Peptide templated AuPt alloyed nanoparticles as highly efficient bi-functional electrocatalysts for both oxygen reduction reaction and hydrogen evolution reaction

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ABSTRACT

Peptide driven metallic nanomaterials fabrication under mild conditions has become a significant research thrust recently, thanks to its capability to generate multifunctional materials with tunable electronic and catalytic properties. Herein, peptide Z1 is employed to fabricate a series of AuPt alloyed nanoparticles with different Au-to-Pt ratios. The morphology is examined by electron microscopic techniques, and well-defined peanut-shape is observed for Au33Pt67 while spherical particles are obtained for the other samples. The AuPt alloys demonstrate effective electrocatalytic activity toward both oxygen reduction reaction (ORR) and hydrogen evolution reaction (HER) and Au33Pt67 exhibits the best bifunctional electrocatalytic performance. Its ORR activity is superior than commercial Pt/C in alkaline media, verified by the positive onset potential (+1.02 V vs. RHE) and large diffusion-limited current density, while its HER activity is comparable with Pt/C, evidenced by the small overpotential at 10 mA cm−2 (−171 mV vs. RHE) and low Tafel slope. Moreover, Au33Pt67 exhibits markedly higher long-term durability than Pt/C for both ORR and HER. The excellent activity and stability for ORR and HER are ascribed to Au alloying Pt induced synergistic effects. The results highlight the advantages of using peptide to fabricate bi-functional or multiple functional electrocatalysts with desirable activity and robust stability.

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1. Introduction

Bimetallic nanoparticles have been continuously gaining increasing research attentions as catalysts and electrocatalysts, mainly thanks to their distinct physical and chemical properties (optical, electronic, magnetic, and so on) as well as the alloying effects [1–3]. Compared with their counterparts monometallic nanoparticles, the synergistic effects during the catalytic process from the bimetallic nanoparticles have empowered them enhanced catalytic activities [4]. Note that, the catalytic properties of the catalyst are highly dependent on their surface structure, and the introduction of another metallic component can significantly modify the size, shape/morphology and surface composition of the catalyst, which could further facilitate fine-tuning of their catalytic performance. For instance, PtM (M=Fe, Co, Ni, Pd, Au etc.) alloys have demonstrated superior catalytic activity than Pt nanoparticles as electrocatalysts toward ORR and HER [5–7]. It is worth noting that, ORR is a key electrochemical process which determines the energy conversion efficiency of fuel cells and metal-air batteries [8–10], while HER is critically important for the production of clean and sustainable energy [11,12].

Among a series of metals to alloy Pt, Au is of particular interest to us in terms of its inertness in the bulk state and high catalytic activity at the nanoscale [13–15]. Previously, AuPt nanostructures...
have already been proved and served as perfect synergistic platforms for catalyzing either ORR or HER or both reactions. Luo et al. examined the catalytic activity and bimetallic composition of carbon supported gold platinum nanoparticles toward ORR, and found the maximized mass activity was achieved with the sample of 60–80% Au [16]. In 2010, Han group reported the synthesis of bimetallic hetero-nanostructure consisting of a dendritic Pt shell and structured Au core by seed growth method, and higher activity and durability than Pt nanoparticles toward ORR were achieved [17]. Moreover, Au–Pt alloyed nanowire networks [18], AuPt alloyed flower-like assembly nanochains [19], Pt–Au superlattice arrays [20], porous AuPt nanoparticles [21] and other AuPt nanostructures have been recorded to possess enhanced ORR activity and stability. In addition, Shao et al. reported the direct synthesis of core-shell AuPt@Pt nanocrystals supported on reduced graphene oxide (RGO) which exhibited enhanced electrocatalytic performance toward HER [22], while such excellent HER activity can also be obtained from RGO supported bimetallic AuPt nanodendrites by Wang and co-workers [23]. Recently, Feng et al. developed an ultrasonication-assisted wet-chemical fabrication of uniform AuPt nanodendrites, and the as-prepared architecture displayed remarkable catalytic performance toward both ORR and HER than commercial Pt black and Pt/C [24].

