Method Article

Synthesis of biodiesel from *Annona muricata* – *Calophyllum inophyllum* oil blends using calcined waste wood ash as a heterogeneous base catalyst

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**A B S T R A C T**

Naturally, biodiesel synthesized from highly viscous and high-density vegetable oil is usually unsuitable as fuel in the internal combustion engine. However, mixing/blending of two or more oils as a feedstock for biodiesel production could produce a low viscous fuel suitable for the engine. This study produced a novel heterogeneous base catalyst from waste wood ash (WWA) and applied it to synthesis of biodiesel from *Annona muricata* and *Calophyllum inophyllum* oilseed blend. The production route was via a two-step process due to the high free fatty acid of the blended oil. Process optimization of the transesterification step was carried out via response surface methodology (RSM). The strength of the developed catalyst was tested through catalyst regeneration and recyclability. The quality of the biodiesel was compared with biodiesel recommended standard.

- Waste wood ash contained a high percentage of calcium carbonate
- Blended oil produced oil of low viscosity
- Two-step production route was used for biodiesel synthesis
- Process optimization via hybrid design produced optimum biodiesel yield.

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**A R T I C L E  I N F O**

*Method name: API gravity blend ratio, Catalyst characterization, transesterification*

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**Specifications Table**

| Subject Area | Energy |
|--------------|--------|
| More specific subject area | Biodiesel |
| Method name | API gravity blend ratio, Catalyst characterization, transesterification |
| Name and reference of the original method | Synthesis of biodiesel from the binary blend of Annona muricata – Calophyllum inophyllum oil using fly ash |
| Resource availability | n/a |

**Background**

Renewable, but sufficiently considered as ecofriendly, nontoxic, economically competitive, and environmentally beneficial are terms used to describe the liquid biodiesel (biofuel) derived from biomass sources that can be synthesized via biological routes in the presence of suitable catalyst [1].

The base catalyst used for biodiesel synthesis is classified mainly as a homogeneous catalyst (hydroxide of Na/K) and heterogeneous catalyst (majorly from solid wastes). The work reported by Ozkan et al. [2] developed calcium oxide (CaO) from waste materials as a heterogeneous catalyst for the synthesis of biodiesel, while [3] employed the use of bio-based catalyst from cocoa pod husk as a heterogeneous catalyst for biodiesel production. Ogunkunle et al. [4] reported the production of biodiesel using both catalysts. The work of [5] produces biodiesel with the help of a heterogeneous catalyst, while [6] described the use of a heterogeneous catalyst in the optimization of the production of rubber seed/palm oil biodiesel, IDI diesel engine performance, and emissions. Milano et al. [7] further used heterogeneous catalysts for the conversion of waste cooking oil-Calophyllum inophyllum to biodiesel, but the work reported by Nath et al. [8] utilized the green heterogeneous base catalysts as an effective bio-based for the conversion of oil to biodiesel. Papasanne et al. [9] reported a novel process for biodiesel production from slum sludge whereas [10] studies the use of catalyst derived from cotton stalk for production of biodiesel from Madhuca indica oil. In another study, Choksi et al. [11] reported the development of catalyst derived from palm-fruit-bunch for biodiesel. However, the conversion of a low-value industrial waste into biodiesel via a catalyst derived from brewery waste was reported by Saravanan et al. [12]. The advantages of using heterogeneous catalysts include ease of recoverability and reusability, readily available, nontoxic, eco-friendly, and low cost [13,14].

Among the low-cost base catalysts is the waste wood ash (WWA), this is a fine powder that is a byproduct of combustion of wood such as burning wood in a home fireplace or an industrial power plant. Researchers have revealed that the major components of the waste wood ash are calcium carbonate (CaCO₃) [15], some find no carbonate but calcium oxide (CaO) [16], some show the presence of about 12% iron oxide due to soil contamination while some show none [17]. However, because of its availability, environmental pollution effects, non-weather resistance, unsuitability for polishing, and brittle structure, it is considered as base-feedstock for biodiesel synthesis from seed oil.

