


 Cite this: *RSC Adv.*, 2024, 14, 36327

Synthesis of vanillin *via* oxidation of kenaf stalks in the presence of CeO₂: tuning the catalytic behaviour of CeO₂ *via* nanostructure morphology†

 Anita Ramli,^a Nur Akila Syakida Idayu Khairul Anuar,^a Normawati Mohamad Yunus^b and Alina Rahayu Mohamed^c

Different CeO₂ nanostructures were synthesized using a hydrothermal method and treated with alkaline NaOH, followed by drying at 120 °C for 16 h and calcined at 400 °C for the direct oxidation of kenaf stalks to vanillin under microwave irradiation. The catalysts were characterized for their physicochemical properties using XRD, BET, Raman spectroscopy, TPR, TPO, and XPS. All synthesized CeO₂ nanostructures show diffraction peaks corresponding to the formation of cubic fluorite, which agrees with Raman spectra of the F_{2g} mode. The N₂ adsorption–desorption isotherms showed that all catalysts possess a type IV isotherm, indicating a mesoporous structure. TPR and TPO analyses display formation peaks corresponding to surface-to-bulk reducibility and the oxidized oxygen ratio, which is responsible for the redox properties of ceria nanostructures. The XPS analysis of CeO₂ nanostructures proved that Ce exists in the Ce³⁺ and Ce⁴⁺ oxidation states. All catalysts were tested for direct oxidation of kenaf stalks under microwave irradiation with the highest vanillin yield obtained by the CeO₂-Nps-400 heterogeneous catalyst at 3.84%, whereas 4.66% vanillin was produced using 2 N NaOH as a homogeneous catalyst.

 Received 12th August 2024
 Accepted 16th October 2024

DOI: 10.1039/d4ra05833j

rsc.li/rsc-advances

Introduction

Vanillin (4-hydroxy-3-methoxy benzaldehyde) is a vital commercial compound with a wide range of applications in the food and perfumery industries, and it is naturally extracted from dried pods of the vanilla plant.¹ The commercial vanillin market is served by three major sources: natural vanilla from vanilla beans (approximately 0.2%); vanillin produced from petroleum-based intermediates, especially guaiacol, which accounts for 85% of the world supply; while the remaining 15% is produced from lignin derived from the wood pulping process.² Synthetic vanillin cannot be labelled as natural but rather as a “synthetic” or “artificial” vanilla flavor.³ However, according to EU regulations, if the process and base material are natural, such as lignin, phenolic stilbenes, isoeugenol,

eugenol, ferulic acid and aromatic acids, the vanillin produced can be considered ‘natural vanilla aroma’.⁴ Therefore, lignin could be used as a natural precursor to meet the growing demand for natural vanillin.

Various studies have been conducted to produce vanillin from lignocellulosic biomass under a catalyzed medium. Deng *et al.*⁵ reported 5.3% vanillin yield obtained from wet aerobic oxidation of lignin from cornstalks under 2 M NaOH as a solution with 5 wt% of LaCo_{0.8}Cu_{0.2}O₃ as a catalyst. Another study by Qu *et al.*⁶ reported that 5% vanillin yield was obtained through direct oxidation of Japanese cedar biomass using microwave irradiation in a 2 N NaOH alkaline medium with peroxide as an oxidant and CuO as a catalyst. Ceria is a well-known catalyst or catalyst promoter for numerous industrial redox processes. The interconversion between the +4 and +3 oxidation states of cerium makes CeO₂ a suitable candidate for redox catalysts.⁷ Both surface and bulk oxygen vacancies in CeO₂ are suitable sites for adsorption. Hence, besides being useful as a support, CeO₂ can actively participate in chemical reactions and provide an active site as an oxygen carrier.⁸

The oxidation of commercial organosolv lignin was carried out using a batch-type high-pressure autoclave reactor at 170–185 °C for 24 h in a MeOH solution medium under continuous stirring with 1–10 bars purged of oxygen and Pd/CeO₂ as a catalyst. The highest vanillin yield of 5.2% was achieved.⁹ However, Rawat *et al.*¹⁰ reported that selective production of vanillin from oxidative depolymerization of alkali lignin

^aHICoE Centre of Biofuel and Biochemical Research (CBBR), Institute of Sustainable Energy & Resources (ISER), Department of Fundamental & Applied Sciences, Universiti Teknologi PETRONAS, Seri Iskandar 32610, Perak, Malaysia. E-mail: anita_ramli@utp.edu.my

^bCentre of Research in Ionic Liquids (CORIL), Institute of Sustainable Energy & Resources (ISER), Department of Fundamental & Applied Sciences, Universiti Teknologi PETRONAS, Seri Iskandar 32610, Perak, Malaysia. E-mail: normaw@utp.edu.my

^cFaculty of Chemical Engineering & Technology, UniMAP, Complex of Academics Jejawi 3, Jejawi, Arau 02600, Perlis, Malaysia. E-mail: alina@unimap.edu.my

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4ra05833j>



catalyzed using MoPO/CeO₂ was carried out with a batch reactor under continuous stirring at 150 °C for 3 h in 2 M NaOH solution with 5 bars of oxygen as an oxidant yielding 9% of vanillin. Our initial work employing Ce and Zr catalysts supported by MgO to oxidize kenaf stalks to produce vanillin directly indicates that CeO₂ is active in vanillin production. The oxidation of lignin from the kenaf stalk using Ce/MgO catalyst under microwave gives a 3.70 yield of vanillin, while bare MgO and Zr/MgO do not give any vanillin product.¹¹ These facts provoked our interest in conducting the present investigation on ceria.

Different shapes of CeO₂ nanostructures give a prevalence of specific crystal facets that display different reactivities in various catalytic processes.¹² By tailoring the shape and size of the particles, certain crystal facets can be exposed, leading to different structural and redox properties, as reported by researchers.^{13–15} Thus, methods that control crystal shape may provide a course to ascertain an insight aspect required to enhance catalytic activity.¹⁶ Theoretically, the presence of particular active facets of the ceria surface affected oxygen vacancy formation and its ability to act as an oxygen donor–acceptor.¹⁷ The oxygen storage/release capacity is capable of explaining the quantitative amount of oxygen that can be substituted by a reducible oxide in the gas phase or with an active metal component. Konsolakis *et al.*¹⁷ and Qiao *et al.*¹⁴ found that the rod-shaped ceria nanostructures prevailing [110] and [100] facets possess a high catalytic conversion of CO oxidation, where they show a consistent trend with the highest oxygen storage capacity (OSC) and surface to bulk ratio compared to nanocubes and nanopolyhedral. Altering the ceria structure generates differences in the surface atomic arrangement and the electronic properties, which also significantly influence the oxygen vacancy formation energy.¹⁸ This research investigated catalytic activity and correlated it with the number of oxygen vacancies and OSC, especially for the oxidation of kenaf stalks to vanillin production. Nevertheless, the reactivity of oxygen vacancies on different crystal facets for vanillin production has rarely been discussed.

The kenaf plant was used as a biomass feedstock. Kenaf is an annual fiber crop plant that is composed of inner core fiber (75–60%) and bast fiber (24–40%)¹⁹ and has a high syringyl to guaiacyl ratio in aromatic composition owing to its lignin characteristic. It is reported that kenaf stalks contain 13–15% lignin.^{20,21} This paper is based on the use of CeO₂ in the oxidation of the lignocellulosic biomass of kenaf to vanillin reported recently. In a previous paper, CeO₂ nanostructures were prepared by varying the calcination temperature from 400 to 600 °C. Catalyst screening was done to choose the ideal calcination temperature based on the performance of vanillin production. It was found that the CeO₂ nanostructure calcined at 400 °C showed a higher production of vanillin due to its high characteristics in textural properties compared to other catalysts.²² The conditions for catalyst screening were 180 °C of microwave heating for 20 min with 10 wt% of a catalyst under 200 W microwave irradiation, and it was found that a calcination temperature of 400 °C gave the highest yield. The chosen catalyst was further utilized to determine the optimum reaction

conditions.²² Our recently published paper shows different active facets on different nanostructures of CeO₂ that have been successfully proven and discussed using TEM and HRTEM. The uniform shape of the CeO₂ nanostructures (nanoparticles, nanorods and nanocubes) from the TEM image was found, while HRTEM images show that the CeO₂ nanostructures are single crystalline in nature.²² Thus, this research discussed the correlation of different CeO₂ nanostructure morphologies for renewable vanillin produced directly from biomass under microwave heating and was further tested for optimized reaction conditions.

Experimental

Materials

Dried kenaf stalks were obtained from the National Kenaf and Tobacco Board, Malaysia. Methanol (CH₃OH), sodium hydroxide pallet (NaOH), ethanol, ethyl acetate, hydrochloric acid (HCl), hydrogen peroxide and vanillin laboratory standard (as standard) were obtained from Aldrich, USA. Sodium hydroxide and cerium hexahydrate (Ce(NO₃)₃·6H₂O) were obtained from Merck, Germany.

Methods

Catalyst preparation. The CeO₂ nanostructures were synthesized following the method reported by Torrente-Murciano *et al.*²³ Typically, to synthesize CeO₂ nanoparticles, 0.6 g of Ce(NO₃)₃·6H₂O was added to 40 mL of 5 M NaOH solution and stirred for 10 min in a PTFE beaker. Then, the mixture was transferred to a Teflon-lined hydrothermal vessel, sealed and heated to 70 °C for 10 h. At the end of the synthesis duration, the hydrothermal vessel was cooled to room temperature before it was unsealed. The solid was filtered and washed several times with deionized water, followed by drying at 120 °C for 16 h and calcination in air at 400 °C for 2 h. The same procedure was repeated to synthesize CeO₂ nanorods except that the hydrothermal temperature was 100 °C. CeO₂ nanocubes were synthesized using the same procedure, except that the concentration of the NaOH solution used was 15 M and the hydrothermal temperature was 180 °C. The synthesized catalyst was denoted as CeO₂-Nps-400 (CeO₂ nanoparticles), CeO₂-Nrs-400 (CeO₂ nanorods) and CeO₂-Ncs-400 (CeO₂ nanocubes).

