A Highly Reactive and Sinter-Resistant Catalytic System Based on Platinum Nanoparticles Embedded in the Inner Surfaces of CeO₂ Hollow Fibers

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The catalytic properties of a supported system are strongly dependent on the types of metal and supporting material involved.\(^{[1–4]}\) Platinum nanoparticles (NPs) supported on ceria (CeO₂) have shown higher catalytic activities for a wide variety of reactions, including water-gas shift,\(^{[5,6]}\) CO oxidation,\(^{[7]}\) and hydrogenation\(^{[8,9]}\) when compared to those supported on other oxides. Therefore, CeO₂ has been widely employed as a support for manufacturing automotive catalysts because of its peerless oxygen storing/releasing capabilities as well as its superior ability to stabilize noble metals.\(^{[10,11]}\) In particular, Pt/CeO₂ catalysts are known to exhibit strong metal–support interaction effects with a potential to enhance the catalytic activities for reactions involving rapid oxygen and/or electron transfer between the metal and the support.\(^{[12–15]}\) Although there are many benefits in utilizing the Pt/CeO₂ system in real-world catalytic applications, great challenges, such as low thermal stability and loss of catalytic activity owing to sintering, still need to be addressed. In catalysts for automotive exhaust treatment, these problems are partially mitigated by combining CeO₂ with ZrO₂ in a solid solution, but precious metal sintering remains a primary deterioration mechanism.

The use of nanostructured composite materials is one strategy to address the sintering issue. Yan and co-workers recently synthesized CeO₂ nanoparticles covered with PtNPs and then SiO₂ shells to protect the PtNPs from aggregation during calcination,\(^{[16]}\) but their approach required multiple steps to generate the SiO₂ shells and then dissolve them. As a major drawback, the Pt/CeO₂ nanocomposites could only withstand calcination up to 450°C after the removal of the SiO₂ shells. Tsang and co-workers demonstrated the synthesis of a Pt/CeO₂ core–shell NPs by a modified microemulsion method.\(^{[17,18]}\) The resultant catalyst showed significantly improved catalytic activity and selectivity in water-gas shift over methanation by controlling the thickness of the CeO₂ protective layer. However, the Pt/CeO₂ core–shell NPs exhibited considerable aggregation in the reaction medium, which may negatively impact their catalytic activity in a practical application.

Herein, we report a simple, template-based procedure for the fabrication of CeO₂ hollow fibers with PtNPs embedded in the inner surfaces (Figure 1). The first step involved the generation of polystyrene (PS) fibers by electrospinning, followed by plasma treatment to render their surfaces hydrophilic for the deposition of PtNPs. The deposited PtNPs were almost completely encapsulated by CeO₂, which may negatively impact their catalytic activity in a practical application.
The nonwoven mat of PS fibers was prepared by electrospinning a 20 wt% solution of PS in a mixture of tetrahydrofuran (THF) and dimethylformamide (DMF) at a mass ratio of 1:1. The average diameter of the as-spun PS fibers was $1.16 \pm 0.22$ μm (Supporting Information, Figure S1a). The PtNPs stabilized by poly(vinyl pyrrolidinone) (PVP) were prepared using a method based on polyol reduction.\textsuperscript{19} The as-synthesized PtNPs were uniform in size (Figure S1b), with an average diameter of 3.5 nm (Figure S1c). The surface of a plasma-treated PS fiber could be easily coated with the PtNPs by immersing the fiber mat in a suspension of PtNPs in ethanol, followed by washing to remove the loosely bound NPs and drying in air (Figure S1d).

