Journal of Materials Chemistry A



PAPER

View Article Online View Journal | View Issue



Cite this: J. Mater. Chem. A, 2023, 11, 14043

Received 20th March 2023 Accepted 6th June 2023

DOI: 10.1039/d3ta01668d

rsc.li/materials-a

Revealing surface fine structure on PtAu catalysts by an in situ ATR-SEIRAS CO-probe method†

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The electrochemical performance of Pt-based catalysts depends on their surface structure. Nevertheless, it is still a challenge to investigate their intricate surface-active sites. Here, PtAu films were utilized as model catalysts to understand the surface fine structure and structure-activity relationship. Significantly, the in situ attenuated total reflectance-surface enhanced infrared absorption spectroscopy (ATR-SEIRAS) CO-probe method was established to reveal the surface-active sites of PtAu catalysts. By employing the second derivative spectrum (SDP) method and Gaussian fitting for the infrared spectra of the CO oxidation reaction (COR), three surface configurations, the PtisoAu, PtadiAu site and PtrichAu site, were unveiled. Additionally, the activity order and reactive species of the COR oxidation peaks were clarified by correlating CO_L intensities with the COR current curves. More importantly, this in situ ATR-SEIRAS COprobe strategy can be extended from PtAu model films to PtAu nanoparticles.

Introduction



Dr Bin-Wei Zhang is an Associate Professor at Center of Advanced Energy Technology and Electrochemistry, Institute of Advanced Interdisciplinary Studies, and School of Chemistry and Chemical Engineering, Chongqing University. He received his bachelor's and master's degrees from Xiamen University in 2012 and 2015, respectively. In 2019, he obtained his PhD degree from Institute for Superconducting and

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Fuel cells, as energy conversion devices, have attracted significant attention due to their high efficiency and environmental friendliness.1-3 Pt-based catalysts usually show excellent performance in fuel cells; however, the high price and scarcity of Pt hinders their widespread application.4,5 To reduce the utilization of Pt and still maintain its outstanding activity, Pt alloys have been extensively studied.6-8 As electrochemical reactions usually take place on the surface of catalysts, it is important to study the surface Pt reactive sites of the Pt alloy. 9-12 X-ray absorption spectroscopy (XAFS) and spherical-aberrationcorrected transmission electron microscopy (SACTEM) have advanced in study of coordination environments and surface atomic structure, but it is still challenging to identify the practical active sites because of the lack of effective research tools to detect intermediate species. 13-15 Therefore, it is highly demanded to develop an effective method to clarify the surface structure-activity relationship of Pt-based catalysts. 16-19

PtAu catalysts have been considered as promising candidates owing to their high activity and chemical stability towards

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[†] Electronic supplementary information (ESI) available: The characterization and cyclic voltammetry of electroless Au films. Additional in situ ATR-SEIRAS spectra, second derivative spectra and Gaussian fitting peaks of COR on Pt_{1.5}Au_{0.5} and Pt_{0.5}Au_{1.5} surfaces. See DOI: https://doi.org/10.1039/d3ta01668d

electrocatalytic reduction reactions and various organic smallmolecule electro-oxidation reactions, including HCOOH, methanol, and ethanol.20-23 For example, PtAu catalysts have been employed as efficient catalysts for oxygen reduction reactions.^{20,24} As these electrocatalytic reactions usually take place on the surface of the electrocatalysts, their surface structure plays an important role in the performance. 10,25,26 Although great efforts have been made to improve their activity by modifying the electronic and geometric structure of the Ptbased catalysts, the intricate surface-active sites of these PtAu catalysts still needed to be studied further. Recently, we have developed CO as a probe molecule to characterize the d-band center of Pt-based catalysts.^{27,28} Although Au does not have a beneficial effect on anchoring either COad or OHad, it will modify the electronic and surface-site structure of the PtAu alloy, which influence the electrochemical performance towards CO electro-oxidation. Additionally, the CO adsorption and activity on Au sites are much weaker than on Pt sites, which is ideal for investigating the influence of the electronic and surface-site structure of the PtAu catalyst. Nevertheless, the surface fine structure and electronic structure of PtAu catalysts are unknown. Thus, it is very necessary to reveal the surfaceactive sites of PtAu catalysts, which can rationally guide the design of PtAu alloy.

In situ attenuated total reflectance-surface enhanced infrared absorption spectroscopy (ATR-SEIRAS) has recently been demonstrated to be a powerful tool to detect surface-adsorbed intermediate species and monitor reaction mechanism. Due to the limited penetration depth of the hidden wave, 100 nm in general, the in situ ATR-SEIRAS method was high surface sensitivity. 29-32 Nevertheless, using the conventional ink drip method, the catalyst layer thickness of carbon nanomaterials is close to the micron level, which may lead to a worse infrared signal ratio due to exceeding the detection depth. Therefore, the development of an efficient strategy to study the surface fine structure of nanocatalysts is required. The electrodeposition and underpotential deposition methods are simple and useful ways to construct nanoscale surface structure by controlling quantity of electricity. 33,34

In this work, the electrochemical in situ ATR-SEIRAS and CO probe molecule method were employed to investigate the surface fine structure on PtAu catalysts from model films to nanocatalysts. PtAu model films with various surface ratios were prepared using the electrodeposition strategy, which presented an enhanced infrared signal for CO intermediate species. Moreover, the second derivative spectrum (SDP) method and corresponding Gaussian fitting of in situ ATR-SEIRAS for CO oxidation were developed to locate the peak positions. Based on the structure characterization and surface content, there are three CO_L sites in the PtAu film catalysts: surface Pt atoms completely isolated by Au atoms (Pt_{iso}Au site), adjacent Pt active sites (PtadiAu site), and abundant continuous Pt atoms without Au atoms (Pt_{rich}Au site). Moreover, the activity order of the different sites was identified. Significantly, a consistent appearance and results were also verified in PtAu nanoparticles, which clarified the surface fine configurations on the nanocatalyst.

