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Pt/Al₂O₃ coated with N-doped carbon as a highly selective and stable catalyst for catalytic hydrogenation of p-chloronitrobenzene to p-chloroaniline†

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Selective catalytic hydrogenation of p-chloronitrobenzene on Pt-based catalysts is a green and high-efficient way for p-chloroaniline production. However, supported monometallic Pt catalysts often exhibit undesirable p-chloroaniline selectivity. We herein reported supported Pt catalysts with N-doped carbon (NC) as an overcoating (Pt/Al₂O₃@NC) to overcome the disadvantage. Three Pt/Al₂O₃@NC catalysts with different NC coating amounts were prepared by in situ carbonization of an ionic liquid. For comparison, Al₂O₃ coated by NC and Pt/Al₂O₃ coated by SiO₂ were also prepared. A combination characterization confirmed that the NC overcoating was successfully formed on Pt/Al₂O₃ surface and Pt particles were completely coated by NC layers when ion liquid amount increased to 25 μ l per g catalyst. Due to the intimate contact of NC layers and Pt particles Pt-NC heterojunctions were effectively formed on the catalyst surface. For the catalytic hydrogenation of p-chloronitrobenzene, Pt/Al₂O₃@NC with 25 μ l ionic liquid as the NC precursor exhibited 100% selectivity to p-chloroaniline at 100% conversion of p-chloronitrobenzene. A lower ionic liquid amount led to decreased selectivity to p-chloroaniline. Furthermore, no deactivation was observed on Pt/Al₂O₃@NC during 5 catalytic cycles. The findings in the study demonstrate that coating noble metal catalysts by N-doped carbon is a promising method to enhance the selectivity and stability for catalytic hydrogenation of p-chloronitrobenzene.

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1. Introduction

As important chemical raw materials, chloroanilines (CANs) are widely used as intermediates in pharmaceuticals, herbicides, pigments and dyes. ¹⁻³ Conventionally, CANs could be produced *via* reduction of corresponding chloronitrobenzenes (CNBs) using stoichiometric reductants such as potassium borohydride and hydrazine hydrate. ²⁻⁴ However, the use of stoichiometric reductants would bring large amounts of harmful waste and severe pollution to the environment. Alternatively, selective catalytic hydrogenation reduction is an eco-friendly and cost-effective method for the conversion of CNBs to CANs.

For catalytic hydrogenation reduction, noble metals such as Pd and Pt are highly active because of their strong abilities for H₂ activation under mild conditions.^{5,6} However, supported noble metal catalysts exhibited low selectivity because dechlorinated byproducts were inevitably formed due to their strong hydrodechlorination capabilities.^{5,7} For example, supported Pt

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catalysts were considered as the most selective ones among supported noble metals for the catalytic conversion of CNBs to CANs, ^{3,8,9} however, Pt/C catalyst only displayed 75.2% selectivity to *p*-CAN at 100% conversion of *p*-CNB. ¹⁰ Additionally, catalyst deactivation often occurs as a result of contamination of metal surface or/and metal leaching during the reaction processes. ^{11,12} For example, Mahata *et al.* ¹¹ found that Pt/CNT deactivated drastically in the catalytic hydrogenation of *o*-CNB during the second and subsequent reaction runs. Hence, exploring effective strategies to improve the catalytic selectivity and stability of supported noble metal catalysts for hydrogenation of CNBs to corresponding CANs is of significant importance.

Many attempts such as addition of a promoting metal or decorating metal particles with organic modifiers have been made to improve the catalytic selectivity for *p*-CNB hydrogenation.^{13–17} For example, Zhu *et al.*¹⁸ prepared a core–shell structural bimetallic Pt@Cu/TiO₂ catalyst with monolayer Cu as the shell for catalytic hydrogenation of *p*-CNB and obtained 100% selectivity to *p*-CAN. Very recently, coating supported catalysts have been proved to be an effective approach to enhance catalytic selectivity. Chen *et al.*¹⁴ prepared a Pt-nanowire catalyst chelated by ethylenediamine on catalyst surface, which exhibited prominent selectivity for the catalytic production of *N*-

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hydroxylanilines from nitrobenzene. Leng et al. 19 reported that controlled and chemoselective hydrogenation of nitrobenzene was achieved on a Ru@C₆₀ catalyst. Notably, coating supported catalysts was also reported as a prominent strategy to enhance catalytic stability through protecting metal particles from leaching and poisoning. For example, Fu et al. 20 illustrated that a Ni/Al_2O_3 catalyst coated with carbon nitride layer provided an exceptional stability for the catalytic hydrogenation of nitrobenzene. Li et al.21 developed a Pt catalyst with Pt particles embedded in N-doped carbon rods and observed a superior catalytic stability in the liquid-phase catalytic hydrogenation of bromate. Recent years, N-doped carbon (NC) materials have attracted tremendous attentions because of their unique and versatile properties. For example, N-doped carbon could transfer electrons to metal particles or vice versa via metal-NC heterojunctions and thus serve as catalytically active sites for the adsorption and activation of H₂ and reactants.²⁰⁻²² Inspired by recent advances, it is reasonable to speculate that coating supported noble metal catalysts with N-doped carbon may probably improve the catalytic selectivity and stability for hydrogenation of CNBs. However, few studies have been conducted for the selective hydrogenation of p-CNB on coated noble metal cata-