Since the catalytic performance is closely associated to the surface structure/composition of the catalyst, to achieve better performance, precise control or subtle modification of the surface structure/composition has arisen as the grandest challenge for bimetallic nanoparticles as electrocatalysts. To fabricate alloyed nanoparticles with well-defined structures, organic molecules [2], dendrimer [25], biomolecules including protein [26], DNA [27] especially peptide [28,29] have all been extensively and intensively employed as ligand/template to tune/control the surface structure. Peptide-based approach exploits such a new avenue for fabricating multifunctional nanomaterials under mild conditions based on the following rationales: First of all, the preparation process was normally conducted in water without using organic solvents at room temperature, almost zero-energy input, and more importantly, environmentally friendly; Secondly and more significantly, these peptide sequences hold specific binding affinities for target metal substrates, and they can direct the nuclei growth and nanoparticle formation hence precisely control the size, shape/morphology, composition as well as subtle surface structure of the metal nanomaterials [26,28,30]. Bedford et al. successfully examined the nanoscale surface segregation of AuPd bimetallic nanoparticles, and surface-dependent catalytic activities were achieved for methanol oxidation reaction [31]. Kim group reported the fabrication and catalytic response to ORR of surface-composition-controlled AuPt bimetallic nanoparticles on carbon nanotubes [32]. Recently and notably, our group also successfully demonstrated the facile fabrication of peptide A4 based AuAg alloyed nanoparticle networks [33] and FlgA3 templated AuPd bimetallic spherical nanoparticles [34], and both bimetallic systems exhibited excellent electrocatalytic activity toward ORR.

Herein, we report a facile fabrication of Z1 templated AuPt bimetallic nanoparticles and their electrocatalytic performance for both ORR and HER. Peptide Z1, with a sequence of KHKHW, has been found as a multi-material binder to Au, ZnS and CdSe by using a yeast surface display system by Belcher group in 2005 [35]. In 2013, Knecht group quantified the binding affinity (Gibbs free energy ΔG) of Z1 to Au surface through quartz crystal microbalance [36], and Z1 templated Au nanoparticles also demonstrated effective catalytic activity toward 4-nitrophenol reduction [37]. In this study, peptide Z1 was employed as a template to stabilize the AuPt alloyed nanoparticles with different Au-to-Pt ratios through a facile wet chemical approach. Electron microscopic studies revealed the well-defined peanut shape of the Au33Pt67 sample, while spherical particles were observed for the other samples in the series. The as-prepared alloys exhibited effective activity toward both ORR and HER. The Au33Pt67 sample showed the best bi-functional activities in alkaline electrolyte with ORR performance superior than Pt/C and HER activity close to Pt/C. Intriguingly, the long-term stability of the Au33Pt67 sample was drastically higher than that of Pt/C in both ORR and HER performances.

2. Materials and methods

2.1. Chemicals

Peptide Z1 (KHKHW, >95%, Top-Peptide, Shanghai, China), Hydrogen tetrachloroauric acid (III) trihydrate (HAuCl4·3H2O, 98%, Energy Chemicals, Shanghai, China), Chloroplatinum acid hexahydrate (H2PtCl6·6H2O, 99%, Energy Chemicals, Shanghai, China), Sodium borohydride (NaBH4, 98%, Aladdin Industrial Corporation, Shanghai, China), Nafion (5 wt%, Alfa Aesar) and Pt/C (20 wt %, Alfa Aesar). All chemicals were used as received without further purification, and ultrapure water (18.2 MΩ cm) was employed in all experiments.

2.2. Synthesis of Z1 templated AuPt nanoparticles

The AuPt alloyed nanoparticles have been fabricated by a modified procedure in our previous reports [33,34,38]. For all the samples, the metal-to-peptide ratio was kept as 30:1, while the Au-to-Pt ratios were set as 100: 0, 83: 17, 67: 33, 50: 50, 33: 67, 17: 83 and 0: 100. The Au33Pt67 sample was prepared as follows. Briefly, 200 μL Z1 aqueous solution (1 mM) was diluted into 1.53 mL H2O, followed by the addition of a mixture of 50 μL HAuCl4 (40 mM) and 100 μL H2PtCl6 (40 mM) aqueous solution under slight agitation. The mixture was incubated for 20 min. Then 120 μL freshly prepared NaBH4 (0.1 M) aqueous solution was added rapidly, and the solution turned from yellow to black immediately. The solution was kept stirring at least 1 h at ambient temperature. The other samples were synthesized in a similar manner by changing the volume of the HAuCl4 and H2PtCl6 solutions correspondingly.

