The Influence of Precursor on the Preparation of CeO$_2$ Catalysts for the Total Oxidation of the Volatile Organic Compound Propane

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Abstract: CeO$_2$ catalysts were prepared by a precipitation method using either (NH$_4$)$_2$Ce(NO$_3$)$_6$ or Ce(NO$_3$)$_3$, as Ce$^{IV}$ or Ce$^{III}$ precursors respectively. The influence of the different precursors on catalytic activity was evaluated for the total oxidation of propane with water present in the feed. The catalyst prepared using the Ce$^{IV}$ precursor was more active for propane total oxidation. The choice of precursor influenced catalyst properties such as surface area, reducibility, morphology, and active oxygen species. The predominant factor associated with the catalytic activity was related to the formation of either CeO$_2$.nH$_2$O or Ce$_2$(OH)$_2$(CO$_3$)$_2$.H$_2$O precipitate species, formed prior to calcination. The formation of CeO$_2$.nH$_2$O resulted in enhanced surface area which was an important factor for controlling catalyst activity.

Keywords: VOC total oxidation; cerium oxide; propane oxidation; cerium precursor

1. Introduction

Volatile organic compounds (VOCs) are associated with various issues affecting the environment and human health [1]. For example, the formation of ground level ozone from the reaction of VOCs with NO$_x$ species in the atmosphere has been linked with respiratory issues in humans, as well as some VOCs being known carcinogens [2]. VOCs are wide ranging in their chemical nature and short chain alkanes, such as propane, are known to be difficult to remove from the atmosphere [3], hence propane is an excellent model compound to study. In addition, the increasing use of liquid petroleum gas (LPG) as a transport fuel has led to a rise in propane emissions [4]. As short chain alkane emissions are linked to mobile and stationary sources including the petroleum industry and vehicle exhausts, other compounds such as water vapour may co-exist in the gas feed [5]. These compounds are thought to affect the removal of VOCs such as propane, therefore recent research has also focused on the impact of the conditions used for VOC removal [6,7].

Control of VOC emissions is of growing concern and mitigation can be achieved in many ways, such as thermal oxidation, catalytic oxidation, adsorption, and absorption [8,9]. However, catalytic oxidation has been identified as a more efficient way to remove VOCs, requiring less energy than thermal oxidation, and unlike adsorption and absorption techniques it is a destructive process. Catalytic oxidation also has the ability to simultaneously remove multiple VOCs from waste streams and treat low levels of VOCs [10]. In addition, catalytic oxidation has the benefit of producing more environmentally benign products in comparison with toxic by-products often created by thermal oxidation [11]. Noble metal catalysts containing Pd and Pt have been widely reported as active catalysts for VOC oxidation; however, there is a driving force to switch to metal oxide catalysts, as they have the advantage of being less expensive and more abundant [12].

Amongst the metal oxide catalysts reported, cerium (IV) oxide (CeO$_2$) is widely regarded as an effective oxidation catalyst, due to many beneficial characteristics. The

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favourable redox properties of CeO₂, in addition to its high oxygen storage capacity (OSC), and ability to form oxygen defects, which enable fast oxygen mobility through the lattice, has made it the catalyst of choice for various oxidation reactions [13–16]. Furthermore, many other metal oxides have been used in conjunction with CeO₂ to enhance their characteristics, resulting in more active catalysts for soot [17] and benzene [18] oxidation. The morphology of CeO₂ based catalysts can also be finely tuned to selectively control the exposed facets. This method has been employed for a range of oxidation reactions, leading to surface interactions improving catalytic activity [19–21]. The impact of adsorbed species, such as water vapour, on different CeO₂ surfaces has also been studied in the literature [22] which is important when considering reaction conditions used. The structure and transformation of cerium precursors to form CeO₂ have been studied, leading to materials with varying characteristics, but few consider the catalytic uses for these materials [23]. Recent research on the influence of the cerium precursor salt for catalysts have mainly been studied in the context of mixed metal oxides [24–26], but single oxide CeO₂ has rarely been investigated.

It is well established that altering the synthesis route for catalyst preparation can drastically influence the activity trends for various reactions [27]. Altering preparation method, aging and calcination conditions have all been shown to result in varied catalyst characteristics, which can improve activity [14]. For example, precipitation allows for the careful control of synthesis parameters such as pH, aging time, solution concentrations and precipitating agent. However, the influence of the metal precursor can sometimes be overlooked.

In this work, two CeO₂ catalysts were prepared by a precipitation method, using either (NH₄)₂Ce(NO₃)₆ or Ce(NO₃)₃ as the precursor cerium source. These catalysts were characterised by X-ray diffraction (XRD), X-ray photoelectron spectroscopy (XPS), Thermal gravimetric analysis-Differential thermal analysis (TGA-DTA), Laser Raman spectroscopy, Temperature programmed reduction (TPR), Electron microscopy (TEM/SEM), and Brunauer–Emmett–Teller (BET) surface area analysis. Catalysts were evaluated for the total oxidation of propane as a model VOC, focusing on the impact the cerium precursor had on the catalytic activity and how it can be related to the structure, redox properties, and surface state of the CeO₂ catalysts.

