Primary oxide spillover DL pseudo-capacitance charging and discharging in the light of the first principle theory

ABSTRACT

Due to the irreversible disproportionation of the primary (Pt-OH), into much more stable surface (Pt=O) oxide, all plain (Pt) and non-interactive supported (Pt/C) noble metals, within a wider (more than 600 mV) potential range between hydrogen and oxygen evolving limits are deprived from the former, and thereby behave partially, but broadly polarizable. Since almost all interactive catalytic hypo-d-oxide supports in wet hydrated state feature continuous and renewable latent storage and spillover of the Pt-OH, such composite catalysts (Pt/WO$_3$, TiO$_2$) enable provided the reversible electrocatalytic properties for the oxygen electrode reactions. Potentiodynamically the latter consists in almost unlimited and reversible DL pseudo-capacitor charging and discharging within the whole potential range in between of oxygen and hydrogen evolving limits. To confirm such DL properties, based and consisted in the Pt-OH dipole spillover, the rigorous First principle analysis has been carried out and yielded with relevant interrelations of broader significance.

Keywords: First Principle; spillover; SMSI {Strong Metal-(Hypo-d-(f)-Oxide) Support Interaction}, primary (Pt-OH) oxide, surface (Pt=O) oxide, DL charging and/or discharging of double layer pseudocapacitance, work function ($\Delta$\Phi); electrocatalysis; heterogeneous catalysis
reversible for the oxygen electrode reactions, but each of them more or less irreversible or polarizable.

Figure 1 - Cyclic voltammograms of polycrystalline Pt scanned in 0.1 M NaOH at sweep rate 100 mV.s⁻¹

In fact, in the absence of the Pt-OH, therefrom imposes such a critical and typical, very pronounced the reaction polarization range (Fig. 1), all along until oxygen potential evolving limits, and in both scan directions, back and forth [4-7]. Such highly reversible (Eq. 1), relative to the subsequent strongly irreversible, and substantially polarizable (Eq. 2) transient and coexistence of two interrelated reaction steps, has been decisive as concerns the overall electrocatalytic properties and behavior of both Pt and Au, and all other noble and even many non-noble transition metal electrodes for the oxygen electrode reactions (cathodic oxygen reduction (ORR) and anodic oxygen evolution OER)). Since the equimolar ratio of the primary and surface oxide concentrations defines the optimal interfering self-catalytic spillover reaction step in cathodic oxygen reduction reaction ↔ mechanism (ORR) [8],

\[
\text{Pt-OH} + \text{Pt=O} + 3 \text{H}_2\text{O} + 3e^- \rightarrow 5 \text{H}_2\text{O} + 2 \text{Pt} \quad (3)
\]

and in particular along the reversible (low slope, 30 mV/dec, or even lower), parts of Tafel line plots, the irreversible disproportionation (Eq. 2) imposes an extremely high reaction polarization barrier that amounts for even more than 600 mVs and then, in absence of the Pt-OH, makes plain Pt (and Au) and non-interactive supported both (Pt/C or Au/C) platinum and gold irreversible for the oxygen electrode reactions within the broader potential range, back and forth. The thermodynamic definition of irreversibility then would state that plain Pt(Au) by no means can feature the reversibility and/or (electro)catalytic activity all along the potential range in aqueous media. In other words, plain and non-interactive (Pt/C and/or Au/C) supported Pt(Au) electrodes themselves by no means can behave electrocatalytic features of the reversible oxygen electrode (ROE) all along the potential axis between hydrogen and oxygen evolving limits. The strongly adsorptive and thence highly polarizable Pt=O (Au=O), deprived from any local and/or the external Pt-OH (Au-OH) source and supply, then defines one of the most pronounced issues of the reaction polarization in the entire electrochemical science: No Pt-OH, means that there is no reversible reaction (Eq. 3). Quite another story arises when nanostructured Pt(Au) electrocatalyst is interactive selective grafting bonded on various, in particular mixed valence hypo-d-(f)-oxide supports.

