Highly Active Carbon Supported Pd–Ag Nanofacets Catalysts for Hydrogen Production from HCOOH

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Supporting Information

ABSTRACT: Hydrogen is regarded as a future sustainable and clean energy carrier. Formic acid is a safe and sustainable hydrogen storage medium with many advantages, including high hydrogen content, nontoxicity, and low cost. In this work, a series of highly active catalysts for hydrogen production from formic acid are successfully synthesized by controllably depositing Pd onto Ag nanoplates with different Ag nano facets, such as Ag{111}, Ag{100}, and the nanofacet on hexagonal close packing Ag crystal (Ag{hcp}). Then, the Pd–Ag nanoplate catalysts are supported on Vulcan XC-72 carbon black to prevent the aggregation of the catalysts. The research reveals that the high activity is attributed to the formation of Pd–Ag alloy nanofacets, such as Pd−Ag{111}, Pd−Ag{100}, and Pd−Ag{hcp}. The activity order of these Pd-decorated Ag nanofacets is Pd−Ag{hcp} > Pd−Ag{111} > Pd−Ag{100}. Particularly, the activity of Pd−Ag{hcp} is up to an extremely high value, i.e., TOF{hcp} = 19 000 ± 1630 h⁻¹ at 90 °C (lower limit value), which is more than 800 times higher than our previous quasi-spherical Pd−Ag alloy nanocatalyst. The initial activity of Pd−Ag{hcp} even reaches (3.13 ± 0.19) × 10⁶ h⁻¹ at 90 °C. This research not only presents highly active catalysts for hydrogen generation but also shows that the facet on the hcp Ag crystal can act as a potentially highly active catalyst.

KEYWORDS: hydrogen production, formic acid, Pd, nanofacet, hexagonal close packing

1. INTRODUCTION

Hydrogen is regarded as a future sustainable and clean energy carrier. However, the production and storage of hydrogen still have some challenging problems, such as low efficiency, high operating temperature, huge volume and weight loading, and excessive CO content. To solve these problems, formic acid (FA) is used, being an outstanding hydrogen storage medium with many advantages, including high hydrogen content (4.4 wt %), nontoxicity, and low cost. To produce hydrogen from FA, it is necessary to dehydrogenate FA through

\[
\text{HCOOH} \rightarrow \text{H}_2 + \text{CO}_2 \quad \Delta G^\Theta = -48.4 \text{ kJ mol}^{-1}
\]  

A catalyst is always needed to dehydrogenate FA at low temperatures (<100 °C). Thus, both homogeneous and heterogeneous catalysts for FA dehydrogenation have been successfully developed. The homogeneous catalysts are mainly some coordination compounds with metal cores such as ruthenium, iridium, and rhodium. These homogeneous catalysts usually show very high catalytic activity. On the other hand, heterogeneous catalysts are also studied alongside the homogeneous ones. The antipoisoning heterogeneous catalysts are usually Pd-based or Au-based nanocatalysts, which can continuously catalyze FA dehydrogenation to H₂ with trace or even without CO. Such hydrogen could be directly fed into a proton exchange fuel cell (PEMFC) to achieve high performance. Recently, researchers have made great achievements with regard to heterogeneous catalysts for the FA dehydrogenation reaction. First, special nanostructures, such as ultrathin palladium nanoparticles and the core–shell structure, were used in an attempt to improve the activity of Pd-based catalysts. Xu et al. found that ultrathin palladium nanoparticles with a small size and a clean surface have a high TOF, up to 7256 h⁻¹ at 60 °C. Tsang et al. found that the Pd shell...
contains terrace sites and is electronically promoted by the Ag core, leading to significantly enhanced catalytic properties.  

Second, other metals or additives, such as Cu,18 Ag,18 Au,19 Co,20 Ni,21 B,22 MnO2,23 rare earth elements,13 and Schiff base,24 were added to Pd-based catalysts to improve the activity. These metals or additives usually have different functions in improving the activity. The addition of other metals, e.g., Cu,18 Ag,5 Au,13,19 Co,20 Ni,21 and B,22 could electronically promote Pd sites to significantly higher catalytic activity, as well as to a better tolerance toward CO poisoning via the synergetic effects between other metals and Pd.5,13,18,22 The addition of metal oxide, e.g., MnO2,23 and rare earth elements,13 could provide sacrificial CO anchoring sites, forming carbonate. As a result, Pd-based active sites remain available for the dehydrogenation of FA for an extended duration. PdAu-MnO2 nanoparticles can produce hydrogen through additive-free FA dehydrogenation, with previously unprecedented activity (TOF = 785 mol H2 mol catalyst−1 h−1 at room temperature).23 Zhang et al. invented a Schiff base modified gold catalyst for efficient H2 production from FA, with a TOF of 4368 h−1 at 50 °C.24 The addition of a Schiff base could provide a synergetic effect for C–H activation between the protonated Schiff base and electronegative gold nanoparticles at the interface and increase the FA dehydrogenation.  