2. Results and Discussion
2.1. Catalyst Precursor Characterisation

The dried catalyst precursors prior to calcination were analysed using powder XRD (Figure 1) and there was a significant difference between the two precipitated precursors. The d-CeO₂ (IV) sample had a structure representative of a cubic CeO₂ phase with low crystallinity, exhibiting reflections at 29°, 33°, 47°, and 57°, corresponding to (111), (200), (220) and (311) lattice planes, respectively. Whereas the d-CeO₂ (III) sample predominantly exhibited reflections representing an orthorhombic Ce₂(OH)₂(CO₃)₂.H₂O structure [28], with other additional reflections identified as hexagonal Ce(OH)(CO₃). Hence, under these synthesis conditions, the Ce⁴⁺ and Ce⁴⁺ precursors follow two different chemical precipitation mechanisms.
Thermal decomposition of the precipitated catalyst precursors were analysed using TGA-DTA (Figure 2a,b). Around 7% mass loss was observed from the d-CeO$_2$ (IV) sample over the range 100–500 °C. In contrast, the d-CeO$_2$ (III) sample showed a mass loss of 23% with a small decrease around 70 °C, followed by a sharp decrease between 250–300 °C. These were associated with the evolution of H$_2$O species and decomposition of carbonate species from the dried sample respectively; confirming that the precipitate formed from the Ce$^{III}$ precursor was a Ce(CO$_3$)$_x$(OH)$_y$ type species [29]. Furthermore, the endothermic peak around 250 °C, characterised by DTA, is attributed to the liberation of CO$_2$ from the decomposition of carbonate species [30]. The broad endothermic peak observed between 50–250 °C, in the d-CeO$_2$ (IV) sample, can be attributed to the evolution of H$_2$O. Previous work carried out by Hirano et al. looked at the precipitation mechanism of different Ce$^{IV}$ and Ce$^{III}$ precursors using urea prepared by hydrothermal synthesis [31,32]. They discovered that when using a Ce$^{IV}$ precursor, hydrated [Ce(OH)$_y$(H$_2$O)$_{n-y}$]$^{4+y}$ ions were formed as a result of the ability to undergo strong hydration from the lower basicity and higher charge of the Ce$^{4+}$ ion. This caused CeO$_2$.nH$_2$O to precipitate rapidly before reaction with carbonate to form Ce(CO$_3$)$_x$(OH)$_y$ type species could occur. Evidence of CeO$_2$ nanoparticle formation from the hydrolysis of ammonium cerium nitrate in aqueous solution without the addition of a base has also been documented by Pettinger et al. [33]. This effect was not observed when using different Ce$^{III}$ precursors, and the products formed were the carbonates, either Ce$_2$O(CO$_3$)$_2$.H$_2$O or Ce(OH)CO$_3$ [31].
Figure 2. Thermal gravimetric and differential thermal analysis of the precipitated catalyst precursors: (a) d-CeO$_2$ (IV) and (b) d-CeO$_2$ (III). Samples heated under flowing air atmosphere from 50 to 750 °C at 5 °C min$^{-1}$.

Based on the bulk phases identified from XRD, the theoretical mass losses were calculated for each thermal decomposition, assuming CeO$_2$ was the final product.

\[
\text{Ce}_2(\text{OH})_2(\text{CO}_3)_2\cdot\text{H}_2\text{O} \rightarrow 2\text{CeO}_2 
\]
(1)

\[
\text{CeO}_2\cdot n\text{H}_2\text{O} \rightarrow \text{CeO}_2 
\]
(2)

For the Ce$^{III}$ precursor, the theoretical mass loss was calculated to be 24% which is in good agreement with the 23% experimental loss identified by TGA-DTA analysis. From the XRD data, the formation of hydrated CeO$_2$ particles in solution was the only product formed using the Ce$^{IV}$ precursor. Therefore, a mass loss of 7% equates to 0.67 H$_2$O.

2.2. Catalyst Characterisation

Table 1 summarises some of the characterisation data for the two CeO$_2$ catalysts. The BET surface areas were significantly different for the two catalysts, with CeO$_2$ (IV) having a surface area roughly four times larger than CeO$_2$ (III). The adsorption–desorption isotherms for the two catalysts differ (Figure 3a,b), indicating diverse pore structures. Both plots represent a type IV isotherm, indicative of a mesoporous-type structure, with the CeO$_2$ (IV) catalyst having an H2 hysteresis loop. This type of hysteresis is indicative of capillary condensation in disordered and ill-defined pore structures, suggesting a higher porosity, which would be consistent with the higher surface area shown for this catalyst. In contrast, the CeO$_2$ (III) catalyst shows an H3 hysteresis loop, which is related to the formation of non-rigid, plate-like particles that form a disordered pore structure with slit-shaped pores [34]. In addition, hysteresis of this kind is linked with the incomplete filling of macro pores, suggesting a higher concentration of macro pores in this catalyst [35].

