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
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Strain-Enhanced Metallic Intermixing in Shape-Controlled Multilayered Core–Shell Nanostructures: Toward Shaped Intermetallics

Abstract

Controlling the surface composition of shaped bimetallic nanoparticles could offer precise tunability of geometric and electronic surface structure for new nanocatalysts. To achieve this goal, a platform for studying the intermixing process in a shaped nanoparticle was designed, using multilayered Pd-Ni-Pt core–shell nanocubes as precursors. Under mild conditions, the intermixing between Ni and Pt could be tuned by changing layer thickness and number, triggering intermixing while preserving nanoparticle shape. Intermixing of the two metals is monitored using transmission electron microscopy. The surface structure evolution is characterized using electrochemical methanol oxidation. DFT calculations suggest that the low-temperature mixing is enhanced by shorter diffusion lengths and strain introduced by the layered structure. The platform and insights presented are an advance toward the realization of shape-controlled multimetallic nanoparticles tailored to each potential application.

Keywords

multilayered core–shell nanostructures, shaped intermetallic nanoparticles, strain-enhanced metallic intermixing

Disciplines

Engineering Physics | Materials Chemistry | Nanoscience and Nanotechnology

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Strain-Driven Metallic Intermixing in Shape-Controlled Multilayered Core-Shell Nanostructures: Toward Shaped Intermetallics

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Supporting Information Placeholder

ABSTRACT: Controlling the surface composition of shaped bimetallic nanoparticles could offer precise tunability of geometric and electronic surface structure for new nanocatalysts. To achieve this goal, we design a platform for studying the intermixing process in a shaped nanoparticle, using multilayered Pd-Ni-Pt core-shell nanocubes as precursors. We find that, under mild conditions, the intermixing between Ni and Pt could be tuned by changing layer thickness and number, triggering intermixing while preserving nanoparticle shape. Intermixing of the two metals is monitored using transmission electron microscopy. The surface structure evolution is characterized using electrochemical methanol oxidation. DFT calculations suggest that the low-temperature mixing is enhanced by strain introduced by the layered structure. The

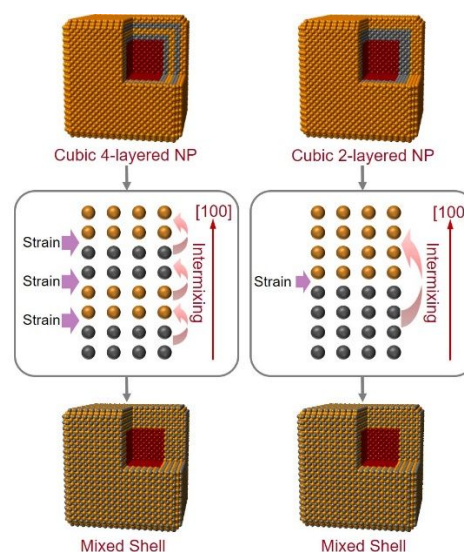
platform and insights presented here represent an advance toward the realization of shape-controlled multimetallic nanoparticles tailored to each potential application.

Nanomaterial design has increasingly focused on developing highly active and stable catalysts for chemical transformations¹⁻⁹. Platinum group metals are particularly active for a diverse set of reactions, but they are expensive and susceptible to surface poisoning. By incorporating nonprecious metals, catalyst cost can be lowered and performance can be improved¹⁰⁻¹⁶. The degree of intermixing between the metals has recently been shown to dramatically affect catalyst performance, but the mechanism of mixing is not well understood.^{2, 17-18} Thus, we aim to study the intermixing process and clarify its effect on catalysis.

Among alloys, the compositions that form intermetallic compounds (IMCs) have attracted recent attention because, in these cases, intermixing is thermodynamically favored. Generally, IMCs are crystalline alloys composed of two or more metallic/metalloid elements with specific stoichiometry and long-range order. More robust and resilient to etching¹⁹, these atomically ordered structures offer unique properties, including shape-memory²⁰, hydrogen storage²¹, and superconductivity²². Further, their tunable surface electronic structures²³⁻²⁵ allow their adsorption and desorption properties to be tailored to specific chemical reactions. In previous works, bulk IMCs have been extensively studied. More recent studies have focused on IMC nanoparticles (NPs) for their tailorable geometries, electronic surface structures and molecular adsorption properties²⁶⁻³⁴. The kinetics of solid-state intermixing to form IMCs, however, has received little attention.

