Highly efficient hydrogen generation from formic acid using a reduced graphene oxide-supported AuPd nanoparticle catalyst†

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Highly dispersed AuPd alloy nanoparticles have been successfully immobilized on reduced graphene oxide (rGO) using a facile non-noble metal sacrificial method, which exhibit the highest activity at 323 K (turnover frequency, 4840 h⁻¹) for hydrogen generation without CO impurity from the formic acid/sodium formate system.

Hydrogen is of critical importance in clean energy applications although effective hydrogen storage technologies still need to be developed.¹ Formic acid (FA) is a promising hydrogen carrier,² which has several remarkable features such as (a) nontoxicity with high hydrogen content (4.4 wt%), (b) high stability as a liquid at room temperature, and (c) easy synthesis via a biomass process, CO₂ reduction or methyl formate hydrolysis.³ Depending on the catalysts, however, FA can be decomposed in two ways: desirable dehydrogenation (HCOOH → H₂ + CO₂) and undesirable dehydration (HCOOH → H₂O + CO), which produces CO impurities that are toxic to the fuel cell catalysts.⁴

So far, homogeneous catalysts have been extensively studied for hydrogen production from FA and very high catalytic activities have been reported.⁵ Along with the ongoing progress in FA dehydrogenation using homogeneous systems, heterogeneous catalysts have attracted tremendous attention due to their advantages of easy separation and recycling. A large number of metal nanoparticle (MNP) catalysts have been studied, most of which, however, suffer from severe aggregation and limited catalytic activities.⁶

Herein, we report a highly active AuPd alloy nanoparticle catalyst prepared using a non-noble metal sacrificial approach (NNMSA), which exhibits the highest turnover frequency (TOF) of 4840 h⁻¹ at 323 K for hydrogen generation from the FA/sodium formate (SF) system.⁷

Cobalt was used as the sacrificial agent. Amorphous Co₃(BO₃)₂ was formed from the reaction between Co(CH₃COO)₂ and NaBH₄, which can be easily dissolved in H₃PO₄ (Fig. S1 and S2, for details see the ESI†).⁸ Graphite oxide (GO) was prepared from graphite using a modified Hummers’ method.⁹ The rGO-supported AuPd nanoparticle (NP) catalyst was synthesized using a wet-chemical process, as illustrated in Scheme 1. Firstly, according to the stoichiometry, the aqueous solution containing HAuCl₄, Co(CH₃COO)₂ and K₂PdCl₄ was added into 50 mL of GO suspension (0.1 wt%). Then the above mixture was reduced by a fresh aqueous solution of NaBH₄. After continuous stirring for 2 h, the mixture was washed and re-dispersed in H₃PO₄ solution for etching (2 h). Finally, the catalyst, denoted as (CoₓAu₁₋ₓPd₁/rGO (ESI†), was obtained by centrifugation and washing with water.

![Scheme 1 Schematic illustration of the synthesis of the (Coₓ)AuₓPd₁₋ₓ/rGO catalyst via a non-noble metal (cobalt) sacrificial approach (NNMSA).](image-url)

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Fig. 1a shows the powder X-ray diffraction (PXRD) pattern of the (Co₃)Eₐu₀.₆Pd₀.₄/rGO catalyst. Except for a low and broad diffraction of rGO at 2θ = 24°, the diffraction peaks, which are located between the fcc Au (JCPDS No. 65-8601) and Pd (JCPDS No. 65-2867), indicate the formation of an AuPd alloy. For comparison, PXRD patterns of the non-etched counterparts Co₃Au₀.₆Pd₀.₄ and Co₃Au₀.₆Pd₀.₄/rGO (Fig. S3, ESI†) show an additional broad band for Co₃(BO₃)₂. After etching with an acid, the characteristic peaks corresponding to AuPd are clearly visible and match well with the directly deposited counterpart of Au₀.₆Pd₀.₄/rGO (Fig. S3, ESI†), confirming the successful removal of cobalt and the formation of AuPd. The N₂ sorption isotherms for (Co₃)Eₐu₀.₆Pd₀.₄/rGO, Co₃Au₀.₆Pd₀.₄/rGO and Au₀.₆Pd₀.₄/rGO are shown in Fig. 1b. Generally, the restacking of rGO can be partially overcome by the direct deposition of NPs. Hence, the Au₀.₆Pd₀.₄/rGO and Co₃Au₀.₆Pd₀.₄/rGO possess a type-IV N₂ sorption curve with Brunauer–Emmett–Teller (BET) surface areas of 189 and 219 m² g⁻¹, respectively. Notably, the surface area of (Co₃)Eₐu₀.₆Pd₀.₄/rGO (321 m² g⁻¹) increases drastically, illustrating that the removal of cobalt by the acid etching has resulted in high porosity, which will favour the diffusion of reactants to metal NPs in catalysis.