Third, supports, such as ZIF-8-RGO,26 nanoporous carbon MSC-30,27 amine-functionalized SiO2,25 TiO2,28 reduced graphene oxide (RGO),29 and macroreticular basic resin,30,31 were also found to be very effective in improving the activity. These supports usually have very high specific surface area, which is advantageous in preparing well-dispersed ultrafine catalyst nanoparticles.26,27 So far, very highly active Pd-based catalysts were achieved by using the Pd/MSC-30 catalyst (2623 h−1 at 50 °C)27 and (Co90Ag10)Pd50/RGO (2739 h−1 at 50 °C).29 The supports can also have strong electron-donating effects on Pd and enhance the adsorption of formate to the catalyst and dehydrogenation from formate.28 The supports with an amine functional group play a positive role in the O–H bond cleavage of FA.32,35 A catalyst on basic resin showed a high reaction rate of more than 6000 mL min−1 g−1 (Pd), with a high TOF of 1900 h−1 based on Pd at 75 °C.30  

However, the facet, one of the most important factors in heterogeneous catalysis,31–34 has rarely been investigated in FA dehydrogenation.35 It is well-known that different facets on a catalyst may yield different reaction activity and selectivity.35,32 Recently, studies of different facets on a nanocrystal have attracted more and more attention in various catalytic reactions.31,33,34,36–40 Some of the studies are focused on facets with monometal or metal oxide.33,41 However, additional attention must be paid to facets with multimetals because such type of facets usually shows better activity and selectivity due to both a mediated electronic structure and surface atomic arrangement.5,42,43 Therefore, it is possible to further improve the activity of a heterogeneous catalyst by modulating the facet on the catalyst for FA dehydrogenation.  

Here, we report a series of highly active catalysts, which are synthesized by controllably depositing Pd onto different facets of Ag nanocrystals. Particularly, the activity of Pd-decorated facets on a hexagonal close packing Ag crystal (Ag{hcp}) reaches an extremely high value of 19 000 ± 1630 h−1 (turnover frequency (TOF)) at 90 °C, which is more than 800 times higher than that of the quasi-spherical Pd–Ag alloy nanocatalyst.

2. RESULTS AND DISCUSSION

2.1. Highly Active Catalysts with Pd-Decorated Ag Nanoplate for FA Dehydrogenation. To make the catalyst, Ag nanoplates with 88 ± 37 nm average edge length and 6.35 ± 1.00 nm average thickness (Figure S1) were prepared according to the literature.44 The Pd-decorated Ag nanoplate catalysts with different nPd:nAg were synthesized by reducing different amounts of PdCl2 with ascorbic acid onto an Ag nanoplate. Then, the nanoplate catalysts with multiple Pd–Ag nanofacets were supported on Vulcan XC-72 carbon black to prevent the aggregation of catalysts. Notably, Figure 1 shows that the Pd–Ag nanoplate catalysts exhibit extremely high activity for FA dehydrogenation at 90 °C. Figure 1a,b also shows that there are two activity peaks of Pd–Ag nanoplate catalysts at nPd:nAg = 8.33 × 10−4 and 0.33. These two activity peaks are, incredibly, up to 64 300 ± 4870 mL min−1 gPd−1 (or 9170 ± 690 h−1) at nPd:nAg = 8.33 × 10−4 and 4430 ± 40 mL min−1 gPd−1 (or 6570 ± 60 h−1) at nPd:nAg = 0.33, respectively (lower limit value is used; see SI.17 for details). Both of them are much higher than our previous report for the quasi-spherical Pd–Ag alloy nanocatalyst (160 mL min−1 gPd−1 or TOF = 22 h−1 at 92 °C).5

2.2. Discover the Highly Active Locations on Pd-Decorated Ag Nanoplate Catalysts. Previous studies show that only one activity peak appears at the optimum Pd content (∼50%) in the Pd–Ag alloy nanocatalyst for FA dehydrogenation.5,16 Either a too high or a too low Pd content will cause lower catalytic activity of a Pd–Ag catalyst. However, this research shows that there are two activity peaks with the
increase of Pd content in Figure 1. In this paper, because a Ag nanoplate has different facets, e.g., Ag{111} and Ag{100}, at different locations, we could suppose that the two activity peaks in Figure 1 originate from different locations of the Ag nanoplate. In this section, by tracking the Pd content at different locations of the nanoplate, we try to determine the highly active locations on Pd-decorated Ag nanoplate catalysts.

Figure 2a shows that the shape and contrast of Pd-decorated Ag nanoplates at the low ratio \( n_{\text{Pd}/n_{\text{Ag}}} = 8.33 \times 10^{-4} \) are similar to those of the as-synthesized Ag nanoplate in Figure S1. However, Figure 2d shows that the edges of the nanoplate have much higher Pd content compared to the mid part, indicating that most of the Pd deposits on the edges of the Ag nanoplate at a low ratio of \( n_{\text{Pd}/n_{\text{Ag}}} \). Moreover, the edges can reach an optimum Pd content (\(~50\%\)) while the mid part has a too low Pd content to have high activity at this moment. Therefore, the first activity peak at the low ratio \( n_{\text{Pd}/n_{\text{Ag}}} = 8.33 \times 10^{-4} \) should be attributed to the edges of the nanoplate.

