

Showcasing research from Professor Hess's laboratory, Department of Chemistry, Technical University of Darmstadt, Germany.

Controlling the dispersion of ceria using nanoconfinement: application to $CeO₂/SBA-15$ catalysts for $NH₃-SCR$

Loading the active phase of catalytic materials in a highly dispersed fashion onto porous supports is still a great challenge. As illustrated for ceria-based catalysts, we show that template P123 can be used to control the crystal growth, with direct implications for the catalytic performance. To elucidate the detailed mechanism of this template-assisted synthesis of SBA-15 supported ceria we have applied *in situ* spectroscopies. On this basis, a rational catalyst synthesis was developed by exploiting nanoconfinement effects, resulting in an improved ceria dispersion and superior NH₃-SCR performance.

rsc.li/materials-advances

Materials Advances

PAPER

Cite this: *Mater. Adv.*, 2021, 2, 7400

Received 28th July 2021, Accepted 7th September 2021

DOI: 10.1039/d1ma00658d

rsc.li/materials-advances

Introduction

In the past decade, mesoporous materials have attracted great attention due to their high specific surface area, large pore volume, and controllable pore sizes between 2 and 50 nm (according to the IUPAC definition) with extensive applications in gas adsorption, drug delivery, and catalysis. 1 Since the first synthesis of mesoporous SBA-15 by Zhao et $al.$ ², the enlarged pore sizes (4.6–30 nm) have allowed larger molecules to enter the pore system, and have also decreased the mass transfer limitations compared to conventional zeolites and microporous molecular sieves.3,4 Due to their inert behaviour in most catalytic reactions, mesoporous silicates are promising support materials for the incorporation of various active compositions, metals or metal oxides, exhibiting a great potential for heterogeneous catalysis.⁵⁻⁸

At present, synthetic approaches leading to catalytically active mesoporous silicates are divided into two classes, the "one-pot" strategy and the "loading" strategy.⁹ In the case of the ''one-pot'' strategy (or pre-synthesis method), the silicon source is mixed with the precursor of the active composition,

Controlling the dispersion of ceria using nanoconfinement: application to $CeO₂/SBA-15$ catalysts for NH₃-SCR⁺

Jun Shen **D** and Christian Hess **D** *

Mesoporous silicas, such as SBA-15, with their high and inert surface area, constitute promising support materials for catalytic applications. However, loading the active phase onto porous materials to give high dispersion and even distribution is still a great challenge. In the presence of the template P123, ceria crystal growth can be controlled by nanoconfinement effects. We have applied in situ Raman, DRIFT, and DR UV-vis spectroscopies to elucidate the mechanism of template-assisted synthesis of ceria/SBA-15 catalysts for NH₃-SCR applications. The co-decomposition of P123 and cerium precursor shows a catalytic interaction, resulting in a change of ceria nucleation and growth behavior during the calcination process. Based on this knowledge, we have developed an improved catalyst synthesis by combining the nanoconfinement effects of P123 and the use of an inert atmosphere, resulting in an improved ceria dispersion on SBA-15 and superior NH₃-SCR performance. Our results highlight the significance of the calcination process for defining catalyst properties and the use of in situ spectroscopy for rational synthesis of supported catalysts. PAPER
 EXECUTE:
 **CONTrolling the dispersion of ceria using

CONFORMITY CONTICUTE CONTICUTE CONTICUTE CONTICUTE CONTICUTE

CARGO CONTICUTE CONTICUTE CONTICUTE CONTICUTE CONTICUTE

CARGO CONTICUTE CONTICUTE CONTICUTE CON**

which then both interact with the template to co-assemble into a porous framework.^{10–12} This method has the advantage of being a simple operation and the formation of an atomic-scale mixture of active sites in the porous materials, while the additional heteroatom in the framework may destroy the ordered mesoporous structure, leading to further collapses during calcination. The ''loading'' strategy (or post-synthesis method) mainly involves the preparation of a stable mesoporous silica matrix, followed by its loading with the active composition.13,14 The bottleneck of this method is the distribution of active species (e.g. nanoclusters of metals or metal oxides) inside the long pore channel, which may be susceptible to sintering effects at high-temperature calcination.¹⁵ However, one may envision an intermediate between the pre- and postsynthesis approaches, denoted med-synthesis, in which the template remains in the pore (see Scheme 1) and which,

Scheme 1 Synthesis approaches for mesoporous silicate materials (e.g. SBA-15) modified by the addition of an active component at different stages.

Eduard-Zintl-Institut für Anorganische und Physikalische Chemie, Technical University of Darmstadt, Alarich-Weiss-Str. 8, 64287 Darmstadt, Germany. E-mail: jun.shen@tu-darmstadt.de, christian.hess@tu-darmstadt.de

[†] Electronic supplementary information (ESI) available: Additional images and characterization. See DOI: 10.1039/d1ma00658d

although largely ignored for a long time, has the potential to combine the advantages of the pre- and post-synthesis approaches, such as good active phase distribution and defined structure.

Recently, the positive effect of surfactants on the loading of active components in mesoporous silicates has attracted considerable attention. To this end, previous studies have shown that 3 nm-sized ceria particles dispersed on SBA-15, as achieved with the assistance of H_3 Cit, caused an enhanced reducibility and defect concentration of the particles, influencing the catalytic performance.¹⁶ Besides, surfactant-assisted impregnation has been widely applied in the context of Ni/SBA-15 catalysts for $CO₂$ reforming of $CH₄$.¹⁷ These studies have revealed that the use of surfactant P123 yields a catalytic performance superior to that of other surfactants, $17-20$ which was explained by the coordination of Ni atoms by the OH groups of P123, thus preventing their aggregation.¹⁹ Considering that P123 is the template of SBA-15, a method was put forward that used as-made SBA-15 for wetness impregnation after partial removal of P123. This method was employed for the synthesis of Cu/ SBA-15 materials, enabling a good dispersion of copper even at high loadings.^{21,22} Yin et al. suggested an improved method to use the P123-contained SBA-15 combined with solid-state impregnation methods to avoid the extra step of partial removal of P123, which was applied to the synthesis of CuO/SBA-15 for thiophene capture,^{23,24} CeO₂/SBA-15 for sulphur capture,^{25,26} Pt/SBA-15 for hydrogen storage, 27 Co/SBA-15 for the oxidation of organic pollutants,²⁸ and Fe/SBA-15 for phenol degradation.²⁹ Also composite catalysts, i.e., Ag-CeO₂/SBA-15, were prepared by using P123-contained SBA-15, showing good dispersion and catalytic performance for room-temperature reduction of 4-nitrophenol.³⁰ Summarizing, previous studies have demonstrated the great potential of the template P123-assisted synthesis of mesoporous catalysts containing highly dispersed metals or metal oxides. Puper

Moreles the commission of the pre- article points of the particles are proposed points and points are article is a supported to the common access Article is a supported to the common access Article is licensed unde

Although the med-synthesis approach towards improved supported catalysts has received increasing attention, a detailed mechanistic understanding of the template P123-assisted synthesis is still lacking. One hypothesis is that P123 remaining in the pores of SBA-15 forms a confined space between the silica walls and residual P123 in the intra-wall pores, $22,25-29$ while another proposes an interaction between P123 and the metal precursors, influencing their decomposition behavior.¹⁹ However, due to lack of direct experimental evidence, particularly in situ analysis, none of the above hypotheses have been verified so far. Besides, it was found that the dispersion effect of P123 was limited to samples with high loadings.²⁶ Thus, elucidating the mechanism of the template assistance will be of great importance to further improve its applicability to catalyst synthesis.

