Rhodium(0) Nanoparticles Supported on Nanocrystalline Hydroxyapatite: Highly Effective Catalytic System for the Solvent-Free Hydrogenation of Aromatics at Room Temperature

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Supporting Information

ABSTRACT: The hydrogenation of aromatics under mild conditions remains a challenge in the fields of synthetic and petroleum chemistry. Described herein is a new catalytic material that shows excellent catalytic performance in terms of activity, selectivity, and reusability in the hydrogenation of aromatics in solvent-free systems under mild conditions. The catalyst, consisting of rhodium nanoparticles supported on nanocrystalline hydroxyapatite, can quantitatively hydrogenate neat benzene to cyclohexane with exceptionally high rates (initial TOF > 10^3 h^-1) at 298 K and 3 bars of initial H2 pressure. This new material maintains its inherent catalytic activity after several reuses. Importantly, catalyst preparation does not require elaborate procedures because the active metal nanoparticles are readily formed from the in situ reduction of Rh3+-exchanged hydroxyapatite while submerged in the aromatic solvent at room temperature under 3 bars of H2 pressure.

INTRODUCTION

The hydrogenation of aromatic molecules is an important reaction in the petroleum and chemical industries. For instance, cyclohexane obtained from the complete hydrogenation of benzene is a high-volume intermediate used in the production of Nylon 6. New regulations call for a drastic reduction in the aromatic content in fuels because these molecules have been shown to be highly detrimental toward both fuel quality and human health. Most studies regarding aromatic hydrogenation have used benzene as the model aromatic feed molecule, and transition-metal nanoparticles and organometallic complexes have been thoroughly investigated as catalysts for the saturation of its C=C bonds. Indeed, benzene is notoriously difficult to hydrogenate when compared to olefins because of the resonance stabilization imparted by the aromatic ring. Thus, hydrogenation reaction conditions often require temperatures >373 K, H2 pressures >30 bar, and/or dilution of the aromatic substrate with a solvent, leading to higher byproduct generation and more energy-intensive product separation procedures. Very few reports exist on catalytic schemes that perform the complete hydrogenation of benzene under mild conditions (i.e., ≤300 K and <10 bars of H2) and under solvent-free conditions with acceptable rates. Unfortunately, most of the catalysts used in these schemes suffer from either low activity, low stability, or difficulty in synthesis procedures (Table S-1 in the Supporting Information). Therefore, the development of a simply prepared, highly active, reusable catalyst that operates under mild and solvent-free conditions remains a difficult challenge in the hydrogenation of aromatics.

Recently, hydroxyapatite ([Ca10(OH)2(PO4)6], HAp) has generated great interest in view of its potential application in a wide variety of fields including in catalysis as a support with attractive properties. Specifically, HAp has the following advantages as a catalyst support: (i) reduced mass transfer limitations due to the absence of structural porosity, (ii) a high ion-exchange ability and adsorption capacity, and (iii) a low surface acidity, thus preventing any side reactions arising from the support itself. These physicochemical properties of HAp prompted us to focus on the use of the HAp matrix in the stabilization of metal nanoparticles. The reduction of the HAp matrix particle size from the microcrystalline to the nanocrystalline regime (from >1 μm to <100 nm) was performed in anticipation of improved activity due to a higher external surface area, a larger number of exchange sites, and lower mass-transfer limitations for liquid-phase hydrogenation at low temperature.

In this letter, we report our results on the preparation and characterization of a catalyst consisting of rhodium(0) nanoparticles (2.7 ± 1.1 nm) supported on a nanocrystalline HAp matrix, hereafter referred to as Rh(0)@HAp, and on its remarkable catalytic performance in terms of the activity, selectivity, and reusability in the hydrogenation of aromatics in solvent-free systems under mild conditions. Rh(0)@HAp can be reproducibly prepared with exceptional simplicity by the ion exchange of Rh3+ ions with Ca2+ ions of the HAp matrix, followed by the reduction of Rh3+-exchanged nanocrystalline hydroxyapatite (Rh3+(@HAp)) in situ during the hydrogenation of an aromatic substrate. After an induction period, the resulting Rh(0)@HAp hydrogenates aromatic...
substances (e.g., benzene, toluene, m-xylene, o-xylene, and p-xylene) to their corresponding saturated alkanes at high rates under solvent-free, mild conditions (at 298 K and under 3 bars of initial H₂ pressure). More importantly, the supported rhodium(0) nanoparticles on the HAp matrix feature notable resistance against agglomeration and leaching. Indeed, when the isolated Rh@HAp catalyst is reused, it retains 92% of its initial catalytic activity after five cycles of quantitative hydrogenation of neat benzene into cyclohexane under the same reaction conditions.

