Construction of hybrid multi-shell hollow structured CeO$_2$–MnO$_x$ materials for selective catalytic reduction of NO with NH$_3$†

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Hollow structured CeO$_2$–MnO$_x$ hybrid materials with up to three shells were prepared successfully by using carbon spheres as the hard template. It was found that the shells could be well controlled by simply adjusting the calcination rate. Elemental mapping results showed Mn and Ce species exhibited a homogeneous spatial distribution and the existence of Mn could improve the diffusion of CeO$_2$ into the carbon spheres during the construction of the multi-shell structure, suggesting the intensive cooperative interaction between Mn and Ce. When triple-shell CeO$_2$–MnO$_x$ hollow spheres were used as a catalyst for selective catalytic reduction of NO with NH$_3$, superior low-temperature catalytic performance was exhibited, compared with traditional CeO$_2$–MnO$_x$ nanoparticles, single-shell and double-shell hollow spheres. Combined with XRD, H$_2$-TPR and XPS characterization, it was indicated that the synergistic effects and surface active species enhanced by the special multi-shell CeO$_2$–MnO$_x$ hollow structures could account for the excellent performances. The results of the present study shed light on the creation of complex and hybrid hollow structured materials for superior performance in catalysis fields, like NO reduction for environmental protection.

1 Introduction

Ceria, as a typical rare earth oxide, owns considerable intrinsic properties such as abundant oxygen vacancy defects, high oxygen storage capacity and alterable valence states,$^1$ which result in its extensive applications in the field of diesel soot oxidation,$^2$ CO oxidation,$^3$ photocatalysis,$^4,5$ NO reduction,$^6,7$ and so on. Nevertheless, pure ceria is rarely used in catalysis due to its inherent poor activity. It is recognized that the combination of CeO$_2$ and transition as well as noble metals, for instance, Cu,$^8$ Mn,$^x$ Nb,$^9$ Pt$^{10}$ and Au,$^{11}$ is beneficial to its properties. Besides, manganese hybrid oxides, with variable valences and resulting intrinsic environmentally benign features as well as excellent catalytic activity, have been considered as promising catalysts for redox reactions.$^{12–14}$ Thus, much attention has been paid to CeO$_2$–MnO$_x$ composites due to their admirable performance in catalysis, for example, the low-temperature dNO$_x$ reaction.$^{6,15,16}$ The advantageous interactions between Ce and Mn species bring about the excellent redox ability, amphoteric property and thermal stability, which are helpful for the catalytic application of CeO$_2$–MnO$_x$ composites.$^{17}$

Generally, compared with the irregular particles, catalysts with well-designed structure are supposed to exhibit amazing functions on account of the size, shape, interface effects, etc.$^{18}$ Thus, considerable researches have been focused on preparing CeO$_2$–MnO$_x$ composites with specific structures. Chen et al.$^{19}$ prepared the Ce–Mn binary oxide hollow cubes by a novel method, which obtained excellent CO oxidation activity and remarkable adsorption capacity of Congo red.$^{20}$ Putla and the co-workers designed a heteronanostructure of CeO$_2$ nanocubes (100) with dispersed MnO$_x$ nanoparticles for diesel soot oxidation and continuous-flow benzylamine oxidation.$^2$ Besides, Wei et al.$^{21}$ reported a kind of MnO$_2$ doped CeO$_2$ catalyst with tailored 3-D channels, which possessed large surface area for chemical absorption, as well as proper channels for diffusion.$^{22}$ Hence, the controlled fabrication of well-structured CeO$_2$–MnO$_x$ composites is of great significance to catalysis field.

Recently, hollow structured materials with features of low density, high permeability and confined inner cavity have captured tremendous attention.$^{23–24}$ Especially, the fabrication of multi-shell hollow spheres with special isolated chambers and controllable shell structures attracted considerable attention.$^{25,26}$ According to the literatures, multi-shell CeO$_2$ hollow spheres have been successfully synthesized.$^{4,11,27}$ Nevertheless, to the best of our knowledge, hybrid multi-shell ceria-based
hollow spheres with diverse compositions have rarely been prepared, and the relationship of their structure and catalytic performance is not reported yet.