The selective catalytic reduction (SCR) of NO with $NH₃$ is an essential process to reduce the emission of NO_x gas, which is known to be harmful to the global environment and human health.³¹ Ceria-based materials have been considered as potential catalysts for the NH3-SCR reaction, due to their high oxygen storage capacity (OSC) and excellent redox properties. 32 Since crystalline $CeO₂$ cannot provide a sufficiently high

specific surface area, especially after high-temperature aggregation, SBA-15 has been employed previously as a support material to disperse and stabilize $CeO₂$ particles,³³ but also other $NH₃$ -SCR-related materials.³⁴

In this contribution, we report how $CeO₂/SBA-15$ catalysts were prepared by solid-state impregnation methods starting from bare and template-containing SBA-15, aiming at a mechanistic understanding of the synthesis. To monitor the calcination process in detail, in situ Raman, DRIFT, and DR UV-vis spectroscopies were applied, revealing details of the transformation of the cerium precursor to the supported oxide. Additional thermogravimetric analysis (TGA) and exhaust gas analysis by FT-IR spectroscopy allowed the temperaturedependent decomposition of the P123 template and the cerium precursors to be monitored. The differently prepared samples were characterized with regard to their structural and surface properties and tested in $NH₃$ -SCR. Based on these combined results, a synthesis mechanism was proposed and used to develop a novel synthesis protocol to yield catalysts with improved ceria dispersion and superior $NH₃$ -SCR performance.

Results and discussion

1. Characterization of dispersed $CeO₂$ on SBA-15

Three $CeO₂/SBA-15$ samples were prepared by calcination of the cerium oxide precursor Ce($\rm NO_3)_3$ ·6H₂O and the SBA-15 support following different routes, as detailed in Scheme 2. The sample tfSBA–CeO₂–air was synthesized by grinding template-free SBA-15 (tfSBA-15) with cerium nitrate and subsequent calcination in air to obtain ceria crystallites grown inside SBA-15 pores. For the sample asSBA–CeO₂–air, P123-containing SBA-15 (asSBA-15) was employed as support material. Using the same calcination step in air results in ceria dispersed inside the SBA-15 pores. Moreover, a third route was designed, which yields the sample asSBA–CeO₂–N₂, by starting from asSBA-15 but changing the gas atmosphere to inert N_2 .

Fig. 1a summarizes the temperature program of the calcination process, which is characterized by two stages, i.e., heating from room temperature to 500 $^{\circ}$ C at a rate of 1.5 $^{\circ}$ C min⁻¹ and keeping the temperature at 500 $^{\circ}$ C for 5 h.

Scheme 2 Different synthesis routes for CeO₂/SBA-15 catalysts using the solid-state impregnation method.

Fig. 1 (a) Temperature program of the calcination process, and (b) XRD patterns of the calcined CeO₂/SBA-15 materials, after start-ing from different SBA-15 matrixes (tfSBA-15 and asSBA-15) and cerium contents (low and high), and applying different atmospheres (air and N₂). For comparison the XRD pattern of bare SBA-15 is shown. The CeO₂-related peaks are marked. XRD patterns are offset for clarity. (c) TEM images of SBA-15 (i), tfSBA-L-CeO₂-air (ii), asSBA–L-CeO₂–air (iii), and asSBA–L-CeO₂–N₂ (iv).

Table 1 Composition, specific surface area, and porosity characteristics of the CeO₂/SBA-15 samples compared to bare SBA-15

Samples	$CeO2$ (wt%)	S_{total}^a [m ² g ⁻¹]	$D_{\rm p}{}^b$ [nm]	$_{10}$ ^c [cm ³ g ⁻¹] $V_{\text{micro}}^{\prime}$	\int_1^d [cm ³ g ⁻¹] V_{total}	$\frac{e}{2}$ [nm] $d_{\rm CeO}$
$tfSBA-15$		616	6.3	0.14	0.64	
t fSBA-L-CeO ₂ -air	39	414	6.3	0.06	0.43	7.8
$asSBA-L-CeO2-air$	39	359	6.3	0.03	0.39	6.4
$asSBA-L-CeO2-N2$	39	280	5.8	0.02	0.32	5.4

^a Total BET surface area. ^b Pore diameter determined from the adsorption branch by NLDFT. ^c Micropore volume. ^d Total pore volume. ^e Calculated from XRD results using the Scherrer formula.

Fig. 1b shows wide-angle XRD patterns of the calcined samples. Starting with bare SBA-15, the broad diffraction peak at $2\theta = 23^\circ$ is assigned to amorphous silica. For the samples with CeO₂ loading, new peaks appear at 2θ values of 28.5° , 33.2° , 47.5° , and 56.3° , which can be attributed to different planes of crystalline CeO₂, *i.e.*, (111), (200), (220), and (311), respectively. Detailed analysis reveals that the peaks show a full width at half maximum (FWHM) in the order tfSBA-L-CeO₂air \langle asSBA–L-CeO₂–air \langle asSBA–L-CeO₂–N₂. Based on a fit analysis of the ceria (111) diffraction peak and application of the Scherrer formula,³⁵ the crystallite sizes are found to decrease as tfSBA–L-CeO₂–air > asSBA–L-CeO₂–air > asSBA– L-CeO₂–N₂ (see Table 1), reflecting the suppression of ceria crystal growth when the use of template P123 is combined with a N_2 calcination atmosphere. Similarly, with increasing $CeO₂$

loading, the crystallite sizes decreased as as SBA-H-CeO₂-air $>$ asSBA–H-CeO₂–N₂, indicating that an inert calcination atmosphere further improves the $CeO₂$ dispersion compared to the effect of the surfactant template alone.

Fig. 1c shows TEM bright field images of the calcined samples. The SBA-15 sample (i) has a regular morphology and ordered structure. For the samples tfSBA-L-CeO₂-air (ii), asSBA–L-CeO₂–air (iii), and asSBA–L-CeO₂–N₂ (iv) CeO₂ nanoparticles with decreasing size are observed outside the SBA-15 matrix, showing that similar to XRD TEM detects larger particles but not the highly dispersed ceria inside the pores.

Table 1 summarizes some physical properties of the $CeO₂/$ SBA-15 samples as determined on the basis of N_2 adsorption isotherms, including the specific surface areas, the micropore and total pore volumes, and the pore sizes. The $CeO₂$ -containing

Fig. 2 (a) N₂ adsorption/desorption isotherms, and (b) NLDFT pore size distributions of template-free SBA-15 and CeO₂/SBA-15 materials prepared by different synthesis routes.

samples exhibit smaller specific surface areas than bare templatefree SBA-15 (tfSBA-15), decreasing in the order tfSBA-L-CeO₂-air > asSBA–L-CeO₂–air > asSBA–L-CeO₂–N₂, owing to micropore blocking and, in the case of as SBA–L-CeO₂–N₂, also the reduction of mesopore diameter by ceria loading onto the inner surface of the porous matrix.

The N_2 adsorption/desorption isotherms of the SBA-15 and $CeO₂/SBA-15$ samples all show profiles typical for mesoporous materials (see Fig. 2a), but with variations in the shape of the hysteresis loop. Bare SBA-15 powder shows the typical shape of a mesoporous material with an H1 hysteresis loop at the relative pressure p/p° of 0.55–0.75, reflecting uniform hexagonal pores with a narrow pore size distribution centred around 6.3 nm diameter. For samples loaded with $CeO₂$, a decrease in the micropore area compared to bare SBA-15 is observed. Also, the closing point of the hysteresis loop shifts to lower relative pressure values of about 0.45, which implies that $CeO₂$ is formed inside the pores, resulting in open and narrowed mesopores.³⁶ The desorption branch of the hysteresis loop of tf -SBA-L-CeO₂-air shows two steps, representing open mesopores (p/p° of 0.53–0.70) and narrowed mesopores (p/p° of 0.45– 0.53), respectively. For as SBA-L-CeO₂-air, a weak feature from open mesopores and a forced closure of the hysteresis loop at $p/p^{\circ} = 0.46$ is observed, which is attributed to ink-bottle type pores. In contrast, the desorption branch of asSBA-L-CeO₂-N₂ is characterized by a smoother decrease, which is somewhat similar to that of SBA-15, representing a more uniform pore size along the channels, while the shift of the curve to lower relative pressure values compared to bare SBA-15 results from the decreased pore size after the coating with $CeO₂$.

