Architecture engineering of nanostructured catalyst via layer-by-layer adornment of multiple nanocatalysts on silica nanorod arrays for hydrogenation of nitroarenes

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Direct consideration for both, the catalytically active species and the host materials provides highly efficient strategies for the architecture design of nanostructured catalysts. The conventional wet chemical methods have limitations in achieving such unique layer-by-layer design possessing one body framework with many catalyst parts. Herein, an innovative physical method is presented that allows the well-regulated architecture design for an array of functional nanocatalysts as exemplified by layer-by-layer adornment of Pd nanoparticles (NPs) on the highly arrayed silica nanorods. This spatially confined catalyst exhibits excellent efficiency for the hydrogenation of nitroarenes and widely deployed Suzuki cross-coupling reactions; their facile separation from the reaction mixtures is easily accomplished due to the monolithic structure. The generality of this method for the introduction of other metal source has also been demonstrated with Au NPs. This pioneering effort highlights the feasibility of physically controlled architecture design of nanostructured catalysts which may stimulate further studies in the general domain of the heterogeneous catalytic transformations.

Catalysts are sine qua non for most of the chemical processes to produce fine chemicals, pharmaceuticals, petrochemicals, among plethora of other fine chemicals1–7. From the industrial point of view, heterogeneous catalysts are often preferred to their homogeneous counterparts because of low cost, easy separation from reaction mixture, stability, and reusability8–10. In addition, the use of heterogeneous catalysts in continuous flow processes have several advantageous features such as easy mass production, automation, higher yields, and quality of products11–14.

Despite these remarkable advantages, one of the critical issues in heterogeneous catalysts is their low catalytic activity and selectivity15–17. In particular, very low catalytic activity and the rapid decay in efficiency are inevitable when they are stabilized in the non-porous solid supports. Therefore, chemical and material engineers have made several strides to design and synthesize heterogeneous catalysts with enhanced catalytic activity and selectivity. As supported heterogeneous catalysts are composed of two important segments: (i) catalytically active metal nanoparticles (NPs), and (ii) support materials18–20, the developed methodologies to enhance the catalytic activity of metal NPs entails tailoring their morphologies, facets, and atomic arrangements21–24. Support materials have also been extensively developed to uncover facile recyclability, good physiochemical stability, high surface area for the accessibility of metal nanocatalysts to various substrates and reactants25–28. In addition, various approaches
have been explored to functionalize the engineered heterogeneous catalysts thus enabling them to accommodate high loading capacity of catalytically active nanocatalysts and ameliorating their performance. For the next generation of heterogeneous catalysts endowed with specific functions and high catalytic activity, it is indispensable to design the architecture of engineered catalytically active NPs to be deposited in the accessible positions. Undeniably, the catalytic systems often developed by wet chemistry, the widely used method to prepare heterogeneous catalysts, cannot immobilize multiple catalytically active NPs on supports in a controllable manner at the designed nanoscale. Even if endeavored, it requires very complicated and time-consuming multi-step process and the deployment of toxic chemical reagents. Therefore, new methods of designing and fabricating the heterogeneous catalysts comprising unique frameworks with various components should be planned.

Physically controlled decoration of catalytically active species on certain well-known and defined supports provides a great potential to fulfill this objective. Compared to the conventional wet-chemical methods, this strategy presents advantages of low-cost, and well-tolerated loading amounts and decoration locations on supports, which is unattainable by existing synthetic techniques. Consequently, we established in our approach the highly tolerable design of benign hosts and catalytically active species. A novel template-free and designable bottom-up grown Pd NPs deposited on vertically aligned SiO2 nanostructured catalysts (v-SiO2 NRs@Pd) is reported for the reduction of nitroarenes. Due to shadowing effect during the GLAD process, SiO2 NRs can be deposited on silicon wafers. As SiO2 NRs have open top structure, accessible Pd NPs can be precisely decorated on the surface of SiO2 NRs in a layer-by-layer architecture benefitting from a multiple-step GLAD method. Furthermore, our strategy enables the combinations of various metal NPs and support materials, and nano-scale controlled affixation of catalyst NPs robustly on nanorod arrays.

Materials and methods
Oxide grains (SiO2 and SnO2) and metal pellets (Pd and Au) with 99.99% purity, used for film deposition, were purchased from Kojundo Chemistry. The other chemicals used in this study were purchased from the Daejung and Samchun Chemical Companies.

