Microporous Organic Network Hollow Spheres: Useful Templates for Nanoparticulate Co₃O₄ Hollow Oxidation Catalysts

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Supporting Information

ABSTRACT: Hollow microporous organic networks (H-MONs) were prepared by a template method using silica spheres. The shell thickness was delicately controlled by changing the synthetic conditions. The H-MONs were used as a template for the synthesis of nanoparticulate Co₃O₄ hollows which showed excellent catalytic performance in H₂O₂ oxidation.

Microporous organic networks (MONs) are a new class of functional materials.1 The MONs have been prepared through various carbon–carbon coupling reactions between organic building blocks.1,2 For example, Cooper research group and others have prepared various microporous organic polymers through Sonogashira coupling between alkynes and arylhalides.3 The porosity and functionalities of MONs were controlled by the tailored use of building blocks.1–3 The resultant materials have been applied for gas adsorbents and catalysts.1–3 In addition to inner porosity, the overall shape of porous materials is also important for their performance. To interact with inner working sites in adsorption or catalysis, guest molecules diffuse into pore structures. The diffusion pathways are dependent on the shape of materials and can be synthetically engineered. As an example of shape effects in material science, hollow spheres showed better efficiencies in electrochemical or photocatalytic performance than nonhollow ones.4 In this regard, various hollow structures were designed and prepared.5 However, to the best of our knowledge, the shape controlled synthesis of hollow MONs is less explored.6

Templated synthesis is a straightforward method for the shape control of materials, because the resultant shapes originating from the sacrificed templates are quite predictable.7 For example, silica has been used as a template because it can be removed easily through chemical etching by reaction with HF or NaOH.8 Thus, the formation of MON on the surface of silica will be quite an interesting research subject because, if successful, the hollow MONs can be obtained through the chemical etching of silica templates.

In addition to inorganic templates, organic ones have been developed and used for the shape controlled synthesis of inorganic materials.9 Because the organic templates can be removed easily through heat-treatment under air, shape-controlled inorganic products can be obtained. If the organic templates have porosity, inorganic precursors can be readily absorbed into the pores to result in the homogeneity of shapes.

Moreover, due to the porosity of templates, particulate walls can be obtained, which can maximize the surface area of materials.

Our research group has studied the synthesis of functional MONs6a,10 and the shape controlled synthesis of inorganic nanomaterials,11 separately. In this work, we report the templated synthesis of hollow MON spheres (H-MONs), their successful use as templates for nanoparticulate cobalt oxide hollow spheres, and the excellent catalytic performance as oxidation catalysts.

Figure 1 shows the synthetic scheme for the H-MONs using a silica template. First, monodisperse silica spheres were prepared by Stöber method12 and well-dispersed in a mixture of toluene and triethylamine. Then, palladium catalyst and copper iodide were added and the reaction mixture was stirred at room temperature for 1 h. Tetra(4-ethyl)phenylmethane and 2 equiv of 1,4-diiodobenzene were added. The reaction mixture was heated at 100 °C for 2 days. The resultant yellow solids were isolated by centrifugation and washed with methylene chloride, acetone and methanol. After drying under vacuum, the materials were treated with HF solution.

Figure 1. Synthesis route for hollow microporous organic networks (H-MONs) using a silica template.

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for 2 h. The resultant materials were isolated by centrifugation, washed with water, methanol and acetone, and dried under vacuum.

The materials isolated during the synthetic process were investigated by scanning (SEM) and transmission electron microscopy (TEM). The silica templates had a narrow distribution with a 545 ± 17 nm average size (Figure S1). According to TEM studies, the MON shells with a lighter contrast were clearly distinguished from inner darker silica. As shown in Figure 2e, the MON coatings on silica spheres were quite homogeneous. After the chemical etching of the inner silica with HF, the resultant materials showed a hollow structure (Figure 2f). Through the screening of various synthetic conditions such as solvent and amount of reagents, synthetic methods for thickness control of MON shells were discovered. The four representative synthetic conditions were chosen and the resultant hollow MONs are displayed in Figure 2a–d.

In the case of the solvent system, as the amount of triethylamine increased from 10 mL (H-MON-1) to 15 mL (H-MON-2) and 30 mL (H-MON-3) with a fixed total amount (30 mL) of solvent, the thickness of MON shells gradually decreased from 81 ± 6 to 51 ± 3 and 40 ± 2 nm, respectively (Figures 2a–c and 3a) Also, as expected, when the amount of silica increased from 300 mg (H-MON-2) to 400 mg (H-MON-4), the thickness of shells decreased from 51 ± 3 to 22 ±2 nm, respectively (Figures 2b,d and 3a) When 500 mg of silica was used with 50 mg of tetra(4-ethyl)phenylmethane, incomplete hollow structures were dominantly observed (Figure S2).

The analysis of N2 sorption isotherms based on the Brunauer–Emmett–Teller (BET) theory showed microporosity (<2 nm pores) (Figures 3b, inset, and S3). The pore size distribution analysis by density functional theory (DFT) method revealed that the MONs have microporosity (<2 nm pores) (Figures 3b, inset, and S3). The solid phase 13C NMR spectra of MONs matched well with the expected structure, showing 13C peaks at 90 and 63 ppm from C=C triple bonds and benzylcarbons, respectively (Figure 3c). Thermogravimetric analysis showed that the MONs start to decompose at 240–280 °C (Figure S4). Powder X-ray diffraction studies (PXRD) showed an amorphous character of H-MONs, as commonly observed for MONs prepared by Sonogashira coupling in the literature (Figure S5).