We note three approaches to the formation of IMC NPs^{29, 35}. The first is a top-down synthesis method, where particles are formed by the mechanical alloying of mixed metallic powders through ball milling and annealing to form IMC phases³⁴. The size and shape of particles synthesized by these methods cannot be controlled, and no information can be gained on the nanoscale intermixing process. The second is direct, bottom-up synthesis by wet-chemistry methods. Shaped IMC NPs have been generated using this method³⁶⁻³⁷, but the need to precisely and simultaneously control the deposition rates of the chosen metals makes extending their scope and understanding their formation kinetics challenging. The third reported strategy involves synthesizing alloyed or heterostructured bimetallic NPs (e.g. core-shell or

dimer) by wet-chemistry and then converting these NPs to the intermetallic phase through intermixing triggered by thermal annealing at high temperatures³⁸. While promising, intermixing under these harsh conditions degrades the NP, generally through sintering and reshaping, preventing the study of intermixing in a specific geometry. The resulting indistinct particle shapes can further convolute catalytic results due to inconsistencies in size, shape/faceting, and surface composition, making the fundamental study of the catalyst surface or active sites difficult. In this work, we aim to optimize this third approach because we believe that intermixing, taking place in well-controlled nanoscale geometries, offers the best opportunity to understand the shape-dependent formation kinetics of IMC phases, which could, in turn, lead to a general synthesis scheme for shaped IMC NPs.



Scheme 1: Schematic representation of the conversion process from shaped Pd-Ni-Pt and Pd-(Ni-Pt)₂

nanoparticle precursors to Pd-Ni₃Pt core-mixed shell structures through atomic diffusion.

Since the solid-solid interdiffusion of metal atoms to form ordered intermetallic structures is the greatest contributor to the need for high annealing temperatures³⁹⁻⁴¹, we focus on lowering the effective diffusion barriers, which we hypothesize can be reduced by the use of strained metallic layers of variable thickness. Moreover, varying the layer thickness could allow us to test the effect of these variables on metallic mixing. Structures with a higher number of thinner layers (and thus higher strain and shorter diffusion lengths), we hypothesize, should show mixing at lower temperature than those with fewer, thicker layers. Previously, our group developed defined overgrowth methods for well-controlled core-shell NPs using mild aqueous phase systems^{12-13, 42}. During these syntheses, variables such as substrate size and shape, layer thickness and compositions, and layer number could be controlled. Notably, our Pd-Ni-Pt system¹³ stands out as an interesting candidate for the study of intermetallic formation, as the incorporation of Ni into Pt NPs has been shown to increase catalytic activity towards small molecule redox reactions^{10-11, 19, 43}. This well-defined platform serves as an ideal arena to study intermetallic formation, because this controlled, layered structure allows for a simple mixing process that can be easily characterized compared to other systems, such as alloy nanoparticles (Scheme 1). Herein, we focus on the PtNi₃ system as it has the highest loading of the nonprecious metal, Ni, relative to the precious metal, Pt. To maintain a bimetallic and more easily studied system, the core particle used to direct the shape of the NPs should be inert to the layer above it. For this reason, the Pd core was chosen; Pd and Ni are not

known to form stable IMC phases under mild annealing temperatures. This lack of a stable IMC phase suggests that there is no thermodynamic driving force for intermixing of Pd and Ni. We have performed DFT analysis to support this hypothesis, which will be discussed in detail below. Further, our experimental results show that the outer shell remains of a consistent thickness despite annealing, indicating limited diffusion between the core and shell. By intermixing the multilayered shell of defined core-shell structures, we can begin to construct a set of design parameters to be extended to other IMC NP structures of various shapes and compositions.

