Evaluation of La-Doped CaO Derived from Cockle Shells for Photodegradation of POME

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Abstract
Photocatalysis has merged to be one of the most promising technology in wastewater remediation. However, the application of photocatalysis in treating palm oil mill effluent (POME) is still limited. Many researches were conducted to explore simple and cost-effective alternatives to replace TiO₂ for various industrial purposes. Therefore, the aim of this study is to synthesize and characterize lanthanum doped calcium oxide (La/CaO) as photocatalyst as well as to evaluate the performance of these photocatalysts in the degradation of POME. The photocatalyst used in this study was converted from cockle shells to transform into calcium oxide (CaO) through calcination process. The CaO produced was doped with 1 wt%, 3 wt%, and 5 wt% of lanthanum (La) using wet impregnation method to enhance its photocatalytic activity. The photocatalysts were characterised using X-ray Diffraction (XRD), Fourier Transform Infrared Spectroscopy (FTIR), Brunauer-Emmett-Teller (BET), Scanning Electron Microscopy (SEM), Energy-Dispersion X-ray (EDX), and Inductively Coupled Plasma Mass Spectrometry (ICPMS). Then, this photocatalyst was performed on POME under UVC in a batch system by using different La/CaO at optimum catalyst dosage of 3.0 g/L. Through this research, it was found that the POME degradation through photocatalytic reaction was increased with the incorporation of La where 3 wt% La/CaO showed the highest POME degradation compared to others. This is due to the larger BET surface area that provides more active sites resulted from the incorporation of La. The findings of this study imply that the contaminants in POME can be reduced by utilizing CaO derived from cockle shells. Copyright © 2019 BCREC Group. All rights reserved

Keywords: Calcium Oxide; Cockle Shell; Palm Oil Mill Effluent; Photodegradation

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1. Introduction
In Malaysia, palm oil industry is one of the leading sectors and it contributes significantly to our economic growth. Being one of the world’s largest palm oil producer, the industry discharged approximately between 48 to 72 million tonnes and 49 to 74 million tonnes of POME in 2013 and 2014, respectively [1]. Freely discharged of POME can bring severe impacts to the environment and increase the likelihood of affecting the aquatic life and decreasing the water quality because it contains high level of pollutants such as suspended solid, residual oil and grease, chemical oxygen demand (COD), and biochemical oxygen demand (BOD). At presence, numerous conventional methods have been used in treating POME such as anaerobic digestion [2-5], membrane technology [6-8], and adsorp-
Photocatalysis has shown potentials as one of the treatment methods used in wastewater remediation. This is due to its ability to diminish the pollutants in the wastewater compared to other methods [14,15]. The highly reactive hydroxyl radicals generated from heterogeneous photocatalysis can completely oxidize the organic matter contained in the aqueous system with existing of semiconductor photocatalyst. Titanium oxide (TiO$_2$) has been used extensively as a photocatalyst in photocatalytic process due to a great potential as a sustainable treatment technology to destruct the contaminants in wastewater [16-18]. However, many researches have been conducted to explore simple, cost-effective and eco-friendly alternatives to replace TiO$_2$ for various industrial purposes such as zinc oxide (ZnO) [19-21], cadmium sulphide (CdS) [22,23], cerium oxide (CeO$_2$) [24,25], and others.

Recently, calcium-based catalysts such as CaO have attracted special attention due to their low cost with high mechanical stability which made it interesting for removal pollutant matter in wastewater [26,27]. CaO has been studied and used as the photocatalyst in the treatment/degradation of Rhodamine 6G [28,29], methylene blue [30,31], congo red dye [32], violet GL2B azo dye [33], ammonia [27], and indigo carmine dye [34]. Most of the preparation method to synthesize CaO are from chemical methods. Therefore, in this study CaO generated from cockle shell waste is used as alternative and environment friendly photocatalyst. In addition, this can turn the annually generated from cockle shell waste is used as alternative and environment friendly photocatalyst.

