Controlling polarization at insulating surfaces: quasiparticle calculations for molecules adsorbed on insulator films

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By means of quasiparticle-energy calculations in the \( G_0W_0 \) approach, we show for the prototypical insulator/semiconductor system NaCl/Ge(001) that polarization effects at the interfaces noticeably affect the excitation spectrum of molecules adsorbed on the surface of the NaCl films. The magnitude of the effect can be controlled by varying the thickness of the film, offering new opportunities for tuning electronic excitations in e.g. molecular electronics or quantum transport. Polarization effects are visible even for the excitation spectrum of the NaCl films themselves, which has important implications for the interpretation of surface science experiments for the characterization of insulator surfaces.

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On the nanoscale, materials often reveal extraordinary features. To harness this potential it is essential to grow or manufacture nanostructures in a controlled way. Ultrathin insulating films are an example for which this has been achieved on metal and semiconductor surfaces. We have recently shown that these films develop new and unique properties as their thinness approaches the limit of a few atomic layers and that such supported ultrathin films should be regarded as new nanosystems in their own rights [1]. Here we go one step further and demonstrate by means of first principles calculations that control over the film thickness means control over the polarization of the film. This in turn gives access to properties on the film’s surface, for example the energy levels of molecular adsorbates, which are relevant in the context of e.g. catalysis, molecular electronics, or quantum transport.

The fact that ultrathin insulator films offer a new perspective of control on the nanoscale is increasingly being recognized. Repp et al. for example, have recently demonstrated that gold atoms adsorbed on NaCl/Cu can be reversibly switched between the neutral and negative charge state [2]. Alternatively, the charge state of Au atoms on MgO films can be controlled by the film thickness [3]. For planar molecules adsorbed on Cu-supported NaCl films, the molecular orbitals can be resolved spatially and energetically by scanning tunneling microscopy (STM) and spectroscopy (STS), and electronic reactions can be followed [4]. Interestingly, STS experiments performed on pentacene molecules adsorbed on NaCl/Cu(111) show a significant influence of the film thickness on the molecular gap [5].

Ultrathin insulator films have also developed into highly valuable and intensively studied model systems for characterizing insulating surfaces. The study of insulator surfaces has proven difficult due to their lack of conductivity, which severely limits the range of applicable surface science techniques. Ultrathin insulator films grown on conducting substrates offer a solution to this dilemma because the films can exchange electrons with the substrate by tunneling [2, 3]. Caution has to be applied, however, when transferring thin-film results to the surfaces of technological interest. The properties of ultrathin films may deviate considerably from those of macroscopic films [1] and the excitation spectrum may be affected by the polarization effects presented here.

In this Letter we will address both of these points by means of \( G_0W_0 \) quasiparticle-energy calculations [3, 10] for the example of a prototypical insulator-semiconductor interface (NaCl/Ge(001)) and CO as a model adsorbate. Supported NaCl films are well-behaved model systems for studying the properties of insulator surfaces [3, 11, 12, 13]. Although they are mostly grown on metals, notably Cu [2, 13], Ge(001) is the substrate of choice for studying insulator/semiconductor interfaces [11, 12, 14, 15, 16]. In recent years, these films have also attracted increasing interest in the context of STM and STS studies of atomic and molecular adsorbates [2, 3, 4, 5, 6].

We use density-functional theory (DFT) in the local-density approximation (LDA) to determine the atomic structure of NaCl films on Ge(001). The electronic exci-
tation spectrum is calculated with many-body perturbation theory in the $GW$ approach as perturbation to the LDA ground state (henceforth denoted $G_0W_0@LDA$). The $GW$ approach has not only become the method of choice for calculating quasiparticle excitations in solids [8, 10] as probed in STS or direct and inverse photoemission, but also includes long-range polarization effects. In a bulk material these encompass the screening of additional charge. At a surface or an interface, however, the abrupt change in dielectric constant gives rise to a net build up of charge, so called image charges. This net polarization acts back on the additional charge and increases in strength with decreasing distance between the additional charge and the interface. These polarization effects are absent from the most common density functionals (such as the local-density or generalized gradient approximation, exact-exchange and hybrid functionals), but enter the $GW$ self-energy ($\Sigma=iGW$) through the screened Coulomb potential $W$. The application of the $GW$ method is therefore necessary to capture these effects [17, 18, 19, 20].

Polarization or image effects are present at any surface or interface, but are most commonly associated with metal surfaces, where the ratio in dielectric constants is largest. In supported ultrathin films, however, a charged excitation (e.g. an electron added to the $2\pi^*$ level of CO on NaCl/Ge(001)) polarizes two interfaces (vacuum/NaCl and NaCl/Ge(001)). The combination of dielectric constants and film thickness therefore controls the strength of the polarization effects from the semiconductor/insulator interface to the insulator surface.