Biomass sources of feedstock for biodiesel production could be derived from vegetable oil or animal fat [9]. The oil could be edible or non-edible oil, depending on the acid value of the oil. Oil can be classified as edible if the acid value of the oil is less than 3.00 mg KOH/g oil (%FFA < 1.5), and can successively undergo transesterification reaction with a base, while the non-edible oil acid value must be greater than 3 mg KOH/g oil (%FFA ≥ 1.5), which required acid esterification (H₂SO₄/HCl) before transesterification with base.

Meanwhile, many fruits bearing seeds are produced all over the nation from under-utilized to the most utilized one. These seeds majorly constitute a nuisance to the environment, which results in environmental pollution due to the problem of disposal. Among the fruits producing seeds are Annona muricata (Soursop) and the Calophyllum inophyllum (Berry). Studies revealed that soursop seed is rich in oil ranging from 22.57% to 34.61%, the oil nature proved edible and contained high unsaturated compounds [18,19]. Berry on the other hand was reported to be rich in oil up to 75% and contained both linoleic (36.0) and oleic (37.6%) fatty acid [14,20].

Response Surface Methodology (RSM) is an optimization software package for variable optimization of complex processes that helps to understand the interaction of the variables, produce several
The results of the experimental runs, determine the level of significance, and predict the optimum conversion of the output. This software has been reported to apply to biodiesel synthesis from various oil feedstocks [13,14,21]. The software is capable of handling design related factors such as Central Composite Design (CCD), Central Composite Rotatable Design (CCRD), Box Behnken Design (BBD), Pentagon, Hexagon, D-Optimal, Distance-based, Modified Distance, 3-Level Factorial, Hybrid, One-Factor, User Design, and Historical Data. When dealing with four process variable factor with three/five levels, only CCD and Hybrid design can be exploited. Since CCD can give a higher number of experimental runs (30) with repetition runs, presence of lack of fit, and give room for outliers, Hybrid design (HD) is preferable to overcome the CCD drawback. HD can easily produce 16 runs with no lack of fit, no repetition, and does not give room for outliers.

Hence, this paper exploits the use of blended oil of *Annona muricata* and *Calophyllum inophyllum* for the synthesis of biodiesel in the presence of waste wood ash (WWA) as a base-catalyst. Blend oil physicochemical properties were determined through AOAC, 1997 [22] standard methods, catalyst characterization of WWA were determined using scanning electron microscopy (SEM), energy dispersive spectroscopy (EDS), X-ray diffraction analysis (XRD), Fourier transforms infrared spectroscopy (FTIR), BET isothermal adsorption, and Hammett indicators. The quality of the biodiesel was ascertained by comparing it with biodiesel standard (ASTM D6551 [23] and EN 14,214 [24]).

### Experimental

**Oil extraction and its physicochemical properties**

Oil was extracted from the cleaned milled seed powder of *Annona muricata* and *Calophyllum inophyllum* using a soxhlet extractor with n-hexane as solvent. The reaction temperature was set at the range of the boiling point of n-hexane, and the reaction time was maintained at 70 min for complete oil extraction from the two powders. The excess n-hexane in the oil was recycled and reused, while the residual cake was used as an animal feed supplement. The physicochemical properties of the two oils were determined to ascertain the suitability of the oil for biodiesel synthesis.

**Oils blending**

The oil’ blending was carried out based on the physicochemical properties of the oil depicted in Table 1. Since the most significant parameters needed to achieve volatile mixed oil are the viscosity, the specific gravity (density), and the API gravity ratio of the oil. In this case, the viscosities of the oils were determined using a viscosimeter and the specific gravity of the oil by using a specific gravity bottle. The API gravity of the oil was estimated using the equation already adopted by Adepoju [25] as

### Table 1

| Properties                      | AMO   | CIO   | Mixed/Blended oil | Total API gravity |
|---------------------------------|-------|-------|-------------------|-------------------|
| Moisture content (%)            | 0.011 | 0.011 | 0.001*            |                   |
| Viscosity @ 40 °C (mm²/s)       | 2.32  | 5.40  | 3.35±              |                   |
| Acid value (mg KOH/g oil)       | 1.56  | 6.84  | 4.10±              |                   |
| % Free Fatty Acid (FFA)         | 0.78  | 2.92  | 2.05±              |                   |
| Peroxide value (meq O₂/kg oil)  | 1.34  | 1.40  | ND                |                   |
| Saponification value (mg KOH/g oil) | 224.63 | 201.00 | ND                |                   |
| Iodine value (g I₂/100 g oil)   | 114.32| 68.56 | ND                |                   |
| Specific gravity                | 0.82  | 0.91  | ND                |                   |
| API gravity                     | 41.05 | 24.00 | ND                |                   |
| API gravity ratio (%)           | 63    | 37    | ND                |                   |
| Simplest Ratio                  | 1.7   | 1     |                   |                   |
| **Blended ratio**               | AMO₂₃; CİO₃₇ |        |                   |                   |

where, am = Value after mixed/blended, NYD = Not Determined.

