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Silver-decorated Palladium on Carbon Catalyst for Enhanced Ammonium Formate Dehydrogenation

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Abstract:

Palladium (Pd)-based catalysts efficiently convert ammonium formate solution to hydrogen at low temperatures (< 100 °C), but they tend to deactivate quickly during stability testing. This manuscript presents a systematic investigation into the catalytic properties of Pd-Ag bimetallic catalysts, focusing on their surface compositions and exploring the mechanisms behind the deactivation of Pd/Ag-based catalysts. This study reports a carbon-supported Pd-Ag bimetallic nanoparticles (NPs) catalyst through a galvanic replacement method, which showed enhanced formate dehydrogenation performance. The best catalyst, Pd₃Ag₁₀/ACA-G (Pd-Ag bimetallic NPs with 3:10 mass ratio loaded on acid-washed activated carbon, prepared by galvanic replacement method), presents the highest activity with the TOF of 5202 h⁻¹ (~2.6-fold of commercial Pd/C). The advantages of smaller nanoparticles size with an enhanced

electron density of Pd-Ag bimetallic nanoparticles and the fact that the Ag surface alloy and Pd weakens the hydrogen adsorption energy and modulates the formate adsorption energy over the Ag/Pd (111) facet led to experimental higher turnover rates of hydrogen production. The changes on the catalyst surface, including surface Ag fraction decrease, NPs size growth, and O-containing species (carboxylate, etc.) adsorption, gradually resulted in the catalyst deactivation.

Keywords: Pd-Ag bimetallic catalyst; LOHCs; formate dehydrogenation; surface composition.

1. Introduction

Hydrogen (H₂) is considered a sustainable clean energy carrier that plays a vital role in the mitigation of greenhouse gas (GHGs) emissions¹⁻⁴. In recent years, many researchers investigated the chemical storage of hydrogen using formic acid/formate salts⁵⁻⁷. The formate salt solution is considered a promising liquid organic hydrogen carrier (LOHC) with the following advantages (i) It is non-volatile, non-corrosive, non-toxic, and provides a convenient reaction medium. (ii) As reported, the Gibbs free energy changes of hydrogen release from the formate salts is close to 0 ($\Delta_r G^0 \approx 1$ kJ/mole for formate salts, Na⁺, K⁺, and NH₄⁺), which makes the bicarbonate/formate-based LOHCs system thermodynamically favorable near ambient conditions⁸. (iii) Avoiding by-product CO poisoning^{9,10}. Even though formic acid and formate salt have a similar molecular structure, the dehydrogenation reaction pathway and mechanism over formic acid and formate salt solution are of difference. Dehydrogenation of formic acid suffers from poisonous by-product of CO, while trace of CO was detected in the ammonium formate dehydrogenation.

Efficient dehydrogenation of a formate solution requires appropriate catalysts to generate high-purity hydrogen. Based on our previous research and others' work, the carbon-supported Pd nanoparticles show high activities for ammonium formate dehydrogenation¹¹. And optimizing the support, such as removing O-functional groups¹² and introducing pyridinic/pyrrolic N species¹⁰ on carbon, is an efficient approach to improve the Pd/C catalysts' activity for formate dehydrogenation. Besides, developing Pd-based bimetallic catalysts (Pd-Au¹³⁻¹⁵, Pd-Ag¹⁶⁻¹⁸, Pd-Ni¹⁹, etc.) had also been investigated to enhance formate dehydrogenation reaction kinetics, among which the Pd-Ag bimetallic catalysts exhibited outstanding catalytic performance. For instance, Zhang et al. developed a composition-controlled synthesis of monodisperse 2.2 nm AgPd NPs which has high activity and stability.²⁰ Huang et al. reported a Pd-Ag alloy catalyst supported on the nitrogen-modified reduced graphene oxide (N-rGO) support for formic acid decomposition. The catalyst system was designed to make Pd electron-rich to enhance its utilization (The Pd and Ag atomic ratio was optimized, and the Pd₁Ag₆/N-rGO catalyst demonstrated a high TOF of 171 h⁻¹)²¹. In a similar work, Nabid et al. reported the synthesis of Ag@Pd catalyst supported on the nitrogen-modified graphene carbon nanotubes by employing Ag as a core and Pd as a shell. The catalyst was tested for formic acid decomposition, showing a TOF of 413 h⁻¹ at 298 K¹⁶.

Thus far, most of the reported work on Pd-Ag systems focused on tuning the Pd/Ag ratios to optimize the formate dehydrogenation performance. In addition, Pd-Ag bimetallic alloy with a specific structure may be highly active in catalyzing hydrogen release from a formate solution. Tsang et al. prepared an Ag-Pd core-shell nanocatalyst, Ag@Pd (1:1), showing a good hydrogen generation efficiency with TOF of 500 h⁻¹ at

70 °C²².

Among the several methods for the synthesis of bimetallic NPs, the galvanic replacement approach is particularly interesting due to advantages: it enables the synthesis of bimetallic hollow nanomaterials in a single reaction step with short reaction time²³; the surface composition and morphology can be controlled by adjusting the molar ratio between the sacrificial template and the guest metal ions²⁴. Galvanic replacement usually takes place between the sacrificial metal (low reduction potential) and guest metal ions (high reduction potential) in the solution²⁵. However, this approach can be optimized by introducing the adsorbed hydrogen atoms. A guest metal ion with a specific reduction potential higher than hydrogen but lower than sacrificial metal could form a thin film over the sacrificial metal template but not corrode it. Thus, it is promising to generate a bimetallic catalyst with core-shell structure.

In this work, we prepare the carbon-supported Pd-Ag bimetallic NPs catalysts through the optimized galvanic replacement and co-impregnation methods. We investigate the impact of various surface compositions on formate dehydrogenation while tuning the Pd/Ag ratios to reach the highest hydrogen formation yield. The results show that the catalyst prepared by galvanic replacement with a 3:10 Pd:Ag mass ratio (Pd₃Ag₁₀/ACA-G) displayed the best dehydrogenation performance with TOF of 5202 h⁻¹ at 80 °C and the lowest activation energy of 33.1 kJ/mol. The characterizations are conducted to reveal the advantages of the catalyst prepared by galvanic replacement and the possible deactivation reasons hindering the application of formate-based hydrogen carriers.

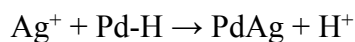
2. Experimental section

Materials: The commercial Norit GSX activated carbon powder (steam activated, acid-washed) (ACA) was purchased from Alfa Aesar (L11860); Sodium hydroxide ($\geq 98\%$), Ammonium formate ($\geq 99.995\%$), Silver nitrate ($\geq 99.0\%$) and Palladium(II) acetate (98%) were purchased from Sigma Aldrich®.

Synthesis of Pd/ACA. 250 mg active carbon washed by acid (ACA) support was added into 20 mL acetone, sonicating to form a uniform suspension. Then dissolve 15.8 mg Pd (II) acetate (3 wt% of Pd loading) into 20 mL acetone and mix these two suspensions. Stir till all the acetone evaporates, and then add 20 mL DI water to make a uniform suspension again. 20 mL NaBH₄ solution (2.22 mg/mL) was added drop by drop into the previous suspension with continuous stirring and stirred for 1 h at room temperature. After the filtration, the solids were washed with DI water and freeze-dried overnight, labeled as Pd3/ACA.

Synthesis of Ag/ACA. With the similar wet impregnation method, 500 mg active carbon washed by acid (ACA) support was added into 20 mL DI water, sonicating to form a uniform suspension. Then dissolve 27.4 mg AgNO₃ (3 wt% of Ag loading) into 20 mL DI water and mix these two suspensions. Stir at room temperature for 3 h and then add dropwise 40 mL NaBH₄ solution (2.22 mg/mL) into the previous suspension with continuous stirring and stir for 1 h at room temperature. After the filtration, the solids were washed with DI water and freeze-dried overnight, labeled as Ag3/ACA.

Synthesis of Pd_xAg_y/ACA-G. Pd-Ag bimetallic catalysts were prepared according to the method described by Epron *et al.*²⁶, where Ag was deposited via the galvanic replacement reaction onto the pre-synthesized Pd/C catalyst under a reducing atmosphere according to the following reaction:



The as-prepared Pd/ACA catalyst was added into 100 mL DI water (previously degassed by hydrogen bubbling) and then continuously stirred in the hydrogen flow for 2.5 h (25 °C). After that, a solution of a well-defined amount of the Ag precursor was added and maintained in contact with the suspension under hydrogen flow for another 2 h. The Pd-Ag bimetallic catalyst denoted as PdxAg_yACA-G (x and y mean the mass ratio of Pd and Ag, respectively. G means the galvanic replacement method) was filtered, washed, and freeze-dried overnight. The schematic diagram is shown in **Figure 1**.