Before we address the polarization effects at the NaCl/Ge interface in detail we briefly describe its atomic structure, which had not been determined previously. The DFT-LDA calculations were performed with the SFHInG code[21]. We employ a plane-wave basis set (40 Ry cutoff) and norm-conserving pseudopotentials. The NaCl/Ge system is modeled in the repeated slab approach with a 6-layer Ge slab at the experimental lattice (110) compared to the free surface (111). Increasing the Ge thickness to 12 layers produces no significant change in the atomic or electronic structure of the NaCl films. In agreement with experimental indications [16], we find that the Ge dimers of the clean Ge(001) surface prevail below the NaCl film, giving rise to a $2 \times 1$ surface lattice [11]. The Ge dimers below the NaCl film remain asymmetric, but with a smaller tilting angle (10°) compared to the free surface (19°). The adhesion of the film is dominated by the electrostatic interaction between the ions in the film and the partial charges developing at the buckled-dimer surface of Ge(001). The relaxation pattern in the bottom layer follows the electrostatic profile of the Ge surface, which attracts the ions next to the dimer, but repels those above the inter-dimer troughs (cf. Fig. 1). The corrugation in the higher layers is induced by the bottom layer and quickly flattens out as the thickness increases.

Next, we use the adsorption of a small molecule to probe the effect of the interface polarization at the film’s surface. For this purpose, we placed a single CO molecule in the $2 \times 1$ surface unit cell (we have no indications for relevant changes at lower coverages). CO physisorsbs perpendicular to the NaCl(001) surface with the C-end down (cf. Fig. 1). The CO axis tilts along (110), i.e. the Ge dimer, by 2.6° for 2 monolayer(ML) NaCl, 0.7° (3 ML), and 0.1° (4 ML), respectively. The adsorption energy of 0.28 eV is the same for the two inequivalent Na sites at the surface and does not depend on the film’s thickness to within 0.01 eV. It also agrees with the value for a thick free-standing NaCl slab. The molecular states give rise to flat bands in the band structure and do not hybridize with NaCl states. In the following, we focus on the molecular gap (given by the $5\sigma-2\pi^*$ splitting) at the $\Gamma$ point.

The $G_0W_0$ calculations for the electron addition and removal spectra were performed with the Gy-st code [23, 24, 25]. For the correlation (exchange) self-energy, a 14 Ry (28 Ry) plane-wave cutoff and a $6 \times 3 \times 1$ k-point sampling was used. State summations included 2500 bands (81 eV above the Fermi level). To correct for artificial polarization effects in the repeated-slab approach, a "finite vacuum" correction was applied [21].

In Fig. 2 we compare the molecular gap of CO computed for LDA and $G_0W_0@LDA$ for films of 2–4 ML thickness. At the level of LDA, we observe a small reduction of the molecular gap with increasing film thickness due to the thickness-dependent structural changes in the surface of the NaCl film. The $G_0W_0$ corrections, however, reverse this trend and introduce a significant increase in the quasiparticle gap from 12.48 eV (for 2 ML) to 12.80 eV (4 ML). The other CO orbitals exhibit analogous thickness-dependent shifts (not shown). These gaps are significantly smaller than for CO on a pure NaCl surface (13.1 eV for 6 ML NaCl, no Ge) or the free molecule (15.1 eV). Similar reductions have been found in $G_0W_0$ calculations for molecules adsorbed on insula-

![FIG. 2: a) $5\sigma$-$2\pi^*$ splitting of CO on NaCl/Ge(001) at the LDA and $G_0W_0@LDA$ levels of theory, and b) $G_0W_0$ corrections to the $5\sigma$ and $2\pi^*$ molecular orbitals as a function of the NaCl film thickness.](image-url)
 tors [27] or semimetals [28] and are a result of surface polarization or charge transfer to the molecule [29].

We will now demonstrate that the reduction of the 5σ-2π* splitting is caused by polarization effects at the two interfaces. Inspection of Fig. (2b) reveals that the thickness-dependent changes are of similar magnitude, but opposite sign. Such a behavior is characteristic for long-range polarization effects [26, 28]. To illustrate this, we split the electron addition/removal into two steps. First, the charged state is created on a free molecule. Secondly, we consider the electronic polarization of the NaCl/Ge substrate. If the lifetime of the charged state on the molecule is long compared to the electronic relaxation time of the substrate, we may treat the rearrangement of charge with classical electrostatics. The polarization lowers the energy of the charged state $E_{N\pm 1}$ by $\frac{1}{2}q\Delta V$, where $\Delta V$ is the polarization-induced change in the electrostatic potential and $q$ the charge. The energy of the hole state ($E_N - E_{N-1}$) thereby increases whereas that of the electron state ($E_{N+1} - E_N$) becomes smaller.

To substantiate this picture, we estimate $\Delta V$ outside the supported NaCl films. For this, the Ge substrate, the NaCl film, and the vacuum region are replaced by homogeneous dielectric media ($\varepsilon=14$, 2.8, and 1, respectively) with abrupt interfaces. The thickness of the NaCl region is taken to be 2.8 Å/layer. The effective polarization is then computed using the image-charge method. For $\Delta V$, we then take the value of the image potential at the position of the CO molecule outside the surface. Since this position is somewhat ambiguous due to the spatial extent of the CO orbitals we determine it by requiring that the reduction of the 5σ-2π* splitting of 2 eV at the bare NaCl surface is reproduced by the model. This yields a value of 1.5 Å, and the model then gives 5σ-2π* splittings of 12.6, 12.7, and 12.8 eV for the 2, 3, and 4 ML films, respectively, in good agreement with the values from our $G_0W_0@LDA$ calculations.