At the high ratio of \( n_{\text{Pd}/n_{\text{Ag}} = 0.33} \), Figure 2b shows that the edges of the Pd–Ag nanoplate are whiter and thicker relative to the mid part, indicating that more Pd prefers to deposit on the edges. The shape and size of the Ag nanoplate can be fixed at the beginning of the synthesis of the PdAg/C nanoplate catalyst. Hence, Figure 2b shows that there is a thin frame around the nanoplate. The EDX measurement in Figure 2e shows that the edges of the Ag nanoplate indeed have higher Pd content compared to the mid part. However, the Pd content on the edges could be up to 90%, which is too high to show high activity. However, at such a ratio, the mid part has an optimum Pd content (\(~50\%\)). Therefore, the second activity peak at the high ratio of 0.33 in Figure 1 should be attributed to the mid part of the nanoplate.

At other ratios of \( n_{\text{Pd}/n_{\text{Ag}}} \), neither the edges nor the mid part has an optimum Pd content (\(~50\%\)) and the Pd–Ag nanoplate will show lower activity compared to the activity peaks. First, at a ratio \( n_{\text{Pd}/n_{\text{Ag}}} \) lower than \( 8.33 \times 10^{-4} \), the edges will have a Pd content lower than 50% and the mid part has a too low Pd content to have high activity. At such a condition, the activity decreases with decreasing ratio \( n_{\text{Pd}/n_{\text{Ag}}} \).

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Figure 2. Controllably deposit Pd on different locations of Ag nanoplate by controlling the mole ratio of Pd to Ag \( (n_{\text{Pd}/n_{\text{Ag}}}) \). (a–c) HADDF-STEM images of Pd–Ag nanoparticles at different \( n_{\text{Pd}/n_{\text{Ag}}} \). (d–f) EDX measurement along the red line of the nanoplate in the inset at different \( n_{\text{Pd}/n_{\text{Ag}}} \). The inset in (a) and (b) show the HRTEM images of the corresponding Pd–Ag nanoplate. More images are found in Figure S2.
Moreover, Figure 3 shows that the fcc and hcp layers can survive at a low ratio of \( n_{Pd} \cdot n_{Ag} = 8.33 \times 10^{-4} \). The hcp crystal structure can be clearly identified and measured in Figure 3b; the interface of hcp and fcc crystals is also clearly marked. Moreover, the area of the nanoplate edges cannot be omitted, although the Ag nanoparticles in this paper have 88 ± 37 nm average edge length and 6.35 ± 1.00 nm average thickness. The total area of the edges is up to ∼20% of the total area of the whole Ag nanoplate. Therefore, it is reasonable to assign the first peak activity at a low ratio of \( n_{Pd} \cdot n_{Ag} \) to the average activity of the three facets, including Pd−Ag{111}, Pd−Ag{100}, and Pd−Ag{hcp}.

Note that there are many difficulties in taking an HRTEM image of a vertically oriented Ag or Pd−Ag nanoplate. First, it is difficult to make Ag or Pd−Ag nanoparticles stand on their edges. Even if we increased the concentration of Ag or Pd−Ag nanoparticles, only a low percentage of nanoparticles will be vertically oriented on the copper grid for HRTEM measurement. Second, the Pd−Ag nanoplate should be in the right orientation during the HRTEM imaging. Otherwise, it is impossible to obtain a high-quality image with atom resolution. Third, it is difficult to obtain a high-quality image for both fcc and hcp crystals in one image because the orientations of fcc and hcp may be different in one nanoplate. Fourth, an hcp crystal is usually not perfect in an Ag or Pd−Ag nanoplate. There are possibly other defects at the interface between the fcc and hcp crystal or in the hcp crystal. All the above difficulties make it extremely hard to obtain a high-quality image for a vertically oriented Ag or Pd−Ag nanoplate. Through several attempts, we overcame the difficulties and took some high-quality HRTEM images for both Ag and Pd−Ag nanoparticles, as in the literature45 (Figure 3, Figures S3 and S4).

The second peak activity (6570 ± 60 h⁻¹) at a high ratio of \( n_{Pd} \cdot n_{Ag} = 0.33 \) should be attributed to the activity of the Pd−Ag{111} facet. From the previous section, we know that the second activity peak at a high ratio of 0.33 in Figure 1 should be attributed to the mid part of the nanoplate. The inset in Figure 2b clearly shows that the mid part is composed of a Pd−Ag{111} alloy facet because the lattice space is 0.230 nm, which is in the middle of Pd (0.220 nm) and Ag (0.240 nm). The following XRD measurements also support this result.

2.4. Track the Variation of the Crystal Structure at Different \( n_{Pd} \cdot n_{Ag} \). To track the variation of the crystal structure with the increase of \( n_{Pd} \cdot n_{Ag} \), a series of XRD measurements were done. Figure 4a shows that the XRD pattern of the Ag nanoplate (black curve) has a strong \{111\} peak and other weaker peaks, i.e., \{200\}, \{220\}, and \{311\} of the fcc Ag metal (JCPDS file: 04-0783). In addition to these, the typical peaks of the fcc Ag crystal, there is a weak, but clear, peak that appears at \( 2\theta \approx 35.9^\circ \) (black curve in Figure 4b), indicating the existence of the hcp Ag crystal structure in the Ag nanoplate. When the ratio of \( n_{Pd} \cdot n_{Ag} \) is increased, the hcp crystal can survive at a low ratio of \( n_{Pd} \cdot n_{Ag} \), i.e., \( 8.33 \times 10^{-4} \) (blue curve in Figure 4b), but will be totally destroyed at a high ratio of \( n_{Pd} \cdot n_{Ag} \), i.e., 0.33 (green curve in Figure 4b). This result is difficult to make Ag or Pd−Ag nanoparticles stand on their edges. Even if we increased the concentration of Ag or Pd−Ag nanoparticles, only a low percentage of nanoparticles will be vertically oriented on the copper grid for HRTEM measurement. Second, the Pd−Ag nanoplate should be in the right orientation during the HRTEM imaging. Otherwise, it is impossible to obtain a high-quality image with atom resolution. Third, it is difficult to obtain a high-quality image for both fcc and hcp crystals in one image because the orientations of fcc and hcp may be different in one nanoplate. Fourth, an hcp crystal is usually not perfect in an Ag or Pd−Ag nanoplate. There are possibly other defects at the interface between the fcc and hcp crystal or in the hcp crystal. All the above difficulties make it extremely hard to obtain a high-quality image for a vertically oriented Ag or Pd−Ag nanoplate. Through several attempts, we overcame the difficulties and took some high-quality HRTEM images for both Ag and Pd−Ag nanoparticles, as in the literature45 (Figure 3, Figures S3 and S4).