Therefore, in this study, by using carbon spheres as a hard template, we reported for the first time the fabrication of CeO$_2$–MnO$_x$ hollow spheres with adjustable shells via simple modification of calcination rate for the first time. The obtained samples were characterized by scanning electron microscopy (SEM), transmission electron microscopy (TEM), X-ray diffractometry (XRD), Brunauer–Emmett–Teller (BET), X-ray photoelectron spectroscopy (XPS), X-ray fluorescence analysis (XRF), H$_2$–temperature programmed reduction (H$_2$–TPR) and tested in the triple-shell sample.

Additionally, traditional CeO$_2$ hollow spheres could be synthesized without another raw material. Then, a hydrothermal treatment of the above mixed solution at several times, respectively. The precursor was dried at 80 °C for 9 h was taken. A brown precipitate was dried at 60 °C and then calcined in air at 600 °C for 3 h with drying rates (2 °C min$^{-1}$, 5 °C min$^{-1}$ and 10 °C min$^{-1}$) to get the anticipated MSCMHSs. Individual cerium and manganese hollow spheres could be synthesized without another raw material. Additionally, traditional CeO$_2$–MnO$_x$ nanoparticles (NP) with identical molar ratio were also prepared through the same method without carbon spheres (heating rate being of 10 °C min$^{-1}$) for comparison purposes.

2.2 Characterizations

Scanning electron microscopy (SEM) images were captured on a Philips XL30 electron microscope operated at beam energy of 10.0 kV.

Transmission electron microscopy (TEM) images were taken on a JEM-2100 instrument with an acceleration voltage of 200 kV. High-resolution transmission electron microscopy (HRTEM) and selected area electron diffraction (SAED) were determined on a JEM-2100F transmission electron microscope with an accelerating voltage of 200 kV. High-angle annular dark field (HAADF)-STEM characterizations were carried out using a JEOL 2100F microscope equipped with an Oxford energy dispersive X-ray (EDX) analysis system with an acceleration voltage of 200 kV.

Fourier translation infrared spectrum (FT-IR) was collected on a NICOLET iS10 FT-IR spectrometer (Thermo Electron Corporation, USA) from 600 to 4000 cm$^{-1}$. The number of scans was 32 at a resolution of 4 cm$^{-1}$.

Powder X-ray diffraction (XRD) patterns were recorded on a Shimadzu XRD-6000 power X-ray diffractometer with Cu K$_\alpha$ radiation ($\lambda = 1.5418$ Å). The intensity data were collected over a 2θ range of 10–80° with a 0.05° step size.

Textural characteristics of the samples were determined by a nitrogen adsorption operation at −196.15 °C on a Micrometrics ASAP-2020 adsorption analyzer by the Brunauer–Emmett–Teller (BET) method. Ahead of each analysis, a pre-processing of the samples was performed under vacuum at 300 °C for 3 h. The surface areas were calculated by the BET equation and the average pore diameter along with pore size distribution were determined by using the equivalent cylindrical model analysis of BJH.

X-ray fluorescence analysis (XRF) was implemented on an ARL-9800 apparatus to clear the bulk composition of the samples. The X-ray tube was operated at 60 kV and 20 mA.

X-ray photoelectron spectroscopy (XPS) analysis was performed on a PHI 5000 Versa Probe system, using monochromatic Al K$_\alpha$ (1486.6 eV) radiation operating at an accelerating power of 15 kW. Prior to the measurement, the samples were outgassed at room temperature in a UHV chamber (<5 × 10$^{-7}$ Pa). All binding energies (BE) of the samples were calibrated with the adventitious C 1s peak at 284.6 eV to compensate the charging effect. The experimental error was within ±0.1 eV.

H$_2$–Temperature programmed reduction (H$_2$–TPR) was carried out with H$_2$–Ar mixture (7% H$_2$ by volume) as reducing agent. Typically, 50 mg sample was used in a quartz U-tube reactor for each measurement. After the sample was pretreated in a N$_2$ stream at 200 °C for 0.5 h, reduction measurement of the sample by H$_2$ began from room temperature to 800 °C with a rate of 10 °C min$^{-1}$.