2. Experimental

2.1 Chemicals and reagents

 $\rm H_2PtCl_6\cdot H_2O$ (wt% Pt: 37.5%), $\rm HAuCl_4\cdot 4H_2O$ (99.9%, trace metal basis), and $\rm HClO_4$ (70% aqueous solution, ACS) were purchased from Aladdin and used without purification. A silicon wafer (99.9%, 10 mm × 10 mm × 0.5 mm) was purchased from TEXCHEM. A ladder-type silicon prism was purchased from Jing Xaun Guang Dian (Shanghai, China). Ar (99.99%) and CO (99.999%) were purchased from Linde Industrial Gas. Deionized water used in electrochemical studies and device cleaning was from a Millipore Milli-Q system (resistivity 18.2 MΩ cm $^{-1}$).

2.2 Structure characterization

The surface morphologies of the electroless Au film and electrodeposited PtAu films on a silicon wafer were studied using field-emission scanning electron microscopy (SEM) (HITACHI-4800, HITACHI, Japan). The crystalline phases of the electrodeposited PtAu films were analyzed using powder X-ray diffraction (XRD) patterns using a Rigaku IV XRD with Cu Kα radiation ($\lambda = 1.54$ Å, step size: 0.02 for PtAu films on silicon wafer; current: 30 mA; and voltage: 40 kV). Surface electronic structures of the electrodeposited PtAu films were analyzed using an ESCALAB 250 XI X-ray photoelectron spectrometer (XPS, Thermo Fisher Scientific) with monochromatic Al Ka radiation (1486.6 eV; current: 12 mA; voltage: 12 kV; diameter beam spot: 500 µm), and the binding energies were calibrated with reference to the C 1s peak at 284.6 eV. The morphology of the PtAu nanomaterials was studied using transmission electron microscopy (TEM, TECNAI F20). The STEM imaging and EDS elemental mapping were conducted using a FEI Themis Z microscope equipped with a probe corrector operated at 300 kV.

2.3 Preparation for electroless plated Au

The configurations and characteristics of the electrochemical IR cell were described in a previous work.35 The method of secondary electroless gold plating on the silicon prism/wafer was detailed in previous reported work: a silicon prism/wafer was ground with 1 μm, 0.3 μm and 0.05 μm Al₂O₃ and cleaned using ultrasonication with water and acetone. After treatment with piranha solution $(H_2SO_4:30\%H_2O_2=7:3)$ for 90 min, the plating bath was dropped on the silicon prism/wafer after being soaked in 40 wt% NH₄F solution for 90 s, and electroless plating was conducted for 150 s at 60 °C. Aqua regia dissolved the unstable gold film and it was gilded again. The electrodeposited PtAu model films on the Au-coated silicon prisms were used for electrochemistry measurement and in situ ATR-SEIRAS measurements. The electrodeposited PtAu model films on the Au-coated silicon wafers were used for XRD and XPS measurement. The chemically synthesized PtAu nanocatalysts were used from our previous work.¹⁷ The cyclic voltammetry curve and structural characterization of the Au film on the Si prism are shown in Fig. S1.†

2.4 Electrochemistry measurements

The PtAu model films with different surface ratios were obtained by electrodepositing different concentration ratios of H₂PtCl₆ and HAuCl₄ at 0.30 V (vs. RHE) under Ar saturation (molar ratios of electrodeposition solution: H_2PtCl_6 : $HAuCl_4$ = 9:1, 3:1, 1:1, 1:3, 1:9, total concentration: 2 mM, supporting electrolyte: 0.1 M HClO₄). The electrodeposited PtAu films on Au-coated silicon wafer were used for analysis of structure and morphology. The electrodeposited PtAu on Au-coated silicon prism and PtAu nanocatalysts were used as the working electrodes for cyclic voltammetry, CO stripping measurement and in situ ATR-SEIRAS measurement in 0.1 M HClO4 solution by a potentiostat/galvanostat, 263A model, Princeton Applied Research, with a saturated calomel electrode (SCE) and a graphite rod as the reference and counter electrodes, respectively. The PtAu nanocatalyst was dispersed in 1 mL mixed solution (495 μL isopropanol + 495 μL water + 10 μL 5 wt% Nafion solution) and ultrasonically dispersed for 1 hour. 25 µL ink was dropped on the electroless Au@Si prism and dried in air for in situ ATR-SEIRAS measurement. In this study, all the indicated potentials were corrected to the reversible hydrogen electrode potential (RHE). After CO had been adsorbed for 8 min at 0.1 V and Ar-purged for 15 min, the electrochemistry and electrochemical in situ ATR-SEIRAS experiments of PtAu COR were conducted.