2.3. Characterizations

The crystal structure of the samples was measured by X-ray diffraction (XRD) using a Bruker D8 diffractometer with Cu Kα radiation (λ = 0.1541 nm). The composition and valence state of the nanoparticles were analyzed by X-ray photoelectron spectroscopy (XPS) with an Escalab 250 photoelectron spectrometer (Thermo Fisher Scientific, USA). The UV−visible absorbance test was carried out on a Shimadzu 2600/2700 UV−visible scanning spectrophotometer with a 1 cm quartz cuvette. The size and morphology of the samples were characterized by a high-resolution transmission electron microscope (JEOL TEM-2010) coupled with an energy dispersive X-ray spectroscopy (EDS) system. The subtle surface microstructure of the Au33Pt67 sample was examined by a high angle annular dark-field (HAADF) scanning transmission electron microscope (STEM) at an acceleration voltage of 200 kV. The elemental distribution in the Au33Pt67 sample was observed by energy dispersive X-ray spectroscopy (EDS) mapping.

2.4. Electrochemical measurements

The ORR measurements were conducted on a CHI 750e electrochemical workstation (CH Instruments Inc.) in both 0.1 M KOH and 0.1 M HClO4 aqueous solution at room temperature with a three-electrode system. The system including a glassy carbon-disk
electrode (diameter 5 mm, geometrical area of 0.196 cm², Pine Instrument Inc., RRDE collection efficiency is 37%) as the working electrode, a AgCl/Ag electrode in 0.1 M KOH and a saturated calomel electrode (SCE) in 0.1 M HClO₄ as the reference electrode and a platinum wire as the counter electrode. An appropriate amount of catalyst aqueous solution containing 20 µg solution was dropwise cast onto the glassy carbon disk electrode after cleaned with polishing with aqueous slurries of 0.2 µm alumina powders on a polishing microcloth, and dried at room temperature. For commercial Pt/C (20 wt %), 2 mg Pt/C was well-dispersed in 1 mL ethanol and 10 µL of the ink was cast onto the glassy carbon disk electrode and dried at ambient temperature. Subsequently, 10 µL diluted Nafion solution (20 µL, Nafion in 1 mL ethanol) was dropped on the dried samples and Pt/C catalyst. Cyclic voltammetric (CV) curves and linear sweep voltammograms (LSV) with various rotation rates from 100 to 2500 rpm were acquired at a scan rate of 10 mV s⁻¹ in oxygen-saturated 0.1 M KOH and 0.1 M HClO₄ solution. The RRDE voltammetric measurements were conducted at the potential range from −0.03 V to +1.17 V and the ring potential was set as +0.5 V. The stability of the catalysts was tested by chronoamperometric measurements at +0.5 V with a rotation rate of 900 rpm and accelerated durability tests (ADT) from +0.6 V to +1.0 V at 50 mV s⁻¹ in O₂-saturated 0.1 M KOH solution before and after 2500 cycles of potential scans.

The HER measurements were performed on a CHI 750E electrochemical workstation (CH Instruments Inc.) in both 0.1 M KOH and 0.5 M H₂SO₄ aqueous solutions with scan rate of 10 mV s⁻¹ at ambient temperature. In both solutions, a glassy carbon electrode (geometrical area of 0.07 cm²) and a graphite rod was employed as the working electrode and the counter electrode, respectively. In 0.1 M KOH, a AgCl/Ag electrode was employed as the reference electrode, while a SCE was used as reference electrode in 0.5 M H₂SO₄. 1.25 times amount of solution and Pt/C catalysts used in ORR test was cast onto the working electrode in HER. The durability of the Au₃₃Pt₆₇ catalysts was assessed by accelerated linear potential sweeps conducted repeatedly on the electrode at a scan rate of 100 mV s⁻¹.

In all electrochemical tests, the potential was referenced to a Reversible Hydrogen Electrode (RHE): \( E_{\text{RHE}} = E_{\text{Ag/AgCl}} + (0.197 + 0.0591 \text{ pH}) \) V, \( E_{\text{RHE}} = E_{\text{SCE}} + (0.24 + 0.0591 \text{ pH}) \) V.

3. Results and discussions

3.1. Electron microscopic image analysis of AuPt bimetallic nanoparticles

Fig. S1 presents the UV–visible absorbance spectra of the peptide Z1 templated AuPt nanoparticles before and after the reduction by NaBH₄. Prior to the reduction, there are two absorbance peaks located at −210 nm and −260 nm, which can be attributed to the ligand-to-metal charge transfer to Au and Pt surface [39,40], respectively. Upon the reduction of NaBH₄, the absorbance arising from peptide coordinated Au³⁺ and Pt²⁺ no longer exist, and the featureless absorbance profile with broad absorption was noted for each material, indicating the formation of nanomaterial [41].

