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Controlled morphology and dimensionality evolution of NiPd bimetallic nanostructures

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Highlights

- Versatile NiPd nanostructures prepared via one-pot aqueous synthesis.
- HCl concentration and reaction aging time plays vital role in the morphology evolution.
- The diverse NiPd nanostructures exhibit enhanced catalytic activity toward the reduction of 4-nitrophenol.

Abstract

Controlling the morphology of noble metal-based nanostructures is a powerful strategy for optimizing their catalytic performance. Here, we report a one-pot aqueous synthesis of versatile NiPd nanostructures at room temperature without employing organic solvents or surfactants. The synthesis can be tuned to form zero-dimensional (0D) architectures, such as core–shell and hollow nanoparticles (NPs), as well as nanostructures with higher dimensionality, such as extended nanowire networks and three-dimensional (3D) nanodendrites. The diverse morphologies were successfully obtained through modification of the HCl concentration in the Pd precursor solution, and the reaction aging time. An in-depth understanding of the formation mechanism and morphology evolution are described in detail. A key factor in the structural evolution of the nanostructures was the ability to tune the reduction rate and to protonate the citrate stabiliser by adding HCl. Spherical core–shell NPs were formed by the galvanic replacement-free deposition of Pd on Ni NPs which can be transformed to hollow NPs via a corrosion process. High concentrations of HCl led to the transition of isotropic spherical NPs into anisotropic wormlike nanowire networks, created through an oriented attachment process. Aging of these nanowire networks resulted in the formation of 3D porous nanodendrites via a corrosion process. The diverse structures of NiPd NPs were anchored onto acid treated-activated carbon (AC) and exhibited improved catalytic efficiency towards the hydrogenation of 4-nitrophenol (4-NP) to 4-aminophenol (4-AP).

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1. Introduction

Metal nanoparticles (NPs) have garnered sustained research interest because of their unique physical and chemical properties that can be controlled and optimized by altering their size, shape, composition, crystal structure, or surface functionality [1–3]. Noble metal NPs such as Pd are employed in a myriad of catalytic applications, including hydrogenations [4], oxidations [5], organic coupling reactions [6] and electrochemical reactions [7]. The limited natural reserves and cost associated with extracting Pd from the ground has led to the tremendous use of bimetallic noble metal NPs in catalytic applications, as noble metals can be combined with less expensive transition metals without affecting catalytic performance.

Bimetallic nanostructures often display remarkably improved catalytic activity and selectivity compared to their monometallic counterparts due to the synergistic effects between the constituent metals [8–10]. In particular, alloying Pd with non-precious first-row transition metals, such as Ni, Co, and Fe can lead to cost-effective catalysts and enables modification of the electronic and geometric structures for enhanced activity and/or selectivity. NiPd NPs have shown exceptional activity and stability towards a range of coupling reactions [11–13], and the oxygen reduction reaction (ORR) [14,15]. PdNi [16] and PdCu NPs [17] have also displayed superior performance in the oxidation of formic acid compared to monometallic Pd NPs.

In addition to alloying, the morphology of bimetallic NPs can be tailored with specific crystal facets to maximize the number of low-index facets [3,28,29]. Engineered hollow nanostructures have however often required multistep reactions in the presence of precursors and cations, including hydrogenations [4], oxidations [5], organic coupling reactions [6] and electrochemical reactions [7]. The limited natural reserves and cost associated with extracting Pd from the ground has led to the tremendous use of bimetallic noble metal NPs in catalytic applications, as noble metals can be combined with less expensive transition metals without affecting catalytic performance.

2. Experimental

2.1. Materials

Trisodium citrate dihydrate (Na3C6H5O7·2H2O, 99%), sodium borohydride (NaBH4, 99.99%), palladium(II) chloride (PdCl2, 99.9%), sodium palladium(II) tetrachloride (Na2PdCl4, 98%), cobalt (II) chloride hexahydrate (CoCl2·6H2O, ≥99%), iron(III) chloride (FeCl3, ≥99%), copper(II) chloride dihydrate (CuCl2·2H2O, ≥99.95%), hydrochloric acid (HCl, 37%), nitric acid (HNO3, 65%), sodium hydroxide (NaOH, 98%), commercial Pd/C (5 wt%) and 4-nitrophenol (O2NC6H4OH, 98%) were purchased from Sigma-Aldrich. Nickel(II) chloride hexahydrate (NiCl2·6H2O, 99.99%) was obtained from Alfa Aesar. All chemicals were of analytical grade and used as received without further purification. Activated charcoal (untreated powder; 100–400 mesh) was purchased from Honeywell FlukaTM. Deionized water (DI H2O) was used as the solvent. All glassware and magnetic stir bars were cleaned with aqua regia, followed by rinsing with DI H2O before drying in an oven.

