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# Journal Name

#### COMMUNICATION

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# On the dual character of charged metal-molecule hybrids and the opposite behaviour of the forward and reverse CT processes J. Roman-Perez, S. P. Centeno, M. R. López-Ramírez, J. F. Arenas, J. Soto, I. López-FORWARD CT: $M \leftrightarrow A$ REVERSE $M \rightarrow A$ $M \leftarrow A$ LUMO

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DFT calculations predict two different electronic structures of metal-molecule hybrids which are selected by the surface charge. While the metal-to-molecule CT states are very sensitive to the charge, the energies of the reverse moleculeto-metal CT processes are surprisingly not modified at all by the charge of the metal.

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Charge transfer (CT) in metal-adsorbate (M-A) hybrids is a key process involved in many technological devices.<sup>1,2</sup> In spite of the large amount of studies dealing with this simple process some basic questions remain not yet well understood. This is the case of the unexpected efficiency of the electrode potential (Ev) in tuning the energy of the forward metal-to-molecule CT states (ECT) observed, for instance, in surface-enhanced Raman scattering (SERS).<sup>3,4</sup> Huge energy gains ( $G=\Delta E_{CT}/\Delta E_V$ ) up to 5 eV/V have been reported from electrochemical SERS experiments. This cannot be understood from a classical view of the CT process (Schemes 1 and S1), given that the energy of the forward or reverse (F/R) metal-molecule CT-processes would be red or blue-shifted ( $\Delta E_{CTF}/\Delta E_{CTR}$ ) as much as -1 or +1 eV, respectively, when the electrode potential shifts  $-1 \text{ V} (-\Delta E_V)$ .<sup>5</sup> The main shortcoming of the very popular CT models analogous to Scheme 1 is that both metal and adsorbate are considered as noninteracting subsystems, what implies the equivalence between the slopes in the Schemes S1b and S1c ( $G_F = -G_R = 1 \text{ eV/V}$ ) irrespective of the nature of M and A.

This conclusion is in contradiction with experimental values of G which are very sensitive to the nature of the complex as well as to other experimental variables. Recently, we demonstrated<sup>6</sup> that two contributions are involved in the energy gain  $G=SC=E_{CT}/E_V$ . C is a

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Scheme 1. Standard energy diagram to visualize metal-molecule M-A CT processes between the Fermi level of the metal and the occupied (HOMO) or vacant (LUMO) orbitals of the adsorbate, showing the effect of a negative shift of the electrode potential (- $\Delta E_{\nu})$  on the Fermi level (E\_{FL,E\nu+} \rightarrow E\_{FL,E\nu-}). This originates a red shift of the forward (F) metal-to-molecule  $CT_F$  energy ( $E_{FL}(M) \rightarrow LUMO(A)$ ) or a blue shift of the reverse (R) molecule-to-metal CT\_R energy (HOMO(A)  $\rightarrow$  $E_{FL}(M)$ ). From these widely used schemes, an equivalence between both energy quantities ( $\Delta E_{CT} = e \Delta E_V$ ) is deduced giving trivial energy gains of  $G = \pm 1 \text{ eV/V}$ , respectively. This is in disagreement with both the experimental results for CTF  $(G_F >> 1 \text{ eV/V})$  and the prediction made in this communication for reverse  $CT_R$ process when the negatively charged isonicotinate is involved ( $G_R \approx 0 \text{ eV/V}$ ).

kind of electrical capacitance that quantifies the capability of the metal in converting the applied macroscopic electrode potential in changing the charge of the surface metal atom to which the molecule is adsorbed ( $C=q/E_V$ ). In turn, S quantifies the dependence of the energy of the CT states of the hybrid system on the atomic surface charge ( $S=E_{CT}/q$ ). As we shall show, S is the most subtle and unforeseeable contribution given that it depends on the particular state, the nature of the metal and the molecule and on the experimental conditions too. Widely used CT diagrams like Schemes 1 or S1 are absolutely incompatible with some experimental evidences and the predictions derived from the present work. In them, the Fermi level of the metal  $(E_{FI,Ev})$  is raised up +1 eV when a negative shift of -1 V is applied to the electrode potential. Moreover,

the CT<sub>F</sub> or CT<sub>R</sub> processes are originated by the transfer of a complete electron between the Fermi level of the metal (E<sub>FI,Ev</sub>) and vacant (LUMO) or occupied (HOMO) molecular orbitals of the adsorbate, respectively. Scheme 1 cannot explain how a  $\Delta E_V = 1$  V can be amplified up to  $\Delta E_{CT,F} = 4$  eV,<sup>6</sup> but it seems even more difficult to explain how the reverse mechanism could be insensitive to the electrode potential. This is the subject of this work.

