].

Three parameters were studied using calcined eggshells as a potential catalyst. Those were the weight of the calcined eggshell catalyst (1.0 wt %, 2.0 wt %, 3.0 wt%, 4.0 wt%, 5.0 wt%), the reaction temperature (35 °C, 45 °C, 55 °C, 65 °C) and the methanol to oil ratio (1:1, 2:1, 3:1, 4:1, 5:1). The reaction temperature was set below 65 °C due to the boiling point of methanol being at 65 °C. Methanol starts to bubble as it boils which would hinder the three-phase interface reaction (methanol, CaO, and oil) to take place[15].

2.6 Separation and purification

After transesterification, the solution mixture underwent a gravity separation overnight in a separatory funnel. After that, the separation was complete with densities and the upper layer as methyl ester, methanol and the bottom layer as glycerine[16].

The upper layer was subjected to centrifugation at 6000 rpm for 15 minutes to remove the residue catalyst; meanwhile the bottom layer was withdrawn and discarded. Next, warm water was required to purify the sample product by rinsing out the impurities such as methanol, glycerol, CaO, and soap. Warm water was introduced slowly to avoid emulsion and soap formation [17]. The washing of biodiesel was stopped until the water appeared transparent and clear compared to the cloudy yellowish solution from previous rinsings.

Then, the sample product was subjected to heating to evaporate the residual moisture. Finally, the biodiesel was cooled to room temperature and sealed with a parafilm kept in a dark place.

2.7 Analysis of fatty acid and methyl esters content

The fatty acid component of used cooking oil and its methyl esters content of sample products were carried out using Agilent 7890 N gas chromatography with an Agilent 7693 mass selective detector. The capillary column applied had a 30 m x 0.25 mm dimension i.d 0.25 lm film thickness (Agilent Technologies) with HP-5MS (5% phenyl methylsiloxane).

The gas chromatograph setting was achieved accordingly using a fatty methyl ester (FAME) procedure. The molecular mass was set at a range from 30 – 300. The desired compounds were matched according to the peak and retention time based on the NIST (National Institute of Standards and Technology) library.

3 Results and Discussion

3.1 Conversion of eggshell into potential catalyst

3.1.1 XRD pattern of different calcination methods of eggshell samples

XRD analysis was carried out to identify the component and crystallinity of eggshell components. Figure 1 represented an XRD pattern of eggshells before calcination and after calcination at two different temperatures ranges, that were 800°C and 1000°C and with two different conditions such as with closed air, open air and with nitrogen flow or without nitrogen flow.

Based on the pattern of the X-ray diffraction on raw eggshells, only calcium carbonate existed in the raw eggshell powder before calcination (A). This depicted that calcium carbonate was the prime compound in the eggshell. Calcium carbonate was the major compound for the eggshells after close atmosphere calcination under temperature 800°C (B) as observed in figure 1. After closed atmosphere calcination at 800°C (B), the intensity
The intensity of the calcium carbonate for calcination at 800°C (B) changed compared to the uncalcined eggshells which was due to the calcium carbonate gradually transforming into calcium oxide[15]. A similar study demonstrated that the calcium carbonate did not completely convert into calcium oxide during the 800°C calcination process[18].

A calcium oxide peak only appeared after calcination at 1000°C with nitrogen gas flow (C) and in open air (D). This was due to the high thermal transition that occurred in the eggshells causing complete decomposition of the calcium carbonate into calcium oxide which occurred at 1000°C.

The XRD pattern for both the open air calcined eggshells at 1000°C (D) and the nitrogen gas flow calcined eggshells at 1000°C (C) showed identical and highly crystallite clear peaks. The crystallite size of eggshells after calcination under 1000°C higher compared calcination under 800°C[19].

The calcination of eggshell waste within the two to four-hour timeframe found the catalyst activities comparative to the transesterification process[21]. Extending the calcination time frame would diminish the catalyst and suppress the active site whereby dropping the catalytic activity of the eggshell in transesterification. Therefore, it was suggested that the timeframe should not exceed 4 hours accordingly with the different calcination temperature.

### 3.1.2 XRD pattern of different calcination methods of Pure CaO samples

The XRD analysis was carried out to identify the pure calcium oxide component and to determine whether it had potential in becoming a catalyst without calcination, or calcination under different conditions. Figure 2 displays an X-ray diffractogram of pure calcium oxide before calcination, and pure calcium oxide after being subjected to different calcination conditions.

Based on the pattern of the X-ray diffraction of pure calcium oxide before calcination (A), two main components appeared to include calcium oxide and calcium hydroxide. The presence of calcium hydroxide was probably due to the uptake of moisture from the surrounding environment. This indicated that the pure calcium oxide should be subjected into in order to get rid of other than calcium oxide component before carried on with transesterification process.

