

Research Article

Electrochemical Glucose Oxidation Using Glassy Carbon Electrodes Modified with Au-Ag Nanoparticles: Influence of Ag Content

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This paper describes the application of glassy carbon modified electrodes bearing Au_x-Ag_y nanoparticles to catalyze the electrochemical oxidation of glucose. In particular, the paper shows the influence of the Ag content on this oxidation process. A simple method was applied to prepare the nanoparticles, which were characterized by transmission electron microscopy, Ultraviolet-Visible spectroscopy, X-ray diffraction spectroscopy, and cyclic voltammetry. These nanoparticles were used to modify glassy carbon electrodes. The effectiveness of these electrodes for electrochemical glucose oxidation was evaluated. The modified glassy carbon electrodes are highly sensitive to glucose oxidation in alkaline media, which could be attributed to the presence of Au_x-Ag_y nanoparticles on the electrode surface. The voltammetric results suggest that the glucose oxidation speed is controlled by the glucose diffusion to the electrode surface. These results also show that the catalytic activity of the electrodes depends on the Ag content of the nanoparticles. Best results were obtained for the Au₈₀-Ag₂₀ nanoparticles modified electrode. This electrode could be used for Gluconic acid (GA) production.

1. Introduction

Glucose electrooxidation, specifically on Au electrodes, has been extensively studied because it is a reaction of scientific and industrial interest [1–11]. Due to the high catalytic activity of modified electrodes bearing nanoparticles, Au nanoparticles have been studied for glucose fuel cells [12–14], for degradation of dyes [15–17], and, last but not least, for glucose sensor in medical applications [18–21]. Furthermore, bimetallic nanoparticles have received more and more attention because their amazing optical, electronic, and catalytic properties, leading to various applications, as,

indeed, sensors, catalysis, or SERS substrates [22–29], and so forth. Among these nanoparticles, Au_x-Ag_y or Au_x-Pt_y bimetallic nanoparticles have been extensively explored for different electrochemical reactions as methanol and glucose electrooxidation [30–34].

On the other hand, Gluconic acid (GA) is a high value added chemical used as intermediate in the food, pharmaceutical, and beverage industries. The only commercial/industrial method to produce GA is by glucose biochemical oxidation [35], whose main disadvantages are slow reaction rate, low space-time, and problematic enzyme-product separation. In the past few years, heterogeneous

TABLE 1: Composition of Au-Ag nanoparticles synthesized.

Sample	Composition	% wt Au	% wt Ag	H ₂ AuCl ₄ (mol)	AgNO ₃ (mol)
M1	Au ₁₀₀	100	0	$2.53 * 10^{-4}$	0
M2	Au ₉₀ Ag ₁₀	90	10	$2.28 * 10^{-4}$	$4.63 * 10^{-5}$
M3	Au ₈₀ Ag ₂₀	80	20	$2.03 * 10^{-4}$	$9.27 * 10^{-5}$
M4	Au ₇₀ Ag ₃₀	70	30	$1.77 * 10^{-4}$	$1.39 * 10^{-4}$

catalysis using Pt, Pd, or Au nanoparticles has attracted much attention [36, 37]. The selective glucose oxidation to GA has been reported using Au clusters as catalyst and O₂ as oxidant giving a yield of 98% at mild conditions, that is, 50 °C and atmospheric pressure. Indeed, higher costs, electrode poisoning, and hard reuse of nanocatalyst make this process far from competitive GA production [38]. Some studies have been oriented to the use of bi- and trimetallic nanoparticles to overcome the noneconomic electrochemical GA production, at least to elucidate the best nanoparticle composition for the electrooxidation of glucose. In this sense, bimetallic Au-Ag nanoparticles show important synergistic effects in the catalytic properties; these properties depend upon metals ratio of nanoparticle, nanoparticle synthesis, nanoparticle size, form, and crystallographic orientation [39, 40]; indeed, it has been shown that the surface morphology and particularly the size of nanoparticles can be easily controlled by experimentally adjusting the synthesis conditions [41, 42].

In our previous work, we reported that Au nanoparticles supported on glassy carbon presented a catalytic activity and selectivity towards glucose oxidation, depending on the particle size and on the crystallographic orientation. Results also suggest that oxidation process in these conditions is taking place with lower poisoning of the surface in the case of the Au nanoparticles than for massive gold that the process is irreversible, and perhaps there are some chemical reactions involved in the overall oxidation process [25].

The aim of this work is to evaluate the catalytic properties of Au_x-Ag_yNPs towards the glucose electrooxidation; hence, we studied mainly the Au-Ag nanoparticles voltammetric response of different Au-Ag ratio.

2. Experimental Section

2.1. Preparation of Au_x-Ag_y Nanoparticles (Au_x-Ag_yNPs). Au_x-Ag_yNPs were prepared according to a previously published procedure [25]. The procedure has been modified in order to prepare bimetallic nanoparticles. Briefly, 30 mL of aqueous solutions containing the respective amount of H₂AuCl₄ and AgNO₃ to give nanoparticles Au₁₀₀ (M1), Au-Ag_{90/10} (M2), Au-Ag_{80/20} (M3), and Au-Ag_{70/30} (M4) was prepared (see Table 1) and was added to 10 mL of toluene (Aldrich) containing 0.34 mM of tetraoctylammonium bromide (TOAB 98%, Fluka) as a phase-transfer agent. Dodecanethiol (Aldrich) was incorporated to these different solutions as a stabilizing agent in an Au-Ag:thiol 2:1 ratio followed by the addition of an excess of NaBH₄ as an aqueous reducing agent. The reaction was allowed to proceed under constant stirring at controlled temperature (80 °C), for

3 h. Finally, colored dispersions were obtained and purified several times in ethanol (J. T. Baker). The resulting Au-Ag nanoparticles were characterized by transmission electron microscopy (TEM), UV-Visible spectroscopy (UV-Vis), X-ray diffraction spectroscopy (XRD), and cyclic voltammetry (CV).