2. Materials and Methods

2.1 Materials

Cockle shells were collected from local market at Kuantan, Pahang. Pre-treated POME was obtained from pond facultative B at palm oil mill based in Felda Lepar Hilir 3 located at Pahang, Malaysia. Lanthanum nitrate (La(NO$_3$)$_3$) with 95% purity was purchased from Sigma-Aldrich was used to be incorporated as a dopant to CaO formed from waste cockle shells. Deionized water was used in all the experiments that was obtained from Ultra Pure Water Sysyem (Model: Elix 5UV).

2.2 Preparation of Photocatalyst

The CaO from cockle shells was prepared based on Jones et al. [38] and Shariffuddin et al. [39]. Firstly, the cockle shells were cleaned thoroughly using hot water to remove dirt and impurities. After that, the cockle shells were crushed into small particles before they were grinded into powder form using Retsch ZM 200 Grinder. The shells powder was then sieved over a mesh screen to obtain fine powder of less than 63 μm. The fine powder obtained was transformed into CaO catalyst via calcination method at 900 °C for 5 hours in a furnace.

Next, the synthesized CaO was doped with 1 wt%, 3 wt%, and 5 wt% of La using wet impregnation method. In a typical preparation, 10 g of CaO was suspended in 40 mL of deionized water. To this suspension, 10 mL La(NO$_3$)$_3$ aqueous solution of desired concentration was added. The resulted slurry was stirred for 2 hours without heat and then evaporated to dryness. The precipitate formed was dried overnight in an oven at 120 °C. Lastly, the precipitate was calcined at 400 °C in furnace with a heating rate of 5 °C/min for 3 hours.

2.3 Characterization of Photocatalyst

XRD of the photocatalyst were obtained through Bruker D8 Advance X-ray diffractometer operating with a 40 kV in the 2θ range of 20° to 80° with a 0.02° step size with a 1s step time. BET was evaluated based on N$_2$ adsorption-desorption isotherm at -196°C using Micromeretics ASAP 2020 system. Samples were
degassed at 110°C overnight before the N₂ adsorption measurements. Then, specific surface areas were calculated from the N₂ adsorption data based on BET method using P/P₀ values between 0.01-0.3.

SEM equipped with EDX were performed to capture the morphology images and to find relative composition and abundance of elements present on the samples using Swift ED3000 in Hitachi TM3030 Plus. The samples were sputter-coated with platinum to prevent charging during the analysis. The functional groups of sample powders were characterized using Nicolet iS5 Model FTIR (Spectrum100) with spectrum collected between the wavenumber of 400 and 4000 cm⁻¹ using OMNIC software. Elemental analysis for La and Ca was also determined by ICP-MS (Agilent technologies ICP-MS Model 7700).

2.4 Photocatalytic Degradation of POME

Prior starting the degradation experiments, POME sample collected from local palm oil mill was filtered to remove the suspended solids. To ensure the degradation of POME was due to photocatalytic degradation, two sets of control experiments which were photolysis and adsorption were conducted using CaO and 3 wt% La/CaO. Firstly, the suspension was magnetically stirred in the dark to ensure the establishment of the adsorption/desorption equilibrium between photocatalyst and POME at room temperature. The second control experiment was done on evaluating the degradation of POME under UVC without presence of photocatalyst to check the ability of photodegrade organic compounds by the UVC.

This was then followed by photocatalytic degradation of POME in the presence of UVC irradiation and CaO photocatalyst. Firstly, 3 g/L of CaO was added into 200 mL of diluted POME solution and the suspension was stirred continuously at 300 rpm to keep the mixture in suspension during the photodegradation. After 30 min, the mixture was irradiated with UVC. During the first hour, 2.0 mL of sample was withdrawn from the reactor vessel at every 15 min interval. While in the next 3 hours, samples were withdrawn at every 30 min interval. The photodegradation experiments were conducted under limited oxygen condition (no oxygen gas was bubbled into the solution). Prior to analysis, the samples were centrifuged to separate the photocatalyst from POME solution. The COD concentration in POME was then examined using Hach DRB-200 reactor. All digestion solutions would be subjected to the digestion chamber at 150 °C for 120 min. The experiment was repeated using La/CaO photocatalyst of different La loadings at 1 wt%, 3 wt%, and 5 wt%.