However, after calcination at 800°C (B), the calcium oxide peak appeared with high crystallinity and exhibited after calcination.
a lesser calcium hydroxide component than before calcination (A). However, calcium hydroxide did not fully convert into calcium oxide under calcination at 800°C (B). Thus, this indicated that the calcium oxide transformation required a higher temperature or longer calcination timeframe to completely remove the moisture content in CaO.

As depicted from figure 2, the calcium oxide peak only appeared after calcination at 1000°C with a nitrogen gas flow (C) and with open air (D). This showed that the calcination at a higher temperature, but shorter time could help eliminate the water content in CaO.

The XRD pattern for both open air calcined eggshells at 1000°C (D) and nitrogen gas flow calcined eggshells at 1000°C (C) showed high crystallite. The method for the open air calcined eggshell at 1000°C (D) was selected to make the potential catalyst as it could be achieved at a reduced cost compared to the nitrogen gas method and the cost for purchasing).

3.2 Optimization of transesterification using calcined eggshell derived catalyst

3.2.1 Effect of methanol to oil ratio towards biodiesel formation

A different methanol to oil molar ratio in this study was investigated from 1:1 to 5:1 as other factors remained constant. The reaction took place at a temperature of under 65°C for two hours, and used a 3wt% calcined eggshell catalyst and 50 ml of used oil.

Figure 3 depicted that the methanol to oil ratio of 1:1 to 5:1, biodiesel yield increase from 2ml to 4ml. Initially, the methanol to oil molar ratio from 1:1 to 2:1 did not have any further change, it stagnated at 2 ml of biodiesel yield. However, the volume of biodiesel gradually increased from 2ml to 4ml, from 2:1 to 4:1 of methanol to oil ratio respectively. As the methanol volume increased so did the biodiesel yield due to calcium methioxide, Ca(CH$_3$O)$_2$, that formed during the pre-treatment of the calcium oxide with methanol before transesterification. This triggered the reaction equilibrium to shift to the right. Calcium methioxide can boost the reaction between oil and methanol [21].

The biodiesel yield no longer increased and stayed at 4ml for the 4:1 to 5:1 methanol to oil ratio. This could be ascribed to the excess of methanol preventing the gravity separation of methyl ester and glycerol fragment into two phases [14]. Researchers demonstrated that the higher methanol to oil ratio would promote a better FAME yield.

However, the further increment in the methanol to oil ratio based on the system, caused the transesterification process to be reversible. The glycerol residues would shift the equilibrium to the left and hence the FAME yield would drop [22].

The biodiesel yield was relatively low compared to other studies; this was probably due to the slightly higher FFA content from used cooking oil. The biodiesel yield was highly related to the FFA content [23]. Researchers suggested that the way to maximize the methyl esters yield was to reduce the FFA content to 2% before carry on with transesterification.

3.2.2 Effect of catalyst concentration towards biodiesel formation

The effect of catalyst concentration, varying quantities from 1 wt% to 5 wt% according to the weight of initial used cooking oil towards the biodiesel formation were investigated. The reaction took place at a temperature below 65°C for two hours, and used 150ml methanol(4:1) and 50 ml used oil. The outcomes were revealed in figure 4.

From figure 4, the conversion rate was identified to be slightly accelerated from 1wt% to 2 wt% of catalyst concentration that was from 3ml to 4ml. This result described that the increment in weight of the catalyst boosted the contact between the reactant and the catalyst[15]. The more the catalyst used, the faster the reaction due to the faster collision between reactants molecules.

However, the for the further increase of catalyst concentration from 2 wt% to 3 wt%, volume of biodiesel drops from 4ml to 3ml, following by no any fluctuation in biodiesel volume for catalyst concentration of 4wt% and 5 wt%.
The drop in biodiesel yield was due to the FFA and the water residue in used cooking oil which tended to react and cause soap formation during the transesterification process. The free fatty acids reacted with the alkaline catalyst that lead to saponification. Saponification depleted the alkaline catalyst inevitably lowering the effectiveness of the catalyst, hence collecting a lower biodiesel yield [15].

In addition, as the CaO used more, the leaching process from the CaO catalyst increased, resulting in more of the CaO to react with FFAs whereby turning into calcium soap during the purification process. Besides, there were certain amounts of biodiesel to form an emulsion during the washing process whereby lowering its yield. The amount of catalyst use was increased.[24].