Catalyst characterization. The crystallinity of the CeO₂ catalysts was determined using the Bruker X-ray diffraction (XRD) model X'Pert3 Powder and Empyrean (PAN Analytical, Billerica, MA, USA). The catalysts were recorded using an X-ray diffractometer system between 2θ value of 20–80° with 227 s per step exposure time and 0.105° per step size. The crystallite size of synthesized CeO₂ nanoparticles was determined according to Scherrer's equation: $D = k\lambda/\beta \cos \theta$. Crystallite size was analyzed using HighScore Plus software (version 3.0), Malvern, UK, with an X-ray wavelength of Cu Kα radiation at $\lambda = 1.54$ Å, where θ is the Bragg angle. β is the full width at half maximum in radians, corresponding to the 2θ value at the (111) plane. The unknown shape factor, k , was assumed to be 0.89, and the reflecting peak at 2θ was chosen for the entire sample.



The surface area and pore size of the catalysts were analyzed using Brunauer–Emmett–Teller (BET) (Micromeritics ASAP 2020, Norcross, GA, USA). The catalysts were degassed at 200 °C for 24 h prior to N₂ adsorption measurement at –77 °C.

The TPR experiment on H₂ consumption and oxygen capacity storage was conducted using a Thermo Scientific TPDRO 1100 with a TCD detector. The samples (0.5 g) were treated by heating from 298 to 1073 K at a rate of 10 K min^{–1} in a flow of 5 vol% H₂ in He. The total gas flow rate was 10 mL (NTP) per min. The TPO experiment was conducted on a Thermo Scientific TPDRO 1100 using a TCD detector. The samples (0.5 g) were treated by heating from 298 to 1073 K at a rate of 10 K min^{–1} in a flow of 5 vol% O₂ in He. The total gas flow rate was 10 mL (NTP) per min.

The Raman spectra of the glasses were collected at room temperature using a Jasco NRS-3300. Raman spectrometer was equipped with a CCD detector (–69 °C) using the 514 nm line of an air-cooled Ar ion laser (Melles Griot), 600 lines per mm grating, 0.1 × 6 mm slit and 100× Olympus objective lens.

Thermo Scientific X-ray photoelectron spectroscopy (XPS) equipment with MgKα X-ray radiation source ($E = 1253.6$ eV) was performed to analyze each element's surface composition and chemical state present in the catalyst structure. The C 1s peak set at 284.6 eV was used as the charge referencing. The atomic concentration ratios in the outer layers of the samples were evaluated from the corresponding XPS area ratios using an effective ionization cross-section of the ejected electrons.²⁴ According to the ratio of the Ce³⁺ ion peak area to that of the total Ce³⁺ and Ce⁴⁺ ion peak areas, the following equations were used to calculate the relative content of Ce³⁺ in the CeO₂-Nps-400 catalyst.²⁵

$$ACe^{3+} = Av' + Au',$$

$$ACe^{4+} = Av + Av'' + Av''' + Au + Au'' + Au'''. \quad (1)$$

$$\text{Relative content of Ce}^{3+} = ACe^{3+}/(ACe^{3+} + ACe^{4+}). \quad (1)$$

The surface chemisorbed oxygen ratio $[R(O_s)]$ can be obtained as follows:²⁶

$$R(O_s) = A(O_s)/[A(O_s) + A(O_b)]. \quad (2)$$

Catalytic oxidation of raw kenaf. Under microwave irradiation, the performance of the synthesized CeO₂ nanostructures was investigated in the direct oxidation of kenaf stalks to vanillin. The dried kenaf stalks (2 g) were soaked for 24 hours in a screw-cap test tube with 20 mL of 0.01 N (normality) NaOH solution before being transferred to a Teflon tube. After that, 15 wt% of catalyst and 1 mL of H₂O₂ were added to the mixture, and the pH recorded was 11.5. The mixture was heated in a microwave (Milestone, MicroSYNTH MA143) at 160–180 °C for 20 minutes after being stirred at 600 rpm for 10 seconds at room temperature using a magnetic stirrer. The liquid was then cooled to room temperature and filtered using filter paper to

eliminate any insoluble materials. The residue was washed twice with a 20 mL solution of 0.01 N NaOH. The filtrate was poured into a glass test tube and sealed. The same procedure was performed by changing the heterogeneous catalyst to a homogeneous sodium hydroxide catalyst. According to the obtained result, 15 wt% was chosen as optimized catalyst loading for further study using 2 N (normality) of sodium hydroxide catalyst. The filtrate was transferred to a glass test tube with a cap. Simultaneously, the catalyst residue was washed several times with ethanol to further investigate catalyst reusability under optimized reaction conditions, followed by a drying process in an oven at 100 °C for 2 h. Concentrated hydrochloric acid (37%) was added to the filtrate in a ratio of 1 : 2, agitated for 15 seconds with a vortex agitator at 5000 rpm, and then centrifuged for 15 minutes at 1000 rpm. The supernatant was then treated with 1 : 1 ethyl acetate. Vanillin and other low-molecular weight molecules were extracted into the organic phase. The mixture was agitated for 60 seconds at 5000 rpm and then centrifuged for 5 minutes at 1000 rpm, separating the mixture into two phases. The upper phase was placed in a vial. At 50 °C and 400 mbar, excess solvent was evaporated using a rotary evaporator.^{11,16}

Product characterization. The product's detailed composition was analyzed using gas chromatography-mass spectrometry as a qualitative analysis (Agilent 7890A) at an ionization voltage of 70 eV on a DB-5MS column (30 m × 0.25 mm i.d.; 0.25 μm film thickness). The oven temperature was programmed at 50 °C for 3 minutes, followed by heating to 300 °C at 8 °C min^{–1} and held for 20 min. The temperature of the injection part was 250 °C, while the temperature of the detector was set at 300 °C. Helium (He) was used as the carrier gas at 1.6 mL min^{–1} with 1 μL injection volume. The split ratio was 1 : 10, and the mass range (m/z) was in the range of 40–600 m/z .⁶

Results and discussions

Catalyst characterization

Fig. 1 shows the XRD patterns of synthesized CeO₂ nanoparticles (CeO₂-Nps-400), nanorods (CeO₂-Ncs-400) and nanocubes (CeO₂-Nrs-400) calcined at 400 °C for 2 h. Multiple sharp

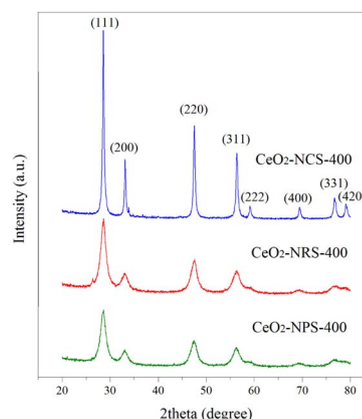


Fig. 1 XRD patterns of CeO₂ nanostructures calcined at 400 °C.



peaks were observed for all catalyst samples, indicating the nanocrystalline nature of CeO₂. The XRD results demonstrated that all prepared CeO₂ nanostructures show the formation of cubic fluorite that is perfectly indexed to ICSD: 81-0792 with different intensities. The diffraction peaks at $2\theta = 28.53, 33.07, 47.46, 56.31, 59.06, 69.38, 76.66, \text{ and } 79.03^\circ$ correspond to the (111), (200), (220), (311), (222), (400), (331) and (420) planes, respectively. These observations agree with those of Lykaki *et al.*,²⁷ who reported that the presence of all diffraction peaks of CeO₂ nanostructures indicates the prominent peak of a pure ceria face-centered cubic fluorite structure (*Fm* $\bar{3}$ *m* symmetry). In general, the results also indicate that the peak intensity for CeO₂-Ncs-400 is higher and has a well-defined sharp peak compared to CeO₂-Nrs-400 and CeO₂-Nps-400, which means an increase in the size and crystallinity of the products.²⁸

Based on the full width at half maximum (FWHM) of the diffraction peak (111), crystallite sizes of CeO₂ nanostructures calcined at 400 °C were estimated *via* the Scherrer equation and are summarized along with textural properties of catalysts (surface area, pore volume and average pore size), as illustrated in Table 1. The obtained nanostructures of CeO₂ in the forms of nanoparticles, nanorods and nanocubes were prepared by adjusting the concentration of sodium hydroxide and hydrothermal reaction temperature.

In particular, the crystallite sizes of CeO₂ are 11.3, 15.0 and 28.7 nm for CeO₂-Nps-400, CeO₂-Nrs-400 and CeO₂-Ncs-400, respectively, while the crystallite size of CeO₂ nanostructures increases in the order of CeO₂-Ncs-400 < CeO₂-Nrs-400 < CeO₂-Nps-400. The large crystallite size of CeO₂-Ncs-400 might be due to the expansion of lattice between nanocubes crystals, which agrees with the report by Hailstone *et al.*²⁹ They also found that the increase in the cluster size was due to lattice expansion ascribed to the decreased amounts of Ce³⁺ at the surface of smaller clusters, which directed the reduced Ce–O electrostatic attraction.

