Work Function Changes Induced by Charged Adsorbates: Origin of the Polarity Asymmetry

Paul S. Bagus,1 Daniel Käfer,2 Gregor Witte,2 and Christof Wöll2

1Department of Chemistry, University of North Texas, Denton, Texas 76203-5070, USA
2Lehrstuhl für Physikalische Chemie I, Ruhr-Universität Bochum, D-44780 Bochum, Germany

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A theoretical analysis of charged adsorbates on a metal surface reveals a pronounced polarity asymmetry between electropositive and electronegative species, thus reproducing a well known but so far not properly understood experimental fact. For ionic adsorbates on metal surfaces, we analyze the several, often canceling, terms that contribute to the change of the interface dipole and, hence, to work-function changes, \( \Delta \phi \). We demonstrate that for the prototypic case of I on Cu(111) the magnitudes and the signs of these terms can be understood on the basis of their physical and chemical origins. An important consequence of their cancellation is that negatively charged adsorbates can lead to a paradoxical \( \Delta \phi < 0 \) rather than the expected \( \Delta \phi > 0 \).

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The adsorption of atomic or molecular species on metal surfaces induces significant changes of the work function, \( \phi \) [1–4]. Historically, such changes have been important for lowering the work function to enhance thermal electron emission, e.g., from W filaments [3], and for monitoring adsorption phenomena [5]. Recently, the precise tailoring of the work function of metal surfaces has become an important task with regard to optimizing charge injection at metal-molecule interfaces, in particular, for applications in organic electronics [6]. For strongly electropositive species, e.g., alkali metal atoms, adsorption is accompanied by an electron transfer to the substrate. As first proposed by Langmuir [3], the resulting dipole layer that is directed positive above the surface should result in a strong lowering of the work function [1]. A reasonable approximation for these changes at low coverages is \( \Delta \phi \sim -N\Delta \mu \), where \( N \) is the adsorbate density and \( \Delta \mu \) is the change in the surface dipole induced by a single adsorbate [1]. For relatively low coverages of alkali metals on metal surfaces this model has been successfully used to explain the experimentally observed large lowerings of the work function by \( 2–3.5 \) eV [1].

In principle, the opposite behavior is expected when an electron acceptor is adsorbed on a metal surface with the charge transfer yielding a dipole layer with the opposite orientation, negative above the surface; hence, \( \phi \) should increase [7,8]. While the expectations derived from the simple Langmuir model have been confirmed many times for electropositive adsorbates, for the opposite case a textbook example, i.e., a case where a simple electronegative adsorbate leads to a work-function increase, is lacking [7,8]. Instead, several cases have been reported [8–10] where an e transfer to the adsorbate is accompanied by a work-function decrease, e.g., I/Ni(100) [9] (\( \Delta \phi = -0.8 \) eV) and I/Pt(111) [10] (\( \Delta \phi \sim -0.7 \) eV). Only a few theoretical works have been presented to explain this rather disturbing asymmetry. In earlier work, Pettersson and Bagus [11] analyzed F– and Cl– on Cu(100) and concluded that polarization of the Cu surface made an important contribution to \( \Delta \phi \). More recently, Michaelides et al. [12] reported that for a charged N+ adsorbed on W(111) the work function is significantly affected by the Pauli exclusion principle. It has been shown earlier by Bagus, Staemmler, and Wöll [13] that this effect explains the surprising lowering of the work function in the case of neutral adsorbates; for rare gas or noble metal systems, see Hückstädt et al. [14] However, N is not a typical electron-acceptor and, accordingly, the concept of charge transfer to explain work-function changes is clearly not appropriate for this adsorbate.

In the following, we present a thorough theoretical analysis comparing Cs, Xe, and I adsorbed on Cu(111). These adsorbates have similar sizes and hence should be suitable to examine the relation between polarity and work-function change. For the electropositive Cs the experimental findings for \( \Delta \phi \) are nicely described by a simple dipole layer model. For the nonpositive Cs the adsorption-induced reduction in \( \phi \) has recently been shown to result from the Pauli exclusion effect [13,15]. However, we shall show that for I, the work function is decreased, in complete contradiction to the expectation of \( \Delta \phi > 0 \) for an electronegative adsorbate. Further, we will show that \( \Delta \phi < 0 \) for I can be directly related to the chemistry of the I-substrate interaction. The theoretical results are validated by our measurements of the work-function changes for I on Cu(111). By using the Helmholtz equation, an interface dipole per I atom, denoted as \( \Delta \mu \), of 1.7 D has been derived from the slope of the measured (using He I UV-photoelectron spectroscopy) \( \Delta \phi(\theta_I) \) curve in the low coverage regime (\( \theta_I < 0.05 \)).