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**References**

[13,14,21,22,23,24,25]
where: $SG$ is the specific gravity of the oil; API gravity is the American Petroleum Institute gravity. Hence, the mixed/blended ratio of the oil was computed by adding up the API gravity of oils, resulting in the total API gravity as stated in Eq. (2) (supplementary file). The blend ratio (BR) of the oil was estimated by dividing the API gravity of individual oil with the total API gravity in Eq. (2), as estimated in Eq. (3). The blended oil obtained was preheated on a magnetic shaker at 50 °C to achieve a homogeneous phase with low viscosity and specific gravity required for biodiesel production.

\[
API\ gravity_{Total} = SG_{AMO} + SG_{CIO}
\]  

(2)

\[
BR = \left( \frac{SG_{AMO}}{API\ gravity_{Total}} \times 100 : \frac{SG_{CIO}}{API\ gravity_{Total}} \times 100 \right)
\]

(3)

Where $API\ gravity_{Total}$ is the total API gravity, $SG_{AMO}$ is the specific gravity of Annona muricata, $SG_{CIO}$ specific gravity of Calophyllum inophyllum.

The% free fatty acid (%FFA) of the oils and the blend was determined using a standard titration method:

0. 1 M of 95% ethanolic potassium hydroxide (KOH) was titrated against the mixture containing 2.0 g of oil in a hot mixture of ethanol and petroleum ether (1:1), with two drops of phenolphthalein as an indicator (Table 1).

Catalyst preparation and characterization

The obtained WWA from the local bakery was soaked in distilled water for 30 mins to remove unwanted particles floating on the surface of the sediment. The unwanted particles were removed by decantation, while the pure wetted waste wood ash (PWWWA) was obtained through filtration. The filtrate was discarded, and the residual PWWWA was oven-dried at 110 °C in an oven until a constant weight is achieved. The dried WWA was calcined in a programmable muffle furnace at 1100 °C for 3 h, the calcined waste wood ash (CWWA) was allowed to cool to room temperature for 48 h, before characterization.

Characterization of the CWWA and WWA was carried out using scanning electron microscopy (SEM) to determine the wide range of surface topography and composition illustrated in Fig. 1. Fourier
Table 2
FTIR sample spectrum analysis of CWWA.

| SN | Wavelength (cm\(^{-1}\)) | Transmittance (%) | Bonding Functional groups |
|----|---------------------------|-------------------|--------------------------|
| 1  | 760.4 to 1077.2           | 71.799 to 69.142  | C-Cl, CO\(_2\)^2-, N-H wagging and twisting, O = C = O bending vibration |
| 2  | 1148.0 to 1636.3          | 69.142 to 81.972  | C=C, C = C, C = N, O = C = O, CHO, C=C, C=N, O = C = O of low energy, O-H, and O–Ca-O bending vibration |
| 3  | 2928.0 to 3272.0          | 83.297 to 74.128  | O-H bending structure, O = O, and N=O |

Table 3
BET-adsorption, XRD analysis, and Hammett indicator value of calcined CWWA.

| Catalyst | N\(_2\)-AA(m\(^2\)g\(^{-1}\)) | TPV(cm\(^3\)g\(^{-1}\)) | %CaO | BS (\(\mu\)mole.g\(^{-1}\)) | TBS | BSD (\(\mu\)mole/m\(^2\)) |
|----------|--------------------------------|--------------------------|-------|-----------------------------|-----|---------------------------|
| CWWA     | 1.10                          | 0.0045                   | 62.83 | 47                           | 162 | 209                        |
|          |                                |                          |       | 400-BS<500                   |     | 190.00                     |

N\(_2\)-AA= nitrogen adsorption analysis, TPV = Total pore volume, BS = Basic site, TBS = Total basic site, BSD = Basic site density.