Many CT processes take place in true M-A complexes where M and A are chemically bonded and cannot be considered as separate moieties. In these cases, CT occurs between the ground electronic state of the hybrid and excited states with CT character. The effect of the electrode potential is just to modify the surface excess of charge of the metal and this process is controlled through the electrical capacitance C. Each surface charge generates a new M-A chemical species and modifies all the electronic structure of the particular hybrid in an unknown, either large or small, extension. This affects the properties of the ground electronic state, like the M-A bond energy, and especially those of the CT excited states. This subtle mechanism is controlled by the slopes *S*.

At the present time, the only operative way to get insight into the complex CT mechanism in true M-A hybrids is to resort to electronic structure calculations. In two previous Communications<sup>7,8</sup> we have introduced a methodology to study theoretically the relationship between the charge of the metal and the electronic structure of the surface complex, i.e., the energy of the CT states. It is based on timedependent (TD) DFT calculations carried out on the ground and excited states of complexes formed by a molecule bonded to linear stick-like metal clusters  $(M_n)^q$  with different sizes (n=2,3,5 and 7) and charges (q=0,±1 a.u.), giving the corresponding metal-adsorbate  $[(M_n)^q-A]$  systems (Figure 1a, bottom). These  $(M_n)^q$  clusters allow to account smoothly for the effect of a fractionary charge excess on the electronic structure of the M-A complex. In all the ground and excited electronic states some amount of charge is interchanged between M and A, but true CT states are easily recognized by comparing the corresponding Mulliken's net charges of the Agn clusters (or the adsorbate) in the ground S<sub>0</sub> and the corresponding singlet excited state S<sub>i</sub>. More complex redistributions of charge between M and A like back-donation, are not discussed here. Theoretical calculations have been carried out by using Gaussian 09 suite or programs<sup>9</sup> and are detailed in ESI where a complete set of selected results concerning the ground S<sub>0</sub> and the CT states are summarized. The mentioned previous works<sup>6-8</sup> were focused on the excited CT states whose energies are linearly dependent on q<sub>eff</sub>=q/n:

$$E_{CT} = S q_{eff} + E^0$$
 (1)

 $q_{eff}$  is a microscopic analogue to the macroscopic charge excess  $q_{M}$  (C/cm<sup>2</sup> of surface) and quantify the atomic charge excess averaged amongst the metallic atoms (a.u./atom of surface), ranging from -0.33 to +0.33 for the respective  $Ag_3^{-1}$  and  $Ag_3^{+1}$  complexes. In the cases of pyridine, pyrazine and isolated or protonated adenine linked to Ag or Rh clusters the CT process is dominated by the forward metal-tomolecule M® A electron transfer in the UV-VIS range of energies.<sup>6-8</sup>

Fig. 1a shows the dependence of the TD-M06-HF<sup>10</sup>/LanL2DZ<sup>11</sup>



**Figure 1.** a) TD-M06-HF/LanL2DZ vertical energies (E) of the forward (CT<sub>F0</sub>, CT<sub>F1</sub>, CT<sub>F1</sub>, CT<sub>F1</sub>, and reverse (CT<sub>R0</sub>, redrawn in c)) charge transfer states of [Ag<sub>n</sub>-Py]<sup>q</sup> complexes and the first excited singlet of pyridine (S<sub>1,Pγ</sub>) vs. the effective charge q<sub>eff</sub>. b) TD-M06-HF/LanL2DZ energies of the same states but referred to the energy of the respective metallic clusters (Ag<sub>n</sub><sup>q</sup>) and the adsorbate (Py) at infinite separation (E<sub>M...A</sub>) (each label is accompanied by the slope *S* (eV/a.u) in eq. 1).