2.2. Preparation of Glassy Carbon Modified Electrodes. To remove the thiol stabilizing agent layer from Au_x-Ag_yNPs, the NPs were mixed with Vulcan XC-72 and were heated at 300 °C for 2 hours under air atmosphere; the temperature was controlled within 2 °C.

Then, Au_x-Ag_yNPs-modified electrodes were prepared as follows: 1 μL aliquot of thiol-Au_x-Ag_yNPs in hexane and Nafion (5% Electrochem) were mixed (1:10 ratio) and cast onto a carbon disk (CD) followed by natural evaporation at room temperature.

2.3. Characterization of Nanoparticles. Synthesized Au_x-Ag_yNPs were characterized using TEM, UV-Vis, XRD, and cyclic voltammetry (CV). TEM characterizations were performed on a TITAN 80-300 FEI microscope and the particle size distribution was measured from about 200 particles using the TEM images. Nanoparticles samples dissolved in hexane were cast onto a carbon-coated copper grid sample holder followed by natural evaporation at room temperature. UV-Vis measurements were carried out on HP spectrophotometer model 8453. XRD measurements were obtained using a Siemens D5000 diffractometer. Spectra were collected from 10 to 80 ° at a speed of 0.002°·seg⁻¹. Cyclic voltammetric measurements were performed in BioLogic VSP Modular 5 channels Potentiostat/Galvanostat using a three-electrode conventional cell. A separated compartment and an Ag/AgCl electrode were used as reference and a Pt wire as the counter electrode; all potentials were referred to this electrode. Glucose 0.01 M in KOH 0.3 M solution was used for the electrochemical characterization. The solution was purged for twenty minutes with high-purity nitrogen before taking measurements.

3. Results and Discussion

3.1. Characterization of Au_x-Ag_yNPs

3.1.1. TEM Characterization. Figures 1(a)–1(d) show TEM micrographs of the synthesized Au_x-Ag_y nanoparticles (labeled as M1 to M4) and their particle size distribution. Synthesis of Au_x-Ag_yNPs can be easily controlled to obtain narrow particle size distribution of spherical nanoparticles

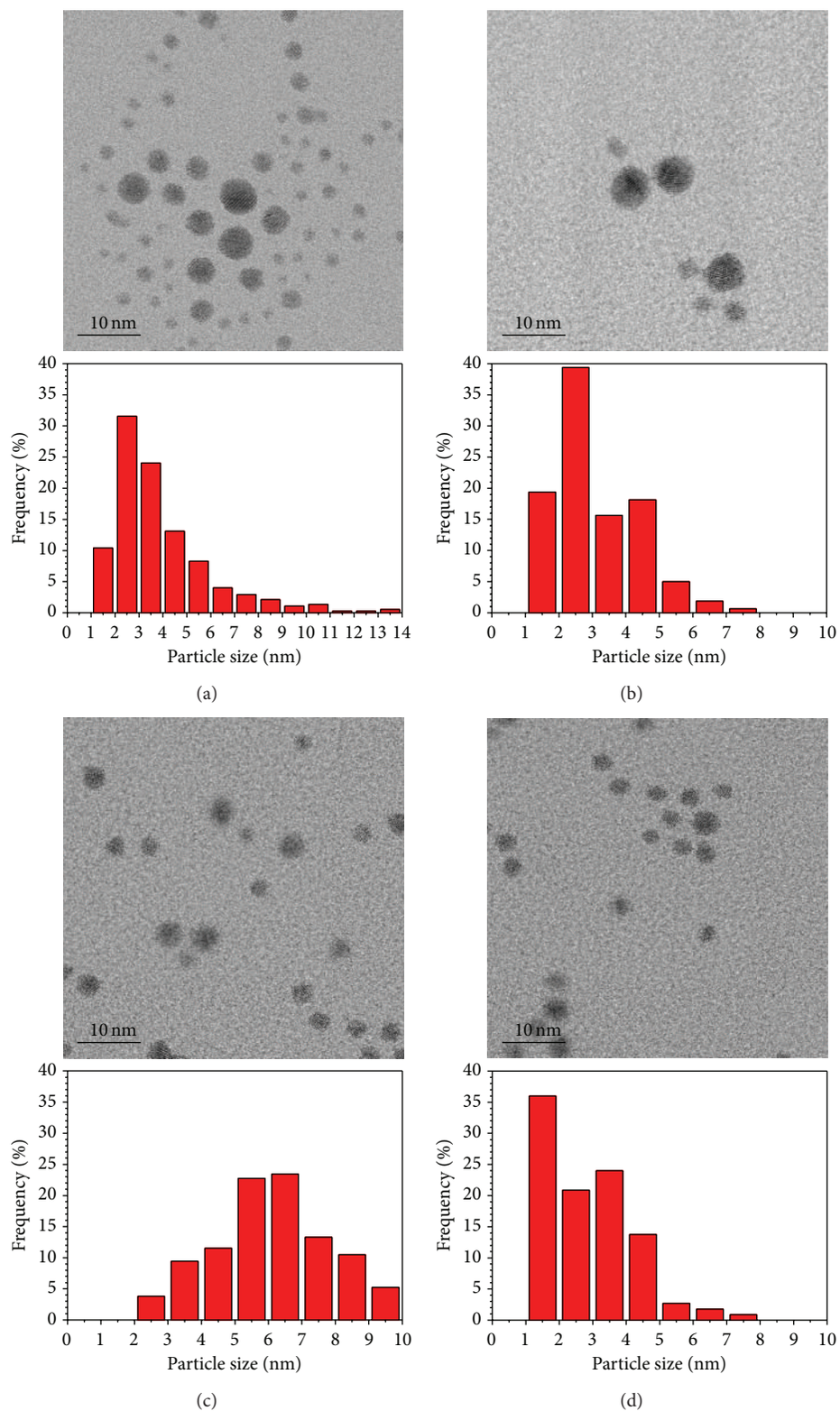


FIGURE 1: TEM images of thiol capped Au-Ag bimetallic nanoparticles. (a) $\text{Au}_{100}\text{Ag}_0$, (b) $\text{Au}_{90}\text{Ag}_{10}$, (c) $\text{Au}_{80}\text{Ag}_{20}$, and (d) $\text{Au}_{70}\text{Ag}_{30}$.

ranging from 1 to 7 nm. Figures 2(a)–2(d) show the Au_x-Ag_y NPs supported in Vulcan after heat treatment for thiol elimination. It can be seen that Au_x-Ag_y NPs keep their spherical form but at the same time there is an increase in size and distribution. The final size ranges from 6 to 40 nm; in spite of the size increase, the Au_x-Ag_y NPs are still small compared to other synthesis methods [33].