The main objective of the present study is to develop highly selective and stable supported Pt catalysts for the catalytic hydrogenation of CNBs to CANs. We prepared three Pt/Al $_2$ O $_3$ catalysts coated by NC layer with different NC amounts by *in situ* carbonization of ion liquid (1-ethyl-3-methylimidazolium dicyanamide, EMIm-dca). For comparison, Al $_2$ O $_3$ coated with NC and Pt/Al $_2$ O $_3$ coated with SiO $_2$ were also prepared. The structural properties of the synthesized catalysts were characterized by a series of techniques and the catalytic performances of the catalysts for hydrogenation of p-CNB were examined.

2. Experimental

2.1 Catalyst preparation

lysts thus far.

Chemical and materials. Ion liquid (EMIm-dca) and hydrogen hexachloroplatinate($_{\rm IV}$) hexahydrate ($_{\rm H_2}$ PtCl $_{\rm 6}$ ·6H $_{\rm 2}$ O) were purchased from Aldrich. Aluminum isopropoxide, oleylamine, sodium silicate, ammonium hydroxide (NH $_{\rm 3}$ ·H $_{\rm 2}$ O), hydrogenchloride (HCl), nitric acid (HNO $_{\rm 3}$), p-CNB, and p-CAN were obtained from Sinopharm Chemical Reagent Co., Ltd., China. All chemicals were used without further purification.

Preparation of aluminum sol. A desired amount of aluminum isopropoxide powder was added slowly into 150 ml distilled water with a molar ratio of water to aluminum isopropoxide of 100:1 in a three-necked round flask under vigorous stirring at 80 °C for controlled hydrolysis of aluminum isopropoxide. After refluxing at 80 °C for 1 h, the mixture was further evaporated at 90 °C for 1 h to remove formed alcohol. The pH of the mixture was then adjusted to 3.0 using 10% HNO₃ solution. Aluminum sol was obtained after the mixture further refluxed at 80 °C for 24 h.

Preparation of Al_2O_3. Al $_2$ O $_3$ was prepared using method reported by Fu *et al.*²⁰ Briefly, 180 g of alumina sol, 14 g of oleylamine and 10 ml of 25% NH $_3$ ·H $_2$ O was dispersed in 150 ml of

distilled water at 80 $^{\circ}$ C in a water bath. After stirring for 2 h at 80 $^{\circ}$ C, the mixture was transferred into the autoclave, sealed and kept at 180 $^{\circ}$ C for 72 h in an oven. The resulting mixture was filtered and washed using ethanol for several times and then oven-dried at 100 $^{\circ}$ C. Finally, the obtained powder was calcined at 550 $^{\circ}$ C in a muffle furnace for 4 h.

Preparation of Pt/Al_2O_3. Pt/Al $_2$ O $_3$ was prepared using a conventional impregnation method with H $_2$ PtCl $_6$ as metal precursor. Briefly, 1 g of grinded and sieved Al $_2$ O $_3$ was impregnated with 2.7 ml of 1.9 \times 10 $^{-5}$ mol ml $^{-1}$ H $_2$ PtCl $_6$ solution. After stirring for 2 h, the mixture was evaporated under 80 °C in a water bath. The sample was oven-dried at 100 °C overnight and calcined in a muffle furnace at 300 °C for 4 h. The nominal content of Pt was 1 wt%.

Preparation of Pt/Al₂O₃(a)NC. Pt/Al₂O₃(a)NC was prepared by a reported method with ion liquid (EMIm-dca) as NC precursor.23 Briefly, 1 g Pt/Al₂O₃ and a desired amount of ion liquid (10, 15 and 25 µl of EMIm-dca, respectively) were added in a 20 ml of beaker containing 5 ml of distilled water. After mixed well by ultrasonic treatment, the slurry was dried at 70 °C in a porcelain boat and calcined in a tubular furnace with a heating rate of 5 °C min⁻¹ up to 400 °C and maintained at this temperature for 1 h under N2 atmosphere. For comparison, Al₂O₃@NC without Pt loading was also prepared. The resulting catalysts were labeled as Pt/Al₂O₃@NC-X or Al₂O₃@NC-X, where X refers to the amount of ion liquid added. The prepared Pt/ Al₂O₃@NC-25 was further calcined at 500 °C in air for 2 h to remove the NC overcoating to clarify the role of NC. This catalyst was labelled as Pt/Al₂O₃@NC-25-calcined. All the catalysts were pre-reduced in H₂ atmosphere at 300 °C for 2 h in prior to the characterization and evaluation of catalytic activity.

Preparation of Pt/Al₂O₃@SiO₂. Pt/Al₂O₃@SiO₂ was prepared using method reported by Li *et al.*¹² Briefly, 1 g of Pt/Al₂O₃ was dispersed in 200 ml of distilled water, which was stabilized at 80 °C under a 30 ml min⁻¹ N₂ flow. Under vigorous stirring, 20 ml of sodium silicate solution (1.0 mol l⁻¹) was added to the suspension dropwise, and then the pH of the mixture was adjusted to 6.0 using a HCl solution (1 mol l⁻¹). The mixture was further stirred at 80 °C for 30 h. The catalyst was obtained after washed with distilled water for several times and dried at 65 °C under vacuum for 10 h.