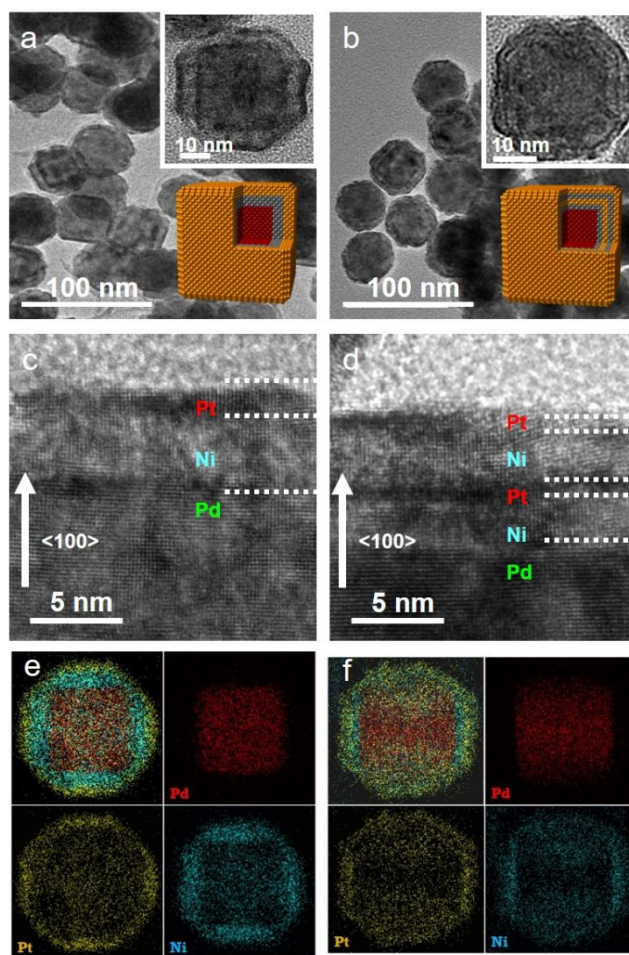


Figure 1: Transmission electron microscopy images of the as-synthesized (a) Pd-Ni-Pt samples at low magnification with the inset showing a single precursor particle and (b) Pd-(Ni-Pt)₂ samples at low magnification with the inset showing a single precursor particle. (c and d) show high magnification images of the Pd-Ni-Pt and Pd-(Ni-Pt)₂ particles, respectively, with the lighter Ni phase “sandwiched” between the darker Pd and Pt phases. (e and f) Energy dispersive x-ray mapping of the Pd, Ni, and Pt phases showing the Pd core and Ni and Pt surface layers.

Representative transmission electron microscopy (TEM) images of the Pd-Ni-Pt (a and c) and the Pd-(Ni-Pt)₂ (b and d) core-multilayered shell NP precursors can be seen in Figure 1. Mapping by energy dispersive x-ray spectroscopy (EDX) clearly shows Pd located in the core cube with Ni and Pt layered in the shell. Each layer shows epitaxial overgrowth, even with the high interfacial energy and strain associated with the metal boundary between the highly mismatched (~10%) lattices of Ni and Pt. This induced strain should contribute to a lower effective diffusion barrier facilitating metal-metal migration, particularly through the open (100) facet (which dominates the surface of our cubic NPs). Calculations for the expected thickness of the metallic layers were performed based on atomic ratios obtained through inductively coupled plasma optical emission spectroscopy (ICP-OES). By scanning electron microscopy (SEM), the base Pd cube is found to be ~30 nm in edge length (Figure S1) and, by TEM, the final layered structure, either two- or four-layered, is ~35 nm in edge length. Lattice constants of 352.4 pm and 392.42 pm were used for Ni and Pt, respectively. For the Pd-Ni-Pt particles, layers of 3.6 nm and 1.4 nm were found respectively for the

Ni and Pt (3.52 and 1.43 were measured by TEM). The calculation for the four-layered particles assumed that both shell layers were of equivalent thickness for each metal. Based on this assumption, thicknesses of 1.7 nm and 0.8 nm (1.78 nm and 0.77 nm were measured by TEM) were found for Ni and Pt, respectively. The experimental ratios were found to be Pd-Ni_{3.1}-Pt for the two-layered samples and Pd-(Ni_{1.3}-Pt_{0.5})₂ for the four-layered samples, as can be seen in Table S1. These values are close to target value of 3:1 and the expected values from the nominal concentration loading.