Table 1. Physiochemical properties of CeO$_2$ (IV) and CeO$_2$ (III) catalysts calcined at 500 °C for 3 h in static air.

| Catalyst   | Surface Area/m$^2$ g$^{-1}$ | Average Crystallite Size/nm | Lattice Parameter/nm | A$_{590}$/A$_{463}$ |
|------------|-----------------------------|-----------------------------|----------------------|---------------------|
| CeO$_2$ (IV) | 81                          | 8.7                         | 0.5409               | 0.018               |
| CeO$_2$ (III) | 19                          | 9.3                         | 0.5408               | 0.019               |
SEM images shown in Figure 4 indicate the different morphologies of the catalysts. This information helps to rationalise the surface area and adsorption–desorption isotherm data previously discussed. The CeO$_2$ (III) catalyst displays clumped aggregates with platelet or needle-like structures, which is representative of the H3 hysteresis loop; whereas, the CeO$_2$ (IV) catalyst forms larger well-defined particles. From the characterisation of the precipitates formed during the synthesis, it can be suggested that the presence of the Ce$_2$(OH)$_2$(CO$_3$)$_2$.H$_2$O phase ensures the formation of these type of non-rigid aggregates seen for CeO$_2$ (III). In contrast, forming the CeO$_2$.nH$_2$O precipitate forms the well-defined structures shown for CeO$_2$ (IV).

TEM images displayed in Figure 5 strengthen conclusions drawn from the SEM data. Differences in large scale morphology are apparent between CeO$_2$ (IV) and CeO$_2$ (III) (Figure 5a,d). However, the small-scale morphology appears similar (Figure 5b,e), with both CeO$_2$ (IV) and CeO$_2$ (III) samples showing agglomerated, small faceted CeO$_2$ particles of 8–10 nm, consistent with the crystallite size determined by XRD. High magnification images representative of the catalyst samples (Figure 5c,f) were used to measure the interplanar distances present on the CeO$_2$ (IV) and CeO$_2$ (III) catalysts, which could be noticed in
different regions (Figure 5b,e). The interplanar distances were 0.31 nm for both samples which is representative of the (111) lattice planes, indicating their preferential exposure.

![Transmission electron microscopy images and selected area electron diffraction patterns](image)

**Figure 5.** Transmission electron microscopy images and selected area electron diffraction patterns of (a–c) CeO$_2$ (IV) and (d–f) CeO$_2$ (III) catalysts.

Data acquired from XRD analysis of the two catalysts shown in Figure 6 indicates a similar bulk structure. XRD patterns of the catalysts only showed the cubic fluorite structure of CeO$_2$, with the lattice parameters of both catalysts within 0.0001 nm of each other. This is also consistent with the structure identified by selected area electron diffraction (SAED) analysis (Figure 5a,d). The crystallite sizes were calculated using the Scherrer equation by taking an average of the values obtained when analysing the peak widths of the four dominant (111), (200), (220), and (311) reflections. The sizes calculated were similar for both catalysts, with the CeO$_2$ (IV) catalyst having a slightly smaller crystallite size on average (8.7 nm verses 9.3 nm, Table 1). However, the average crystallite sizes calculated are within the experimental error (±1.2 nm), hence no significant difference can be identified.
It is well established that CeO$_2$ catalysts exhibit an intense Raman band around 460 cm$^{-1}$, relating to the F$_{2g}$ vibrational mode. Other weaker bands are active, such as a band around 600 cm$^{-1}$, which is associated with the presence of defect sites [36]. It has been previously established that the ratio of these two peaks can be used to estimate the concentration of defect sites in the material, which can then be used to greater understand the redox properties and oxygen mobility through the lattice of these types of catalysts [37]. This factor is especially important for propane total oxidation as it is thought that it takes place via a Mars-van Krevelen mechanism [12]. The Raman spectra shown in Figure 7 matches that of the cubic fluorite CeO$_2$ structure, consistent with that determined by XRD. The intense band at 463 cm$^{-1}$ denotes the F$_{2g}$ vibrational mode, and a small band around 590 cm$^{-1}$ indicated the presence of some defect species. The ratio of bands at 590 cm$^{-1}$ and 463 cm$^{-1}$ ($A_{590}/A_{463}$) are shown in Table 1, and they are very similar for both catalysts, which implies that the defect concentrations of both catalysts detectable by Laser Raman spectroscopy are similar.
From the TPR profiles shown in Figure 8, only one main reduction peak was observed for both catalysts. This peak occurred at similar temperatures for both catalysts, 478 °C for CeO₂ (III) and 498 °C for CeO₂ (IV). CeO₂ has two main types of reduction features, these are the reduction of surface species and the reduction of bulk lattice species [38]. Bulk reduction occurs at temperatures above 700 °C, whilst surface reduction occurs around 500 °C, therefore, the features present in the TPR profiles relate to the reduction and removal of oxygen species from the catalyst surface. Table 2 shows the H₂ consumption normalised for surface area and mass for both catalysts. Both catalysts exhibit good redox ability; however, the CeO₂ (III) catalyst had a higher H₂ consumption normalised for surface area. This factor, in combination with the lower temperature peak for the surface reduction, suggests CeO₂ (III) had a greater extent of surface reduction, and that reduction was slightly more facile compared to CeO₂ (IV). These differences between CeO₂ (III) and CeO₂ (IV), could possibly arise as a result of different structures, identified by SEM, originating from the different synthesis precursors and the subsequent transformation into the CeO₂ catalysts.