2.5 In situ ATR-SEIRAS measurements

Electrochemical in situ ATR-SEIRAS measurements were conducted using a Nicolet-8700 spectrometer equipped with a liquid-nitrogen-cooled MCT-A detector. A 263A model potentiostat was used to control the electrode potential. The electrodeposited PtAu on an Au-coated silicon prism was electrochemically cleaned until stable in Ar-saturated 0.1 M HClO₄ solution. The multi-step FTIR method (MS-FTIR) was used to collect the spectrum from 0.1 V to 1.3 V at 0.05 V intervals. The spectra were recorded from 4000 cm⁻¹ to 1000 cm⁻¹ with a spectral resolution of 8 cm⁻¹ and a scan number of 200. The spectra were calculated from the relative change in reflectivity ($\Delta R/R$) with eqn (1) below. The $R(E_S)$ and $R(E_R)$ represent the single-beam spectrum recorded at the sample setting potential $E_{\rm S}$ and the reference potential $E_{\rm R}$, respectively. In order to more intuitively display the changes in the spectra, the original infrared spectra were rotated and the direction of peaks were opposite.

$$\frac{\Delta R}{R} = \frac{R(E_{\rm S}) - R(E_{\rm R})}{R(E_{\rm R})} \tag{1}$$

3. Results and discussion

Structural characterization of the PtAu films

The PtAu film model catalysts were prepared through the electrodeposition method at 0.3 V in solutions with different H₂- $PtCl_6$: HAuCl₄ ratios of 1.8: 0.2, 1.5: 0.5, 1: 1, 0.5: 1.5, and 0.2: 1.8 with a total concentration of 2 mM. The total deposited

charge was 0.10C to ensure the same thickness for all PtAu films (Fig. 1a). The obtained samples were labeled as Pt_{1.8}Au_{0.2}, $Pt_{1.5}Au_{0.5}$, Pt_1Au_1 , $Pt_{0.5}Au_{1.5}$, and $Pt_{0.2}Au_{1.8}$, respectively. The cyclic voltammetry curves of the PtAu films with different ratios were shown in Fig. 1b. By integrating the hydrogen underpotential deposition region charge (210 μ C cm⁻²) and Au redox peak charge (340 μ C cm⁻²), the surface compositions of the PtAu films were calculated and were shown in Table S1.† 17,36 The surface ratios of Pt for Pt_{1.8}Au_{0.2}, Pt_{1.5}Au_{0.5}, Pt₁Au₁, Pt_{0.5}Au_{1.5}, and Pt_{0.2}Au_{1.8} were 0.92, 0.86, 0.68, 0.48 and 0.16, respectively.

The powder X-ray diffraction (XRD) patterns of the PtAu films were shown in Fig. 1c and S2.† The magnified band details from 37.00° to 41.00° were shown in Fig. 1c to clarify the structure change. Compared with the Au(111) diffraction peak at 38.18° in the standard cards, the peak of the electroless-plated Au shifted positively to 38.20° due to lattice mismatch between Au and Si. Alloying with Pt, the diffraction peak positively shifted to 38.36° in the Pt_{0.2}Au_{1.8} film. Moreover, new diffraction peaks appeared in the PtAu model films from Pt_{0.5}Au_{1.5} to Pt_{1.8}Au_{0.2}. It gradually positive-shifted from 38.71° to 39.56° in the PtAu alloy phase with Pt ratio increasing. The Pt_{1.8}Au_{0.2} film has the highest peak at 39.56°, which was close to the Pt(111) diffraction peak.17 This result indicated these PtAu model films have various surface structures. Moreover, we used Vegard's law to study the surface structures and the calculated lattice constants of the different diffraction peaks at different angles, as shown in Fig. S2.† 37-39 It could clearly be seen that the lattice constants of the Pt_{0.2}Au_{1.8} model film were well fitted by the linear relation of Vegard's law (red dotted line), indicating the formation of alloy

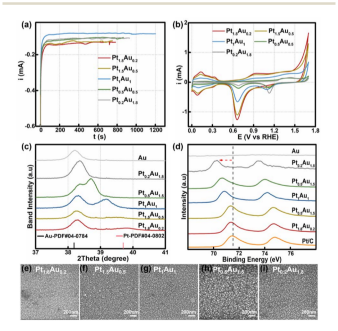


Fig. 1 (a) Electrodeposition current-time curves for the various concentration ratios at 0.30 V. (b) Cyclic voltammetry curves of the PtAu film catalysts. (c) Magnification of the fine XRD patterns of the PtAu film model catalysts from 37° to 41°. (d) Pt XPS spectra of the PtAu films. (e)-(i) SEM images of the PtAu films.

phase. For the $Pt_{0.5}Au_{1.5}$ to $Pt_{1.5}Au_{0.5}$ model films, their high diffraction peaks were close to Vegard's law, also suggesting the formation of an alloy phase. Additionally, their lattice constants at the low-angle diffraction peaks showed mismatch with Vegard's law in Fig. S2d.† These results indicated that there were two alloy phases. Moreover, the lattice constants of the low-angle diffraction peak of the $Pt_{0.5}Au_{1.5}$ to $Pt_{1.5}Au_{0.5}$ model films were close to that of the $Pt_{0.2}Au_{1.8}$ model film, indicating that they may share the same PtAu phase, *i.e.*, Au-rich PtAu alloy phase structure.