**EXPERIMENTAL SECTION**

**Materials.** Synthetic hydroxyapatite nanocrystalline powder ([(Ca₅(OH)(PO₄)₃]ₓ, ≥99.995% trace metals basis), hydroxyapatite powder ([(Ca₅(OH)(PO₄)₃]ₓ), benzene (Chromasolv Plus, for HPLC, ≥99%), toluene (anhydrous, ≥99.8%), m-xylene (anhydrous, ≥99%), o-xylene (anhydrous, ≥99%), p-xylene (anhydrous, ≥99%), cyclohexene (puriss. p.a., ≥99.8%), and carbon disulfide (CS₂, anhydrous, ≥99%) were purchased from Sigma-Aldrich and used as received. Rhodium(III) chloride trihydrate (RhCl₃·3H₂O) and d-chloroform (CDCl₃) were purchased from Strem Chemicals and Cambridge Isotope Laboratories, respectively. Deionized water was distilled with a water purification system (Thermo Scientific Barnsted Nanopure System). All glassware and Teflon-coated magnetic stir bars were cleaned with acetone, followed by copious rinsing with doubly deionized water under ultrasonication, and finally dried in an oven at 423 K.

**Preparation of Rh³⁺@HAp and General Procedure for the In Situ Generation of Rh(0)@HAp During the Hydrogenation of Aromatics.** Rhodium(III) cations were introduced into Ca²⁺@HAp by the ion exchange of 500 mg of Ca²⁺@HAp in 20 mL of an aqueous solution of 25.3 mg of RhCl₃·3H₂O (0.09 mmol) for 4 h at room temperature. The sample was then filtered by suction filtration using a Whatman-1 filter (Ø = 9 cm), washed three times with 100 mL of deionized water, and dried at 353 K in the oven. The in situ formation of Rh(0)@HAp and the concomitant hydrogenation of olefinic or aromatic substrates were performed in a Fischer–Porter (F–P) pressure bottle connected to a line through Swagelock tetrafluoroethylene (TFE)-sealed quick connects and to an Omega-PX209-100GI pressure transducer interfaced through an Omega-UWPC-2-NEMA wireless transmitter to a computer using an Omega-UWT-REC2-D-V2 wireless receiver. The progress of an individual hydrogenation reaction was followed by monitoring the pressure of H₂ gas on the Omega-data logging and using the recording software program. In a typical experiment, 100 mg of Rh³⁺@HAp (4.86 μmol Rh) was weighted and placed in a new 22 × 175 mm pyrex culture tube containing a new 1/8 μm inner diameter stir bar. The culture tube was then placed inside the F–P bottle, and the entire setup was placed inside a constant-temperature circulating water bath thermostatted at 298 K and purged/evacuated for at least 30 min to remove any trace of oxygen and water. Next, 1 mL of substrate was added to the F–P bottle rapidly via a syringe with a long needle under hydrogen purging. Afterwards, the bottle was purged 10 times with H₂ (15 s per purge with stirring at 1000 rpm), and the reaction timer was started (t = 0 min). When no more hydrogen uptake was observed, the experiment was stopped, the F–P bottle was closed and disconnected from the line, and the hydrogen pressure was released. Finally, a small aliquot from the reaction solution in the culture tube was withdrawn for ¹H NMR analysis.

**Data Handling and Curve Fit of Hydrogenation Data.** The raw pressure versus time data collected with the computer-interfaced transducer were exported from Omega software program and imported into OriginPro 8, which was then converted into the equivalent concentration loss of substrate using the known stoichiometry (for example; 3:1 for H₂/C₆H₆ in the complete hydrogenation of benzene). The curve fitting of the [substrate] versus time data to the Finke–Watzky two-step mechanism was performed as described elsewhere using the software package OriginPro 8, which is a nonlinear regression subroutine and uses a modified Levenberg–Marquardt algorithm.