Table 2. H₂ Consumption of CeO₂ (IV) and CeO₂ (III) catalysts determined from temperature programmed reduction.

| Catalyst  | TPR Cycle | H₂ Consumption Per Surface Area/μmol m⁻² | H₂ Consumption Per Mass/μmol g⁻¹ |
|-----------|-----------|------------------------------------------|----------------------------------|
| CeO₂ (IV) | 1         | 0.435                                    | 35.20                            |
|           | 2         | 0.145                                    | 11.73                            |
| CeO₂ (III)| 1         | 4.575                                    | 86.93                            |
|           | 2         | 0.464                                    | 8.81                             |

To assess oxygen storage capacity (OSC), TPR-TPO cycles were performed. Data in Table 2 affirms the trend shown in the TPR analysis, with CeO₂ (III) having a higher H₂ consumption than the CeO₂ (IV) catalyst. However, upon re-oxidising and then reducing the catalysts again, the H₂ consumption for the CeO₂ (III) catalyst decreased significantly. The results for the H₂ consumption per surface area indicated a decrease by a factor of 10 for the CeO₂ (III) catalyst, whilst the decrease was only three-fold for the CeO₂ (IV), bringing the values closer together for both catalysts. This analysis also suggests that the
first TPR analysis may not be representative of the redox properties of the catalyst under reaction conditions. Furthermore, it has been suggested that the re-oxidation of the catalyst is the rate determining factor [38], hence there may not be a direct relationship between propane oxidation activity and H₂ consumption.

Core-level Ce 3d photoelectron spectra for both CeO₂ (III) and (IV) catalysts are shown in Figure 9. Given the large number of final states arising from photoemission, Ce 3d XPS spectra are recognised as being difficult to analyse; however, it is generally accepted a total of 10 peaks are present for CeO₂ relating to mixed (III)/(IV) states. These peaks are divided into the Ce³⁺ and Ce⁴⁺ oxidation states with peaks denoted v₀, v’, u₀, u’ being used to calculate the concentration of Ce³⁺ and v, v”, v’”, u, u”, u’” used to represent the Ce⁴⁺ oxidation state [39]. The ratio of surface Ce³⁺ to Ce⁴⁺ was calculated using the integrated peak areas of each relative fitting of the two oxidation states (Table 3). The CeO₂ (IV) catalyst shows a higher ratio, indicating a higher concentration of reduced Ce³⁺ species, which is initially counterintuitive using a precursor with cerium in the +4 oxidation state.

It is proposed that the higher concentration of Ce³⁺ surface species directly relates to an increased amount of surface defect sites, which can affect catalytic activity [21,37,38,40]. The increased amounts of reduced Ce³⁺ surface species in the CeO₂ (IV) catalyst can also be linked to the lower surface H₂ consumption determined by TPR analysis. The lower quantity of reduction, defined by H₂ consumption, for the CeO₂ (IV) catalyst could be a direct result of the higher concentration of Ce³⁺ species identified, hence the CeO₂ (IV) catalyst would initially have a more reduced surface.

![Figure 9](image-url)

**Figure 9.** X-ray photoelectron spectra of Ce 3d region for (a) CeO₂ (IV) and (b) CeO₂ (III) catalysts, where fitted peaks refer to Ce³⁺ (blue) and Ce⁴⁺ (red) states.

| Catalyst   | XPS Ce:O:Na Ratio | O⁰/O⁻ Ratio | Ce³⁺/Ce⁴⁺ Ratio | EDX Ce:O:Na Ratio |
|------------|-------------------|-------------|----------------|------------------|
| CeO₂ (IV)  | 29:68:3           | 0.438       | 0.155          | 30:68:2          |
| CeO₂ (III) | 22:67:11          | 0.289       | 0.086          | 34:59:7          |

Catalytic activity may also be influenced by the surface oxygen species. Figure 10 shows the fitted O 1s core-level spectra for both catalysts. Two distinct oxygen species/environments can be extracted from the spectra, which we ascribe as O⁰ (531 eV) and O⁻ (529 eV) states, reported to be characteristic of defect oxygen and lattice oxygen species respectively [37,41]. There is some controversy on the labelling of the O⁰ region; it is thought that this region could also relate to the presence of hydroxyl and carbonate...
oxygen species [37]. However, some researchers have linked it to the appearance of low co-ordination oxide ions [41,42]. As a result, the species represented by the $O^β$ region have been referred to as surface oxygen defect sites. The ratio of $O^β/O^α$ shown in Table 3 indicates the CeO$_2$ (IV) catalyst has a higher proportion of surface oxygen defect sites compared to the CeO$_2$ (III) catalyst, which is self-consistent with the higher quantity of Ce$^{3+}$ on the surface determined by XPS analysis of the Ce 3d region. In contrast, data acquired by Laser Raman spectroscopy showed little difference in defect concentration between the catalysts and suggests the defects are highly localised in the near surface region, due to the surface sensitivity of XPS, and the bulk sampling of Laser Raman.