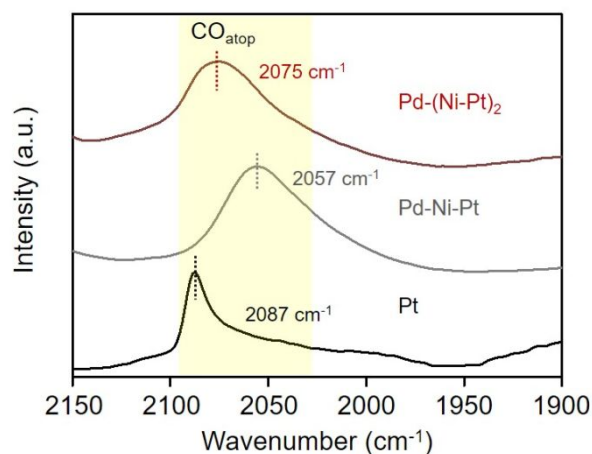


Figure 2: Normalized CO-DRIFTS spectra showing a shift in the energy of the characteristic C-O stretching peak. A larger shift in energy is seen for the thicker-layered Pd-Ni-Pt than the thinner-layered Pd-(Ni-Pt)₂.

CO-adsorbed monitored diffusion reflectance infrared spectroscopy (CO-DRIFTS) was used to monitor the initial structure of the NP surface (Figure 2)⁴⁴⁻⁴⁵. In each case, the NPs are supported on SiO₂. For pure Pt NPs, a characteristic C-O stretch is seen around 2087 cm⁻¹, corresponding to CO adsorbed on atop Pt⁴⁶⁻⁴⁷. When Ni is introduced below the Pt surface in the two-layered structure, a shift in the C-O peak to 2057 cm⁻¹ is seen.

This shift could be attributed to the ligand effect. Ni has a Fermi level energy of 9.46 eV while the Fermi level of Pt sits at 8.79 eV⁴⁸. As such, charge is transferred from the inner Ni layer to the outer Pt layer. This filling of the Pt d-band lowers its energy, which changes its interaction with CO and shifts the C-O stretch frequency.⁴⁹ By the same logic, we should observe a larger shift for the four-layered samples because the Pt layer is thinner and thus the surface Pt is closer to the Ni layer. However, the band of Pd-(Ni-Pt)₂ shifts only 15 cm⁻¹ from original Pt. This observation could be attributed to the effect of increased strain in the four-layered system. Compressive strain from Ni serves to shift the Pt d-band higher in energy, offsetting some of the shift induced by the ligand effect. This result suggests that Pd-(Ni-Pt)₂ shows increased strain compared with Pd-Ni-Pt.

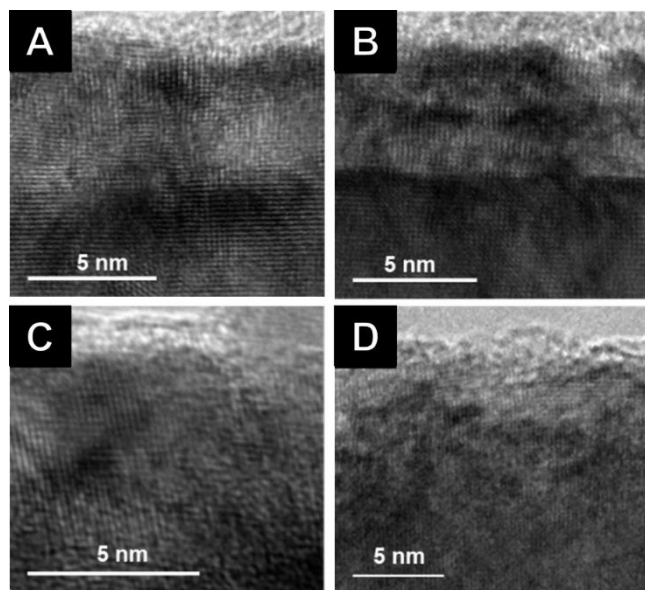


Figure 3: Representative high-resolution TEM images from which the quantitative lattice spacing data are calculated. (a and b) Pd-Ni-Pt and Pd-(Ni-Pt)₂, respectively, after annealing at 200 °C. Layering is still

visible by contrast in the image. (c and d) Pd-Ni-Pt and Pd-(Ni-Pt)₂ after annealing at 400 °C. One mixed layer can be seen for the Pd-(Ni-Pt)₂ sample.