Fabrication of the v-SiO2 NRs@Pd nanostructured catalyst: A p-type Si wafers were cleaned in acetone, isopropanol, and deionized water under ultrasonication followed by drying in a nitrogen atmosphere. To fabricate SiO2 NRs, the evaporation was performed at a glancing angle with a rotation speed of 80 rpm; the wafer was located 50 cm away from the crucible, tilted at 80°. The base pressure was maintained at 1.0 × 10−6 Torr and growth rate was 1.0 Å s−1. After 250 nm-thick SiO2 NRs deposition, 3 nm-thick Pd film was sequentially deposited on the wafer at the original position (0°). These processes were repeatedly carried out until the ~1 μm-thick v-SiO2 NRs were obtained. All the fabricated catalysts were annealed at 300 °C for 30 min in a reducing atmosphere to improve the crystallinity of Pd NPs and infixed them on the support promoting their catalytic activity and durability for recycling.

The morphology of the v-SiO2 NRs@Pd nanostructured catalyst was characterized using a field-emission scanning electron microscopy (FESEM, Zeiss SUPRA 55VP). The transmission electron microscopy (TEM) and cross-sectional high-resolution TEM images were obtained using a 200 kV filed-emission TEM (JEM-2100 F, JEOL). A focused ion beam (FIB, Helios 650) was utilized to prepare the samples for TEM analysis. The energy-dispersive X-ray spectroscopy (EDS) analysis was conducted to investigate the chemical mapping of the v-SiO2 NRs@Pd nanostructured catalyst. Grazing-incidence X-ray powder diffraction (GIXRD) was performed using PANalytical X-pert Pro X-ray diffractometer. The X-ray photoelectron spectroscopy (XPS) was carried out using AXIS Ultra DLD instrument (Kratos, U.K.) operating at a base pressure of 1.6 × 10−10 mbar at 300 K. An inductively coupled plasma atomic emission spectrometer (ICP-AES, Varian 730ES) was used for the measurement of Pd amounts. The products from the catalytic reactions were analyzed by a gas chromatography mass spectrometer (GC-MS) and using an Agilent Technologies 7693 Autosampler. The structures of the chemical products were identified using a nuclear magnetic resonance (NMR, 600 MHz, High Resolution NMR Spectrometer, AVANCE Bruker).

Results and discussion
The fabrication process for the v-SiO2 NRs@Pd nanostructured catalyst using an electron beam evaporator is illustrated in Fig. 1. A multiple-step GLAD was utilized to deposit v-SiO2 NRs on a p-type Si wafer and effectively adorning Pd NPs on v-SiO2 NRs, thus substantially improving the previously reported methods. SiO2 was deposited on a p-type Si wafer tilted at 80° and rotated with a speed of 80 rpm. After deposition of a 250 nm-thick layer of SiO2 NRs, a 3 nm-thick Pd layer was deposited at a tilt angle of 0° on the v-SiO2 NRs. These steps were performed repeatedly until the total length of the v-SiO2 NRs reached to 1 μm. Finally, the fabricated catalysts were annealed at 300 °C under a reducing atmosphere to furnish robustly stabilized Pd NPs.

As shown in Fig. 2a and b, v-SiO2 NRs@Pd nanostructured catalyst was uniformly deposited on 4-inch Si wafer. The cross-sectional scanning electron microscopy image in Fig. 2b shows highly one-dimensional structures of v-SiO2 NRs@Pd catalyst on the Si wafer substrate. The nanorods were separated from each other by elongated pores and their average diameter was about 60 nm, improving the approachability of the Pd NPs and diffusion of the reactants and products. Due to their accessible nanostructure, the reagents can efficiently permeate into and interact with the v-SiO2 NRs@Pd nanostructured catalyst. To clarify the existence of Pd NPs in the v-SiO2 NRs, GIXRD was performed which revealed the peaks assignable to the (111) and (200) reflections of the face-centered cubic (FCC) Pd NPs (Fig. 2c). XPS analysis was also carried out to confirm the oxidation state of Pd species in v-SiO2 NRs@Pd nanostructured catalyst (Fig. 2d). The peaks corresponding to Si, O, and Pd were clearly observed in XPS spectra (Fig. S1). The Pd 3d5/2 and Pd 3d3/2 binding energies were found to be at 335.8 and 341.1 eV, respectively thus indicating the formation of zero-valent Pd species in v-SiO2 NRs@Pd nanostructured catalyst.
To further analyze the nanostructure of v-SiO$_2$ NRs@Pd catalyst, TEM images were obtained as shown in Fig. 3a; four dense layers of Pd NPs were seen clearly deposited on v-SiO$_2$ NRs. The Pd NPs were also observed in the side walls of the v-SiO$_2$ NRs. The energy dispersive spectroscopy (EDS) mapping on the v-SiO$_2$ NRs@Pd nanostructured catalyst was conducted. Figure 3b–d show the chemical mapping images of Si, O, and Pd elements for the selected areas in Fig. 3a (orange dashed rectangle). For Si and O elements, a uniform distribution across the v-SiO$_2$ NRs has been observed. However, Pd NPs were distributed in a layer-by-layer architecture with total of four layers. The Pd element observed in between the layers has been attributed to the Pd NPs on sidewalls of v-SiO$_2$ NRs. Figure 3e–g shows the high resolution TEM (HRTEM) images of Pd NPs on the surface (yellow rectangle) of the SiO$_2$ NRs; Pd NPs were crystallized with the (111) preferred orientation. From the above results, it can be concluded that Pd NPs are efficiently decorated on the whole surface area of SiO$_2$ NRs and immobilized at desired locations on SiO$_2$ NRs through multiple-step GLAD method to realize a favorable spatial confinement. In other words, the catalytic active sites, Pd NPs, are well-refined on the v-SiO$_2$ NR supports, resulting in more accessible catalytically active sites on the v-SiO$_2$ NRs@Pd nanostructured catalyst.