The samples were then examined by electron microscopic techniques. The representative TEM and HR-TEM image of the Au₁₇Pt₈₃ sample were shown in Fig. 1a and b. Peanut shape was observed for the Au₁₃₃Pt₆₇ sample, and some peanuts connect with each other. Such peanut morphology can be more readily identified in HAADF-STEM image inset in Fig. 1c and Black field-TEM in Fig. 1d. Continuous lattice fringes can be easily recognized in the HR-TEM image (Fig. 1b), and the d-spacing of the adjacent fringe is 0.232 nm. Note that, this spacing value is in the between of pure Au (0.234 nm, JCPDS-04-0784) and pure Pt (0.227 nm, JCPDS-04-0802), indicating the formation of AuPt alloyed structure. From the HR-TEM image of a single Au₁₃₃Pt₆₇ particle, the cross-sectional line-scan profile was conducted. As shown as Fig. 1c, the Au element hold almost identical distribution trend with the Pt element, further manifesting the AuPt alloy formation. To closely examine the Au and Pt elemental distribution of the Au₃₃Pt₆₇ sample, elemental energy dispersive X-ray spectroscopic (EDS) mapping was conducted and presented in Fig. 1e–g. It can be noted that, both the Au and Pt species were homogenously distributed and exhibited random elemental inter-mixing. The uniform Au–Pt overlapped pattern further confirms that homogeneous AuPt alloys were obtained.

The typical TEM images of the Au₈₃Pt₁₇, Au₆₇Pt₃₃, Au₅₀Pt₅₀, Au₁₃₃Pt₆₇, and Pt₁₀₀ samples can be found in Fig. S2. For the Pt₁₀₀ sample, bulky aggregates cluster together, while for the Au₁₃₃Pt₆₇ sample, slight aggregation can also be observed. Besides that, well-defined spherical nanoparticles can be easily identified for the other samples, and the average diameter was calculated as 4.05 ± 0.56 nm, 3.46 ± 1.13 nm, 3.45 ± 0.59 nm, 3.72 ± 0.81 nm and 31.66 ± 8.94 nm for Au₈₃Pt₁₇, Au₆₇Pt₃₃, Au₅₀Pt₅₀, Au₁₃₃Pt₆₇, and Pt₁₀₀ (Table S1), respectively.

3.2. XRD and XPS analysis

The structure of the Au₁₀₀, Au₃₃Pt₆₇, Pt₁₀₀ samples was then studied by X-ray diffraction (XRD) measurements. From Fig. 2a, it can be seen that, there are four well-defined peaks located at 2θ = 39.06, 45.39, 66.31, and 80.21, which can be assigned to the (111), (200), (220), and (311) planes of the fcc alloyed structure. In addition, the diffraction peaks are located between the Au₁₀₀ sample (JCPDS-04-0784) and the Pt₁₀₀ sample (JCPDS-04-0802) [20], further implying the alloyed structure was successfully obtained.

The composition and valence state of the series was probed by X-ray photoelectron spectroscopy (XPS). Fig. S3 shows the survey scan spectra of the Au₁₀₀, Au₃₃Pt₆₇ and Pt₁₀₀ samples. From XPS, the Au-to-Pt ratio can be estimated as 30.2: 69.8 for the Au₃₃Pt₆₇ sample, which approximately agrees with the initial loading ratio (1: 2). The core-level XPS spectra of the Au₄f and Pt₄f electrons can be found in Fig. 2b and c. For the Pt₄f electrons, the binding energies at −71.3 eV and −74.7 eV are attributed to Pt₄f₇/2 and Pt₄f₅/2 electrons, and both peaks can be de-convoluted into doublet peaks as well. The strong doublets at the binding energies of 71.1 eV and 74.4 eV are assigned from metallic Pt, while the other doublets at the binding energies of 72.3 eV and 76.1 eV are attributed to Pt(II). Based on the integrated area from the Pt₄f₇/2 electrons, the percentage of metallic Pt and Pt(II) was estimated as 83.6% and 16.4%, indicating that metallic Pt was the main species. Note that, the binding energy of Pt₄f₅/2 electron decreased about 0.2 eV, compared to metallic Pt (74.6 eV) in Pt/C [6,42,43]. However, for the Au₄f spectra, two sharp peaks with binding energies at −87.6 eV and −83.8 eV can be attributed to the Au₄f₇/2 and Au₄f₅/2 electrons, and both binding energy values agree well with bulk gold (Au³⁺, −87.6 eV and −83.8 eV) [44–46], indicating barely binding energy shift. The decrease of the Pt₄f binding energy indicated the charge transfer occurred from Au atoms to Pt atoms, which have been documented in previous reports [20,46,47]. Furthermore, the negative shift of the binding energy suggests the decrease of the d-band center of Pt, which could weaken the surface adsorption capability of poisoning intermediate, hence facilitating the reaction kinetics and significantly enhancing the catalytic activity [47].

