PtPd hollow nanocubes with enhanced alloy effect and active facets for efficient methanol oxidation reaction

An alternating-reduction approach by galvanic replacement and co-reduction, which enables incorporation of Pd into Pt shell, is developed to synthesize PtPd hollow nanocubes with highly catalytically-favoured (100) facets and enhanced alloy effect for efficient methanol oxidation reaction.

We report an alternating-reduction approach by galvanic replacement and co-reduction to enable incorporation of Pd into Pt shell, and the obtained PtPd hollow nanocubes with an enhanced alloy effect and highly active (100) facets show high catalytic activity and superior durability in the methanol oxidation reaction.

Highly cost-effective Pt catalysts should increase the atom utilization efficiency together with controllable nanostructures and facets for high activity and selectivity.\(^1\)\textsuperscript{-3} Hollow nanocubes with (100) facets represent an emerging nanostructure in high-performance Pt design, which exhibit greatly reduced Pt usage and high activity and selectivity of the (100) facets.\(^4\)\textsuperscript{-8} Nanocages with very few atom layers of Pt thus show significant enhancement of electrocatalytic activity and stability.\(^9\)\textsuperscript{-13} It has to be pointed out that this is an elegant idea for the fabrication of catalysts with minimum Pt usage and maximum exposure of highly active (100) facets, in spite of loss of the alloying effect. Bimetallic nanostructured Pt-based metals not only present the physicochemical properties of Pt, but also usually show a superior performance compared with pure Pt counterparts.\(^14\)\textsuperscript{-16}\n
The catalytic activity of bimetallic Pt-based nanoparticles can be greatly enhanced by maximizing the expression of active facets and/or sites toward a specific reaction.\(^17\)\textsuperscript{-19} However, by exploiting Pd nanocubes with exposed {100} facets as a template, it is technically difficult to increase the Pd amount in the Pt shell because a relatively high energy barrier limits the diffusion of Pd.\(^5\) Therefore, an enhancement of the Pd content in PtPd hollow nanocages with dominating {100} facets is rarely reported. Herein, an alternating-reduction approach by galvanic replacement and co-reduction, which we developed, enables incorporation of Pd into a Pt shell. The resultant PtPd hollow nanocubes after etching (denoted as PtPd HNCs-E) possessing an enhanced alloy effect and the highly active {100} facets show catalytic activity and stability in the methanol oxidation reaction (MOR).

![Fig. 1](image-url) Scheme for the formation of enhanced alloy effect in PtPd HNCs-E. (a) Pd nanocube and its surface atoms. (b) Growth of Pt atoms by galvanic replacement. (c) Formation of the alloy shell in PtPd HNCs by co-reduction of Pt- and Pd-salts by DMF. (d) Mixed PtPd atoms in PtPd HNCs-E after removal of Pd.

\[ \text{[PtCl}_6\text{]}^{2-} + 6\text{I}^- \rightarrow \text{[PtI}_6\text{]}^{2-} + 6\text{Cl}^- \quad (1) \]
\[ 2\text{Pd} + \text{[PtI}_6\text{]}^{2-} + 2\text{I}^- \rightarrow \text{Pt} + 2\text{[PdI}_4\text{]}^{2-} \quad (2) \]
\[ \text{[PtI}_6\text{]}^{2-} \text{DMF} \rightarrow \text{Pt} + 6\text{I}^- \quad (3) \]
The co-reduction of Pt(IV) and Pd(II) by DMF would occur (eqn (3)) resulting in the coordination of I⁻ to Pt(IV). Moreover, the addition of I⁻ will facilitate the galvanic replacement between Pd and Pt. Galvanic replacement would lead to the growth of Pt around the Pd nanocube surface and Pd⁴⁺ dissolves to form [PdI₄]²⁻ in the presence of I⁻ (eqn (1)). The coordination effect can be visualized through the color change from pale yellow to dark color when colorless I⁻ was added to [PtCl₆]³⁻ (Fig. S1a–c, ESI†), indicating the formation of [PtI₆]²⁻, which could alter the reduction kinetics of Pt(IV). Then, a co-reduction of Pt(IV) and Pd(II) by DMF would occur (eqn (3) and (4)). Such an alternating-reduction by galvanic replacement and co-reduction will incorporate Pd atoms into the Pt shell (Fig. 1c). The dissolution of the Pd core and deposition of Pd into a growing Pt shell will be repeated several times, resulting in the hollow structure of the PtPd nanocubes (denoted as PtPd HNCs). Finally, the unstable Pd template in the interior of the PtPd HNCs would be removed by acid etching, resulting in PtPd HNCs-E with highly active {100} facets (Fig. 1d). To gain more insight into the formation of the PtPd alloy shell, zeta potentials were measured to characterize the surface charges of nanomaterials. The value of the zeta potential was changed during the synthesis process of PtPd-HNCs, starting from −8.42 mV (Pd nanocubes), to −23.2 mV (10 min), −12.4 mV (30 min), −13.6 mV (60 min), −14.2 mV (90 min) and −14.0 mV (120 min) (Fig. S2, ESI†). Compared to the zeta potential of Pt nanocrystals (−33.6 mV), the increased value at 10 min indicates the formation of surface Pt, and the decreased value at 30 min indicates the formation of a PtPd alloy. The zeta potential value was kept almost unchanged from 60 to 120 min, which illustrates a nearly unchanged structure and stable state of the PtPd shell.