vertical energies of selected singlet excited states of silver-pyridine complexes [Agn-Py]<sup>q</sup> on qeff. Four series of forward metal-to-molecule CT<sub>F</sub> states below 6 eV have been drawn.<sup>7</sup> All of them show a linear dependence on qeff but very different slopes. The two lowest ones, CT<sub>F0</sub>;B1 and CT<sub>F1</sub>;A2 are very sensitive (S=6.71 and 7.43 eV/a.u. respectively) to q<sub>eff</sub> being stabilized by some 4 eV when q<sub>eff</sub> varies from +0.33 to -0.33 ( $\Delta q_{eff}$ =-0.66). For the sake of comparison, the energies of the first excited singlet of pyridine S1, Py; B1 are also represented. This excitation is mainly localized in the adsorbate (HOMO→LUMO in Scheme 1) and therefore, no dependence on q<sub>eff</sub> should be expected given the very small transferred charge in this case  $(\Delta q_t = q_{Py,Si} - q_{Py,S0})$  which amounts less than 10% of the corresponding CT<sub>Fi=0,1</sub> series. In spite of this, S<sub>1,Py</sub> shows an S value much larger than the secondary series of true CT<sub>Fi=1,0</sub>' states. The differentiated behaviour of the five states is quantified through the respective slopes. All of them are positive but numerically very different, ranging from  $S_F=0.39$  up to 7.43 eV/a.u. On the contrary, the first series of molecule-to-metal CT<sub>R0</sub>;A<sub>2</sub> states lie above 6 eV, showing a negative slope as expected for reverse processes ( $S_R$ =-4.05 eV/a.u.), but much smaller than CT<sub>Fi</sub>. This complex and unexpected behaviour cannot be understood at all on the simple basis of Scheme 1 given that S is very dependent on the particular CT (or even non-CT) considered state. Fig. 1a also indicates that Py does not undergo redox processes in the ground So state in the overall range of scanned  $\Delta q_{eff}=0.66$ , but it can be transitorily photoexcited up to a CT<sub>F</sub>;Ag<sup>+</sup>-Py<sup>-</sup> state at q<sub>eff</sub>£-0.15 a.u. under 514.5 nm VIS radiation, what accounts for the selective enhancement of Raman lines due to a resonant metal-to-molecule CTF process detected in the SERS recorded at negative electrode potentials.8

Figure 1b shows the energies of the same states but referred to a common zero namely, the energy of the metallic clusters  $(Ag_n^q)$  and the adsorbate (Py) at infinite separation  $(E_{M...A})$ . This makes evident the stabilities (bond energy) of the respective complexes in



**Figure 2.** Dependence on q<sub>eff</sub> of: a) TD-M06-HF/LanL2DZ vertical energies (eV) of the forward (CT<sub>Fi=0,1</sub>) and reverse (CT<sub>Ri=0-2</sub>) charge transfer states of silver-isonicotinate complexes [(Ag<sub>n</sub>)<sup>q</sup>-Din']<sup>q-1</sup> (each label is accompanied by the slope S (eV/a.u) in eq. 1), b) TD-M06-HF/LanL2DZ energies of the same states but referred to the energy of the metallic clusters (Ag<sub>n</sub><sup>q</sup>) and the adsorbate (in-) at infinite separation (E<sub>M...A</sub>). Arrows correspond to the discussed forward (solid) and reverse (hollow) CT transitions from the ground electronic state S<sub>0</sub> and the colours are arbitrarily assigned. c) C<sub>2</sub> complexes of isonicotinate (Ag-Oin').

their ground state  $S_0$  which are also very dependent on  $q_{eff}$ . As expected, a positive  $q_{eff}$  stabilizes the adsorption of Py on the surface through the nitrogen atom giving a perpendicular surface orientation. The M-A bond energy in  $S_0$  amounts to -1.7 eV (-38.8 Kcal/mol) at  $q_{eff}$ =+0.33, while the  $[Ag_3-Py]^-$  complex should be weakly bonded. The slopes for each series of states are modified from those of Figure 1a, showing either positive (CT<sub>Fi=0,1</sub>) or negative values (CT<sub>R0,</sub> CT<sub>Fi=0,1</sub>',  $S_0$  and  $S_{1,py}$ ). As a conclusion,  $q_{eff}$  modifies smoothly the electronic structure of Ag-Py surface complexes by tuning the energies of CT and even non-CT states in a continuous way in the range  $\Delta q_{eff}$ =0.66.