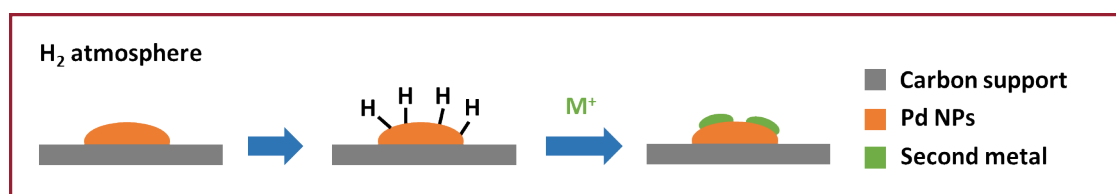


Figure 1. Schematic diagram of the synthesis of the Pd-based bimetallic catalysts via galvanic replacement reactions.

Synthesis of PdxAg_y/ACA-C. The Pd-Ag bimetallic catalysts were also prepared through wet impregnation and followed by co-reduction with the NaBH₄ aqueous solution labeled as PdxAg_y/ACA-C (x and y mean the mass ratio Pd and Ag, respectively. C means the co-reduction methods). Typically, Pd₃Ag₁₀/ACA-C was prepared by dispersing 500 mg ACA support into 40 mL DI water, then mixed with the Pd and Ag precursor aqueous solution (32.48 mg Pd(NO₃)₂ · xH₂O and 91.25 mg AgNO₃ dissolving into 40 mL DI water). After magnetic stirring for 3 h at room temperature, the NaBH₄ aqueous solutions (231.6 mg dissolved into 40 mL DI water) were dropped into the metal precursor and ACA support mixture. After 1 h stirring at room temperature, the Pd₃Ag₁₀/ACA-C particles were collected back through vacuum

filtration, 3 times DI water washing, and freeze-dried overnight.

Characterizations. XPS measurements were performed using a Thermo Fisher NEXSA spectrometer with a 125 mm mean radius, full 180° hemispherical analyzer, and 128-channel detector. This system uses a focused monochromatic Al K α X-ray (1486.7 eV) source for excitation and an electron emission angle of 60 degrees. The narrow scan spectra were collected using a passenergy of 50 eV with a step size of 0.1 eV. For the Ag 3d5/2 line, these conditions produced an FWHM of 0.84 eV \pm 0.02 eV. The binding energy (BE) scale is calibrated using the Cu 2p3/2 feature at 932.62 \pm 0.05 eV and Au 4f7/2 at 83.96 \pm 0.05 eV. The morphology and structure of the samples were analyzed by FEI Quanta 200F field emission scanning electron microscope (SEM), JEM-2100F transmission electron microscopy (TEM) and Tecnai G2 F20 S-Twin High-revolution transmission electron microscopy (HRTEM). A Micromeritics Auto Chem II system measured the CO pulse chemisorption and temperature program decomposition. The powder X-ray diffraction (XRD) was carried out on a Rigaku Miniflex II X-ray diffractometer with 40 kV and 15 mA for CuK α (λ =0.154178 nm), with a scan range between 5~90° at the speed of 4° min⁻¹. The ATR-FTIR analysis was performed on the Bruker Tensor II spectrometer with Harrick Horizon Multiple Reflection ATR Accessories and a Harrick temperature controller ATC-024-4. A 2.89 mL liquid sampling cell with a ZnSe horizon trough (HON-LSP-J) was used to carry out the catalytic reactions. The *Operando* ¹³C solid-state magic angle spinning (MAS) nuclear magnetic resonance (NMR) experiments were carried out using a Varian Inova wide-bore 300 MHz NMR spectrometer equipped with a 7.5 mm commercial ceramic probe. Briefly, 53 mg of catalyst and 200 μ L 1M ¹³C labeled sodium formate were loaded into a homemade *in situ* rotor capable of completely sealing a mixture containing solid, liquid and gaseous phases at elevated temperature and pressure^{27,28}. The Larmor

frequency is 75.42 MHz and a $\pi/4$ pulse width of 3 μs with a 4 s recycle delay was used in ^{13}C single pulse experiments. As for ^1H - ^{13}C cross-polarization, a $\pi/2$ pulse width of 8 μs for ^1H with a 4 s recycle delay and a contact time of 1.5 ms was applied. The sample temperature was calibrated using ethylene glycol as previously reported²⁹.

Dehydrogenation performance evaluation. The formate decomposition reaction experiments were carried out in a 100 mL Parr reactor. The gas outlet was connected to a NaOH solution trap (7 M). The trap was connected to an H_2 digital flowmeter. The NaOH trap is used to absorb CO_2 generated from the bicarbonate decomposition. Before the reaction, the system was charged with N_2 gas 3 times to ensure no O_2 was in the reaction system. After the set reaction temperature was reached, the released gas was passed through the NaOH trap. Its volume was monitored using the H_2 digital flowmeter (M-500SCCM-D-DB9M/CM from Alicat Scientific, Inc.). The catalytic decomposition reaction for the release of hydrogen was initiated by stirring the mixture of the aqueous suspension of catalyst (0.05g) in 20 ml NH_4HCO_2 solution (1 M) at 80 °C after N_2 purging. After the dehydrogenation reaction, the spent catalyst was collected back by filtration and washed with DI water 3 times followed by freeze drying for overnight. The stability test was carried out using spent catalyst in subsequent cycles with the same process. Hot filtration experiments were conducted with the same procedure as reported before¹².

Computational Methods. All DFT based simulations were performed using Vienna ab initio Simulation Packages (VASP) code^{30–33} with the RPBE functional³⁴. Projector-augmented wave formalism^{35,36} wave potentials were used for core-valence treatment. The plane-wave cutoff energy was set to 400 eV and the Brillouin zone sampling was performed with a Monkhorst-Pack k-point grid of $4\times 4\times 1$. The Methfessel-Paxton scheme was used to smear the Fermi level with a Gaussian width (σ) of 0.1 eV for the

surface. Geometry optimizations were conducted with a self-consistent electronic convergence limit of 1×10^{-6} eV and an ionic convergence limit of 0.05 eV \AA^{-1} for unconstrained atoms. DFT calculations were performed for all geometry relaxations except for computing the gas phase reference energy of hydrogen and formic acid. Periodic surface slabs of Pd(111) and Ag(111) were constructed using a 3×3 supercell with a thickness of four layers. For the Ag on Pd surface alloy (termed Ag/Pd) construction, the top layer of the $3 \times 3 \times 4$ Pd(111) surface slab was replaced with Ag atoms. The bottom two layers of all surface slabs were held fixed at the optimized bulk lattice constant, while the top two layers were allowed to relax until convergence. Stable configuration site sampling of surface -adsorbed formate and hydrogen was performed on the surfaces. The adsorption energies of hydrogen and formate were calculated on the Pd, Ag, and Ag/Pd surfaces by evaluating the binding strength of hydrogen and formate on several high-symmetry binding sites. Hydrogen adsorption was evaluated on the three-fold face-centered cubic (FCC) and hexagonal-close-packed (HCP) hollow sites. Formate adsorption was computed with a bidentate configuration of formate wherein the formate oxygen was bound to neighboring on-top sites of Pd (Ag). Hydrogen adsorption was referenced to 0.5 H_2 gas. Formate ion adsorption was computed using both a thermodynamic cycle method as previously described¹² and additionally using an implicit solvent constant-potential method described below.

To mimic the effect of solution at the catalyst-liquid interface, the VASPsol implicit solvation method^{37,38} was utilized to describe the solvated interface with a polarizable continuum model that treats the ions in solutions using a linearized Poisson-Boltzmann (LPB) distribution. A dielectric constant of 80 and a Debye length of 3 \AA were used, corresponding to a 1 M monovalent solution, neglecting the cavitation energy contribution. The potential of zero charge (PZC) for each surface model was computed

using a constant potential calculation based on the methodology described in Duan's work³⁹, wherein the Fermi level was set relative to the vacuum level. The Fermi level corresponds to an applied potential value vs. the computational standard hydrogen electrode (SHE). When the net charge in the simulation cell was zero (surface charge is zero), the applied potential value obtained corresponds to the potential of zero charge (U_{PZC}). Adsorption energies of formate and hydrogen were recomputed at the U_{PZC} value of each surface composition.