To more quantitatively track the degree of mixing for each temperature, high-resolution TEM (HRTEM) images were taken to study the change in lattice parameter through the different layers of the NP. Figure 3 shows representative HRTEM images taken of each sample at three different temperatures in the [200] direction. For pristine Pd, Ni, and Pt, the d-spacing measurements in the [200] direction are 0.1945 nm, 0.1760 nm, 0.1960 nm, respectively. (XRD) With the migration of the metal atoms, the lattice spacing is expected to change due to the different lattice constants. The larger Pt lattice will compress due to epitaxy with and incorporation of Ni while Ni will expand. Once completely mixed and stable, the lattice measurements throughout the mixed shell should be the same (~0.1875 nm). Figure 4 shows a graphical representation of the changes in the lattice parameters from the Pt shell down to the Pd core. At 200 °C a clear difference can be seen between the mixing in the two structures. The four-layered, strained structure shows mixing initiated at the Ni/Pt interfaces, while clear, separated Ni and Pt layers are seen in the two-layered structure. At 300 °C, the four-layered structure begins to approach full mixing and the two-layered shows the beginning of it at the Ni/Pt interface. At 400 °C, the shell of Pd-(Ni-Pt)₂ is fully mixed, while Ni and Pt layers are still visible in Pd-Ni-Pt. Overall, this analysis reinforces our idea that increased strain in Pd-(Ni-Pt)₂ allows for mixing at lower temperatures.

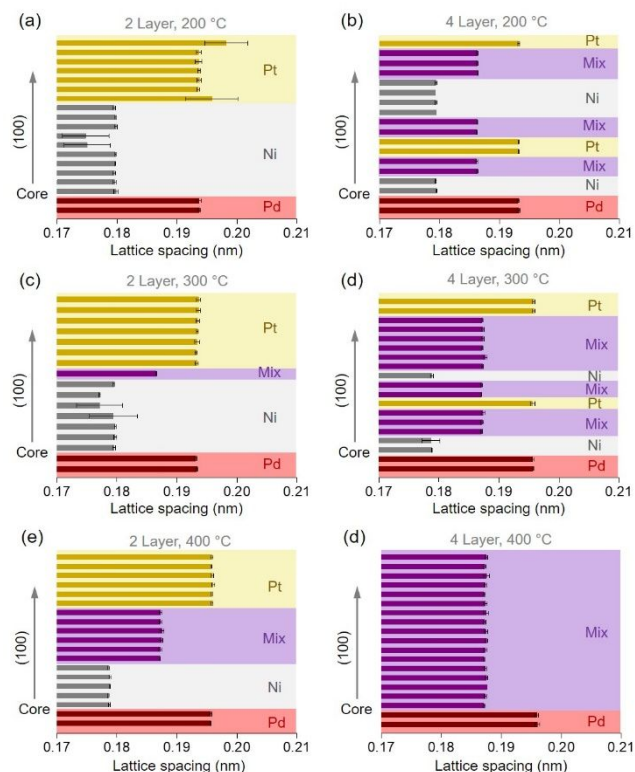


Figure 4. Lattice spacing measurement of Pd (red), Ni (grey), Pt (gold), and intermixed layered (purple). The intermixing is more pronounced at lower annealing temperatures for four-layered samples compared to the two-layered sample.

These quantitative data agree well with our post-annealing characterization by EDX (Figure 5). Mixing can be seen in the outer shell layers after annealing at 400 °C. Importantly, the Pd cube is clearly visible in the core of the mixed particle. This supports our hypothesis that Pd will not migrate into the shell layers at the mild temperatures chosen for our annealing.