3. Results and Discussion

3.1 X-ray Diffraction Studies

The phase structure of CaO and La/CaO (1 wt%, 3 wt%, and 5 wt%) were investigated by XRD and shown in Figure 1. The XRD pattern for CaO in Figure 1(a) shows peaks approximately at 2θ = 32°, 38°, 54°, 64°, and 68°. All intensified and narrowed peaks were dominated by calcium oxide in cubic phase with apparently no calcium carbonate retained in the form of calcite [40,41]. This is due to the polymorphic transformation of aragonite polymorph to calcium oxide following the calcination process at a temperature of 900 °C for 5 hours. The XRD patterns of 1 wt%, 3 wt%, and 5 wt% La/CaO (Figure 1(b), (c) and (d)) show peaks of hexagonal portlandite. The existence of portlandite in the samples were due to the conversion of CaO into Ca(OH)₂ during wet impregnation process [42].

The minor peaks indicated by 2θ = 55°, 63°, and 65° in 1 wt% La-Ca corresponded to CaO polymorph [43]. Similarly, the XRD patterns of La/CaO for 3 wt% and 5 wt% show the same generation of peaks that are related to CaO polymorph. This shows that, CaO still exist in La/CaO photocatalyst after doping with La elements. The peak intensity of CaO and La/CaO was also determined by ICP-MS (Agilent technologies ICP-MS Model 7700).

![Figure 1. XRD patterns indicating the change in crystal structure and chemical composition of (a) CaO calcined at 900 °C, (b) 1 wt% La/CaO, (c) 3 wt% La/CaO and (d) 5 wt% La/CaO.](image-url)
are different because additional of La can reduced their degree of crystallinity as reported by Shaveisi et al. [27]. Besides, the diffraction peaks of La/CaO in Figure 1 shows the peaks shifted to lowest 2θ due to lattice expansion [44] which indicated the incorporation of La ion into CaO lattice structure. Meanwhile, the small peak approximately around 2θ = 30° in 1 wt%, 3 wt%, and 5 wt% La/CaO were detected to be lanthanum oxide [45,46]. The XRD patterns for all the La/CaO samples show the absence of lanthanum carbonate as shown in Figure 1(b), (c), and (d).

3.2 FTIR Spectra of Photocatalyst

The functional group of photocatalysts produced were analysed using FTIR spectroscopy. Figure 2 shows the FTIR spectra of CaO and La/CaO in the range of 500-3900 cm⁻¹. The bottom spectrum as shown in Figure 2(a) represented the CaO was calcined at 900 °C. Only a weak band at around 3630 cm⁻¹ was detected and attributed to hydroxyl stretching vibration which appeared resulting from small amount of water adsorbed in the crystalline CaO [47,48]. There are no other impurity peaks related to other groups are observed in this spectrum.

The spectra obtained for samples after the calcined shell was incorporated with La ions at 1 wt%, 3 wt%, and 5 wt% were as shown in Figure 2(b), (c), and (d). The sharp intense peaks at 3640 cm⁻¹ in principle can be assigned to −OH groups which attached to CaO to form Ca(OH)₂ as reported by Shaveisi and Sharifnia [27], and Kaur et al. [42]. This −OH groups was formed due to CaO was wet impregnated with La(NO₃)₃ solution which showed similar finding with XRD analysis in Figure 1(b), (c), and (d). The two new bands appeared at 1450 cm⁻¹ and 858 cm⁻¹ were corresponded to the peaks of bending vibration of O−Ca−O groups and stretching of C−O bond from carbonate [49]. These bands were emerged possibly due to the reaction of photocatalysts with CO₂ in the air during sample preparation [50,51].

3.3 Surface Area and Crystallite Size

The crystallite size of catalysts can be determined from full width of half maximum (FWHM) of the most intense peaks using Scherrer’s equation as indicated in Equation (1):

\[ d = \frac{K \lambda}{\beta \cos \theta} \]  