3. Results and discussions

3.1. Characterizations

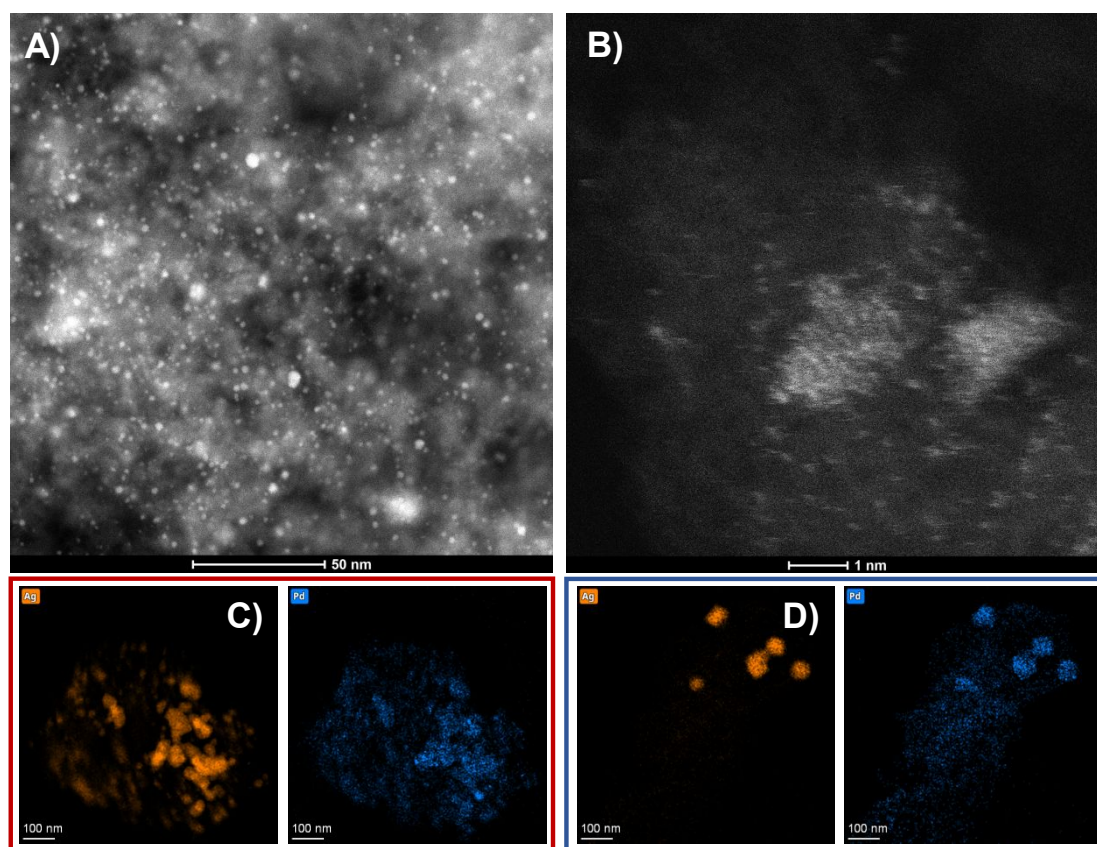


Figure 2. TEM images of Pd₃Ag₁₀/ACA catalysts, A-C) Pd₃Ag₁₀/ACA-G; D)

Pd₃Ag₁₀/ACA-C (Orange are Ag atoms and Blue are Pd atoms).

HRTEM and HAADF-STEM-mapping investigations examined the morphology, particle sizes, and composition of the Pd₃Ag₁₀/ACA-G and Pd₃Ag₁₀/ACA-C catalysts. The HRTEM images of Pd₃Ag₁₀/ACA-G shown in **Figures 2A** and **2B** reveal the Pd-Ag nanoparticles are well-dispersed on the carbon support. The mean diameter of Pd₃Ag₁₀/ACA-G is ~1.62 nm, smaller than that of Pd₃Ag₁₀/ACA-C (~2.50 nm), shown in **Figure S1**. This may be due to different formation mechanisms. The surface-decorated Ag atoms follow the morphology of small Pd NPs and would not significantly change the as-synthesized particle size, while both Pd and Ag species are more easily reduced simultaneously to form larger NPs in the co-reduction process. **Figures 2C** and **2D** display the corresponding element mapping of Pd₃Ag₁₀/ACA-G and Pd₃Ag₁₀/ACA-C, respectively. The Pd and Ag elements are homogeneously dispersed on carbon support by a galvanic replacement approach, but they are loaded with large particle size through co-reduction methods. The high-angle annular dark-field images (HAADF) of Pd₃Ag₁₀/ACA-G and Pd₃Ag₁₀/ACA-C, are shown in **Figure S2**. The lattice distance of NPs in the Pd₃Ag₁₀/ACA-G and Pd₃Ag₁₀/ACA-C are 2.21 nm and 2.36 nm assigned to Pd(111) and Ag(111), respectively. The HAADF-STEM images of the Pd₅/ACA and Ag₃/ACA catalysts are shown in Figure S3. For Pd₅/ACA, the Pd nanoparticles were well-dispersed, and even single atoms were observed. However, the bimodal size distribution was exhibited: after statistical calculations, the small and large Pd particle sizes are within the range of 1-2.5 nm and 15-20 nm, respectively. In contrast, no single atoms were observed in the Ag₃/ACA catalyst. Moreover, the Ag nanoparticles were larger and exhibited a broader size distribution with the range of 10-100 nm.

The XRD patterns of the Pd₃/ACA, Ag₃/ACA, Pd₃Ag₁₀/ACA-G, and Pd₃Ag₁₀/ACA-C samples were recorded to examine the crystal structures and phases, shown in **Figure**

S4. No noticeable peaks related to metallic Pd species could be found in the XRD pattern of Pd3/ACA, suggesting Pd NPs are highly dispersed on the carbon support. The sharp peaks at 38.1°, 44.3°, and 64.4° correspond to metallic Ag species (JCPDS No. 04-0783). The Ag peaks can be observed in Ag3/ACA, Pd3Ag10/ACA-G, and Pd3Ag10/ACA-C samples, but the peak intensities of Ag3/ACA and Pd3Ag10/ACA-C samples are higher than Pd3Ag10/ACA-G. According to the mechanism of galvanic replacement synthesis method we used in this manuscript, the Ag atoms would deposit on Pd NPs surface, and replace the adsorbed H atoms to form a microstructure that Pd NPs surface decorated with Ag atoms. Therefore, the Pd-Ag alloy particle size should be very similar with the original small Pd particle size, which is too small to show a clear diffraction peak in the XRD pattern. The Ag (111) peak shown in the **Figure S4** is related to excessive Ag precursor, which is reduced to form metallic Ag NPs loaded on the carbon support.

The XPS analysis reveals that the Pd3Ag10/ACA-G and Pd3Ag10/ACA-C catalysts contain C, O, Pd, and Ag elements without other significant impurities shown in **Figure S5A**. **Table 1** summarizes surface element atomic fractions, indicating that the Pd3Ag10/ACA-G has a higher surface Ag concentration (6.00% vs. 2.31%) but lower Pd fractions (0.62% vs. 3.65%) compared with Pd3Ag10/ACA-C. The atomic ratio of Pd and Ag was 1.6 when we synthesized Pd3Ag10/ACA-C through the co-reduction method. However, the galvanic replacement efficiently decreases the surface Pd/Ag atomic ratio from 1.6 to 0.1. Thus, most Pd species were decorated with Ag atoms after galvanic replacement. Besides, the O atomic fractions of Pd3Ag10/ACA-C and Pd3Ag10/ACA-G are 5.74% and 6.25%, respectively.

In the Pd 3d spectra (**Figure 3A**) of Pd₃Ag₁₀/ACA-C and Pd₃Ag₁₀/ACA-G, two main peaks associated with metallic Pd(0) 3d_{5/2} and Pd(0) 3d_{3/2} are located at 335.9 eV and 341.1 eV, respectively. Two peaks at higher binding energies (337.6 eV and 342.6 eV) correspond to the Pd(II) species^{40,41}. The peaks at 344.3 eV in both samples are related to the satellite structure highly correlated to the electrons in the partially filled *d* orbitals⁴². The Pd(0)/Pd(II) ratios in both samples were calculated using the peak areas of Pd 3d_{5/2}. The Pd₃Ag₁₀/ACA-C has a higher Pd(0)/Pd(II) ratio of 2.62 compared with Pd₃Ag₁₀/ACA-G (1.15), suggesting Pd(II) ions consist of surface Pd species. In other words, the Ag atoms are favorable to deposited on the metallic Pd surface due to no adsorbed H on Pd(II) surface, which confirms the galvanic replacement mechanism (shown in **Figure 1**) that Ag(I) ions replace the adsorbed H over Pd surface to form Ag(0) atoms. Besides, the exact Pd(II) 3d_{5/2} binding energy of Pd₃Ag₁₀/ACA-G is 337.4 eV, lower than that of Pd₃Ag₁₀/ACA-C (342.6 eV). More electrons were transferred to Pd(II), resulting in Pd(II) species of Pd₃Ag₁₀/ACA-G being electron-rich. The Pd(0) 3d_{5/2} binding energy of Pd₃Ag₁₀/ACA-G is close to but slightly lower than that of Pd₃Ag₁₀/ACA-C (335.7 eV vs. 335.9 eV). It indicates that metallic Pd species are in a similar surface condition, but metallic Pd species over Pd₃Ag₁₀/ACA-G also represent trend to enriched electrons.