2.3 Catalytic activity test

The catalytic performance of the samples for NH$_3$–SCR reaction was determined under the reaction conditions as follows: 500 ppm NO, 500 ppm NH$_3$, 5 vol% O$_2$, 5 vol% H$_2$O (when used) and balance N$_2$. The total flow rate was 100 mL min$^{-1}$ and
100 mg sample was put to use in a fixed-bed quartz reactor tube to get a space velocity of 60 000 mL g⁻¹ h⁻¹. Before the mixture reactant gases were fed to the system, a preprocessing for the samples at 200 °C for 0.5 h was implemented to remove existent impurities, using high-purity N₂. The concentration of NO at designated temperature during the reaction was detected by an online Thermo fisher IS10 FTIR spectrometer equipped with a 2 m path-length gas cell (250 mL volume), through which NO conversion of the samples could be calculated in accordance with the following equations:

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

\[
\text{N₂ selectivity (\%) } = \frac{[\text{NO}]_{\text{in}} - [\text{NO}]_{\text{out}} + [\text{NH}_3]_{\text{in}} - [\text{NH}_3]_{\text{out}} - [\text{NO}_2]_{\text{out}} - 2[N_2]_{\text{out}}}{[\text{NO}]_{\text{in}} - [\text{NO}]_{\text{out}} + [\text{NH}_3]_{\text{in}} - [\text{NH}_3]_{\text{out}}} \times 100
\]

3 Results and discussion

3.1 Morphology and structure

As presented in Fig. S1 (ESI†), uniform carbon spheres (CSs) with size of about 600 nm were obtained under hydrothermal condition. During hydrothermal process, dehydration, intermolecular cross-linking and nucleus growth are supposed to occur. As a result, carbon spheres with abundant superficial active functional groups (FT-IR, Fig. S2†) and accessible pore channels were formed. Thus, target ions could be trapped easily on the surface of carbon spheres due to electrostatic force and diffuse inward through the channels. Sequentially, hollow structures were obtained after the following calcination process to remove the template. It was widely reported that different constituents and situations could give birth to varied hollow structures. In the present work, mono-component and hybrid oxide hollow structures of manganese and cerium were prepared to explore the interaction between the constituents. Furthermore, the calcination rates were investigated for the tunable inner structures of the hybrid oxide hollow spheres.

The morphologies of as-synthesized samples were observed by SEM and TEM characterizations. As exhibited in Fig. 1b–d, after the calcination process under different conditions, multi-shell CeO₂–MnOₓ hollow spheres (MSCMHSs) with sphere-in-sphere morphology were obtained. It was interesting that the inner structures of the hollow spheres could be controlled by changing the calcination rate, i.e., single-shell (2 °C min⁻¹), double-shell (5 °C min⁻¹) and triple-shell (10 °C min⁻¹). The average diameter of the obtained MSCMHSs was about 400 nm, and the average shell thickness was 40 nm. Specially, the external, middle and internal shells of the triple-shell hollow spheres were approximately estimated as 400 nm, 150 nm and 50 nm, respectively. To further explore the elemental distribution of the multi-shell hollow spheres, HAADF-STEM was conducted on a single triple-shell hollow sphere as an example. Element mapping analysis in Fig. 2a showed that the triple-shell hollow structure could be discriminated clearly, in which the elements of Mn, Ce O were distributed uniformly. It is worthy to be noted that the homogeneous elemental distribution of Mn and Ce was favorable to the synergistic effects between them. The HRTEM image and inset SAED pattern of triple-shell hollow sphere (Fig. 2b) demonstrated that the shell was polycrystalline. The lattice fringes with \( d \) spacing of 0.31 nm, 0.28 nm and 0.19 nm were ascribed to (111), (200), (220) crystal planes of fluorite-type CeO₂, respectively. Moreover, the lattice fringe of 0.25 nm was indexed to (202) crystal plane of Mn₃O₄. The diffraction rings and the inset XRD patterns accorded well with the crystalline phase results.