2.2 Catalyst characterization

Pt contents in the samples were precisely determined by an inductively coupled plasma-atomic emission spectrometer (ICP-AES) (J-A1100, Jarrell-Ash, USA). The XRD patterns of the samples were collected on a Rigaku D/max-RA powder diffraction-meter using a Cu K α radiation (Rigaku, Tokyo, Japan). The carbon and nitrogen contents in the catalysts were determined using an elementary analyzer CHN-O-rapid apparatus (Heraeus, Germany). The Brunauer-Emmett-Teller (BET) surface areas of the catalysts were measured using a Micromeritics ASAP 2200 instrument (Micromeritics Instrument Co., Norcross, GA). High-angle annular dark field-scanning transmission electron microscope (HAADF-STEM) and elemental mapping images were acquired using an ARM-200F

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transmission electron microscope (JEOL Co., Japan.) equipped with an X-MAX energy dispersive spectrometer (Oxford, UK). Xray photoelectron spectroscopy (XPS) spectra were recorded on a PHI5000 VersaProbe with a monochromatized Al Ka X-ray source ($h\nu = 1486.6$ eV) (ULVAC-PHI, Japan). The binding energy was calibrated by measuring the C 1s peak at 284.6 eV.

Temperature programmed oxidation (TPO) was conducted on an AMI-300 setup (Altamira Instruments, USA) with a gas analysis system (Thermo Star, Germany). Typically, 50 mg of catalyst was loaded in a U-shaped quartz tube, which was pretreated at 120 °C for 1 h using a 25 ml min⁻¹ of He flow in a heating furnace. After cooling down, the catalyst was in situ oxidized with a 5% O₂/He flow (25 ml min⁻¹) from ambient temperature to 600 °C at a rate of 5 °C min⁻¹. The evolved gases were monitored with intensities of m/z 17 (NH₃), 28 (CO), 30 (NO), 44 (CO₂) and 46 (NO₂), respectively.

2.3 Catalytic hydrogenation of p-chloronitrobenzene

The catalytic hydrogenation of p-CNB over the catalysts was conducted in a high pressure stainless steel autoclave with a controlling system. Briefly, 20 mg catalyst, 100 mg p-CNB and 20 ml ethanol was added to the autoclave. Before hydrogenation reaction, the autoclave was flushed with N2 for 5 times to replace air in the reactor under ambient condition. The reaction was conducted under 1 MPa pressure of H₂ at 80 °C with stirring rate fixing at 1000 rpm. Samples were taken periodically and detected by a 6980 gas chromatograph equipped with a FID detector and a DB-5 capillary column (Agilent Technologies, US), and *n*-hexane was used as the inner standard substance. For catalyst reuse, the used catalyst was prepared under the same reaction conditions but without sampling. After reaction for 180 min, the catalyst was collected by filtration, rinsing with ethanol and distilled water and drying at 60 °C for 12 h. The catalyst was then subjected to the subsequent reaction test.

The conversion of p-CNB and selectivity to p-CAN and aniline were calculated following equations below:

Conversion =
$$\frac{m_{(p\text{-CNB})0} - m_{(p\text{-CNB})t}}{m_{(p\text{-CNB})0}} \times 100\%$$
 (1)

Selectivity_(p-CAN) =
$$\frac{m_{(p-CAN)t}}{m_{(p-CAN)t} + m_{(aniline)t}} \times 100\%$$
 (2)

Selectivity_(aniline) =
$$\frac{m_{(\text{aniline})t}}{m_{(p\text{-CAN})t} + m_{(\text{aniline})t}} \times 100\%$$
 (3)

where m refers to the molar amount.

3. Results and discussion

Catalyst characterization

Table 1 shows Pt content of the catalysts determined by ICP-AES. For Pt/Al₂O₃, Pt content was determined to be 0.96 wt%. As for Pt/Al₂O₃@NC, Pt content were in the range of 0.71-0.78 wt%, which were lower than that of Pt/Al₂O₃ due to successful loading of NC overcoating.

The compositions of C and N elements in the catalysts were determined by elemental analysis and the results are

| - | - | , | | | | | | | | |
|--|-------------------------------|--|------------------------------|---|------------------|---|----------------------------------|--|--|--------------|
| Catalysts | Pt content ^a (wt%) | Pt content ^a C content ^b (wt%) | N content ^b (wt%) | Surface concentration of N ^c (at%) | ${ m N_1/N_2}^c$ | BET surface area ^{d} N ₁ /N ₂ ^{c} (m ² g ⁻¹) | d ₁ ^e (nm) | CO adsorption amount d_1^e (nm) $(\mu \text{mol } g^{-1})$ | Coverage of Pt d_2^f surface (%) (nn | d_2^f (nm) |
| Pt/Al ₂ O ₃ | 96.0 | BDL | BDL | BDL | BDL | 103 | 4.0 | 12.0 | I | 4.3 |
| Pt/Al ₂ O ₃ @NC-10 | 0.78 | 0.33 | 0.17 | 0.49 | BDL | 100 | 3.9 | 3.7 | %69 | |
| Pt/Al ₂ O ₃ @NC-15 | 0.71 | 0.63 | 0.30 | 0.65 | 0.92 | 101 | 4.0 | 0.5 | %96 | I |
| Pt/Al ₂ O ₃ @NC-25 | 0.78 | 0.76 | 0.39 | 0.79 | 0.71 | 66 | 4.1 | 0 | 100% | |
| Al_2O_3 (a) NC-25 | BDL^g | 0.58 | 0.24 | 0.61 | 0.47 | 110 | I | BDL | 1 | |