X-ray photoelectron spectroscopy (XPS) was used to analyze the surface ratios of the PtAu films, as shown in Fig. 1d and S3.† Their sub-subsurface component analyses were shown in Table S1,† and corresponded with the surface ratios calculated from the cyclic voltammetry results.27 Compared with Pt/C, the Pt binding energy peaks of PtAu film were negatively shifted, and with Pt ratio increasing, the Pt 4f binding energy of the PtAu film became close to that of Pt/C. This phenomenon also could be found in Au XPS results for the PtAu film catalysts in Fig. S3.† The shift in the XPS results is affected by the electronegativity and structure effect. 40-42 As the electronegativity of Pt (2.1) is lower than that of Au (2.3), electrons should transfer from Pt to Au.43 In contrast, the atomic radius of Au is larger than Pt, and thus would generate tensile strain during the alloying of Au with Pt. This result corresponds with XRD results, i.e., the lattice constants gradually increase with Au content increasing. Significantly, the strain effect of PtAu is a long-range interaction, and would reduce the orbital overlap between the Pt atoms and increase the electron energy level of the d orbitals. Therefore, the binding energy of Pt would be negatively shifted and the d-band centre of Pt positively shifted, coinciding with the previous reported work. 44,45 The morphologies of the PtAu films were studied using SEM as shown in Fig. 1e-i. It could clearly be seen that all PtAu surfaces consisted of uniform-size nanoparticles without any special shapes.

3.2 In situ ATR-SEIRAS of COR on PtAu film model catalysts

The COR curves of PtAu film model catalysts were measured in 0.1 M HClO₄, as shown in Fig. 2a. There was one obvious CO oxidation peak at 0.71 V in the Pt electrode.46 However, in the Au electrode, there was no obvious oxidation peak, which may be attributed to CO being unable to stably adsorb on the Au electrode (Fig. S4†). For the PtAu model film catalysts, there were a pair of CO oxidation peaks in the COR curves, which was in agreement with previous works. 47-49 These were usually assigned to the low/high potential oxidation peaks as the preoxidation and post-oxidation peak. A summary of the peak potentials of the pre-oxidation peaks and post-oxidation peaks were shown in Fig. 2b. Compared with those of the Pt electrode, the CO oxidation peaks of the PtAu model films gradually positively shifted. As the Pt ratio was reduced from Pt_{1.8}Au_{0.2} to Pt_{0.5}Au_{1.5}, the pre-oxidation peak shifted positively from 0.73 V to 0.84 V; interestingly, the pre-oxidation COR peak disappeared in the Pt_{0.2}Au_{1.8} film. In contrast, the post-oxidation peak was independent of the Pt content at 1.02 V. These results suggested that there were various reactive sites for CO oxidation on the

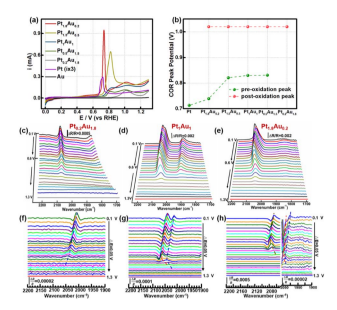


Fig. 2 (a) CO oxidation reaction (COR) measurement of the PtAu film catalysts, Pt electrode and Au electrode in 0.1 M HClO₄; the current of the Pt COR curve has been multiplied by 3 for clear display. (b) Oxidation potential of the pre-oxidation and post-oxidation peaks of (a). (c)–(e) CO oxidation infrared spectra of the PtAu model films: (c) Pt_{0.2}Au_{1.8}, (d) Pt₁Au₁, and (e) Pt_{1.8}Au_{0.2}. (f)–(h) Corresponding second-derivative spectra of (c)–(e) at 2200–1900 cm⁻¹; the right part of (h) is the magnified spectra at 2050–1900 cm⁻¹.

PtAu model films. Additionally, these PtAu film model catalysts showed different activity towards COR: Pt_{1.8}Au_{0.2} (0.95 $mA@0.73 V) > Pt_{1.5}Au_{0.5} (0.65 mA@0.82 V) > Pt_1Au_1 (0.15)$ $mA@0.83 V) > Pt_{0.5}Au_{1.5} (0.095 mA@0.83 V) > Pt_{0.2}Au_{1.8} (0.10)$ mA@0.83 V). This result also indicated that their COR mechanisms were different. In general, the CO_{ad} was oxidized by adjacent adsorbed OH* species (OH*) dissociated from H2O at free sites, i.e. the Langmuir-Hinshelwood mechanism (L-H mechanism).50,51 The emerging free sites originating from the oxidized CO sites could produce new OH* to accelerate COad oxidation. For the continuous Pt surface sites of the model film, the free Pt sites near the Pt-CO_{ad} sites could provide OH* to oxidize other CO_{ad} active sites. It was interesting that with Pt ratio increasing, the pre-oxidation peaks of the PtAu film catalysts were negatively shifted. Moreover, the post-oxidation peaks of these PtAu film catalysts were almost unchanged. This can be attributed to different OH generation potentials.