To shed light on the possible forming process of MSCMHSs, the morphology of CSs@CeO₂–MnOₓ precursor before calcination was investigated. For that matter, the solvent-thermal treatment was believed as the critical step for MSCMHSs formation, which could efficiently enhance the adsorption and diffusion of manganese and cerium cations into the carbon spheres through the pore channels. As shown in Fig. S3† for the SEM image of the precursor after the solvent-thermal process, the spheres were uniformly dispersed with the size of approximately 600 nm as carbon spheres. The rough surface of the spheres containing tiny particles indicated the deposition of CeO₂–MnOₓ precursor on the carbon spheres. Corresponding TEM image (Fig. 1a) displayed the core-shell architecture with the evident interface between CSs and CeO₂–MnOₓ precursor. Notably, the MSCMHSs (Fig. 1b–d) exhibited smaller diameters than CSs@CeO₂–MnOₓ precursor, which
revealed a shrinked process in accompany with combustion of carbon spheres during calcination. Besides, TG-DTA curves of the precursor were manifested in Fig. 3. Effects of different heating rates on the thermal process were investigated. There were two main exothermal peaks in the DTA curves for all cases. The exothermal peak around 250 °C and the sharp peak concentrated at around 320 °C derived from the combustion of carbon spheres.\(^{11}\) Apparently, with the increasing heating rate, the intensity of exothermal peak for carbon spheres combustion increased. In particular, when the heating rate rose to 10 °C min\(^{-1}\) eventually, greatly enhanced exothermal intensity for carbon spheres combustion was observed. The results indicated that higher heating rate resulted in faster combustion of carbon spheres. As was reported,\(^{32}\) the construction of multi-shell structure was mainly determined by the rate differences between oxide shell formation and carbon shrinkage, which

\[\text{Fig. 2 (a) HAADF-STEM mapping images and (b) HRTEM image of the triple-shell hollow sphere. Insets are the corresponding XRD pattern and selected area electron diffraction (SAED) image.}\]

\[\text{Fig. 3 TG-DTA curves of CSs@CeO}_2–\text{MnO}_x \text{ precursor with different heating rates of (a) 2 °C min}^{-1}, (b) 5 °C min}^{-1} \text{ and (c) 10 °C min}^{-1}.\]
might cause separation. Therefore, with the increasing heating rate, a kind of heterogeneous contraction existed and facilitated the combustion of carbon spheres as well as the diffusion, penetration and redistribution of cations inward the carbon spheres. Thus, accompanied by different degrees of crystallization and penetration for manganese and cerium cations, controllable internal structures were obtained through the non-equilibrium calcination process, with the removal of carbon spheres. The schematic diagram for synthesis of the MSCMHSs was illustrated in Scheme 1.

Additionally, the hollow spheres with single component (i.e., Mn$_2$O$_3$ and CeO$_2$; confirmed by XRD patterns in Fig. 4) under calcination rate of 5 °C min$^{-1}$ were synthesized (Fig. S4†). We could observe from the SEM image (Fig. S4a†) that Mn$_2$O$_3$ showed ruptured spherical structure with holes on the shell, which might be caused by airflow impact during calcination process. The ruptured structure and TEM image (Fig. S4c†) demonstrated its hollow characteristic, and it was noteworthy that double-shell hollow structures appeared. On the contrary, CeO$_2$ sample (Fig. S4b†) exhibited smoother surface and more complete spherical in shape with respect to those of Mn$_2$O$_3$, indicating stronger resistant ability to airflow impact for CeO$_2$. However, corresponding TEM image (Fig. S4d†) suggested that no double-shell structures were formed in CeO$_2$ hollow spheres. The result showed distinct behaviors between Mn$_2$O$_3$ and CeO$_2$ during the removing process of the templates. Otherwise, in the identical situations, CeO$_2$–MnO$_x$ hybrid oxide possessed more complete spherical double-shell hollow structures than Mn$_2$O$_3$ and CeO$_2$, which indicated that hybrid oxide was conducive to forming complete multi-shell hollow structures due to synergistic interaction. Besides, considering of the structural properties of Mn$_2$O$_3$ and CeO$_2$ hollow spheres, it was proposed that the diffusivity between manganese and cerium elements inward carbon spheres was different. Occasionally, HAADF-STEM mapping images of one triple-shell hollow sphere with non-dispersed particles were captured. As shown in Fig. S5†, an aggregation state was detected from the non-dispersed particles outside the shells, which mainly consisted of O and Ce, with scarcely any Mn element. Therefore, combined with the widespread homogeneous dispersion of the elements in triple-shell hollow spheres, it was deduced that, under the synthesis process, the existence of Mn element was synergistically helpful for CeO$_2$ to diffuse inward carbon spheres, and then the multi-shell hollow structure was formed.