3.3. The ORR activity comparison of the series of samples in alkaline media

The as-prepared samples were first subjected to the
electrocatalytic test toward ORR. Table S1 summarizes the ORR activity results of the series of samples. The CV curves of all the samples can be found in Fig. S4a. From the RRDE voltammograms in Fig. 3a, it can be seen that, the Au100 sample exhibited relatively weak ORR activity, besides that, all the samples demonstrated effective activity. Notably, the onset potential and diffusion-limited current density at +0.2 V varied drastically with the change of Au-to-Pt ratio among the series. As summarized in Table S1, the onset potential first increased then decreased with the decreasing of Au-to-Pt ratio, while besides the Au100 sample, all the samples exhibited at least comparable or more positive onset potential than Pt/C. The Au33Pt67 and Au50Pt50 samples showed the same and most positive onset potential value (1.02 V), much higher than that of Pt/C (0.98 V). Similar change trend was also observed for the diffusion-limited current density with regard to the Au-to-Pt ratio. Besides the Au100 sample, the diffusion-limited current density of the other samples was slightly lower than Pt/C, however, the Au33Pt67 sample exhibited the largest value (5.3 mA cm⁻²), slightly higher than Pt/C (5.2 mA cm⁻²). The above results indicated that the Au33Pt67 sample possessed the best activity among the series, which is further compared with Pt/C in Figs. S4b and 3b. As shown in Fig. S4b, no obvious redox peak was observed for the Au33Pt67 sample and Pt/C in N₂-saturated 0.1 M KOH solution, but a well-defined cathodic peak is clearly observed at ~0.88 V when the 0.1 M KOH solution was saturated with O₂, implying the excellent electrocatalytic activity for ORR. As depicted in Fig. 3b, the Au33Pt67 sample exhibited comparable diffusion-limited current density with Pt/C but much higher onset potential than Pt/C, suggesting higher ORR activity than Pt/C.

Furthermore, based on the RRDE data (Fig. 3b), the electron transfer number (n) and the yield of H₂O₂ in ORR can be calculated through the equations of

\[
n = \frac{4I_d}{I_r/N + I_d}
\]  

\[
H_2O_2\% = \frac{200I_r/N}{I_r/N + I_d}
\]

where \(I_d\) is the disk current, \(I_r\) is the ring current, and \(N\) is the RRDE collection efficiency (0.37). The calculated results for the Au33Pt67 sample and Pt/C can be found in Fig. 3c. In the potential range of +0.2 V to +0.80 V, the electron transfer number was 3.95–3.99 and 3.96 to 3.99 for the Au33Pt67 sample and Pt/C, respectively, both values are very close to 4, indicating a 4-electron transfer pathway was adopted. Correspondingly, the H₂O₂ yield was 0.39%–2.48% and 0.39%–1.64% for the Au33Pt67 sample and Pt/C, both of them were less than 5%, suggesting negligible byproduct was produced during the catalytic process. The remarkable high electron transfer number and extremely low H₂O₂ yield indicated both of them can catalyze electroreduction of oxygen very efficiently. Moreover, the \(n\) and H₂O₂ yield of the Au33Pt67 sample was comparable with Pt/C, together with the higher onset potential, comparable diffusion-limited current density, all the findings attest that the Au33Pt67 sample possessed superior ORR activity than commercial Pt/C.

Next, the reaction kinetics of the Au33Pt67 sample was disclosed and compared with Pt/C. From Figs. S4c and d, it can be observed that the voltammetric currents of the Au33Pt67 sample and Pt/C increased with the increasing of the electrode rotation rate for both samples. In addition, the corresponding Koutecky-Levich (K-L) plots of the Au33Pt67 sample and Pt/C can be found in Fig. S5. A great linearity was obtained with a quite consistent slope for both samples, implying a first reaction-kinetics with regard to the oxygen concentration in the solution. The K-L plots of the other samples in the series can be found in Fig. S6, and a good linearity was exhibited for all the samples in the potential from 0.42 V to 0.63 V.