Electrochemical *in situ* ATR-SEIRAS was employed to investigate the molecular behavior at the electrochemical interface and understand the structure–activity relationship of the PtAu catalyst. The infrared spectra of COR were recorded from 0.10 V to 1.3 V and the reference potential was set at 1.3 V. Pt_{0.2}Au_{1.8}, Pt₁Au₁, and Pt_{1.8}Au_{0.2} were assigned as the research objects, and their spectra were shown in Fig. 2c–e. The bands at 2000–2100 cm⁻¹ and 1900–1800 cm⁻¹ were assigned as linear adsorbed CO (CO_L) and bridge-adsorbed CO (CO_B) on the PtAu films, respectively.⁵² No signal of CO₂ generated by COR on the PtAu model films was detected in Fig. S5†.^{48,53} This was attributed to the generated CO₂ rapidly diffusing to the solution

phase due to the structural features of the electrolytic cell; the in situ ATR-SEIRAS method is insensitive to solution phase species. 16,54,55 In order to investigate the CO oxidation on the Au catalysts, the spectra and peak intensity change of Au-CO also showed that CO was weakly adsorbed on the Au film at 2100 cm⁻¹ and desorbed after Ar purging (Fig. S4†).³² This meant that CO was difficult to stable adsorption on Au site in the PtAu model films after Ar purged. It was interesting that with Au content increasing, the CO_L band red-shifted from 2063 cm⁻¹ to 2008 cm⁻¹. This could be ascribed to the enhancement of the electronic effect of Pt-CO by Au atoms. 47 Au would affect the electronic structure of Pt, which would result in the positive shift of the d-band centre for Pt, and thus enhanced the $d-\pi^*$ feedback effect between Pt and CO.²⁸ The result was conducive to weakening the bond energy of C≡O and coupling with OH* to oxidize CO to CO₂. 51 It was noteworthy that both the Pt_{0.2}Au_{1.8} and Pt_{1.8}Au_{0.2} catalysts presented one CO_L peak in their spectra; in contrast, the Pt₁Au₁ film showed two CO_L peaks. The different CO_L peaks indicated different CO adsorption sites on the surface of the PtAu films, which was confirmed by the COR curves and structure characterization. However, it was difficult to identify the CO_L peak sites of the CO bands. Therefore, it was necessary to clarify the CO peak sites and quantitative band intensity to study the surface fine configurations and activity of the PtAu catalysts.

The SDP method is convenient and effective for locating peak positions by mathematically calculating the extreme points. Hereby, SDP method was employed to identify the CO_L peaks on PtAu catalysts. The second-derivative spectra of these PtAu films were shown in Fig. 2f-h and S6.† In the secondderivative spectra at 0.10 V, Pt_{0.2}Au_{1.8} showed one CO_L peak at 2005 cm⁻¹ (Fig. 2f) and Pt₁Au₁ presented a pair of CO_L peaks at 2038 cm⁻¹ and 2013 cm⁻¹ (Fig. 2g), which were coincident with the original spectra. For both the Pt_{1.5}Au_{0.5} and Pt_{0.5}Au_{1.5} films, two CO_L peaks could be observed in the original spectra and second-derivative spectra (Fig. S4†). It was noteworthy that there were three peaks at 2063 cm⁻¹, 2043 cm⁻¹, and 2005 cm⁻¹ for the Pt_{1.8}Au_{0.2} film in the second-derivative spectrum (Fig. 2h). This suggested that there should be three CO_L peaks in the original unipolar spectra; however, the original spectra of Pt_{1.8}Au_{0.2} only showed two peaks. This indicated that the COR mechanism should be different from the others, which suggested that the surface fine structures of these PtAu films were diverse.

To investigate the COR mechanism on these PtAu films, Gaussian fitting was employed on the ATR-SEIRAS spectra, as shown in Fig. 3 and S7.† The fitted CO_L peaks and a summary of the CO_L fitting curves of Pt_{0.2}Au_{1.8}, Pt_{1.5}Au_{0.5}, and Pt_{1.8}Au_{0.2} at 0.10 V was shown in Fig. 3a-c. For the Pt_{0.2}Au_{1.8} film catalyst, the surface composition was Au-rich (surface ratio of Pt: 16%), and there was only one CO_L peak at 2005 cm⁻¹ (Fig. 3d) without CO_B. Therefore, there may be only one active site in Pt_{0.2}Au_{1.8}, i.e., its surface Pt atoms may be completely isolated by Au atoms (named as PtisoAu). The COad pattern on PtisoAu corresponding to its surface fine structure diagram was shown in Fig. 3d (left figure). For the Pt1Au1 film, there were two COL peaks at 2038 cm⁻¹ and 2013 cm⁻¹. This indicated that there were two

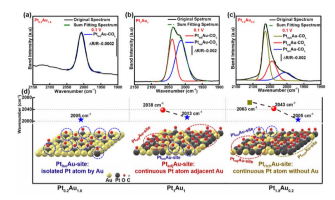


Fig. 3 (a)-(c) Peak fitting of the CO_L bands of the PtAu model films at $0.1\,V$: (a) $Pt_{0.2}Au_{1.8}$, (b) Pt_1Au_1 and (c) $Pt_{1.8}Au_{0.2}$. (d) Wavenumbers of the fitting peak positions and CO_{ad} diagrams of the surface fine structures of the PtAu model films: Pt_{0.2}Au_{1.8} left figure, Pt₁Au₁ middle figure, Pt_{1.8}Au_{0.2} right figure.