Transforms infrared spectroscopy (FTIR) to determine the chemical bonds and the angle of resolution within the wavelength band displayed in Fig. 2. The spectra give rise to the shape of the CWWA and WWA as an individual molecular pattern that can be used to screen and scan the powders for different transmittances as explained in Table 2. X-ray diffraction analysis (XRD) was used to quantitative and qualitatively estimated the elemental compositions in the sample. BET isothermal adsorption to determine the surface area, the pore volume, and the total basic density of the powder, while Hammett indicator was used to determine the basic strength of the CWWA displayed in Table 3.

Biodiesel production

Esterification: acid treatment

The acid value of the blended oil was determined to be 4.10 mg KOH/ g oil (%FFA = 2.05), acid value reduction was carried out through the esterification process using a 250 ml three-necked reactor in a temperature control hot plate with a magnetic stirrer. The reaction temperatures were varied between 50 and 90 °C at a reaction time of 40–80 min with H\(_2\)SO\(_4\) concentration of 1.0–3.0% (v/v).
Table 4

| Reaction temperature (°C) | Reaction time(min) | H₂SO₄ conc.(% v/v) | Acid value(mg KOH/ g oil) |
|---------------------------|--------------------|--------------------|--------------------------|
| -                         | -                  | -                  | 4.10                     |
| 50                        | 40                 | 1.0                | 3.70                     |
| 60                        | 50                 | 1.5                | 3.40                     |
| 70                        | 60                 | 2.0                | 2.80                     |
| 80                        | 70                 | 2.5                | 2.50                     |
| 90                        | 80                 | 3.0                | 2.02                     |

A total of five experiments (Table 4) were performed to establish a perfect process condition for the lowest acid value of 2.02 mg KOH/g oil (%FFA = 1.01). A known amount of acid concentration was mixed with methanol in the ratio of blended oil/methanol ratio, the reaction was allowed till completion at a specific temperature and time. At the end of the reaction, the resulting mixture was allowed to settle for phase separation. The bottom layer was removed which contained impurity, H₂SO₄, and excess methanol, while the top layer (esterified oil) was subjected to a transesterification reaction process after washing to remove the leftover acid.

Transesterification: heterogeneous base treatment (calcined waste wood ash)

Synthesis of biodiesel from the esterified oil with the lowest acid value was carried out as follows: the reaction was carried out in a three-necked batch reactor flask. 120 ml of esterified oil was preheated at 70 °C for 60 min, 2.0 (wt.%) calcined waste wood ash (CWWA) was added to the flask containing 50 ml methanol (MeOH), the mixture was partially soluble and was homogenized by using a magnetic shaker. The partially soluble preheated esterified oil mixture was adjusted to a reaction temperature of 60 °C for completion of the reaction. At the end of the reaction, the products were left to stand for 24 h to cool and to enhance phase separation. The used CWWA was recovered from biodiesel by decantation, while the methanol phase was separated from biodiesel by separating funnel. The leached CWWA in the biodiesel was removed through washing with the hot mixture of 2.0 g Na₂CO₃ dissolved in 40 ml methanol. The washed mixture was filtered and then washed with distilled water twice before separation over gravity settling. The wet biodiesel was further dried over anhydrous CaCl₂ and filtered to obtain dry biodiesel. The percentage of biodiesel yield was calculated using Eq. (4). These steps were repeated according to the number of experimental runs (16) generated by Hybrid design (HD).

\[ \text{BY (\% wt.)} = \frac{\text{Weight of pure biodiesel obtained}}{\text{Weight of blended oil used}} \times 100 \]  

Here, BY represents the experimental biodiesel yield.

Experimental design for transesterification of biodiesel

For transesterification of esterified blended oil to biodiesel, four variable factors were considered for process modeling and optimization, this includes reaction time, reaction temperature, CWWA weight (wt.%), and methanol/oil molar ratio (CH₃OH/OMR), respectively. A Hybrid Design (HD) with a minimal point design for 3, 4, 6, and 7 factors with 5 levels each were used for experimental design, which produced a total of sixteen experimental runs (16 runs) without repetitions. Table 5 displayed the five-level-four-variable- factors, the units, and the symbol used for the HD design (Table 5).