The crystal structures of the samples were characterized by XRD, and the obtained patterns were displayed in Fig. 4. It was noted that the diffraction peaks of single component oxides of manganese and cerium were assigned to the characteristic reflections of Mn$_2$O$_3$ (bixbyte-C, JCPDS 31-0825) and CeO$_2$ (cubic fluorite, JCPDS 04-0593), respectively. While the hybrid multi-shell CeO$_2$–MnO$_x$ hollow spheres (single, double and triple referred to the hollow structured samples with corresponding shells, respectively) all exhibited cubic fluorite-type CeO$_2$ phase (JCPDS 04-0593) together with a weak diffraction peak of tetragonal hausmannite Mn$_3$O$_4$ (JCPDS 08-0017) phase at around 36.0°. The result was in good consistency with the research finding upon CeO$_2$–MnO$_x$ composites that when mole fraction of Mn was...
Discrepancy of manganese crystalline phase for MSCMHSs ($\text{Mn}_3\text{O}_4$: $\text{Mn}^{3+}$, $\text{Mn}^{3+}$) and mono-component $\text{Mn}_2\text{O}_3$ hollow spheres ($\text{Mn}^{3+}$) might be caused by synergistic interactions between Mn and Ce elements ($\text{Mn}^{3+} + \text{Ce}^{3+} \leftrightarrow \text{Mn}^{2+} + \text{Ce}^{4+}$). Meanwhile, the constituents and calcination rates also had an effect on the crystallinity of the samples. For the identical calcination rates, $\text{Mn}_3\text{O}_4$ and $\text{CeO}_2$ hollow spheres both exhibited higher crystallinity compared with double-shell $\text{CeO}_2$-$\text{MnO}_2$ hollow spheres. While with the identical constituents for MSCMHSs, higher calcination rates resulted in higher crystallinity. As listed in Table 1, crystallite sizes of samples were calculated by Scherrer equation based on the main diffraction peaks. It could be observed that the crystallite sizes for MSCMHSs were much smaller than that of $\text{Mn}_3\text{O}_4$ and $\text{CeO}_2$ hollow spheres due to the broader diffraction peaks. Furthermore, crystalline structure of the traditional $\text{CeO}_2$-$\text{MnO}_2$ nanoparticles (NP) was investigated as well. As discerned in Fig. 4, except for the $\text{CeO}_2$ and $\text{Mn}_3\text{O}_4$ phase, NP possessed extra characteristic lines ascribed to $\text{Mn}_3\text{O}_4$ phase compared to MSCMHSs, indicating that template-assisted method could promote the dispersity of manganese species. Besides, the crystallite size of NP was estimated to be slightly larger than MSCMHSs.