2.2. Synthesis

A one-pot synthesis was used to fabricate the NiPd bimetallic NPs. Firstly, the sacrificial Ni NPs were synthesized using the borohydride reduction method. Briefly, in a two-necked 100 mL round bottom flask, 4.7 mg of NiCl2·6H2O (0.02 mmol), and 29.4 mg of trisodium citrate (TSC) (0.1 mmol) were dissolved in 40 mL of DI H2O at room temperature (~20 °C). The solution was stirred and purged with N2 for 30 min to eliminate the dissolved O2. Then, 2 mL of freshly prepared aqueous NaBH4 solution (7.5 mg) was added dropwise into the solution under magnetic stirring at 550 rpm. The appearance of a dark yellow color in the solution indicated the formation of Ni NPs. After 10 min, 40 mL of H2PdCl4 solution (0.02 mmol) was injected into the Ni NPs with a rate of 1.4 mL/min at room temperature. The reaction mixture was allowed to stir for another 3 h. To validate the critical role of the aging time, the colloidal solutions of Ni NPs were stored for 24 h under ambient conditions (air, ~20 °C) without any agitation. For comparison, the same procedure was done using Na2PdCl4 (0.02 mmol) solution as a Pd precursor without extra HCl addition, while other conditions were kept unvaried. For Co-Pd, Fe-Pd, and Cu-Pd systems, the same protocol was used except replacing NiCl2·6H2O with CoCl2·6H2O, FeCl3, and CuCl2·2H2O, respectively with the same molar amount.

2.2.1. Preparation of H2PdCl4 solution (0.02 mmol)

Stock solutions of 10 mM H2PdCl4 were prepared by dissolving 88.65 mg of PdCl2 in 50 mL of a 0.2 M, 0.1 M and 0.02 M HCl H2PdCl4 solutions (0.02 mmol) were prepared by diluting 2 mL of stock solutions to 40 mL with DI H2O to give 0.02 mmol H2PdCl4/10 mM HCl solution, 0.02 mmol H2PdCl4/5 mM HCl and 0.02 mmol H2PdCl4/1 mM HCl solutions, respectively.

2.3. Immobilization of NiPd NPs on acid-treated activated carbon

For improving the absorbing ability of activated carbon (AC), it was treated with HNO3 acid. 1 g of AC was refluxed in 10 mL of 6 M HNO3 solution while stirring for 6 h at 95 °C. The AC was collected by centrifuge and washed until neutral pH of the filtrate.
To immobilize the NPs, the AC was added to the as-prepared NPs solution and sonicated for 10 min. The mixture was stirred at room temperature for 2 h and collected by centrifugation. The catalyst was washed with DI water and acetone several times to remove the residual impurities and then dried at 50 °C overnight.

2.4. Characterization

_Microscopy_. Low-resolution transmission electron microscopy (TEM) images were recorded using a JEOL 2100 transmission electron microscope operated at 200 kV. High-resolution (HR)TEM images, scanning transmission electron microscopy (STEM), and energy dispersive X-ray spectroscopy (EDX) mapping and line scanning were performed on the FEI Titan transmission electron microscope at an operating voltage of 300 kV. For electron microscopy grids, as-prepared NPs solution was directly dropped on a carbon-coated Cu grid (300 mesh, Agar Scientific), followed by drying under ambient condition (air, ~20 °C). X-ray Photoelectron Spectroscopy (XPS) was carried out using a KRATOS AXIS ULTRA spectrometer equipped with an Al Kx (hv = 1486.58 eV) X-ray source. All spectra were calibrated using the C 1s peak at 284.8 eV. Inductively coupled plasma optical emission spectroscopy (ICP-OES) was conducted for determining the accurate amounts of metals deposited on the AC support using Varian 710-ES ICP Optical Emission Spectrometer. Working parameter: the power of 1 kW with the auxiliary of 1.5 L/min and nebuliser pressure of 200 kPa. The emission lines used to measure the Ni and Pd concentrations were 231.604 nm and 340.458 nm, respectively. The microwave-assisted digestion method using nitric acid and hydroperoxide was performed to prepare the samples for the ICP measurements. UV–Vis spectroscopy was performed for observing the catalytic reduction of 4-nitrophenol using Genesys 10 s UV–Vis Thermo scientific spectrometer.