pyridinic N amount to pyrrole N amount on catalyst surface. ^a Calculated from BET. Determined by ICP. ^b Determined by elemental analysis. ^c Determined by XPS, N₁/N₂ refers to the ratio of j Open Access Article. Published on 07 April 2020. Downloaded on 1/5/2026 7:19:49 AM.

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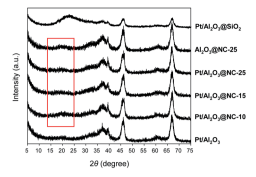


Fig. 1 XRD patterns of the catalysts.

summarized in Table 1. For Pt/Al₂O₃@NC-10, Pt/Al₂O₃@NC-15 and Pt/Al₂O₃@NC-25, the C contents were 0.33%, 0.63% and 0.76% respectively, exhibiting increased C content with the increase of ion liquid loading amount from 10 µl to 25 µl. Consistently, N contents in Pt/Al2O3@NC catalysts increased from 0.17% to 0.39% with ion liquid loading. For Al₂O₃@NC-25, the C and N content were determined to be 0.58% and 0.24% respectively, lower than those of Pt/Al₂O₃@NC-25, probably because the presence of Pt particles on Al₂O₃ surface stabilized the NC matrix formed by pyrolysis of ion liquid via evoking interactions with NC matrix. The molar N/C ratios in the catalysts were in the range of 0.35-0.44 which was lower than 0.63 of ion liquid, this can be ascribed to more prominent loss of unstable carbonization N species upon high temperatures.24,25

Fig. 1 shows the XRD patterns of the catalysts in the range of 20-80°. Diffraction peaks centered at 37.3°, 39.5°, 45.7°, and 67.1° indexed to the γ-Al₂O₃ phase, were clearly observed on all the catalysts. No diffraction peaks characteristic of Pt were found in the XRD patterns probably due to low metal contents of the catalysts.²⁶ Notably, in the range of 15-25°, a weak and broad peak centered at 19.5° were observed on Pt/Al₂O₃@NC catalysts. The very weak peak was ascribed to amorphous Ndoped carbon, which was lower than that of amorphous carbon (23.5°) due to the increase of inter-layer distance of carbon upon N doping.27 As for the XRD pattern of Pt/Al2O3@-SiO₂, the broad peak centered at 23.1° was assigned to amorphous silica. In comparison with Pt/Al₂O₃@NC, Pt/Al₂O₃@SiO₂ had markedly weakened intensities of peaks of γ-Al₂O₃ phase, likely resulting from the thicker SiO₂ overcoating than NC overcoating.

Fig. S1, ESI† shows the HAADF-STEM images of the catalysts. Al₂O₃ had uniform rhombus morphology with diameter in the range of 30-60 nm. The TEM images showed that Pt particles were spherical and were well dispersed on Al₂O₃ surface. The average Pt particle diameter was measured based on the surface area weighted diameter:28,29

$$\overline{d_{s}} = \sum n_{i} d_{i}^{3} / \sum n_{i} d_{i}^{2} \tag{4}$$

where n_i is the number of selected Pt particles having diameter of d_i and the selected total particle number $(\sum n_i)$ exceeds 100.

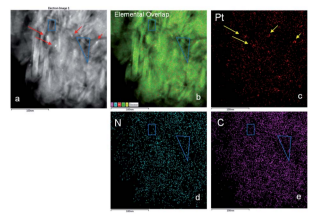


Fig. 2 (a) HAADF-STEM image of Pt/Al₂O₃@NC-25 and the corresponding EDS elemental mapping of (b) the overlapping of Pt, C, and N element, (c) Pt element (red), (d) N element (blue), and (e) C element (purple).

Results are listed in Table 1. The average sizes of Pt particles of Pt/Al_2O_3 , Pt/Al_2O_3 @NC-10, Pt/Al_2O_3 @NC-15 and Pt/Al₂O₃@NC-25 were 4.0, 3.9, 4.0 and 4.1 nm, respectively, indicating that NC coating did not affect the structure of Pt/Al₂O₃. However, NC overcoating layer was not clearly visible in TEM images, probably because of the very low NC contents (as indicated by elemental analysis results). To verify the presence of NC overcoating on the catalysts, EDS mapping analysis was conducted on Pt/Al₂O₃@NC-25 and the images are presented in Fig. 2. In Fig. 2a light particles with diameter around 5-7 nm could be clearly observed as marked by red arrows. Consistently, Pt signal was very dense in the same area in the mapping image in Fig. 2c, reflecting that the lightness variation was due to the presence of Pt particles on Al₂O₃ support. The EDS mapping of elemental C and N in Pt/Al₂O₃@NC-25 (see Fig. 2d and e) showed that elemental C and N were evenly distributed on Al2O3 surface. The clear spatial correlation among elemental Al, C and N was indicative of a close contact of NC with Al2O3 support, verifying the presence of NC overcoating on Al2O3 surface.