3.1.2. UV-Visible. It is well known that nanoparticles have surface plasmon (SP) resonance absorption bands in the visible region. SP resonance bands are strongly dependent on the size, shape, composition, and medium dielectric properties [43–46].

Figure 3 shows UV-Visible spectra from Au_x-Ag_y nanoparticles dispersed in hexane. Au and Ag nanoparticles are known to have plasmon bands in the visible region located at 520 and 420 nm, respectively. In the figure, distinct peaks are observed clearly at 530, 520, 512, and 506 nm; all these peaks are located at intermediate positions between the Au and Ag plasmon bands. The plasmon band is blue-shifted as the silver amount increases.

Two plasmon bands would be expected for a physical mixture of Au and Ag nanoparticles and the formation of an Au-Ag alloy could be deduced from the fact that the optical absorption spectrum shows only one plasmon band for all the samples [47, 48].

The formation of spherical Au_x-Ag_y NPs was demonstrated by TEM (Figure 1). Indeed, upon heat treatment for thiol elimination, the Au_x-Ag_y NPs, in spite of slight size increase, keep the spherical form as shown in Figure 2. Results are in agreement with the literature [25].

3.1.3. XRD Characterization. Figure 4 shows the XRD patterns of the different NPs synthesized. XRD analysis revealed in all the samples the presence of four representative diffraction peaks of face-centered cubic (fcc) structure that correspond to (111), (200), (220), and (311) planes. The angles of these diffraction peaks in all the samples were greater than those for pure Ag (38.11, 44.27, 64.42, and 77.47°) and lower than those for pure Au (38.18, 44.39, 64.57, and 77.54°) according to the standards for Ag [JCPDS 04-0783] and Au [JCPDS 04-0784]. These results confirm again the formation of an alloy of Au-Ag. No other impurity peak is detected in the samples.

The intensity of each signal respect to the other signals means the proportion of each orientation present in that particular nanoparticle. For nanoparticles synthesized by our method, the main factor influencing the orientation plane is the Au-Ag: thiol ratio [25]. In this case, we used a constant Au-Ag: thiol ratio of 2:1 in order to obtain a (111): (200) plane ratio of 2.3–2.8.

3.1.4. Electrochemical Characterization. We have used cyclic voltammetry for the Au_x-Ag_y NPs electrochemical characterization. Figure 5 shows the response of both electrodes, Au polycrystalline (Poly-Au) (Figure 5(a)) and carbon disk Au_{100} nanoparticle-modified electrode (Figure 5(b)) in 0.3 M KOH. We can see the typical Au voltammetric response, that is,

the formation and reduction of Au oxides. The surface area of the Au_{100} NPs electrode was larger than that of a Poly-Au electrode by integrating the reduction charge of gold oxide monolayer. It indicated that there was nanostructured gold on the Au NPs electrode. The Au_{100} NPs electrode exhibited the same behavior of Poly-Au electrode, having higher current densities of relative peaks compared to that of Poly-Au electrode. Poly-Au and Au_{100} NPs electrodes exhibit the anodic peak located at 0.28 V corresponding to the oxidation of gold. The cathodic peak is located at 0.07 V for Au_{100} NPs and 0.1 V for Poly-Au corresponding to the subsequent reduction of the gold oxides. Nevertheless, there was a weak cathodic peak located at -0.2 V for Au_{100} NPs and -0.18 V for Poly-Au attributed to desorption of hydroxide anions.

As the content of Ag in nanoparticles increases ($Au_{80}Ag_{20}$ and $Au_{70}Ag_{30}$, Figures 5(c) and 5(d)), the voltammogram changes to take the form of that of Ag nanoparticles in alkaline media (Figures 5(e) and 6(f)) [49]. The cyclic voltammograms of the silver disk electrode or silver nanoparticle-modified electrode in alkaline media show three anodic peaks (Ox_1 , Ox_2 , and Ox_3) appearing at 0.25, 0.33, and 0.65 V, respectively (Figures 5(e) and 6(f)). During a reverse scan, two cathodic peaks (R_1 and R_2) appeared at 0.38 and 0.06 V, respectively, as reported in previous studies [49]. The first anodic peak (Ox_1) is due to the electroformation of the monolayer of Ag_2O . The second anodic peak (Ox_2) is due to the formation of the multilayer of Ag_2O . The third peak (Ox_3) is due to the oxidation of Ag_2O to AgO or direct oxidation of Ag to AgO.

The cathodic current peak R_1 is due to the reduction of AgO to Ag_2O , and R_2 corresponds to the reduction of Ag_2O to Ag. A small anodic peak (Ox_4) at 0.63 V appears during the cathodic scan, but it was dependent on the scan rate and occurred only when the scan rate is faster than 50 mV/s. Stonehart has suggested that peak Ox_4 is associated with autocatalytic oxidation process to convert Ag_2O to AgO [50]. When the nuclei of AgO formed at the Ag_2O surface during the anodic scan, further formation of AgO became easy and occurred at a lower potential than the initial formation of AgO nuclei. Therefore, during the cathodic scan, further formation of AgO resulted in the appearance of a small anodic peak.

It can be seen in Figure 5(c) ($Au_{80}Ag_{20}$) that the anodic scan shows three peaks probably due to the oxidation of gold (Ox_1), the electroformation of the monolayer of Ag_2O (Ox_2), and the oxidation of Ag_2O to AgO (Ox_3). The cathodic scan shows also three peaks probably corresponding to the reduction of AgO to a multilayer of Ag_2O (R_1), the reduction of Ag_2O to Ag (R_2), and the reduction of Au oxides (R_3).