Statistical analysis of biodiesel production

Statistical analysis of the transesterification process of biodiesel production was carried out based on experimental results and predicted values by the HD with the variables factors as the constraints via fit summary. The model second order was used to establish the relationship between the biodiesel yield and the four-variable factors. Analysis of Variance (ANOVA) was used to test
Table 5
Experimental design variables for transesterification.

| Variables         | Units    | Symbol | Levels |
|-------------------|----------|--------|--------|
| Reaction time     | (min)    |        | −2     | −1     | 0      | 1      | 2      |
| Reaction temp.    | (°C)     |        |        |        |        |        |        |
| CWWA (wt.%)       |          |        |        |        |        |        |        |
| CH₂OH/OMR (vol./vol.) |        |        |        |        |        |        |        |

Table 6
Biodiesel yield, predicted and residual values of transesterification of esterified oil.

| Std | Run | X₁ (min) | X₂ (°C) | X₃ (wt.%) | X₄ (vol./vol.) | BY (%v/v) | PBY (%v/v) | σ   |
|-----|-----|----------|---------|-----------|---------------|-----------|------------|-----|
| 1   | 9   | 0.000    | 0.000   | 0.000     | 1.732         | 92.00     | 92.00      | 0.000 |
| 2   | 4   | 0.000    | 0.000   | 0.000     | −0.269        | 91.00     | 91.00      | 0.000 |
| 3   | 16  | −1.000   | −1.000  | −1.000    | 0.604         | 93.43     | 93.42      | 2.50E-003 |
| 4   | 3   | 1.000    | −1.000  | −1.000    | 0.604         | 93.57     | 93.58      | −2.50E-003 |
| 5   | 15  | −1.000   | 1.000   | −1.000    | 0.604         | 95.17     | 95.18      | −2.50E-003 |
| 6   | 10  | 1.000    | 1.000   | −1.000    | 0.604         | 96.62     | 96.62      | 2.50E-003 |
| 7   | 11  | −1.000   | −1.000  | 1.000     | 0.604         | 92.68     | 92.68      | −2.50E-003 |
| 8   | 2   | 1.000    | −1.000  | 1.000     | 0.604         | 94.42     | 94.42      | 2.50E-003 |
| 9   | 13  | −1.000   | 1.000   | 1.000     | 0.604         | 95.82     | 95.82      | 2.50E-003 |
| 10  | 1   | 1.000    | 1.000   | 1.000     | 0.604         | 98.82     | 98.82      | −2.50E-003 |
| 11  | 5   | 1.518    | 0.000   | 0.000     | −1.050        | 98.76     | 98.75      | 0.001 |
| 12  | 14  | −1.518   | 0.000   | 0.000     | −1.050        | 96.00     | 96.00      | 0.000 |
| 13  | 7   | 0.000    | 1.518   | 0.000     | −1.050        | 99.14     | 99.15      | −0.001 |
| 14  | 6   | 0.000    | −1.518  | 0.000     | −1.050        | 91.22     | 91.20      | 0.000 |
| 15  | 12  | 0.000    | 0.000   | 1.518     | −1.050        | 96.59     | 96.60      | −0.001 |
| 16  | 8   | 0.000    | 0.000   | −1.518    | −1.050        | 88.64     | 88.64      | 0.000 |

σ = residual value, X₁ = reaction time, X₂ = reaction temperature, X₃ = catalyst weight, X₄ = methanol/oil molar ratio, BY = biodiesel yield, PBY = predicted biodiesel yield.

the model significance and estimate the fit of the model depicted in Table 7. The contour and the three-dimensional plots were used to determine the interactive effects of the linear variables on the biodiesel yield as illustrated in Figs. 3 and 4. The probability value (p-value) and factor value (F-value), the variance inflation factor (VIF), and the degree of freedom (df) were also elucidated for model significance. The R-square coefficient of determination (R²), the R-square predicted coefficient of determination (R²ₚₑ₀ᵈ), and the R-square adjusted coefficient of determination (R²ₐ₀ᵈ), were estimated arithmetically and used to check the model suitability. The final model correlation relationship that expressed vividly the response for every change of variables concerning optimum yield was presented in terms of the second-order differential equation expressed as in Eq. (5).