The PS fiber decorated with PtNPs was then employed as a substrate for the in situ growth of a CeO\textsubscript{2} sheath by a process involving reaction of cerium(III) acetate with 6-aminocaproic acid (6ACA) in an aqueous medium in air.\textsuperscript{20} As shown in Figure 2a, the CeO\textsubscript{2} coating was deposited uniformly on the surface of each Pt/PS fiber and no interfiber connection was observed. The average diameter of the CeO\textsubscript{2} sheath was estimated to be about 100 nm in thickness. Based on an XRD pattern (Supporting Information, Figure S2a), the as-deposited CeO\textsubscript{2} coating was crystallized in a cubic fluorite structure ($Fm\overline{3}m, a = 5.411$ Å, JCPDS Card No. 34-0394). It is remarkable that a complete sheath made of crystalline CeO\textsubscript{2} nanoparticles could be formed on the PS fiber under relatively mild reaction conditions (60°C for 2 h). In this case, the PtNPs played an important role in the nucleation and growth of CeO\textsubscript{2} nanocrystals in the presence of 6ACA. Because the PVP-stabilized PtNPs were negatively charged with a zeta potential of $-0.83$ mV under a slightly basic pH (7.5), the Ce\textsuperscript{3+} cations could easily adsorb onto the PtNPs to facilitate the nucleation and growth of CeO\textsubscript{2} nanocrystals. Although the details for CeO\textsubscript{2} growth still need to be fully investigated, our TEM observations did provide some evidence to support this proposed mechanism. The organic components in the as-prepared CeO\textsubscript{2}/Pt/PS structures, including PS, PVP, and 6ACA, could be removed by calcination in air at an elevated temperature to generate CeO\textsubscript{2} hollow fibers with PtNPs embedded in the inner surfaces. The SEM image in Figure 2b and the TEM image in the inset clearly show a hollow structure for the CeO\textsubscript{2} fibers. The loading of Pt in the as-prepared Pt\textsubscript{encap}/CeO\textsubscript{2} hollow fibers was about 0.98%, as determined by inductively coupled plasma mass spectrometry (ICP-MS).

Figure 3 shows TEM images of the Pt\textsubscript{encap}/CeO\textsubscript{2} hollow fibers after they had been further treated at different temperatures in air for 2 h. The products obtained at 400 and 800°C still maintained a tubular structure without being collapsed, demonstrating a rather good thermal stability for the relatively thin CeO\textsubscript{2} sheath. Furthermore, the PtNPs were effectively prevented from sintering up to 700°C as a result of the CeO\textsubscript{2} barrier surrounding each PtNP. At 800°C (Figure 3f), both sintering of PtNPs and growth of CeO\textsubscript{2} crystallites started to occur, which was manifested by the formation of Pt large agglomerates and a substantial increase in grain size for CeO\textsubscript{2}. The X-ray powder diffraction pattern (Supporting Information, Figure S2a) taken from these samples confirmed that the CeO\textsubscript{2} phase remained to be in a cubic fluorite structure. For the product obtained at 800°C, a weak peak at 39.6 degrees appeared, which could be assigned to the (111) diffraction of Pt, indicating the formation of Pt large agglomerates by sintering. The changes in size for the embedded PtNPs (measured from the TEM images) and the CeO\textsubscript{2} crystallites (derived from the XRD patterns using the Debye–Scherrer equation) after the samples had been
treated in air at different temperatures are shown in the Supporting Information, Figure S2b. Up to 400°C, no significant change in size was observed for both the PtNPs and CeO2 crystallites. When the temperature was increased from 400 to 700°C, the average sizes of the PtNPs and CeO2 crystallites were increased from 3.5 to 4.1 nm and from 3.2 to 5.1 nm, respectively. As the temperature was further increased to 800°C, the PtNPs grew to 12 nm and the CeO2 crystallites also grew to 11.2 nm, which is more than triple relative to their original sizes. The missing diffraction peak at 39.6° for Pt for all the samples until 700°C also suggested that the new Pt/encap/CeO2 system could resist thermal sintering up to this temperature. Above 700°C, the simultaneous sintering of CeO2 crystallites may have induced and facilitated sintering of the PtNPs, leading to a quick increase in size for the PtNPs. We also tried to resolve possible changes to the morphology of PtNPs and the platinum–oxide interface by high-resolution TEM (Supporting Information, Figure S3), but it was very difficult to obtain any meaningful data owing to the presence of a polycrystalline CeO2 matrix around each PtNP.

We measured the surface areas of the PtNPs embedded in the Pt/encap/CeO2 hollow fibers by CO titration of chemisorbed oxygen (Supporting Information, Table S1). The titrations were performed at −20°C to minimize oxygen spillover to Pt sites from CeO2. For the samples treated at 400, 600, 700, and 800°C in air for 2 h, the accessible Pt surface areas and the corresponding Pt dispersions were found to be in the ranges of 0.3–4.1 m² g⁻¹ and 0.1–1.7%, respectively. Unexpectedly, CO titration failed to give a measurable Pt surface area and dispersion for the Pt/encap/CeO2 sample calcined at only 400°C. It is possible that organic residues from the incomplete combustion of PVP and PS might have blocked the channels in the CeO2 sheath so that the O₂/CO analytes could not access the surface of PtNPs. After calcination at 600°C, which presumably allowed more of the organic residues to be burned away, CO titration gave a Pt surface area of 2.6 m² g⁻¹ and dispersion of 1.2%. The highest accessible Pt surface area (3.4 m² g⁻¹) and dispersion (1.7%) were obtained after calcination at 700°C. By further increasing the calcination temperature to 800°C, the surface area and dispersion dropped to 0.8 m² g⁻¹ and 0.4%, respectively, which is consistent with the sintering of PtNPs observed by TEM and XRD.