Where \( d \) is the average of crystal size; \( K \) is constant values which related to the shape of the polycrystals (0.9); \( \lambda \) is the X-ray wavelength of Cu Kα radiation (1.5418 Å); \( \beta \) is FWHM of the XRD peak value and \( \theta \) is the maximum of Bragg’s angle in radians [52,53]. By using these data, the estimated crystallite sizes for CaO and La/CaO are shown in Table 1. The crystallite size, \( d \) for CaO sample (507.44 nm) was found to be larger than La/CaO. Meanwhile, the crystallite size of

| Sample          | 2θ (deg) | β (rad) | d (nm) | BET Specific Surface Area, m².g⁻¹ | Average Pore Diameter (nm) |
|-----------------|----------|---------|--------|---------------------------------|-----------------------------|
| CaO             | 37.57    | 0.0027  | 507.44 | 1.12                            | 69.87                       |
| 1wt% La/CaO     | 34.24    | 0.0030  | 483.82 | 3.88                            | 96.69                       |
| 3wt% La/CaO     | 34.41    | 0.0037  | 388.50 | 4.08                            | 85.22                       |
| 5wt% La/CaO     | 34.41    | 0.0035  | 411.61 | 6.73                            | 67.02                       |

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La/CaO was decreased with increasing of La content up to 3% and then increased. Previous research by Shakir et al. [20], explained this observation due to expansion of ZnO lattice caused by radius of La$^{3+}$ that was larger than Zn$^{2+}$. Besides, the distortion of lattice structure in CaO sample occurred when a foreign impurity dopant was introduced. As a result, the nucleation and subsequent growth rate had decreased because the size of Ca and La ions are different [54].

Table 1 also shows BET specific surface area of the photocatalysts. Based on the results obtained, the BET specific surface area has increased upon the incorporation of lanthanum. This result was consistent with the outcomes of crystallite size obtained from XRD analysis and was in agreement with the literature described by Yang et al. [55]. The increase in BET surface area when increasing dopant percentage can be explained due to the deposited of La that has collectively contributed to the overall BET specific surface area of the CaO photocatalyst. The higher specific surface area may benefit the photocatalytic activity due to enhanced adsorption of irradiation photons and organic pollutants. Nesic et al. [36] stated that the increasing of La content in TiO would affect pore diameter by decreasing the diameter. This finding is similar with the result in Table 1 which shows the decreasing of pore diameter with the increasing of La from 1 wt% (96.69 nm), 3 wt% (85.22 nm) and 5 wt% (67.02 nm).

3.4 Morphology and Composition Analysis

Figure 3 illustrates the surface morphology of undoped (CaO) and La/CaO samples studied using SEM. It can be clearly observed from the close-up SEM micrographs that there are

![Figure 3](image-url)
Figure 4. SEM micrographs and EDX spectrum of (a) CaO, (b) 1 wt% La/CaO, (c) 3 wt% La/CaO and (d) 5 wt% La/CaO.
structural changes that occur after the incorporation of dopant element in catalyst. The cockle shell powder calcined at 900 °C is irregular in shape and bonded together as aggregates [56]. Based on Figure 3(a), there are some relief lines on the particle surfaces due to high temperature calcination process. When the calcined shell is further impregnated in lanthanum nitrate solution and activated at 400 °C, the morphologies La/CaO photocatalyst of catalyst are demonstrated in Figure 3(b), (c), and (d). The SEM micrograph results show a honeycomb-like porous network produced due to the water molecular (H₂O) in form of gaseous is released from decomposition of Ca(OH)₂ which created large number of voids [57]. Several small particles within the grains were discovered and the crystallites were observed to have no uniform shape. Again, these findings were in good agreement with XRD studies which revealed that the crystallinity of photocatalyst declined when increasing dopant percentage.

Figure 3 also represents the EDX spectra of CaO, 1 wt% La/CaO, 3 wt% La/CaO, and 5 wt% La/CaO. Based on the EDX analysis as shown in Figure 3 (a), it indicates that calcium element is allocated with highest weight percentage, which is 48.19%, followed by oxygen and carbon. Trace amount of carbon detected in the sample denoted that lime has formed with an unconverted calcium carbonate residual. Since the quantity of carbon is comparably small, the calcium carbonate peak in the form of calcite has been flattening out in the XRD pattern as shown in Figure 1(a). Meanwhile, to prevent the electrons bombarded during SEM analysis from staying embedded in place where they hit the non-conductive calcined shell, a thin layer of metal was coated on the sample surface [58]. This explained the fact that platinum element presented in the samples as a coating material.