As a consequence of the increase in Ag content, the anodic scan of $Au_{70}Ag_{30}$ NPs shows two peaks. The first one (Ox_1) is clearly due to the simultaneous formation of Au oxides and the formation of the Ag_2O monolayer because practically these two processes take place at the same potential resulting in one peak; the second anodic peak (Ox_2) is due to the oxidation of Ag_2O to AgO. The cathodic scan shows four peaks: R_1 is due to the reduction of AgO to a multilayer of Ag_2O , R_2 is due to the reduction of Ag_2O to Ag, and

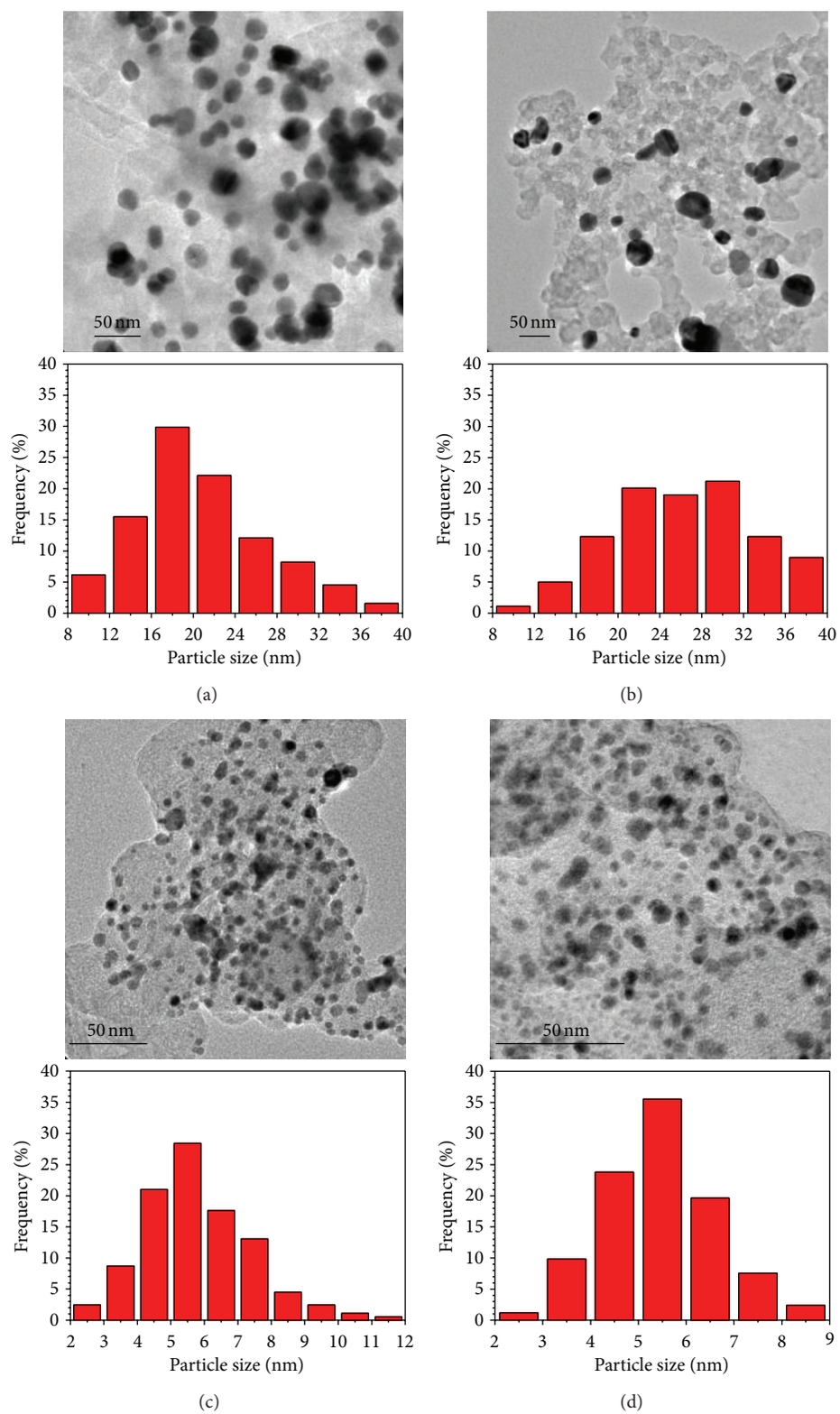


FIGURE 2: TEM images of Au-Ag bimetallic nanoparticles supported in Vulcan after heat treatment for thiol elimination. (a) $\text{Au}_{100}\text{Ag}_0$, (b) $\text{Au}_{90}\text{Ag}_{10}$, (c) $\text{Au}_{80}\text{Ag}_{20}$, and (d) $\text{Au}_{70}\text{Ag}_{30}$.

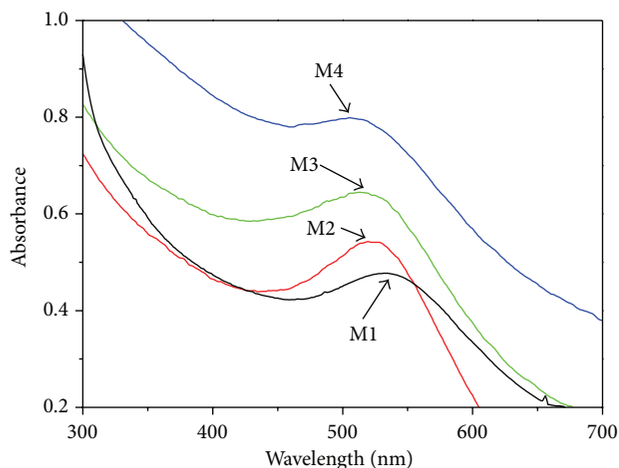


FIGURE 3: UV-vis spectra of bimetallic nanoparticles at different Au-Ag molar compositions. (M1) $\text{Au}_{100}\text{Ag}_0$, (M2) $\text{Au}_{90}\text{Ag}_{10}$, (M3) $\text{Au}_{80}\text{Ag}_{20}$, and (M4) $\text{Au}_{70}\text{Ag}_{30}$.