Figure 4 compares the TOFs of CO oxidation reaction obtained for two samples of Pt/encap/CeO2 hollow fibers and a Pt/SiO2 catalyst reported by Cant and co-workers.[21] The Pt/SiO2 TOFs fit the same Arrhenius curve extrapolated from higher-temperature measurements under the same conditions on clean Pt(100) and clean, well-defined Pt nanoparticles of different sizes supported on planar SiO2 substrates,[22] and thus represent the expected structurally insensitive activity of normal Pt sites. Also shown are TOFs for a Pt/TiO2 catalyst prepared in the same way as the structure shown in Figure 2b,[23] with a Pt dispersion of 31% by CO titration. These TOFs are almost identical to Pt/SiO2, as expected for the normal metallic Pt sites. The Pt/encap/CeO2 hollow fibers exhibited TOFs three orders of magnitude higher than those of the Pt/SiO2 and Pt/TiO2 catalysts within the comparable temperature range of 384–476 K, providing a direct evidence to support its far superior catalytic activity. These TOFs were calculated based on the Pt dispersion measured by CO titration (Supporting Information, Table S1), which were essentially unchanged after these catalytic rate measurements (not shown), but which were about 10-fold less than what was measured from the size of the PtNPs using TEM. Even considering the worst possible experimental errors in dispersion, the activity of the Pt/encap/CeO2 hollow fibers was still more than 100-fold higher than the Pt/SiO2 and clean Pt model catalysts. Also, if we used the average particle size of 4.1 nm measured by TEM after calcination at 700°C to calculate the Pt surface area (assuming no poisoned Pt sites), it gave 40 m² of Pt. Based on this area, we still calculate a TOF that was ca. 100-fold higher than that of the Pt/SiO2. Moreover, it is worth pointing out that the Pt/SiO2, Pt/Al2O3, and Pt/CeO2/Al2O3 catalytic systems studied by other groups[24] all exhibited much lower TOFs than that of Pt/encap/CeO2 hollow fibers under the same conditions with O₂/CO ratio = 0.5. In studies of CO oxidation over traditionally prepared Pt/CeO2 catalysts, it was reported that the CO oxidation activity of Pt, which ended up partially covered and/or encapsulated by a thin layer of CeO2, was higher than that of Pt itself.[25,26] However, those prior studies only showed a 3-4-fold increase in TOF, which is much less than the 1000-fold increase observed here. The CeO2 layer on Pt was suggested to accelerate the overall reaction rate by participating in accepting and donating oxygen atoms, which compensated the limited accessibility of reactants to the catalytic sites.[27,28] A roughly 100-fold increase in Pt activity owing to CeO2 was reported when the CO/O2 ratio was about 1000, which could be attributed to oxygen spillover to the Pt from the CeO2 if the Pt sites were too saturated by...
CO to dissociate O₂ but no increase owing to CeO₂ was observed in that study at the stoichiometric ratio used here.[29] For the hollow Ptencap/CeO₂ fibers described here, the CeO₂ nanocrystalline sheath tightly surrounding the PtNPs with a maximized Pt–CeO₂ interface greatly benefitted the catalytic CO oxidation rate. Further studies on the mechanism of CO oxidation using the Ptencap/CeO₂ system are required to understand this phenomenon.

In summary, we have demonstrated a method for embedding Pt NPs in the inner surfaces of CeO₂ hollow fibers with open ends to generate a new catalytic system that could resist thermal sintering up to 700 °C. The TOF of the as-synthesized Ptencap/CeO₂ hollow fibers for CO oxidation was 2–3 orders of magnitude higher than those based on the conventional Pt/SiO₂ catalytic system, which could be attributed to a unique structure and composition for maximizing the Pt encap/CeO₂ interface. Future work will involve the introduction of a protective layer based on oxides, such as SiO₂ or Al₂O₃, to interface. Further studies on the mechanism of catalytic CO oxidation rate. Further studies on the mechanism of CO oxidation using the Ptencap/CeO₂ system are required to understand this phenomenon.

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