Fig. 2b shows the pore size distributions calculated from the adsorption branches of the isotherms. The samples treated in air, tfSBA-L-CeO₂-air and asSBA-L-CeO₂-air, exhibit a maximum at a pore size of around 6.3 nm, *i.e.*, at the same diameter as the original SBA-15, implying that a considerable amount of surface area inside the pores is uncoated. Especially the pore

sizes of tfSBA–L-CeO₂–air are distributed over a wider range towards lower values, suggesting the presence of $CeO₂$ crystallites inside the pores. In contrast, as SBA-L-CeO₂-N₂ shows a maximum at a decreased pore diameter of 5.8 nm, as a result of a more even coating of the pores with nano-scale $CeO₂$. For illustration, Fig. S1 in the ESI† provides schematic diagrams of the pore structures of these three samples, as proposed on the basis of the above analysis. In summary, for tfSBA-L-CeO₂-air, the growth of crystalline ceria results in the presence of narrowed pores, while in the case of as SBA–L-CeO₂–air the limited diffusion of the cerium precursor leads to the formation of ink-bottle pores. By contrast, for as SBA–L-CeO₂–N₂, extensive diffusion of cerium ions results in an evenly coated porous matrix.

2. Mechanism of $CeO₂$ dispersion during calcination

Previously, the good dispersion on asSBA-15 has been proposed to be due to the existence of template P123, resulting in confined spaces for $CeO₂$ crystal growth, however, without providing direct evidence. To elucidate the interaction between the SBA-15 support and the ceria species during calcination, TG analysis and online gas-phase FTIR spectroscopy, as well as in situ Raman, DR UV-vis, and DRIFT spectroscopies were applied.

First, we applied TG analysis to gain detailed insight into the decomposition of the template-containing SBA-15 during calcination, compared to bare asSBA-15 and the cerium precursor $Ce(NO₃)₃·6H₂O$ as reference (see Fig. 3a), following the temperature program given in Fig. 1a. The observed thermal decomposition of bare as-made SBA-15 shows a single weight loss of 47 wt% at about 120 \degree C, which is associated with the decomposition of triblock copolymer P123 slightly embedded in the siliceous pore walls.³⁷ Previous studies have addressed the decomposition mechanism of bare cerium nitrate $\text{Ce}(\text{NO}_3)_3$ -6H₂O in air by TGA, as summarized in Fig. S2 $(ESI[†])^{38,39}$ Here, by combining TG and DTG analysis (see Fig. 3a and Fig. S3, ESI†), we

Fig. 3 TGA profiles (a), and IR intensities of exhaust NO₂ (b), and CO₂ (c) during calcination of bare asSBA-15, bare cerium nitrate, and cerium nitrate/ SBA-15 mixtures, following the protocol given in Fig. 1a.

identified 221 $^{\circ}$ C as a critical temperature point, attributed to the onset of cerium nitrate decomposition. Weight losses observed below 221 \degree C are assigned to the release of surface water and structural water, while at temperatures above 221 $^{\circ}$ C the decomposition of nitrate resulted in the release of NO_x gas (see below, Fig. 3b) and a further substantial weight loss of 35 wt%. Sample $tfsBA-L-CeO₂$ calcined in air shows a substantial decrease in weight below 200 \degree C due to lower onset temperatures for the release of water and cerium nitrate decomposition compared to bare cerium nitrate.

However, because, for the mixed samples (cerium nitrate/ SBA-15), the weight loss reflects a more complex interplay of different processes, the exhaust was continuously monitored by gas-phase FTIR spectroscopy. To this end, Fig. 3b depicts the temperature-programmed evolution of NO₂ gas (at 1629 $\rm cm^{-1})^{40}$ resulting from cerium nitrate decomposition. As a reference, the $\rm NO_2$ evolution of bare Ce($\rm NO_3)_3$ ·6H $_2$ O salt is shown (see light-blue curve), which is characterized by a sharp increase at 240 $^{\circ}$ C and a slower decline up to \sim 500 °C, in agreement with the literature.³⁹ There is an additional small peak at about 120 \degree C, indicating the evaporation of aqueous acid azeotrope, as discussed in the context of Fig. S2 (ESI†). Compared to the bare cerium salt, the $NO₂$ signal of tfSBA–L-CeO₂–air appears within a much narrower temperature range, suggesting the influence of SBA-15 on the decomposition of cerium nitrate. The lack of $NO₂$ features at lower temperatures, *i.e.*, at around 120 °C and at 190–230 °C, suggests the inhibition of azeotrope formation by dry SBA-15. In contrast, the sample asSBA–L-CeO₂–air shows an increasing signal at $160-250$ °C, which is attributed to template P123 decomposition, producing water to form the $HNO₃$ azeotrope, as well as a stronger nitrate decomposition peak with an onset at \sim 250 °C. For asSBA-L- $CeO₂-N₂$, the cerium nitrate precursor is observed to decompose within a very narrow temperature window (250 $^{\circ}$ C–300 $^{\circ}$ C), indicating differences in the interaction between precursor and template as a function of the calcination conditions.

Fig. 3c depicts the temperature-programmed evolution of exhaust CO₂ gas (at 2350 $\rm cm^{-1})^{40}$ resulting from the decomposition of template P123. For bare as-made SBA-15, the

decomposition starts at around 155 °C and extends up to 500 °C. Sample tfSBA15-L-CeO₂-air shows a small feature at 240–350 \degree C, originating from residual P123 after calcination. By comparison of Fig. 3b and c, it can be seen that the temperature range of the CO_2 -related band overlaps with that of the $NO₂$ band, possibly leading to synergetic effects in the decomposition and oxide formation processes. The sample asSBA–L-CeO₂–air exhibits several CO_2 -related peaks. Compared to the single broad $CO₂$ band detected for bare asSBA-15, these more discrete features indicate the presence of different intermediate states during P123 decomposition. In contrast, when the calcination is performed in N_2 , a significant amount of CO_2 is released within a very narrow temperature window $(270-275 \text{ °C})$, strongly overlapping with that of the $NO₂$ release (see Fig. 3b), reflecting an interaction between the decomposition of P123 and that of cerium nitrate, while the $CO₂$ emission at higher temperatures can be ascribed to the decomposition of oxygen-containing fragments of the template.

To gain more detailed insight into the decomposition processes, Fig. 4 shows the temporal evolution of all exhaust gases during calcination, as detected by online gas-phase IR analysis. Features at around 1150-1250 cm^{-1} are assigned to C-O-C stretching modes,⁴¹ those at 1350 cm⁻¹ and 2800-3000 cm⁻¹ to C-H bending and stretching modes, respectively.⁴¹ and those at 1750 and 3638 cm^{-1} to C=O and O-H stretching modes, respectively.⁴⁰

For bare asSBA-15, the P123 decomposes directly to gaseous $CO₂$ as well as C1 or C2 fragments, starting at around 170 °C. In the case of tfSBA-L-CeO₂-air, the decomposition of cerium nitrate, leading to $NO₂$ formation, and small amounts of $CO₂$ and C_n ketone are observed from 260 °C onwards. The exhaust gas evolution of asSBA-L-CeO₂-air is characterized by four stages, beginning at about 160 (I), 240 (II), 300 (III), and 350 °C (IV), respectively, consistent with the discrete $CO₂$ peaks in Fig. 3c: In stage I, C_n fragments (including ketone, alcohol, and ether) are formed, while $NO₂$ shows a gradual increase, indicating that P123 fragments may affect the decomposition of cerium nitrate, considering the higher onset temperature for

Fig. 4 Online IR detection of exhaust gases during calcination of the samples (a) asSBA15-air, (b) tfSBA-L-CeO₂-air, (c) asSBA-L-CeO₂-air, and (d) asSBA–L-CeO₂–N₂. The low-temperature features at around 1600 and 3750 cm⁻¹ can be attributed to water released from cerium nitrate. For details see text.

decomposition observed for tfSBA-L-CeO₂-air. In stage II, which corresponds to the temperature range of the original cerium nitrate decomposition, a facilitated decomposition of P123 compared to bare asSBA-15 is observed, which may be attributed to a catalytic effect by the formed ceria (see below). In stage III, most of the residual P123 is catalytically converted by ceria to $CO₂$ and $H₂O$, while in stage IV more stable fragments are decomposed.