Another method to calculate the electron transfer number is based on the K-L theory, specifically on Equation (3) [48,49], which describes the current density behavior on RDE.

\[
\frac{1}{j} = \frac{1}{j_k} + \frac{1}{j_a} = \frac{1}{B^{1/2}} + \frac{1}{j_k}
\]  

where \(j\), \(j_k\), and \(j_a\) are the measured, kinetic-limited, and mass
transfer-limited current densities, respectively. $j_k$ is assumed to be a constant at a certain potential, $j_f$ is proportional to the square root of angular velocity ($\omega$) of the RDE. The proportionality coefficient (B) is

$$B = 0.62D^{2/3}v^{-1/6}nF C^*$$

where $n$ is the electron transfer number, $F$ is the Faraday constant (96,485 C mol$^{-1}$), $v$ is the kinematic viscosity (0.01 cm$^2$ s$^{-1}$), $C^*$ is the bulk concentration of O$_2$ ($1.2 \times 10^{-6}$ mol cm$^{-3}$), $D$ is the diffusion coefficient of O$_2$ in 0.1 M KOH (1.9 $\times$ 10$^{-5}$ cm$^2$ s$^{-1}$). Based on the above two equations, the electron transfer number of the Au$_{33}$Pt$_{67}$ sample at +0.45 V is calculated as 3.95, which is in good accordance with the $n$ value (3.96) that obtained from RRDE measurements.

Fig. 3d displays the Tafel plots of the Au$_{33}$Pt$_{67}$ sample and Pt/C. It has been well documented that the Tafel slopes are typically found at 60 or 120 mV dec$^{-1}$ for ORR, where 60 mV dec$^{-1}$ corresponds to a pseudo-two-electron process as the rate-determining step [6,15,43] and 120 mV dec$^{-1}$ suggests that the rate-determining step is probably a first-electron transfer to oxygen molecule, while the subsequent reduction and O=O bond-breaking steps are relatively facile [50–52]. In the high overpotential range, the Tafel slope of the Au$_{33}$Pt$_{67}$ sample was calculated as 76.2 mV dec$^{-1}$, close to that of Pt/C (67.8 mV dec$^{-1}$), suggesting a similar reaction mechanism with Pt/C was adopted for Au$_{33}$Pt$_{67}$, that is, the ORR process is a pseudo-two-electron reduction step. While in the low overpotential range, the Tafel slopes of the Au$_{33}$Pt$_{67}$ sample and Pt/C were determined as 119.8 mV dec$^{-1}$ and 120 mV dec$^{-1}$, indicating the ORR reaction rate was dominated by the first electron transfer to oxygen. [10,15,53,54]

### 3.4. Durability comparison of the Au$_{33}$Pt$_{67}$ sample and commercial Pt/C for ORR in alkaline media

The long-term durability of the Au$_{33}$Pt$_{67}$ sample was then evaluated and compared with Pt/C by the chronoamperometric measurements. As displayed in Fig. 4a, after continuous operation of about 2,000 s, the cathodic current retained about 83.2% of its initial value for Pt/C, in contrast, the Au$_{33}$Pt$_{67}$ sample only exhibited a loss of 9.3% (90.7% retained), suggesting a much higher stability than that of commercial Pt/C. Accelerated durability test (ADT) is another important approach that can assess the long-term durability of the catalysts, and Fig. 4b and c present the ADT results of Pt/C and the Au$_{33}$Pt$_{67}$ sample by cycling the catalysts over the potential range from +0.2 V to +1.1 V in an O$_2$-saturated 0.1 M KOH solution. After 2500 cycles, the half-wave potential shifted negatively about 18 mV for Pt/C, while only 9 mV shift was observed for the Au$_{33}$Pt$_{67}$ sample. The results further confirmed that Au$_{33}$Pt$_{67}$ possessed markedly superior long-term stability than Pt/C in the electrocatalytic process of ORR in alkaline media.