The results showed that CeO₂ nanoparticles have the highest BET surface area, pore volume and average pore size with increasing order of CeO₂-Ncs-400 < CeO₂-Nrs-400 < CeO₂-Nps-400. In addition, the surface areas found in the present study correlate with the crystallinity of the catalysts, as shown in the XRD results. According to Lykaki *et al.*,²⁷ these differences in the distribution of textural properties are linked to different ceria morphologies and can be mainly accounted for by the observed variations in the BET surface area. Moreover, it is noteworthy that the BET surface area follows the reverse order of the crystallite size of CeO₂ (111) phases (Table 1). This shows that the larger the crystallite size, the lower the BET surface area. Fig. 2 shows the N₂ adsorption–desorption isotherms of the nanostructures of CeO. According to the International Union of Pure

and Applied Chemistry (IUPAC) classification, all the synthesized CeO₂ nanostructures resemble type IV isotherm with type H1 desorption hysteresis according to IUPAC classification, which indicates the formation of a well-developed mesoporous structure.³⁰ The hysteresis loop of type H1 is commonly found in highly ordered mesoporous materials, such as MCM-41, MCM-48 and SBA 15.³¹

Raman spectroscopy was used to characterize the oxygen vacancy abundances of ceria's exposed planes on the structural defects. Fig. 3 shows the Raman spectra of the CeO₂ nanostructures catalyst. The spectra are dominated by a strong peak at 463 cm⁻¹ from the F_{2g} mode of the CeO₂ *Fm* $\bar{3}$ *m* cubic fluorite, which agrees with the XRD patterns (Fig. 2). Kainbayev *et al.*³² reported that the observed peak inherent to cerium oxide ranges from around 460 to 463 cm⁻¹, which is in accordance with the F_{2g} mode corresponding to the symmetric vibrations of oxygen ions around Ce⁴⁺ ions in cerium oxide, while the D band observed at 598–615 cm⁻¹ is assigned to structural defects caused by the perturbations in cerium lattice.²⁷ For CeO₂-Nps-400, the Raman band is detected at 610 cm⁻¹ with high intensity compared to CeO₂-Nrs-400 and CeO₂-Ncs-400 catalyst, which is also assigned to the presence of oxygen atom detachment, resulting from Ce⁴⁺ transforming to Ce³⁺ partial reduction.³³ As presented in Table 1, the intensity of the peak and FWHM values is the highest for CeO₂-Nps-400, followed by CeO₂-Nrs-400 and CeO₂-Ncs-400. According to Ndifor *et al.*,³⁴ the FWHM of CeO₂ increased with a decreasing crystallite size and/or a greater concentration of oxygen vacancies. The Raman shifts in the mode corresponded to the particle size effect, lattice constant, and crystallite size of the catalyst.²⁷ Lattice deformation and oxygen vacancies of CeO₂ nanostructures were measured using the A_D/A_{F_{2g}} ratio, following the order of CeO₂-Nps-400 > CeO₂-Nrs-400 > CeO₂-Ncs-400, which interestingly

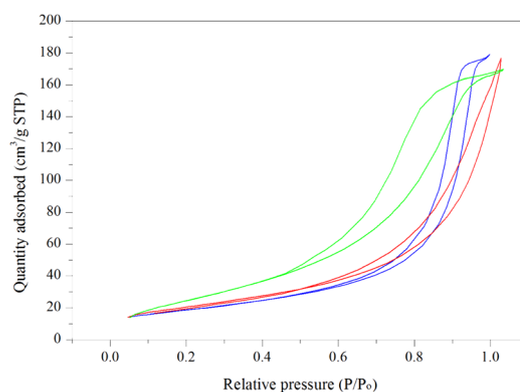


Fig. 2 N₂ adsorption–desorption of CeO₂ nanostructures.

Table 1 Structural and textural properties of CeO₂ nanostructures

Catalyst	S_{BET} (m ² g ⁻¹)	S_{V} (cm ³ g ⁻¹)	S_{P} (nm)	S_{size} (nm)	FWHM (cm ⁻¹)	$A_{\text{D}}/A_{\text{F}_{2\text{g}}}$
CeO ₂ -Nps-400	66.1	0.275	27.5	11.3	50	0.136
CeO ₂ -Nrs-400	57.8	0.145	24.0	15.0	46	0.113
CeO ₂ -Ncs-400	22.5	0.134	17.1	28.7	19	0.065



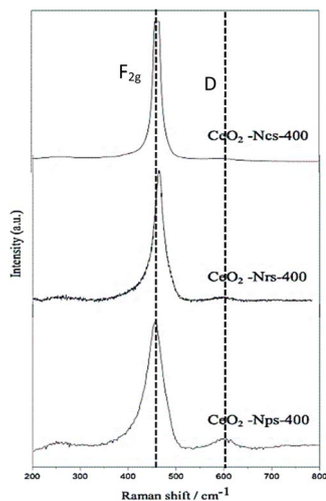


Fig. 3 Raman spectra corresponding to different CeO₂ nanostructures.

agrees with the reducibility order of TPR analysis for CeO₂ nanostructures.

As depicted in Fig. 4, hydrogen temperature programmed reduction (H₂-TPR) was further used to study the reducibility of these ceria nanostructures. However, temperature programmed oxidation (O₂-TPO) was chosen for further characterization to determine the oxidation behaviors of ceria nanostructures, as shown in Fig. 5. TP profiles of ceria particles reveal a reduction peak at about 350–500 °C assigned to consumption of surface oxygen (O_s),³⁵ where the total consumption in this region is well correlated with the polycrystalline of CeO₂ surface area.³⁶ On the contrary, the reduction peaks above 800 °C are attributed to the reduction of oxygen species in bulk CeO₂ (O_b),³⁷ resulting from diffusion of O out of the CeO₂ bulk to the CeO₂ surface, where it

can react with adsorbed or gas phase H species.³⁸ Consequently, measurements of OSC were determined from lower temperature reduction regions, as it mainly reflects on surface oxygen.

As observed in Fig. 4, the O_s peak is smaller for ceria nanocubes compared to nanorods and nanoparticles. This is attributed to the smaller amount of easily reduced oxygen species available in the cubic samples, which can also be closely related to the lowest lattice deformation and oxygen vacancies ($A_D/A_{F_{2g}}$) and exposed surface facets. A few studies have also found that exploring the role of exposed surface facets using ceria nanostructures of different morphologies revealed differences in reduction behavior.^{39,40} These results agree with a study by Lykaki *et al.*,²⁷ which reported that ceria nanocubes exhibit a smaller O_s peak compared with nanorods and nanopolyhedra, indicating that the lower reducibility of the cube-shaped sample possesses the smallest population of weakly bound oxygen species.^{39,40} The reduction of ceria through a temperature-programmed reduction in hydrogen (H₂-TPR) also indicates a superior behavior of CeO₂-Nps-400 compared to that of CeO₂-Nrs-400 and CeO₂-Ncs-400, as shown in Table 2. This is evidenced by the anticipation of the onset of surface Ce⁴⁺ reduction and by the increase in reduction degree at low temperatures, which can be associated with easier oxygen removal from exposed [100] and [111] surfaces and the higher density of surface defects present in CeO₂-Nps-400.⁴¹ The results in Table 2 demonstrate that H₂ at lower temperatures removed surface oxygen as the lattice strain increased, suggesting that catalytic activity was associated with the reversible oxygen storage capacity of the catalysts.³⁸ Consumption of H₂ resulting from surface reduction occurred at lower temperatures over nanoparticles and nanorods than over the nanocubes, corroborating the ease of reducibility of CeO₂ surface facets exhibited by each nanostructure.

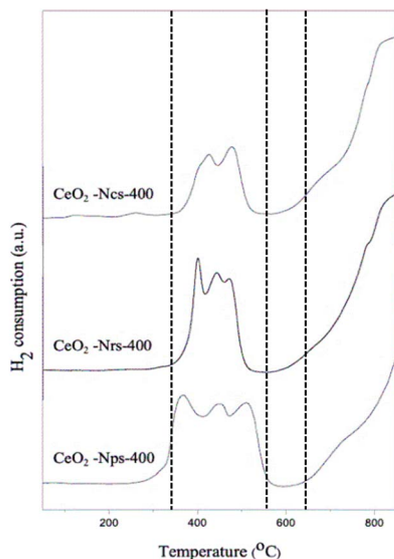


Fig. 4 H₂-TPR profiles of CeO₂ nanostructures.

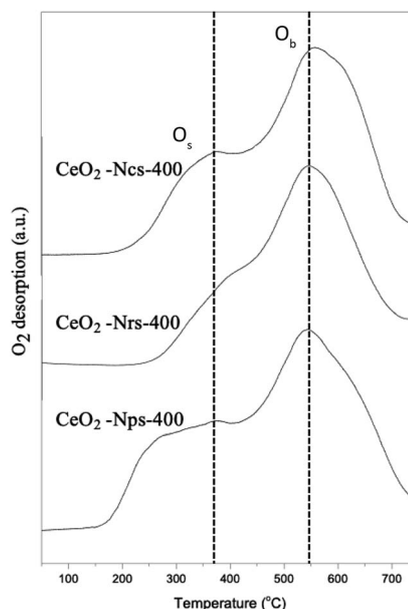


Fig. 5 O₂-TPO profiles of CeO₂ nanostructures.



