RESPONSE OF THE POLYCRYSTALLINE 
GOLD/ALKALINE SOLUTION INTERFACE TO 
COMPLEX POTENTIODYNAMIC PERTURBATIONS 
WITHIN THE POTENTIAL RANGE OF THE 
THERMODYNAMIC STABILITY OF WATER

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Abstract - The potentiodynamic characteristics of Au/alkaline solution interfaces within the potential range 
of the thermodynamic stability of water, under different complex potential perturbation conditions are 
described. The composite E/I contours are interpreted in terms of a reversible initial process involving OH- 
electrosorption, followed by the formation of Au(III) surface species when the Au/Au(III) equilibrium 
potential is reached. The latter process clearly involves the simultaneous participation of ageing effects 
during the course of the reaction. The results correlate with optical data reported in the literature.

INTRODUCTION

Most of the electrochemical data related to gold 
electrodes, particularly concerning the oxygen electro-
sorption and electrodesorption processes refer to acid 
aqueous electrolytes[1,2]. Contrarily these reactions 
appear to have been less studied in alkaline aqueous 
electrolytes than in acid electrolytes[3-9]. The poten-
tiodynamic E/I displays in the potential range 
assigned to the water thermodynamic stability exhibit 
a relatively large number of anodic and cathodic 
current peaks and shoulders[10-11]. The characteris-
tics of the cathodic current peaks are sensitive to the 
type of perturbation applied to the interface. This fact 
is due at least in part to the ageing of the species 
anodically formed. The ageing of the electrochemical 
interfaces can be promoted by different types of 
potential perturbations[12-14]. Thus, open circuit, 
potentiostatic and potentiodynamic induced ageing 
effects can be distinguished[14]. The information 
derived from the ageing effects reveals interesting 
details of the electrochemical formation of layers on 
different electrodes.

It has been recently demonstrated[15,16] that the 
potentiodynamic E/I profiles related to the elec-
 troreduction of oxygen-containing layers, either within 
of the order of a monolayer thickness or thin layers, 
can be drastically changed when a repetitive triangular 
potential sweep confined to a certain potential range is 
 applied for a definite length of time. Consequently, the electrodeposition current peak potential shifts in the 
negative potential direction. This is expected for a reactant which, owing to the intermediate pertur-
bation, became more stable than that involved in the 
conventional triangular potential sweep without the intermediate perturbation. This effect known as the 
potentiodynamic aging of the film, has already been 
described for various electrochemical reactions[16-18] including that of the oxygen-containing 
monolayer on gold in different acid electrolytes[15].

The present paper also includes the electrochemical 
response of the gold/alkaline interface to triangularly 
modulated triangular potential sweeps[19,20]. The latter perturbation technique renders straightforward 
information on the degree of reversibility of possible 
conjugate redox couples occurring in the course of the 
reaction including, under certain circumstances, redox 
couples involving intermediate species.

EXPERIMENTAL

The electrolysis cell is the same as that already 
described in previous publications[21,22]. Specpure 
polycrystalline gold wires of 0.38 cm² apparent area 
working electrodes were used. The electrode surface 
was firstly mirror polished with the purest 300 mesh 
alumina powder, then the electrode was potential 
cycled in the alkaline solution to attain the conven-
tional E/I display under repetitive triangular potential 
sweeps covering the potential range of the water 
thermodynamic stability[23]. Finally, the E/I displays 
run with each one of the different potential/time programmes were recorded.