surface configurations, in agreement with the XRD results. The low-wavenumber CO_L peak (2013 cm⁻¹) was close to that of the Pt_{iso}Au site in Pt_{0.2}Au_{1.8} (2005 cm⁻¹). This result suggested that the Pt1Au1 film also possessed the PtisoAu sites. The highwavenumber CO_L peak of 2038 cm⁻¹ suggested that it may have another active site. Notably, Pt1Au1 showed a new COB peak at 1856 cm⁻¹, which was usually attributed to the presence of continuous Pt atoms, that was, adjacent Pt active sites (named as Pt_{adi}Au site). Therefore, the Pt₁Au₁ film catalyst has Pt_{iso}Au sites and Pt_{adi}Au sites, and its surface fine structure corresponding with the CO_{ad} diagram was shown in Fig. 3d (middle figure). Pt_{0.5}Au_{1.5} and Pt_{1.5}Au_{0.5} share similar active sites with Pt₁Au₁, as shown in Fig. S7.† For the Pt_{1.8}Au_{0.2} film, three were three CO_L peaks at 2005 cm⁻¹, 2043 cm⁻¹ and 2063 cm⁻¹. The peaks at 2005 cm⁻¹ and 2043 cm⁻¹ could be attributed to CO adsorbed on PtisoAu sites and PtadiAu sites, respectively. The highest-wavenumber CO_L peak (2063 cm⁻¹) was close to that of CO adsorbed on pure Pt, which could be attributed to the high Pt surface ratio (92%).56,57 This indicated that it has a Pt-rich surface configuration, that was, abundant continuous Pt atoms without Au atoms (named as PtrichAu site). This corresponds with the results of the XRD pattern of the Pt_{1.8}Au_{0.2} film. The CO_{ad} diagram and surface fine structure diagram of Pt_{1.8}Au_{0.2} were shown in Fig. 3d (right figure).

To further verify the conformations of the PtAu sites of these PtAu films, the formic acid (HCOOH) electrooxidation reaction (FAOR) was carried out, as shown in Fig. S8.† As the FAOR is sensitive to the active site conformation, it is widely used to characterize the surface structure of Pt-based catalysts.58,59 HCOOH will go through the indirect pathway to produce CO via the dehydration reaction on contiguous Pt atom sites. For the isolated Pt sites, HCOOH is oxidized via the direct pathway without CO and its activity is higher than in the indirect pathway. There were two oxidation peaks at around 0.6 V (peak 1) and around 0.95 V (peak 2) in the forward scan curve in Fig. S8a.† This were attributed to the oxidation of HCOOH to CO/CO₂ and oxidation of CO to CO₂, corresponding to the direct and indirect oxidation pathways, respectively. 60-62 Therefore, the

values of peak 1 and peak 2 could be used to evaluate the surface structure of the PtAu films. The current densities of peak 1 and peak 2 are summarized in Fig. S8b.† It can be clearly seen that the Pt_{0.2}Au_{1.8} model film only presents one peak at around 0.6 V, indicating that HCOOH goes through the direct pathway. This result demonstrated the PtisoAu site of Pt_{0.2}Au_{1.8} model film. For the Pt₁Au₁ model films, the current density of peak 1 was 2 mA cm⁻² and that of peak 2 was 2.3 mA cm⁻², respectively. The increased activity of peak 2 indicated that it possessed the indirect pathway and direct pathway towards FAOR. This suggested continuous Pt sites, i.e., PtadiAu sites, for the Pt1Au1 model films. When the Pt content was increased to give the Pt_{1.8}Au_{0.2} model films, the current density of peak 2 (1.5 mA cm⁻²) was higher than peak 1 (1.2 mA cm⁻²). This was beneficial to generate CO intermediate species via the indirect pathway due to the greater amount of continuous Pt atom sites, corresponding to the PtrichAu sites in the structure diagram of the Pt_{1.8}Au_{0.2} model films.

3.3 Surface fine structure and CO oxidation activity

To further study the relationship between the surface-active sites of the PtAu films and the CO oxidation activity, the SDP method located peak position and Gaussian fittings of the various potential $\rm CO_L$ peaks were shown in Fig. 4 and S9–S11.† The fitted spectra at a potential interval of 0.10 V and the changes in the $\rm CO_L$ integrated intensities were shown in Fig. S9–S11.† As the onset potential of CO was at 0.1 V, the change in the CO integrated intensity and COR curves of

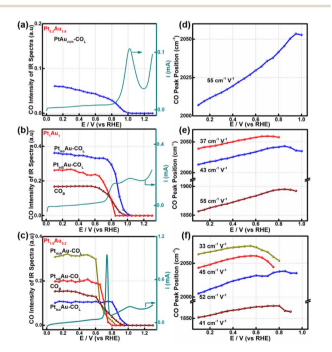


Fig. 4 Integrated peak intensities and peak positions of the various CO species in the COR process plotted against electrode potential change. (a) and (d) $Pt_{1.8}Au_{0.2}$, (b) and (e) $Pt_{1.4}Au_{1.8}$, (c) and (f) $Pt_{0.2}Au_{1.8}$. $Pt_{rich}Au - CO_L$ (dark yellow line), $Pt_{adj}Au - CO_L$ (red line), $Pt_{iso}Au - CO_L$ (blue line), CO_B (wine-colored line). The cyan solid lines are the COR curves of the PtAu samples in (a), (c) and (e).