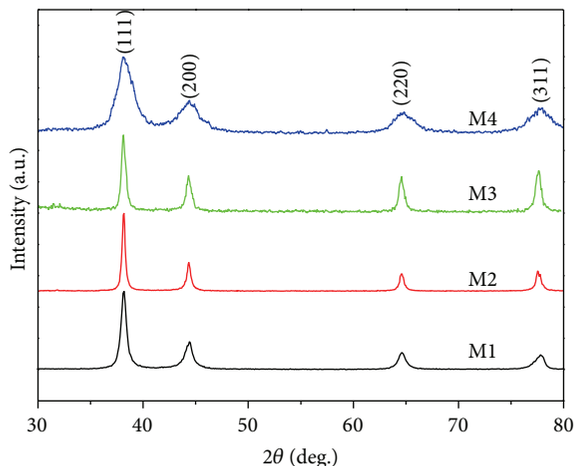


FIGURE 4: XRD patterns of different nanoparticles. (M1) $\text{Au}_{100}\text{Ag}_0$, (M2) $\text{Au}_{90}\text{Ag}_{10}$, (M3) $\text{Au}_{80}\text{Ag}_{20}$, and (M4) $\text{Au}_{70}\text{Ag}_{30}$.

peaks R_3 and R_4 are due to the reduction of the gold oxides and the desorption of hydroxide anions. These results are in agreement with others works [49, 51].

3.1.5. Electrochemical Oxidation of Glucose on AuNPs and Au-AgNPs. Figures 6(a)–6(e) compares the voltammograms obtained for the electrochemical oxidation of 10 mM glucose at the Poly-Au electrode and the $\text{Au}_x\text{-Ag}_y\text{NPs}$ electrodes in 0.3 M KOH solution. The Poly-Au and the $\text{Au}_x\text{-Ag}_y\text{NPs}$ electrodes demonstrated a typical voltammetric response for glucose. Interestingly, the current density of oxidation of glucose on the $\text{Au}_x\text{-Ag}_y\text{NPs}$ electrodes was higher than that on Poly-Au electrode; it could be attributed to the high surface area of $\text{Au}_x\text{-Ag}_y\text{NPs}$. However, the only $\text{Au}_x\text{-Ag}_y\text{NPs}$ electrode that did not show better performance is the M4 electrode ($\text{Au}_{70}\text{Ag}_{30}$); hence, we discard it for further experiments and analysis.

TABLE 2: Oxidation peak and current density as a function of Ag content.

Sample	Potential, E/V versus Ag/AgCl/KCl sat'd	Current density (mAcm^{-2}) @ 0.49 V versus Ag/AgCl/KCl sat'd
Poly-Au	-0.52 ± 0.01	1.0 ± 0.1
M1 ($\text{Au}_{100}\text{-Ag}_0$)	-0.39 ± 0.03	1.2 ± 0.1
M2 ($\text{Au}_{90}\text{-Ag}_{10}$)	-0.41 ± 0.04	0.82 ± 0.1
M3 ($\text{Au}_{80}\text{-Ag}_{20}$)	-0.49 ± 0.04	1.9 ± 0.3
M4 ($\text{Au}_{70}\text{-Ag}_{30}$)	-0.54 ± 0.04	0.5 ± 0.1

There is also a shift of the glucose potential oxidation on the $\text{Au}_x\text{-Ag}_y\text{NPs}$ M1, M2, and M3 electrodes with respect to the Poly-Au electrode (see Table 2); the shift is to less-negative potentials and reveals that the $\text{Au}_x\text{-Ag}_y\text{NPs}$ electrodes could catalyze the oxidation of glucose. Indeed, we show in Table 2, for comparison purposes, the current density under the same potential for the electrode tested. These results are similar to that of Qiaofan's work on glucose oxidation in an alkaline electrolyte obtained at bimetallic Au-Ag/RGO/GC (bimetallic Au-Ag/reduced graphene oxide/glassy carbon) nanoparticles modified electrode [51].

The CV curves of the $\text{Au}_x\text{-Ag}_y\text{NPs}$ electrode presented a typical two-step oxidation process and three obvious oxidation peaks in the positive scan were observed. As the mechanism for the electrocatalytic oxidation of glucose at $\text{Au}_x\text{-Ag}_y\text{NPs}$ electrodes is a multistep one, the first step mediated the chemisorptions of hydroxide anions onto the electrode surface, leading to the formation of hydrous gold-silver oxide ($\text{Au}_x\text{-Ag}_y\text{OH}_{\text{ads}}$) which was believed to be the catalytic component of $\text{Au}_x\text{-Ag}_y\text{NPs}$ electrodes [52].

The glucose oxidation highly depends on the quantity of $\text{Au}_x\text{-Ag}_y\text{OH}_{\text{ads}}$ and the premonolayer oxidation was enhanced at $\text{Au}_x\text{-Ag}_y\text{NPs}$ surface. The first peak, Ox_1 , located between -0.54 and -0.39 V could be attributed to the electrosorption of glucose to form adsorbed intermediate gluconolactone, releasing one proton per glucose molecule. At around these potentials, there could be a very limited number of $\text{Au}_x\text{-Ag}_y\text{OH}_{\text{ads}}$ sites on the $\text{Au}_x\text{-Ag}_y\text{NPs}$ surfaces due to the reductive nature of the electrodes potential. Glucose molecules interacted with the $\text{Au}_x\text{-Ag}_y\text{OH}_{\text{ads}}$ sites to give rise to the intermediate, gluconolactone [53, 54]. Nevertheless, the accumulation of gluconolactone blocks the active sites of the $\text{Au}_x\text{-Ag}_y\text{NPs}$ electrode surface inhibiting the direct oxidation of glucose, leading to the decrease of current density. The second oxidation peak Ox_2 appeared at a more positive potential located between 0.18 and 0.26 V depending on the Ag content in the electrode. When the potential reaches those values, the population of $\text{Au}_x\text{-Ag}_y\text{OH}_{\text{ads}}$ sites on the $\text{Au}_x\text{-Ag}_y\text{NPs}$ electrode increased; hence, catalytic oxidation of the intermediates takes place. A further positive increase in the potential to around 0.6 V gives rise to the formation of gold-silver oxides on the surface of $\text{Au}_x\text{-Ag}_y\text{NPs}$. The formation of Au-Ag oxides results in a decrease of the number of the $\text{Au}_x\text{-Ag}_y\text{OH}_{\text{ads}}$ sites on the surface $\text{Au}_x\text{-Ag}_y\text{NPs}$ and impedes the electrocatalytic oxidation of glucose and by-products. The