The counter electrode was also made of pure gold. A 
Hg/HgO reference electrode was used although the 
electrode potentials given in the text are referred to the 
reversible hydrogen electrode (rhe) at the solution pH. 
The following electrolyte solutions (ionic strength ca 1.3) were employed: 1 M NaOH + 0.1 M Na2SO4 ; 
0.1 M NaOH + 0.4 M Na2SO4 ; 0.01 M NaOH + 
0.43 M Na2SO4, . These solutions were made from AR 
chemicals and triply distilled water. The electrolytic 
solutions were continuously kept under purified ni-	ron. Extreme precautions were taken during the 
experimental work to avoid impurities from the labo-
atory atmosphere. Taps and ground joints were 
lubricated with the same electrolyte solution and the 
gas tubing to be used under a continuous positive 
pressure was made in part with thick wall PTFE and in 
part with glass.
Water purity was systematically tested through the characteristics of the voltammogram of the Pt/I M H₃SO₄ interface in the region of hydrogen adatoms using platinum electrodes[24]. The purity criterion was to keep the hydrogen adatom electrooxidation multiple peaks unaltered during the repetitive triangular potential sweeps at 0.05 V/s for 10 min in the cathodic switching potential, Eₐ, = 0 V, to the anodic switching potential, Eₐ, = 1.5 V, range.

Experiments were made at 25°C. The gold/alkaline interfaces were perturbed by the use of the appropriate conventional electronic circuitry including a careful ohmic drop compensation in the following ways. Firstly, a repetitive triangular potential sweep was run between 0.15 V ≤ Eₐ,c ≤ 0.30 V and 0.8 V ≤ Eₐ,a ≤ 1.55 V, at equal anodic and cathodic potential sweep rate (vₐ = vₐ) to attain a stable E/I profile. Then, immediately after the interface was perturbed with the intermediate repetitive triangular potential sweep between Eₐ,c > Eₐ,a and Eₐ,a at vₐ = vₐ (v > vₐ) during the time τ. Finally, after the last intermediate potential sweep (FIPS), a single negative going linear potential sweep was run (Fig. 1). The duration of the intermediate perturbation, τ, was varied.

The triangularly modulated repetitive triangular potential scanning (TMTPS) technique was also independently applied. The following perturbation characteristics were then used: base signal sweep rate 0.1 V/s ≤ vₐ ≤ 2 V/s; base signal switching potentials 0 V ≤ Eₐ,c ≤ 0.2 V and 0.93 V ≤ Eₐ,a ≤ 1.79 V; modulating signal sweep rate 20 V/s ≤ vₐ ≤ 80 V/s; modulating signal amplitude 0.030 V ≤ ΔEₐ ≤ 0.390 V. The complex E/I contours were photographed from a 502A Tektronix oscilloscope screen.

RESULTS

The E/I displays obtained with a triangular potential perturbation in the potential range of the thermodynamic stability of water exhibits a variety of complementary anodic and cathodic current peaks and shoulders[3,10,11]. The shoulders become more clearly distinguished at potential sweep rates greater than 10 V/s. For the sake of simplicity the E/I plane can be divided at the potential E₉, the Au/Au(III) equilibrium potential, so that the potential range below E₉ is denoted as the non-oxide region and that above E₉, as the oxide region[11].

As seen in Figs. 2 and 3 there is a noticeably different behaviour of the electrochemical system whether it covers the non-oxide or the oxide region. In the former case (Fig. 2) for the experiment run at 0.1 V/s between Eₐ,c = 0.03 V and Eₐ,a = 0.96 V just before perturbing the electrochemical interface with the intermediate triangular potential sweeps, the E/I displays exhibit flat anodic and cathodic current envelopes and Qₑ, the total cathodic charge, is slightly greater than Qₐₑ, the total anodic charge. During the intermediate triangular potential perturbations, the Qₑ/Qₐₑ ratio of each potentiodynamic cycle is always slightly larger than one. This is also the case for the following negative going potential sweep from Eₐ,a to Eₐ,c. For the latter, the anodic E/I profile is slightly modified while the cathodic one is more extensively changed as the charge tends to accumulate on the negative potential side.