2.5. Catalytic evaluation of nanostructures

The catalytic reduction of 4-nitrophenol (4-NP) was chosen as a model reaction to assess the catalytic efficiency of NiPd nanoparticles with different morphologies and commercial Pd/C. Due to the low solubility of 4-NP in water, 4-NP was first dissolved in NaOH. The resulting 4-nitrophenolate ion was then reduced by NaBH₄ to a solution of Ni NPs. Fig. 1A shows the formation of 4-nitrophenolate ion conversion to 4-aminophenol (4-AP) was monitored by UV–Vis spectrometer by recording the absorption spectra of the reaction mixture with a 30 s time intervals at ambient temperature.

3. Results and discussion

Fig. 1 illustrates the shape evolution of NPs upon the addition of a H₂PdCl₄/10 mM HCl solution into a solution of Ni NPs. Fig. 1A shows that TEM analysis of NPs after 3 h had a predominately solid spherical morphology. Detailed size distribution analysis showed a bimodal diameter distribution with modes centered at 5.5 and 15.7 nm (Fig. S1A). In addition, there was a minor contribution of hollow spheres with a diameter of 19.1 ± 3.4 nm (Fig. S1B) and a shell thickness of 4.3 ± 1.8 nm (Fig. S1C). EDX mapping and line scans of the solid NPs (15.7 nm), shown in Fig. 1(B, C), indicate that they consist of a core–shell structure. The typical chemical composition of the NPs, as determined by EDX analysis showed a Ni core with monometallic Pd located in the shell. This core–shell structure was also evidenced by XPS analysis, displayed in Fig. S2A, which showed no peaks for the Ni 2p core level due to their attenuation from the Pd overlayer. EDX analysis (Fig. S3) confirmed that the smaller diameter NPs (5.5 nm) were composed of Pd only, attributed to a minor degree of homogeneous nucleation of Pd.

Fig. 1D shows the morphological evolution of the core–shell NPs to a hollow nanostructure on aging the solution for 24 h due to etching of the Ni core. The mean diameter of the hollow nanospheres was determined to be 14.4 ± 2.4 nm (Fig. S4A) with a shell thickness of 2.5 ± 0.5 nm (Fig. S4B). EDX mapping and line scans of the hollow nanospheres (Fig. 1(E, F)) confirmed etching of the Ni core to leave a shell consisting mostly of Pd with an estimated composition of Ni₄Pd₉₆₅ also supported by XPS analysis (Fig. S2A), however, it could not be quantified accurately due to detection limits. HRTEM analysis of the hollow NPs (the inset of Fig. 1D) reveals the lattice fringe with a d spacing of 0.234 nm, matching the (111) plane of face-centered cubic Pd [35].

Conducting the same synthesis procedure at a lower pH by using a H₂PdCl₄/10 mM HCl solution allows assess to nanostructures with 2D and 3D architectures. After 3 h, Fig. 2A shows the formation of 2D wormlike nanowire networks. Despite being overlapped, folded and twisted, these nanowire networks were well dispersed, highly interconnected and extended to hundreds of nanometers in length as observed in low-resolution TEM (Fig. S3A). The typical diameter of the wormlike nanowire was 5.6 ± 1.1 nm (Fig. S3B) and displayed excellent uniformity throughout the nanowire networks. These nanowire networks are potentially highly effective catalysts due to their inherent anisotropic structures and the reactivity of the surface atoms at the different facets. Some of these nanowires were connected with a hollow head forming a tadpole structure. TEM analysis of an individual nanowire showed that they consisted of a polycrystalline assembly of nanoparticles made up of nodules (marked by circles in Fig. 2B) connected through slightly narrower stems (marked by rectangles in Fig. 2B). High-resolution TEM and EDX were carried out to further understand the structure and composition of the nanowire-tadpole morphologies. EDX line scans, taken from various positions along the nanowire (Fig. 2D), revealed different chemical compositions in the stem and nodes of the nanowire. The mean alloy composition, detailed in Fig. 2E, shows that the stems contained a greater amount of Ni while the nodes were predominately Pd. The presence of Ni was also supported by the XPS analysis (Fig. S6), however, it could not be quantified accurately due to detection limits. HRTEM of a typical tadpole (Fig. 3) showed the lattice fringes of its head and tail (node) with a measured d spacing of 0.233 nm, attributed to (111) plane of face-centered cubic (fcc) Pd.