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is consistent with the observation of the HRTEM image in Figure 3 and Figure S4.

Figure 4c shows that the intensity ratios of \{hcp\} peak to \{111\} peak \(I_{\text{hcp}}:I_{\text{111}}\) remain at \(\approx 0.05\) at \(n_{\text{Pd}}:n_{\text{Ag}} < 0.01\), indicating that either the fcc or the hcp crystal structure of the Ag nanoplate does not have an obvious change at a low ratio \(n_{\text{Pd}}:n_{\text{Ag}} < 0.01\). With increasing \(n_{\text{Pd}}:n_{\text{Ag}} I_{\text{hcp}}:I_{\text{111}}\) increases and reaches a maximum value at the ratio of \(n_{\text{Pd}}:n_{\text{Ag}} = 0.083\), indicating that the fcc crystal structure is destroyed much faster than that of the hcp. This may be because the hcp layer is protected by the two fcc crystal layers, which sandwich the two sides of the hcp layer. When \(n_{\text{Pd}}:n_{\text{Ag}}\) is higher than 0.33, the hcp crystal structure will be totally destroyed and the ratio \(I_{\text{hcp}}:I_{\text{111}}\) will be zero in Figure 4c.

Figure 4a shows that 2\(\theta\) at {111} for the Pd–Ag nanoplate only has a single peak at the location between the standard peaks of pure Ag (2\(\theta_{\text{Ag}}\) = 38.1°), and Pd (2\(\theta_{\text{Ag}}\) = 40.1°), indicating that a Pd–Ag alloy forms when Pd is deposited onto the Ag nanoplate. Thus, the high catalytic activity is due to the synergetic effect of Pd and Ag.\(^\text{47}\) Moreover, Figure 4d shows that 2\(\theta\) at {111} for the Pd–Ag nanoplate increases as the ratio of \(n_{\text{Pd}}:n_{\text{Ag}}\) increases. Interestingly, the curve in Figure 4d obviously has two parts, which are separated at \(n_{\text{Pd}}:n_{\text{Ag}} = \approx 0.1\). This implies that the deposition of Pd on the triangular face (or body) of the Ag nanoplate dramatically happens from \(n_{\text{Pd}}:n_{\text{Ag}} = \approx 0.1\). Therefore, it is supported again that the activity originates from the edges at low \(n_{\text{Pd}}:n_{\text{Ag}}\) but from the mid part at high \(n_{\text{Pd}}:n_{\text{Ag}}\).

2.5. Activity of the Nanofacets Pd–Ag\{111\}, Pd–Ag\{100\}, and Pd–Ag\{hcp\}. Because the high activity \(\text{TOF}_{\text{111}} = 9170 \pm 690 \text{ h}^{-1}\) at 90 °C is the average activity of three nanofacets, including Pd–Ag\{111\}, Pd–Ag\{100\}, and Pd–Ag\{hcp\}, we consider which one of them dominates the high activity. Currently, we know that the activity on the Pd–Ag\{111\} facet is \(\text{TOF}_{\text{111}} = 6570 \pm 60 \text{ h}^{-1}\) at 90 °C, which is, however, much lower than \(\text{TOF}_{\text{111}} = 9170 \pm 690 \text{ h}^{-1}\) at 90 °C. Thus, the high activity should be mainly due to the Pd–Ag\{100\} or Pd–Ag\{hcp\} facet rather than the Pd–Ag\{111\} facet. However, it is difficult to directly synthesize the catalyst purely with the facet on the hcp Ag crystal due to the relative instability of the hcp Ag structure. However, it is much easier to synthesize the Pd–Ag nanowire catalyst mainly with the Pd–Ag\{100\} facet on its side wall.\(^\text{48}\) If we can obtain the activity \(\text{TOF}_{\text{100}}\) on the Pd–Ag\{100\} facet, then it is possible to know which one of the three facets is the most active one.