The changes produced by the intermediate fast potential perturbation in the non-oxide region are relatively less than those found in the oxide region (Fig. 3). The changes produced in the negative potential going E/I profile when Eₐ,a extends into the oxide region where the Au(OH)₃ species predominates[25], depend considerably on Eₐ, the cathodic switching potential of the fast triangular potential perturbation. Thus, when the potential difference Eₐ,a - Eₐ,c involves the removal and reforming of ca 30 per cent of either the anodic or the cathodic charge, a net accumulation of cathodic charge is observed which is
mostly distributed towards the negative potential side of the $E/I$ display overlapping to a large extent with the charge corresponding to the non-oxide region. Then, the height of the main cathodic current peak not only slightly increases but its shape becomes broader than that of the conventional stabilized cathodic current peak\cite{11}. Under these circumstances the electrochemical interface appears relatively active during the intermediate fast perturbation. However, the situation changes according to the $E_{m,c}$ values. Thus, as $E_{m,c}$ is fixed at more positive values, the charge increase during the negative going excursion is considerably reduced. The shape of the cathodic current peak is very symmetric and thin and the cathodic current peak potential is located at a potential more negative than that of the current peak obtained through a conventional triangular potential scan\cite{11}. The charge obtaining during the fast perturbation decreases as the number of successive cycles increases.

The preceding description also extends to more positive $E_{m,c}$ values, but the largest effect apparently occurs when $E_{m,c}$ coincides with the potential of the first anodic current peak located in the oxide region\cite{4,11}. Nevertheless, as $E_{m,c}$ becomes more positive, namely $E_{m,c} - E_{c,c}$ decreases, the electrochemical interface tends to passivate. Furthermore, the influence of the potentiodynamic ageing in the oxide region on the $E/I$ display in the non-oxide region is considerably diminished as $E_{m,c}$ is more positive. The effect of the potentiodynamic ageing is more noticeable as the time of the fast potential perturbation increases.

In conclusion, the effects produced by the potentiodynamic ageing of the gold/alkaline solution interface can be summarized as follows.

(i) An increase of the cathodic charge in both regions of the $E/I$ display.

(ii) A definite change of the cathodic $E/I$ profile which depends on the duration of the intermediate perturbation.

(iii) A shift of the negative potential going $E/I$ profile towards more negative potentials as should be expected for the appearance of more stable species than those produced under conventional perturbations.
Additional information about the gold/alkaline solution interfaces, particularly that concerning the number of anodic and cathodic current peaks as well as their kinetic responses can be derived through the TMTPS experiments. The corresponding \( E/I \) displays obtained with a small modulating signal amplitude (\( \Delta E_m = 0.03 \) V) (Fig. 4) exhibits the following features. During the positive going potential excursion in the potential range AB (Fig. 4a) only the periodic charging and discharging of the double layer is observed. In the potential range BC there is a reversible faradaic contribution together with the dl charging and discharging but the \( E/I \) display exhibits no particular fine structure. Finally, in the potential range CE, a rather sluggish faradaic process is involved. During the reverse scan in the DE region the dl charging and discharging is predominantly observed. Within the EF potential range a slow faradaic process is noticed and the remaining portion of the negative potential going excursion approaches the shape of the \( E/I \) contour already described for the positive going potential excursion within the same potential range.

Further details are observed when modulating signal amplitudes greater than 0.1 V are employed. Thus, for \( \Delta E_m = 0.120 \) V at \( v_m = 20 \) V/s (Fig. 4b) the positive going potential excursion confirms all the anodic current peaks and shoulders already described. However the difference between the potentials of current peaks \( I_a \) and \( I_c \) is larger than that found at lower \( \Delta E_m \) values (Fig. 4a). The positive going potential excursion (Fig. 4b) shows two anodic shoulders preceding current peak \( V_a \), together with their complementary cathodic currents, namely a cathodic shoulder at 1.07 V, a cathodic current peak at 1.13 V and another cathodic shoulder at 1.19 V. Then, during the positive going potential excursion the \( E/I \) display
in the 0.9 V – 1.6 V range exhibits reasonable reversible electrochemical behaviour. On the other hand, the successive negative going potential excursion in the 1.6 V – 1.2 V range displays only a small faradaic contribution. The main cathodic current peak appears at 0.89 V and at potentials lower than 0.7 V the E/I contour approaches the characteristics already described for the preceding positive going potential excursion. The irreversibility of the overall electrochemical process is more noticeable when the potential is swept from 1.6 V downwards, although it is likely that under these circumstances the cathodic envelope depicted during the negative going potential excursion is to some extent influenced by traces of molecular oxygen produced at the highest E∞ values.