### 3.5. The ORR activity and stability comparison of the Au$_{33}$Pt$_{67}$ sample with Pt/C in acid media

Moreover, the ORR activity of the Au$_{33}$Pt$_{67}$ sample was also evaluated and compared with Pt/C in acid electrolyte of 0.1 M HClO$_4$. Fig. 5a illustrates the polarization curves of the Au$_{33}$Pt$_{67}$ sample and Pt/C in O$_2$-saturated 0.1 M HClO$_4$ solution at a rotation rate of 1600 rpm. The onset potential of the Au$_{33}$Pt$_{67}$ sample was estimated as 0.98 V, which slightly higher than that of Pt/C (0.96 V). Meanwhile, the diffusion-limited current density for the Au$_{33}$Pt$_{67}$ sample was 5.7 mA cm$^{-2}$, close to that of Pt/C (6.1 mA cm$^{-2}$). These results suggest that the Au$_{33}$Pt$_{67}$ sample possessed comparable ORR activity with Pt/C in acid media. Additionally, the LSV polarization curves at different rotation rates as well as the corresponding K-L plots in 0.1 M HClO$_4$ for the Au$_{33}$Pt$_{67}$ sample and Pt/C can be found in Fig. S7. It can be noted that, great linearity with consistent slope was observed for the Au$_{33}$Pt$_{67}$ sample and Pt/C, suggesting a first reaction kinetics with respect to oxygen concentration in 0.1 M HClO$_4$ for both catalysts. The Tafel plots were then extrapolated and shown as the inset graph in Fig. 5a. The Au$_{33}$Pt$_{67}$ sample exhibited a Tafel slope of 75.6 mV dec$^{-1}$, very close to that of Pt/C, indicating the ORR also proceed with a 4-electron pathway as Pt/C. Furthermore, after continuous operation of 10,000 s, the current remained 70.2% of its initial value (Fig. 5b), about 10% higher than that of Pt/C (60.2%), suggesting a superior long-term durability than Pt/C in acid media.
3.6. The HER activity comparison of the series of samples with Pt/C

In addition to ORR, the electrocatalytic HER activity of the AuPt bimetallic series was also tested in comparison with Pt/C. Fig. 6a presents the polarization curves for HER in 0.1 M KOH solution. Pt/C exhibited the best HER activity, with a nearly zero onset potential. Interestingly, negligible HER activity was observed for the Au100 sample. Other than that, the AuPt alloys and the Pt100 sample displayed effective HER activity, however, the activity varied with the change of Au-to-Pt ratio. To afford a current density of 20 mA cm$^{-2}$, the required overpotential was -217 mV for Au33Pt67, -204 mV for Au50Pt50, -172 mV for Au67Pt33, -171 mV for Au50Pt50, -186 mV for Au17Pt83, -227 mV for Pt100, and -165 mV for Pt/C, respectively. The Au33Pt67 sample displayed the best HER activity, as the smallest overpotential was observed. Such overpotential value is also close to that of Pt/C, indicating a comparable HER activity. Tafel slope is an inherent property for an electrocatalyst, and it can be obtained by fitting the linear portions of the polarization curves with the Tafel equation:

$$\eta = b \log j + a$$

(5)

where $j$ is the current density and $b$ is the Tafel slope. The Tafel slopes of the Au33Pt67 sample and Pt/C was calculated as 81 mV dec$^{-1}$ and 73 mV dec$^{-1}$ (Fig. 6b), further confirming that the HER activity of the Au33Pt67 sample was close to Pt/C. Then the HER
stability of the two samples were probed and compared in alkaline electrolyte. Fig. 6c and d present the polarization curves of Pt/C and the Au33Pt67 sample before and after 4000 cycles of potential scans. By using a current density of 10 mA cm$^{-2}$ as a metric, 29 mV potential shift was observed for Pt/C, while in contrast, after 4000 cycles, almost no potential shift can be found in the Au33Pt67 sample, suggesting a markedly superior stability in alkaline media.

Furthermore, the HER activity in acid electrolyte of 0.5 M H$_2$SO$_4$ of the AuPt alloys was also tested and the results are shown in Fig. S8. From the polarization curves in Fig. S8a, the Au100 sample barely exhibited HER activity, and as expected, Pt/C displayed the best activity. To afford a current density of 20 mA cm$^{-2}$, the required overpotential was 53 mV for Au50Pt50, 49 mV for Au50Pt50, 44 mV for Au100Pt50, 61 mV for Pt100, and 33 mV for Pt/C. Au100Pt50 demonstrated the best HER activity in acidic media as well among the series. The calculated Tafel slope was 34 mV dec$^{-1}$ for Au33Pt67 and 29 mV dec$^{-1}$ for Pt/C (Fig. S8b), and once again confirmed that Au33Pt67 exhibited comparable HER activity with commercial Pt/C in acid electrolyte. Finally, the stability of Au33Pt67 was tested and compared with Pt/C in 0.5 M H$_2$SO$_4$. The polarization curves before and after 4000 cycles of potential scan for Pt/C and Au33Pt67 can be found in Fig. S8c and d. To afford a current density of 20 mA cm$^{-2}$, the required overpotential increased about 115 mV while only 19 mV potential increase was observed for Au33Pt67. The results strongly attest that Au33Pt67 possessed much higher stability than
Pt/C for HER in acid media as well.