Summarizing, from the above in situ gas-phase results we conclude that P123 fragments facilitate the decomposition of cerium nitrate, while in turn the formed ceria catalyses the decomposition of P123. Interestingly, for sample asSBA-L-CeO₂- N_2 , P123 and cerium nitrate decompose almost simultaneously within a narrow temperature window at about 280 \degree C, which equals the decomposition temperature of cerium nitrate, according to the tfSBA–L-CeO₂–air and asSBA–L-CeO₂–air results. Thus, we can conclude that the inert atmosphere inhibits the decomposition of P123 at low temperatures (see also Fig. S4, ESI†), allowing an increased catalytic interaction between cerium nitrate and P123 at elevated temperature, boosting the nucleation of nano ceria.

The in situ Raman spectra of tfSBA–L-CeO₂–air, asSBA–L- $CeO₂$ –air, and asSBA–L-CeO₂–N₂ during calcination are mainly characterized by Raman features at 457, 745, and 1046 cm^{-1} (see Fig. S5, ESI†). The band at about 460 cm^{-1} is assigned to the triply degenerate symmetrical stretching mode (F_{2g}) of $ceria,$ ^{42,43} and its position and shape are known to depend on the detailed ceria lattice structure.⁴⁴⁻⁴⁶ The Raman features at 745 cm^{-1} and 1046 cm^{-1} originate from bending and stretching vibrations of free nitrate, respectively.⁴⁷ Fig. 5 compares the temperature-dependent transformation of cerium nitrate into ceria for the three samples, focusing on the nitrate stretching mode at 1046 cm^{-1} to monitor the precursor decomposition (a) and the F_{2g} mode at 457 cm⁻¹ to describe formation of crystalline ceria (b). For tfSBA–L-CeO₂–air, the cerium nitrate features disappear at 300 $^{\circ}$ C, while those for ceria start to appear at 250 \degree C, implying simultaneous cerium nitrate decomposition and ceria formation. Initially, a broad F_{2g} -related band is detected at about 450 cm^{-1} , which undergoes a narrowing and red-shift to 457 cm^{-1} with increasing temperature, thus indicating the growth of ceria crystallites (see Fig. S5a, ESI†).

Fig. 5 Temperature-dependent Raman intensities (514.5 nm) at 1046 cm⁻¹, representing cerium nitrate (a), and at 457 cm⁻¹, representing CeO₂ (b), recorded during calcination, following the protocol given in Fig. 1a. The corresponding Raman spectra are shown in Fig. S5 (ESI†).

The additional smaller features at about 250 and 279 cm^{-1} are attributed to nitro- and ceria-related vibrations and discussed in the context of Fig. S5 (ESI†). In the case of the asSBA–L-CeO₂ samples, temperature windows of about 250–300 $\mathrm{^{\circ}C}$ (in air) and 250–450 °C (in N_2) were detected. Sample asSBA–L-CeO₂–air shows the F_{2g} band only at about 300 °C, suggesting an inhibited ceria growth process. In contrast, in the case of asSBA– L-CeO₂-N₂, a F_{2g} band is only indicated, consistent with the presence of amorphous ceria and/or very small ceria crystallites.

Previously, a possible ion-exchange reaction on silicate adsorbent has been reported:^{19,48}

$$
Ce-O-NO2 + OH- \rightarrow Ce-OH + NO3- (1)
$$

Thus, one may deduce that the dispersion of cerium within the pores is related to the extent of migration of soluble Ce^{3+} , stabilized by hydroxyl groups of P123, prior to forming solid $CeO₂$ crystallites at higher temperatures. Thus, a wide

temperature window is expected to facilitate the migration process of cerium ions within the pores, resulting in a better dispersion.

In situ DR UV-vis spectra were recorded to gain information about changes in the coordination environment of cerium ions during the transformation from $Ce(NO₃)₃$ to $CeO₂$ (see Fig. 6). The initial DR UV-vis spectra of tfSBA–L-CeO₂–air, asSBA–L- $CeO₂$ –air, and asSBA–L-CeO₂–N₂ at low temperature are characterized by typical features of $Ce(NO₃)₃$ at 218, 255, and 314 nm, which correspond to a $4f^1$ -5d¹ electronic transfer of Ce^{3+} , a charge transfer (CT) of O^{2-} 2p to Ce^{3+} 4f/5d, and a CT of Ce^{3+} 5d to O^{2-} 2p, respectively.⁴⁹ At higher temperature, spectra show a broad absorption band at 220–320 nm, which can be attributed to a CT band of O^{2-} to Ce⁴⁺ (at \sim 270 nm) and interband transitions (at \sim 310 nm) of ceria. With increasing temperature from room temperature to 500 $^{\circ}$ C, tfSBA-L-CeO₂air (see Fig. 6a) shows an increase of the bands at 320 nm and 255 nm, but a decrease of the band at 220 nm, which is

Fig. 6 In situ DR UV-vis spectra during the calcination of (a) tfSBA-L-CeO₂-air, (b) asSBA-L-CeO₂-air, and (c) asSBA-L-CeO₂-N₂, following the protocol given in Fig. 1a.

attributed to the transformation of nitrate to oxide. In addition, there is a shift of the absorption edge from 370 nm for $Ce(NO₃)₃$ to 470 nm for $CeO₂$. For the samples asSBA-L-CeO₂-air (see Fig. 6b) and as SBA–L-CeO₂–N₂ (see Fig. 6c) similar overall changes were observed, i.e., the band at 260 nm increased, while the shoulder band at 320 nm disappeared. The presence of noise signals in the UV region can be explained by the strong absorption of the carbon-containing P123 template. Similarly, the absorption edge shifted from 370 nm to 450 nm for asSBA– L-CeO₂ calcined in air or in N₂/air. The intensity at about 380 and 250 nm has been reported to be proportional to the amount of Ce^{4+} ions in an octahedral/polymeric and a tetrahedral coordination environment of $CeO₂$, respectively.⁵⁰ Thus, we can conclude that there is more tetrahedrally coordinated cerium present in asSBA-L-CeO₂-air/N₂ than in tfSBA-L-CeO₂air after calcination (see Fig. S6, ESI†).