When the solution of these nanowire networks was aged for 24 h under ambient conditions, a shape transition from wormlike nanowire networks to 3D nanodendrites was observed, as shown in Fig. 4A. These nanodendrites had a bimodal distribution with a mean particle diameter modes centered at 30 nm and 95.6 nm (Fig. S7). HRTEM analysis, shown in the inset of Fig. 4A, confirmed the polycrystalline, porous nature of the nanodendrites with a d spacing of 0.233 nm, assigned to the (111) plane of Pd. EDX elemental mapping and line scans (Fig. 4B, C) revealed that the nanodendrites were composed of Pd with trace amounts of Ni, which was also supported by the XPS analysis (Fig. S6).

The Pd 3d core level of the core–shell, hollow, nanowire and nanodendrite morphologies supported on activated carbon are shown in Fig. 5. The nanosstructures were composed primarily of metallic Pd(0) with the Pd 3d₅/₂ and Pd 3d₃/₂ doublet located at a binding energy of 335.7 and 341.0 eV, respectively. For the core–shell, hollow and nanowire network structures, there was a small yet clearly observable positive shift of ~0.15 eV in the B.E. of the Pd 3d doublet associated with the presence of Ni. This higher binding energy can be attributed to the presence of Pd⁺⁺⁺ species generated by charge transfer from Pd to Ni [14,36]. A minor degree of
PdOx species was also indicated by the presence of a peak at a binding energy of 337.8 eV.

3.1. Mechanism

The distinct change in the morphology of the NiPd NPs with varying HCl concentration in the H2PdCl4 solution suggests the essential role of HCl in the system. To further investigate the role of HCl concentration, the synthesis was carried out at a lower HCl concentration using a H2PdCl4/1 mM HCl solution. This HCl concentration yielded predominately small diameter partially agglomerated Pd NPs with a small contribution of hollow NPs, as shown in Fig. S8A formed through a GRR [14]. No core–shell NPs was observed using 1 mM HCl. After aging for 24 h, no significant shape transition was observed (Fig. S8B).

When the synthesis was performed using Na2PdCl4 as the precursor without the addition of HCl (Fig. S8C), small diameter NPs were exclusively observed, indicating that a suitable HCl concentration in addition to pH was a critical factor to tailor the structure of bimetallic NPs. To understand the pH-dependent reaction pathways occurring at different HCl concentrations, it is necessary to take into consideration the strong pH-dependent factors such as the reduction potential of NaBH4 [37–39], Pd(II) complexes with chloride and/or hydroxide as the coordinating ligands [40–42], and the pH-dependent stabilizing ability of the trisodium citrate capping agent [43–45]. NaBH4 is typically regarded as a strong reducing agent, however, its reducing capacity is strongly dependent on the solution pH with a redox potential of $\Delta E_{0.48}$ V at pH 1 decreasing to $\Delta E_{1.24}$ V vs. SHE at pH 14, therefore its reducing power is enhanced by decreasing pH [38,39]. The deprotonation and effectiveness of the stabilizer, trisodium citrate, is also pH-dependent. The pKa values of citric acid are 3.2, 4.8, and 6.4, respectively, therefore when the pH is above the third pKa, the fully deprotonated C2H3− is the dominating form of citrate. Decreasing the pH results in protonation of citrate ions thereby lowering its stabilizing ability.