To further investigate the composition of Pt/Al₂O₃@NC-25, temperature-programed oxidation of Pt/Al2O3@NC-25 was

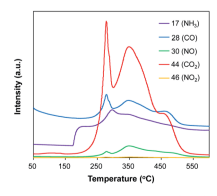


Fig. 3 Temperature programmed oxidation profile of Pt/Al₂O₃@NC-25.

conducted, and the dependence of gas release on oxidation temperature are shown in Fig. 3. In Fig. 3, CO and CO₂ exhibited very similar evolving profiles, in which a strong and broad peak with two shoulders presented in the range of 200–550 °C. In parallel, marked release of NH₃ and NO was also observed in the range of 180–530 °C. The presence of CO₂, CO, NH₃ and NO₂ could be ascribed to the oxidation of C and N containing species, confirming the presence of NC overcoating on the catalyst surface.³⁰ Notably, the oxidation temperature of N species was slightly lower than that of C species, reflecting that N species were more prone to lose than C species upon high temperature, in line with elemental analysis results.

The effective decoration of NC overcoating on Pt particle could be further verified by CO chemisorption (see Table 1). The CO adsorption amount on Pt/Al₂O₃ was determined to be 12.0 μmol g⁻¹. Accordingly, the average diameter of Pt particles of Pt/ Al₂O₃ could be calculated by CO chemisorption amount with a chemisorption stoichiometry of CO/Pt = 1. The diameter was calculated to be 4.3 nm, in good agreement with that measured by TEM. After NC coating, the CO adsorption amount markedly decreased. For example, CO adsorption amount of Pt/ Al_2O_3 @NC-10 and Pt/Al_2O_3 @NC-15 was 3.7 µmol g^{-1} and 0.5 μmol g⁻¹ respectively, much lower than that of Pt/Al₂O₃. Because CO could chemically adsorb only on exposed Pt sites of the catalyst, the decreased chemisorption amount of CO clearly reflected lowered exposure of Pt sites of the catalysts as a result of NC overcoating.31 Further increasing NC coating amount even led to the absence of CO adsorption on Pt/Al₂O₃@NC-25, indicating that Pt sites were completely blocked by NC overcoating in the catalyst. For Pt/Al₂O₃@NC-25 calcined at 500 °C in air, the CO chemisorption amount increased to 7.0 μ mol g⁻¹

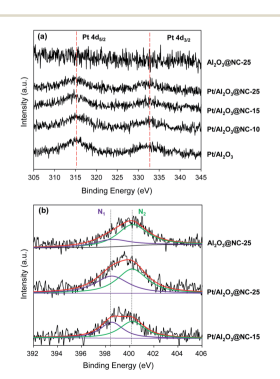


Fig. 4 XPS spectra of catalysts in the regions of (a) Pt 4d and (b) N 1s.

(see Table 3). Because calcination led to the oxidation of NC matrix of the catalyst and thus resulted in the exposure of blocked Pt sites for CO chemisorption. Notably, the CO chemisorption amount of Pt/Al_2O_3 @NC-25-calcined was lower than that of Pt/Al_2O_3 , probably due to incomplete oxidation of NC overcoating upon calcination.

Fig. 4 shows the XPS spectra of the catalysts. Because Pt 4f region was overlapped by the strong Al 3p peak, the surface Pt compositions of the catalysts were analyzed by XPS spectra in Pt 4d region. In Fig. 4a, two peaks around 315.2 and 332.7 eV were found for Pt/Al₂O₃, characteristic of Pt 4d_{5/2} and 4d_{3/2}, respectively (see Fig. 4a).32,33 Upon NC coating, slightly lowered binding energies of Pt 4d were observed. For example, the binding energies of Pt $4d_{5/2}$ was 315.0, 314.8 and 314.7 eV for Pt/ Al₂O₃@NC-10, Pt/Al₂O₃@NC-15 and Pt/Al₂O₃@NC-25 respectively. The gradual red shift of the binding energy of Pt 4d_{5/2} with NC coating amount in the catalysts may be ascribed to the electron transfer from electronic-enriched N sites to Pt due to their strong interactions.34,35 Fig. 4b shows XPS spectra of the catalysts in the N 1s region. Deconvolution of the N 1s peak gave rise to two peaks located at 398.6 and 400.2 eV, characteristic of pyridinic N and pyrrole N respectively, further confirming the successful formation of NC overcoating layers. The ratio of pyridinic N to pyrrole N of the catalysts were calculated and the results are summarized in Table 1. Notably, the ratio of pyridinic N (N_1) to pyrrole N (N_2) markedly differed with samples. For Al₂O₃@NC-25, the ratio of N₁/N₂ was 0.47, which was much lower than that of Pt/Al₂O₃@NC-25 (0.71). In comparison with Pt/Al₂O₃@NC-25, Pt/Al₂O₃@NC-15 with