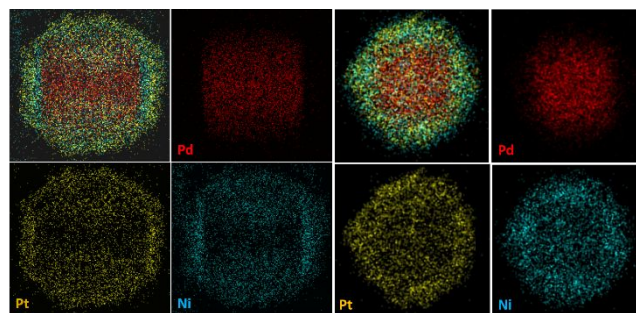


Figure 5. Energy dispersive x-ray images of Pd-(Ni-Pt)₂ (left) before and (right) after annealing, showing the core Pd cube and the mixing of the outer layers.

Next, we discuss theoretical insights into energetics relevant to this lower-temperature intermixing in this system using Density Functional Theory (DFT) analysis. Additional details and results are provided in the SI. As a preliminary analysis, using the PBEsol functional, the result support our hypothesis that intermixing of Ni and Pt to form Ni₃Pt is thermodynamically favored by 0.27 eV (per 3 Ni + 1 Pt), but intermixing of Ni and Pd to form Ni₃Pd is disfavored by 0.13 eV (per 3 Ni + 1 Pd). We have also checked that intermixing of Ni with Pt is favored, and with Pd is disfavored, for other stoichiometries (Table S2). Regarding the kinetics of intermixing, we first performed a benchmark analysis for the vacancy-mediated diffusion of Ni in bulk Pt. Here, a vacancy diffuses through Pt to a site adjacent to the Ni impurity, thereby allowing Ni to hop to that adjacent site. The effective barrier, E_{eff} , for such a process is the sum of a barrier for an atom hopping into an adjacent vacancy, E_{dv} , and a formation energy for creating the vacancy, E_{fv} .⁵⁰ Nuances are described below⁵¹. Here, we only report results from the PBEsol functional, which has been shown to be reliable for vacancy diffusion⁵². For the diffusion of Ni in Pt, the barrier for the Ni impurity to hop into an adjacent vacancy is found to be $E_{\text{dv}} = 0.93$

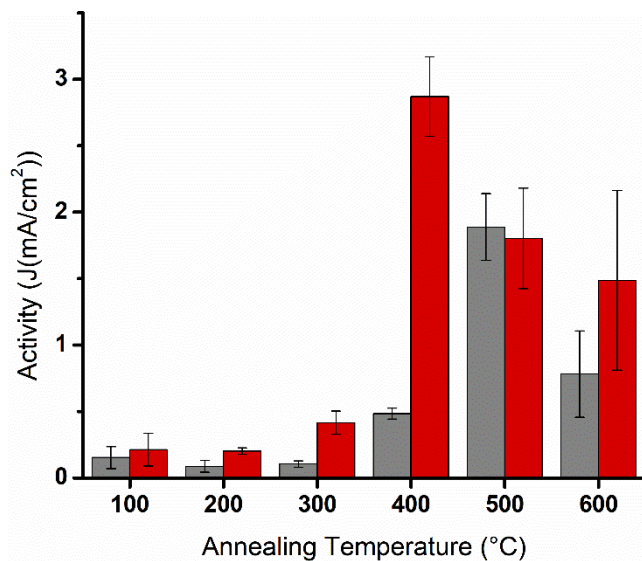
1 eV (versus $E_{dv} = 1.4$ eV for a Pt atom to hop into an
2 adjacent vacancy in Pt). Thus, diffusion of the vacancy
3 through Pt to reach a site adjacent to the Ni impurity
4 (rather than the hopping of Ni into the vacancy) is the
5 rate-controlling step. Consequently, the higher $E_{dv} =$
6 1.4 eV determines E_{eff} . The formation energy for a
7 vacancy next to a Ni impurity in Pt is $E_{fv} = 0.94$ eV
8 (versus $E_{fv} = 0.84$ eV for formation of a vacancy in pure
9 Pt), and the former higher value determines E_{eff} .
10 However, DFT underestimates E_{fv} due to so-called
11 “surface intrinsic error”, so, guided by previous analysis
12 for pure Pt⁵⁰, we use $E_{fv} = 1.2$ eV (rather than 0.94 eV)
13 to obtain $E_{eff} \approx 1.4 + 1.2 = 2.6$ eV for Ni diffusion in Pt.
14 The corresponding effective rate for hopping of the Ni
15 impurity, assuming an Arrhenius form with a typical
16 prefactor of $10^{13}/s$ is about $10^{-6.5}/s$ at 400 °C, too low
17 for effective intermixing on the experimental timescale.