Differences in the surface area of ceria can also influence the overall H₂-TPR profiles by changing the order of reactivity between the CeO₂ nanostructures. During H₂-TPR experiments, the reduction of Ce⁴⁺ to Ce³⁺ indicated oxygen removal and oxygen vacancy formation. In contrast, O₂-TPO experiments were thus conducted to present the oxygen uptake and storage expedition charge transfer of ligand (O 2p) to metal (Ce 4f) by the primary photoionization process. The lower binding energy of v° and u° spin-orbit doublet located at 889.7 and 908.6 eV associated capacity.⁴² The TPO profile exhibits two main O₂ consumption areas with broad peaks that contributed to the oxidation of CeO₂ species in sequence from low to high temperatures at around 350 and 540 °C for all CeO₂ nanostructured catalysts. As shown in Fig. 5, it is proposed that the first stage (200–450 °C) might be attributed to oxygen uptake as a surface lattice to oxygen (O_s) and the second (450–700 °C) could be attributed to oxygen uptake as bulk oxygen (O_b).⁴³ From the result, it can be observed clearly that the intensity of O₂ consumption was increased in the order of CeO₂-Ncs-400 < CeO₂-Nrs-400 < CeO₂-Nps-400, which revealed a good agreement with the measured reduction of CeO₂ nanostructures in TPR analysis. Due to the highly oxidized particle-shaped surface of CeO₂, CeO₂-Nps-400 exhibits the highest population of weakly bound oxygen.^{44,45}

Subsequently, the samples were further characterized using XPS analysis for a comparative assessment of each CeO₂ nanostructure, which gives insight into the impact of morphology on elemental chemical states and surface composition, as depicted in Fig. 6. The XPS analysis on CeO₂ nanostructure was performed to obtain more profound insight into the valence state of the chemical compound and chemical composition contained in the sample. Fig. 6 shows the Ce 3d core level spectra of different CeO₂ nanostructure catalysts. The XPS survey spectra indicated that the samples included no components besides C, Ce, and O, confirming the exceptional chemical purity of the CeO₂ nanoparticles (Fig. 10), which is consistent with the conclusion made by Bortamuly *et al.*⁴³ More specifically, the

spectra of Ce 3d revealed that Ce exists in the +4 and +3 oxidation states for all samples. This could be determined as Ce 3d_{5/2,3/2} spin-orbit doublet peaks, splitting at around 27 eV. Based on Burroughs *et al.*⁴⁴ studies, the Ce 3d spectrum can be represented as u and v doublet peaks, indicating the 3d_{3/2} and 3d_{5/2} spin-orbit states, respectively. The v''' and u''' doublets at 906.2 and 925.2 eV with 19 eV separation correspond to the primary photoemission O 2p orbital to an empty core level of Ce 4f orbital with Ce 3d⁹4f¹ O 2p⁵ and Ce 3d⁹4f² O 2p⁴ final state shake-down satellite features, respectively.^{23,39,45} According to Anandan *et al.*,⁴⁶ the satellite features are due to the photoionization from Ce³⁺ with Ce 3d⁹4f² O 2p⁵ final state. However, the v' and u' spin-orbits at 898.2 and 915.2 eV indicate the

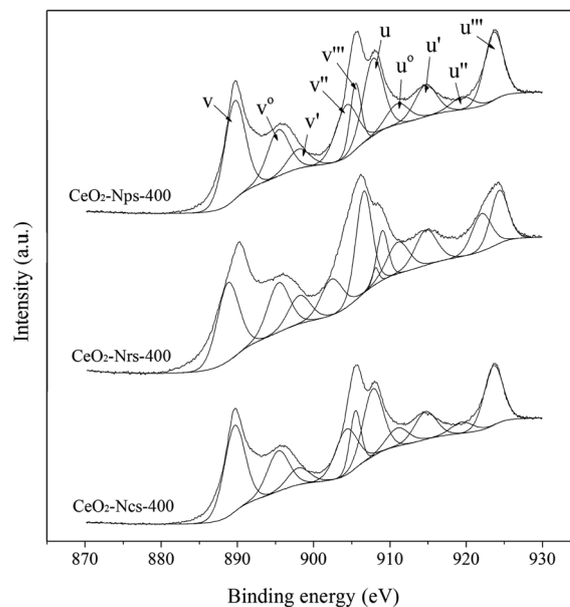


Fig. 6 Ce 3d core level XPS spectra of the CeO₂ nanostructure catalyst.

Table 2 Redox properties of CeO₂ nanostructures

Catalyst	H ₂ consumption (μmol H ₂ per g)			O _b	Total	O _s /O _b	Peak temperature (°C)	
	Peak 1	Peak 2	Peak 3				O _s	O _b
CeO ₂ -Nps-400	7219.7	6262.0	6400.8	11 112.3	30 994.8	1.79	418–484	847
CeO ₂ -Nrs-400	5348.8	4603.0	4233.1	9293.8	23 478.7	1.53	401–492	842
CeO ₂ -Ncs-400	6535.0	6101.2	—	20 791.2	835.3	0.61	386–509	863

Catalyst	O ₂ consumption (μmol O ₂ per g)			Total	O _s /O _b	Peak temperature (°C)	
	O _s	O _b	O _s			O _b	
CeO ₂ -Nps-400	4035.0	4453.0	8488.00	0.91	378	546	
CeO ₂ -Nrs-400	4248.5	4882.5	9131.0	0.87	401	553	
CeO ₂ -Ncs-400	2362.4	3676.8	6039.2	0.64	372	556	



photoemission of Ce^{3+} from $Ce\ 3d^{9}4f^{1}\ O\ 2p^{6}$ final state.^{47,48} Stetsovych *et al.*⁴⁹ reported that the Ce^{3+} phase is buried under the CeO_2 layer and the fully oxidized CeO_2 layer is grown on top of the interfacial phase.

The XPS scan of the O 1s core level for different CeO_2 nanostructure catalysts is depicted in Fig. 7. The spectra of the samples were deconvoluted into three peaks at binding energies of around 536.2, 538.1, and 540.3 eV. According to ref. 50, the lowest binding energy peak at 536.2 eV is attributed to the lattice oxygen in CeO_2 ; the binding energy peak appearing at 538.1 eV is ascribed to the surface oxygen, while the observed binding energy at 540.3 eV is due to the hydroxyl on the surface of CeO_2 . According to Zhang *et al.*,⁵¹ surface oxygen content influences the catalytic performance of CeO_2 materials. Moreover, the surface oxygen vacancy adsorbs and activates oxygen molecules to produce adsorbed oxygen species. This mechanism promotes the redox properties of CeO_2 .⁵² The obtained deconvoluted peaks differ in binding energy values around 4–5 eV compared to the literature findings. This result is consistent with the findings reported by Sudarsanam *et al.*⁵³ and Hillary *et al.*,⁵⁴ which state that the formation peak at a lower binding energy corresponds to the lattice oxygen of the metal oxide phases (O^{2-}). However, the peak occurring at a higher binding energy is ascribed to surface chemisorbed oxygen, including adsorbed oxygen (O^-/O_2^{2-}), adsorbed water, hydroxyl (OH^-), and carbonate (CO_3^{2-}) species.

The calculated values of relative Ce^{3+} content and surface chemisorbed oxygen ratio of the CeO_2 nanostructure are shown in Table 3. Because oxygen vacancy is generated when Ce^{4+} is reduced to Ce^{3+} , it is crucial to calculate the relative Ce^{3+} content and chemisorption oxygen ratio to evaluate the generation of oxygen vacancy on the catalyst surface. Both the relative contents of Ce^{3+} and the surface chemisorbed oxygen ratio of CeO_2 display an increasing pattern in the order of CeO_2 -Nps-400 > CeO_2 -Nrs-400 > CeO_2 -Ncs-400. Overall, the nanostructured ceria's direct oxidation intrinsic catalytic activity increases as

the amount of Ce^{3+} increases, and the presence of Ce^{3+} ions demonstrates the formation on the surface of the non-stoichiometric CeO_2 .⁵⁵ The Ce^{3+} ions accompanying the oxygen vacancies play a crucial role in the oxidation activity, which is also associated with the surface chemisorption of oxygen and activation of oxygen surface vacancies on the material and the migration of oxygen toward the surface material, as confirmed by observing the catalytic performance of the CeO_2 -Nps-400 catalyst that recorded high production of vanillin compared to CeO_2 -Nrs-400 and CeO_2 -Ncs-400 catalysts.⁵⁶ The Ce^{3+} and O_s species on the CeO_2 surface play major roles in controlling catalytic oxidation for vanillin production, which are directly related to oxygen vacancies and the oxygen storage capacity of the catalyst. Based on the summarized data in Table 3, the results regarding the chemisorption oxygen of CeO_2 -Nps-400 agree with those of the relative Ce^{3+} content.

Catalytic oxidation of raw kenaf

Effects on catalyst loading. Fig. 8 shows the vanillin yield obtained on different catalyst loadings (5 wt%, 10 wt%, 15 wt% and 20 wt%) produced with 2 g of dried kenaf stalks heated in a 300 W microwave power output for 20 minutes at 170 °C. Optimization of reaction duration, temperature and microwave power output was discussed in a previous paper.²² The study discovers that vanillin production increases as power output increases from 100 to 300 W due to a notable amount of thermal energy converted to microwave energy and the intensity of the electromagnetic radiation increases simultaneously. These conditions result in a narrow distribution for producing aromatic monomers by selectively cleaving the phenyl side chain of C_{α} - C_{β} . Specifically, molecular motion and diffusion coefficient increased considerably when the temperature increased rapidly, while the mixture's boiling point reached more quickly. In addition, the use of CeO_2 -Nps-400 as a catalyst in this reaction gradually increases the reaction rate. Moreover, it has a more significant number of specific surface-active sites, initiating the partial oxidation of lignin components in biomass. In addition to the context mentioned earlier, the Ce species acts as an electron acceptor in the rate-limiting step and facilitates the conversion of phenolate anions into phenoxy radicals, which catalyze vanillin production.⁵⁷

The outcome in Fig. 8 demonstrates the increased number of yields with the increased amount of catalyst loading for 15 wt% with a vanillin yield of 3.84% (CeO_2 -Nps-400), 3.07% (CeO_2 -Nrs-400) and 2.96% (CeO_2 -Ncs-400). This result agrees with the study conducted by Prado *et al.*,⁵⁸ which states that vanillin yield increases from 0.015 to 0.22% when catalyst loading increases

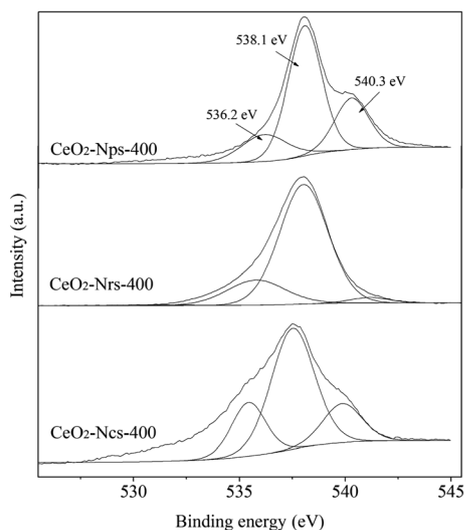


Fig. 7 O 1s core level XPS spectra of the CeO_2 nanostructure catalyst.