The final equation in terms of coded factors

\[ \text{BY(\%wt.)} = +90.87 + 0.84X₁ + 1.94X₂ + 1.19X₃ - 0.33X₄ + 0.32X₁X₂ + 0.39X₁X₃ \\
- 0.071X₁X₄ + 0.34X₂X₃ - 0.65X₂X₄ - 1.36X₃X₄ + 2.40X₁² + 1.45X₂² \\
+ 0.34X₃² + 0.57X₄² \]  

(5)

All the experiments were duplicated and the optimum predicted yield was validated in triplicate. The optimum yield (Table 8) was validated in triplicate, and average biodiesel yield was obtained.

Quality of the biodiesel

The quality of the product was examined by determining the properties of the biodiesel such as cetane number, higher heating value, density, viscosity, moisture content, peroxide, saponification, and
Table 7
Anova and test of significant for biodiesel statistical analysis.

| Source | Sum of Squares | df | Mean Square | F Value | Prob > F |
|--------|----------------|----|-------------|---------|----------|
| Model  | 140.63         | 14 | 10.04       | 2.009E+005 | 0.0017   |
| $X_1$  | 8.79           | 1  | 8.79        | 1.759E+005 | 0.0015   |
| $X_2$  | 47.25          | 1  | 47.25       | 9.449E+005 | 0.0007   |
| $X_3$  | 17.90          | 1  | 17.90       | 3.579E+005 | 0.0011   |
| $X_4$  | 1.37           | 1  | 1.37        | 27.438.83  | 0.0038   |
| $X_1^2$| 35.29          | 1  | 35.29       | 7.059E+005 | 0.0008   |
| $X_2^2$| 12.79          | 1  | 12.79       | 2.557E+005 | 0.0013   |
| $X_3^2$| 0.70           | 1  | 0.70        | 13.943.38  | 0.0054   |
| $X_4^2$| 1.64           | 1  | 1.64        | 32.763.02  | 0.0035   |
| $X_1X_2$| 0.81          | 1  | 0.81        | 16,129.00  | 0.0050   |
| $X_1X_3$| 1.23           | 1  | 1.23        | 24,649.00  | 0.0041   |
| $X_1X_4$| 0.040          | 1  | 0.040       | 797.23    | 0.0225   |
| $X_2X_3$| 0.94           | 1  | 0.94        | 18,769.00  | 0.0046   |
| $X_2X_4$| 3.39           | 1  | 3.39        | 67,676.35  | 0.0024   |
| $X_3X_4$| 1.86           | 1  | 1.86        | 2.973E+005 | 0.0012   |
| Residual| 5.000E-005  | 1  | 5.00E-005   |         |          |
| Cor Total| 140.63       | 15 |             |         |          |

Fits statistics

| Std. Dev. | R-Squared | Adj R-Squared | Pred R-Squared | Adeq Precision |
|-----------|-----------|---------------|----------------|----------------|
| 7.071E-003| 0.9999    | 0.9997        | 0.9998         | 1535.084       |

![Predicted vs Experimental biodiesel plots](image1)

![Quadratic model plot](image2)

Fig. 3. Model plots.
Fig. 4. 3-dimensional plots.
iodine value were determined via [22], and the values were compared with biodiesel recommended standards [23,24] (Table 9).

**Performance estimation of CWVA**

The strength of the catalyst developed was tested through catalyst recyclability and reusability test. The CWVA was recycled at the end reaction, because of the impurity at the surface of the regenerated catalyst; the washing with alcohol to remove impurity was carried out before being centrifuged at 4000 rpm for 25 min, and then filtered to obtain wet CWVA. The wet CWVA was then oven-dried at 150 °C for 120 min, allowed to cool at room temperature before reusing. The degeneration in catalyst strength was established by graphical plot, with a reduction in 5th and 6th cycles. Henceforward, the catalyst reusability test was stopped at the 4th cycle (Fig. 5).