The enhanced long-term stability for both ORR and HER of the Au33Pt67 sample is quite encouraging, and such robust stability might be attributed to the following factors: 1. The alloying effects in the product. It has been widely acknowledged that, the introduction of another metal component can tune and/or modify the surface geometric and electronic structure of the metal architectures, hence significantly improve the stability of the alloyed nanoparticles [5,8]. 2. Peptide Z1 serves as the stabilizer and capping agent, and it can prevent the strong metal-metal interaction for aggregation and/or coalescence. [35]

3.7. The ORR and HER activity comparison with other AuPt nanostructures

The remarkably excellent activity toward both ORR and HER from the Au33Pt67 sample was higher than recently reported AuPt alloys, Au@Pt core-shell structures and other AuPt bimetallic nanostructures either in alkaline or acid electrolyte. The comparison results are compiled in Table S2. For examples, in 0.1 M KOH for ORR test, the onset potential was 0.90 V, 0.96 V and 0.91 V for PtAu superlattice arrays [20], AuPt nanodendrites [24] and Pt@Au nanorods on Pyridine Graphene [55], respectively, all of them were lower than that of the Au33Pt67 sample (1.02 V). In addition, when the electrolyte was switched into 0.1 M HClO4, the Au33Pt67 sample exhibited an onset potential of 0.98 V, which is also much more positive than Au75Pt25 alloyed nanodendrites (0.89 V) [56], PtAu/C-250 (0.89 V) [57], as well as Imp-PtAu/C (0.5 M HClO4, 0.87 V) [58]. Furthermore, when employing 0.5 M H2SO4 as electrolyte for HER, to afford a current density of 10 mA cm−2, the required overpotential was 30 mV for the Au33Pt67 sample, lower than that of AuPt nanostructures supported on carbon fiber (235 mV) [59] and AuPt nanodendrites (48 mV) [24], and close to that of AuPdPt-WC/C (25 mV) [60]. It is worth noting that, the Au33Pt67 sample also demonstrated great capability to produce hydrogen in alkaline media, of which rare examples can be found to compare.

The remarkably high catalytic activity toward ORR and HER of the Au33Pt67 sample can be attributed to the Au–Pt alloying induced synergistic effects, namely the geometric effects and the electronic effects. The combination of two metals can lead to formation of hetero-atom bond, hence resulting in generation of new catalytic active sites [2,4], and this is so called geometric effect. Meanwhile, the introduction of Au atoms to Pt nanoparticles (versa vice) can largely alter and modify the electronic structure of the metal nanoparticles, evidenced by the electron transfer phenomenon observed in XPS measurement for the Au33Pt67 sample. Such electronic effect can be crucial for the enhancement of the electrocatalytic activity. Take ORR as an example, Pt has strong binding affinity to oxygenated intermediates, while the introduction of a relatively weak binder (Au) can “dilute” the strong interaction at the atomic level. The weakened binding interaction led to the enhanced activity with a maximum of the Au33Pt67 sample. Moreover, further increase of the Au content or the Pt content can result in the interaction either too strong or too weak, both of which would diminish the ORR activity. The maximum activity of Au33Pt67 sample represents the optimal balance. Such perfect alloying effects can be further evidenced by the HAADF-STEM images with homogeneously well-mixed Pt–Au patterns and well-maintained Pt crystalline. As a note, previous studies have shown that the bimetallic AuPt synergistic effects can significantly promote the electrokinetics of ORR and HER. [19,20,23,24]

4. Conclusions

In summary, a series of bimetallic AuPt alloyed nanoparticles were fabricated by employing Z1 as template/ligand. Au33Pt67 exhibited well-defined peanut shape, while spherical particles were obtained for the other samples in the series. The series of AuPt bimetallic nanoparticles demonstrated effective electrocatalytic activities toward both ORR and HER. Au33Pt67 exhibited the best bifunctional catalytic activities among the series, of which the ORR activity was superior than commercial Pt/C while the HER activity is close to Pt/C in alkaline media. Such remarkable bifunctional electrocatalytic activities are ascribed to the Au alloying Pt induced synergistic effects. Moreover, Au33Pt67 demonstrated markedly higher long-term stability than Pt/C in both ORR and HER. The findings can shed light upon further rational design and engineering of peptide templated bimetallic nanomaterials with premium activity and robust stability for dual or even multiple electrocatalytic reactions.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.electacta.2017.11.057.

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