Table 3 Relative the Ce^{3+} content and surface chemisorption oxygen of CeO_2 nanostructures

Sample	Relative to Ce^{3+} content	Surface chemisorption oxygen
CeO_2 -Nps-400	0.197	0.139
CeO_2 -Nrs-400	0.126	0.111
CeO_2 -Ncs-400	0.119	0.098



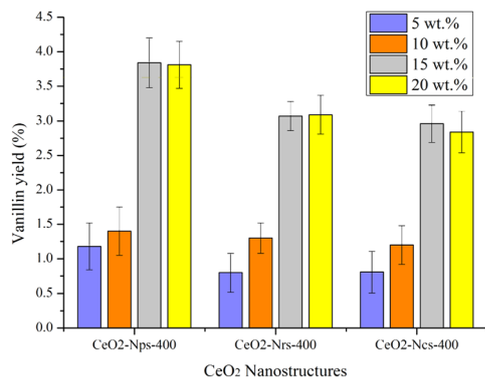


Fig. 8 Reaction conditions: 2 g of dried kenaf stalks, 20 mL of 0.01 N NaOH solution, and 1 mL of H₂O₂ under 300 W microwave heating at 170 °C for 20 min.

from 1 to 20%. This is attributed to the higher total contact frequency of catalysts with lignin, which is also consistent with the study conducted by Qu *et al.*⁶ They stated that the higher the catalyst loading, the higher the conversion of kenaf stalks. According to Chen *et al.*,¹⁸ it was found that the catalysts had a greater overall contact frequency with lignin. It is also believed that higher catalyst loading made more active sites accessible to break down lignin into aromatic aldehyde compounds. Based on the surface area of the CeO₂ nanostructures catalyst, CeO₂-Nps-400 possesses a high surface area among all CeO₂ nanostructures. Due to its high surface area, CeO₂-Nps-400 provided many active surfaces for the oxidation reaction of lignin to occur compared to other CeO₂ nanostructured catalysts. Furthermore, the surface oxygen provided by CeO₂-Nps-400 promoted oxidation as the catalyst loading increased. This showed a considerable effect on the vanillin yield, greatly improving the selectivity of the conversion of lignin to vanillin. According to Zhang *et al.*,⁵⁹ materials in the nanoscale possess a surfeit of oxygen defects and improved catalytic performance as the crystallite decreases. In fact, the fine-tuning of the ceria with textural and defect structural characteristics exhibited by nanoparticles ceria as a catalyst proves its strong dependence between support morphology and catalytic activity, as reported in previous studies.^{60,61} However, as the catalyst loading increased to 20 wt%, the vanillin produced a slight decrease. These findings agreed with those of Behling *et al.*, who found that increased loading of Co₃O₄ from 1 to 2 wt% led to increased yield of vanillin from around 13% to 19%. However, the vanillin yield shows a slightly decreased pattern when the catalyst loading further increases to 20 wt%, which might be due to the over-oxidation of vanillin that is completely formed with available oxidizing species in the media. Excessive loading of the catalyst might also result in increased side reactions due to the overcrowding of the active sites.⁶²

Conversion through a chemical transformation in heterogeneous catalysts is usually generated on specific sites on the catalyst surface, and it is generally referred to as active sites and intrinsic activity on a per site basis. The intrinsic activity of catalysts can normalize with different parameters that normally

provide information on certain features of a catalyst, such as overall mass (catalyst loading), the mass of a specific component or surface area. In this study, various masses of catalyst loadings were used to show the normalized rate of CeO₂ nanostructure on the production of vanillin. Direct oxidation tests from kenaf stalks for each CeO₂ nanostructure catalyst were performed from 5 wt% to 20 wt% of catalyst loading with a constant 20 min reaction duration, 170 °C temperature, and 300 W microwave power output. Different catalyst loadings were used as parameters to observe the normalized pattern for each catalyst to allow for a direct comparison of the reaction rate and its effect on vanillin production on CeO₂ nanostructures. The slope trends of CeO₂-Nps-400 depicted in Fig. 9 are 0.256, 0.094, 0.246 and 0.178 for 5, 10, 15 and 20 wt%, respectively, where it shows a decreasing pattern from 5 to 10 wt% and an increasing pattern at loading 15 wt%, followed by a decreasing normalized value for loading greater than 15 wt%. This indicates that the production of vanillin is independent of 10 and 20 wt% of catalyst loading, which can be explained by the mass transfer limited of the catalyst, where it showed a decreasing order to the factor of three and two, respectively.⁶² In contrast, normalized for 15 wt% shows an increasing normalized rate approximately to the factor of three. Meanwhile, CeO₂-Nrs-400 and CeO₂-Ncs-400 displayed the same pattern as CeO₂-Nps-400.⁶³ It could be concluded that catalyst loadings of 5 and 15 wt% show a significant normalized value to the production of vanillin through the direct oxidation test and the least significant for catalyst loadings at 10 and 20 wt%, respectively, as illustrated by the transition slopes in Fig. 10.

Comparison of catalytic activity using heterogeneous and homogeneous catalyst. The previous section showed that vanillin was detected in the liquid produced from the direct oxidation of the kenaf stalks in the presence of all CeO₂ nanostructured catalysts. Specifically, CeO₂-Nps-400 shows the highest yield of vanillin production from the direct oxidation of kenaf stalks compared to other CeO₂ nanostructures. Using the optimized process parameters, this research further studied catalytic activity in vanillin production using a different

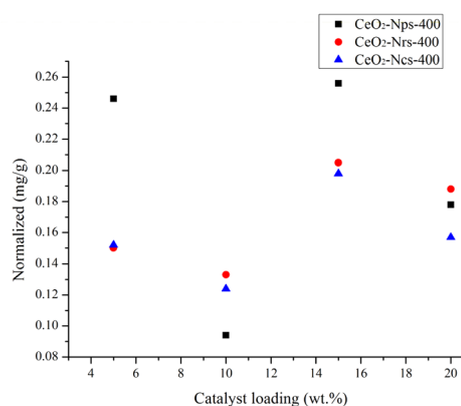


Fig. 9 Comparison of the normalized rate of the production of vanillin to the loading of each type of catalyst from 5–20 wt%.



physical catalyst that is homogeneous (2 N of sodium hydroxide) and heterogeneous ($\text{CeO}_2\text{-Nps-400}$) for the production of vanillin through the direct oxidation of kenaf stalks. Based on previous optimization, the maximum vanillin yield was achieved at 170 °C and 20 min under 300 W of microwave power output in the presence of 15 wt% of catalyst loading.²² Therefore, this parameter is used to study the catalytic activity of the chosen $\text{CeO}_2\text{-Nps-400}$ heterogeneous catalyst and 2 N of sodium hydroxide (NaOH), as shown in Fig. 10. This study aimed to compare their catalytic performance and yield on vanillin production.

The yield of vanillin obtained for the $\text{CeO}_2\text{-Nps-400}$ heterogeneous catalyst was 3.84%, while 4.66% vanillin was produced using 2 N (normality) NaOH as a homogeneous catalyst. This is noteworthy because vanillin production was directly from biomass without extracting the lignin from the hemicellulose and cellulose components. This result agrees with a study conducted by Khairul *et al.*¹¹ in which a 4.45% vanillin yield was obtained utilizing 2 N of NaOH as a homogeneous catalyst. In comparison, 2.90% vanillin was produced using 30Ce/MgO-48 as a heterogeneous catalyst at 170 °C for 20 min. They also reported that although heterogeneous catalysts possess high selectivity and conversion percentages for vanillin production, there are drawbacks throughout the reaction process due to surface saturation/deactivation in the formation of alcohols and aromatic compounds. On the contrary, the homogeneous catalyst is considered a flexible catalyst that allows it to freely contact the CAr-O bonds without high steric limitations because the reaction occurs solely in a liquid phase that also reacts as a reaction medium simultaneously, as reported in previous studies.^{11,64}

Catalyst reusability. For more insight into catalyst stability, the heterogeneous CeO_2 nanoparticle catalyst was used to undergo further optimization to study catalyst reusability for vanillin production. According to Takale *et al.*,⁶⁵ nanoparticles (NPs) have emerged as an appealing option for catalysis in organic syntheses. This is due to their expansive reactive

surfaces, which offer greater potential for selectivity. Consequently, reactions can occur efficiently under mild conditions. Additionally, the ease of isolating products and the ability to reuse the catalyst are supplementary benefits of nanocatalyzed multicomponent reactions.⁶⁶ Thus, in this part, the reusability of the $\text{CeO}_2\text{-Nps-400}$ catalyst was determined by the repeated use of the catalyst for the direct oxidation of kenaf stalks to vanillin production three times with the new reactants and under optimized reaction conditions. From the results, the $\text{CeO}_2\text{-Nps-400}$ catalyst is highly stable and useable up to three times with a slight drop in sustained activity to produce vanillin compared to the yield from the first use (Fig. 11). The vanillin yield recorded for the first use was 3.84% and 3.48% for the second cycle, which is a 0.36% difference in yield values. For the third cycle, the vanillin yield showed no significant difference in terms of yield, with 3.38% of vanillin. The results obtained in this study were similar to the other findings reported for MgO-based Ce catalysts. A study showed that the Ce/MgO catalyst could be used for up to three cycles with a slightly significant loss in vanillin production.¹¹ The difference in vanillin yield between the first and second cycles might be due to a saturated catalyst surface caused by biomass side products, along with a significant lignin-like oligomer from vanillin oxidative polymerization of vanillin, which could explain the small reduction in vanillin and a slight decrease in vanillin synthesis.^{65,67,68} Moreover, the gradual loss in catalytic activity through consecutive reaction cycles could be due to surface poisoning and pore filling by other particles (impurities) that participated during the reaction, where the active sites were blocked by the catalyst for subsequent usage.⁶⁹