For bare CeO₂, the direct band gap energy $(E_{\rm g})$ can be calculated from the DR UV-vis spectra according to 51

$$
\alpha h v = A (h v - E_{\rm g})^{1/2} \tag{2}
$$

where α is the absorption coefficient, which is described by the absorbance here. Using Tauc's method, the evolution of the E_{φ} value during calcination can be derived (see Fig. S7, ESI†). In the low-temperature range, the gradual wavelength blueshift in the UV absorption (*i.e.*, E_g decrease) is due to the removal/ dissociation of crystal water and chelated $\mathrm{NO_3}^-$ according to 52

$$
Ce(NO3)3·xH2O \rightarrow Ce(NO3)3(s) \rightarrow CeONO3(s) \rightarrow CeO2-x(s)
$$

The E_g values decrease rapidly for tfSBA–L-CeO₂–air to reach a relatively stable state at about 275 °C, while asSBA-L-CeO₂-air and asSBA–L-CeO₂–N₂ require higher temperatures of \sim 375 °C and 500 \degree C to reach their final state, respectively. In addition, with increasing temperature from 300 to 500 \degree C, the band gap energy increases from 2.90 eV to 3.12 eV for tfSBA-L-CeO₂-air, suggesting the growth of $CeO₂$ crystallites. In contrast, for asSBA–L-CeO₂–air, the band gap reaches its minimum value of 2.85 eV at about 375 \degree C and then stays unchanged at higher

temperature, indicative of a stable ceria crystallite size. Finally, for asSBA-L-CeO₂-N₂, the determined E_g value suggests that there is no dominant ceria crystal phase formation with N_2 at 500 \degree C, while subsequent calcination for 2 h in air at 500 \degree C induces an E_g red-shift to 2.71 eV, which may indicate the formation of tiny $CeO₂$ crystallites.

Fig. 7 shows in situ DRIFT spectra of tfSBA-L-CeO₂-air, asSBA–L-CeO₂–air, and asSBA–L-CeO₂–N₂ during the calcination process. The assignments for the observed IR features are summarized in Table S1 (ESI†). Briefly, in the low-wavenumber region (1400-1800 cm^{-1}), mainly bands of nitrogen-containing species, and in the high-wavenumber region $(>3000 \text{ cm}^{-1})$ hydroxyl-related features are detected. In addition, there are features attributed to vibrational modes of organic groups, such as $C-O$, $C=O$, and $C-H$, originating from P123 in the case of the asSBA-15-based samples, and from a small amount of residual P123 in the case of the tfSBA-15-based sample. As can be seen in Fig. 7, all three samples exhibit a distinct change in vibrational structure at a critical temperature. In detail, at low temperatures, sample tfSBA-L-CeO₂-air shows a broad band at about 3200-3600 cm^{-1} , originating from H-bonded hydroxyl groups, which disappears when the temperature is raised from 225 \degree C to 275 \degree C, suggesting the removal of crystal water from $\text{Ce}(\text{NO}_3)_3$ \cdot 6H₂O. Simultaneously, a strong decline of the nitrate features (at $1400-1800$ cm⁻¹) is detected, 52 indicating the decomposition of cerium nitrate. Similarly, for asSBA-L-CeO₂-air and asSBA-L-CeO₂-N₂, there are rather sudden spectral changes within $225-275$ °C and $275-300$ °C. These observations further prove that the inert gas causes a delay in the decomposition of the cerium precursor. The asymmetric peak at 3740 cm^{-1} is typical for Si-OH groups, while the feature at 3700 cm^{-1} is attributed to Ce–OH.⁴⁵ It is noteworthy that tfSBA–L-CeO₂–air shows a peak at 3740 cm⁻¹ over the whole temperature range, providing evidence for the presence of uncovered silica, even after $CeO₂$ formation. For asSBA–L-CeO₂–air, at the beginning no peak at 3740 cm⁻¹ is detected, but at temperatures $>$ 275 °C, indicating that surface Si-OH is initially covered by the template, but emerges after the Puper
 Starting definites are commonly are alternative on 09 September 2021. The decomposition of the commonly of the september 2021. The decomposition of the september 2021. The definite is like the september 2021. The

Fig. 7 In situ DRIFT spectra during calcination of (a) tfSBA–L-CeO₂–air, (b) asSBA–L-CeO₂–air, and (c) asSBA–L-CeO₂–N₂, following the protocol given in Fig. 1a.

removal of P123. At low temperatures, the sample asSBA–L- $CeO₂-N₂$ shows spectra similar to those of asSBA-L-CeO₂-air, but the appearance of the Ce–OH peak at 3700 cm^{-1} evidences that the silica surface is coated by ceria. Based on these changes of the surface hydroxyl groups, we can conclude that P123 inside the pores shrinks to the pore wall when the temperature is increased. After reaching a critical temperature, surface Si–OH is exposed. On the other hand, the observation of Ce–OH in the absence of silanol features clearly shows that the formed ceria coats the silica surface.

3. Physical role of P123 in nanoconfinement formation

Previously, it had been proposed that the template could form a confined space to hinder the growth of larger crystallites. An unverified hypothesis was that the confined space was formed between the P123 micelles and the inner wall of SBA-15. As part of our detailed analysis, in situ DRIFT spectra show Si–OHrelated signals at about 3734 cm^{-1} , which appear after the removal of P123, strongly suggesting that P123 shrinks to the wall with increasing temperature. Especially, at low temperatures, the P123 chains are stabilized by their integration into the silica wall, while in the shrinkage stage, the water loss would break the central micelle structure (see Scheme 3). Thus, P123 tends to shrink to the inner wall, forming a confined space in the pore centre, and cerium ions entering the pore are coordinated by P123, as shown in Scheme 3, overturning previous hypotheses.²⁹ In the further course of the synthesis, cerium, initially coordinated by P123 in the central confinement,¹⁹ is dispersed onto the silica surface when the template is removed at higher temperatures (see Scheme 3). Besides shrunk P123, carbon (deposited during calcination at higher temperatures) may contribute to the suppression of $CeO₂$ agglomeration by further confining the space for the ceria growth. **Materials Advances**

COL M₃ ARTIONS ARE UN

4. Chemical role of P123 for co-catalytic interaction with ceria species

Based on the findings described above, the calcination process involves three steps, i.e., decomposition of cerium nitrate, migration of cerium species, and nucleation/growth of ceria. The final dispersion of ceria is shown to depend on the relative

Scheme 3 Schematic diagram of the template-assisted processes leading to ceria dispersion within the mesopores matrix.

Scheme 4 Calcination of cerium nitrate with (left) or without (right) the assistance of template P123.

decomposition temperatures of P123 and cerium nitrate. The TG and FTIR results have revealed that the decomposition behaviour of cerium nitrate/SBA-15 mixtures significantly deviates from that of bare cerium nitrate and P123 containing SBA-15, confirming the presence of interactions between cerium nitrate and the P123 template.

Calcination of mixed samples in air with the assistance of P123 results in the coordination of cerium ions by P123, while cerium nitrate migrates into the pores.¹⁹ Due to the catalytic reaction between cerium ions and P123, ceria nanoparticles are formed by oxidation in air, while without P123, thermal decomposition of cerium nitrate occurs over a wide temperature range, accompanied by nucleation and growth of ceria crystallites (see Scheme 4). The catalytic reaction facilitates the oxidation of P123-coordinated cerium to ceria nuclei at even lower temperature than the critical decomposition temperature of bare $Ce(NO₃)₃·6H₂O$. Meanwhile, reduction by P123 consumes lattice oxygen, which slows down the growth of ceria nanocrystals, thus increasing dispersion of ceria on SBA-15.

Based on the knowledge of the catalytic effect of P123, the synthesis protocol for $CeO₂/SBA-15$ was modified by application of an inert N_2 atmosphere to further improve the ceria dispersion. Considering the low thermal stability of P123, which decomposes at a temperature of about 115 \degree C (TGA) in air, the nitrogen atmosphere could delay the decomposition of P123 into carbon chain fragments, thus inhibiting the catalytic oxidation induced formation of ceria nuclei at an early stage until the decomposition temperature of cerium nitrate. Reaching this temperature results in an explosive nucleation of ceria and catalytic decomposition of P123, which is favourable for the formation of a homogeneous nanoscale grain structure.