**Analysis of spent catalyst**

Analysis of the spent catalyst was studied after reusability and recyclability to depict its morphological characteristic by BET and the effect of reaction on the CaO by XRD. Observation showed that the $S_{\text{BET}}$ (Surface area) and the total pore volume increased as the reusability continued, while the basic site (BS), the total basic site (TBS), and the basic site density (BSD) of the spent catalyst decreased (Table 10). This is an indication that catalyst recyclability, purification, and reusability decreased the basic strength of the catalyst during the reaction process. The main aim of catalyst regeneration is targeted at restoring the high surface area and high dispersal of the element in the catalysts [26]. Furthermore, the effect of leaching within the reactor can easily lead to coverage of active surface sites hereby decreasing the biodiesel yield. The results displayed by XRD analysis alongside with the BET displayed that the percentage decrease to (25.43%) in CaO derived from CWVA

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**Table 8**

Optimum selected condition for biodiesel yield.

| Number | X1 (min) | X2(deg. C) | X3(wt. %) | X4 (vol/vol) | Biodiesel Yield (% wt) | Desirability |
|--------|----------|------------|-----------|--------------|------------------------|--------------|
| 1      | 0.89     | 0.92       | 0.39      | –0.78        | 99.1498                | 1.000        |
| 2      | 0.99     | 0.98       | 0.35      | –0.46        | 99.1499                | 1.000        |
| 3      | 0.95     | 0.93       | 0.38      | –0.66        | 99.1502                | 1.000        |
| 4      | 1.00     | 1.00       | 0.79      | 0.24         | 99.071                 | 0.992        |
| 5      | –1.00    | 0.94       | 0.46      | –1.00        | 97.898                 | 0.881        |
| 6      | –1.00    | 1.00       | 0.34      | –0.94        | 97.7156                | 0.864        |
| 7      | –0.99    | 1.00       | 1.00      | –0.07        | 97.1149                | 0.806        |
| 8      | –1.00    | 1.00       | –0.10     | –1.00        | 96.7254                | 0.764        |
| 9      | –1.00    | 0.27       | 1.00      | –1.00        | 96.5662                | 0.754        |
| 10     | 0.99     | –1.00      | –1.00     | 1.00         | 94.5515                | 0.562        |

**Table 9**

Quality of mixed oil and biodiesel.

| Parameter          | Mixed oil         | Biodiesel | astm d6751 | en 14,214 |
|--------------------|-------------------|-----------|------------|-----------|
| Color® 27 oc       | yellowish-brown   | brownish  | –          | –         |
| Density (kg/m³) @ 25 °C | 880               | 860       | –          | 860–900   |
| Viscosity @ 40 °C (mm²/s) | 3.35             | 2.02      | 1.9–6.0   | 3.5–5.0   |
| Moisture content (%) | 0.001             | <0.001    | <0.03     | 0.02      |
| %FFA (as oleic acid) | 1.01              | 0.21      | 0.40 max   | 0.25 max  |
| Acid value (mg KOH/g oil) | 2.02             | 0.42      | 0.80 max   | 0.50 max  |
| Iodine value (g I₂/100 g oil) | 98.70          | 77.24     | –         | 120 max   |
| Saponification value (mg KOH/g oil) | 214.30         | 120.20    | –         | –         |
| Peroxide value (meq O₂/kg oil) | 1.40            | 1.20      | –         | 12.85     |
| HHV (MJ/kg)        | 39.16             | 43.34     | –         | –         |
| Cetane number      | 49.56             | 74.33     | 57 min    | 51 min    |
| API                | 29.30             | 33.03     | 39.95 max | –         |
| Diesel index       | 54.94             | 89.35     | 50.4 min  | –         |
was very high after reusability and regeneration. It was observed that the percentage of CaO decreased from 62.83% to 25.43%.

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**Declaration of Competing Interests**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper. This work receives no fund from University, Private organization, or Government body.

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**Table 10**

Characteristic of a spent CWWA.

| Catalyst | N$_2$-AA (m$^2$g$^{-1}$) | TPV (cm$^3$g$^{-1}$) | %CaO | BS (μmole.g$^{-1}$) | TBS | BSD (μmole/m$^2$) |
|----------|--------------------------|----------------------|-------|---------------------|-----|-------------------|
| CWWA     | 1.40                     | 0.0080               | 25.43 | -                   | 86.00 | 86.00 | 61.43            |

N$_2$-AA = nitrogen adsorption analysis, TPV = Total pore volume, BS = Basic site, TBS = Total basic site, BSD = Basic site density.
Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.mex.2020.101188.

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