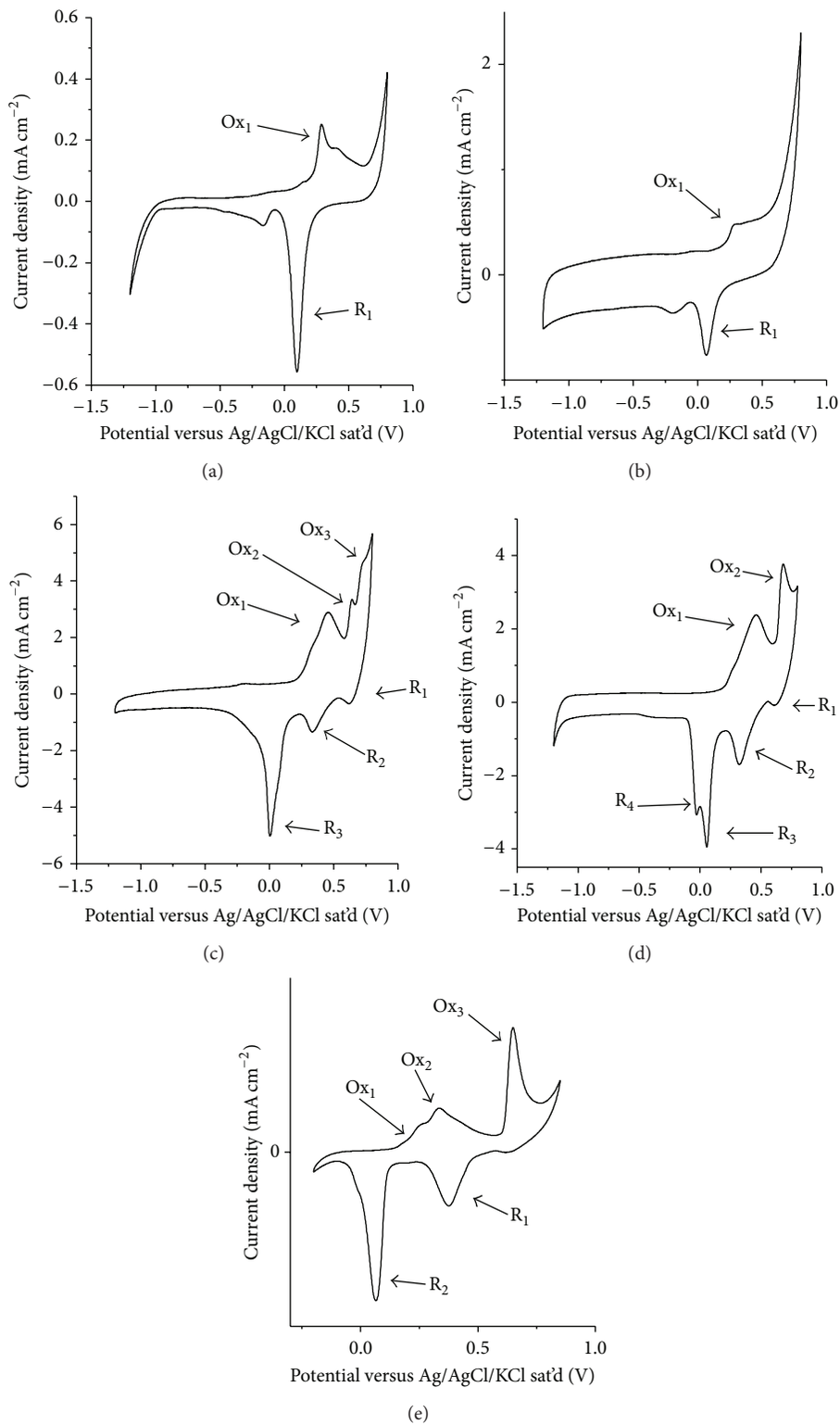


FIGURE 5: CV curves in 0.3 M KOH solution for (a) solid Au electrode, (b) Au₁₀₀, (c) Au₈₀Ag₂₀, (d) Au₇₀Ag₃₀, and (e) Au₀Ag₁₀₀. Scan rate, 50 mV s⁻¹. Scan direction →.