In the Ag 3d spectra (**Figure 3B**) of Pd₃Ag₁₀/ACA-G, two main peaks at 368.4 eV and 374.4 eV are assigned to metallic Ag species^{43,44}. No Ag(I) related peaks were observed, all the Ag species are in metallic states. In comparison, the Ag(0) 3d peaks of Pd₃Ag₁₀/ACA-C shift to lower binding energy 368.2 eV and 374.2 eV, respectively,

which is the opposite of the binding energy shift trend in the Pd 3d spectra (**Figure 3A**). As shown in **Table 1**, the surface Ag/Pd ratio of Pd3Ag10/ACA-G is 9.68, ~15-fold higher than that of Pd3Ag10/ACA-C (0.63). Like the report of Mustafa et al., the core-level shifts in the binding energy of Ag 3d_{5/2} showed a good linear relationship with the number of Pd atoms neighbors around Ag atoms. Higher binding energies correspond to more contiguous Ag ensembles with a lower number of Pd neighbors per Ag atom⁴⁵. In our case, more contiguous Ag atoms are formed on the Pd3Ag10/ACA-G surface, leading to the Ag 3d_{5/2} peak shifting to higher binding energies.

In the high-resolution XPS spectra of O 1s (**Figure S5B**), the asymmetric peaks suggest different forms of O²⁻ existing in Pd3Ag10/ACA-C and Pd3Ag10/ACA-G. The peaks at 532.1 eV and 533.4 eV are assigned to the C-O bonds of adsorbed hydroxyl groups on the carbon support and -OH of hydroxyl groups, respectively. Moreover, the broad adsorbed oxygen peaks were detected at 537.4 eV. The same binding energy of C-O bonds and O-H bonds of hydroxyl groups and similar O fractions of Pd3Ag10/ACA-G and Pd3Ag10/ACA-C samples suggest the O-containing groups are not significantly changed during co-reduction and galvanic replacement methods.

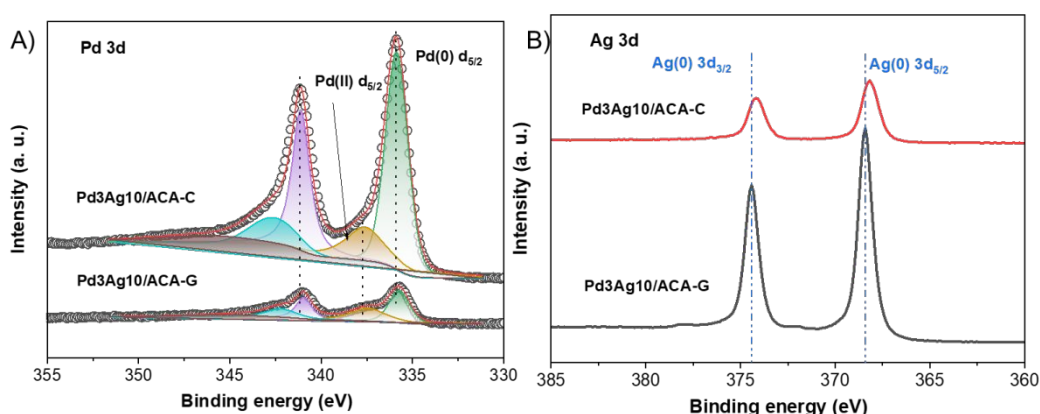


Figure 3. A) Pd 3d, B) Ag 3d XPS spectra of Pd3Ag10/ACA-C and Pd3Ag10/ACA-G.

Table 1. Surface elements containing in Pd3Ag10/ACA-C, Pd3Ag10/ACA-G, and Spent Pd3Ag10/ACA-G.

Catalyst	Surface elements atomic fraction			
	C (%)	O (%)	Pd (%)	Ag (%)
Pd3Ag10/ACA-C	88.30	5.74	3.65	2.31
Pd3Ag10/ACA-G	87.12	6.25	0.62	6.00
Spent Pd3Ag10/ACA-G	88.94	7.12	0.85	3.09

3.2. Activity evaluations

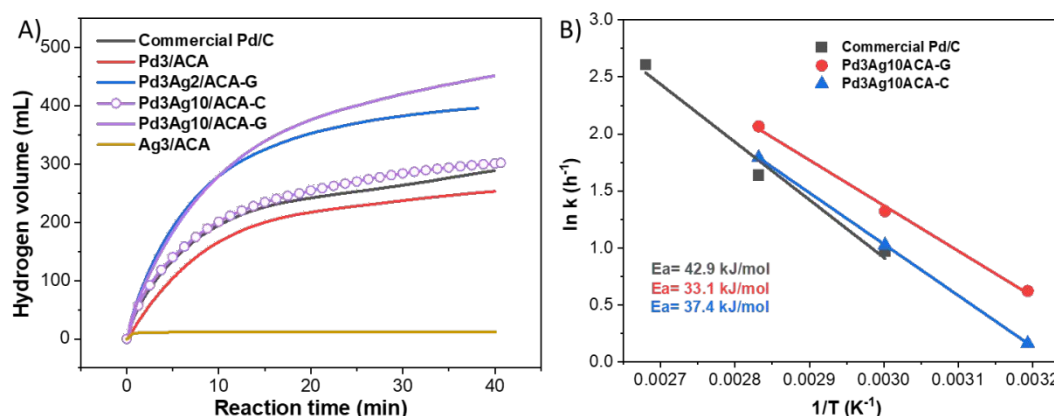


Figure 4. Formate dehydrogenation performance over A) Pd-Ag catalysts with increasing Ag ratios and Pd-Ag catalysts with different structures. Reaction conditions: 0.05 g catalyst, 20 mmol Ammonium formate, 80 °C, 40 min, 1 atm N₂. B) Kinetics of Arrhenius plots over Pd3Ag10/ACA-G, Pd3Ag10/ACA-C, and commercial Pd/C (the initial reaction rates were averaged from 0 to 5 min).

To investigate the catalytic performance of formate dehydrogenation over Pd-Ag bimetallic nanoparticles supported on carbon support catalysts, we evaluated the activities of the Pd-Ag/ACA catalysts with different Pd/Ag ratios. **Figure S6** shows that only a trace amount of CO₂ and no ammonia were detected in the gas phase products. Contrarily, hydrogen is the dominant gaseous product and accounts for 99.6

v% (**Table S1**). As shown in **Figure 4A**, under the same reaction conditions (0.05 g catalyst, 20 mL 1 M ammonium formate, 80 °C, 1 atm of N₂, 40 min), the released H₂ volumes (or H₂ yields) over the Pd₃/ACA, Pd₃Ag₂/ACA-G, Pd₃Ag₁₀/ACA-G, and Ag₃/ACA catalysts are 253 mL (50%), 396 mL (79%), 451 mL (90%), and 12 mL (2%), respectively. The hydrogen volume over Pd₃/ACA monometallic catalyst is lower than that of the commercial Pd/C (5 wt% Pd loading) (288 mL H₂ or 57% yield of H₂) due to a lower Pd loading. However, both Pd-Ag/ACA-G catalysts have improved hydrogen yields over commercial Pd/C even though with lower Pd loading. It suggests that surface-decorated Ag can significantly facilitate dehydrogenation performance. With an increasing amount of Ag species introduction, the hydrogen yields enhanced from 78.6% to 89.5%, further confirming the specific contribution of surface-decorated Ag species. Ag₃/ACA catalyst generates little hydrogen (12 mL, 2%) compared with other Pd-based catalysts, demonstrating that the metallic Ag nanoparticles are not the active sites of formate dehydrogenation. Thus, when more surface-decorated Ag species were added, the interaction between Pd-Ag species was enhanced, leading to higher activity instead of adding more active sites. Therefore, attention should be paid to studying Pd-Ag interaction to understand the mechanisms of Pd-Ag catalysts efficiently converting formate to bicarbonate.