On the other hand, the EDX spectra of La/CaO samples as shown in Figure 3 (b), (c), and (d) were observed to exhibit similar elements in CaO, with addition of La element in small amount. The amount of La element presented in 1 wt% La/CaO, 3 wt% La/CaO, and 5 wt% La/CaO were increased to 1.43 wt%, 2.16 wt%, and 3.46 wt%, respectively. The wt% of La in the samples was not the same with the wt% of dopant due to the dissimilar distribution of La spread in the photocatalyst. However, from the EDX spectra line, it can be concluded that La element was successfully impregnated into the crystal structure of CaO particles.

Therefore, to verify the exact amount of the La and Ca (in ppm) for each sample were analysed by using ICP-MS measurements and listed in Table 2. The result shows that La ele-

| Parameter     | CaO          | 1 wt% La/CaO | 3 wt% La/CaO | 5 wt% La/CaO |
|---------------|--------------|--------------|--------------|--------------|
| Calcium (Ca)  | 134077.0     | 404144.1     | 223793.0     | 505884.5     |
| Lanthanum (La)| -            | 4136.7       | 12762.0      | 21285.1      |

**Figure 5.** The experiments of POME degradation for (a) control experiment of adsorption at CaO, photolysis for POME and adsorption at 3 wt% La/CaO, and (b) the C/C graph of organic material in POME for photocatalytic degradation over different lanthanum concentration (photocatalyst = 3.0 g/L, stirring speed = 300 rpm, UVC, time = 240 min).

**Table 2.** The amount of La and Ca in photocatalysts (in ppm) using ICP-MS
Figure 6. SEM micrographs of photocatalyst (a) 1 wt% La/CaO before photocatalytic treatment, (b) 1 wt% La/CaO after photocatalytic treatment (c) 3 wt% La/CaO before photocatalytic treatment, (d) 3 wt% La/CaO after photocatalytic treatment, (e) 5 wt% La/CaO before photocatalytic treatment, (f) 5 wt% La/CaO after photocatalytic treatment.
ment was present into the catalyst after impregnated. This shows the agreement between the EDX results and ICP-MS measurement on the successful doping of La into CaO through wet impregnation method. From the ICP-MS results, it is shown that the amount of La increase as expected.

3.5 Effect of Lanthanum Concentration

Before starting the experiment, blank run experiments were conducted to ensure the degradation of POME is due to photocatalytic degradation. Therefore, Figure 4 (a) and (c) shows the adsorption process by using CaO and 3 wt% La/CaO where the maximum POME degradation after 30 min was about 10.9 % and 13.73 %, respectively. After that, the reaction was remained constant for the rest of reaction time (240 min). Therefore, adsorption can be regarded as negligible since the degradation is in minimal percentage presumably a minor for POME removal mechanism. This maximum removal of POME during the adsorption (dark condition) within 30 min has interpreted that the experiments was necessity to start stirred in the dark for at least 30 min before photocatalytic reaction undertaken next remaining of process time. The photolysis result in Figure 4 (b) shows no degradation occurred during the reaction time studied. Therefore, the removal of organic substances in POME by photolysis is insignificant. Hence, photolysis alone will not able to degrade the POME.

In this study, La was chosen as a dopant materials due to its ability to enhance the

![EDX spectrum](image1)

![EDX spectrum](image2)

![EDX spectrum](image3)

**Figure 7.** EDX spectrum of (a) 1 wt% La/CaO photocatalyst after photocatalytic treatment, (b) 3 wt% La/CaO photocatalyst after photocatalytic treatment, (c) 5 wt% La/CaO photocatalyst after photocatalytic treatment.
photocatalyst properties as reported in other studies by Nesic et al. [59] and Thi & Lee [60]. Figure 5(b) shows the degradation of POME by using different type of La loadings. The removal of organic substances in POME by using CaO shows the lowest degradation compared to other photocatalysts. Subsequently, when increasing the percentage of La in the photocatalyst, the removal of pollutants in POME through photocatalysis has increased significantly. It can be clearly seen that when CaO was doped with 1 wt%, 3 wt%, and 5 wt% of La, the COD concentration was reduced by 43.4 %, 54.0 %, and 33.9 %, respectively. The increase in the degradation with the increment of La loadings is possibly due to the lattice deformation that occurred after dopant ions (La$^{3+}$) were introduced which could hinder the recombination of electron-hole pairs and therefore enhance the photoactivity [61-63]. In addition, the incorporation of La ions have increased the BET specific surface area as shown in Table 1, thus allowing an extensive amount of organic compounds adsorbed on the photocatalyst surface. On the other hand, the reduction of photocatalytic degradation for 5 wt% La/CaO may be due to the excess doping metal into photocatalyst which created recombination center for photogenerated charge carriers [37].