Pt_{0.2}Au_{1.8}, Pt₁Au₁, and Pt_{1.8}Au_{0.2} were summarized in Fig. 4a-c to reveal the relationship between the surface-active Pt sites and COR. For Fig. 4a, there were only PtisoAu-COL sites in the Pt_{0.2}Au_{1.8} model film and the corresponding post-oxidation peak in the COR curve. The CO intensity of PtisoAu-COL decreased slightly at 0.20-0.85 V, which was in agreement with the previous work.29 As shown in Fig. 4d-f, the Stark tuning rate of Pt_{iso}Au-CO_L was near 55 cm⁻¹ V⁻¹ due to effect of Au in the film. It was more susceptible to potential variation than Ptrich-Au-CO. Based on the site structure, the PtisoAu-COL may induce conformational changes or convert intermediate species. In Fig. 4b, it was interesting that when the intensity of Ptadi Au-COL decreased at 0.70 V, an oxidation current of the pre-oxidation peak appeared in the COR curve. At 0.85 V, the PtadiAu-COL was completely oxidized and the pre-oxidation disappeared. This could be attributed to PtadiAu-COL as the reactive species of the pre-oxidation peak. Furthermore, when the voltage exceeded 0.85 V, the intensity of Pt_{iso}Au-CO_L gradually decreased, corresponding to the oxidation of the post-oxidation peak. This indicated that PtisoAu-COL was the main reactive species of post-oxidation peak. The only CO peak and postoxidation peak in Fig. 4a also confirmed the result. As depicted in Fig. 4c, similar to the Pt₁Au₁ film, Pt_{iso}Au-CO_L was also the main CO peak for the post-oxidation peak when the voltage surpassed 0.85 V. However, in the potential range of 0.65-0.80 V, the PtadiAu-COL and PtrichAu-COL were almost completely oxidized. This meant the PtrichAu-COL was also the reactive species of the post-oxidation peak. Due to the higher CO oxidation ability of the Pt_{rich}Au-CO_L, the pre-oxidation peak of Pt_{1.8}Au_{0.2} shifted negatively 0.10 V to 0.73 V, compared with the other samples. The result also verified the site structure and mechanism speculation of the COR curves. The PtrichAu site was oxidized in the early potential range as free sites and dissociated H₂O to OH*. The new OH* of the Pt_{rich}Au site contributed to oxidation of the adjacent COad of the PtadiAu sites. However, for the Pt atom isolated by Au of the PtisoAu-COL sites, the steric hindrance of Au may hinder OH* transfer to adjacent sites to oxidize PtisoAu-COL. The oxidation mechanism of PtisoAu-COL may be attributed to the Au product OH* at high potential to oxidize CO_{ad}. Compared with the Pt₁Au₁ and Pt_{1.8}Au_{0.2} model films in Fig. 4b-c, the intensity of Pt_{iso}Au-CO_L, Pt_{adi}Au-CO_L and Pt_{rich}Au-CO_L decreased to 0.65 V, 0.70 V and 0.85 V, respectively. This meant that the CO oxidation reactivity of the Pt surface could be expressed in the order Pt_{rich}Au-CO_L > Pt_{adj}Au- $CO_L > Pt_{iso}Au-CO_L$.

The Stark tuning rates of $Pt_{rich}Au-CO_L$, $Pt_{adj}Au-CO_L$ and $Pt_{iso}Au-CO_L$ in the different PtAu films were shown in Fig. 4d-f. In Fig. 4f, before CO oxidation, the Stark tuning rates of $Pt_{rich}Au-CO_L$, $Pt_{adj}Au-CO_L$ and $Pt_{iso}Au-CO_L$ gradually increased from 33 cm⁻¹ V⁻¹ to 45 cm⁻¹ V⁻¹ and 52 cm⁻¹ V⁻¹. The change in the Stark tuning rates could be attributed to the effect of Au. Compared to the $Pt_{rich}Au$ sites, the Pt atoms were decorated with Au atoms in the $Pt_{adj}Au$ sites and were isolated by Au atoms in the $Pt_{iso}Au$ sites. The heightened electronic effect results in an increase in the Stark tuning rate. Additionally, the various CO_L were changeable in different surface-active and model

films. This may be affected by the effect of CO_{ad}-CO_{ad} repulsions in the different active sites.29

3.4 In situ ATR-SEIRAS for studying the fine structure of the PtAu nanoparticles

To investigate the generalizability of the COR mechanism on the multiple active sites of the PtAu catalyst, we also prepared PtAu nanoparticles (NPs) using the NaBH₄ reduction method reported in previous work.17 The structure characterization and electrochemistry tests of the PtAu NPs are shown in Fig. 5 and S12.† The XRD patterns of the PtAu NPs showed typical facecentered cubic (fcc) features in Fig. S12a,† without phase separation. The PtAu NPs had a uniform particle size and were homogeneously dispersed on the carbon substrate (Fig. S12b†). Atomic-resolution high-angle annular dark field scanning transmission electron microscopy (HAADF-STEM) images of the PtAu NPs (Fig. 5a) showed that the lattice spacing was 0.23 nm, which corresponded to PtAu(111). In addition, the elemental mapping in Fig. 5b confirmed that Pt and Au were homogeneously distributed in the PtAu NPs. As the PtAu model films present an island structure without special structure (Fig. 1), the uniform spherical structure of the PtAu nanoparticles ensures the reasonableness of extending the in situ ATR-SEIRAS CO-probe method.

In the cyclic voltammetry tests (Fig. S12c†), by integrating the hydrogen underpotential deposition region charge and Au redox peak charge, the surface composition of the PtAu NPs was

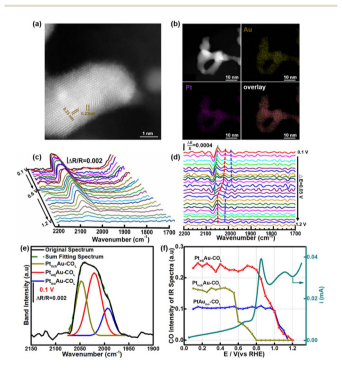


Fig. 5 (a) HAADF-STEM image of the PtAu NPs. (b) Elemental mapping of PtAu NPs. (c) CO oxidation infrared spectra and the reference potential at 1.2 V. (d) Second-derivative spectra of the COR spectra; the dotted lines shows the peak position changes. (e) Fitting peaks of the CO_L band at 0.1 V. (f) Integrated peak intensities of the various CO species in the COR and COR curve.