![Figure 10](image)

**Figure 10.** X-ray photoelectron spectra of O 1s peaks for calcined (a) CeO$_2$ (IV) and (b) CeO$_2$ (III) catalysts, where fitting shows the $O^α$ (red), $O^β$ (blue) oxygen species as described in the main text and yellow and green refer to the overlapping Na Auger peaks.

In addition to using XPS to identify surface oxygen and cerium species, it is also important to look at other surface species that could be present, which could affect catalytic activity. Alkali metals, such as Na, present from precipitation using Na$_2$CO$_3$, have been proposed as a catalyst poison for certain oxidation reactions by metal oxides [43]. From the XPS data shown in Table 3, the surface content of Na is much higher on CeO$_2$ (III) compared with CeO$_2$ (IV). This can also be evidenced from the presence of the Na Auger signal (green curve, Figure 10) in the O 1s spectra for CeO$_2$ (III). This effect is consistent with the EDX data, which shows that the CeO$_2$ (III) catalyst also has a higher bulk Na content (Table 3). As both catalyst precursors underwent the same extensive washing procedure, the precursor carbonate phase of the CeO$_2$ (III) appears to be more efficient in retaining Na.

### 2.3. Catalyst Performance

Catalyst performance for propane total oxidation is shown in Figure 11 for CeO$_2$ (IV) and CeO$_2$ (III) catalysts. The main reaction product detected was CO$_2$ and both catalysts maintained a carbon balance of >98%. Selectivity to CO$_2$ was >99% across the temperature range for CeO$_2$ (IV), whilst between 500–600 °C it was 96% for CeO$_2$ (III), due to the formation of low levels of propene. It is clear that the CeO$_2$ (IV) catalyst was more active for propane total oxidation across the temperature range. Data shown in Table 4 indicates that, when normalised for surface area, both catalysts had similar activity for propane oxidation.
Catalyst activity for propane total oxidation in the presence of water vapour: (Circle) CeO$_2$ (IV) and (Triangle) CeO$_2$ (III). Conditions: 5000 ppm propane in air, GHSV = 45,000 h$^{-1}$, 5% Water Saturation.

Table 4. Mass normalised and surface area normalised catalytic activity of CeO$_2$ (IV) and CeO$_2$ (III) catalysts for propane total oxidation.

| Catalyst  | Propane Conversion/% | Surface Area Normalised Propane Oxidation a/mol s$^{-1}$ m$^{-2}$ | Mass Normalised Propane Oxidation a/mol s$^{-1}$ g$^{-1}$ |
|-----------|----------------------|---------------------------------------------------------------|----------------------------------------------------------|
| CeO$_2$ (IV) | 50                   | 3.73 $\times$ 10$^{-3}$                                      | 3.02 $\times$ 10$^{-1}$                                   |
| CeO$_2$ (III) | 4                    | 3.68 $\times$ 10$^{-3}$                                      | 6.99 $\times$ 10$^{-2}$                                   |

*a* Calculated at 525 °C.

A comparison between wet and dry conditions, where the addition of 5% water was not included in the gas feed, was also assessed using the different CeO$_2$ catalysts (Figure S1). The activity trends shown for propane oxidation under dry conditions mirror those under wet conditions, with the CeO$_2$ (IV) catalyst remaining by far the most active. However, conversion to CO$_2$ was slightly increased for both catalysts under dry conditions indicating the inclusion of 5% water to the gas feed slightly inhibited propane oxidation. Marécot et al. reported that the addition of water inhibited propane and propene oxidation over Pt and Pd catalysts due to a decrease in the number of active surface sites [44]. Furthermore, researchers have previously reported the addition of water negatively impacted propane oxidation over metal oxide catalysts, which was thought to occur from competitive adsorption between water and propane on the catalyst surface [5,45]. Water inhibition has also been observed for other VOCs, such as toluene [2]. It was proposed that the addition of water created competition for adsorption sites with the VOC, leading to surface active sites being blocked by water [46,47]. Our current data and these other studies emphasise the importance of considering water in the VOC effluent.

It is shown from XRD and Raman characterisation that both catalysts had the common cubic fluorite CeO$_2$ structure. In addition, the lattice parameter, crystallite size and bulk defect concentration calculated using these techniques were very similar for both catalysts. Furthermore, whilst differences in large-scale morphology were identified by TEM and
SEM, the small-scale morphology for both catalysts were similar, with the (111) lattice plane exposed preferentially for both catalysts.

The redox properties of CeO$_2$ catalysts are thought to be important for oxidation reactions that occur via a Mars-van Krevelen mechanism, often showing a relationship between activity and increased H$_2$ consumption and more facile reducibility [48]. The ability to easily remove active oxygen from the catalyst surface facilitates C-H bond activation, which is known to be the rate determining step for propane oxidation [49]. From the data shown (Figure 11), the catalytic activity is greater for the CeO$_2$ (IV) catalyst. This suggests that the slightly enhanced redox behaviour of the CeO$_2$ (III) catalyst is less significant for controlling activity, and the surface areas of the catalysts are a far more important parameter.