When the frequency of the modulating signal increases (Fig. 4c) the anodic current peaks potentials shift slightly towards more positive values while the cathodic current peaks potentials change to the reverse direction. In general, the distribution of the current peaks previously described through the triangular potential sweep E/I displays either in the absence of any ageing [11] or with potentiodynamic ageing covering different potential ranges. Furthermore, the TMTPS experiments give a more clear understanding about the reversibility of the initial processes related to the electroformation of the oxygen-containing film and the increasing irreversibility of the electrochemical reaction as the potential sweep moves towards the positive potential side. However, the greatest irreversibility is observed in the first half portion of the TMTPS envelope when the potential excursion reverses towards the negative potential side.

**DISCUSSION**

The potentiodynamic E/I profiles for the formation and reduction of surface oxide films on gold in alkaline electrolytes exhibit a number of similar features with those studied more in detail in acid electrolytes [21, 22, 26–28]. The similarities may be both of fundamental and general significance in explaining the early stages of the metal electrooxidation, the stability of the different species formed in the course of the film forming reaction and, eventually, the particular influence of anions on the processes generally described under the heading, ageing effects.

It is already well established that the potentiodynamic E/I profile of the gold/alkaline solution interfaces exhibit an appreciable faradaic current contribution in the 0.45 V – 1.0 V, that is in the non-oxide region, where the gold/water solvent interface is stable in terms of the thermodynamics derived from the bulk properties of gold and water. The processes occurring there, at the positive side of the potential of zero charge of the metal in these solutions [5], should be ascribed to the same initial stage which has been observed in acid electrolytes [21, 22, 27, 28], namely the reversible formation of OH-electrosorbed species. In this case, however, the reaction should be put forward in terms of OH− species instead of water molecules as in the acid electrolyte [29]. Although the experimental results are usually less distinctive than simple models, the kinetics of the electroadsorption reaction of either charged or neutral substances on a metal surface can be described in terms of partial charge transfer reactions [30–33]. Then, the reaction in the non-oxide region can be formally presented according to:

\[
\{x\text{Au}\} + y\text{OH}^- = \{[\text{Au}_x][\text{OH}]_y\}^{(z/2)-} + (y-z)\text{e} \ (1)
\]

where \(x\) refers to the number of available gold surface sites and \(x \geq y\). When \(x = y\) a monolayer of OH species are formed. According to (1) the electrosorbed species may be partially charged and eventually when \(z = 0\), the maximal monolayer charge corresponding to the OH species is formed. Therefore, (1) may involve a partial charge transfer electroadsorption of OH− species. The fact that the overall charge taking part during the process occurring in the potential range of the “non-oxide” region is, in principle, smaller than that expected for the OH monolayer, may be related to the fact that surface saturation is achieved for \(y\)-values smaller than \(x\) or, to the fact that although \(x = y\), the value of \(z\) is larger than \(0\). The monolayer charge is arbitrarily defined by comparison with the hydrogen adatom monolayer on polycrystalline platinum [34] taking into account that the roughness factor of polycrystalline gold is 1.3 [37–39]. The latter value, however, can be larger than 3 for etched gold electrode [40]. At any rate, the absence of any ageing process within the non-oxide potential region and the fast response under the TMTPS perturbation indicate that (1) has to be considered as a fast electrode process. Spectroscopy reflectance [4] and ellipsometric [7–9, 41] data support the existence of surface species in the non-oxide region.