In our calculations—for details see Refs. [13,15,16]—the unrelaxed Cu(111) surface is modeled by a three layer, 32 atom Cu cluster, Cu32, with 19 atoms in the first, 12 in the second, and one in the third layer. This cluster has been used successfully to describe the adsorption of C6H6 and C6H12 on Cu(111) [15]. A Cs, Xe, or I adsorbate, denoted
X, is placed directly above the central first layer atom of Cu(111) to form Cu32X. We chose to place the adsorbates at on-top sites to more easily allow a direct comparison of the various contributions to $\Delta \phi$. The close agreement of our calculated and measured $\Delta \mu$ for I/Cu(111) justifies this approach [17]. We then used \textit{ab initio} methods to obtain wave functions (WFs), energies, and other properties of the clusters. The 1s to 4d core electrons of the I, Xe, and Cs atoms are represented using pseudopotentials with parameters taken from Refs. [18–20], respectively; the basis set for I was augmented with diffuse functions to describe the polarizability [21]. Electron correlation effects were included using second order Moller-Plesset (MP2) perturbation theory [22], because MP2 gives a reliable description of dispersion, or van der Waals, forces [23,24]. The interaction of Cs or I with a metal surface is dominantly an ionic bond, with Cs and I being largely Cs$^+$ and I$^-$; however, we will show that a weak dative covalent bond is formed by donation from I$^-$ to Cu and that this bond contributes significantly to the work-function change induced by I. For convenience, the calculations have been carried out for the isoelectronic clusters of Cu32I-, Cu32Xe, and Cu32Cs$^+$. First, the equilibrium distance, $z_e$, of the adsorbate above Cu(111) was determined from MP2 calculations with the geometry of the copper cluster fixed and only the height of the adsorbate above the surface allowed to vary.

Subsequently, the different contributions to the metal-adsorbate interaction and the work-function changes have been obtained by analyzing the properties of the wave functions at the self-consistent field (SCF) level as done successfully in previous work [13,15,25,26]. An additional analysis has also been carried out for I kept at the position of Cs. In Table I, we list our calculated $z_e$ and $\Delta \mu$ for the three adsorbates. The least strongly bound of the three, Xe, is at the largest distance from Cu(111) and has the smallest $\Delta \mu$; the calculated $z_e$ is only slightly larger than the experimental $z_e = 3.6$ Å [27]. For this case, the sizable reduction of $\Delta \phi$ is dominated by the Pauli exclusion effect [13]. The Cs$^+$ adsorbate is substantially, ~0.5 Å, closer to Cu(111) than Xe. This is expected, since Cs$^+$ is electrostatically attracted by its “image” charge inside the metal [7,28,29]. The large positive $\Delta \mu$ for Cs in Table I is consistent with the large maximum lowerings of $\phi$, by ~3 eV, observed for Cs on metal surfaces [1].

The CSOV steps considered [15] are the following: (1) The frozen orbital interaction, denoted Pauli-FO, where the WFs of the separated entities (Cu cluster and adsorbate X) are superimposed to form an antisymmetric total WF and only physical changes in the WF (i.e., Pauli exclusion) are taken into account. For the charged adsorbates, we also consider the $\mu$ at an initial CSOV step, denoted point charge (PC)–FO or CSOV step(0), where the adsorbate is represented as a point charge of PC = +1 (Cs$^+$) or PC = −1 (I$^-$). The $\mu$ for a PC is different from that for an extended ion at the Pauli-FO step because there is no Pauli exclusion for a PC at CSOV step(0). (2) V(Cu), where the adsorbate orbitals are held fixed but the orbitals of the Cu32 cluster are allowed to vary in the presence of the adsorbate. (3) V(adsorbate), where the substrate orbitals are held fixed after the V(Cu) variation and the adsorbate orbitals are allowed to vary. (4) The final CSOV step is a full SCF calculation, where no constraints are applied. The changes at this step provide a measure of the non-additivity of the different CSOV contributions where small changes indicate that the decomposition is fully additive.

For Xe/Cu(111), the CSOV decomposition (see Ref. [13]) reveals that the dominant contribution to $\Delta \mu$ comes from the Pauli exclusion; the other CSOV terms contribute only about 30%. In the case of the charged Cs$^+$, the very large dipole seen in steps (0) and (1) is strongly

<table>
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<tr>
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<th>$z_e$ (Å)</th>
<th>$\Delta \mu$ (Debye)</th>
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<tbody>
<tr>
<td>Xe</td>
<td>3.86</td>
<td>+0.23</td>
</tr>
<tr>
<td>Cs</td>
<td>3.31</td>
<td>+5.41</td>
</tr>
<tr>
<td>I</td>
<td>2.74</td>
<td>+1.50</td>
</tr>
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reduced by the polarization of the Cu substrate (step 2) and the $\Delta \phi$ for an 0.1 ML coverage of Cs$^+$ is estimated to be $\approx -3.4$ eV, consistent with experimental data [1,32]. However, this situation changes considerably for I/Cu. In Table II, we show the CSOV analyses for I/Cu at $z_e = 2.74$ Å and at $z = 3.31$ Å, which is the $z_e$ for Cs/Cu. For the full SCF variation at the longer distance, $\Delta \mu = -3.1$ D is negative, exactly as expected from the CT model. However at the shorter $z_e$ for I/Cu(111), the work-function change is, unexpectedly, positive, $\Delta \mu = +1.5$ D. A detailed investigation of the CSOV steps establishes the origin of this apparently paradoxical change of sign in $\Delta \mu$.