To gain more insight into the Pd-Ag species interactions, we evaluated the formate dehydrogenation performance over Pd₃Ag₁₀/ACA-G and Pd₃Ag₁₀/ACA-C at the same reaction conditions (**Figure 4A**). After 40 minutes, the Pd₃Ag₁₀/ACA-G catalyst shows a higher hydrogen yield (451 mL, 90%) than Pd₃Ag₁₀/ACA-C (301 mL, 60%) and Pd₃/ACA (253 mL, 50%), indicating that both kinds of Pd-Ag bimetallic active sites are more beneficial to monometallic Pd active sites, following an efficiency order

that Pd-based surface decorated structure > bulk alloy structure > monometallic active site. As shown in **Table 2**, the Pd₃Ag₁₀/ACA-G has a 2.6-fold higher turnover frequency (TOF, the detailed TOF calculation is shown in Supporting Information) of 5202 h⁻¹ than the commercial Pd/C (1966 h⁻¹). The TOFs of the Pd₃Ag₁₀/ACA-C and Pd₃/ACA are 3395 h⁻¹ and 2804 h⁻¹, respectively, suggesting that the Pd-Ag bimetallic structure plays a more vital role than the composition ratio. Indeed, the Pd₃Ag₁₀/ACA-G has a higher TOF than most Pd-Ag-based catalysts reported in the literature under comparable reaction conditions (**Table 2**). Zhao et al.⁴⁶, Yin et al.⁴⁷, Song et al.⁴⁸, and etc. calculated TOF per unit of the total amount of loaded metal species. In this report, Ag species have no activity for formate dehydrogenation. Thus, we calculated the TOF based on the total amount of Pd loading. The catalysts with the reported TOF (turnover frequency) data are labeled with various superscripts to distinguish the different TOF calculation methods. Our as-developed Pd₃Ag₁₀/ACA-G displayed enhanced TOFs compared with most reported Pd-Ag bimetallic catalysts, as shown in Table 2. For instance, Yin et al.⁴⁷ mentioned the TOF, based on total metal loading, of Pd_{0.7}Ag_{0.3}/CeO_x-NPC was 875 h⁻¹ in dehydrogenation of the formic acid and sodium formate mixture. It was revised to 1250 h⁻¹ when calculated through the same method as we used, i.e., normalized by the Pd loading amount, which is still lower than that of Pd₃Ag₁₀/ACA-G (1790 h⁻¹) at 40 °C. An exception is Masuda et al.⁴⁹ reported TiO_x shell-modified core-shell Ag@Pd supported on TiO₂, which exhibited a high TOF of 6499 h⁻¹ at 75 °C, calculated over the total amount of Pd loading. The presence of Ag sites and TiO_x shell in the vicinity of Pd likely promoted the rate-limiting C–H bond dissociation step during dehydrogenation. Nevertheless, the Pd₃Ag₁₀/ACA-G catalysts are easily utilized in practical applications due to their robust structure and relatively simple synthesis method compared to the TiO_x shell-modified core-shell Ag@Pd

catalysts.

It is interesting to compare the dehydrogenation of ammonium formate (AF), formic acid (FA), and their mixtures (AF + FA) over the Pd₃Ag₁₀/ACA catalyst. We found that FA dehydrogenation alone was slower than that of AF under identical reaction conditions (Figure S7). Considering the mixture of AF and FA (50:50), the dehydrogenation of AF was adversely affected by FA in terms of the overall H₂ yields. It is not surprising that the dehydrogenation of AF yielded ammonium bicarbonate and hydrogen ($\text{HCOO}^- + \text{H}_2\text{O} \rightarrow \text{HCO}_3^- + \text{H}_2$) with the Pd₃Ag₁₀/ACA catalyst. However, ammonium bicarbonate further reacted with FA in the solution to be converted back to AF and co-produce CO₂ ($\text{HCO}_3^- + \text{HCOOH} \rightarrow \text{HCOO}^- + \text{H}_2\text{O} + \text{CO}_2$). The presence of CO₂ might inhibit AF dehydrogenation as both FA and AF/FA dehydrogenation yielded only approximately 50% of H₂ from AF dehydrogenation after 40 minutes of the reactions. This phenomenon should be further investigated.

The kinetic rate measurements of as-developed Pd₃Ag₁₀/ACA-G, Pd₃Ag₁₀/ACA-C, and commercial Pd/C were conducted at 40 °C -100 °C, as shown in **Figure S8**. The calculated TOFs of Pd₃Ag₁₀ACA-G from 40 °C to 80 °C increased from 1799 h⁻¹ to 5202 h⁻¹, indicating higher reaction temperatures benefits the dehydrogenation kinetics. The corresponding Arrhenius plots were obtained according to the first-order power law kinetic model¹² and were shown in **Figure 4B**. The activation energies (E_a) of Pd₃Ag₁₀/ACA-G and Pd₃Ag₁₀/ACA-C were 33.1 kJ/mol and 37.4 kJ/mol, lower than that of commercial Pd/C (42.9 kJ/mol). It means the formate dehydrogenation can be performed with a lower energy barrier over Pd-Ag bimetallic active surface, in which the one prepared by galvanic replacement is more favorable than the one prepared by the co-reduction method.

Table 2. The catalytic activity of different Pd-Ag based catalysts for the formate dehydrogenation.

Catalysts	Substance	T (°C)	TOF (h ⁻¹)	Ref.
Commercial Pd/C	NH ₄ HCO ₂	80	1966 ^a	
Pd3/ACA	NH ₄ HCO ₂	80	2804 ^a	
Pd3Ag10/ACA-C	NH ₄ HCO ₂	80	3395 ^a	This work
Pd3Ag10/ACA-G	NH ₄ HCO ₂	80	5202 ^a	
		40	1790 ^a	
PdAg/rGO	H ₂ CO ₂ /NaHCO ₂	25	164 ^b	46
Pd _{0.7} Ag _{0.3} /CeO _x -NPC	H ₂ CO ₂ /NaHCO ₂	30	875 ^b	47
PdAg@ZrO ₂ /C/rGO	H ₂ CO ₂ /NaHCO ₂	60	4500 ^b	48
PdAg-MnO _x /N-SiO ₂	H ₂ CO ₂	25	1430 ^b	50
C-PdAg BNSs	H ₂ CO ₂ /NaHCO ₂	50	378 ^b	51
PVPI@PdAg NWs	H ₂ CO ₂	25	312 ^b	52
Ag@Pd@TiO _x (TTIP)/TiO ₂	NaHCO ₂	75	6499 ^a	49
PdAg/CN-3%	H ₂ CO ₂	30	254 ^b	53

* TOF was calculated based on different sites. a: based on total Pd atoms; b: based on total metal atoms;

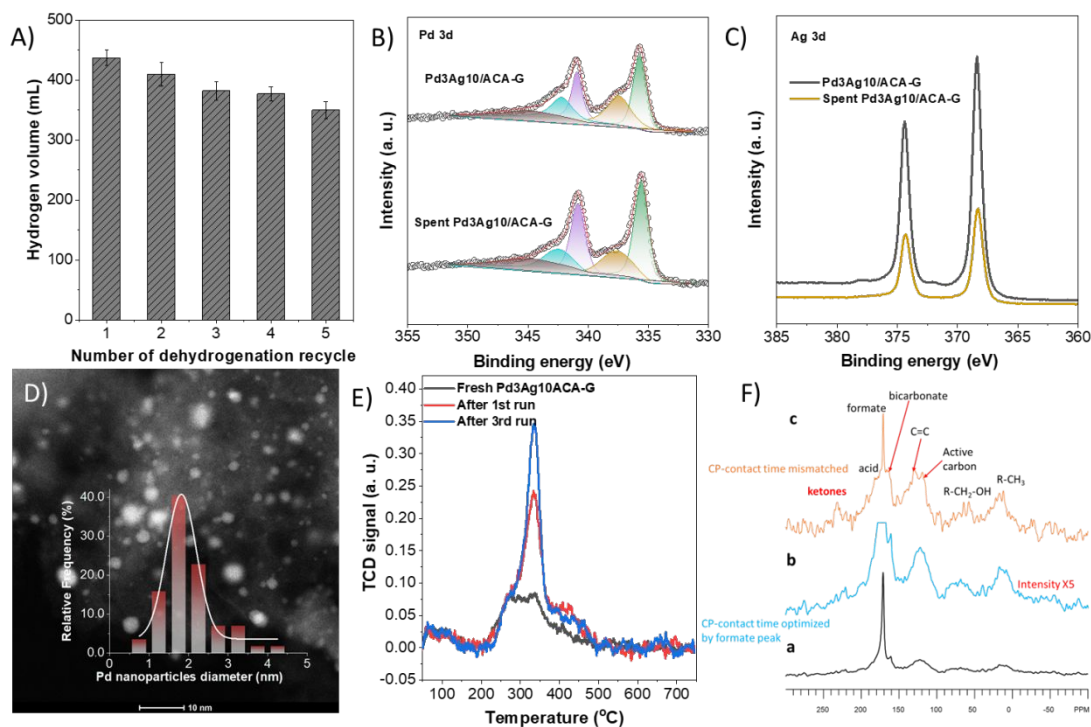


Figure 5. A) recyclability of Pd₃Ag₁₀/ACA-G (Reaction conditions: 0.05 g catalysts, 20 mmol ammonium formate, 80 °C, 1 atm N₂, 40 min). B) Pd 3d; C) Ag 3d XPS spectra comparison of fresh and spent Pd₃Ag₁₀/ACA-G. D) HRTEM image of spent Pd₃Ag₁₀/ACA-G and NPs size distribution (inset). E) TPD curves of fresh and spent Pd₃Ag₁₀/ACA-G after different cycles. F) ¹H-¹³C cross-polarization NMR spectra of formate decomposition over Pd/Ag/AC catalyst (a) optimized CP condition with respect to the formate peak (full intensity spectrum); (b) vertically expanded by 5 times of (a); (c) mismatched conditions with respect to formate but more matched for the other peaks.