To further demonstrate the advantages of this physical architecture design, we fabricated v-SiO$_2$ NRs decorated with Au and Pd alternately (v-SiO$_2$ NRs@Pd/Au), using the same protocol. The four dense layers of metal NPs were observed on the v-SiO$_2$ NRs@Pd/Au nanostructured catalyst (Fig. S2). The EDS mapping on the v-SiO$_2$ NRs@Pd/Au nanostructured catalyst revealed that the Pd and Au NPs were layer-by-layer deposited on v-SiO$_2$ NRs by turns. The HRTEM images also confirmed that the metal NPs were single crystalline in each layer for both, the Au and Pd NPs. To confirm the oxidation state of Pd and Au NPs, XPS analysis was conducted. The peaks observed at 84.1, 87.8, 335.8, and 341.2 eV corresponded to Au 4$f^{7/2}$, Au 4$f^{5/2}$, Pd 3$d^{5/2}$ and Pd 3$d^{3/2}$, respectively. This indicated that the Pd and Au NPs on the v-SiO$_2$ NRs@Pd/Au nanostructured catalyst were in zero-valent state. The similar catalyst was also synthesized on SnO$_2$ supports using our strategy to broaden various type of supporting materials. As shown in Fig. S3, Pd NPs were well-decorated on v-SnO$_2$ NRs. These results illustrated that metal NPs could be stabilized on the designed sites on various supports using the multiple-step GLAD physical method.

Compared with previously reported fabrication processes for nanostructured catalysts, this approach has several advantages including favorable design, precise architecture engineering, and accessible metal nanocatalysts for the reactants, etc. Using wet chemical methods, mainly applied for nanostructured catalysts fabrication, catalytic sites are randomly distributed due to the lack of controllability and the need for complicated functionalization processes. In addition, it is very difficult to combine two or more metal NPs into the nanostructured support. However, this approach circumvents the limitations of previous reports for the catalysts on silicon.

![Figure 1. Synthetic procedure of v-SiO$_2$ NRs@Pd nanostructured catalyst.](image-url)
wafers as the supporting materials via top-down fabrication process which often is accompanied by harmful
and toxic HF etching. The present strategy provides the varying catalytically active sites precisely integrated
on the support and their location can be controlled on nanoscale, thus improving the approachability of the
catalytic NPs to reactants.

To verify the catalytic activity and chemoselectivity of the v-SiO$_2$ NRs@Pd nanostructured catalyst, we
conducted the hydrogenation of various substituted nitroaromatics at room temperature; a reaction that has
important impact in many industries, including environmental safety. Our designed nanostructured catalyst
presented superb catalytic activity for the selective hydrogenation of nitroaromatics in aqueous solution (Table
1). The procedure was accomplished under eco-friendly mild conditions, attaining high yields for the hydrogena-
tion of substituted nitroaromatics to the corresponding aminoaromatics. The hydrogenation of nitroaromatics
bearing two reducible groups could be achieved selectively to the amino compound as the only product, while
another reducible group was retained unchanged (Table 1, entries 4–6). In addition, achieving highly selective
reduction of nitro groups of aromatic molecules possessing olefin bonds is very challenging as there is high
probability for both the functional groups to be readily hydrogenated. Importantly, this process proceeded with
high selectivity (99%), reducing only nitro group (Table 1, entry 8) as affirmed by nuclear magnetic resonance
spectroscopy (Fig. S4). Furthermore, a comparison of the catalytic performance study between the v-SiO$_2$ NRs@
Pd nanostructured catalyst and previously reported noble metal-comprising catalysts shows a highly comparable
catalytic activity (Table S1).