A similar study has been carried out on the isonicotinate anion (in- = 4-pyridinecarboxylate), a negatively charged derivative of pyridine in which the reverse molecule-to-metal CT<sub>R</sub> process is expected at much lesser energy than in the case of neutral adsorbates like pyridine. This species can be bonded to silver through the nitrogen as in pyridine (Nin-), or through the oxygen atoms of the carboxylate (Oin-), giving the respective  $[(Ag_n)^q\text{-}Nin\text{-}]^{q\text{-}1}$  or  $[(Ag_n)^q\text{-}$  $Oin-]^{q-1}$  complexes which have been also constrained to  $C_{2v}$ symmetry (Fig. 2c). Fig. 2a shows the firsts series of TD-M06-HF CT states for the most stable  $[(Ag_n)^q\mbox{-}Oin\mbox{-}]^{q\mbox{-}1}$  complex where a completely opposite behaviour of the forward and reverse processes can be seen. Under positive or zero  $q_{\text{eff}}$  the first two series of metalto-molecule forward CT<sub>Fi=0,1</sub> states are continuously tuned as in pyridine, showing large slopes of  $S_F$ =4.90 and 6.17 eV/a.u. respectively, and therefore providing high energy gain  $G_{F}$ . Excitation up to these states originates the transient formation of the respective radical dianion of the adsorbate (Oin<sup>•2-</sup>), a species far less stable than in the case of neutral ones like Py (Py<sup>1</sup>). In spite of the repulsion between the two negative charges,  $CT_{F0}$  is also reachable under 514.5 nm excitation at negative  $q_{\text{eff}}$  as we observed in the SERS of the analogous benzoate anion.<sup>12</sup> Forward CT<sub>F</sub> states disappear at positive surface charges, being replaced by two series of near degenerate reverse  $CT_{Ri=0-1}$  states at ca. 4.50 eV ( $\Delta q_t$ =0.35/0.55 a.u.) plus another one with lesser CT character (CT<sub>R2</sub>: 4.75 eV and  $\Delta q_t$ =0.2/0.25 a.u.). As expected,  $CT_{Ri}$  for the anion Oin<sup>-</sup> are much more stable than for

neutral Py, but the most remarkable result is the surprising lack of dependence of the respective energies on  $q_{eff}$ , i.e., the surface charge does not tune at all the reverse molecule-to-metal electron transfer processes. Therefore no energy gain  $G_R=CS_R\approx 0$  is predicted given that  $S_R\approx 0$ .

By inspection of Fig. 2a a qualitative change in the behaviour of the M-A complex can be realized. Negative or positive surface charge selects one of the two different electronic structures of the M-A complex which have been named as P- or C-hybrids, respectively. This structural change is highlighted in Fig. 2b where the energies of the CT and the S<sub>0</sub> states are referred to both fragments M and A at infinite separation (E<sub>M...A,</sub> bond energy). At zero or negative charges  $(q_{eff} \pounds 0)$  the stability of the system in its ground state  $(E_{M...A}, S_{0-})$  varies continuously from +38 to -35 Kcal/mol. The largest positive value of +38 Kcal/mol points out to a metastable complex located in a relative minimum where the strong M-A chemical affinity prevents the repulsive dissociation (Fig. S1, ESI). As in pyridine, these [(Ag<sub>n</sub>)<sup>q</sup>-Oin<sup>-</sup>]<sup>q-</sup> <sup>1</sup> weakly bonded complexes at q<sub>eff</sub>£0 can be considered as a case of physisorption (P-hybrid). On the contrary, the bond energies of complexes containing positive silver clusters (E<sub>M.,A</sub>,S<sub>0+</sub>) are less dependent on q<sub>eff</sub> and much more stable, with typical bond energies of chemisorbed species (C-hybrid) ranging from -119 to -133 kcal/mol. Therefore, the electronic structure of M-A systems show very complex dependence on the metal charge what overrides any standard discussion exclusively based on simple Coulombic interactions. In the case of adsorbates with negative charge as in<sup>-</sup>, the electrostatic repulsion between the anion and the metal at negative potentials destabilizes the structure of the surface complex, and the attractive interaction stabilizes it at positive Ev. This causes a disruption in the more or less continuous trend shown by the properties of the pyridine-silver system (Fig. 1b). The electronic structure of Ag-Oin<sup>-</sup> undergoes a qualitative gap when the surface charge changes its sign  $(q_{eff} \ge 0 \leftrightarrow q_{eff} \ge 0)$  as shown in Fig. S3, where the main molecular orbitals involved in the CT states are drawn. A quantitative gap of the corresponding energies can be appreciated when q<sub>eff</sub> is positive/zero or negative which is much less pronounced in the case of pyridine (Fig. S2), what allows to consider the Ag-Py complex as P-hybrid in the overall range of qeff.