In addition to various application fields reported for nonhollow MONs in the literature,1–3 the hollow MONs can be applied for new purposes. In this study, considering the existence of pores in walls, the hollow MONs were applied as an organic template for inorganic hollows (Figure 4a). With an organometallic precursor (cobalt octacarbonyl), cobalt species were introduced into the pores of H-MONs. While we studied the shell thickness effect of H-MON-2 and H-MON-4 as organic templates, the resultant Co3O4 hollows showed no significant differences in physical properties such as nanoparticulate shape of walls and surface area (60–67 m2/g). Thus, we used the H-MON-2 as a representative template. The cobalt oxide/H-MON composites were calcinated at 500 °C for 5 h under air to remove the organic template. The TEM analysis of the hollow cobalt oxides showed the interconnected nanoparticulate structure of walls (Figure 4b,c). High resolution (HR) TEM analysis clearly showed the nanoparticulate

**Figure 2.** TEM images of hollow MONs (H-MONs) prepared using 50 mg of tetra(4-ethylphenyl)methane, 80 mg of 1,4-diiodobenzene, 300 mg of silica, and 20 mL/10 mL (a, H-MON-1), 15 mL/15 mL (b, H-MON-2), and 0 mL/30 mL (c, H-MON-3) of toluene/triethylamine. TEM image of H-MON prepared using 50 mg of tetra(4-ethylphenyl)methane, 80 mg of 1,4-diiodobenzene, 400 mg of silica (d, H-MON-4), and 15 mL/15 mL of toluene/triethylamine. TEM image (e) of silica/MON core/shell materials for H-MON-2. (f) Magnified TEM image of H-MON-2.

**Figure 3.** (a) Thickness distribution diagram of H-MON shells. (b) N2 adsorption/desorption isotherms at 77 K, pore size distribution diagram (DFT method, inset) of H-MON-2. (c) Solid-phase 13C NMR spectra of H-MONs.
domains and high crystallinity. (Figure 4c). The major (311) crystalline planes with 0.24 nm distance of Co$_3$O$_4$ were dominantly observed. According to N$_2$ sorption isotherm analysis, 64 m$^2$/g of surface area and 0.32 cm$^3$/g pore volume were measured (Figure 4d). The high surface area is attributed to the nanoparticulate walls. The average size of particles in walls was calculated as 7.6 ± 1.2 nm by measuring 231 particles (inset Figure 4d). PXRD showed that the obtained materials are cubic Co$_3$O$_4$ (JCPDS #74-2120) (Figure 4e). X-ray photoelectron spectroscopy (XPS) showed Co 2p$_{3/2}$ orbital peak at 780.3 eV, matching well with the values of Co$_3$O$_4$ materials (inset Figure 4e).

Recently, nanostructured cobalt oxides have been applied for environmental oxidation catalysts. For example, cobalt oxides have been applied as oxidation catalysts for the removal of gas pollutants such as CO and NO. In addition, the cobalt oxides have been used for H$_2$O$_2$ oxidation for the ultimate removal of organic pollutants in water through the reaction of resultant radical (·OH) species. Considering the high surface area of the nanoparticulate Co$_3$O$_4$ hollows, we studied their catalytic performance for H$_2$O$_2$ oxidation (0.20 M H$_2$O$_2$ aqueous solution, 25 mL). Figure 5 summarizes the results. Commercial Co$_3$O$_4$ nanoparticles (20 mg) showed 54 mL (89% of theoretical O$_2$ volume) of oxygen generation for 120 min with 2.4 × 10$^{-2}$ min$^{-1}$ rate constant. In comparison, the nanoparticulate Co$_3$O$_4$ hollows (20 mg) showed 57 mL (93% of theoretical O$_2$ volume) of oxygen generation within 30 min with 1.8 × 10$^{-2}$ min$^{-1}$ rate constant. The recovered Co$_3$O$_4$ catalysts retained catalytic activities in the successive runs. Even in the fifth run (57 mL O$_2$ generation after 30 min, 1.7 × 10$^{-1}$ min$^{-1}$ rate constant), there was no significant difference in catalytic activity, compared with that of the first run. The Co$_3$O$_4$ hollows recovered after the five runs showed nearly no change in PXRD, XPS, and TEM studies (Figure S6). Thus, the use of hollow MONs for the synthesis of nanoparticulate inorganic oxide materials is quite a successful synthetic strategy for the development of new oxidation catalysts.

In conclusion, this work suggests a synthetic method for shape controlled MONs. Using silica templates, monodisperse hollow MONs could be engineered through the chemical etching of a silica template. Moreover, the hollow MON spheres could serve as an organic template for nanoparticulate inorganic hollows. The resultant hollow Co$_3$O$_4$ showed excellent catalytic activities in H$_2$O$_2$ oxidation. We believe that the hollow MONs in this work can be applied for more diverse purposes including drug delivery. In addition, more diverse inorganic materials can be engineered through screening transition metal sources.

**ASSOCIATED CONTENT**

*S Supporting Information*

Experimental procedure, TEM image of silica templates, additional TEM image for MONs, N$_2$ adsorption/desorption isotherms, pore size distribution diagrams, TGA curves, and PXRD patterns of H-MONs, characterization of the nanoparticulate Co$_3$O$_4$ hollow spheres before and after five runs in catalytic H$_2$O$_2$ oxidation. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.
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(13) Unfortunately, underlying reasons are not clarified yet.


(17) Co3O4 nanopowder (<50 nm, Aldrich Co., Cat. No. 637025) was used.

(18) Recently, the mesoporous Co3O4 showing 1.1 × 10−11 min−1 rate constant at 25 °C was reported (ref 16a).