### 3.2 Catalytic performance results

Selective catalytic reduction of NO with NH$_3$ was utilized to evaluate the catalytic performances of MSCMHSs as well as NP samples. The reaction was carried out at the space velocity of 60 000 mL g$^{-1}$ h$^{-1}$ over the temperature range from 100 °C to 275 °C. As disclosed in Fig. 5a, the NO conversion of all samples increased as the temperature increased and then decreased with different activated temperature windows. The MSCMHSs reached more than 90% conversion at temperatures as low as 125 °C, and it was noteworthy that hollow spheres with different number of shells exhibited different catalytic abilities. Among them, the triple-shell hollow spheres with a maintained conversion at 100% from 150 °C to 250 °C had the best performance. The NO conversion of double-shell hollow spheres was similar to that of the triple-shell ones from 100 °C to 200 °C but then decreased with the temperature increasing. Moreover, the single-shell hollow spheres showed the worst activity compared with the double-shell and triple-shell hollow spheres. However, it still displayed superior performance in contrast with NP, which could just achieve 84.3% conversion as the maximum activity value at 200 °C. Thus, an apparent enhancement in NO conversion for MSCMHSs than NP was observed within the range of testing temperature, indicating the positive role of multi-shell hollow structures in the reaction of NH$_3$-SCR, especially at low temperatures (100–150 °C). Otherwise, the N$_2$ selectivity of the samples declined gradually along with the rising temperature (Fig. 5b), this may due to the strong redox behavior of Mn-based catalysts, which could lead to the oxidation of NH$_3$ by O$_2$ to the by-product, for instance, NO$_2$ and N$_2$O.$^{14}$

As discussed above, the catalytic performances of the samples ranked as the order of triple-shell > double-shell > single-shell > NP. Interestingly, the order of performances showed an inversed correlation with the crystalline sizes of the corresponding samples, just as the reports that smaller crystalline size was good for the enhanced activities.$^{35,36}$ Moreover, the superior catalytic activity of MSCMHSs might be attributed to the synergetic effect between active species of Mn and Ce elements.

To understand the H$_2$O resistance performance of the samples, the NH$_3$-SCR reaction under the condition with water...
at 150 °C was performed. As displayed in Fig. 6a, after the H2O was added into the mixture reactant gases, a decrease from 80% to less than 40% of NO conversion for NP samples was detected. At the same time, the NO conversion of the MCMSHSs declined as well but maintained at around 70% for more than 14 h, exhibiting better H2O resistance performance than NP samples. It was obvious that the addition of H2O caused deactivation of the samples in some degree, which could be explained that the adsorption of H2O on the surface of the catalyst led to the formation of hydroxyl, resulting in a decline in activity.37–39 Besides, an enhancement in N2 selectivity (Fig. 6b) was also observed among the samples after the addition of H2O, and this could be ascribed to the inhibition of H2O to the oxidation of NH3 by O2.38 To further investigate the key factor for the improved NH3-SCR activity and the effects of special multi-shell hollow structures on the properties of the catalysts, characterizations of different samples were deeply examined.

3.3 N2-Sorption characterization (BET)

N2-Sorption isotherms were measured to get insights into the textural properties of the samples. It could be seen from Fig. 7a that all the isotherms of hollow spheres belonged to typical type-IV in the light of IUPAC classification, indicating the existence of mesoporous structure.40 The observed H2-type hysteresis loops of CeO2 hollow spheres manifested uniform pore structures. However, Mn2O3 and MCMSHSs possessed H3-type hysteresis loops, which might symbolize an irregular pore size distribution, just in keeping with pore distribution curves in Fig. 7b. It was noted that MSCMHSs samples showed similar broad pore size distribution with the average pore diameters being at around 30.0 nm. While Mn2O3 and CeO2 hollow spheres exhibited narrower pore size distributions and smaller average pore diameters centered at 3.50 nm and 24.0 nm, respectively. As summarized in Table 1, there were no significantly differences among single-shell (36.0 m2 g−1), double-shell (30.0 m2 g−1) and triple-shell (35.0 m2 g−1) hollow spheres, which were smaller than 78.4 m2 g−1 of CeO2 but larger than 22.6 m2 g−1 of Mn2O3. Thus, compared to monocomponent oxides, the composite of manganese and cerium resulted in modifications on not only the inner structures but also the textural properties.