To obtain the activity of the Pd–Ag\{100\} nanofacet, we synthesized a series of Pd–Ag nanowire catalysts by depositing Pd onto the side wall of a silver nanowire through the same synthesis strategy as for the Pd–Ag nanoplate catalyst because the side wall of the Ag nanowire is composed of the \{100\} facet (Figure S8).\(^\text{48}\) Figure S11 shows that the maximum activity \(\text{TOF}_{\text{100}} = 4080 \pm 280 \text{ h}^{-1}\) at 90 °C of Pd–Ag nanowire catalysts appears at the ratio of \(n_{\text{Pd}}:n_{\text{Ag}} = 8.33 \times 10^{-2}\). However, it is also much lower than \(\text{TOF}_{\text{111}} = 9170 \pm 690 \text{ h}^{-1}\). Thus, the Pd–Ag\{hcp\} facet plays a crucial role in the high catalytic activity of \(\text{TOF}_{\text{111}} + \text{TOF}_{\text{100}} + \text{TOF}_{\text{hcp}}\) rather than the Pd–Ag\{111\} or Pd–Ag\{100\} facet.

To obtain the exact activity of Pd–Ag\{hcp\}, some calculations are required. Because \(\text{TOF}_{\text{111}} + \text{TOF}_{\text{100}} + \text{TOF}_{\text{hcp}}\) is the average activity of the three facets

\[
\begin{align*}
\text{TOF}_{\text{111}} + \text{TOF}_{\text{100}} + \text{TOF}_{\text{hcp}} &= a_1 \text{TOF}_{\text{111}} + a_2 \text{TOF}_{\text{100}} + a_3 \text{TOF}_{\text{hcp}} \\
&= a_1 + a_2 + a_3 = 1
\end{align*}
\]

where \(a_1\), \(a_2\), and \(a_3\) are the fractions of the thicknesses of the two fcc crystal layers and one hcp crystal layer, respectively, which are calculated using \(d_1 = t_1/d, d_2 = t_2/d, \) and \(d_3 = t_3/d\). The parameter \(d\) is the total thickness of the nanoplate, and \(d_1, d_2\), and \(d_3\) are the thicknesses of the two fcc crystal layers and one hcp crystal layer, respectively. The parameters \(d_1, d_2\), and \(d_3\) are also marked on the nanoplate in Figure 3a.

\[
d = d_1 + d_2 + d_3
\]

Then, the activity of Pd–Ag\{hcp\} is

\[
\text{TOF}_{\text{hcp}} = (\text{TOF}_{\text{111}} + \text{TOF}_{\text{100}} + \text{TOF}_{\text{hcp}}) - a_1 \text{TOF}_{\text{111}} - a_2 \text{TOF}_{\text{100}}/a_3
\]

According to our measurements (Figure S1) and the literature,\(^\text{45}\) the thickness of the hcp layer \(d_1 = 1.78 \pm 0.50 \text{ nm}\) and the thickness of the two fcc crystal layers \(d_2, d_3\) where \(d_1 = d_2d_3 = 2.28 \pm 0.25 \text{ nm}\). By using eq 5, we can determine the activity of Pd–Ag\{hcp\}, which is an extremely high value, i.e., \(\text{TOF}_{\text{hcp}} = 19000 \pm 1630 \text{ h}^{-1}\) at 90 °C. Therefore, Pd–Ag\{hcp\} dominates the high activity of \(\text{TOF}_{\text{111}} + \text{TOF}_{\text{100}} + \text{TOF}_{\text{hcp}}\).

In this work, the activity of Pd–Ag\{hcp\} reaches 9790 h\(^{-1}\) at 60 °C and 19 000 h\(^{-1}\) at 90 °C, which is similar to the results in the literature, e.g., 7256 h\(^{-1}\) at 60 °C\(^\text{13}\) and 2739 h\(^{-1}\) at 50 °C.\(^\text{29}\) Therefore, this paper shows that the Pd-decorated Ag nanofacet is a type of highly active catalyst for the FA dehydrogenation reaction.

It is well-known that the hcp structure is not a natural crystal structure for metal silver and is less stable than the fcc structure. Thus, it is extremely difficult to study the catalyst made by the hcp Ag crystal. However, our work here shows that the catalyst with the hcp crystal can be synthesized by carefully controlling the synthesis conditions. The activity of the catalyst can reach a plate after 15 min (Figure S13).

2.6. Reaction Kinetics of FA Decomposition on Different Nanofacets. Why can these nanofacets show such high catalytic activity? The analysis of the reaction kinetics of FA decomposition on these facets may give us some answers. The FA decomposition on the Pd–Ag catalyst is a very complex reaction process, which mainly includes FA dehydrogenation, poisoning, and the refreshment process of the catalyst, as follows:\(^\text{51}\)

\[
\text{HCOOH} + * \rightarrow * + \text{H}_2 + \text{CO}_2
\]

(6)

\[
\text{HCOOH} + * \rightarrow * - \text{CO} + \text{H}_2\text{O}
\]

(7)

\[
* - \text{CO} + k_\text{t} \rightarrow * + \text{CO}
\]

(8)

where \(k_\text{t}\) is the FA dehydrogenation rate constant, \(k_\text{t}\) is the poisoning rate constant, and \(k_\text{t}\) is the refreshment rate constant. We adopt the model from our previous work\(^\text{51}\) to describe the relationship between \(V\) and \(t\):
The C–H bond. This is one of the reasons for the high activity of the Pd–Ag(hcp) facet.