Whereas hydrogen molecules undergo spontaneous, and thence reversible adsorptive dissociation on plain Pt (Pt/C), yielding H-adatoms (Pt-H) to establish thermodynamic equilibrium of the RHE (Reversible Hydrogen Electrode, \(\text{Pt} (\text{H}_2)/\text{Pt}-\text{H}/\text{H}_2\text{O}^-\)), within the above identified critical potential range Pt (Au) is deprived (Eq. 2), from the Pt-OH (Au-OH) to provide reversible properties for the ROE ((\(\text{Pt}(\text{O}_2)/\text{Pt}-\text{OH},\text{Pt}=\text{O})/\text{OH}^-\)), (Eq. 3), and both together assemble the reversible hydrogen fueled LMT (Low and Medium Temperature) PEMFCs. In other words, the Pt-OH (or Au-OH) plays the same thermodynamic constitutional role for establishing the equilibrium for the ROE, as Pt-H does for the RHE, and thence, unavoidably imposes continuous need and requires a permanent source of the former to overcome the reaction polarization by the latter, and enables the reversible electrocatalytic properties of the ROE [6,7]. The substance is that Pt-OH (Au-OH) is insoluble in aqueous media and represents surface species able only for heterogeneous reactions and spillover spreading. Thus, to learn and define behavior and properties of the primary oxides (Pt-OH, Au-OH) for DL charging and overall adsorptive and interactive features, one indispensably needs the First Principle search and analysis.

2. MEMBRANE PROPERTIES OF CATALYTIC HYPO-D-OXIDE SUPPORTS

In such a respect, the first main step in the concept towards the ROE implies that catalytic hyper-d-metals (Pt/Au,Ru) establish with hypo-d-(f)-oxides (or their mixed valence compounds, like Nb₂O₅, TiO₂, CeO₂ or Ta₂O₅, TiO₂, CeO₂) the interactive Brewer type (d-d or d-f) bonding effect [9,10], otherwise well known in heterogeneous catalysis as Tauster [11-13] SMSI (Strong Metal-Support Interaction), one of the strongest in the entire chemistry (Fig.3 in ref. [4]) [5]. Consequently, this way interactive inter-bonded composite electrocatalysts (example, Pt/Nb₂O₅, TiO₂), while...
strong bonding, impose remarkably stretched d-orbitals, and thereby exhibit much weaker adsorptive inter-bonding strengths of intermediates (Pt-H, Pt-OH) in the RDSs (Rate Determining Steps), thence correspondingly facilitated cleavage of the latter, and thereby, increased the catalytic activity both for hydrogen and oxygen electrode reactions. Meanwhile, some other accompanying effects, in particular the ones associated with the Pt-OH spillover, play even more significant role for the latter. In such a respect, the dramatically pronounced Pt-OH spillover effect has earlier and primarily been noticed as the remarkable increased CO-tolerance (Fig. 7 in ref. [4]). First, hypo-d-oxides and their mixed valence compounds, as based on typical d-d-(f)-metallic bonds, exhibit extra high stability in both acidic and alkaline media, and many of them pronounced (above 300 S/cm) electron conductivity (Magneli phases even approaching 1,000 S/cm). Furthermore, majority of hypo-d-(f)-oxides and in particular of higher allervalent numbers, feature prevailingly high percentage of dissociative water molecules adsorption [14,15], (Eq.(1-a)), and thence, the mostly enhanced surface membrane type of hydroxyl ions migration mass transfer [16-18], (Eq. (1-b) and (1-c)),

\[
\begin{align*}
\Sigma Pt + 2 H_2O & \leftrightarrow Pt-OH + H_3O^+ + e^- \quad (1) \\
\end{align*}
\]

ending up with the prevailing electron transfer to the interactive supported metallic catalyst, so that the Pt-OH behaves as a pronounced dipole species [19], and thus, exhibits the strong spillover surface repulsion, transfer and distribution. At the same time, the highly pronounced reversible potentialydomatic peaks testify for the extremely fast overall spillover reaction (Eq. 1), in both directions [2,3], primarily used for DL charging and discharging, and then being ready and available for fast heterogeneous electrocatalytic reactions. Such an oxide network, in particular of polyvalent (high allervalent capacity) hypo-d-elements, when in hydrous state, behaves as an ion exchange membrane [16-18]. In fact, gels (aero and xerogels) are biphasic systems in which solvent molecules are trapped inside an oxide network, and such a material can be considered as a water-oxide membrane composite [16-18].