In addition, the stability tests were performed using the recycled Pd₃Ag₁₀/ACA-G catalysts (**Figure 5A**). We observed that the activities gradually decreased during the formate dehydrogenation reactions throughout five repeated cycles. The TOFs at different cycles were calculated and summarized in Table S3. Similar with the total hydrogen volume decrease, the TOFs calculated by first 10 min data gradually reduce from 5202 h⁻¹ to 3242 h⁻¹, indicating an intrinsic deactivation occurs over

Pd₃Ag₁₀/ACA-G catalyst. In parallel, other researchers reported the deactivation of Pd-Ag bimetallic catalysts, as well. Zhao et al, demonstrated the Pd_{0.9}Ag_{0.1}B/rGO lose obvious activity in the third run, which was attributed to the aggregation of rGO during the reaction process⁴⁶. Yang et al, reported the AgPd NPs were well dispersed on the graphene without no obvious agglomeration but the catalytic activity slightly decreases in the third run⁵⁴. Multiple possible reasons make it unclear to account for deactivation. Herein, a systematic investigation of deactivation mechanism was conducted to have an insight on that. The ICP-MS analysis of the spent Pd₃Ag₁₀/ACA-G (**Table S4**) and the hot filtration experiment (**Figure S9**) revealed no Pd leaching during the reaction. The XPS spectra of the spent Pd₃Ag₁₀/ACA-G were compared with the fresh Pd₃Ag₁₀/ACA-G to elucidate the possible deactivation mechanism. The Pd 3d spectra (**Figure 5B**) display that the metallic Pd species of spent Pd₃Ag₁₀/ACA-G maintain the same binding energy as fresh Pd₃Ag₁₀/ACA-G, indicating Pd species are stable during dehydrogenation. And the Pd(0)/Pd(II) ratio of the spent catalyst is 2.08, higher than that of the fresh catalyst (1.15), demonstrating more Pd(II) species were reduced to the metallic state, which would be beneficial to keep high activity. The Ag 3d spectra of the spent catalyst (**Figure 5C**) show the same binding energy but a lower peak intensity, meaning the Ag electronic conditions are stable, but the quantity of Ag species decreases. **Table 1** reveals the specific Pd and Ag atomic fraction changes after dehydrogenation. The Pd fractions increase from 0.62% to 0.85%, while the Ag fractions decrease from 6.00% to 3.09%. It means the surface structure is reconstructed after the reaction, around half of Ag species leave from the surface decorated Pd-Ag nanoparticles, and a little more Pd are exposed. This surface structure change possibly led to deactivation due to the Pd monometallic NPs support on carbon showing a relatively low activity. As shown in **Figure 5D**, the HRTEM image of the spent

Pd₃Ag₁₀/ACA-G catalyst present means Pd NPs size is 2.00 nm, larger than that of fresh catalyst (1.62 nm). The slight growth of Pd NPs size may also be one of the deactivation reasons. **Table S4** showed that the Pd dispersion tested by CO pulse chemisorption of Pd₃Ag₁₀/ACA-G decreases from 21.4% to 19.2% and 17.9% with the cycling reaction. It is attributed to the larger NPs size tested by HRTEM and occupation of chemically bonded carboxylate species tested by TPD. TPD spectra of fresh and spent catalysts after different cycles are shown in **Figure 5E**. Compared with fresh Pd₃Ag₁₀/ACA-G, two peaks at 335 °C and 420 °C appear after one-time dehydrogenation. After a three-cycles reaction, the peak at 335 °C increased further, while the peak at 420 °C maintains the same peak intensity. According to our previous research, the multiple O-containing species on the carbon support will decompose during the temperature ramping. The carboxylate species decompose from 250 °C to 350 °C, and the lactone species decompose from 350 °C to 450 °C. The adsorbed carboxylate species are dominant component negatively impact the formate dehydrogenation activity. On the Pd₃Ag₁₀/ACA-G surface, carboxylate related species with 335 °C decomposition temperature, accumulate on the catalyst surface with more cycle's operation, high possibly leading to the deactivation. Besides, it is well accepted that Pd NPs are mobile and unstable on the carbon surface, resulting in aggregation and change of surrounding bonding conditions during the reaction, which may lead to deactivation. Therefore, multiple factors, such as the surface decorated Pd-Ag nanostructure change, growth of NPs size, and intermediate poisoning adsorption, lead to the deactivation of Pd₃Ag₁₀/ACA-G in the formate dehydrogenation reaction. In addition, to further unravel the origin of deactivation, the *Operando* NMR ¹H-¹³C cross-polarization was employed to identify surface-adsorbed species, shown in **Figure 5F**. By optimizing the contact time of cross-polarization with regard to the formate

resonance, the formate peak at 171 ppm displays the highest intensity. Other relatively lower peaks are observed clearly as well by expanding the spectrum five times. By mismatching the optimized conditions, besides formate and bicarbonate peaks, those small peaks are still observable, including other minor visible peaks. These peaks between 220-240, 180-190, 100-140, 50-90, and 0-30 ppm are attributed to ketones, acid, C=C, oxygenated carbon, and aliphatic carbon, respectively⁵⁵. They are tightly bound on the surface such that ^1H - ^{13}C cross-polarization can catch these resonance signals, evidencing that these surface-adsorbed carbon species are responsible for the fast deactivation. Based on the signature peaks associated with ketones, acid, C=C, oxygenated carbon, and aliphatic carbon, a model of, likely a large molecule containing the identified functional groups by NMR, should be able to be proposed to describe the surface adsorbed chemical species that are responsible for the catalyst deactivation.

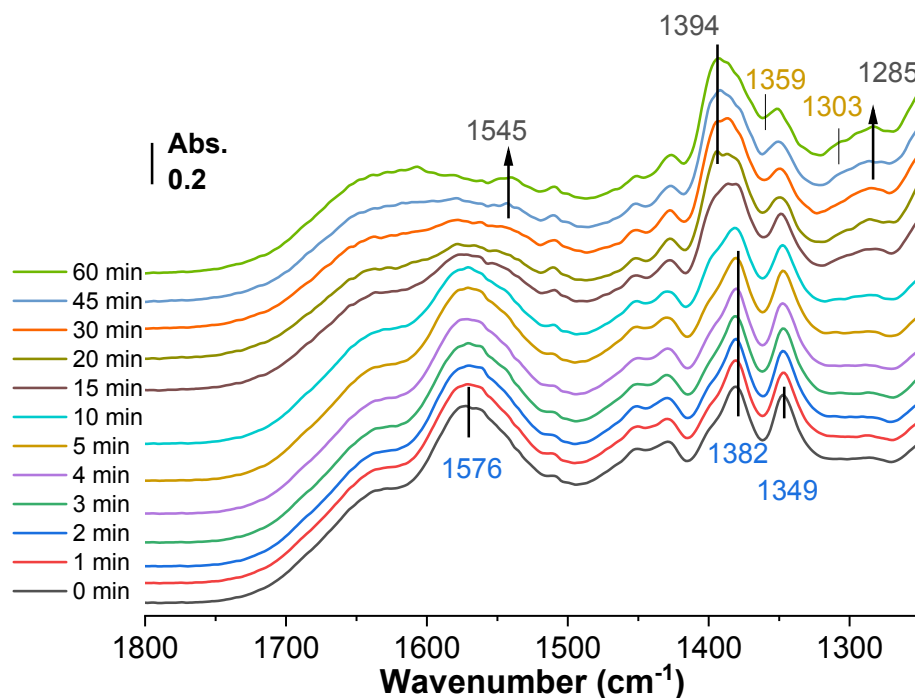


Figure 6. Selected *in-situ* ATR-IR spectra on Pd₃Ag₁₀/ACA-G catalyst at 40 °C as a function of time recorded after rinsing a 2 M ammonium formate solution to the spectral cell.

The reference spectrum was recorded in the air before the rinse.