The catalyst's physicochemical properties for pre-reaction and post-reaction were compared to identify the structural integrity of the catalyst as the active oxygen sites on the catalyst's surface are exposed and play the primary role during the direct oxidation reaction. The $\text{CeO}_2\text{-Nps-400}$ was used for up to three cycles under 170 °C microwave heating for 20 min at 300 W power output. The catalyst was treated for reusability studies, and the third cycle catalyst was used for further characterization using XRD and Raman analyses. Fig. 12 displays the (a) XRD and Raman (b) analyses of $\text{CeO}_2\text{-Nps-400}$ for pre-reaction and post-reaction after the three cycles used. As

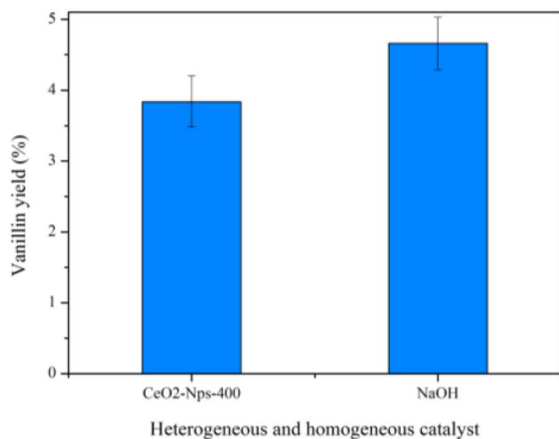


Fig. 10 Reaction conditions: 2 g of dried kenaf stalks, 20 mL of 0.01 N NaOH solution, 15 wt% of catalyst, and 1 mL of H_2O_2 under 300 W of microwave heating at 170 °C for 20 min.

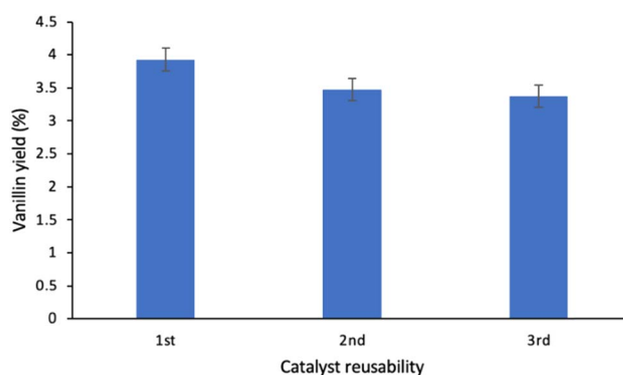


Fig. 11 Catalyst reusability of $\text{CeO}_2\text{-Nps-400}$.



observed in the XRD figure of CeO₂-Nps-400, there are no significant changes in the formation of peaks with identical peaks of the ceria before and after the use of the catalyst. However, there are slight differences in diffraction intensity as the post-reaction catalyst showed a slight decrease in diffraction intensity peak compared to a pre-reaction catalyst for diffraction peak at higher $2\theta = 30^\circ$. This was proved by comparing the crystallite size, where the post-reaction catalyst was 10.1 nm with a 1.2 nm difference from the pre-reaction catalyst (11.3 nm). There is a slight peak shift to the right with a shifted value of 2θ around 0.1 to 0.01° for the post-reaction catalyst compared to the original diffraction peaks before the reaction. The different intensities of the diffraction peaks indicate that there are structural changes between the crystal lattices. This might be caused by the agglomeration of the particles at a higher product composition owing to uneven particle distribution, which is evident by the differences in the peak intensity and crystallite size of the catalysts.⁶⁸

The same goes for Raman analysis, where the absorption peak intensity and peak shift for post-reaction can be detected at the F_{2g} and D modes with slightly different pre-reaction catalysts. The defect oxygen (D) Raman for post-reaction demonstrated an intensity decrease compared to pre-reaction CeO₂-Nps-400 at 601 cm⁻¹. In contrast, there are no significant differences in Raman intensity for the observed peak inherent to cerium oxide at 463 cm⁻¹. The Ce oxidation state reduces as oxygen is released and oxygen vacancies are formed. As the concentration of oxygen vacancies increased, the non-stoichiometry F_{2g} in the lattice becomes asymmetric and the shoulder at band ~600 cm⁻¹, which is associated with defect species, broadens and shades at high temperatures as the lattice becomes more mobile for the post-reaction catalyst due to the regeneration of lattice Ce–O species to replenish oxygen vacancies simultaneously in the reduced state of metal oxide.^{69,70}

Correlation between OSC of the CeO₂ nanostructures and direct oxidation of kenaf stalks to vanillin. It is noteworthy that the yield of the catalytic conversion of vanillin from direct oxidation of kenaf stalks for CeO₂ nanostructures follows the same trend as BET, relative oxygen vacancy concentration and lower temperature OSC. The results presented in Table 3

confirm that the surface of CeO₂ nanoparticles is saturated with Ce³⁺ species. This concentration leads to the greatest amount of lattice distortion and the strongest tendency for a shift to Ce³⁺ from Ce⁴⁺ with the formation of more oxygen vacancies.⁴⁵ The quantitative analysis to determine the OSC value of the synthesized samples uses hydrogen consumption based on the low-temperature peak area in the H₂-TPR profiles. The lower temperature OSC can be used to study the amount of oxygen species available in the oxides (surfaces) that contribute to the reoxidation of the catalyst by a continuous redox cycle.⁷¹ Table 3 shows a considerable increase in the low-temperature OSC for the CeO₂-Nps-400 compared with CeO₂-Nrs-400 and CeO₂-Ncs-400 counterparts. In the following H₂-TPR process, H₂ reacts with the samples by reducing Ce⁴⁺ to Ce³⁺. Therefore, the increase in Ce³⁺ and oxygen vacancy concentrations increased the OSC of the ceria nanostructures.⁴⁵ The results are consistent with the BET surface area and lattice defect in Raman analysis, as the OSC is well related to the surface area and lattice defect formation of nanostructures.

The CeO₂ nanostructures also showed a clear contradiction pattern between XRD and BET analyses, where it showed an increasing crystallite size in the order of CeO₂-Nps-400 > CeO₂-Nrs-400 > CeO₂-Ncs-400.⁷² Simultaneously, the surface area of the catalyst follows the reverse order. It was found that by decreasing crystallite size, the CeO₂ nanostructures were exposed to the high OSC.⁷³ Thus, the large surface area to volume ratio existing in a catalyst permits CeO₂ to react differently, resulting in unique properties. A larger particle size results in lattice expansion in the crystal structure. The expansion of the lattice decreases its oxygen release and reabsorption capabilities, as reported by Hailstone *et al.*²⁹ This is proved by observing the catalytic performance of the CeO₂-Ncs-400 catalyst, which produced the least amount of vanillin yield compared to the other catalysts. In particular, high surface area-engaged nanostructures are considered perfect for the oxidation reaction by providing a greater tendency for active species to come into contact with reactants and improve catalytic performance.⁷⁴ The relationship between the specific area and OSC was investigated by several authors,^{73,75,76} considering that

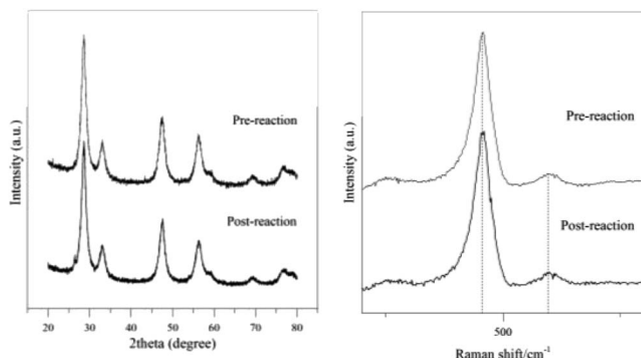


Fig. 12 Characterization of CeO₂-Nps-400 using XRD and Raman analyses for the post-reaction.

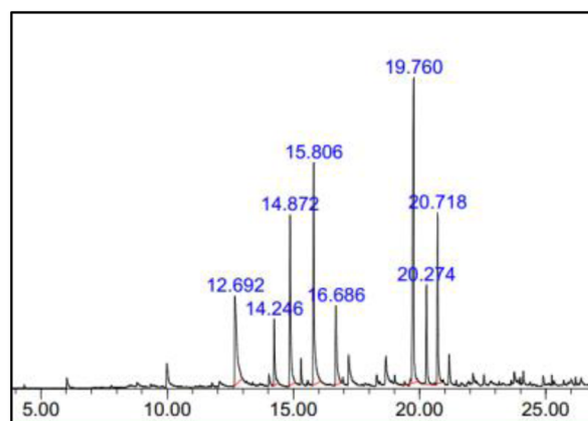


Fig. 13 Gas chromatogram of degradation wood products.



materials with high specific areas display more active sites to adsorb oxygen from external sources, such as oxidants that supply more active oxygen species.⁷⁷ The summarized results in Table 1 show that the CeO₂-Nps-400, CeO₂-Nrs-400 and CeO₂-Ncs-400 whose areas correspond to 66.1, 57.8 and 22.5 m² g⁻¹, respectively, while the OSC values are 1664.6, 1340.4 and 835.3 μmol O₂ per g, respectively. As the BET surface area of the materials increased, the OSC value also increased. The comparable OSC values in different nanostructures of CeO₂ indicate that specific surface area and OSC have a high relation and depend on each other as the vanillin yield also shows an increasing pattern with increasing BET surface area and lower temperature OSC.