In such an overall constellation, individual hypo-d-(f)-oxides, and even more so their mixed valence composite compounds, when in expanded (sol-gel synthesis with thorough liquid \( \text{CO}_2 \)) supercritical drying) hydrated surface state (Fig. 2), behave as unlimited latent storage and spillover sources of the primary oxide [4-7,20,21], continuously being renewed and recovered simply by water vapor supply and instantaneously proceeding with spontaneous dissociative adsorption of water molecules.

\[
\begin{align*}
\text{M} + \text{H}_2\text{O} & \leftrightarrow \text{M-OH} + \text{H}^+ + e^- \quad (1) \\
\end{align*}
\]

The overall summation (\( \Sigma \)) for the ion exchange properties of hypo-d-oxide hydrated membrane transferring matrix, under directional electric field effect or polarization, yields the reversible Pt-OH adsorptive and/or desorptive spillover upon, or from metal electrocatalyst reactive surface, resulting that way with the characteristic pronounced reversible effusion peaks of extremely fast electrode reaction [2-7], (Eq. 1-a, 1-b and 1-c, yielding by such a summation the same former Eq. 1),

\[
\begin{align*}
\text{M} + \text{H}_2\text{O} & \leftrightarrow \text{M-OH} + \text{H}^+ + e^- \quad (1) \\
\end{align*}
\]

The starting first principle thermodynamic relation for such a SMSI (examined as M/TiO\(_2\)) system and just defined general reversible peak reaction, Eq. 1, arises therefrom in the following form,

\[
\bar{\mu}_- = -\Delta \mu_{m-M-OH} - \bar{\mu}_e + \mu_{H_2O} + \mu_m \quad (4)
\]

while from the main general self-explanatory relation for electrochemical potential definition,

\[
\bar{\mu}_e = \mu_e - e\varphi = \mu_e - e\psi - e\chi \quad (5)
\]
and particularly for hydronium ion (H$_3$O$^+$, abbreviated for simplification as H$^+$), there from Eq. 5 further follows an alternative relation,

$$\bar{\mu}_H = \mu_H - e\phi_H$$  \hfill (6)

To avoid and eliminate electrochemical potentials ($\bar{\mu}_H$, $\bar{\mu}_e$), by equalizing the right-hand sides of Eq. 4 and 6, and replacing the corresponding value from Eq. 5, there straightforward yields the following relation,

$$\mu_H = -\mu_{M-OH} + \mu_M +$$
\[ + \mu_{H_2O} + e\phi - e\phi_H \]  \hfill (7)

Meanwhile, since $\phi = \phi_H + \chi = \psi + \chi$, and thereby, $\phi_H = \psi - \chi$, there further follows,

$$\mu_H = -\mu_{M-OH} + \mu_e +$$
\[ + \mu_M + \mu_{H_2O} + 2e\psi + e\chi \]  \hfill (8)

Finally, by taking the derivative of Eq. 7, because $\Delta \mu_e = 0$, and since $\mu_{M-OH}$, $\mu_{H_2O}$, and $\psi = \text{const.}$, while $\Delta \mu_H + << e\Delta \chi$, there straightforward results,

$$\Delta \mu_{M-OH} = e\Delta \chi$$  \hfill (9)

exactly as cyclic voltammograms (Fig. 2) reveal: Every change in spillover of the M-OH, or of its chemical potential $[(\Delta \mu)] \downarrow (M-OH)$, as the driving force of the effusion, or, simply its concentration variations, corresponds to the increment of surface potential (Δχ), or the change in capacity of DL because of M-OH dipole adsorption package, or desorption (as the spillover capacity), both of them being fast and substantially reversible, define DL charging or discharging (in its pseudo-capacity). Namely, since $\Delta \mu_H +$ is logarithmic function of hydronium ions concentration, and according to the spillover electrode reaction (Eq. 1), primary oxide (M-OH) and H$_2$O$^+$ ions become simultaneously spent or produced in equivalent (1:1) amounts, while the contribution of surface energy is the linear function of surface potential (prevailing dipole effect), e $\Delta \chi$, then the inequality conclusion, $\Delta \mu H+ << e\Delta \chi$, emerges quite logically.