On the experimental side, a gap reduction as a function of film thickness has been reported for STS experiments on pentacene molecules adsorbed on NaCl/Cu(111) [3]. In STS, the molecular states give rise to tunneling resonances and the observed tunneling gap amounts to 3.3, 4.1, and 4.4 eV for NaCl films of 1, 2, and 3 ML in thickness (the gap of pentacene in the gas phase is 5.3 eV). The overall trend as well as the magnitude agree very well with our $G_0W_0@LDA$ calculations for CO/NaCl/Ge considering that the ratio of dielectric constants is much larger in the NaCl/Cu case. A tunneling resonance gap will be hard to observe experimentally for CO/NaCl/Ge, however, because the highest occupied molecular orbital (5σ) is located $\sim$8.5 eV below the Ge valence band maximum (vbm) in $G_0W_0@LDA$ and thus even below the NaCl valence band (4-7 eV below the Ge vbm, cf. Fig. 3).

Our results highlight that surface and interface polarization effects are important for adsorbed atoms and molecules. Supported ultrathin films thereby offer unprecedented opportunities for controlling these effects by tailoring both the film’s thickness and dielectric constant to the desired properties. These additional parameters may expedite the design of devices in molecular electronics or quantum transport, where the distance of a molecular state to the Fermi level is an important quantity.

We will now demonstrate that the same polarization effects also affect excitations inside the ultrathin film and discuss the implications of this for the characterization of insulator surfaces. Fig. 3 shows the density of states (DOS) projected onto the NaCl film for films with 2–4 ML for LDA with and without the $G_0W_0$ corrections. The most important change when comparing the LDA and $G_0W_0$ DOS is the shift of the NaCl bands [30] relative to the Ge states. This is not too surprising, as the $G_0W_0$ corrections to the bulk band gaps are much larger for NaCl (3.3 eV) than for Ge (0.7 eV). Including the $G_0W_0$ corrections, the top of the film’s valence band lies $\sim$4.2 eV below that of the Ge substrate in excellent agreement with ultraviolet photoelectron spectroscopy [31]. More remarkable, however, is the change in the shape of the NaCl-derived features when going from LDA to $G_0W_0@LDA$ and from 2 to 4 layers. This indicates that the $G_0W_0$ shifts for the NaCl states are not uniform and that corrections derived from bulk calculations are not easily transferable to thin films. Excited states are instead subject to additional thickness-dependent and substrate-specific changes that are neither encompassed by a ground-state perspective nor easily derivable from bulk $G_0W_0$ calculations for the separate fragments alone. These thickness-dependent variations should be observable in high-resolution spectroscopic experiments.

In LDA the DOS of bulk NaCl (not shown) is already attained at a thickness of only three layers. A similar behavior has previously been observed for ultrathin silica, hafnia and alumina films [1]. However, this is no longer the case when charged excitations (e.g. photoemission or tunneling) are treated appropriately. The $G_0W_0@LDA$
calculations demonstrate clearly that the DOS of ultrathin films differs from that of bulk NaCl (which is identical in shape to the LDA DOS for a 4 layer film). This implies that caution has to be applied when interpreting spectroscopic results. Spectra of ultrathin films are not representative of bulk samples.

The non-uniform $G_0W_0$ shifts are also a result of the interface polarization effects. The position-dependence of the self-energy ($\Sigma = iG_W$) can be understood with the same dielectric slab model introduced above. Solving the image-charge model yields the induced potential $\Delta V(z)$ as a function of the position in the film $z$. From this, we build the following model self-energy for the occupied NaCl states

$$\Sigma(z) = \Delta\Sigma - \frac{1}{2}\Delta V(z). \quad (1)$$

The constant $\Delta\Sigma$ encompasses all $GW$ self-energy effects that are not due to image effects. Its actual value is not important for analyzing the position dependence of the self-energy. Based on our $G_0W_0$ calculations, we estimate it to be $-1.05 \text{ eV}$ for Cl 3$s$ and $-1.3 \text{ eV}$ for Cl 3$p$. In order to assess the local variation of the self-energy in the full $G_0W_0$ calculation, the valence bands are projected onto atomic orbitals. For states that are predominantly composed of orbitals from a single atomic layer (>80%) we observe a linear dependence between the $G_0W_0$ correction and the projection weight. By extrapolating to 100%, the orbital-dependent quasiparticle corrections shown in Fig.4 are obtained. The variation of the quasiparticle shifts throughout the NaCl film ($G_0W_0$ DOS in Fig.3) is reproduced well by a projection-weighted sum of these local-orbital contributions. In Fig.3 we compare the extrapolated shifts to our model self-energy. The good agreement illustrates that the interface polarization effects in the Ge/NaCl system are indeed the cause for the position dependece in the $G_0W_0$ self-energy.

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Since a meaningful analysis of the NaCl's conduction bands is aggravated by their very strong hybridisation with Ge states, we will focus on the valence states only.

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