Different structural characteristics exposed to different active facets established a direct correlation between lower-temperature OSC and the presence of defects.⁷⁸ Generally, there are three low-index lattice planes on the surface of CeO₂ nanostructures that are facet (100), (110) and (111). The stability and excess charge localization of all three facets follow the sequence (111) > (110) > (100), whereas the activity follows the reverse order.^{78–81} According to initial research conducted by Ramli *et al.*,²² the reported TEM and HRTEM results show that CeO₂ nanoparticles promoted the exposure of Ce(111) and Ce(100) facets, while CeO₂ nanorods preferentially enclosed (110) facets,^{82–84} followed by ceria nanocubes promoting the exposure of Ce(100) facets. The (110) and (100) facets are considered more active than the (111) facets in terms of catalytic activity. However, in this study, it was found that nanoparticles promote (100) and (111), which contradicts the activity order of the facets for ceria nanostructures. Moreover, a nanocube was found to promote (100), but its reactivity on the conversion of vanillin shows a poorer yield among other ceria nanostructures. Konsolakis and Lykaki *et al.*²⁷ reported that in terms of the relation to oxygen storage capacity (OSC) obtained for bare ceria nanostructures in their study, the order shows (111) > (100) > (110), which also can correlate with the energy formation of anionic vacancies. In this study, the facets (100) and (111) enclosed by ceria nanoparticles are found to express significantly enhanced redox properties required for higher catalytic activities in converting vanillin from direct oxidation of kenaf stalks,²² which also reconfirms the involvement of the active crystal facets in chemical reactions. They also said that both the (100) and (111) facets facilitate the regeneration of active sites due to their improved oxygen exchange kinetics, thus favoring the regeneration of active sites²⁷ for the oxidation reaction.

Vanillin characterization. Gas chromatography-mass spectrometry (GC-MS) is utilised to examine the detailed composition of the product. Fig. 13 displays the chromatogram corresponding to the varied composition of product presence obtained from the direct oxidation of kenaf stalks identified with profiles from the NIST14.L library. The results only reported major compounds discovered by GC-MS with an area greater than 4%. Syringaldehyde and vanillin were the most abundant aromatic monomers discovered during wood degradation at retention times of 15.806 and 19.760 min, followed by coumaran, 4-vinylguaiacol, syringol, 4-propylguaiacol, aceto-syringone, and non-aromatic monomers, such as glutaric acid.

Based on the GC-MS findings, vanillin was proven to be produced *via* the direct oxidation of the kenaf stalks.

Despite the inclusion of additional aromatic and non-aromatic monomer chemicals in the products, this is remarkable because vanillin is extracted without even isolating the lignin components from cellulose and hemicellulose in kenaf stalks. According to Qu *et al.*,⁶ direct vanillin production from Japanese cedar wood yielded 5% vanillin when CuO is present as a catalyst, and degradation products detected the presence of other aromatic monomer compounds, such as guaiacol, acetovanillone, vanillic acid, syringaldehyde, and syringic acid. This showed that syringaldehyde was identified with a higher peak area than vanillin. This is because hardwood lignin contains both guaiacylpropane and syringylpropane units. Thus, the alkaline oxidation of angiosperm lignin generates large amounts of syringaldehyde. Syringaldehyde has the most similar applications as vanillin because both possess almost the same chemical structure and properties, which is also one reason for the difficulty of separating syringaldehyde from pure vanillin. Vanillin purification is also challenging due to contaminants, such as carbohydrates, various side products, and unreacted lignin.⁸⁴

Conclusions

Vanillin was successfully produced from the direct oxidation of kenaf stalks under microwave heating using H₂O₂ as an oxidizing agent in the presence of CeO₂-Nps-400, CeO₂-Nrs-400 and CeO₂-Ncs-400 catalysts at pH 11.5. A vanillin yield of 3.84% was achieved using CeO₂-Nps-400 as a heterogeneous catalyst, while 4.66% was obtained when NaOH was used as a homogeneous catalyst. The heterogeneous CeO₂-Nps-400 catalyst is chemically stable and can be repeated use for 3 times. Based on the catalytic evaluation, vanillin production showed increasing yield in order of CeO₂-Ncs-400 < CeO₂-Nrs-400 < CeO₂-Nps-400, which also follows the same trend as characteristic morphology in textural and redox properties of CeO₂ nanostructures. The synthesized CeO₂ nanoparticles proved that the highest vanillin yield was affected by its morphology, where it was recorded with high- and low-temperature OSC value, easily weakly bound of oxygen species through TPR and TPO and exceptional high oxygen vacancies from Raman and XPS analyses.

Data availability

Data are present within the article and its ESI.†

Author contributions

Conceptualization, A. R.; methodology, N. A. S. I. K. A.; software, N. A. S. I. K. A.; validation, A. R., N. M. Y. and A. R. M.; formal analysis, N. A. S. I. K. A.; investigation, N. A. S. I. K. A.; resources, A. R., N. M. Y. and A. R. M.; data curation, N. A. S. I. K. A.; writing—original draft preparation, N. A. S. I. K. A.; writing—review and editing, A. R.; visualization, A. R., N. M. Y. and A. R. M.; supervision, A. R., N. M. Y. and A. R. M.; project administration, A. R.; funding acquisition, A. R., N. M. Y. and A.



R. M. All authors have read and agreed to the published version of the manuscript.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

The authors would like to thank Malaysian Ministry of Higher Education under the Fundamental Research Grant Scheme (FRGS/1/2020/STG04/UTP/02/2, cost centre 015MA0-125, funder ID 10.13039/501100003093) for the research grants awarded to conduct the research.

Notes and references

- M. Fache, B. Boutevin and S. Caillol, *ACS Sustain. Chem. Eng.*, 2015, **4**, 35–46.
- N. J. Walton, M. J. Mayer and A. Narbad, *Phytochemistry*, 2003, **63**, 505–515.
- E. B. da Silva, M. Zabkova, J. D. Araújo, C. A. Cateto, M. F. Barreiro, M. N. Belgacem and A. E. Rodrigues, *Chem. Eng. Res. Des.*, 2009, **87**, 1276–1292.
- F. S. Chakar and A. J. Ragauskas, *Ind. Crops Prod.*, 2004, **20**, 131–141.
- H. Deng, L. Lin and S. Liu, *Energy Fuels*, 2010, **24**, 4797–4802.
- C. Qu, M. Kaneko, K. Kashimura, K. Tanaka, S. Ozawa and T. Watanabe, *ACS Sustain. Chem. Eng.*, 2017, **5**, 11551–11557.
- D. Mukherjee, R. Singuru, P. Venkataswamy, D. Damma and B. M. Reddy, *ACS Omega*, 2019, **4**, 4770–4778.
- F. Liu, L. Chen, J. K. Neathery, K. Saito and K. Liu, *Ind. Eng. Chem. Res.*, 2014, **53**, 16341–16348.
- W. Deng, H. Zhang, X. Wu, R. Li, Q. Zhang and Y. Wang, *Green Chem.*, 2015, **17**, 5009–5018.
- S. Rawat, P. Gupta, B. Singh, T. Bhaskar, K. Natte and A. Narani, *Appl. Catal., A*, 2020, **598**, 117567.
- N. A. S. I. Khairul Anuar, A. Ramli and L. Jun Wei, *Catalysts*, 2021, **11**, 1449.
- W. I. Hsiao, Y. S. Lin, Y. C. Chen and C. S. Lee, *Chem. Phys. Lett.*, 2007, **441**, 294–299.
- M. Melchionna and P. Fornasiero, *Mater. Today*, 2014, **17**, 349–357.
- Z. A. Qiao, Z. Wu and S. Dai, *ChemSusChem*, 2013, **6**, 1821–1833.
- Z. Wu, M. Li and S. H. Overbury, *J. Catal.*, 2012, **285**, 61–73.
- W. Huang, *Top. Catal.*, 2013, **56**, 1363–1376.
- M. Konsolakis and M. Lykaki, *Catalysts*, 2021, **11**, 452.
- D. Chen, D. He, J. Lu, L. Zhong, F. Liu, J. Liu, J. Yu, G. Wan, S. He and Y. Luo, *Appl. Catal., B*, 2017, **218**, 249–259.
- Y. Ohtani, B. B. Mazumder and K. Sameshima, *J. Wood Sci.*, 2001, **47**, 30–35.
- G. Al-Naqeb, M. Ismail, G. Bagalkotkar and H. A. Adamu, *Food Res. Int.*, 2010, **43**, 2437–2443.
- N. Saba, M. T. Paridah and M. Jawaid, *Constr. Build. Mater.*, 2015, **76**, 87–96.
- A. Ramli, N. A. S. I. K. Anuar, N. A. A. Bakhtiar, N. M. Yunus and A. R. Mohamed, *Molecules*, 2023, **28**, 4963.
- L. Torrente-Murciano and R. S. Chapman, *Phys. Chem. Chem. Phys.*, 2016, **18**, 15496–15500.
- M. Kovacevic, B. L. Mojet, J. G. van Ommen and L. Lefferts, *Catal. Lett.*, 2016, **146**, 770–777.
- J. Hierso, Ö. Sel, A. Ringuedé, C. Laberty-Robert, L. Bianchi, D. Grosso and C. Sánchez, *Chem. Mater.*, 2009, **21**, 2184–2192.
- L. Ma, D. Wang, J. Li, B. Bai, L. Fu and Y. Li, *Appl. Catal., B*, 2014, **148–149**, 36–43.
- M. Lykaki, S. Stefa, S. A. Carabineiro, P. K. Pandis, V. N. Stathopoulos and M. Konsolakis, *Catalysts*, 2019, **9**, 371.
- F. Gao, Q. Lu and S. Komarneni, *J. Nanosci. Nanotechnol.*, 2006, **6**, 3812–3819.
- R. K. Hailstone, A. G. DiFrancesco, J. G. Leong, T. D. Allston and K. J. Reed, *J. Phys. Chem. C*, 2009, **113**, 15155–15159.
- M. Thommes, K. Kaneko, V. Neimark, P. Alexander, J. R. F. Olivier, J. Rouquerol and S. W. K. Sing, *Pure Appl. Chem.*, 2015, **87**, 1051–1069.
- J. Arnaldo, *Microporous Mesoporous Mater.*, 2020, **291**, 109698.
- N. Kainbayev, M. Sriubas, D. Virbukas, Z. Rutkuniene, K. Bockute, S. Bolegenova and G. Laukaitis, *Coatings*, 2020, **10**, 432.
- M. Yang, G. Shen, Q. Wang, K. Deng, M. Liu, Y. Chen and Z. Wang, *Molecules*, 2021, **26**, 6363.
- E. N. Ndifor, T. Garcia, B. Solsona and S. H. Taylor, *Appl. Catal., B*, 2007, **76**, 248–256.
- J. Shyu, *J. Catal.*, 1989, **115**, 16–23.
- J. D. Kammert, J. Moon and Z. Wu, *Chin. J. Catal.*, 2020, **41**, 901–914.
- G. Ranga Rao and B. G. Mishra, *Bulletin of Catalysis Society of India*, 2003, **2**, 122–134.
- Tana, M. Zhang, J. Li, H. Li, Y. Li and W. Shen, *Catal. Today*, 2009, **148**, 179–183.
- L. Liu, Y. Cao, W. Sun, Z. Yao, B. Liu and F. Gao, *Catal. Today*, 2011, **175**, 48–54.
- T. Désaunay, G. Bonura, V. Chiodo, S. Freni, J. P. Couzinié, J. Bourgon and M. Cassir, *J. Catal.*, 2013, **297**, 193–201.
- B. Murugan and A. V. Ramaswamy, *J. Am. Chem. Soc.*, 2007, **129**, 3062–3063.
- S. Hamoudi, F. Larachi, G. Cerrella and M. Cassanello, *Ind. Eng. Chem. Res.*, 1998, **37**, 3561–3566.
- R. Bortamuly, G. Konwar, P. K. Boruah, M. R. Das, D. Mahanta and P. Saikia, *Ionics*, 2020, **26**, 5747–5756.
- P. Burroughs, A. Hamnett, A. F. Orchard and G. Thornton, *Dalton Trans.*, 1976, **17**, 1686.
- M. Pan, S. Zhang, Y. Xu and R. Li, *Appl. Surf. Sci.*, 2018, **448**, 435–443.
- C. Anandan and P. Bera, *Appl. Surf. Sci.*, 2013, **283**, 297–303.
- E. Bêche, P. Charvin, D. Perarnau, S. Abanades and G. Flamant, *Surf. Interface Anal.*, 2008, **40**, 264–267.
- B. M. Reddy, A. Khan, Y. Yamada, T. Kobayashi, S. Loridant and J. C. Volta, *J. Phys. Chem.*, 2003, **107**, 5162–5167.