From the TPR-TPO cycles carried out, it was shown that the surface H$_2$ consumption significantly decreased, roughly ten-fold, for the CeO$_2$ (III) catalyst in the second cycle, compared with CeO$_2$ (IV). The H$_2$ consumption of the second cycle was more comparable between the two catalysts, with CeO$_2$ (III) having only a slightly increased value. As previously mentioned, this suggests that the initial TPR may not be representative of the redox properties of the catalyst under reaction conditions.

The surface areas resulting from the different catalyst precursors are considered to be very influential when understanding the catalytic activity. It has been well documented by various researchers that the high surface area of CeO$_2$ is a key factor for improved catalytic activity of aromatic VOC oxidation such as naphthalene oxidation [14,50,51]. Similar conclusions can be drawn from the data presented in Figure 11, as the high surface area CeO$_2$ (IV) catalyst was more active for propane total oxidation. Previous research has also shown increased surface area of metal oxide catalysts improved the catalytic activity of propane oxidation due to the increased amount of active sites available [52,53]. When normalised for surface area, both catalysts have similar activity showing a linear relationship between catalytic activity and surface area, indicating this factor to be the most influential when understanding catalyst activity.

The identification of surface oxygen defect sites by XPS analysis showed that the CeO$_2$ (IV) catalyst contained a higher proportion of these sites compared to the CeO$_2$ (III) catalyst. Previous research using the same preparation precursor has indicated a strong correlation between the increased amount of surface oxygen defect sites and improved catalytic activity for propane oxidation [12]. However, the influence of catalyst precursor in this study identifies the varying surface areas as the most influential factor when determining catalytic activity.

As mentioned previously, Na is known to be a poison of metal oxide catalysts for certain oxidation reactions [43]. This factor is particularly important when considering catalytic activity for the total oxidation of propane, as Tang et al. demonstrated that Na greatly hinders this process [54]. It was proposed that Na negatively impacts oxygen mobility by supressing oxygen desorption at lower temperatures, in addition to accumulating high amounts of surface carbonate species at higher temperatures. The impact of poor oxygen mobility from Na poisoning is consistent with extended studies for the oxidation of propane [55]. The impact of increased Na content on the catalyst surface did not impact the oxidation of propane in this work.

The most important factor influencing propane total oxidation activity was the catalyst surface areas, as a relationship can be concluded from data presented in Table 4. Hence, it is important to highlight the influence that the catalyst precursor has on the final catalyst, as a relatively subtle change of the cerium precursor oxidation state is significant. As shown from TGA-DTA and XRD analysis, the use of different precursors results in different precipitation products which, on calcination, form cubic fluorite CeO$_2$ with subtle differences that are important for catalytic performance.

The formation of a CeO$_2$·nH$_2$O precipitate greatly increases the surface area of the resulting catalyst as evidenced by the CeO$_2$ (IV) catalyst. This effect has also been shown in previous research for the synthesis of Sm-doped CeO$_2$ mixed oxide catalysts [56]. Fur-
thermore, studies comparing either the cerium precursor or precipitating agent have drawn similar conclusions, with the Ce\textsuperscript{IV} precursor or the CeO\textsubscript{2}.H\textsubscript{2}O precipitate resulting in the highest surface area [24,57]. Transformation of the CeO\textsubscript{2}.H\textsubscript{2}O precipitate to the CeO\textsubscript{2} catalyst is likely to be topotactic, with very little alteration to the crystal structure from the removal of H\textsubscript{2}O or \textsuperscript{-}OH species. In contrast, the structure of the orthorhombic Ce\textsubscript{2}(OH)\textsubscript{2}(CO\textsubscript{3})\textsubscript{2}.H\textsubscript{2}O precipitate differs significantly from the cubic fluorite structure of CeO\textsubscript{2}, and decomposition occurs via an intermediate oxycarbonate species [30,58]. This disruptive decomposition phase transformation mechanism will result in the decreased surface area shown by the CeO\textsubscript{2} (III) catalyst. Furthermore, the Ce\textsubscript{2}(OH)\textsubscript{2}(CO\textsubscript{3})\textsubscript{2}.H\textsubscript{2}O precipitate contains cerium in the Ce\textsuperscript{3+} oxidation state, requiring an oxidation step to produce the CeO\textsubscript{2} catalyst. Spiridigliozzi et al. found that under hydrothermal conditions, the initially formed metastable Ce(OH)(CO\textsubscript{3}) precipitate undergoes structural changes, as well as oxidation in solution after a long period of time [59]. However, the decomposition of the Ce(OH)(CO\textsubscript{3}) precipitate by heat treatment also induces the oxidation of the Ce\textsuperscript{3+} species to Ce\textsuperscript{4+}. It is stated that the thermal decomposition of the Ce(OH)(CO\textsubscript{3}).H\textsubscript{2}O species results in a combination of endothermic decomposition from dehydration and carbonate decomposition, in addition to an exothermic reaction from the oxidation of Ce\textsuperscript{3+} to Ce\textsuperscript{4+} [30]. This exothermic reaction could increase the driving force to form the thermodynamically stable CeO\textsubscript{2} structure with the increased Ce\textsuperscript{4+} surface concentration observed for the lower activity CeO\textsubscript{2} (III) catalyst.