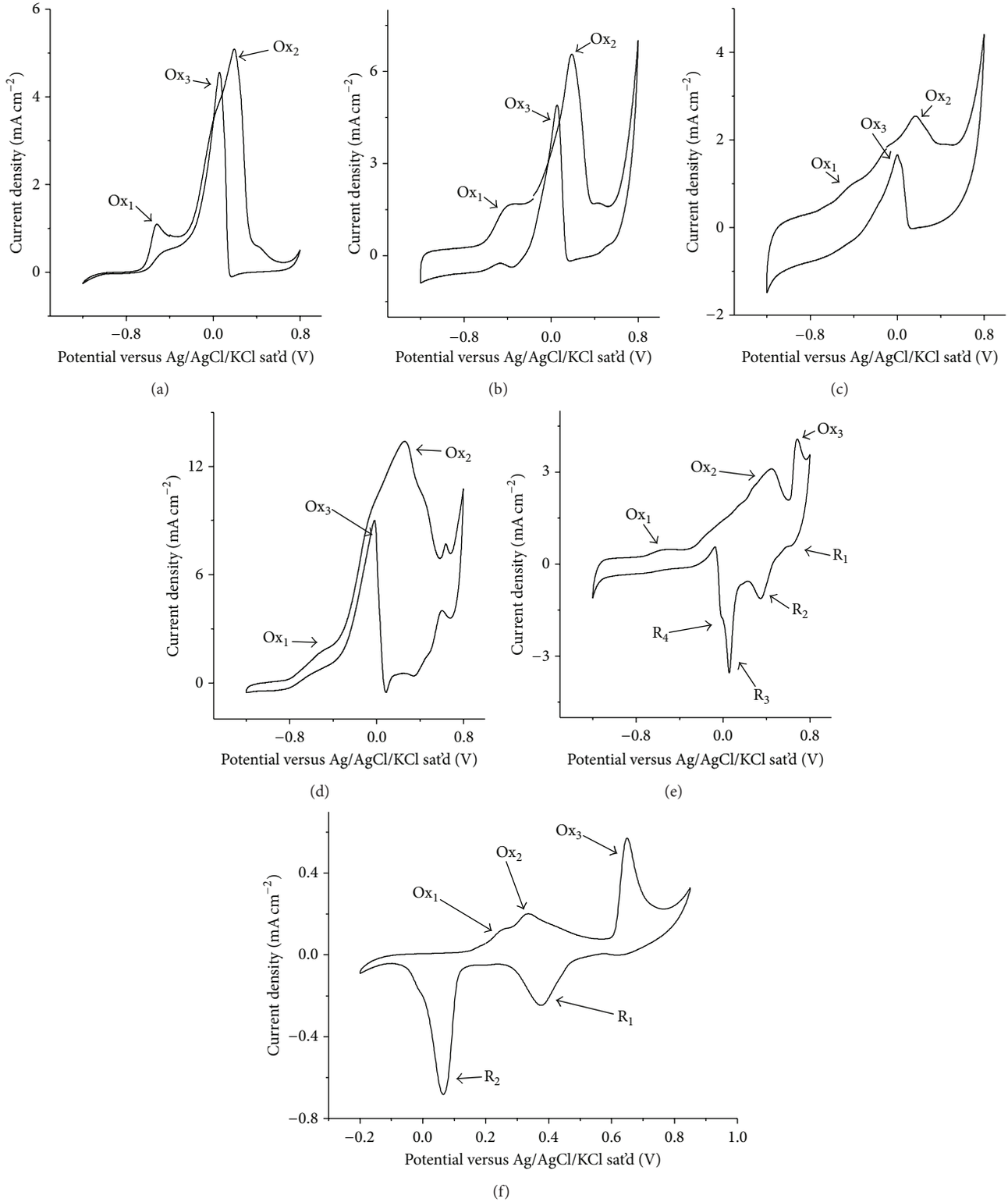


FIGURE 6: CV curves in 0.3 M KOH + 10 mM glucose solution for (a) solid Au electrode, (b) Au₁₀₀NP electrode, (c) Au₉₀Ag₁₀, (d) Au₈₀Ag₂₀, (e) Au₇₀Ag₃₀, and (f) Au₀Ag₁₀₀. Scan rate, 50 mVs⁻¹. Scan direction →.