Since the fundamental definition of work function (Φ) states [22,23] that

$$e\Phi = -\mu_e + e\chi$$  \hfill (10)

and taking into account Eq. 9, by derivation of the latter (Eq. 10), there straightforward follows the broader fundamental relation for M-OH spillover adsorption and/or desorption, which is the same as the DL pseudo-capacitance charging and/or discharging,

$$e\Delta \Phi = e\Delta \chi = \Delta \mu_{M-OH}$$  \hfill (11)

Simple combination of Eq. 10 with Eq. 5, then yields the alternative first principle relation between work function and electrochemical potential,

$$\bar{\mu}_e = -e\Phi - e\psi$$  \hfill (12)

which, for the considered issue of electrochemical cell ($U_{WR}$), assembled between working (W) and stable reference (R) electrode, can be straightforward written to read,

$$\bar{\mu}_e = \Delta U_{WR} = -e\Delta \Phi - e\Delta \psi$$  \hfill (13)

The main contribution both to changes in work function (ΔΦ) and primary oxide dipole spillover (ΔμM–OH), comes from the surface potential increment (Δχ) itself (Eq. 9), as the prevailing part of inner or Galvani potential (Δψ). Namely, Vayenas et al. [24,25] have proved on plentiful systems in solid state electrolyte [26], aqueous media [27-29] and PEM Nafion 112 [30] the basic NEMCA or EPOC promotion relation in heterogeneous catalysis,

$$\Delta U_{WR} = e\Delta \Phi$$  \hfill (14)

that could also be considered as fundamental both in electrode kinetics [31-33] and spillover phenomena [34]. Meanwhile, the latter (Eq. 14), when correlated with the preceding relation (Eq. 13), unambiguously and consequently yields that

$$\Delta \psi = 0 \text{ or } \psi_w = \psi_R = \text{const.}$$  \hfill (15)

which is also implied in above preceding equations (Eq. 8 and 9) [35,36]. Such conclusive and far-reaching statement is in particular reliably clear for the electrode steady state and with high concentration supporting electrolyte.

The whole thermodynamic equilibrium for the general issue of interactive (SMSI) metal/hypo-dioxide supported catalyst, such as M/TiO$_2$, is now illustratively displayed in all interconnected energy details by Fig. 11 in ref. 8. As a consequence, any decrease of the M-OH chemical potential $[(\Delta \mu)] \downarrow (M-OH)$, causes the work function to decrease correspondingly, and vice versa. In other words, any consumption of adsorbed M-OH upon metallic catalyst surface causes to decrease its chemical potential, and this way imposes imbalance within the established equilibrium. Thus, to keep constant both ΔΦ and cell voltage between working (W) and reference (R) electrode ($\Delta U_{WR}$),
the titania phase, by its enriched latent storage of the primary oxide, automatically reacts by its membrane transferring mechanism and supplies hydroxide ions (Scheme I), to emanate further as the primary oxide spillover adsorptive species on exposed metallic catalyst surface by corresponding electrocatalytic reaction (Eq. 1), both from hydrated anatase structure and by reacting of continuously supplied water molecules.