**RESULTS AND DISCUSSION**

The progress of Rh(0)@HAp formation and the concomitant hydrogenation of neat aromatics at room temperature were followed by monitoring changes in the hydrogen pressure and converting them into an equivalent substrate concentration loss using known hydrogenation stoichiometries (e.g., 3:1 H₂/substrate for the complete hydrogenation of benzene). As observed in Figure 1, the hydrogenation of neat benzene starting with the Rh³⁺@HAp precatalyst at 298 K and 3 bars of H₂ pressure follows sigmoidal kinetics that fit well with the Finke–Watzky (F–W) two-step nucleation and autocatalytic growth mechanism.⁹ The F–W mechanism models the formation of nanoparticles by considering a slow, continuous nucleation step (A → B, rate constant k₁) followed by an autocatalytic surface growth step (A + B → 2B, rate constant k₂) that results in a sigmoidally shaped hydrogenation curve.⁷ Thus, the complete hydrogenation of neat benzene catalyzed by Rh(0)@HAp containing 0.5 wt % Rh (as determined by inductively coupled plasma mass spectroscopy [ICP-MS] with 14.1 and 36.8 wt % P and Ca₃, respectively) was achieved within 5 h after an induction time period of ~6 h, as judged by the cessation of H₂ uptake and the quantitative production of cyclohexane (>99% yield confirmed by ¹H NMR). The excellent fit to F–W kinetics is indicative of the formation of metal(0) nanoparticles from the ion-exchanged precatalyst in the presence of a reducing agent at room temperature.¹⁰ We note that none of the partially hydrogenated benzene reduction products (e.g., cyclohexene) were observed during the reaction. No hydrogenation of benzene was observed when the reaction was carried out using rhodium-free nanocrystalline HAp, indicating that the support is catalytically inactive for this reaction.
After the complete hydrogenation of benzene was achieved, the catalyst was isolated by simple filtration and characterized using several analytical techniques. Powder X-ray diffraction (PXRD) patterns of Rh(0)@HAp are identical to that of the host HAp framework (Figure SI-1), thus indicating that no new phases appear after the formation of Rh(0)@HAp and that, after reaction, no observable alterations in the framework lattice or a loss in crystallinity is detected.13,14 Transmission electron microscopy (TEM) images reveal the presence of Rh(0) nanoparticles on the HAp matrix in the range of 1.0−4.0 nm with a mean diameter of 2.7 ± 1.1 nm, which corresponds to Rh(0)nanoclusters (Figure 2).15 High-resolution TEM images of the supported Rh(0) nanoparticles show an interplanar spacing of ∼0.23 nm, conforming to the {111} plane of cubic Rh nanoparticles (Figure 2e).16 The high-angle annular dark-field scanning TEM (HAADF-STEM) images (Figure 3a–d) of Rh(0)@HAp mapped for Ca, P, and Rh indicate that the Rh(0) nanoparticles are well dispersed throughout the hydroxyapatite matrix. The oxidation state of rhodium and the surface composition of Rh(0)@HAp were investigated by X-ray photoelectron spectroscopy (XPS). The survey-scan XPS spectrum of Rh(0)@HAp (Figure SI-2a) suggests that rhodium is the only element detected in addition to the HAp framework elements (Ca, P, and O). The high-resolution Rh 3d XPS spectrum of Rh(0)@HAp (Figure SI-2b) gives two prominent bands at 308 and 312.4 eV that are readily assigned to Rh(0) 3d₅/₂ and Rh(0) 3d₃/₂, respectively, thus indicating the complete reduction of Rh(III) on HAp to Rh(0) by H₂ during the hydrogenation of benzene.17 Compared to the value of metallic rhodium 3d peaks, the slight shift (ΔEₚ = +0.6 eV) observed in Rh(0)@HAp toward a higher energy value can be attributed to the peculiar electronic properties of the HAp matrix and surface oxidation during the XPS sampling procedure.18 Additionally, the interaction of rhodium(0) nanoparticles with the framework oxygen of the HAp matrix induces a positive charge on the rhodium surface, which would increase the Rh 3d binding energy of rhodium(0) nanoparticles.18 The N₂ adsorption isotherms of both HAp and Rh(0)@HAp (Figure SI-3) show type III isotherms reflecting the absence of micropores (<2 nm).19 Surface areas of 19 and 35 m² g⁻¹ were calculated for Rh(0)@HAp and the parent HAp, respectively. The reduction in surface area for Rh(0)@HAp is associated with the presence of rhodium nanoparticles on the HAp surface.