The NiPd and Pd nanostructures synthesized under different solution pH and aging time are schematically shown in Scheme 1. Ni NPs were firstly synthesized through the redox reaction of the Ni salt and NaBH4 forming a basic solution (pH = 9) due to the presence of NaBH4. Oxidation of Ni NPs by the atmospheric oxygen was inhibited by bubbling the solution with N2. The Pd precursor solution containing a certain concentration of HCl was injected into the Ni NPs and consequently, the pH of the solution gradually changes over the course of the reaction, as detailed in Table 1. Injection of the H2PdCl4 precursor into the Ni NPs solution gives rise to competing reactions involving Ni NPs or NaBH4, and the rate of these competing reactions is controlled by tuning the solution pH. Using the H2PdCl4/1 mM HCl solution, the GRR of the Ni NPs dominates (with a rate of $R_{gal}$) to generate the hollow nanostructure (Pathway A, Scheme 1).

Using the H2PdCl4/5 mM HCl solution, the lower pH gives rise to a faster reduction of Pd(II) by NaBH4 (with a rate of $R_{red}$). Galvanic replacement is quenched due to rapid reduction [28,29] and nucleation of Pd NPs which undergo deposition on the surface of the pre-existing Ni NPs generating core–shell NiPd nanostructures (Pathway B, Scheme 1). To investigate this growth mechanism, TEM analysis of the NPs formed at 15 min using different HCl concentrations showed that the core–shell structures were formed.
concentrations was evaluated (Fig. S9). Analysis of the H$_2$PdCl$_4$/1 mM HCl solution confirmed galvanic replacement occurring at an early stage in the reaction (Fig. S9(A, B)). In contrast, TEM analysis of the H$_2$PdCl$_4$/5 mM HCl solution after 15 min (Fig. S9C) was dominated by the formation of solid NPs, supporting that the higher HCl concentration affected the reaction kinetics and blocked the galvanic reaction. This inhibition of GRR is attributed to the template surface passivation discussed by Gilroy et al [46].

Increasing the HCl concentration further using the H$_2$PdCl$_4$/10-mM HCl resulted in the transition from nanoparticle to wormlike nanowire networks and indicated a significant change in the growth mechanism. Two features are important when understanding the growth mechanism of these structures. Firstly, EDX identified the variable alloy composition of the nanowires having NiPd alloy segments and Pd segments. The lower pH of the solution resulted in the rapid reduction of the Pd precursor due to the higher reducing capacity of NaBH$_4$ at lower pH. This rapid reduction resulted in the formation of NiPd alloy NPs with the Ni NP seeds and homogeneously generated Pd nuclei. These NPs are ther-

Fig. 2. (A), (B) TEM images of NiPd wormlike nanowire-tadpoles obtained from H$_2$PdCl$_4$/10 mM HCl after 3 h. The blue dashed cycles represent the nodes while the white rectangles represent the stems. (C) Elemental EDX mapping of tadpole [orange denotes Ni and green denotes Pd]. (D), (E) EDX line scan of a tadpole. Scale bar in (C), (D) is 10 nm. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Fig. 3. TEM image and HR-TEM image of a tadpole with the calculated lattice fringes for the head and tail (node).

Fig. 4. (A) TEM images of Pd nanodendrites obtained from H$_2$PdCl$_4$/10 mM HCl after aging for 24 h. (B) Elemental EDX mapping of nanodendrite [orange denotes Ni and green denotes Pd]. (C) EDX line scan of nanodendrite. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
modynamically unstable and to minimise the total surface free energy of the system, nanowire networks grow through aggregation of NiPd NPs undergoing coalescence with homogeneous nucleated Pd NPs giving a nanowire with variable chemical composition as observed by EDX analysis (Pathway C, Scheme 1). Protonation of citrate capping agents at high HCl concentration weakens electrostatic repulsion of the capping agent promoting the attachment of the NPs [47]. Preferential adsorption of citrate on Pd {1 1 1} facets may also play a role in directing the anisotropic growth. The fast formation of nanowires was confirmed by TEM analysis 15 min after the addition of the H2PdCl4 solution (Fig. S10A). In order to validate the pivotal role of Ni on the formation of the nanowire networks, the synthesis was performed in the absence of the Ni seeds. TEM analysis, shown in Fig. S10B, illustrates the formation of Pd nanodendrites verifying that the presence of Ni seeds is necessary to drive the 2D growth of the wormlike nanowire networks. Previous studies on semiconductors [48] and metal nanowires [49–51] suggested the formation of dipolar moments and van der Waals attraction forces within the NPs as the driving force for the alignment of the small nanoparticles into nanowires. The formation of chains due to dipole–dipole attraction is quite plausible due to the presence of both NiPd and homogeneously nucleated Pd NPs present in solution, giving rise to different reactivity on the crystallographic surfaces. The second important feature in the wormlike nanowire networks is the presence of the tadpole nanostructures, these structures likely form early in the reaction where more basic pH conditions favour the GRR. The presence of hollow and solid tadpole heads and short nanowire tails was observed by TEM analysis after 15 min (Fig. S10A). As the pH decreased with the further addition of the Pd precursor, the enhanced reduction capacity of NaBH4 led to the rapid reduction and nucleation of Pd NPs. The faster reduction kinetics coupled with the lower stabilizing power of the citrate resulted in the formation of nanowire networks through unidirectional aggregation.