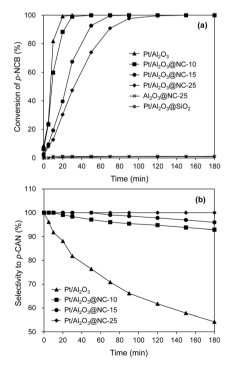


Fig. 5 Catalytic hydrogenation of p-CNB on the catalysts. Reaction conditions: catalyst dosage: 20 mg, alcohol: 20 ml, p-NCB amount: 100 mg, reaction temperature: 80 °C, H₂ pressure: 1 MPa.

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a lower NC content has a higher N_1/N_2 ratio of 0.92. Notably, binding energy in N 1s region characteristic of covalently bound N–Pt species was found to be 398.6 eV, which was identical to the binding energy of pyridinic N.³6 Hence, the much higher N_1/N_2 ratio in Pt/Al $_2O_3$ @NC-25 than that of Al $_2O_3$ @NC-25 could be attributed to the formation of additional N–Pt species when ion liquid was pyrolyzed on catalyst surface. The results clearly confirmed strong interactions between Pt and N atoms in NC matrix as also indicated by elemental analysis results. Consistently, the higher N_1/N_2 ratio of Pt/Al $_2O_3$ @NC-15 can be explained in terms of the formation of more Pt–N species as a result of the higher Pt : N ratio in Pt/Al $_2O_3$ @NC-15 than that of

Pt/Al₂O₃@NC-25 assuming identical Pt sites in Pt/Al₂O₃@NC-15

3.2 Catalytic hydrogenation of p-CNB

to those in Pt/Al₂O₃@NC-25.

The catalytic performances of the catalysts for hydrogenation of p-CNB are presented in Fig. 5 and the reaction constants are compared in Table 2. Aniline and p-CAN were identified as products in all the reactions. Control experiment showed that Al₂O₃@NC-25 exhibited very low activity for catalytic hydrogenation of p-CNB with p-CNB conversion of 0.6% within 180 min of reaction time, reflecting the very weak capability of NC overcoating and Al2O3 for H2 activation under reaction conditions. In contrast, Pt/Al₂O₃ exhibited a much higher catalytic activity with 100% of p-CNB conversion within 30 min indicating the high activity of Pt particles for H2 activation and p-CNB reduction. Upon NC coating, the conversion of p-CNB over Pt/Al₂O₃@NC gradually decreased with NC coating amount. For example, p-CNB was completely converted at 50, 90, and 150 min on Pt/Al₂O₃@NC-10, Pt/Al₂O₃@NC-15 and Pt/ Al₂O₃@NC-25 respectively (see Table 2). The deteriorated catalytic activity of Pt/Al₂O₃@NC was likely due to blocking Pt sites by NC overcoating from p-CNB as evidenced by CO chemisorption results. Accordingly, decreased catalytic activity as a result of coating catalyst surface had been reported previously. 37,38 Notably, effective hydrogenation of p-CNB was found on Pt/Al₂O₃@NC-25 although Pt sites were completely blocked by NC matrix (see CO chemisorption results). This results suggested that H2 was capable of being activated and p-CNB could

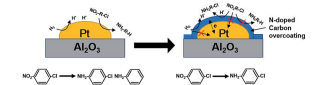
 Table 2
 Catalytic performances of the catalysts^a

| Catalysts | Reaction time (min) | Conversion (%) | p-CAN selectivity (%) | Aniline selectivity (%) |
|---|---------------------------|----------------|-----------------------|-------------------------|
| Pt/Al ₂ O ₃ | 30 | 100.0 | 81.8 | 18.2 |
| Pt/Al ₂ O ₃ @NC-10 | 50 | 100.0 | 97.1 | 2.9 |
| Pt/Al ₂ O ₃ @NC-15 | 90 | 100.0 | 98.6 | 1.4 |
| Pt/Al ₂ O ₃ @NC-25 | 150 | 100.0 | 100.0 | 0.0 |
| Al ₂ O ₃ @NC-25 | 180 | 0.6 | 100.0 | 0.0 |
| Pt/Al ₂ O ₃ @SiO ₂ | 180 | 1.1 | 100.0 | 0.0 |

 $[^]a$ Reaction conditions: catalyst dosage: 20 mg, alcohol: 20 ml, p-nitrochlorobenzene: 100 mg, reaction temperature: 80 °C, reaction time: 60 min; $\rm H_2$ pressure: 1 MPa.