Prior to kinetic measurements, the absence of external and internal mass transfer limitations was confirmed by performing the Madon–Boudart test and by obtaining a Weisz–Prater (WP) parameter value of ≪0.3, following the methodology suggested by Mukherjee and Vannice.20 A range of initial turnover frequency (TOF) values was calculated using the number of active surface sites determined through multiple techniques. CO chemisorption experiments on Rh(0)@HAp indicate CO/Rh uptake values of 1.78 ± 0.1. Indeed, surface species in Rh nanoparticles can form both mono- and dicarbonyl complexes with CO, thus explaining CO/Rh uptake values that are greater than 1.23 Although CO chemisorption values cannot be used to quantify the exact number of surface Rh species, these data suggest that Rh is highly dispersed on the support. Thus, a sensible assumption of a 100% metal dispersion translates into a remarkable lower bound initial TOF for benzene hydrogenation of 1770 h⁻¹. However, different surface sites (i.e., high-coordination-number terrace atoms and low-coordination-number edge and corner atoms) may feature drastically different reactivity profiles. Carbon disulfide (CS₂) poisoning experiments were performed to identify the number of active surface species. Specifically, a series of independent benzene hydrogenation experiments with varying [CS₂]/[Rh] ratios were performed to determine the amount of CS₂ required to deactivate the Rh(0)@HAp catalyst.

It was found that 0.032 mol equiv of CS₂ is sufficient to poison
all of the available active Rh sites in the catalyst and completely stop the reaction (Figure SI-4). Coupling this result with a 1:5 CS$_2$/Rh stoichiometry, as suggested by Hornstein and co-workers based on single-crystal data, results in 16% of the total number of Rh atoms being catalytically available and active for the hydrogenation of benzene. Using this value, an exceptional higher bound initial TOF of 11,000 h$^{-1}$ is obtained.

Although the precise CS$_2$/Rh stoichiometry remains under debate, the poisoning experiments indicate that only a fraction of the surface sites are responsible for catalyzing the hydrogenation reaction. Adopting the model of full-shell nanoparticles for cubic-octahedral Rh crystallites allows for the calculation of total ($N_T$) and surface ($N_S$) (high-coordination [$N_{HS}$] and low-coordination [$N_{LS}$]) atoms using eqs 1–5

$$d_{sph} = 1.105d_{at}N_T^{1/3} \tag{1}$$

$$N_T = \frac{10m^3 - 15m^2 + 11m - 3}{3} \tag{2}$$

$$N_S = 10m^2 - 20m + 12 \tag{3}$$

$$N_{HS} = 6(m - 2)^2 + 4(m - 3)(m - 2) \tag{4}$$

$$N_{LS} = 24(m - 2) + 12 \tag{5}$$

where $d_{sph}$ is the mean diameter of the Rh crystallites obtained from TEM analysis and $d_{at}$ is the atom diameter of Rh (0.268 nm). Thus, it is observed that highly reactive low-coordination-number atoms ($N_{LS}$) correspond to 16.1% of the total number of Rh atoms for 2.7 nm nanoparticles, which is in good agreement with the results obtained from CS$_2$ poisoning experiments.

Reusability experiments geared to assessing the stability of Rh(0)@HAp indicated that the catalyst is very robust and does not undergo deactivation after several hydrogenation cycles. After the complete hydrogenation of benzene in the first run, Rh(0)@HAp was isolated as a dark-brown powder by simple filtration and dried under vacuum at room temperature. The Rh(0)@HAp sample can be safely bottled and stored under ambient conditions. Furthermore, when reused, Rh(0)@HAp retains 92% of its initial catalytic activity after five cycles, achieving the complete hydrogenation of benzene in each cycle without jeopardizing the selectivity (Figure 4). This translates into a lower-bound turnover number [TON] of 11,520. The slight decrease in the initial catalytic activity can be attributed to a decrease in the number of active surface atoms due to a small increase in the size of the Rh(0) nanoparticles from 2.7 ± 1.1 nm (Rh(0)-274 nanoclusters) to 3.2 ± 0.9 nm (Rh(0)-1245 nanoclusters), as measured for Rh(0)@HAp recovered at the end of the first and third cycles, respectively. Importantly, no rhodium leaching was detected in the filtrate solution collected at the end of the reaction (analyzed by ICP-MS with a detection limit of 30 ppb for Rh). A control experiment showed that the hydrogenation of benzene did not continue after Rh(0)@HAp was removed shortly after the reaction had started, confirming no leaching of active catalyst to the solution.