The adsorption–desorption isotherms and pore distribution curves of CeO2–MnO2 nanoparticles (NP) was shown in Fig. 8. It was observed that, compared with MSCMHSs, NP showed a narrower pore size distribution while the same main distribution location. It could be explained by the removal of carbon spheres for the precursor during calcination, which might cause non-uniform pore size distribution of the obtained MCMSHSs, containing interstitial pores between the particles. Generally, the surface area of catalysts acted an important role in catalytic performance. However, despite the larger surface area (43.5 m2 g−1), NP exhibited poorer activity compared to MSCMHSs, which suggested that multi-shell hollow structures played more crucial roles in catalytic performances.

3.4 Bulk and surface analyses of the catalysts (XRF and XPS)

XRF characterization was performed to discover the bulk composition of the samples, and the result was summarized in Table 2. It could be found that the actual ratios of the two metal elements were approximately the same as the theoretical value for each sample, indicating the feasibility of the preparation process with negligible mass losses.

XPS was conducted to investigate the surface composition and existential state of active species on the samples. Herein, XPS spectra of the MSCMHSs and NP in Ce 3d, Mn 2p and O 1s regions were displayed in Fig. 9, with the peaks being fitted to several designated peaks based on the Gauss–Lorentz curves. The complicated Ce 3d spectrum of all the samples was separated into eight components with the allocation defined in Fig. 9a. The series of peaks labeled u were attributed to Ce 3d3/2 spin–orbit states, whereas those labeled v belonged to Ce 3d5/2 spin–orbit states. Specially, the two peaks signed as u’ and v represented Ce3+ species, and the other six peaks (u, v, u”, v”, u‴, u‴”)
and $v''$) were ascribed to Ce$^{3+}$ species. It was obvious that both Ce$^{3+}$ and Ce$^{4+}$ species were co-existed in all the samples and the Ce$^{4+}$ was the predominant chemical valence. In general, the existence of Ce$^{3+}$ often led to the increase of active oxygen species on the surface of catalysts, which were in favor of the redox reaction. Based on the fitting areas of the peaks, the ratios of Ce$^{3+}$/Ce$^{4+}$ of the samples were calculated and listed in Table 2, which ranked in the order of triple-shell > double-shell > single-shell > NP. The results suggested that, with the multi-shell hollow structures, MSCMHSs possessed more active Ce$^{3+}$ species than NP, thus leading to the enhanced catalytic performance.

Fig. 9b exhibited two main peaks in Mn 2p spectrum of the samples at about 653.2 eV and 641.5 eV. There were mixed valences in manganese species on the surface of the samples, and the binding energies were overlapped. The peak at 653.2 eV was attributed to Mn 2p$^1/2$ spin–orbit states, while the peak at 641.5 eV was due to Mn 2p$^3/2$ spin–orbit states. As for Mn 2p$^3/2$ spectrum, the peak was fitted to three characteristic peaks representing Mn$^{2+}$ (640.7 eV), Mn$^{3+}$ (642.1 eV) and Mn$^{4+}$ (643.6 eV) species, respectively. The existence of Mn$^{2+}$ species for MSCMHSs proved the synergistic interaction between manganese and cerium oxide species, which was in good conformity with the above discussion.
with the results of XRD. Previous studies revealed that manganese species of higher oxidation state usually acted more important roles in redox reaction. From Table 2, it was observed that the surface Mn⁴⁺ concentration of MSCMHSs were higher than NP. Especially, the triple-shell hollow spheres showed higher Mn⁴⁺ concentration than double-shell and single-shell hollow spheres. The result indicated the superiority of multi-shell hollow structures on surface active species, which was evidenced by the catalytic performances.

O 1s spectra of the samples (Fig. 9c) were ascribed to two overlapped peaks denoted as Oₐ (531.2-531.4 eV) and O₈ (529.2-529.4 eV), respectively. Peak Oₐ with the higher binding energy was referred to the surface chemisorbed oxygen, correlated to the surface defects of samples. While peak O₈ was associated with lattice oxygen, bonding to the manganese or cerium cations. Due to its remarkable mobility, Oₐ was regarded to be the most active oxygen species on the surface of catalysts. Besides, the higher content of Oₐ relative to total oxygen concentration was able to bring better activity in oxidation reaction of NO to NO₂, which could accelerate the NH₃-SCR reaction through a “fast SCR” process. The listed values of Oₐ/(Oₐ + O₈) in Table 2 stated that the chemisorbed oxygen species of MSCMHSs were higher than NP just as that order of surface Ce³⁺ contents as well, accordant with the report that the existence of Ce³⁺ could trigger the formation of active oxygen species.