### 3.2.3 The reaction temperature profile towards biodiesel formation

The reaction temperature was set higher than the room temperature such as 35 °C, 45°C, 55°C, and 65°C. The reaction took place using 150ml (4:1) methanol and 50 ml used oil and 3 wt% catalyst concentration for two hours. Figure 5 revealed that a slightly increase in temperature resulted in a higher biodiesel yield. At the beginning, the FAME yield was 3ml although the temperature increased from 35°C to 45°C. This result showed that the low temperature did not favor a transesterification reaction. However, the additional increase of the reaction temperature from 45°C to 55°C, lifted the volume of biodiesel from 3ml to 4ml, subsequently no variation in the biodiesel volume was noted for a further increase of the reaction temperature from 55°C to 65°C.

The further increase in biodiesel volume in alignment with the increase of the reaction temperature attributed to the solubility of the methanol towards triglycerides.

However, increasing the reaction temperature further did not influence the biodiesel yield [25]. This is due to the boiling point of methanol at nearly 68°C, the evaporation of methanol and the bubbling which would adversely hinder a reaction if the temperature were higher than 65°C [15].

### 3.3 Product analysis using GC-MS

Figure 6, 7 and 8 show the chromatograms of the initial used cooking oil after transesterification using calcined eggshells and after transesterification using calcined CaO. The chromatogram peaks exhibit the outcome of separation of particular compounds, which were detected by the gas chromatogram. Chromatograms of natural derived CaO from calcined eggshells and pure calcined CaO after transesterification display some similarity in terms of peak shape and size as revealed in figures 7 & 8. The chromatogram peak and its components using calcined eggshells and using calcined pure CaO are exactly the same in table 1 & 2.

There are four methyl ester components in both chromatograms such as methyl tetradecanoate, hexadecanoic acid, methyl ester, 9-Octadecenoic acid, methyl ester, and 5,8,11,14-Eicosatetraenoic acid, methyl ester.

Besides, with the aid of mass spectrum, the selection of resemble mass to charge ratio of each possible compounds could be done more precisely. The desired compounds were selected based on the NIST library that comprises a database of mass spectra. The mass spectrum and its respective structure are displayed in used cooking oil, oil after transesterification using calcined eggshells and oil after transesterification using calcined CaO are displayed in figure 9 & 10.
Figure 6: The chromatograms of used cooking oil before transesterification.

Figure 7: The chromatograms of used cooking oil after transesterification using calcined eggshell.

Figure 8: The chromatograms of used cooking oil after transesterification using calcined CaO.

Table 1: GC-MS analysis of sample product using calcined eggshell as the catalyst.

<table>
<thead>
<tr>
<th>No</th>
<th>Systemic name</th>
<th>Common name</th>
<th>Formula</th>
<th>Molecular mass</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>9-Octadecenoic acid, methyl ester, (E)</td>
<td>Methyl 9-octadecenoate</td>
<td>C₁₉H₃₂O₂</td>
<td>296</td>
</tr>
<tr>
<td>2</td>
<td>Hexadecanoic acid, methyl ester</td>
<td>Methyl palmitate</td>
<td>C₁₇H₃₄O₂</td>
<td>270</td>
</tr>
<tr>
<td>3</td>
<td>Methyl tetradecanoate</td>
<td>Methyl myristate</td>
<td>C₁₅H₃₀O₂</td>
<td>242</td>
</tr>
</tbody>
</table>
The three major components obtained from the mass spectrum analysis report were tabulated accordingly in tables 1 & 2. From table 1, methyl 9-octadecenoate, methyl palmitate and methyl myristate were formed as a biodiesel product after transesterification using calcined eggshells. Additionally, table 2 depicted methyl 9-octadecenoate, methyl palmitate and methyl myristate that were formed as a biodiesel product after transesterification using calcined CaO. In brief, the natural eggshells derived CaO can be used to replace the chemical CaO in future prospect.

### 4 Conclusion

In this study, different calcination methods on eggshell waste were developed to identify the calcium oxide component that acted as a catalyst and which was used in transesterification. Among different calcination methods, the open air calcination was the best option compared to the nitrogen gas flow method. The former did not require an extra cost in purchasing the nitrogen gas. The eggshell derived catalyst consisted only of CaO as an end product.
during the calcination process which was comparable to the calcined pure calcium oxide based on the XRD pattern. The optimal transesterification carried out using calcined eggshells was by the 4:1 methanol to oil ratio, 2wt % of catalyst weight and a reaction temperature of 65°C for 2 hours. Pure calcined calcium oxide acted as the experimental control and was subjected accordingly to the optimum calcined eggshell parameter. Both methyl ester components appeared almost at the same. It can be assumed that the natural derived CaO was capable of showing its catalytic activities during the transesterification process. Hence, it is assumed that the waste calcined eggshell can be replaced by the chemical calcium oxide. The cost of the biodiesel production could reduce the synthesizing catalyst from the waste materials.

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References