The in situ generation of Rh(0)@HAp and its subsequent use as a hydrogenation catalyst was also demonstrated using various other substrates, including cyclohexene, toluene, m-xylene, o-xylene, and p-xylene (Table 1 and Figures SI-7–SI-11). In all cases, Rh(0)@HAp provided high hydrogenation rates and TOF values with quantitative yields of the corresponding saturated product under solvent-free conditions at 298 K and 3 bars of initial H$_2$ pressure. As expected, cyclohexene hydrogenation rates are faster when compared to benzene hydrogenation rates because the resonance stabilization energy is typically absent in olefins. Also, slower hydrogenation rates are observed with increasing methyl substituents in the aromatic ring, in agreement with previous reports showing that methyl substituents on the benzene ring act as electron-contributing groups that stabilize π complexation, thereby increasing the energy barrier toward hydrogenation. The observed trend in the hydrogenation rate of xlyenes ($p$-xylene $> m$-xylene $> o$-xylene) can be explained by the steric hindrance resulting from the position of the methyl groups that hinder the accessibility of one of the C=C bonds.
to the active rhodium sites, which is most prominent in the case of o-xylene.27

■ CONCLUSIONS

Novel Rh(0)@HAp has been reproducibly prepared from the in situ reduction of Rh3+@HAp in pure olefinic or aromatic substrates. Rh(0)@HAp is a highly active and selective catalyst for the hydrogenation of olefins and arenes under solvent-free, mild reaction conditions (298 K and 3 bars of initial H2 pressure). This material demonstrates exceptional catalytic activity (TOF = 11 000 h⁻¹) and reusability (92% of inherent activity after the fifth reuse) for the hydrogenation of benzene. Several advanced analytical tools have been used to characterize the material, highlighting the role of the Rh(0) nanoparticles as the active species. Understanding the existing synergistic effects between the metal and HAp support remains an active area of research in our group. High catalytic activity, reusability, and simple preparation and isolation procedures make Rh(0)@HAp a very attractive catalyst for performing selective aromatic hydrogenation reactions.

■ ASSOCIATED CONTENT

$Supporting Information

Includes the experimental procedure, results of the characterization of Rh(0)@Hap, and the hydrogenation of different substrates. This material is available free of charge via the Internet at http://pubs.acs.org.

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Author Contributions

M.Z. designed and performed the experiments. M.Z. and Y.R.-L. discussed the results and wrote the article. Y.Z. performed TEM, HRTEM, STEM and HAADF-STEM analyses.

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■ REFERENCES

(4) The decrease in the intensity of Bragg peaks (or the absence of some of them) in Rh(0)@HAp can be explained by the changes in the charge distribution and electrostatic fields as result of (i) the existence of Rh(0) nanoparticles on the surface and interaction of their electrophilic surface with framework oxygen atoms, (ii) the occupancy of Ca2+ ions' sites by H2 ions (formed the reduction of Rh3+ by dihydrogen; Rh3+ + 3/2 H2 → Rh0 + 3H+) at the end of the formation of Rh(0)@HAp. Nevertheless, most peaks are not changed and do not change after reaction. Additionally, the low-resolution-TEM images of HAp taken before and after reduction of Rh3+@HAp show the same crystalline morphology (no agglomeration that can lead to amorphous phase).
(5) Using the equation N = NpV/101.1, where Np = 6.022 × 1023, p = 12.4 g cm⁻³ and V = (4/3)πr³, the number of metal atoms in spherical 2.7 and 3.2 nm rhodium(0) nanoclusters were estimated to be 748 and 1245, respectively.
(6) The TOF and TON reported herein are defined as TOF = (mol of H consumed)/(total mol of Rh)/time; TON = (TOF) × (time). That is, the TOF and TON values reported are not corrected for the amount of surface metal atoms or the number of true active sites.