18 However, motivated by the proposed importance of
19 strain effects in intermixing, we consider the effect of
20 compressive Pt strain, $\varepsilon < 0$, on E_{fv} , and E_{dv} . (Recall
21 that epitaxy at the Ni-Pt interface within the NP, as well
22 as Ni incorporation into Pt, should induce compressive
23 strain.) Significantly, we find a strong decrease in the
24 vacancy formation energy, E_{fv} , with increasing strain
25 magnitude, $|\varepsilon|$, for triaxial strain, and, in fact, E_{fv}
26 vanishes when $|\varepsilon|$ increases to only 2%. A weaker
27 decrease occurs for biaxial strain. On the other hand,
28 E_{dv} increases with $|\varepsilon|$ to 1.2 eV (1.7 eV) for Ni (Pt)
29 hopping into a vacancy where $|\varepsilon| = 2\%$ for triaxial strain
30 (with a lesser increase for biaxial strain). Assuming that
31 compressive strain makes E_{fv} negligible, then $E_{eff} \approx E_{dv}$
32 ≈ 1.6 eV. Then, the effective hop rate for the Ni impurity
33 is about $10/s$, which is sufficiently high for effective
34 intermixing.

Second, we briefly consider the vacancy-mediated
diffusion of Pt in bulk Ni. PBEsol predicts that $E_{dv} = 1.4$
eV (1.2 eV) for a Pt (Ni) atom to hop into an adjacent
vacancy in Ni. PBEsol also predicts that $E_{fv} = 1.5$ eV
for vacancy formation next to a Pt impurity in Ni and E_{fv}
 $= 1.6$ eV in pure Ni (which might be increased by ~ 0.2
eV accounting for surface intrinsic error). Thus, we
conclude that $E_{eff} \approx 1.4 + 1.8 = 3.2$ eV, too high for this
intermixing process to be effective. Additional studies
to explore the effect of tensile strain on Ni (induced by
epitaxy at the Ni-Pt interfaces and Pt incorporation in
Ni) reveal that E_{fv} actually increases with strain (at least
up to $\sim 6\%$). Although E_{dv} increases somewhat, there is
no net decrease in E_{eff} . Thus, we conclude that
intermixing in the Ni-Pt system is asymmetric and
associated with strain-enhanced diffusion of Ni into Pt,
rather than by Pt diffusion into Ni, which is in
agreement with previous experimental results⁵³.

To further investigate surface structure, each sample
was tested for the alkaline methanol electro-oxidation
reaction (MOR), a typical small molecule oxidation
reaction with potential in fuel cell applications.
Electrochemical measurements illuminate the surface
electronic structure of the catalyst by testing its ability
to perform desired transformations. Pt-Ni composites
have been shown to display high MOR activity, as
charge transfer brings the metal d-band to an optimal
energy level⁵⁴. The performance and durability of MOR
catalysts can be negatively impacted by the (often
irreversible) adsorption of a CO intermediate, but it has
been shown that intermetallic structures are less prone
to CO poisoning⁵⁵. They are also more resistant to
etching in acidic media, showcasing higher stability as
a result. Each sample was loaded on Vulcan XC-72
carbon and dried under nitrogen for 24 hours. The