Finally, in the non-oxide region a third relatively minor current contribution is observed as a very small cathodic current peak at 0.53 V during the cathodic excursion in the E/I displays resulting immediately after the intermediate perturbation, but at present no explanation for this small charge can be advanced.

The main faradaic process represented by (1), as revealed through the TMTPS runs is, however, a composite one, as in the non-oxide region at least two apparently successive faradaic processes are distinguished. These results confirm the complex E/I displays previously obtained from fast potentiodynamic E/I displays run with a progressive increase of the anodic switching potential \(E_{\text{sw}}\) [11].

The charge estimated for each one of the faradaic contributions of the E/I profile in the non-oxide region is about the same. Therefore, if one assumes that the overall processes occurring in the non-oxide region yields an OH monolayer when \(z = 0\) in (1) and \(x = y\), it is possible to explain the two conjugate current peaks through the following successive reactions taking place along the positive potential going excursion:

\[
\{x\text{Au}\} + x\text{OH}^- = \{[\text{Au}_x][\text{OH}]_x\}^{(z/2)-} + (x\text{e}/2) \ (1a)
\]

and

\[
\{[\text{Au}_x][\text{OH}]_x\}^{(z/2)-} = \{[\text{Au}_x][\text{OH}]_x\} + (x\text{e}/2) \ (1b)
\]

where the parentheses denote the surface species which
are formed during the two successive partial transfer processes. The stability of each species can be measured through the corresponding current peak potential. Both (1a) and (1b) behave as reversible electrochemical processes. However, these reactions must be considered only as limiting processes since, as described further on, the $z = 0$ condition is apparently attained when the potential exceeds $E_T$. When $E_{1,a}$ is increased beyond 1.0 V a gradual change of the $E/I$ display run towards the negative going potential direction is noticed. As the distribution of the different metal–oxygen species formed on the metal can be deduced from the shape of the negative potential going $E/I$ contour (Fig. 3), one concludes that the largest number of species is formed when the potentiodynamic ageing involves a charge which is slightly larger than one-half the overall charge obtained in the whole potentiodynamic $E/I$ profile run between $E_{1,a}$ and $E_{1,a}$. Under these circumstances the overall electroreduction $E/I$ profile, and the main electroreduction current peak are shifted towards the negative potential side. However, the overall cathodic charge recorded immediately after the intermediate perturbation becomes larger than that found in the initial triangular potential sweep $E/I$ display. This effect diminishes noticeably when $E_{1,a}$ moves towards more positive potential, so that the cathodic charge recorded before and after the intermediate triangular potential perturbations becomes almost alike. This suggests that the more drastic effect promoted by the intermediate perturbation when $E_{1,a}$ is more negative can be assigned to a possible penetration of oxygen beyond the first atomic layer of gold. In this way more strongly bound oxygen-containing species in the metal can be produced, which are more difficult for electroreducing. A similar type of surface rearrangement has been recently discussed for the case of platinum/acid electrolytes interfaces under the same type of perturbation conditions[16].

These results give further support to the idea that the ageing process, especially that related to place exchange reactions at the metal surface should be ascribed mainly to the species produced after the OH electrosorption process, that is, to metal–oxygen species. The latter species through these processes are able to attain more stable configurations.

The oxide region involves the occurrence of three different electrooxidation processes which should start from the electrosorbed OH species. These reactions can be formally written as follows:

$\{[\text{Au}_{x}][\text{OH}]_y\} + (z + y)e = \{[\text{Au}_{x}][\text{OH}]_y\} + (z + y)e$ (2)

$\{[\text{Au}_{x}][\text{OH}]_y\} + y\text{H}^+ = \{[\text{Au}_{x}][\text{OH}]_{2y}\} + ye$ (3)

$\{[\text{Au}_{x}][\text{OH}]_{2y}\} + y\text{OH}^- = \{[\text{Au}_{x}][\text{OH}]_{3y}\} + ye$ (4)

where $y + y' = x$.