- 49 V. Stetsovych, F. Pagliuca, F. Dvořák, T. Duchoň, M. Vorokhta, M. Aulická and V. Matolín, *J. Phys. Chem. Lett.*, 2013, **4**, 866–871.
- 50 S. Niroumandrad, M. Rostami and B. Ramezanzadeh, *Appl. Surf. Sci.*, 2015, **357**, 2121–2130.
- 51 B. Zhang, Y. Huyan, J. Wang, W. Wang, Q. Zhang and H. Zhang, *J. Am. Ceram. Soc.*, 2018, **102**, 2218–2227.
- 52 Z. Su, W. Yang, C. Wang, S. Xiong, X. Cao, Y. Peng, W. Si, Y. Weng, M. Xue and J. Li, *Environ. Sci. Technol.*, 2020, **54**, 12684–12692.
- 53 P. Sudarsanam, B. Hillary, M. H. Amin, N. Rockstroh, U. Bentrup, A. Brückner and S. K. Bhargava, *Langmuir*, 2018, **34**, 2663–2673.
- 54 B. Hillary, P. Sudarsanam, M. H. Amin and S. K. Bhargava, *Langmuir*, 2017, **33**, 1743–1750.
- 55 J. M. López, A. Gilbank, T. García, B. Solsona, S. Agouram and L. Torrente-Murciano, *Appl. Catal., B*, 2015, **174**, 403–412.
- 56 A. I. Y. Tok, S. Du, F. Boey and W. K. Chong, *Mater. Sci. Eng., A*, 2007, **466**, 223–229.
- 57 X. Xu, P. Li, Y. Zhong, J. Yu, C. Miao and G. Tong, *J. Biol. Macromol.*, 2023, **243**, 125203.
- 58 R. Prado, X. Erdocia, G. F. de Gregorio, J. Labidi and T. Welton, *ACS Sustain. Chem. Eng.*, 2016, **4**, 5277–5288.
- 59 D. Zhang, X. Du, L. Shi and R. Gao, *Dalton Trans.*, 2012, **41**, 14455–14475.
- 60 M. Zabilskiy, P. Djinović, E. Tchernychova, O. P. Tkachenko, L. M. Kustov and A. Pintar, *ACS Catal.*, 2015, **5**, 5357–5365.
- 61 M. Lykaki, E. Pachatouridou, S. A. C. Carabineiro, E. Iliopoulou, C. Andriopoulou, N. Kallithrakas-Kontos, S. Boghosian and M. Konsolakis, *Appl. Catal., B*, 2018, **230**, 18–28.
- 62 R. Behling, G. Chatel and S. Valange, *Ultrason*, 2017, **36**, 27–35.
- 63 J. C. Walter, A. Zurawski, D. M. Montgomery, M. Thornburg and S. T. Revankar, *J. Power Sources*, 2008, **179**, 335–339.
- 64 A. M. da Costa Lopes, J. R. Gomes, J. A. Coutinho and A. J. Silvestre, *Green Chem.*, 2020, **22**, 2474–2487.
- 65 B. S. Takale, M. Bao and Y. Yamamoto, *Org. Biomol. Chem.*, 2014, **12**, 2005.
- 66 S. Santra, A. K. Bagdi, A. Majee and A. Hajra, *RSC Adv.*, 2013, **47**, 24931.
- 67 M. J. Sampaio, A. Benyounes, P. Serp, J. L. Faria and C. G. Silva, *Appl. Catal., A*, 2018, **551**, 71–78.
- 68 N. J. Abd Rahman, PhD thesis, Universiti Teknologi PETRONAS, Perak, Malaysia, 2020.
- 69 A. Younis, D. Chu and S. Li, *Funct. Nanomater.*, 2016, **3**, 53–68.
- 70 G. Sharma, D. D. Dionysiou, S. Sharma, A. Kumar, A. H. Al-Muhtaseb, M. Naushad and F. J. Stadler, *Catal. Today*, 2019, **335**, 437–451.
- 71 D. Wei, M. Yue, S. Qin, S. Zhang, Y. Wu, G. Xu, H. Zhang, T. Zhang and J. Li, *J. Am. Chem. Soc.*, 2021, **143**, 15635–15643.
- 72 Y. Xiao, H. Li and K. Xie, *Angew. Chem., Int. Ed.*, 2021, **60**, 5240–5244.
- 73 J. Wang, M. Shen, J. Wang, J. Gao, J. Ma and S. Liu, *Catal. Today*, 2011, **175**, 65–71.
- 74 V. P. Santos, M. Pereira and J. Órfão, *Appl. Catal., B*, 2010, **99**, 353–363.
- 75 A. Migani, G. N. Vayssilov, S. T. Bromley, F. Illas and K. M. Neyman, *J. Mater. Chem.*, 2010, **20**, 10535–10546.
- 76 Z. Minwei, M. Shen and J. Wang, *J. Catal.*, 2007, **248**, 258–267.
- 77 N. Kamiuchi, M. Haneda and M. Ozawa, *Catal. Today*, 2014, **232**, 179–184.
- 78 M. Cui, Y. Li, X. Wang, J. Wang and M. Shen, *J. Rare Earths*, 2013, **31**, 572–576.
- 79 E. Mamontov, T. Egami, R. Brezny, M. Koranne and S. Tyagi, *J. Phys. Chem. B*, 2000, **104**, 11110–11116.
- 80 D. C. Sayle, S. A. Maicaneanu and G. W. Watson, *J. Am. Chem. Soc.*, 2002, **124**, 11429–11439.
- 81 Y. Jiang, J. B. Adams and M. van Schilfhaarde, *J. Chem. Phys.*, 2005, **123**, 64701.
- 82 Y. Chen, P. Hu, M. H. Lee and H. Wang, *Surf. Sci.*, 2008, **602**, 1736–1741.
- 83 T. X. T. Sayle, M. Cantoni, U. M. Bhatta, S. C. Parker, S. R. Hall, G. Möbus, M. Molinari, D. Reid, S. Seal and D. C. Sayle, *Chem. Mater.*, 2012, **24**, 1811–1821.
- 84 R. Zhang, R. Maltari, M. Guo, J. Kontro, A. Eronen and T. Repo, *Ind. Crops Prod.*, 2020, **145**, 112095.