1 electrochemical conversions were performed using a
2 standard three-electrode system: Pt wire as the
3 counter electrode, saturated calomel electrode as the
4 reference electrode, and a glassy carbon electrode
5 loaded with 5 μL of the synthesized nanocatalysts
6 supported on Vulcan carbon as the working electrode.
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28 Figure 6. Electrochemical activity for methanol
29 oxidation catalyzed by Pd-Ni-Pt and Pd-(Ni-Pt)₂ NPs
30 annealed at various temperatures. Grey bars represent
31 the two-layered Pd-Ni-Pt and red bars represent the
32 four-layered Pd-(Ni-Pt)₂.
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37 From Figure 6, a comparison of each sample (two- and
38 four-layered) at the designated temperatures can be
39 seen. At 200 °C, both samples perform nominally the
40 same as the as-synthesized samples. At 300 °C, each
41 sample has a slight increase in activity, but the four-
42 layered sample, with its higher lattice strain and thus
43 greater degree of mixing, does see a somewhat larger
44 increase. By 400 °C, as strain allows complete mixing
45 throughout the multilayered shell, the four-layered
46 sample has a significant jump in activity. This complete
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mixing can be seen in the lattice spacing
measurements (Figure 4), and the mixed shell yields a
more active surface. However, at the same
temperature, the two-layered sample has only a minor
increase in activity, on par with the four-layered sample
at 300 °C, as it is only partially mixed, which we
attribute to the lesser strain in its thicker layers. We
moved to higher temperatures to see if we could induce
the same degree of migration in the two-layered
sample as seen in the four-layered. By 500 °C, a larger
jump in activity is observed. The four-layered sample,
meanwhile, decreases in activity. This deactivation of
the four-layered catalyst is attributed to the degradation
of the sample (i.e. NP sintering) at such high annealing
temperatures. To test whether the two-layered
structure would follow suit at higher temperatures, an
annealing temperature of 600 °C was used; the same
dip in activity was observed. Each sample and
temperature profile was run in triplicate, leading to the
errors bars depicted. We again attribute the large error
bars at high temperatures to sample degradation at
these high temperatures, which is not a reproducible
process. Moreover, the change in the surface structure
also causes breaks in the consistency of the active
sites, which would also lead to a higher degree of error.

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37 The temperatures chosen for IMC synthesis beg two
38 questions: First, is the annealing time of one-hour ideal
39 for the mixing process? Perhaps a longer annealing
40 time at a lower temperature could prevent aggregation
41 and preserve fidelity to the initial cubic NP shape.
42 Second, can the Pd-Ni-Pt NPs achieve an activity
43 comparable to the Pd-(Ni-Pt)₂ NPs if given more time
44 to mix? These questions were addressed using
45 extended time studies, annealing the NPs for longer
46 times at each temperature, as shown in Figure S6.
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1 Annealing for longer than one hour at 200 °C appears
2 to have little effect on the catalytic activity, for both the
3 Pd-Ni-Pt and Pd-(Ni-Pt)₂ NPs, implying that the low
4 annealing temperature is unable to overcome the
5 kinetic barriers associated with mixing, irrespective of
6 the annealing time. This experiment agrees well with
7 our DFT results and supports the need to increase
8 annealing temperatures in order to induce metallic
9 mixing. The extended annealing at 400 °C shows two
10 distinct effects for the Pd-Ni-Pt and Pd-(Ni-Pt)₂ NPs.
11 For Pd-(Ni-Pt)₂, extending the annealing time to two
12 hours led to no change in activity, so further studies
13 were not carried out. This result makes sense, as the
14 outer shell is already fully mixed after 1 h. For Pd-Ni-
15 Pt, activity continues to increase up to a five-hour
16 annealing time. This result suggests that, given enough
17 time, the thicker Ni and Pt layers can achieve mixing
18 similar to the thinner layers in Pd-(Ni-Pt)₂. After more
19 than five hours, the activity decreases, which we
20 attribute to surface roughening and sintering after such
21 a long annealing process at high temperature.

22 In conclusion, we have used a variety of techniques to
23 investigate metallic mixing on the nanoscale. Through
24 TEM and HRTEM analysis, we have shown that atomic
25 diffusion is dependent on the thickness of the layers
26 that make up the mixing metals. Through CO-DRIFTS,
27 we have shown that our shell layers increase strain on
28 the outer Pt surface, which DFT calculations suggest
29 is essential for low-temperature metallic mixing.
30 Through MOR, we have shown that this mixing
31 changes the electronic structure of the active metal
32 surface, with clear differences again seen between the
33 layered Pd-Ni-Pt and Pd-(Ni-Pt)₂ nanostructures.
34 Overall, the insight gained represents an important
35 step toward the development of shape-controlled

bimetallic and IMC NPs. In the future, our platform can
be extended to more exotic crystal structures beyond
the *fcc*, d-block metals presented here, with an eye
toward the ultimate goal of generating a set of design
rules for the synthesis of nanocatalysts with custom-
made geometric and electronic surface structure.

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