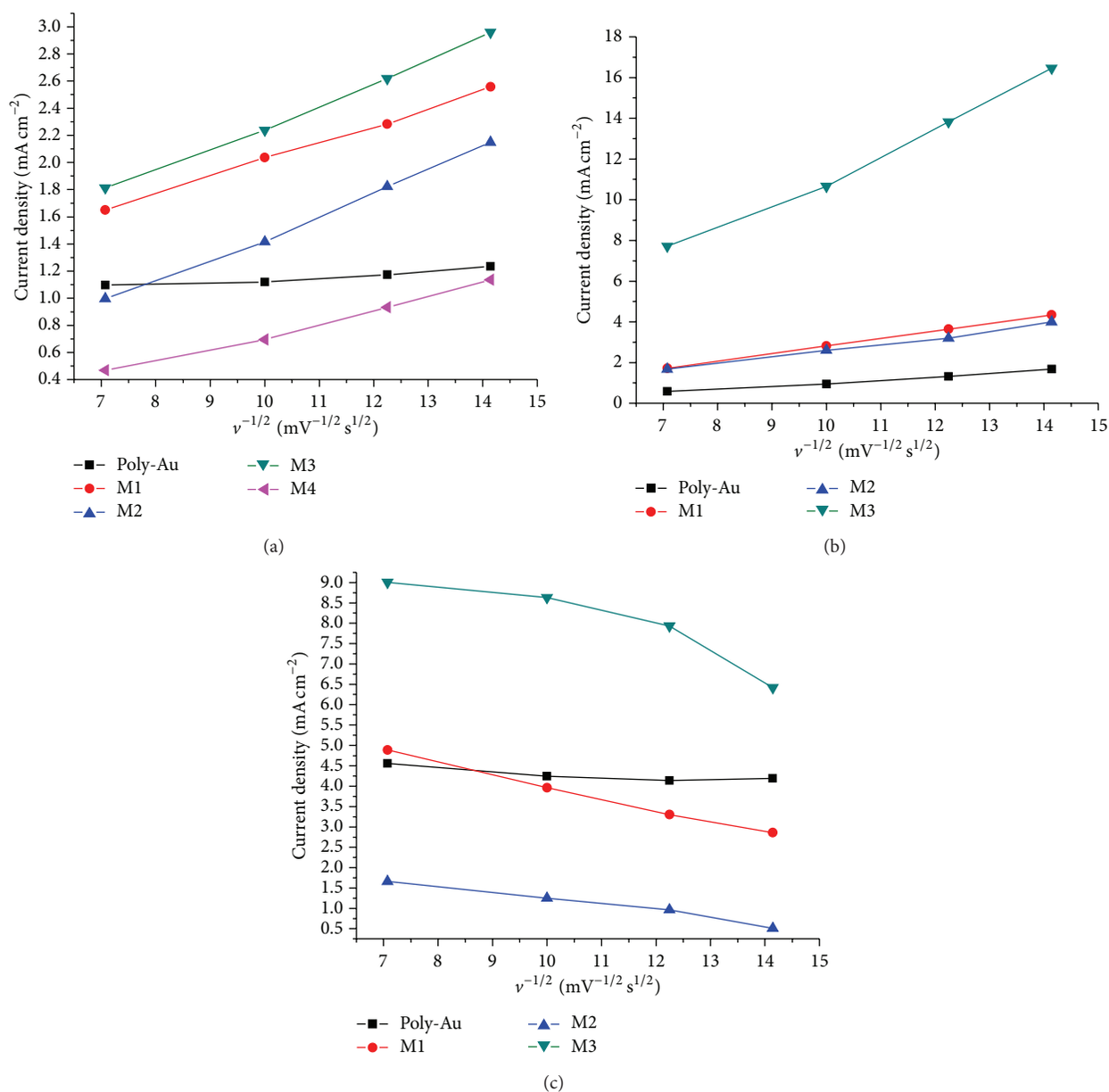


FIGURE 7: Plots of current density versus $V^{-1/2}$ (a) first oxidation scan peak, (b) second oxidation scan peak, and (c) oxidation cathodic peak.

current density, then, decreases dramatically at more positive values than 0.60 V, due to formation of gold and silver oxides. In the negative potential scan, the reduction of the surface gold and silver oxides would occur at the potential more negative than 0.16 V. Then, enough surface active sites would be available for the direct oxidation of glucose, resulting in a sharp increase in anodic current showing a peak (Ox_3), located between 0.001 and 0.15 V, depending on the Ag content in the electrode. Tang's group had reported a similar result; that is, there is a sharp increase in anodic current with a peak at 0.1 V [53]. The surface structure of a gold electrode influenced the electrooxidation of glucose via the influence on the adsorption of OH^- anions and gold oxide surface monolayer formation [55]. The as-prepared Au_x-Ag_y NPs

electrode exhibited better performance on electrocatalytic oxidation toward glucose than a Poly-Au electrode.

To evaluate the kinetics of the glucose oxidation on the Au_x-Ag_y NPs, the CVs of the electrodes in a 0.3 M KOH solution containing 10 mM glucose at different scan rates were recorded (figure not shown). The peak Ox_1 in the positive scan, located between -0.54 and -0.39 V, corresponds to the direct oxidation of glucose and the current density is proportional to the square root of scan rate for all the electrodes used (Poly-Au, M1, M2, M3, and M4, Figure 7(a)); this fact suggests a linear diffusion-controlled process. Indeed, it is in agreement with the proposed mechanism for peak Ox_1 , in which sufficient active sites on the hydroxyl coated gold surface allowed direct oxidation of

glucose to generate gluconolactone [25, 56]. We can see that the order from better to worst with respect to the glucose oxidation is M3 > M1 > M2 > Poly-Au > M4. As expected, the current density of all electrodes increases as scan rate rises but that of Poly-Au remains practically unchanged. Furthermore, the slope of Au_x-Ag_yNPs increases depending on the Ag content in the NPs. The Au₈₀-Ag₂₀ electrode has the highest slope, that is, the highest glucose oxidation rate.

In Figure 7(b), we can see the plot of current density versus square root of scan rate corresponding to the further oxidation of gluconolactone located between 0.18 and 0.26 V (Ox₂) on the same electrodes except that of M4. Figure 7(b) suggests again that the oxidation of gluconolactone is also a diffusion-controlled process. The order from better to worst is M3 > M1 > Poly-Au > M2. This time, the slope of Poly-Au increases as expected; we are working on the matter.

Finally, Figure 7(c) shows the plot of current density versus square root of scan rate for the oxidation peak Ox₃. Interestingly, as could be seen, peak Ox₃ corresponds to the direct oxidation of glucose in cathodic scan and is also linear versus scan rates. Indeed, this is a typical adsorption-controlled process corresponding to a fast electron-transfer behavior. In the negative potential scan, the reduction of the gold surface oxides provides enough surface active sites for the direct oxidation of glucose; they could account for the appearance of peak Ox₃. The order from better to worst is M3 > Poly-Au > M1 > M2. This time, the slope of Poly-Au remains again unchanged.

None of the straight lines of plots in Figures 7(a), 7(b), and 7(c) go through origin indicating a previous chemical step in the oxidation process; as noted above, this step is probably due to the chemisorption of hydroxide anions onto the gold and silver surface, leading to the formation of hydrous oxides (Au_x-Ag_yOH_{ads}), which are believed to be the catalytic component of Au_x-Ag_yNPs electrodes, particularly Au₈₀-Ag₂₀NPs electrode.

4. Conclusion

The nanoparticles applications, for instance, glucose oxidation, critically depend on their specific catalytic activity; unfortunately, it is still poorly understood [57]. Experimental evidence that can be found in literature points towards several variables affecting it, that is, nanoparticle size [57], nanoparticle crystal orientation [56], and composition of nanoparticle [51]. It is admitted that the glucose oxidation depends on the presence of what is known as monolayer or premonolayer of AuOH_{ads} or AgOH_{ads} [48]; at higher potential, the formation of Au and Ag oxides impedes the glucose oxidation. Nevertheless, Mikhlin et al. [57] found that the formation of AgO proceeded readily at the “primary” oxide for nanoparticles of about 12 nm but was retarded at the smaller ones. These findings are in agreement with our previous work where we have found a maximum in the catalytic activity for Au nanoparticles of about 6 nm and could explain the present findings; that is, smaller nanoparticles have the highest catalytic activity towards glucose oxidation.

Regarding nanoparticle composition, the experimental evidence strongly suggests that the best composition is around 80% Au and 20% Ag.

Indeed, we have performed the present study for a (111):(200) plane ratio of 2.3–2.8. Studies to optimizing the NPs size, Au-Ag ratio, and crystallographic orientation are underway.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

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