The *in-situ* ATR-IR of formate dehydrogenation over Pd₃Ag₁₀/ACA-G catalyst to further understand the ammonium formate dehydrogenation mechanism, enabling the time-resolved monitoring of formate conversion and bicarbonate formation. The formate-specific bands are at 1576, 1382, and 1349 cm⁻¹, while the peaks at 1359 and 1303 cm⁻¹ are related to bicarbonate bond vibrations¹². **Figure 6** shows the selected *in-situ* ATR-IR spectra recorded in the air at 40 °C for 1 h of the formate conversion on the Pd₃Ag₁₀/ACA-G catalyst. (The *in-situ* ATR-IR tests at different temperatures are shown in **Figure S10**.) At the initial stage, the specific peaks of formate can be observed at 1576 cm⁻¹, 1382 cm⁻¹, and 1349 cm⁻¹ after the catalyst film is rinsed by a 2 M ammonium formate aqueous solution. Then the formate peaks at 1576 cm⁻¹ and 1382 cm⁻¹ decreased significantly after 15 min of reaction, while the bicarbonate peaks at 1353 cm⁻¹ are so weak to be overlapped by formate peaks at 1349 cm⁻¹, and the bicarbonate peaks at 1303 cm⁻¹ are also barely to be ignored. Interestingly, a new peak at 1545 cm⁻¹ is related to carbonate, one possible intermediate of formate dehydrogenation to bicarbonate. Besides, two new peaks are enhanced during the dehydrogenation at 1394 cm⁻¹ and 1285 cm⁻¹, which are assigned to carboxylate species symmetric stretching. This result is consistent with the TPD peak at 335 °C shown in **Figure 5E**, further confirming that the intermediate carboxylate species are generated from formate but strongly adsorbed on the catalyst surface and poisoned the surface-active site. In addition, the *Operando* ¹³C single pulses NMR of formate decomposition over the Pd₃Ag₁₀/ACA-G catalyst was operated to study the reaction mechanism. The reaction was taken as the temperature increases, shown in **Figure S11**. The dominating peaks observed at about 171 ppm are ascribed to formate reactant, while the increasing

peak at 160 ppm with temperature is assigned to bicarbonate product⁵⁶. The fact that bicarbonate is generated at such a sluggish pace at even 93 °C reveals that there is a severe deactivation occurring during the operando NMR study in the spinning rotor. As the intensity vertically is enlarged by 20 times, the CO₂ peak at 125 ppm is clearly observed during the operando study⁵⁷. However, when comparing intensities of bicarbonate and CO₂, it is evident that the product bicarbonate is a very stable species under this environment, with only less than 3% of bicarbonate decomposing to CO₂.

3.3. Computational results

To examine the role of Ag in the Pd-based catalyst for formate dehydrogenation, the energies of hydrogen and formate adsorption on Pd (111), Ag (111), and Ag/Pd (111) surfaces were computed using first-principles DFT calculations. The representative structures for optimized configurations of adsorbed formate and hydrogen are displayed in **Figure 7**. Model selection was based on the XPS results which showed that the Ag surface fraction (6.00 %) was 10 times higher than Pd surface fraction (0.62 %) of the as-prepared Pd₃Ag₁₀/ACA-G. Thus, the Ag/Pd surface was constructed as a surface alloy of Pd (111) with an Ag on-top layer, as shown in **Figure 7**.

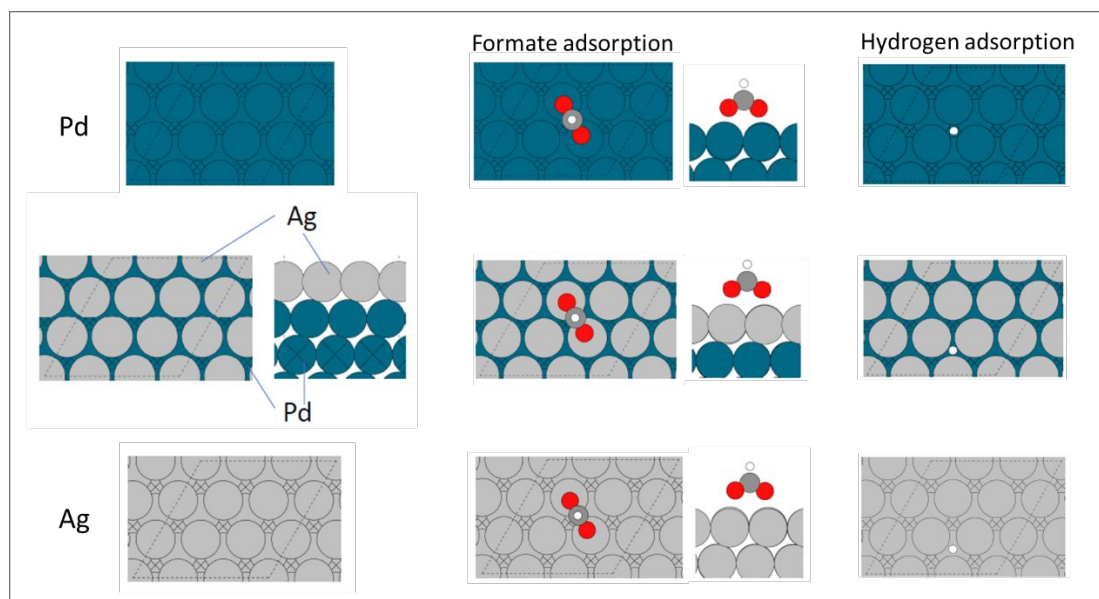


Figure 7. View of Pd (111), Ag (111), and Ag/Pd (111) surface alloy models, and the representative configurations for formate and hydrogen bound to positions of Pd, Ag, and Pd-Ag bimetallic surface. Atom color representations: Pd (teal), Ag (grey), O (red), C (dark gray), and H (white).

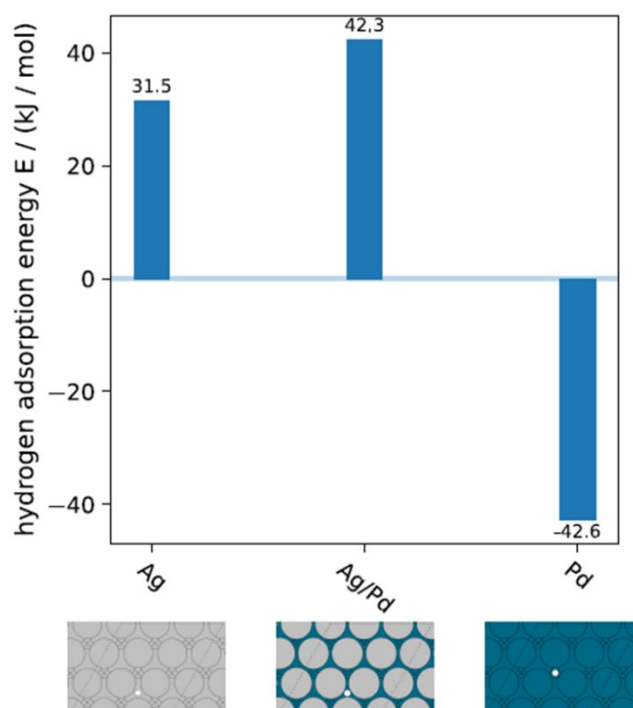


Figure 8. Adsorption energies (in kJ/mol) of hydrogen on Ag (111), Ag/Pd (111) and Pd (111) surfaces. Atom color representations: Pd (teal), Ag (grey), O (red), C (dark gray) and H

(white).

Figure 8 shows the adsorption energies of hydrogen on the different surface models. The hydrogen adsorption energy on Pd (111) surface is -42.6 kJ/mol, much lower than Ag (111) surface (31.5 kJ/mol) indicating strong binding of hydrogen to Pd compared to the Ag surface. The binding is further weakened on Ag/Pd (42.3 kJ/mol) indicating that the electronic and compressive strain on Ag overlayer (lattice constant Ag \sim 4.09 Å, Pd \sim 3.89 Å) in the Ag/Pd surface can play an important role in tuning the binding energy of hydrogen and formate. **Figure 9** further highlights the synergistic effect of Ag and Pd on formate binding. **Figures 9A** and **9B** show the influence on the binding energy with inclusion of solvation and potential considerations in the calculations. When solvation and potential of zero charge (PZC) are accounted for (**Figure 9B**), the formate adsorption is much stronger on Pd (111) surface (-88.9 kJ/mol) and weaker on Ag (111) surface (-21.3 kJ/mol). The formate adsorption strength is intermediate on the Ag/Pd surface alloy (-42.6 kJ/mol). This trend can be primarily rationalized by the potential of zero charge U_{PZC} of these surfaces that is intrinsically linked to the surface charge transfer coefficient and modulated by the catalyst composition, adsorbed species, and the microenvironment at the catalyst/solution interface. **Figure S12** shows the computed U_{PZC} of Ag is -0.62 V vs. SHE, lower than that of Ag/Pd surface alloy (-0.50 V vs. SHE) and Pd surface (0.24 V vs. SHE); consistent with previously published results^{58,59}. The higher U_{PZC} value of Pd intrinsically makes Pd have a higher affinity towards negatively charged adsorbates like formate ions relative to Ag. The U_{PZC} of Ag/Pd is lower than Pd and plays a critical role in modulating the strength of formate adsorption. While the capacitance effect shown here will likely weaken as the coverage of formate increases, the results in **Figure 9B** and **Figure S12** highlight the electronic role of the metal/liquid interphase in modulating catalytic events such as adsorption and

dehydrogenation. Following Sabatier's principle, we rationalize that the temperate bonding strength between the formate reactant and catalytic active site results in a higher TOF and enhanced dehydrogenation performance as observed experimentally with Pd₃Ag₁₀/ACA-G. Additionally, weak hydrogen binding on the Ag/Pd surface alloy can promote associative desorption of H₂ from the catalyst surface.