be reduced on NC matrix on catalyst surface. Considering the negligible catalytic activity of Al₂O₃(a)NC-25, a synergistic effect between Pt and NC overcoating could be deduced. Previous reports proved that supported metal catalysts with carbonaceous materials as the overcoating could catalyze hydrogenation reactions. 20,21,37 It was proposed that intimate contact between metal and carbon matrix would cause a metal-carbon matrix heterojunction, through which H₂ could be adsorbed and activated and electrons could be transferred to reactants. Accordingly, the mechanism well explained the hydrogenation of p-CNB on Pt/Al₂O₃@NC-25. XPS results confirmed strong interactions between Pt particles and NC matrix, forming Pt-NC heterojunction. Furthermore, the catalytic performance of the catalyst with non-conductive overcoating also confirmed this mechanism. For example, Pt/Al₂O₃ coated by SiO₂ exhibited negligible catalytic activity for hydrogenation of p-CNB (1.1% p-CNB conversion in 180 min reaction time) (see Table 2). This was because that SiO₂ is an inert insulator, on which electron transfer was substantially suppressed, thus resulting in a negligible catalytic activity than that of Pt/Al₂O₃@NC-25. Similarly, Li et al.12 observed a much higher catalytic activity of Pd/CNT with N-doped carbon as overcoating than that of SiO₂ coated Pd/CNT for catalytic reduction of Cr (vi) in liquid phase.

Besides catalytic activity coating Pt/Al₂O₃ substantially impacted the selectivity in catalytic hydrogenation of the p-CNB. For example, Pt/Al₂O₃ displayed a catalytic selectivity of 81.8% to p-CAN at 100% of p-CNB conversion, suggesting that Pt sites were active for both reduction of NO₂ group and activation of C-Cl bond in p-CNB. In contrast, Pt/Al₂O₃@NC exhibited significantly improved p-CAN selectivity, and the selectivity to p-CAN was strongly dependent on NC overcoating amount. Specifically, increasing ion liquid amount to 25 µl per g catalyst resulted in p-CAN selectivity increasing monotonically from 81.8% to 100% at 100% conversion of p-CNB, indicating effective inhibition of hydrodechlorination owing to NC coated on the surface of Pt particles. Notably, because Cl group in p-CNB would increase the electron density of NO₂ group, the reduction of NO2 group in p-CNB was favored by the presence of Cl group.3,39 Accordingly, p-CAN is the main production in the hydrogenation of p-CNB. After p-CNB was 100% converted, however, the catalytic selectivity to p-CAN markedly decreased on all catalysts except Pt/Al₂O₃@NC-25. For example, the selectivity to p-CAN on Pt/Al₂O₃, Pt/Al₂O₃@NC-10 and Pt/ Al₂O₃@NC-15 decreased to 54.2%, 92.8% and 95.9% after 180 min of reaction time, respectively. In contrast, Pt/ Al₂O₃@NC-25 maintained 100% selectivity to p-CAN throughout the catalytic hydrogenation process even when only p-CAN left in the reaction system, clearly indicating that the catalytic dechlorination of p-CAN was completely inhibited on Pt/ Al₂O₃@NC-25. In parallel, CO chemisorption showed that the coverage of Pt surface were 69%, 96% and 100% for Pt/ Al₂O₃@NC-10, Pt/Al₂O₃@NC-15 and Pt/Al₂O₃@NC-25 respectively (see Table 1), confirming that C-Cl bond could only be cleaved on exposed Pt sites and NC overcoating was incapable of catalytic hydrodechlorination. Accordingly, the catalytic hydrogenation of p-CNB on Pt/Al2O3 and Pt/Al2O3@NC-25 were depicted in Scheme 1.



Scheme 1 Schematic illustration of catalytic hydrogenation of p-CNB on Pt/Al₂O₃ and Pt/Al₂O₃@NC-25.

To further explore the effect of NC overcoating on the catalytic performances of the coated catalysts. Pt/Al₂O₃@NC-25 was calcined at 500 °C in air for 2 h to remove NC coating and the catalytic hydrogenation of p-CNB was conducted on the calcined catalyst. The results are presented in Fig. 6. In comparison with Pt/Al₂O₃@NC-25, Pt/Al₂O₃@NC-25-calcined exhibited a significantly enhanced catalytic activity, which was very close to that of Pt/Al₂O₃. The increased catalytic activity of Pt/Al₂O₃@NC-25calcined can be explained by the removal of NC overcoating on catalyst surface via calcination as reflected by TPO and CO chemisorption results. However, the catalytic selectivity of Pt/ Al₂O₃@NC-25-calcined was still much higher than that of Pt/ Al₂O₃, with a slightly decreased selectivity of 97.9% after 180 min reaction time in comparison with Pt/Al₂O₃@NC-25. CO chemisorption suggested that partial NC matrix remained on Pt/Al₂O₃@NC-25-calcined surface. Hence, the high selectivity of Pt/Al₂O₃@NC-25-calcined was probably related with NC matrix left on catalyst surface. On one hand, the remaining NC matrix would block partial Pt sites from catalytic hydrodechlorination. On the other hand, the interactions between Pt sites and residual NC matrix impacted the catalytic selectivity of Pt/Al₂O₃@NC-25-calcined. XPS results suggested that strong interactions and electron transfer existed between NC overcoating and Pt, causing negatively charged Pt sites. Because NH₂ groups in *p*-CAN tended to donate electron to Cl groups, which would give rise to electrostatic repulsion between Cl group and negatively charged Pt sites and thus suppressed the cleavage of C-Cl bond. ^{10,40,41} Similarly, Shi *et al.* ³⁵ reported that Pt supported on N-doped carbon nanotube exhibited high selectivities to haloanilines for catalytic hydrogenation of halonitrobenzenes because of the electron-rich chemical state of Pt particles.