5. DeNO_x performance

Fig. 8 summarizes the catalytic performance of $CeO₂/SBA-15$ samples in NH₃-SCR within a temperature range of 100-450 $^{\circ}$ C. Sample tfSBA–L-CeO₂–air shows the lowest NO conversion $(30%) over the whole temperature range (see Fig. 8a) and$ the lowest N_2 selectivity in the high-temperature range (see Fig. 8b). The low reactivity in $NH₃$ -SCR is in agreement with that of bare CeO₂ reported previously.⁴² Sample asSBA-L-CeO₂air, which is based on as-made SBA-15 and is calcined in air atmosphere, shows a dramatic improvement in NO conversion and N_2 selectivity, exhibiting a maximum in NO conversion of about 60% at 300 $^{\circ}$ C. Combining the template interaction with a calcination in inert N_2 atmosphere yields sample asSBA-L- $CeO₂-N₂$, which shows a further significant increase in NO

Fig. 8 Catalytic performance of $CeO₂/SBA-15$ samples in $NH₃-SCR$. (a) NO conversion and (b) N_2 selectivity. Feed: 500 ppm NO, 500 ppm NH₃, 5% O₂, balanced with N₂; total gas flow: 50 N ml min⁻¹, GHSV = 30 000 h^{-1} . The data points were determined based on an average of five gas-phase IR spectra.

conversion compared to as SBA–L-CeO₂–air, and even the N₂ selectivity slightly improves. Thus, the $NH₃$ -SCR performance of these three samples follows the order as SBA–L-CeO₂–N₂ $>$ asSBA–L-CeO₂–air \gg tfSBA–L-CeO₂–air, which resembles the extent of ceria dispersion on the SBA-15 surface, as discussed above. In addition, Fig. S9 (ESI†) provides results from a longterm test (20 h) at 300 $^{\circ}$ C, demonstrating the stability of these samples for NH₃-SCR. At high ceria loading, similar trends were observed (see Fig. S8, ESI†), with overall higher NO conversions compared to low cerium loading. Especially, the N_2 selectivity of asSBA–H-CeO₂–N₂ was significantly higher than that of asSBA–H-CeO₂–air, demonstrating that an inert calcination atmosphere could further improve the $CeO₂$ dispersion and thus the catalytic performance even at high ceria loading. Table S2 (ESI†) summarizes reactivity data of ceria-based catalysts for $NH₃$ -SCR. In general, pure ceria shows a lower NO_x performance

than silica supported ceria, prepared by controlled nanoconfinement thus further improving the ceria dispersion.

Experimental section

1. Materials preparation

asSBA-15 and tfSBA-15. Mesoporous SBA-15 was synthesized according to the methods reported by Zhao et al ² Briefly, 4 g of pluronic $EO_{20}PO_{70}EO_{20}$ (P123) was dissolved in 150 ml of 1.6 M HCl solution and stirred at 35 $^{\circ}$ C and a speed of 250 rpm for 2.5 h. Then, 11 ml of tetraethylorthosilicate (TEOS) was added and the obtained mixture was stirred at 35 \degree C at an increased speed of 400 rpm for 20 h. The stirred mixture was then transferred to a Teflon bottle for hydrothermal treatment and crystallization at 85 °C for 24 h. Vacuum filtration of the suspension at room temperature yielded a white powder, which is referred to as as-made SBA-15 (asSBA-15) and which contains the templating agent P123 in the mesoporous matrix. Calcination of asSBA-15 at 550 \degree C for 12 h with a heating rate of 1.5 \degree C min^{-1} in air yields a white powder, referred to as template-free SBA-15 (tfSBA-15), from which the templating agent P123 has been removed.

 $aSBA-L/H-CeO₂-air/N₂$. Cerium nitrate was employed as the cerium source for the preparation of $CeO₂/SBA-15$ catalysts. The precursor $Ce(NO₃)₃·6H₂O$ was mixed with asSBA-15 (see above) via solid-state grinding at room temperature for 30 min. The thoroughly mixed powder was then calcined in a muffle furnace at 500 °C for 5 h (heating rate: 1.5 °C min $^{-1}$) in air (denoted by –air) or N_2 (denoted by – N_2) to remove the P123 template and form the $CeO₂$ phases. To explore the effect of $CeO₂$ loading, $CeO₂/SBA-15$ catalysts with low and high $CeO₂$ loading were prepared. The sample resulting from a mixture of 0.173 g of $Ce(NO₃)₃·6H₂O$ and 0.225 g of asSBA-15 will be referred to as L-CeO₂, while that resulting from a mixture of 0.346 g of $Ce(NO₃)₃·6H₂O$ and 0.225 g of asSBA-15 will be labelled as H-CeO₂. Thus, as an example, mixing 0.173 g of Ce(NO₃)₃.6H₂O with 0.225 g of asSBA-15, followed by calcination in air, gave the sample as SBA-L-CeO₂-air.

tfSBA–L/H-CeO₂–air/N₂. Similarly, tfSBA-L/H-CeO₂–air/N₂ sam-ples were prepared by following the above protocol, except that asSBA-15 was replaced by tfSBA-15. To ensure comparable $CeO₂$ loadings, 0.225 g of asSBA-15 was calcined yielding 0.108 g of tfSBA-15. Thus, as an example, mixing 0.173 g of $Ce(NO₃)₃·6H₂O$ with 0.108 g of tfSBA-15 followed by calcination in air resulted in the sample tfSBA-L-CeO₂-air.

Prior to catalytic testing in NH₃-SCR, all samples treated in N_2 during synthesis were calcined in air for 2 h at 500 °C to remove residual template.

2. Characterization

XRD. X-ray diffraction experiments were carried out in transmission geometry on an X-ray powder diffractometer (StadiP, Stoe & Cie GmbH) with a Mythen 1K (Dectris) detector. For the measurements, Cu K α 1 radiation (λ = 1.540598 Å) and a Ge[111] monochromator were used.

N2 adsorption isotherm. Nitrogen adsorption–desorption measurements were carried out on a NOVA 3000e (Quantachrome) to determine the surface area and pore size of the mesoporous samples. The Brunauer–Emmett–Teller (BET) method was employed to calculate the specific surface area, and nonlocal density functional theory (NLDFT) was used to calculate the adsorption branch to obtain the porosity characteristics.

Transmission electron microscopy (TEM). The transmission electron microscope (JEOLJEM-2100F, Tokyo, Japan) is equipped with a Schottky field emitter and was operated at a nominal accelerate voltage of 200 kV. The powder samples were dispersed in ethanol and placed on a carbon grid (Plano).

Thermogravimetric analysis (TGA). TGA was performed on a TGA/SDTA851-e (Mettler Toledo). In order to record TGA curves, samples were exposed to a flow of synthetic air or pure N_2 gas $(100 \text{ ml min}^{-1})$ while being heated from room temperature to 500 $^{\circ}$ C at a rate of 1.5 $^{\circ}$ C min⁻¹. Samples were then calcined at 500 \degree C for 5 h. At the same time, derivative thermogravimetry (DTG) profiles were obtained.

Gas-phase Fourier transform infrared (FTIR) spectroscopy. FTIR spectroscopy was performed on a Bruker Tensor 27 equipped with a DLaTGS (deuterated and L-alanine-doped triglycine sulfate) detector, and a low volume gas cell (25 ml, 0.5 m path-length, Axiom). The gas cell was heated to 120 $^{\circ}$ C to avoid the condensation of water produced by the catalytic reaction. IR spectra were continuously recorded every minute with a resolution of 4 cm^{-1} . The backgrounds used for the analysis were recorded under pure nitrogen gas at room temperature. For online IR gas-phase detection, the Tensor was connected to the SCR reactor (commercial CCR1000 catalyst cell, Linkam Scientific Instruments). The gas-phase concentrations were determined based on a set of calibration curves.