In addition to the solution pH, the aging time was another essential parameter in influencing the shape evolution of the nanostructures. Upon aging the NiPd core–shell NPs, Cl− ions could...

Fig. 5. Pd 3d XPS spectra of the four different nanostructures supported on activated carbon.

Scheme 1. Schematic illustrating the synthesis of NiPd bimetallic NPs and the structure evolution relying on the solution pH and aging time via; (A) galvanic reaction route, (B) seeded growth route, and (C) crystallite coalescence route.
combine with $O_2$ from air to form an oxidative etchant for the Ni core resulting in the formation of hollow NPs [52]. In an effort to understand how the core–shell NiPd NPs evolved into hollow spheres, time-resolved TEM analysis was carried out at aging times of 6 h (Fig. S11A) and 12 h (Fig. S11B) confirming that the formation of the hollow structure is a stepwise process. Oxidative etching started with local pitting on the surface of the solid NPs followed by the gradual dissolution of the Ni core generating a hollow structure with a rough shell having a thickness of 6.5 ± 1.8 nm (Fig. S11C). As the Cl$^-$/$O_2$ oxidative pair had the ability to also etch Pd, longer aging times resulted in smoothing of the shell to a thickness of 2.5 ± 0.5 nm (Fig. S4B). The etching ability of the Cl$^-$/$O_2$ pair is influenced by the pH of the solution as the oxidative strength of $O_2$ is significantly enhanced under acidic conditions [53]. To further understand the role of the Cl$^-$ concentration, and pH in the etching mechanism of the core–shell NiPd NPs to form hollow NPs, the synthesis was carried out replacing HCl with NaCl, keeping the total Cl$^-$ concentration equal to that of the 5 mM HCl solution. TEM analysis (Fig. S12) showed limited etching of the core–shell NPs after 24 h of aging, indicating that a combined effect of Cl$^-$ ions and low pH was essential to achieve the shape transition from core–shell NPs to hollow NPs.

When using a Pd precursor solution with a higher HCl concentration, i.e. $H_2PdCl_4$/10 mM HCl solution, aging resulted in the evolution from wormlike nanowire networks to nanodendrites. As previously described, EDX analysis revealed that the composition of the nanowires consisted of NiPd alloy regions and Pd regions. The nanodendrites produced on aging were composed primarily of Pd with trace amounts of Ni detected by EDX and XPS analysis. The high density of defect sites at the particle–particle interface on the nanowires is susceptible to oxidative etching from Cl$^-$/$O_2$. Faster etching of the NiPd regions resulted in the breakdown of the extended nanowire network (Fig. S13, A, B). Pd NPs aggregation driven by the diffusion-controlled deposition and surface energy minimization results in the formation of nanodendrites [54–57].

We further investigated the use of other first-row transition metal seeds in the shape evolution of Pd-based bimetallic NPs. Co-Pd, Fe-Pd, and Cu-Pd systems were investigated by changing the metal precursors (see Experimental section for details). As shown in Figs. S14, S15, and S16, under low HCl concentration ($H_2PdCl_4$/5 mM HCl solution) core–shell structures could not be formed when Ni$^{2+}$ was substituted by equal molar amounts of Co$^{2+}$, Fe$^{3+}$, and Cu$^{2+}$. In comparison, using a higher HCl concentration ($H_2PdCl_4$/10 mM HCl solution) showed the same trend in the shape evolution of forming nanowires and nanodendrites on aging, thus this methodology appears to have wider applicability to other Pd-metal systems. The switch of metal from Ni to Co, Fe, and Cu involved changes to several parameters, including the redox potential of metal ions, the binding strength of the capping agent, and the degree of lattice mismatch between these metals and Pd.