The stability of catalyst is of important significance for supported noble metal catalysts. For stability test, the consecutive catalytic hydrogenation of p-CNB was examined on Pt/ Al₂O₃@NC-25 and results are shown in Fig. 7. It was very interesting to find that in the initial two cycles markedly increased activity of Pt/Al₂O₃@NC-25 from 47.3% to 100% at 30 min reaction time was observed, likely due to the presence of exposed Pt sites as a result of partial elimination of NC overcoating via washing catalyst surface by reaction solution. Accordingly, previous studies indicated that surface coke of used catalysts could be partially removed via washing by organic solvents.42 Consistently, characterization results showed that CO chemisorption amount of Pt/Al₂O₃@NC-25 was 4.3 µmol g⁻¹ after 5 catalytic cycles (see Table 3), reflecting a portion of Pt sites were again exposed. After 5 catalytic cycles, the catalytic activities remained constant and no catalyst deactivation was observed indicative of a very high catalytic stability of Pt/ Al₂O₃@NC-25. HAADF-STEM and ICP results showed that the

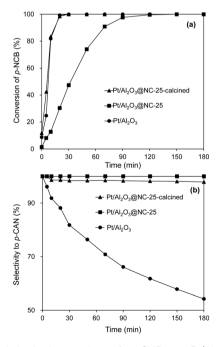


Fig. 6 Catalytic hydrogenation of p-CNB on Pt/Al $_2$ O $_3$ @NC-25-calcined in comparison with Pt/Al $_2$ O $_3$ @NC-25 and Pt/Al $_2$ O $_3$. Reaction conditions: catalyst dosage: 20 mg, alcohol: 20 ml, p-NCB amount: 100 mg, reaction temperature: 80 °C, H $_2$ pressure: 1 MPa.

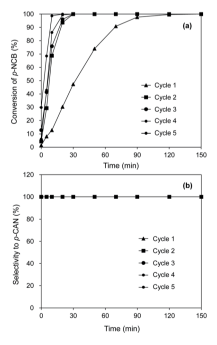


Fig. 7 The reuse of Pt/Al $_2$ O $_3$ @NC-25 for catalytic hydrogenation of p-CNB. Reaction conditions: catalyst dosage: 20 mg, alcohol: 20 ml, p-NCB amount: 100 mg, reaction temperature: 80 °C, H $_2$ pressure: 1 MPa.

Table 3 Structural properties of the catalysts

| Catalysts | CO adsorption amount ^{a} (μ mol g ^{-1}) | d^b (nm) | Pt content ^c (wt%) |
|---|---|------------|-------------------------------|
| Pt/Al ₂ O ₃ @NC-25-calcined | 7.0 | _ | _ |
| Pt/Al ₂ O ₃ @NC-25-cycled | 4.3 | 3.8 | 0.77 |

^a Calculated by CO chemisorption. ^b Calculated from HAADF-STEM results. ^c Determined by ICP.

particle size of Pt and Pt content of Pt/Al₂O₃@NC-25-cycled were very similar to those of the fresh Pt/Al₂O₃@NC-25, further confirming the high stability of Pt/Al₂O₃@NC-25 (see Table 3). Similarly, enhanced stability of supported metal catalyst νia carbon coating was also reported previously. Notably, although enhanced catalytic activity of Pt/Al₂O₃@NC-25 was observed and partial Pt sites were exposed on the catalyst surface after 2 reaction cycles, the catalytic selectivities to p-CAN were not accordingly decreased and remained 100% after 180 min reaction time in 5 reaction cycles, confirming that the electronic effect between Pt and NC overcoating played an important role in the high selectivity of the used Pt/Al₂O₃@NC-25.

4. Conclusions

In the present work, we prepared a series of Pt/Al₂O₃@NC catalysts with different ion liquid amount and investigated selective catalytic hydrogenation of p-CNB on the catalysts. Characterization results showed that NC matrix was formed as overcoating on Pt/Al₂O₃ surface, which blocked exposed Pt sites effectively. Furthermore, strong interactions existed between Pt sites and N atoms in NC matrix and their interactions led to electron-enrichment of Pt atoms. For the selective catalytic hydrogenation of p-CNB to p-CAN, Pt/Al₂O₃@NC-25 exhibited marked hydrogenation activity and 100% selectivity to p-CAN, owing to the electroconductivity of NC overcoating and the complete blockage of surface Pt sites from catalytic dechlorination. The cleavage of C-Cl bond was also suppressed on Pt/ Al₂O₃@NC-25 after calcination treatment due the electrostatic repulsion between Cl groups and negatively charged Pt sites resulting from Pt-N interactions. Consecutive cycle test showed that Pt/Al₂O₃@NC-25 displayed a good catalytic stability. Findings in this work reveal that surface coating of supported noble metal by N-doped carbon is a promising strategy for fabricating highly selective and stable catalysts for catalytic hydrogenation reduction of *p*-CNB.

Conflicts of interest

There are no conflicts to declare.

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