![Scheme I - a & b. Model presentation for the SMSI effect, resulting in the spillover transfer of H-adatoms within the bronze nanostructure (Pt/H_{0.35}WO_3, Scheme Ia), and/or the primary oxide (M-OH) effusion, as a dipole along the hydrated counterpart (Pt/W(OH)_6, Scheme Ib), further continuously transferring them upon the metallic part (Pt) of catalyst, otherwise both originating from the hypo-d-oxide continuously fed by moisture stream to such a composite (M/TiO_2,WO_3) electrocatalyst structure and maintaining them in the reversible interchangeable equilibrium (Pt/H_{0.35}WO_3 ↔ Pt/W(OH)_6), (constructed by Dimitris Tsiplakides).](image)

The overall conclusion associated with otherwise broad experience in electrochemical promotion (NEMCA, EPOC) [22-25] of various chemical reactions in heterogeneous catalysis and particularly for Doebereiner (Ertl interpretation [37]) oxidation of hydrogen in water solutions, for catalysts interactive (SMSI) supported upon hypo-d-oxides, OH - ions play the main membrane transferring and decisive promotion role, while M-OH, as defined dipole state, undertakes the spillover and features as the substantially reacting species all over the metallic catalyst surface, enabling that way to keep established such thermodynamic equilibrium. In fact, there is no catalytic promotion for hydrogen oxidation in acidic aqueous media and apparently hydroxide ions impose such a substantial promoting efficiency, in particular within the interfering (600 – 900 mV vs. RHE) potential range, when NEMCA approaches promotion effects of two orders of magnitude [26,27]. Therefore, any disturbance of such an equilibrium, such as the consumption of spillover adsorbed M-OH species (ΔH\text{H} (M-OH)), or simple imposition of an external polarization, as just analytically shown above, reflects as the driving force (ΔU_{WR} = eΔΦ), and instantaneously tends to reestablish the former equilibrium, or to keep its steady state.

The present analysis is best reflected and proved by comparison of Fig. 1 and 2, once when missing Pt-OH spillover (Pt-OH → 0, Pt=O → 1), and when enriched spillover enables enormous primary oxide adsorptive deposition and reverse desorption, exactly as big capacitors do, and when present all over potentiodynamic cycluses [4-7].

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4. REFERENCES

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IZVOD

PUNJENJE I PRAŽNJENJE PSEUDO-KONDENZATORA ELEKTRičNOG DVOJNOG SLOJA PRELIVANJEM PRIMARNOG OKSIDA U SVETLU TEORIJE PRVOG ZAKONA TERMODINAMIKE

Usled ireverzibilne disproporcionacije primarnog (Pt-OH), u znatno stabilniji površinski (Pt=O) oksid, svi čisti (Pt) i neinteraktivno naneti (Pt/C) plemeni metali, unutar šireg (više od 600 mV) potencijalnog intervala između granica katodnog razvijanja vodonika i anodnog izdvajanja kiseonika, lišeni su prisustva ovog prvog (Pt-OH), i stoga se ponašaju delimično polarizabilno u širem opsegu potencijala. Budući da gotovo svi interaktivni hipo-d-oksidni katalitički nosači u svom vlažnom hidratnom stanju, ispoljavaju neprekidnu i obnovljiva latentna svojstva skladištenja i prelivanja primarnog (Pt-OH) oksida, takva vrsta kompozitnih (binarnih) katalizatora (Pt/VO₃, TiO₂), obezbeđuje reverzibilna elektrokatalitička svojstva za kiseonične elektrodne reakcije. Potenciodinamički ovo se sastoji u gotovo neograničenom i reverzibilnom punjenju i pražnjenu dvojnoslojnog (DL) pseudo-kondenzatora unutar čitavog potencijalnog opsega između granica razvijanja kiseonika i vodonika. Da bi se dokazala i potvrdila takva električna dvojnoslojna svojstva, koja se zasnivaju i sastoje iz dipolnog Pt-OH prelivanja, izvedena je dosledna analiza posredstvom Prvog zakona termodinamike, koja je dovela do pouzdanih i relevantnih medjuzavisnosti od šireg značaja.

Ključne reči: Prvi zakon termodinamike; prelivanje; SMNM (Snažan Metal-(Hipo-d-(f))-Oksidni Nosač (uzajamnog) Međuelovanja); primarni (Pt-OH) oksid, površinski (Pt=O) oksid, DL (Dvojnoslojno punjenje (ili pražnjenje) pseudokapaciteta); Izlazni rad (funkcija izlaznog rada elektrona (∆Φ)); elektrokataliza, heterogena kataliza.

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