Reaction (2) represents the residual discharge of the OH species and it is assigned to the reversible conjugate couple which is observed in ca 1.2 V only during the positive potential excursion TMTPS $E/I$ contour (Fig. 4). The relative charge related to (2) is small and appears as a small shoulder on the negative potential side of the main anodic current peak under the triangular potential sweep $E/I$ display. This reaction is not observed during the negative going potential excursion in the TMTPS runs. Since then no OH-containing species remain on the electrode surface once the $E_{1,a}$ value has been reached. Reactions (3) and (4) are related to the formation of a surface layer approaching the Au(OH)$_3$ stoichiometry[25] and correspond to processes appreciably much slower than those represented by (1) and (2).

The shift of the main electroreduction current peak due to ageing of the species anodically formed in the oxide potential region indicates that the $\{[\text{Au}_{x}][\text{OH}]_{3y}\}$ species acquires a more stable configuration through chemical reactions which can be expressed as:

$\{[\text{Au}_{x}][\text{OH}]_{3y}\} = \{[\text{Au}_{x}][\text{OH}]_{3y}\} + n\text{H}^+$. (5)

Reaction (5) would involve an oxygen exchange in the film, where the place exchange mechanism is assisted by the OH$^-$ ion at the interface. Reaction (5) attempts to represent an average type process related to the ageing effects at the gold/alkaline electrolyte interface. This leaves open the possibility that different types of aged species can be produced.

Since the anodic reaction yields different species the electroreduction process is also a composite one involving different reactants whose contributions during the potentiodynamics excursion depend upon the perturbation programme applied to the electrochemical interface. These reactions can formally be written as:

$\{[\text{Au}_{x}][\text{OH}]_{3y}\} + (2x)e = \{[\text{Au}_{x}][\text{OH}]_{3y}\} + (2x)e$ (6)

$\{[\text{Au}_{x}][\text{OH}]_{3y}\} + n\text{H}_2\text{O} = \{[\text{Au}_{x}][\text{OH}]_{3y}\} + (2x)e$ (7)

and

$\{[\text{Au}_{x}][\text{OH}]_{3y}\} + xe = \{x\text{Au}\} + x\text{OH}^-$. (8)

The largest irreversibility of (7) should involve not only the place exchange mechanism in the film but also the water rearrangement at the electrochemical interface as part of the elementary steps of electrosorption processes[42]. Furthermore, the place exchange-mechanism related to the oxide film formation is influenced by the nature and concentration of the anions present in the electrolyte. The usual result is the local stabilizing influence of anions on the ageing effect in the metal surface[43].

The present results correlate with the fact that complex changes of the experimental ellipsometric data for the Au/I N KOH interface, especially in the monolayer region, should be explained by the formation of surface species whose properties are both potential and time dependent[7-9].

From the slopes of the individual intermediate potentiodynamic fast triangular potential perturbations at each potential the change of resistance of the film can be followed. Thus, when the $E_{1,a} - E_{1,a}$ potential difference is located towards more positive potential, the slope of the $E/I$ display progressively
decreases. This indicates that the electrical resistance increases as the charge of the oxygen containing layer also increases. This correlates with the conclusions reported for the acid electrolyte that the mechanism of growth of the oxygen containing layer changes with increasing overvoltage[39]. The reason for this change is, however, not clear and deserves further investigation, especially in that which concerns the alkaline electrolyte.

Finally, the influence of the increasing concentration of SO\textsubscript{2}\textsuperscript{-} ions is similar to that already reported for the acid electrolyte[43], namely to compete through specific adsorption with other ions and solvent molecules to block the surface. The larger the SO\textsubscript{2}\textsuperscript{-} ion concentration the larger the overvoltage to accomplish similar effects while a poorer resolution of the E/I profile is observed. These effects are, in any case, less noticeable in the alkaline than in the acid electrolytes.

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