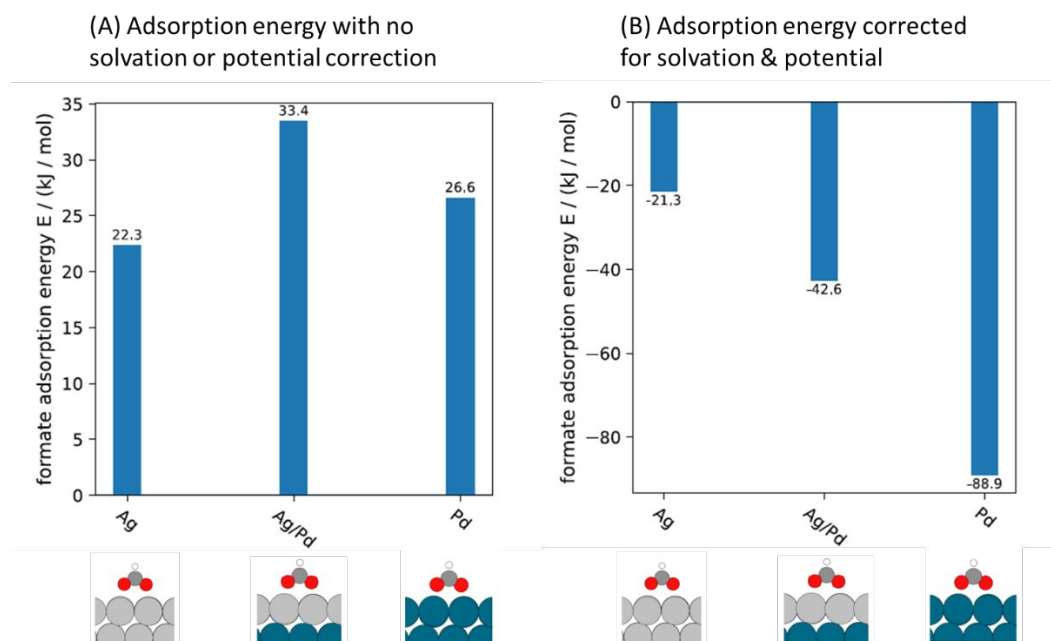


Figure 9. Adsorption energies (in kJ/mol) of formate on Ag (111), Ag/Pd (111) and Pd (111) surfaces (A) without and (B) solvation and potential correction. Adsorption energies in (A) use a vacuum/metal interface. Energies in (B) use 1 M monovalent equivalent implicit solution with the surface set at its U_{PZC} (net surface charge = 0). Atom color representations: Pd (teal), Ag (gray), O (red), C (dark gray) and H (white).

3.4. Catalyst structure-performance discussion

According to the formate dehydrogenation evaluations (**Figure 4A**), the Pd-Ag bimetallic catalysts prepared by galvanic replacement method, Pd₃Ag₂/ACA-G and Pd₃Ag₁₀/ACA-G, displayed improved hydrogen production (396 mL and 451 mL) compared to monometallic catalyst Pd₃/ACA (253 mL). However, the hydrogen

production increased from 253 mL to 396 mL (56.5 % improvement) only with 2 wt% Ag atoms doping, while that increased from 396 mL to 451 mL (12.2 % improvement) with the Ag loading increased from 2 wt% to 10 wt%. It means that the existence of the Pd surface decorated with Ag atoms can significantly improve the formate dehydrogenation performance. DFT simulations further substantiate the role of surface Ag in decreasing the electron affinity of Pd thereby weakening the hydrogen adsorption energy (**Figure 8**) and modulating formate binding with intermediate adsorption strength (**Figure 9B**). As Dong Yun Shin et al. reported, the potential limiting step in formate dehydrogenation was the final step of $\text{H}_2(\text{g})$ desorption⁶⁰. The weakest hydrogen adsorption energy over Ag/Pd (111) facet enhances hydrogen generation efficiently. And as the first step of formate adsorption, it should be a moderate to balance the adsorption rate and energy barrier of the following steps to reach a robust formate dehydrogenation. When more Ag atoms are loaded on the Pd/C surface, only the Ag atoms those form Ag/Pd structure can be involved in the dehydrogenation reaction due to the Ag/C species having no activity (**Figure 4A**). It accounts for the slightly improved hydrogen release from Pd3Ag2/ACA-G to Pd3Ag10/ACA-G. The hydrogen production comparison of Pd3Ag10/ACA-G (451 mL) and Pd3Ag10/ACA-C (301 mL) indicates the specific surface condition existing on Pd3Ag10/ACA-G is of more catalytic efficiency even though an identical Pd/Ag ratio. Firstly, as shown in TEM images (**Figure S1**), the Pd3Ag10/ACA-G has a smaller NPs size (~1.62 nm), while that of the Pd3Ag10/ACA-C sample is around 2.50 nm. The highly dispersed NPs of Pd3Ag10/ACA-G give it more chance to convert formate to bicarbonate and release hydrogen. Besides, the XPS results of Pd 3d spectra (**Figure 3A**) present that the metallic Pd 3d_{5/2} of Pd3Ag10/ACA-G has lower binding energies (335.7 eV) than that of Pd3Ag10/ACA-C (335.9 eV), indicating the metallic Pd over Pd3Ag10/ACA-G

has higher electronic density. As analyzed in Ag 3d spectra (**Figure 3B**), the metallic Ag atoms of Pd₃Ag₁₀/ACA-G have higher binding energies and possess higher electronic density. Combined with Pd and Ag spectra, the electron density of Pd-Ag bimetallic NPs as a whole active species is enhanced in the Pd₃Ag₁₀/ACA-G sample. According to publications, the electron-rich Pd species are more catalytic efficient for hydrogen regeneration from formate solution^{10,21,60}. The hydrogen yield gradually decreases during the reusability test (**Figure 5A**), possibly because of the surface composition change. Multiple characterizations, such as XPS analysis of spent Pd₃Ag₁₀/ACA-G (**Figure 5B, C**), TEM images (**Figure 5D**), TPD (**Figure 5E**), and ¹H-¹³C cross-polarization NMR (**Figure 5F**) revealed that the surface Ag fraction decrease, NPs size growth, O-containing species (carboxylate, ketones, etc.,) adsorption result in the catalyst deactivation.

4. Conclusion

In summary, ammonium formate solution is a promising liquid organic hydrogen carrier since both hydrogenation and dehydrogenation reactions are thermodynamically favorable, in which case hydrogen can be stored and released through the bicarbonate/formate cycles at mild conditions. However, it suffers from kinetic challenges and unclear Pd catalyst degradation mechanisms. In this report, we prepared carbon-supported Pd-Ag bimetallic NPs catalysts with two kinds of microstructures through galvanic replacement and co-reduction methods for catalyzing ammonium formate dehydrogenation. The best performed catalyst, Pd₃Ag₁₀/ACA-G, presents the highest activity with the TOF of 5202 h⁻¹ at 80 °C (~2.6-fold higher than 5% commercial Pd/C) and the lowest activation energy of 33.1 kJ/mol. Furthermore, the smaller NPs size, the enhanced electron density of Pd-Ag bimetallic NPs, the weakest

hydrogen adsorption energy, and intermediate formate adsorption energy over the Ag/Pd (111) facet calculated by DFT corroborate the superior formate dehydrogenation performance over the Pd₃Ag₁₀/ACA-G sample. However, changes on the catalyst surface, including surface Ag fraction decrease, NPs size growth, and O-containing species (carboxylate, ketones, etc.) adsorption, gradually deactivated the catalyst. The structural and composition parameters identified for the developed Pd-Ag bimetallic catalysts in relation to their catalytic properties towards formate dehydrogenation reaction can provide valuable guidelines for developing an efficient heterogeneous catalyst for formate-based hydrogen carrier system applications.

CRedit authorship contribution statement

Zhun Dong: Conceptualization, Investigation, Writing – original draft. **Ahmad Mukhtar:** Writing – review & editing. **Thomas Ludwig:** Software, Writing – original draft. **Sneha A. Akhade:** Conceptualization, Writing – review & editing. **Wenda Hu:** Writing – original draft. **Jian Zhi Hu:** Resources, Writing – review & editing. **Katarzyna Grubel:** Validation, Writing – review & editing. **Mark Engelhard:** Resources. **Brandon C. Wood:** Resources. **Tom Autrey:** Validation, Writing – review & editing. **Hongfei Lin:** Conceptualization, Funding acquisition, Supervision, Writing – review & editing.

Notes

The authors declare no competing financial interests.

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