The nanostructure of PtPd HNCs-E can be visualized by transmission electron microscopy (TEM) images in Fig. 2a and b, exhibiting an obviously conformal and hollow structure with the average diameter of 24 nm. Considering the average diameter of 20 nm for the original Pd nanocubes (Fig. S3a and b, ESI†), the thickness of the shell can be calculated as 4 nm. Two selected regions in Fig. 2b are enlarged as shown in Fig. 2c and e, and their corresponding fast Fourier transform (FFT) results are shown in Fig. 2d and f, respectively. PtPd HNCs-E exhibited a lattice spacing of 0.195 nm, which is between 0.194 and 0.196 nm, corresponding to the {100} interplanar distance of face-centered cubic (fcc) palladium and platinum, respectively. Compared to the spacing of 0.194 nm of Pd nanocubes (Fig. S3c, ESI†) and 0.195 nm of PtPd HNCs (Fig. S4, ESI†), the PtPd HNCs-E showed an alloy structure of PtPd and unchanged shell structure from PtPd HNCs. Moreover, two sets of spots in FFT can be identified as the reflections of the PtPd (200) and (400) planes. All these results suggest that the PtPd HNCs-E are dominantly enclosed by {100} facets. Fig. 2g and h show the high-angle annular dark-field scanning TEM (HAADF-STEM) images with different magnifications. Under dark field, the hollow structure can be seen more clearly. Fig. 2i–l show the elemental mapping of PtPd HNCs-E measured by HAADF-STEM energy-dispersive X-ray spectroscopy (HAADF-STEM-EDX). As can be seen, Pt and Pd elements are homogeneously dispersed throughout the nanoparticles except for very small amounts of Pt elements dispersed in the outer surface, indicating the high alloying degree with a Pt-rich surface in PtPd HNCs-E. According to the results of inductively coupled plasma-atomic emission spectrometry (ICP-AES), the atomic ratio of Pt: Pd was 2: 1 in the PtPd HNCs-E. Notably, the Pd content in the PtPd HNCs-E is much higher than those reported for etched PtPd nanocages. The remaining Pd content indicates a stable alloy structure of PtPd HNCs-E which can resist the further etching of HNO₃ even lasting for five days.

To investigate the crystal structure changes among Pd nanocubes, PtPd HNCs, and PtPd HNCs-E, X-ray diffraction (XRD) measurements were conducted. As shown in Fig. 3a, five peaks around 40°, 46°, 68°, 82°, and 87° can be readily indexed to the (111), (200), (220), (311) and (222) reflections of fcc Pt.
of Pt$^{2+}$ is changed from 23% to 33% and Pd$^{2+}$ from 29% to 47% in PtPd HNCs-E owing to the surface oxidation after nitric acid treatment.