The poisoning step mainly relates to the breakage of C–H and C–OH bonds in eq 7. Because Pd–Ag(hcp) is extremely active in activating the C–H bond, the rate-determining step of the poisoning step should be the activation of the C–OH bond. Moreover, the poisoning rate constant \( k_2 \) mainly reflects the ability of the catalyst in activating the C–OH bond. Although \( k_2 \) of Pd–Ag(hcp) is also the maximum one compared to other facets, it is only ~3 times higher than others. Thus, the poisoning step will not significantly affect the activity of the Pd–Ag(hcp) facet. Furthermore, the selectivity, i.e., \( k_1/k_2 \), of Pd–Ag(hcp) is up to 2070, which is also much higher than others.

The strength of the Pd–C bond determines the dissociation rate of CO\(_{ad}\) in eq 8. However, the refreshment rate constant \( k_3 \) of Pd–Ag(hcp) is similar to that of Pd–Ag(100). Thus, the strength of the Pd–C bond on the Pd–Ag(hcp) facet is not so high and CO\(_{ad}\) can dissociate at a relatively high rate. More active surface will be exposed for catalysis through the refreshment process. This may be another reason for the high activity of the Pd–Ag(hcp) facet.

2.7. More Discussion about the Relationship between Activity and Nanofacets. Recently, the studies of different facets on a nanocrystal have attracted more and more attention in various catalytic reactions\(^{31,33,34,36,40}\) because the crystal facet is one of the most important factors in the activity and selectivity in heterogeneous catalysis.\(^{31–33}\) The atomic arrangement of the surface atoms can strongly affect the binding affinity of reactants, intermediates, and products during catalysis. If the atomic arrangement can match the geometry of adsorbed molecules, the adsorption will be strong. As we all know, either too weak or too strong binding of reactants will induce lower reaction activity.

Pd-based catalysts have high activities due to synergistic effects between Pd and other metals.\(^{28,47,51}\) Pure Pd is active for formic acid decomposition, whereas pure Ag has no or only very weak activity. Previous studies showed that either too high or too low Pd content will cause lower catalytic activity of the Pd–Ag catalyst, and an activity peak will appear at the optimum Pd content (~50%) in the Pd–Ag alloy nanocatalyst for FA dehydrogenation.\(^{51,56}\) Moreover, other metals or additives, such as Cu,\(^{13,19}\) Au,\(^{15,19}\) Co,\(^{20}\) Ni,\(^{21}\) B,\(^{22}\) MnO\(_2,\) rare earth elements,\(^{23}\) basic resin,\(^{29}\) and Schiff base,\(^{27}\) were added into Pd-based catalysts to improve the activity. However, Ag has not been found to have high activity when it is combined with other metals and additives. Thus, Pd atoms are the crucial element for the active sites in the Pd-based catalysts. The activity of a Pd-based active site is greatly improved when other metals and additives are involved. Some studies show that the drastic dehydrogenation activity enhancement is likely caused by their small size and the synergistic effect between Ag and Pd in the alloy structure, which inhibits the adsorption of CO on Pd.\(^{47}\) The synergetic effects may be due to the electronic effect, which has been studied via X-ray photoelectron spectroscopy (XPS).\(^{15}\) Then, the electronic structure, especially the d-band center position of the surface atoms, is modified.

In this paper, we controllably deposit Pd onto different facets of Ag nanocrystals to form Pd-based active sites. The synergetic effects between Pd and Ag clearly induce the activity peaks at different values of \( n_{\text{Ag}}/n_{\text{Ag}} \) in Figure 1. Figure S21a shows that the Pd 3d spectra of the Pd–Ag nanoplate catalyst \((n_{\text{Ag}}/n_{\text{Ag}} = 0.33)\) could be fitted into four symmetric peaks, including the
existence of two states of Pd species. The Pd 3d_{5/2} peak at 335.1 eV and Pd 3d_{3/2} peak at 340.4 eV are attributed to the metallic Pd(0), while the binding energy peaks shown at 335.7 and 341.0 eV are attributed to 3d_{5/2} and 3d_{3/2} of oxidic Pd(II). The higher peak area of Pd(0) indicates that the metallic state Pd is the dominant species on the surface of the Pd−Ag nanoplate catalyst ($n_{Pd/PdAg} = 0.33$). Figure S21b shows that the 3d_{5/2} and 3d_{3/2} peaks of Ag appear at 368.1 and 374.1 eV, indicating that most of the Ag will be in oxidic state Ag(I). Figure S21c,d shows that the catalyst has no chlorine element but has abundant oxygen element. Therefore, the oxidic state Ag(I) is possibly a type of silver oxide. Figure S21e,f shows that the peak area of Pd(II) is much smaller than that of Pd(0), indicating that metallic Pd dominates on the surface of the Pd−Ag/C nanoplate catalyst ($n_{Pd/PdAg} = 8.33 \times 10^{-4}$). The content of Pd is so low that the XPS signal is noisy. Nevertheless, we can still identify the typical peaks for Pd 3d. Then, we tried to fit the Pd 3d spectra with four symmetric peaks. Figure S21f shows that the peak area of Pd(II) is much smaller than that of Pd(0), indicating that metallic Pd is the dominant species on the surface of the Pd−Ag/C nanoplate catalyst ($n_{Pd/PdAg} = 8.33 \times 10^{-4}$). The content of Pd is so low that the XPS signal is noisy.