Therefore the electrode potential is able to select two kinds of surface hybrids for  $[(Ag_n)^q-Oin^-]^{q-1}$ : P-hybrid characterized by a weak surface complex with properties tunable in a continuous way (q<sub>eff</sub>£ 0), and C-hybrid, a strongly bonded system (q<sub>eff</sub>>0) whose electronic structure is hardly modified by E<sub>v</sub>.

The ground  $S_0$  state of all the discussed M-A complexes shows a net donation of negative charge from the adsorbate to the silver cluster ( $q_{M \leftarrow A,S0}$ ), even at negative  $q_{eff}$  (Fig. S4a). In the case of pyridine, a rather smooth dependence is observed, while for the charged Oin- a dual behaviour can be seen with significant slopes at  $q_{eff}$  20, but being almost insensitive at positive charges. In this sense, the  $S_{0+}$  ground states of C-hybrids at  $q_{eff}$ >0 already show CT<sub>R</sub> character given that, for instance, Oin<sup>-</sup> donates ca. -0.35 a.u. to any of the Ag<sub>n</sub><sup>+</sup> clusters. Finally, Fig. S4b shows the very good correlation between the strength of the M-A bond ( $E_{M...A}$ ) and the charge donated by Py or in<sup>-</sup> to the Ag<sub>n</sub> cluster in S<sub>0</sub> ( $q_{M \leftarrow A,S0}$ ). Once again both kinds of hybrids are easily recognized.

In order to confirm these results a similar study has been carried out on the less stable  $[(Ag_n)^{q_-}Nin^{-}]^{q_{-1}}$  complex. The M06-HF results for both complexes as well as those obtained by using the long-range corrected  $\omega$ B97X-D<sup>13</sup> functional are compared in Tables S1-S3 and Fig. S5. All the results predict very similar CT<sub>F</sub> states for the P-hybrid. On the contrary, the reverse CT<sub>R</sub> states of the C-hybrid are dependent on the type of coordination (Ag-Oin<sup>-</sup> or Ag–Nin<sup>-</sup>) and on the used functional, being the  $\omega$ B97X-D CT energies ca. 2 eV more stable than the M06-HF ones. However, all of them point out to the same conclusions: i) the existence of two surface kind of hybrids and ii) the lack of sensitivity of the energies of any CT<sub>R</sub> state to the surface charge, i.e. on the electrode potential. A detailed inspection of the CT<sub>R0</sub> state of Ag-Py also reflects the presence of two different slopes with little dependence on q<sub>eff</sub> (Fig. 1c).

Summarizing, this Communication discusses for the first time the existence of different kind of hybrids in complexes formed by metals bonded to charged molecules corresponding to different electronic structures which are selected by changing the sign of the charge of the metal. Furthermore, the dependence of the forward and reverse CT processes on the surface charge show an opposite dependence on q<sub>eff</sub>, predicting either a huge gain or an almost zero efficiency when the electrode potential is used to tune the energy of the respective CT process. This allows for selecting, opening or closing different CT channels acting in opposite electron transport directions. In spite of this work deal with small metallic clusters, the conclusions could open new perspectives to a better understanding of the complex behavior of molecules bonded to nanometer or atomic size metallic clusters detected in SERS, single molecule SERS, tipenhanced Raman scattering (TERS) or in molecular electronics. It must be stressed that our methodology based on standard DFT calculations is able to predict very surprising behaviours depending on the nature and the respective charges of the metal and the adsorbed molecule. This will allow to classify in a comprehensive way the different types of metal-molecule hybrids according to their respective electronic structures.

Finally, the dual electronic structure of M-A suggests a differentiated mechanism for the oxidation and reduction what could play a key role in electrochemistry, adsorption or heterogeneous catalysis, providing alternative explanations to some results discussed so far on, for instance, molecular reorientation.

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# **GRAPHICAL ABSTRACT**

Charged molecules bonded to metals shows two kind of electronic structures depending on the charge of the metal