The electrocatalytic activities of PtPd HNCs-E for the MOR were investigated in comparison to PtPd HNCs and commercial Pt/C under the same conditions. The mechanism of the MOR on the PtPd alloy is depicted in Fig. S6, ESI.† The electrochemically active surface areas (ECSAs), which were calculated based on charges involved in the adsorption of hydrogen in the cyclic voltammograms (CVs) recorded in 0.1 M KOH (Fig. 4a), were 25.5, 23.5, and 51.7 m$^2$/g$_{metal}$ for PtPd HNCs-E, PtPd HNCs, and commercial Pt/C, respectively. Fig. 4b shows the CV curves of these electrocatalysts for the MOR. The onset potential of PtPd HNCs-E is much lower than that of both PtPd HNCs and commercial Pt/C, indicating that the oxidation of methanol is much easier to be triggered on the surface of PtPd HNCs-E.28,29 The mass activities and specific activities for these three catalysts are summarized in Fig. 4c and Table S1, ESI.† PtPd HNCs-E exhibit a mass activity of 1.7 mA g$_{metal}^{-1}$, which is 1.9 and 3.5 times that of PtPd HNCs (0.89 mA g$_{metal}^{-1}$) and commercial Pt/C catalysts (0.49 mA g$_{metal}^{-1}$), respectively. In addition, PtPd HNCs-E exhibit the highest specific value of 6.7 mA cm$^{-2}$, which is 1.8 and 7.1 times that of PtPd HNCs (3.8 mA cm$^{-2}$) and commercial Pt/C catalysts (0.95 mA cm$^{-2}$), respectively. Notably, the catalytic activities of PtPd HNCs-E towards the MOR are also superior to that of other previously reported Pt-based catalysts (Table S2, ESI†). Moreover, PtPd HNCs-E also exhibit the best MOR activities performed in 0.5 M KOH.
and 2 M methanol (Fig. S7, ESI†). The improved electrocatalytic properties of PtPd HNCs-E can be attributed to the catalytically preferential {100} facets of hollow cubic structures and the downshift of the d-band center of the Pt electronic structure induced by an electronic interaction between the PtPd alloy. Furthermore, CO stripping measurement results showed that PtPd HNCs-E exhibited the best CO tolerance ability (Fig. S8, ESI†).

Chronoamperometric curves were recorded to probe the stability of the catalysts (Fig. 4e). PtPd HNCs-E show a higher specific current density and slower current decay than PtPd HNCs and commercial Pt/C during the whole test process, demonstrating the highest electrocatalytic stability of PtPd HNCs-E for the MOR. Furthermore, the durability of PtPd HNCs-E and commercial Pt/C was studied by performing accelerated durability tests (ADTs) at room temperature in 0.1 M KOH. After 6000 cycles of ADT, a slight drop of current density can be found for PtPd HNCs-E with increasing CV cycle numbers (Fig. S9a, ESI†), while a conspicuous drop for commercial Pt/C is seen (Fig. S9b, ESI†). After 2000, 4000, and 6000 cycles, the specific ECSA of PtPd HNCs-E only dropped by 13%, 18%, and 23%, whereas the specific ECSA of the commercial Pt/C catalyst sharply decreased by 28%, 41%, and 54% (Fig. S10 and Table S3, ESI†). Moreover, the characterization after 6000 cycles of the ADTs indicates the higher MOR durability and stable morphology of PtPd HNCs-E compared to commercial Pt/C (Fig. 4f and Fig. S11–13, ESI†). These results demonstrate that in addition to high MOR activity and CO tolerance, PtPd HNCs-E also exhibit superior electrochemical durability.

In summary, we have presented an effective strategy to synthesize PtPd HNCs-E with exposed {100} active facets and optimized alloy structures. PtPd HNCs-E exhibit 3.5 times the mass activity and a remarkable 7.1 times of the specific activity, as well as much higher CO tolerance and durability than a commercial Pt/C catalyst in the MOR. The enhanced electrocatalytic performance of PtPd HNCs-E could be ascribed to the cubic morphology with catalytically-favoured {100} facets, to the hollow structure with more exposed Pt active sites, and to the synergistic effects from the enhanced alloy effect of PtPd bimetallics. This work displays new possibilities for designing novel nanomaterials with high activity and superior stability towards methanol electrooxidation or other practical applications.

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Conflicts of interest

There are no conflicts to declare.