The above discussion shows that the atomic arrangement of different facets and the electronic structure can affect the adsorption of molecules. Although this research cannot give the binding affinities of every species, Table 1 shows that the refreshment rate constant $k_r$ of Pd−Ag{hcp} is similar to that of Pd−Ag{100}, indicating that the strength of the Pd−C bond on the Pd−Ag{hcp} facet is similar to that of Pd−Ag{100}. Namely, the poisonous intermediate CO$_{ad}$ has a comparable binding affinity on both the Pd−Ag{hcp} facet and Pd−Ag{100}. Therefore, the comparable binding affinity of CO$_{ad}$ is one of the reasons for the extremely high activity of the Pd−Ag{hcp} facet.

As we all know, Pd and Ag classically have an fcc crystal. However, in this paper, the Pd−Ag alloy with an hcp crystal, i.e., Pd−Ag{hcp}, has been synthesized at room temperature because of the nanosize effect. Moreover, Pd−Ag{hcp} shows extremely high activity for formic acid dehydrogenation. This result indicates that the variation of crystal phase can strongly affect the catalytic activity of the nanocatalyst. There is rising interests in crystal phase-controlled synthesis, properties, and applications of noble metal nanomaterials. For example, the 4H/fcc Au@PdAg nanoribbons exhibit much higher electrocatalytic activity toward the hydrogen evolution reaction compared to that of Pd black, which is even quite close to that of the commercial Pt black. Therefore, this research again shows that the activity and selectivity can be greatly modified by finely controlling the crystal phase of the nanocatalyst.

3. CONCLUSION

Pd was successfully deposited onto different nanofacets of triangular-shaped Ag nanocrystals. Vulcan XC-72 carbon black was used to prevent the aggregation of Pd−Ag nanofacets catalysts. The Pd-decorated Ag nanofacets exhibit very high activity in FA dehydrogenation, with an order Pd−Ag{hcp} > Pd−Ag{111} > Pd−Ag{100} at the optimum Pd content (~50%). Particularly, the activity of Pd−Ag{hcp} is up to an extremely high value, i.e., TOF$_{hcp} = 19\,000 \pm 1630\,\text{h}^{-1}$ at 90°C, which is much higher than the activity of other Pd-based catalysts. The initial activity of Pd−Ag{hcp} even reaches (3.13 ± 0.19) × 10^6 h$^{-1}$ at 90°C. This research not only presents highly active catalysts for hydrogen generation but also shows that the nanofacet on the hcp Ag crystal is a potentially highly active catalyst.

4. EXPERIMENTAL SECTION

4.1. Materials and Characterizations. All commercial materials were used as received unless specified. All aqueous solutions were prepared using deionized water in air. X-ray diffraction (XRD) patterns for the catalysts were recorded using a Bruker D8 X-ray diffractometer with the Cu−Kα ($\lambda = 1.5406 \text{ Å}$) radiation source operating at 40 kV and 40 mA. The nanostructure and composition of silver nanoflakes, Ag nanowires, and Pd−Ag catalysts were characterized by TEM, high-resolution TEM (HRTEM), high-angle annular dark-field scanning transmission electron microscopy, (HADDF-STEM) and EDX (EDAS) (200 kV, Tecnai G2 F20 S-TWIN; FEI) at CAS-Platform for Characterization & Test in the Suzhou Institute of Nanotech and Nanobionics.

4.2. Synthesis of Ag Nanoflakes. In a typical synthesis of silver nanoflakes, an aqueous solution of silver nitrate (Aladdin, 99.8%) (0.1 M, 4 mL) and trisodium citrate (Sinopharm Chemical Reagent Co., Ltd.) (75 mM, 80 mL) were added into 1920 mL of pure water with vigorous stirring at room temperature. Then, sodium borohydride (Sinopharm Chemical Reagent Co., Ltd., 96%) (NaBH$_4$, 0.1 M, 40 mL) was rapidly injected into the mixture, followed by injecting H$_2$O$_2$ (Sinopharm Chemical Reagent Co., Ltd.) (30 wt %, 14 mL) drop by drop. In the next several minutes, the solution color changed from yellow to blue, indicating that the morphology changed from sphere to nanoplate. The resulting solution was kept stirring for 4 h at room temperature.

4.3. Synthesis of Ag Nanowires. Silver nanowires were synthesized by the method presented by Xia et al. Typically, 0.1 mol of NaCl (Enox, China) and 2.24 g of poly(vinyl pyrrolidone) (PVP, MW = 1 300 000) (Alrich) were added to ethylene glycol (EG) (Sinopharm Chemical Reagent Co., Ltd.) and heated to obtain a clear solution. AgNO$_3$ solution (0.1 M, in EG) was added drop by drop to the solution with vigorous stirring for 10 min. The mixed solution was poured into a Teflon-lined stainless-steel autoclave, and was heated at 160°C for 6 h. The product was diluted with ethanol (Sinopharm Chemical Reagent Co., Ltd.) or water and centrifuged at 4000 rpm for 5 min several times to remove the surfactant. Finally, the purified nanowires were dispersed in water. The as-synthesized Ag nanowires with 50.1 ± 8.5 nm diameter and 6.3 ± 3.6 μm length were characterized by TEM, SEM, and optical dark-field microscopy.