Diffuse reflectance (DR) UV-vis spectroscopy. DR UV-vis spectra were recorded on a Jasco V-770 UV-visible/NIR spectrometer with a praying mantis mirror cell and a high-temperature reaction chamber (Harrick Scientific Products Inc.). Halogen and deuterium light sources were adopted to transmit wavelengths in the visible and ultra-violet, respectively, and a Peltiercooled PbS detector was employed. Spectra were recorded from 800 to 200 nm with a spectral resolution of 0.5 nm. MgO was used as the white standard.

Raman spectroscopy. Raman spectra were recorded in a backscattering geometry. For excitation, an argon ion laser (514.5 nm, Melles Griot) was employed and for collection of the backscattered light a transmission spectrometer (Kaiser Optical) equipped with a charge-coupled device (CCD) detector was employed. The spectrometer was calibrated using the emission lines of a standard argon lamp. The resolution of the spectrometer was 5 cm^{-1} and the wavelength stability was better than 0.5 cm^{-1} . The laser power was set to 2.5 mW and measured by a power meter (Ophir). The Raman features were fitted by Voigt functions using a Gaussian linewidth of 6 cm^{-1} to account for the instrumental broadening. In situ Raman spectra were recorded by using a commercial CCR1000 catalyst cell (Linkam Scientific Instruments).

Diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS). DRIFT spectra were recorded on a Vertex 70 (Bruker) equipped with a liquid N_2 cooled mercury cadmium telluride (MCT) detector and a commercial Harrick cell. As a background standard, KBr powder was used. The spectral resolution of DRIFTS was 1 cm⁻¹, the temporal resolution was 60 s. In situ DRIFT spectroscopy was combined with gas-phase analysis, by analyzing the gas phase at the outlet of the in situ cell by gas-phase FTIR spectroscopy (see above).

3. Catalytic tests

The SCR performance of the catalysts was measured in a commercial CCR1000 reactor (Linkam Scientific Instruments) in combination with quantitative gas-phase FTIR spectroscopy (see above). The flow rate of the inlet gases was adjusted by mass flow controller, by employing the following gas sources: 2000 ppm NO/N₂ (\pm 0.25% abs.), 2000 ppm NH₃/N₂ (\pm 0.25%) abs.), O_2 (\geq 99.999%), and N₂ (\geq 99.999%). For catalytic testing in $NH₃$ -SCR, the concentrations of NO, $NH₃$, and $O₂$ in the feed gas mixture were set to 500 ppm, 500 ppm, and 5%, respectively, and balanced with N_2 . The weight of the catalyst samples loaded into the reactor was 15 mg; the feed gas mixture passed through the catalyst bed at a total flow rate of 50 ml min^{-1} , resulting in a gas hourly space velocity (GHSV) of 30 000 h^{-1} . Materials Advances
 Nationalizer Access Are the Creative Common Contrast) Distribution (September 2021) (Since the Case Contrast) Distribution and Since the Creative Common Common Common Common Common Common Common Commo

Using the detected concentrations of NO, $NO₂$, $N₂O$, and $NH₃$ in the outlet gas flow, and the feed gas concentrations, the N_2 concentration can be calculated based on conservation of mass, assuming that besides N_2 nitrogen is present only in the form of the gases listed above. NO_x conversion and N_2 selectivity were calculated using the following equations:

NO conversion =
$$
\frac{[NO]_{in} - [NO]_{out}}{[NO]_{in}} \times 100\%
$$
 (3)

$$
N_2 \text{ selectivity} = \frac{[N_2]_{out}}{[N_2]_{out} + [N_2O]_{out}} \times 100\% \tag{4}
$$

where the subscripts in and out indicate the inlet and outlet flows of the reactor, respectively.

Conclusions

Herein, we have reported our studies of the synthesis mechanism of $CeO₂/SBA-15$ catalysts by the solid-state impregnation method using a combination of TG and IR gas-phase analysis, as well as in situ Raman, DR UV-vis, and DRIFT spectroscopies. These studies have revealed that ceria formation on mesoporous SBA-15 during calcination involves first cerium precursor decomposition, then migration of cerium species, and finally ceria formation at high temperature. Based on our findings we can explain the roles of the template inside the mesoporous matrix: on one hand P123 acts as a physical barrier to prevent the growth of ceria, on the other, P123 acts as a chemical reactant, which interacts with cerium ions, thus enabling the catalytic decomposition of the cerium precursor.

With the gained mechanistic knowledge of the formation of CeO₂/SBA-15 materials, a new calcination protocol was developed based on an intended retardation of the decomposition of the template and the cerium precursor, leading to a higher dispersion of ceria inside the pores and thus a significant improvement of catalytic performance in NH3-SCR.

An important finding of the study is that the precursor itself can facilitate the synthesis of the catalyst, as demonstrated here for a ceria-based catalyst, by using both the template and the gas atmosphere to control the catalytic reaction conditions. Our results underline the importance of the details of the calcination process for defining the catalyst properties. Application of in situ spectroscopy allows one to monitor the preparation process, including calcination, thus facilitating a rational synthesis of supported catalysts as shown here in the context of NO_x abatement. The outlined approach is expected to trigger more studies on the development of improved catalysts based on a careful design of the catalytic preparation reactions. Publish downloaded on an interded created on of the energy article. This article is licensed in the energy article is licensed under the energy of the same of the energy of the energy of the same of the energy of the ener

Author contributions

The manuscript was written through contributions of both authors. Original idea, C. H.; conceptualization, J. S. and C. H.; methodology, J. S.; formal analysis, J. S.; writing – original draft preparation, J. S.; writing – review and editing, C. H.; supervision, C. H.; project administration, C. H.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors thank Martin Brodrecht for N_2 adsorption– desorption measurements, Kathrin Hofmann for XRD analysis, Stefan Lauterbach for TEM analysis, and Karl Kopp for XPS analysis and technical support. Financial support by the Deutsche Forschungsgemeinschaft (DFG) is gratefully acknowledged.