Fig. 9 XPS spectra of the samples in (a) Ce 3d, (b) Mn 2p and (c) O 1s regions.

Fig. 10 H₂-TPR profiles of the samples as a function of temperature.
As manifested in above results of surface analysis, it was worth noting that the surface active species contents of the samples, which were beneficial to redox process, possessed the identical order with their catalytic performances. Hence, it seemed that the enriched surface active species of MSCMHSs could be essential for their better activities in SCR reaction compared with NP.

### 3.5 Redox performance

Redox performance of the catalysts that plays an important role in NH₃-SCR activity was measured by H₂-temperature programmed reduction (H₂-TPR) characterization. As demonstrated in Fig. 10, two series of broad reduction peaks could be obtained. In the case of lower temperature peak from 150 °C to 600 °C, two kinds of reduction were obtained, with one attributed to the step-by-step reduction of MnO₂ species, and the other reduction of surface CeO₂ species.²⁴ Besides, the peak at higher temperature from 700 °C to 800 °C corresponded to the reduction of bulk CeO₂ species.²⁴ By means of the Gauss–Lorentz fitting, the lower-temperature broad reduction peak of all the samples was fitted to three reduction peaks characterized by apparent maxima at around 249, 330 and 410 °C, respectively. As reported,¹⁶,²⁴,²⁵ the low-, middle-, and high-temperature reduction peak could be respectively attributed to the reduction of MnO₂ into Mn₂O₃, Mn₂O₃ into Mn₃O₄ and Mn₃O₄ into MnO₂ overlapped by reduction peak of surface Ce⁴⁺. The discrepancy of redox performance showed that different preparation procedures brought different interaction between manganese oxide and cerium oxide in the samples.²⁵ Compared with NP samples, the MCMSHSSs, especially triple-shell ones, exhibited more superficial manganese species in a higher oxidation state (Mn⁴⁺) and much broader reduction temperature range, which was consistent with the XPS analyses. Moreover, NP samples exhibited much higher Mn₂O₃ manganese species, just as shown in XRD characterization.

Moreover, the effect of unique hollow spheres on gases diffusion also acted an inspiring role in catalytic performances.²⁷ In contrast with NP, despite the smaller surface areas, it was considered that multi-shell hollow spheres with porous structures could promote gases diffusion into internal spaces. Thus, thorough contact of reactive gases with both internal and external of spheres was obtained, and then enhanced reactive surfaces were achieved. In addition, hollow spheres with more shells were demonstrated to be more effective to enhance the catalytic performances than those with fewer shells, which should be attributed to multiple collisions of reactive gases among different shells. As illustrated in Fig. 11, more effective collisions tended to appear in hollow structures with more shells, which exhibited richer surface active species as well. As a result, CeO₂–MnO₂ catalysts with multi-shell hollow spherical morphology showed admired performances in NH₃-SCR reaction. The present study demonstrated that the well-designed morphology and structure offered an excellent strategy to the creation of highly efficient catalysts.

### 4 Conclusions

MSCMHSs with the average size of 400 nm were prepared successfully by the hard template method of carbon spheres, during which it was demonstrated that different calcination rate brought hollow spheres with different inner shell structures. The composite of manganese and cerium resulted in more complete hollow structures compared to monocomponent oxides, accompanied by modifications of crystalline phases, valence states and textural properties. Especially, the existence of Mn element could be helpful for CeO₂ to diffuse inward carbon spheres to form multi-shell hollow structures under experimental conditions. Furthermore, superior activities of MSCMHSs for NH₃-SCR reaction than traditional nanoparticles were displayed, and especially, triple-shell hollow spheres exhibited better performance than double-shell and single-shell ones, which could be attributed to the special structure and abundant surface active species, providing a new kind of material for low-temperature NH₃-SCR reaction.

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