4.4. Synthesis of Pd−Ag Nanoplate and Nanowire Catalysts. First, ascorbic acid (Anresco) (AA, 65 mM, 5 mL) as the reductant and 10 mM PdCl$_2$ (Nanjing Precious Metal Factory, China) as the Pd source were successively injected into 340 mL of Ag nanoflakes solution, which was heating at 40°C in an oil bath for 2 h. The amount of Ag nanoflakes was kept constant in the solutions while the amount of PdCl$_2$ was varied to achieve the different molar ratios of Pd to Ag ($n_{Pd/PdAg}$) from 4.17 $\times 10^{-4}$ to 1.0. Subsequently, the resulting solution was poured into 200 mL of the suspension ($V_{Vulcan}:V_{water} = 1:1$) with 100 mg of Vulcan XC-72 carbon black (Cabot Co.). The Pd−Ag nanoflakes were spaced by the carbon black to prevent aggregating. Finally, the suspension was filtered and washed with distilled water, and then dried in oven at 80°C overnight. The Pd−Ag nanowire catalysts were prepared with different molar ratios of $n_{Pd/PdAg}$ by using the same method as above.

4.5. Measurement of HCOOH Dehydrogenation Reaction. To measure the HCOOH (Sinopharm Chemical Reagent Co., Ltd., 98%) dehydrogenation reaction, 30 mg of catalyst was added into 5 mL of solution with 5 M formic acid and 2.5 M sodium formate. The reaction temperature was controlled at 90°C by a thermostat bath. The product gas was collected by a 100 mL injector, and was monitored by a camera.

4.6. Calculation of Catalytic Activity. This paper will mainly focus on the catalytic activity for the steady state after reaction for 10 min. The catalytic activity was calculated through two ways, including...
The volume rate per Pd weight ($V_{CO,45}$, mL min$^{-1}$ gPd$^{-1}$) and turnover frequency (TOF, h$^{-1}$). The volume is at the standard situation. The $V_{CO,45}$ was calculated by dividing the volume rate by the total weight of Pd in the reactor. The volume rate was calculated by dividing volume by the corresponding time. TOF was calculated according to the literature.$^{13}$

$$\text{TOF} = \frac{\text{number of produced hydrogen molecules}}{(\text{reaction time} \times \text{number of Pd atoms on the surface})}$$

(10)

In the above equation, the number of produced hydrogen molecules and reaction time are easy to obtain from experiment. However, the number of Pd atoms on the surface needs to be specifically calculated for different cases. On the basis of the TEM research, we found that Pd prefers depositing onto the edges of the Ag nanoplate at low ratio $n_{Pd}/n_{Ag} (<0.0417)$, while homogeneously covering the entire surface of the Ag nanoplate at high ratio $n_{Pd}/n_{Ag} (>0.0417)$. For the Ag nanowire, Figure S8 shows that Pd homogeneously covers the side wall of Ag nanowire according to TEM research. In addition, Pd–Ag alloy is formed on the nanoplates according to XRD research. Then, the number of Pd on the surface is calculated accordingly. Please refer to the Supporting Information (SI.17) for more detailed information.

4.7. Reaction Kinetics of FA Decomposition on Different Nanofacets. The reaction kinetics of FA decomposition on different facets was carried out according to the literature.$^{13}$ Some interesting kinetics parameters could be obtained by fitting the gas volume to time ($V$–$t$) curves for different nanofacets by the kinetics model in the literature.$^{13}$

In order to do the kinetics analysis, it is necessary to obtain $V$–$t$ curves for different nanofacets first. From experiment, we could directly obtain the $V$–$t$ curves for Pd–Ag{111} and Pd–Ag{100} from the Pd–Ag nanoplate at $n_{Pd}/n_{Ag} = 0.33$ and the Pd–Ag nanowire at $n_{Pd}/n_{Ag} = 8.33 \times 10^{-3}$, respectively. However, we need to do some calculations to obtain the $V$–$t$ curve for Pd–Ag(hcp).

From experiment, we could obtain the $V$–$t$ curve for the edge of the Pd–Ag nanoplate, which includes the facets Pd–Ag{111}, Pd–Ag{100}, and Pd–Ag(hcp). Therefore

$$V(t)_{(111)+{100}+{hcp}} = a_1V(t)_{(111)} + a_2V(t)_{(100)} + a_3V(t)_{(hcp)}$$

(11)

where $V(t)_{(111)}$, $V(t)_{(100)}$, $V(t)_{(hcp)}$, and $V(t)_{(111)+{100}+{hcp}}$ are the gas volume generated by Pd–Ag{111}, Pd–Ag{100}, Pd–Ag(hcp), and Pd–Ag{111} + Pd–Ag{100} + Pd–Ag(hcp), respectively, at the time $t$. Then

$$V(t)_{(hcp)} = (V(t)_{(111)+{100}+{hcp}} - a_1V(t)_{(111)} - a_2V(t)_{(100)})/a_3$$

(12)

By the above equation, $V(t)_{(hcp)}$ could be calculated, since $V(t)_{(111)}$, $V(t)_{(100)}$, and $V(t)_{(111)+{100}+{hcp}}$ are already known. Then, the kinetics model in the literature was used to fit the $V$–$t$ curves for different nanofacets to obtain some interesting kinetics parameters.$^{13}$ These parameters could help us understand the mechanism of the catalysis.

ASSOCIATED CONTENT

4 Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.6b08091.

Experimental details and additional results (PDF)

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Notes

The authors declare no competing financial interest.

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