We consider first the CSOV for Cu$_{32}$I$^-$ with $z(I) = 2.74$ Å. The energetic repulsion for the Pauli-FO WF is $-1.8$ eV, considerably larger than the small repulsion of $-0.03$ eV at this CSOV step for Cs/Cu. The repulsion between the Cu surface charge and I$^-$ leads to a major reduction of charge in the space between Cu and I at the next CSOV step(2) as shown graphically in Fig. 1(a). The Cu conduction band charge flows mainly into the bulk. The variation of the I$^-$ charge at CSOV step (3) leads to substantial changes that are much larger than the changes at this CSOV step for Cs/Cu. At this step, $\mu$ increases by 3.4 D, an increase that is $\approx 10 \times$ larger than that for Cs/Cu, and $\Delta \mu$ for I/Cu(111) becomes positive. The changes in the charge density due to the variation of the I$^-$ orbitals are shown in the $\Delta \rho = \rho[V(I)] - \rho[V(Cu)]$ in Fig. 1(b). There is a loss of density around I, which is due to a combination of polarization of the spherical I$^-$ charge by the positive “image charge” in the surface and by a covalent donation of charge into empty Cu surface orbitals. The buildup of charge in the region between I and the Cu surface is a clear demonstration of a covalent bond. The relatively small number of positive contours reveal that the bond is not particularly strong. Further evidence for a donation of charge from I$^-$ comes from the projection [33] of the anion orbitals on the WF for I/Cu(111).

![Fig. 1](color online). Density difference plots for Cu$_{32}$I showing (a) the Cu polarization and (b) the I polarization and donation. Solid (blue) lines are density increases and dashed (red) lines are density decreases. The plotting plane passes through I and the adsorption site Cu atom; the positions of the Cu atoms, projected onto the plotting plane, are shown as shaded circles. (a) $\Delta \rho = \rho[V(Cu)] - \rho[Pauli FO]$, (b) $\Delta \rho = \rho[V(I)] - \rho[V(Cu)]$.!
that gives 5.9 5p electrons, slightly reduced from the value of 6 for a perfect ionic I⁻. We have used refinements of the CSOV variations [31] to show that the relative contributions to Δμ from I⁻ polarization and from donation are comparable. For both the shorter z(I) and the longer z(I) = 3.31 Å, all of the CSOV variations act to increase Δμ above the point charge value and, at z(I) = 2.74 Å, they are sufficiently large so that Δμ changes sign. At the larger z(I), the increases in Δμ are smaller and Δμ remains Δμ < 0 as expected from the CT model.

In summary, we have shown that, in addition to the dipole layer arising from the charge transfer between an ionic adsorbate and the surface, two other important effects, not symmetric with respect to polarity, must be considered when predicting Δϕ. First, a purely quantum-mechanical effect, exchange, or Pauli, exclusion, leads to an adsorbate induced distortion of the metal charge which, for both polarities, tends to reduce the work function. For electronegative adsorbates, like Cs, the effect of the dipole layer is amplified by this effect while for electronegative adsorbates, the two effects act in opposite directions. Second, and more importantly, the anionic adsorbate is much more polarizable, since it has an extra electron, than the cationic adsorbate, which lost an electron. Thus, (weak) covalent chemical bonds are likely to form between the anion and the metal surface. While it was recognized earlier [12] that this covalent bond affects, to some extent, the work function, we show that the main, very important, effect of the covalent bond is a substantial reduction in bond length. At the shorter bond length, the dipole moment created by the negative charge and its positive charge inside the metal is significantly reduced. The combination and partial cancellation of these effects may then even result in a work-function reduction, as demonstrated here for the prototypic case of iodine, thus yielding the opposite of the prediction of the simple electrostatic model [3].

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[16] For I and Cs adsorbates, there is a minor modification for the treatment of Cu_{32} described in Ref. [13]. Since these adsorbates are closer to the surface than Xe, the adsorption site Cu atom is treated as an all electron atom to ensure that no artifacts due to the pseudopotential are introduced.
[17] Calculations for I at a threefold site of Cu(111) show a similar ionicity as at an on-top site although the distance of I above the surface is slightly shorter at the threefold site.