3. Materials and Methods

3.1. Catalyst Preparation

Two CeO\textsubscript{2} catalysts were prepared by a precipitation method, utilising a Metrohm 902 Titrandt auto-titrator (Metrohm, Cheshire, UK). Aqueous (NH\textsubscript{4})\textsubscript{2}Ce(NO\textsubscript{3})\textsubscript{6} (Acros Organics, Geel, Belgium) and Ce(NO\textsubscript{3})\textsubscript{3} (Merck, Gillingham, UK) were used as Ce\textsuperscript{IV} and Ce\textsuperscript{III} sources respectively, with aqueous Na\textsubscript{2}CO\textsubscript{3} (Merck, Gillingham, UK) as the precipitating agent. In a typical synthesis, 100 mL of nitrate solution (0.25 M) and a solution of Na\textsubscript{2}CO\textsubscript{3} (1 M) were added simultaneously to a thermostatically water heated jacketed vessel (80 °C). Addition rate of the nitrate solution was maintained constant (2 mL min\textsuperscript{-1}), whilst addition of Na\textsubscript{2}CO\textsubscript{3} solution was automatically computer controlled to maintain a constant pH of 9. The precipitate formed was held at 80 °C for 1 h before being collected by vacuum filtration and washed with 2 L of hot deionised water.

The precipitates were dried at 110 °C for 16 h. Dried catalyst precipitates prior to calcination were denoted as d-CeO\textsubscript{2} (IV) and d-CeO\textsubscript{2} (III). Samples denoted as CeO\textsubscript{2} (IV) and CeO\textsubscript{2} (III) were the final catalysts prepared by calcination of the precipitates in static air at 500 °C for 3 h, with a ramp rate of 5 °C min\textsuperscript{-1}.

3.2. Catalyst Characterisation

Thermal gravimetric analysis (TGA) and differential thermal analysis (DTA) were performed using a Setaram Labsys 1600 instrument (SCIMED, Cheshire, UK). Approximately 30 mg of sample was loaded into an alumina crucible and heated to 700 °C at a rate of 5 °C min\textsuperscript{-1} in a flow of synthetic air (50 mL min\textsuperscript{-1}). For all TGA-DTA analysis, blank runs were subtracted from the relevant data to account for any buoyancy effects.

Powder X-ray diffraction (XRD) patterns were collected using a Panalytical X’Pert diffractometer (Malvern Panalytical, Worcestershire, UK) equipped with a Cu X-ray source operating at 40 kV and 40 mA. Analysis was carried out between 20 values of 5–80°. Phase identification was achieved by matching patterns against the ICDD standard database. Application of the Scherrer equation was used to estimate crystallite size, comparing the experimental line widths of the four most dominant reflections ((111), (200), (220), (311)) against a highly crystalline silicon standard. An average crystallite size was then determined, with the (111) diffraction peak used to calculate lattice parameters.

A Quantachrome Quadsorb Evo Analyser (Quantachrome, Hook, UK) was used for surface area analysis. Prior to analysis, catalysts were degassed under vacuum for 16 h at
Surface areas of the catalysts were determined from twenty-point N\textsubscript{2} adsorption–desorption isotherms measured at \(-196\) °C. The Brunauer–Emmett–Teller (BET) method was used to treat the data.

Raman spectra were obtained using a Renishaw inVia confocal Raman microscope (Renishaw, Gloucestershire, UK) with an Ar\textsuperscript{+} visible green laser (514 nm). Spectra were collected in a reflective mode from samples mounted on a steel holder by a highly sensitive charge couple device (CCD) detector. The defect ratio (A\textsubscript{590}/A\textsubscript{463}) was calculated using the area of the bands at 590 cm\textsuperscript{-1} and 463 cm\textsuperscript{-1}.

Temperature programmed reduction and oxidation (TPR, TPO) were performed using a Quantachrome ChemBET (Quantachrome, Hook, UK). Pre-treatment of catalysts were carried out under a flow of He for 1 h at 120 °C. Reduction profiles were obtained by analysing approximately 50 mg of catalyst under a flow of 10% H\textsubscript{2}/Ar (50 mL min\textsuperscript{-1}), over the temperature range 50–700 °C, with a heating rate of 10 °C min\textsuperscript{-1}. H\textsubscript{2} consumption was calculated by calibration against a CuO standard.

TPR-TPO cycles were run using the same conditions stated above. Once the initial TPR was completed, the catalyst sample was cooled under flowing He. TPO profiles were then attained under a flow of 10% O\textsubscript{2}/He (50 mL min\textsuperscript{-1}), over the temperature range 50–700 °C, with a heating rate of 10 °C min\textsuperscript{-1}. Subsequently, a second TPR was completed.