found to be Au rich (Pt: Au = 0.3: 0.7). Surprisingly, there were also three obvious CO oxidation peaks in the PtAu NPs (Fig. S12d†). Their onset oxidation potentials were 0.60 V, 0.75 V, and 0.90 V, respectively. The result indicated that the PtAu alloy was heterogeneous on the surface, although there was not phase separation in the XRD pattern and TEM characterization. The similar appearance maybe exist in other binary and ternary Pt-based alloy nanocatalyst. In particular, the similar CO oxidation feature peaks in the COR curves in the PtAu model films and PtAu nanoparticles enabled development of this in situ ATR-SEIRAS CO-probe method. The CO probe molecule can not only characterize the d-band center of the surface, but also investigate the uniformity of surface for Ptbased catalysts. Using in situ ATR-SEIRAS, the surface fine structure of the PtAu NPs was analyzed in depth. The COR spectra of the PtAu NPs were recorded from 0.10 V to 1.2 V, and the reference potential was set at 1.2 V (Fig. 5c). The results present one CO_L band with multiple peaks around 2050 cm⁻¹, without CO_B. The ATR-SEIRAS could detect detailed structural information due to the higher sensitivity for surface-adsorbed species, compared with only one CO_L peak in the in situ FTIR method. Moreover, second derivative spectra and Gaussian fitting were used to study the surface fine structure. In Fig. 5d, at 0.10 V, three obvious extreme points were observed at 2047 cm^{-1} , 2021 cm^{-1} and 1993 cm^{-1} . Similar to the PtAu model film, these could be attributed to three CO active sites, and their wavenumbers were close to those of Pt_{rich}Au-CO_L, Pt_{adi}Au-CO_L, and Pt_{iso}Au-CO_L, respectively.

The fitting spectra at 0.10 V and other potentials were shown in Fig. 5e and S13.† Comparing the integrated intensities of the three active sites, the CO onset oxidation potentials were 0.60 V, 0.75 V, and 0.95 V, respectively (Fig. 5f). The electrochemistry characteristics of the three active sites were also similar to Pt_{rich}Au-CO_L, Pt_{adi}Au-CO_L, and Pt_{iso}Au-CO_L. Therefore, based on the similar wavenumbers and CO oxidation ability, the three active sites of the PtAu NPs at 2047 cm⁻¹, 2021 cm⁻¹, and 1993 cm⁻¹ could be assigned as Pt_{rich}Au-CO_L, Pt_{adi}Au-CO_L and PtisoAu-COL, respectively. Moreover, the fitting spectra at a potential interval of 0.10 V and the changes in the CO_L integrated intensity were shown in Fig. S11† and 5f. Comparing the COR curve, for the first oxidation peak at 0.60-0.80 V, the intensity of PtrichAu-COL gradually decreased. This meant that Pt_{adi}Au-CO_L was the main active species for the first oxidation peaks. When the voltage exceeded 0.80 V, PtadiAu-COL was gradually oxidized, corresponding to the second oxidation peak. At 1.2 V, Pt_{adi}Au-CO_L was completely oxidized. In addition, the intensity of PtisoAu-CO_L gradually decreased at 1.0 V, close to the third oxidation peak. The result indicated that the Pt_{adi}Au-CO_L was attributed as the reactive species for the second oxidation peak and the third oxidation peak, and PtisoAu-COL was the main active species for the third oxidation.

4. Conclusions

In summary, we successfully utilized the in situ electrochemical ATR-SEIRAS and CO-probe method to classify the surface fine structure of PtAu model films and nanocatalysts. The electrodeposition method was used to construct model PtAu films with $Pt_{iso}Au$ sites, $Pt_{adj}Au$ sites and $Pt_{rich}Au$ sites. Additionally, the COR mechanism on these PtAu surface sites was revealed using *in situ* ATR-SEIRAS. By utilizing the SDP method located peak positions and Gaussian fitting to separate the complex CO band, three CO_L peaks in the COR spectra were disclosed. More importantly, correlating the changes in the integrated intensities of CO_L with the potentials, the CO oxidation reactivity could be placed in a sequence: $Pt_{rich}Au-CO_L > Pt_{adj}Au-CO_L > Pt_{iso}Au-CO_L$. The CO active sites could be assigned to the respective oxidation peaks in the COR curve. Finally, the surface fine configurations of the Au-rich PtAu nanoparticles were also determined.

Author contributions

Guang Li: conceptualization, methodology, investigation, validation, data acquisition & analysis, visualization, software, writing – original draft. Zheng-Chao An: data curation, writing – original draft. Jian Yang: writing – review & editing. Jin-Hong Zheng: methodology, investigation, data acquisition & analysis. Li-Fei Ji: methodology, investigation. Jun-Ming Zhang: methodology, investigation. Jin-Yu Ye: methodology, investigation, writing – review & editing. Bin-Wei Zhang: methodology, investigation, writing – review & editing. Yan-Xia Jiang: conceptualization, methodology, funding acquisition, supervision, writing – review & editing. Shi-Gang Sun: conceptualization, methodology, funding acquisition, supervision, writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was granted by the National Key Research and Development Program of China (2017YFA0206500) and the National Natural Science Foundation of China (22172134, 22288102 and 22279011).

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