### Table 1

| HCl Conc. in $H_2PdCl_4$ | Solution pH | Citrate ion form [44] |
|--------------------------|-------------|----------------------|
|                          | 1           | 2                    | 3                  | B/A                          |
| 1 mM HCl                 | 7           | 9                    | 8                  | Cl$^-$/Cl$^3^-$              |
| 5 mM HCl                 | 7           | 9                    | 6.5–7              | Cl$^-$/HCl$^2^-$/Cl$^3^-$   |
| 10 mM HCl                | 7           | 9                    | 6                  | Cl$^3^-$/HCl$^2^-$/HCl$^--$  |

1. The solution of NiCl$_2$ and TSC mixture.
2. The solution of NiCl$_2$, TSC, and NaBH$_4$ mixture.
3. The solution of NiCl$_2$, TSC, NaBH$_4$, and $H_2PdCl_4$ mixture.
B Before injection of $H_2PdCl_4$.
A After injection of $H_2PdCl_4$.

The catalytic efficiency of NPs is strongly dependent on their composition, size and morphology [58]. Hence, the catalytic performance of the obtained NiPd NPs with different structures was evaluated using hydrogenation of 4-nitrophenol (4-NP) to 4-aminophenol (4-AP) in the presence of NaBH$_4$ as a model reaction system. The four nanostructures, core–shell, hollow, nanowire networks and nanodendrites, were supported on acid-treated activated carbon (AC) with a Pd metal loading determined by ICP-OES as shown in Table 2.

The conversion of 4-nitrophenolate ion to 4-AP was monitored by UV–Vis spectroscopy through the disappearance of the 4-NP absorbance peak at 400 nm, which is accompanied by the gradual appearance of a new peak at 312 nm representing the formation of 4-AP [59,60]. Figs. 6, S17 exhibits the time-dependent UV–Vis absorption spectra assessing the catalytic activity of core–shell NPs, hollow NPs, nanowire networks and nanodendrites structures and commercial Pd/C for 4-nitrophenol reduction. The corresponding linear plots of normalized absorbance $-\ln(A_t/A_0)$ of the 4-nitrophenolate ions at 400 nm versus reaction time are also shown. With the exception of the nanodendrite catalyst, all the other catalyst morphologies displayed a short induction period (~30 s) associated with the adsorption of the reactant species on the catalyst surface. The reduction reaction obeyed pseudo-first-order kinetics with respect to the concentration of 4-NP as NaBH$_4$ was used in excess in the reaction. From eq.: $-\ln(A_t/A_0) = K_{app} t$, where $A_t$ and $A_0$ are the absorbance of the 4-nitrophenolate ions at 400 nm at time $t$ and at 0 sec, respectively. The value of the apparent rate constant ($K_{app}$) was obtained from the slopes of the best-fit lines (excluding the $t_{ind}$ region). A comparison of the catalytic performance of the obtained NiPd NPs with different structures was evaluated using hydrogenation of 4-nitrophenol (4-NP) to 4-aminophenol (4-AP) in the presence of NaBH$_4$ as a model reaction system. The four nanostructures, core–shell, hollow, nanowire networks and nanodendrites, were supported on acid-treated activated carbon (AC) with a Pd metal loading determined by ICP-OES as shown in Table 2.

### Table 2

| Conc. of HCl | Catalyst                | Pd content (X $10^{-3}$ mg/mg) | $K_{app}$ (X $10^{-3}$ s$^{-1}$) | Normalized rate constant $K_{norm}$ |
|-------------|-------------------------|---------------------------------|----------------------------------|-------------------------------------|
|             |                         | Pd                             |                                 | Pd-mg$^{-1}$ s$^{-1}$ | Pd-mmol$^{-1}$ s$^{-1}$         |
| 5 mM HCl    | Core-shell NPs/AC       | 6.63$^a$                        | 7.80                             | 1.1465                             | 125.201                          |
|             | Hollow NPs/AC           | 3.99$^a$                        | 6.27                             | 0.9950                             | 105.886                          |
| 10 mM HCl   | Nanowire networks/AC    | 21.12$^a$                       | 10.98                            | 0.7889                             | 81.824                           |
|             | Nanodendrites/AC        | 14.28$^a$                       | 9.28                             | 0.4975                             | 33.852                           |
|             | Commercial Pd/C         | 14.28$^a$                       | 9.28                             | 0.9950                             | 105.886                          |