3.6 Characterization of Spent Photocatalysts

Figure 6 demonstrates the morphology of 1 wt%, 3 wt%, and 5 wt% of La/CaO photocatalyst respectively before and after the photocatalytic treatment. By comparing the morphology of spent catalyst to the freshly prepared catalyst in Figure 6, it was found that the spent catalyst consists of irregular shapes and various sizes distribution which are in tiny aggregates. Meantime, the microstructure of spent photocatalyst appeared to be less porous due to the adsorption of organic pollutants in POME onto the active sites of photocatalyst.

The elemental composition of 1 wt%, 3 wt%, and 5 wt% La/CaO spent catalysts are shown in Figure 6 (a), (b), and (c), respectively. The presence of additional magnesium (Mg), aluminium (Al), chlorine (Cl), and potassium...
(K) elements in the spent doped photocatalysts indicate that the degradation intermediates of POME were attached to the catalyst surface during photoreaction. This is parallel with statement by Rupani et al. [64] which described POME as a viscous brownish liquid that contains substantial amount of minerals such as Mg, Al, K, and so forth. Moreover, chlorine which act as a highly efficient disinfectant may have added into POME to poison the disease-causing pathogens and also came from the freshwater used in the processing of palm oil. This explained the fact that chlorine was detected in the EDX spectrum in small amount, which is less than 0.1 wt%.

Comparative FTIR spectra of the catalysts before and after photocatalytic treatment with POME are shown in Figure 7. It was observed that the FTIR spectra of 1 wt%, 3 wt%, and 5 wt% of spent La/CaO in Figure 7 (a), (b), and (c), retained almost similar patterns for the functional groups (C–O and O–H) as the freshly prepared La/CaO photocatalyst even after the photoreaction. However, the FTIR spectra for spent and fresh CaO (Figure 7 (a)), shows there are changes in terms of the patterns and functional groups. This shows that by doping the CaO with La, the photocatalyst are much more stable compared to the undoped (CaO). In addition, comparative spectra for the doped fresh and spent photocatalysts reveal that the significant changes in these sample powders are the changes in the intensity of the functional groups (C–O and O–H). It can be observed that, these functional groups decreased after the photocatalytic reactions. The decreased in intensity of the functional groups indicates that these groups are being photo-dissolved in order to allow a greater amount of oxygen from the lattice to participate in the degradation of POME to form OH radicals where this mechanism also is known as Mars Van Krevelen mechanism. Similar results were reported by Ali et al. [62] on the degradation of methylene blue using ZnO as a photocatalyst.

4. Conclusions

Four samples of photocatalysts were successfully synthesized using CaO derived from cockle shells by adding La as dopant to treat POME through photocatalytic reaction. The results showed that POME was successfully degraded with 3 wt% La/CaO functioned most effectively, followed by 5 wt%, 1 wt% of La/CaO, and CaO photocatalysts. It was observed that the degradation of POME increased with the increment of La loading where the maximum degradation occurred at 3 wt% La/CaO. The results from XRD concluded that the incorporation of La into CaO showed lower crystallinity and crystallite size, while from BET results showed that higher surface area was produced with the increment of La as dopant. These results showed that the incorporation of La provided more active sites for photocatalytic reaction. The emerging of this finding is believed to be able to recover and recycle cockle shells in huge amount while providing an environment-friendly solution to wastewater remediation system. From this study, although La/CaO can be used as an alternative material for photocatalyst, the degradation rate is not as efficient compared to TiO₂. Therefore, further study will be needed to improve the performance of La/CaO as a photocatalyst.

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