Notes and references

- 1 W. Li, J. Liu and D. Y. Zhao, Nat. Rev. Mater., 2016, 1, 1–17.
- 2 D. Y. Zhao, J. L. Feng, Q. S. Huo, N. Melosh, G. H. Fredrickson, B. F. Chmelka and G. D. Stucky, Science, 1998, 279, 548–552.
- 3 F. Khanmohammadi, M. A. Molina, R. M. Blanco, S. N. Azizi, C. Marquez-Alvarez and I. Diaz, Microporous Mesoporous Mater., 2020, 309, 1–6.
- 4 S. Inagaki, S. Guan, Y. Fukushima, T. Ohsuna and O. Terasaki, J. Am. Chem. Soc., 1999, 121, 9611–9614.
- 5 J. Nakazawa, B. J. Smith and T. D. P. Stack, J. Am. Chem. Soc., 2012, 134, 2750–2759.
- 6 S. F. Zhang, B. Zhang, H. J. Liang, Y. Q. Liu, Y. Qiao and Y. Qin, Angew. Chem., Int. Ed., 2018, 57, 1091–1095.
- 7 P. Hongmanorom, J. Ashok, G. H. Zhang, Z. F. Bian, M. H. Wai, Y. Q. Zeng, S. B. Xi, A. Borgna and S. Kawi, Appl. Catal., B, 2021, 282, 119564.
- 8 C. Hess, in Comprehensive Inorganic Chemistry II, ed. J. Reedijk and K. Poeppelmeier, Elsevier, Amsterdam, 2013, vol. 7.
- 9 V. Chaudhary and S. Sharma, J. Porous Mater., 2017, 24, 741–749.
- 10 P. Hongmanorom, J. Ashok, S. Das, N. Dewangan, Z. Bian, G. Mitchell, S. Xi, A. Borgna and S. Kawi, J. Catal., 2020, 387, 47–61.
- 11 R. R. Chada, S. S. Enumula, K. S. Reddy, M. D. Gudimella, S. R. R. Kamaraju and D. R. Burri, Microporous Mesoporous Mater., 2020, 300, 110144.
- 12 O. Daoura, G. Fornasieri, M. Boutros, N. El Hassan, P. Beaunier, C. Thomas, M. Selmane, A. Miche, C. Sassoye, O. Ersen, W. Baaziz, P. Massiani, A. Bleuzen and F. Launay, Appl. Catal., B, 2021, 280, 119417.
- 13 J. W. Fan, X. F. Niu, W. Teng, P. Zhang, W. X. Zhang and D. Y. Zhao, J. Mater. Chem. A, 2020, 8, 17174–17184.
- 14 R. A. Mitran, D. C. Culita and I. Atkinson, Microporous Mesoporous Mater., 2020, 306, 110484.
- 15 R. Manno, P. Ranjan, V. Sebastian, R. Mallada, S. Irusta, U. K. Sharma, E. V. Van der Eycken and J. Santamaria, Chem. Mater., 2020, 32, 2874–2883.
- 16 N. N. Mikheeva, V. I. Zaikovskii and G. V. Mamontov, Microporous Mesoporous Mater., 2019, 277, 10–16.
- 17 C. Chong, A. H. K. Owgi, N. Ainirazali, S. Y. Chin and H. D. Setiabudi, Mater. Today: Proc., 2018, 5, 21644–21651.
- 18 S. N. Bukhari, A. H. K. Owgi, N. Ainirazali, D. V. N. Vo and H. D. Setiabudi, Mater. Today: Proc., 2018, 5, 21620–21628.
- 19 W. W. Yang, H. M. Liu, Y. M. Li and D. H. He, Microporous Mesoporous Mater., 2016, 228, 174–181.
- 20 W. W. Yang, H. M. Liu, Y. M. Li, H. Wu and D. H. He, Int. J. Hydrogen Energy, 2016, 41, 1513–1523.
- 21 B. Dragoi, G. Laurent, S. Casale, T. Benamor, B. Lebeau, C. Boissiere, F. Ribot, M. Selmane, P. Schmidt, D. Kreher and A. Davidson, J. Sol–Gel Sci. Technol., 2019, 91, 552–566.
- 22 B. Dragoi, I. Mazilu, A. Chirieac, C. Ciotonea, A. Ungureanu, E. Marceau, E. Dumitriu and S. Royer, Catal. Sci. Technol., 2017, 7, 5376–5385.
- 23 Y. Yin, W. J. Jiang, X. Q. Liu, Y. H. Li and L. B. Sun, J. Mater. Chem., 2012, 22, 18514–18521.
- 24 Y. Yin, P. Tan, X. Q. Liu, J. Zhu and L. B. Sun, J. Mater. Chem. A, 2014, 2, 3399–3406.
- 25 Y. Yin, D. M. Xue, X. Q. Liu, G. Xu, P. Ye, M. Y. Wu and L. B. Sun, Chem. Commun., 2012, 48, 9495–9497.
- 26 Y. Yin, Z. H. Wen, X. Q. Liu, A. H. Yuan and L. Shi, Adsorpt. Sci. Technol., 2018, 36, 953–966.
- 27 Y. Yin, Z. F. Yang, Z. H. Wen, A. H. Yuan, X. Q. Liu, Z. Z. Zhang and H. Zhou, Sci. Rep., 2017, 7, 4509.
- 28 Y. Yin, H. Wu, L. Shi, J. Q. Zhang, X. Y. Xu, H. Y. Zhang, S. B. Wang, M. Sillanpaad and H. Q. Sun, Environ. Sci.: Nano, 2018, 5, 2842–2852.
- 29 Y. Yin, L. Shi, W. L. Li, X. N. Li, H. Wu, Z. M. Ao, W. J. Tian, S. M. Liu, S. B. Wang and H. Q. Sun, Environ. Sci. Technol., 2019, 53, 11391–11400.
- 30 A. Taratayko, Y. Larichev, V. Zaikovskii, N. Mikheeva and G. Mamontov, Catal. Today, 2021, 375, 576–584.
- 31 L. P. Han, S. X. Cai, M. Gao, J. Hasegawa, P. L. Wang, J. P. Zhang, L. Y. Shi and D. S. Zhang, Chem. Rev., 2019, 119, 10916–10976. 32 A. Trovarelli, Catal. Rev., 1996, 38, 439–520.
- 33 J. Strunk, W. C. Vining and A. T. Bell, J. Phys. Chem. C, 2011, 115, 4114–4126.
- 34 J. Shen and C. Hess, Catalysts, 2020, 10, 1386.
- 35 M. T. Weller, Inorganic materials chemistry, Oxford University Press, Oxford; New York, 1994.
- 36 J. P. Thielemann, F. Girgsdies, R. Schlogl and C. Hess, Beilstein J. Nanotechnol., 2011, 2, 110–118.
- 37 S. Y. Chen, T. Mochizuki, Y. Abe, M. Toba, Y. Yoshimura, P. Somwongsa and S. Lao-ubol, Appl. Catal., B, 2016, 181, 800–809.
- 38 E. A. Cochran, K. N. Woods, D. W. Johnson, C. J. Page and S. W. Boettcher, J. Mater. Chem. A, 2019, 7, 24124–24149.
- 39 W. Kang, D. O. Ozgur and A. Varma, ACS Appl. Nano Mater., 2018, 1, 675–685.
- 40 N. Ulagappan and H. Frei, J. Phys. Chem. A, 2000, 104, 7834–7839.
- 41 Y. L. Liu and H. J. Kim, Sensors, 2017, 17, 1469.
- 42 B. L. Zhang, S. G. Zhang and B. Liu, Appl. Surf. Sci., 2020, 529, 147068.
- 43 W. H. Weber, K. C. Hass and J. R. Mcbride, Phys. Rev. B: Condens. Matter Mater. Phys., 1993, 48, 178–185.
- 44 T. Suzuki, I. Kosacki, H. U. Anderson and P. Colomban, J. Am. Ceram. Soc., 2001, 84, 2007–2014.
- 45 A. Filtschew, K. Hofmann and C. Hess, J. Phys. Chem. C, 2016, 120, 6694–6703.
- 46 C. Schilling, A. Hofmann, C. Hess and M. V. Ganduglia-Pirovano, J. Phys. Chem. C, 2017, 121, 20834–20849.
- 47 A. Filtschew, D. Stranz and C. Hess, Phys. Chem. Chem. Phys., 2013, 15, 9066–9069.
- 48 J. Seo, J. W. Lee, J. Moon, W. Sigmund and U. Paik, ACS Appl. Mater. Interfaces, 2014, 6, 7388–7394.
- 49 V. Nicolini, E. Varini, G. Malavasi, L. Menabue, M. C. Menziani, G. Lusvardi, A. Pedone, F. Benedetti and P. Luches, Mater. Des., 2016, 97, 73–85. **Naterials Advances**

11 L.P. Hans, X. Cali M. Goog, Divished on 2021. This article. Common Monte Pays, 1973. Hans and the Common Monte Pays, 1974, American Cali M. This article. The spin of the spin of the spin of the spi
	- 50 C. M. Aiube, K. V. de Oliveira and J. L. de Macedo, Catalysts, 2019, 9, 377.
	- 51 S. Phokha, S. Pinitsoontorn, P. Chirawatkul, Y. Poo-arporn and S. Maensiri, Nanoscale Res. Lett., 2012, 7, 425.
	- 52 C. X. Wang, D. Z. Ren, J. C. Du, Q. G. Qin, A. M. Zhang, L. Chen, H. Cui, J. L. Chen and Y. K. Zhao, Catalysts, 2020, 10, 143.