$^a$ Pd content measured by ICP-OES.
reaction rates of the different nanostructures and commercial Pd/C is summarized in Table 2. The mass normalized rate constants of the nanostructures were all higher than that of commercial Pd/C. The core shell NPs displayed 6 times the activity demonstrating the increased utilization efficiency associated with the Pd located only on the shell. Similarly, the hollow NPs displayed over five times the mass normalized rate, attributed to increased surface area in the hollow interior.

Fig. 6. Representative time-dependent absorption spectra showing the reduction of 4-NP to 4-AP using the four nanostructures supported on AC and their corresponding normalized absorbance of the 4-NP at 400 nm versus reaction time.
For nanostructures prepared using the H$_2$PdCl$_4$/5 mM HCl solution, a comparison of the rate constants normalized to the Pd content shows that the core–shell structures displayed slightly better catalytic activity than that of the hollow structures, which can be attributed to the synergistic alloy effect and the lattice strain effect created between the core and shell [61,62]. In the case of using the H$_2$PdCl$_4$/10 mM HCl solution, the nanodendrites showed superior catalytic activity. The mass normalized rate coefficient of the nanodendrites was 2.4 times greater than that of the nanowire networks. Despite the excellent catalytic efficiency of nanowire networks previously reported [63,64] owing to their unique anisotropic structures, they exhibited the lowest catalytic performance in our study compared to other morphologies. The open structure of the nanowire networks and unfavourable interactions with the activated carbon support may hinder reactivity. All nanostructures catalysed the reaction to completion in very short reaction times. The existence of an electronic effect, confirmed by the observed shift in the B.E. of the Pd 3d doublet, promotes the adsorption of hydrogen onto the catalyst surface and facilitates the electron transfer from NaBH$_4$ to 4-NP [59,65]. Table S1 provides details on literature reports using this catalytic reaction.

4. Conclusions

In summary, we have achieved the shape-controlled synthesis of NiPd bimetallic NPs by simply tuning the HCl concentration in the H$_2$PdCl$_4$ solution and by the reaction aging time. While NaBH$_4$ is conventionally regarded as a strong reducing agent resulting in the rapid reduction of the metal precursor, we demonstrate that modulation of the pH can alter the reducing power of NaBH$_4$ allowing different NP morphologies to be easily synthesized under aqueous and ligand-free environment. The NiPd core–shell nanostructures could be assigned to quenching of the GRR between Ni NPs and H$_2$PdCl$_4$ through enhanced reduction power of NaBH$_4$. Simple aging of these NPs in solution results in the formation of hollow Pd NPs due to the oxidative etching of the Ni core. Conducting the reaction at a higher concentration of HCl gave rise to worm-like nanowire networks which extended for hundreds of nanometers. The formation of these 2D architectures formed at a lower pH was attributed to the enhanced reduction rate of the Pd precursor, insufficient capping of citrate ions and low surface charge of the NPs formed at the early stage. Subsequent aging of these nanowire networks results in the formation of Pd nanodendrites due to the oxidative etching of Ni, followed by the self-aggregation of small NPs through the diffusion-limited mechanism generating the dendritic structure. Therefore in this study, the concentration of HCl in the Pd precursor (H$_2$PdCl$_4$) exhibited a significant role in tailoring the structure of NiPd bimetallic NPs and provides new insights into the growth mechanism when using NaBH$_4$ as a reducing agent. The ability to rapidly synthesize nanostructures with diverse morphology and dimensionality is highly advantageous to catalytic and electrocatalytic applications.

CRediT authorship contribution statement

Mai Maize: Conceptualization, Methodology, Formal analysis, Investigation, Writing - original draft, Funding acquisition. Hanaa A. El-Boraey: Writing - review & editing. Mohamed I. Ayad: Writing - review & editing. Justin D. Holmes: Conceptualization, Writing - review & editing, Supervision, Visualization, Funding acquisition, Supervision. Gillian Collins: Conceptualization, Writing - review & editing, Visualization, Funding acquisition, Supervision.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary material

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