To clarify the correlation between accessibility of Pd nanocatalysts and their activity, as a proof-of-concept,
control experiments were carried out using various nanostructured catalysts in the hydrogenation of nitroben-
zene as a model reaction under identical reaction conditions (Table S2). Initially, only Pd NPs were deposited
on silicon wafer by GLAD method aiming for simple recycling though dipping and removing of the silicon
substrate possessing stable nanocatalysts. The reaction produced a low yield of (15%) of aniline under optimized
reaction conditions. FESEM images of this catalyst showed that the surface of the silicon wafer substrate was
thoroughly covered by stacked Pd NPs without approachable opening for the diffusion of the reactants, result-
ing in low catalytic activity (Fig. S5). In addition, micro- and nano-space channels provide permeable paths to
organic reagents, thus improving the accessibility of the Pd nanocatalysts. We also fabricated and evaluated

Figure 2. (a) A photographic image of v-SiO$_2$ NRs@Pd nanostructured catalyst on p-type Si wafer (left) and
bare p-type Si wafer. The diameter of the p-type Si wafers is 4 inches. (b) Cross-sectional SEM image of the
v-SiO$_2$ NRs@Pd nanostructured catalyst. (c) GI-XRD pattern of v-SiO$_2$ NRs@Pd nanostructured catalysts on p-Si
wafer. (d) XPS core level spectrum of Pd 3d for v-SiO$_2$ NRs@Pd nanostructured catalyst.
the catalytic activity of Pd NPs on SiO2 arrays having ~ 200 nm height (Fig. S6). Interestingly, the SiO2 columns exposing Pd NPs could hydrogenate the nitro group of nitrobenzene in relatively higher yields (35%). Although the prototype short length SiO2 arrays were partly aggregated, from this result we found that the Pd NPs stabilized on sturdy SiO2 support functioned to some extent effectively, which can be attributed to internal surface and pore diffusivity. Consequently, we designed and fabricated SiO2 arrays and aligning them separately from each other with ~ 600 nm length possessing Pd NPs on their scaffolds as depicted in FESEM image (Fig. S7). Although the Pd NPs were located irregularly on the SiO2 arrays, their combination improved their interactions with the solvent and reactants via the accessible pores, promoting efficient catalytic reactions (Table S1, entry 3). Finally, we designed and developed fully arranged and assorted v-SiO2 NRs@Pd catalyst having permeable tunnel paths and elongated pores between SiO2 columns with four strata of attainable Pd NPs, and maximizing their catalytic activity in the given geometric configuration. Similarly, the alternative four dense layers of Pd NPs and Au NPs in the v-SiO2 NRs@Pd/Au nanostructured catalyst provided high catalytic activity in the model reduction. The combination and the activity of different nanocatalysts can easily be tuned and optimized by this method, controlling their locations on supports and distance between the nanocatalysts and the arrays. The controlled pores in the nanostructure provided facile adsorption of reactants to the Pd NPs and subsequent desorption of products from them. Additionally, a larger amount of Pd NPs can be decorated on the whole surface of SiO2 NRs exposing large amount of accessible Pd sites for the catalytic reactions. Thus, v-SiO2 NRs@Pd nanostructured catalyst could expedite Suzuki coupling reaction of various aryl halides with phenylboronic acid (Table 2); reaction of aryliodides and arylbromides with phenylboronic acid proceeded with excellent conversion in 3 h and 5 h, respectively.

Recycling and reuse of the catalysts are crucial parameters because of the environmental impact and practical applications. Although many heterogeneous catalysts have been developed for the reduction of nitroaromatics, they generally entail very arduous recycling procedures such as centrifugation, filtration, laborious work-up purification, etc. However, v-SiO2 NRs@Pd nanostructured catalyst could be easily separated being withdrawn
| Entry | Substrate | Product | Yields (%)<sup>a</sup> |
|-------|-----------|---------|------------------------|
| 1     | \( \text{NO}_2 \) | \( \text{NH}_2 \) | 99                     |
| 2     | \( \text{NO}_2 \) | \( \text{NH}_2 \) | 92                     |
| 3     | \( \text{NO}_2 \) | \( \text{NH}_2 \) | 93                     |
| 4     | \( \text{NO}_2 \) | \( \text{NH}_2 \) | 91                     |
| 5     | \( \text{NO}_2 \) | \( \text{NH}_2 \) | 94                     |
| 6     | \( \text{NO}_2 \) | \( \text{NH}_2 \) | 92                     |
| 7     | \( \text{NO}_2 \) | \( \text{NH}_2 \) | 95                     |
| 8     | \( \text{NO}_2 \) | \( \text{NH}_2 \) | 94                     |