The morphologies of the Au₀.₆Pd₀.₄/rGO, Co₃Au₀.₆Pd₀.₄/rGO and (Co₃)Eₐu₀.₆Pd₀.₄/rGO catalysts were analysed by transmission electron microscopy (TEM). As shown in Fig. S4 and S5 (ESI†), Au₀.₆Pd₀.₄/rGO and Co₃Au₀.₆Pd₀.₄/rGO possess larger particles with severe aggregation (ESI†). In contrast, the AuPd NPs in the (Co₃)Eₐu₀.₆Pd₀.₄/rGO catalyst are well dispersed and immobilized on the rGO support (Fig. 2a). The high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) images further reveal the uniform distribution of AuPd NPs in the (Co₃)Eₐu₀.₆Pd₀.₄/rGO catalyst with an average particle size of 3.9 ± 0.9 nm (Fig. 2b and c), indicating that the sacrifice of the cobalt compound can prevent the primary AuPd NPs from aggregation, which will benefit their catalytic performance. Energy dispersive X-ray (EDX) analyses (Fig. S6–S8, ESI†) show that Co₃Au₀.₆Pd₀.₄/rGO is rich in cobalt, whereas no signals for cobalt are detected in (Co₃)Eₐu₀.₆Pd₀.₄/rGO, clearly confirming that cobalt has been eliminated by the H₃PO₄ etching. The inductively coupled plasma optical emission spectroscopic analysis reveals that the molar ratio of Co : Au : Pd in (Co₃)Eₐu₀.₆Pd₀.₄/rGO to be 0.04 : 0.6 : 0.4, indicating that almost all the co-precipitated Co₃(BO₃)₂ can be removed by H₃PO₄. The X-ray photoelectron spectroscopic (XPS) measurements show no signals for Co in (Co₃)Eₐu₀.₆Pd₀.₄/rGO before and after Ar sputtering (Fig. 3a). The observed Au 4f and Pd 3d spectra reveal that (Co₃)Eₐu₀.₆Pd₀.₄/rGO is composed of metallic Au and Pd (Fig. 3b and c), further confirming the successful formation of AuPd alloy NPs.

The catalytic activities of the (Co₃)Eₐu₀.₆Pd₀.₄/rGO catalysts in hydrogen generation from the FA/SF system were evaluated. Fig. 4a shows the gas generation from the FA/SF system in the presence of (Co₃)Eₐu₀.₆Pd₀.₄/rGO and Au₀.₆Pd₀.₄/rGO. Unquestionably, the sacrificial agent plays a dominant role, by decreasing the size of NPs, in the promotion of catalytic performance. Under our evaluation conditions, the (Co₃)Eₐu₀.₆Pd₀.₄/rGO catalyst shows an extremely high catalytic activity for the complete dehydrogenation of FA. Fig. 4b shows the kinetic curves of the FA/SF system in the presence of (Co₃)Eₐu₀.₆Pd₀.₄/rGO and Au₀.₆Pd₀.₄/rGO. The observed Au 4f and Pd 3d spectra reveal that (Co₃)Eₐu₀.₆Pd₀.₄/rGO is composed of metallic Au and Pd (Fig. 3b and c), further confirming the successful formation of AuPd alloy NPs.

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of FA with the release of 146 mL gas of H₂ + CO₂ in 0.62 min (n_{AuPd}/n_{FA} = 0.02 and n_{SF}/n_{FA} = 2.5) at 323 K, giving a TOF as high as 4840 h⁻¹ (Fig. S9, ESI†), the highest value reported thus far (Table S1, ESI†). After that, the generated gas with a slow speed is attributed to the decomposition of SF. The catalytic activities of the (Co₃)EAu₀.₆Pd₀.₄/rGO catalysts with lower metal loadings have not shown significant changes (Fig. S10, ESI†).

Gas chromatography (GC) measurements demonstrate that the produced gas consists of H₂ and CO₂ without a trace of CO (below 5 ppm), implying that the as-synthesized (Co₃)EAu₀.₆Pd₀.₄/rGO catalyst has a high selectivity for the dehydrogenation of FA into H₂ and CO₂ (Fig. S11, ESI†). Such excellent catalytic performance for FA dehydrogenation can meet the requirements of practical applications. Herein, it should be noted that the molar ratio of SF/FA has an obvious influence on the gas release rate (Fig. S12, ESI†). The gas release rate increases rapidly until the molar ratio of SF/FA increases up to 2.5. The further increase, however, makes no actual impression on the FA decomposition. On the other hand, the catalytic performance of (Co₃)EAu/rGO is affected by the molar ratio of Au/Pd (Fig. S13, ESI†). The mono-metallic (Co₃)Au/rGO has no activity while (Co₃)EPd/rGO has a low activity. However, the introduction of Au to Pd leads to a dramatically enhanced efficiency, which reaches its best value at n_{Au}/n_{Pd} = 6/4, indicative of the excellent synergistic effects between Au and Pd for FA decomposition. Additionally, rGO is an excellent support for the catalysis, which is demonstrated by the quite low activities of their support-free counterparts (Fig. S14, ESI†).

Further, the increased hydrogen release rates from the FA/SF systems have been observed at elevated temperatures (Fig. S15, ESI†). The activation energy (E_a) in this process is calculated to be 39.77 kJ mol⁻¹, which is lower than most of the reported heterogeneous and homogeneous systems for FA dehydrogenation. After the catalytic reaction, the catalyst was recollected by centrifugation, washed with water and recycled for further use. As shown in Fig. 4b and Table S2 (ESI†), no significant loss in activity was observed over five cycles. The structure and
size distribution of metal NPs after the catalytic reactions have been found to be identical to those of the pristine catalyst (Fig. S16 and S17, ESI†). In contrast, the CO poisoned (Co3)Au0.6Pd0.4/rGO, prepared by exposing the catalyst to a CO atmosphere showed no catalytic activity (Fig. S18, ESI†). It is clear that the (Co3)Au0.6Pd0.4/rGO catalyst possesses high selectivity, durability and stability during the FA dehydrogenation.

In conclusion, a (Co3)Au0.6Pd0.4/rGO catalyst has been synthesized using a simple non-noble metal sacrificial method. The obtained catalyst shows efficient catalytic activity for hydrogen generation from the FA/SF system with the highest TOF of 4840 h⁻¹ at 323 K and possesses high selectivity, durability and stability over 5 cycles. It is believed that the way to the utilization of FA as a hydrogen carrier will be paved by the practical use of the presently provided catalyst.

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Notes and references