A Kratos Axis Ultra DLD system (Kratos Analytical, Manchester, UK) was used for X-ray photoelectron spectroscopy (XPS). Spectra were collected using a monochromatic Al K\textsubscript{α} X-ray source operating at 140 W (10 mA and 14 kV). Pass energies of 160 eV for survey spectra and 20 eV for the high-resolution scans were employed for data collection, with step sizes of 1 eV and 0.1 eV respectively. The system was operated in the Hybrid mode, utilising both magnetic immersion and electrostatic lenses for high sensitivity. Spectra were acquired using the ‘slot’ aperture which defines an analysis area of approximately 700 × 300 µm\textsuperscript{2}. Sample surface charging was minimised by a magnetically confined low energy electron charge compensation system, with all spectra taken at a 90° angle. A base pressure of ca. 1 × 10\textsuperscript{-9} Torr was maintained during data collection. Data was calibrated to the C 1s line of adventitious carbon (248.8 eV) and analysed using CasaXPS v2.3.24 [60] after subtraction of a Shirley background and using modified Wagner sensitivity factors, which take in to account the electron escape depth, as supplied by the manufacturer. Peak fits were performed using Voigt type functions and where applicable, modelled on line shapes derived from bulk standards, such as stoichiometric CeO\textsubscript{2}.

Scanning Electron Microscopy (SEM) was obtained from a Tescan MAIA3 field emission gun scanning electron microscope (Tescan, Cambridge, UK) (FEG-SEM) with secondary and backscattered electron detectors. Energy-dispersive X-ray (EDX) analysis was performed using an Oxford Instruments XMax\textsuperscript{N} 80 detector (Oxford Instruments, Abingdon, UK). The Point and ID function on the Oxford Aztec software was used to perform EDX analysis. Catalysts loaded onto carbon tape were sputter coated with 15 nm Au/Pd to prevent charging. A minimum of three areas were analysed across multiple particles and averaged to produce atomic %.

Transmission electron microscopy (TEM) was performed using a JEOL JEM 2100 (JEOL UK, Welwyn Garden City, UK) operating at 200 kV. Samples were prepared by dry dispersion over a 300 mesh copper grid coated with holey carbon film.

### 3.3. Catalyst Testing

Catalyst performance for the total oxidation of propane was assessed using a continuous flow fixed bed microreactor. A constant volume of catalyst was secured in a \(\frac{1}{8}\) inch stainless steel tube between two plugs of quartz wool. A premixed cylinder of 5000 ppm propane in air was used, with the gas flow regulated at 50 mL min\textsuperscript{-1} by electronic mass flow controllers. A water saturator was attached and held at constant temperature to produce a 5% water saturation in the gas feed. A constant powdered catalyst volume of 0.067 mL was used, resulting in a catalyst mass of approximately 0.1 g and 0.3 g for CeO\textsubscript{2} (III) and CeO\textsubscript{2} (IV) respectively, to give a gas hourly space velocity (GHSV) of 45,000 h\textsuperscript{-1}.
The temperature range 200–600 °C was used to measure catalyst activity, with the temperature maintained and controlled by a K-type thermocouple placed into the catalyst bed. The reactor temperature was increased incrementally and allowed to stabilise until steady state was attained at each temperature before analysing the reactor effluent. The reaction effluent was analysed by an online gas chromatograph (Agilent 7890B) with two detectors in series (Agilent Technologies LDA UK Ltd., Stockport, Cheshire, UK). A thermal conductivity detector (TCD) was used to analyse O₂ and N₂. In addition, a flame ionisation detector (FID) equipped with a methaniser was used to analyse CO, CO₂, and hydrocarbons. Using HayeSep Q (80–100 mesh, 1.8 m × 3.2 mm) and MolSieve 13 X (80–100 mesh, 2 m × 3.2 mm) packed columns, with a series/by-pass valving configuration, separation was achieved. Analyses were performed at each temperature until three consistent sets of analytical data were obtained, indicating steady-state operation. Propane conversion was calculated by comparing the measured counts at each temperature to the blank analysis before each test.

4. Conclusions

CeO₂ catalysts were prepared by a precipitation method using either Ce³⁺ or Ce⁴⁺ nitrate precursors to determine the influence on the catalytic activity for propane total oxidation. The CeO₂ catalyst prepared using the Ce⁴⁺ precursor had improved performance for propane total oxidation. The catalytic activity was related to the formation of different precipitated species during synthesis of the catalyst precursor and their transformation to the CeO₂ catalyst. Formation of CeO₂·nH₂O from the Ce⁴⁺ precursor salt resulted in a higher surface area catalyst with increased concentration of surface oxygen defect sites and reduced cerium species. Formation of the Ce₂(OH)₂(CO₃)₂·H₂O precipitate from the Ce³⁺ precursor salt resulted in slightly improved redox properties but had a very low surface area. A direct relationship between catalyst surface area and catalytic activity was found to be the most influential factor for the total oxidation of propane.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/catal11121461/s1, Figure S1: Catalyst activity for propane total oxidation under wet and dry conditions.

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