Table 1. Heterogeneous reduction of substituted nitroarenes catalyzed by v-SiO<sub>2</sub> NRs@Pd nanostructured catalyst. Reaction conditions: Substituted nitroarenes (0.1 mmol), NaBH<sub>4</sub> (0.12 mmol), v-SiO<sub>2</sub> NRs@Pd nanostructured catalyst (1 mol% Pd), H<sub>2</sub>O (20 mL), room temperature, and 1.5 h. <sup>a</sup> Yields were determined by GC–MS.
from the reaction mixture by tweezers, as they have a monolithic structure, a salient advantage of our catalyst. The durability and reusability of our catalyst was investigated for the hydrogenation of nitrobenzene. After completion of the reaction, the v-SiO$_2$ NRs@Pd nanostructured catalyst was simply removed from the reaction mixture, rinsed with water, and reused in the next cycle. The activity of the catalyst remained stable even after seven consecutive reaction cycles, though there was a minuscule decline in activity of the catalyst (Table 3). As the catalytic activity of v-SiO$_2$ NRs@Pd nanostructured catalyst originates from its special architecture engineering of one body nanostructure, we precisely characterized the recovered v-SiO$_2$ NRs@Pd nanostructured catalyst by EDX, TEM, and HRTEM techniques (Fig. S8), affirming the constant stability of the designed catalyst after the recycling. Consequently, Pd leaching could be the reason for this catalytic decrease, which was supported by ICP-AES analysis; negligible Pd species was detected.

**Conclusions**

The novel synthesis of v-SiO$_2$ NRs@Pd nanostructured catalyst described in this study exhibited promising catalytic characteristics for the hydrogenation of nitroaromatics and the Suzuki cross–coupling reaction. The vertically aligned SiO$_2$ NRs provided an excellent platform for Pd nanocatalysts to be firmly cohered to create a monolithic structure, which is superior in the realization of a simple recycling process. Because this fabrication procedure is based on physical vapor deposition under vacuum conditions, it can be applied to various combinations of metal nanocatalysts and supporting nanostructures without complicated synthesis procedures often deployed in wet chemical methods. The effective design of nano-architectures for a monolithic structure.

| Entry | Substrate | Time (h) | Yields (%)$^a$ |
|-------|-----------|----------|---------------|
| 1     | I         | 3        | 97            |
| 2     | I         | 3        | 94            |
| 3     | Br        | 5        | 95            |
| 4     | Br        | 5        | 91            |
| 5     | Br        | 5        | 87            |

Table 2. Suzuki cross–coupling reactions catalyzed by v-SiO$_2$ NRs@Pd nanostructured catalyst. Reaction conditions: Aryl halide (0.1 mmol), phenylboronic acid (0.12 eq.), v-SiO$_2$ NRs@Pd nanostructured catalyst (1 mol%), K$_2$CO$_3$ (1.5 eq.), DMF/H$_2$O (5:1), 100 °C. $^a$Yields were determined by GC–MS.
nanocatalysts-attached SiO2 NRs will expectedly open new perspectives toward the development of highly recyclable nanocatalysts for other catalytic reactions.

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Table 3. Reuse of v-SiO2 NRs@Pd nanostructured catalyst in the heterogeneous reduction of nitrobenzene. Reaction conditions: Substituted nitroarenes (0.1 mmol), NaBH4 (0.12 mmol), v-SiO2 NRs@Pd nanostructured catalyst (1 mol% Pd), H2O (20 mL), room temperature, and 1.5 h. aYields were determined by GC–MS.
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Author contributions

K.H. and J.M.S. contributed equally to this work. M.S. and H.W.J. designed and conceived the idea. M.S. and T.H.L. assisted the transmission electron microscopy characterization and analysis. M.S., K.H., and J.M.S. assisted synthesis and analyses. M.S., H.W.J., S.R., S.H.C., and R.S.V. developed the project and revised the manuscript. M.S., H.W.J., and R.S.V. supervised, monitored, and directed the project. All authors contributed in scientific discussions and preparing the manuscript.

Competing interests

The authors declare no competing interests.
Additional information

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