Simulation of spin-polarized scanning tunneling spectroscopy on complex magnetic surfaces: Case of a Cr monolayer on Ag(111)

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We propose a computationally efficient atom-superposition-based method for simulating spin-polarized scanning tunneling spectroscopy (SP-STS) on complex magnetic surfaces based on the sample and tip electronic structures obtained from first principles. We go beyond the commonly used local density of states (LDOS) approximation for the differential conductance \( dI/dV \). The capabilities of our approach are illustrated for a Cr monolayer on a Ag(111) surface in a noncollinear magnetic state. We find evidence that the simulated tunneling spectra and magnetic asymmetries are sensitive to the tip electronic structure, and we analyze the contributing terms. Related to SP-STS experiments, we show a way to simulate two-dimensional differential conductance maps and qualitatively correct effective spin-polarization maps on a constant current contour above a magnetic surface.

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I. INTRODUCTION

The scanning tunneling microscope (STM) and its spectroscopic mode (STS) proved to be extremely useful for studying local physical and chemical phenomena on surfaces since the invention of the STM 30 years ago.\(^{1,2}\) The progress of experimental techniques in the last two decades was remarkable, thus more sophisticated theoretical models and simulation tools are needed to explain all relevant details of electron tunneling transport measurements.\(^{3,4}\) STS theory and applications are recently focused on extracting surface local electronic properties from experimental differential conductance \((dI/dV)\) data.\(^{5–9}\) The role of the tip electronic structure has been identified to be crucial on the \(dI/dV\) tunneling spectra, see, e.g., Refs. 7 and 10, and a theoretical method has been proposed to separate the tip and sample contributions to STS.\(^{11}\)

An emerging research field in surface science is the investigation of magnetism at the nanoscale and atomic scale with the aim of achieving ultrahigh information density for data storage purposes.\(^{12,13}\) Spin-polarized scanning tunneling microscopy (SP-STM)\(^{14}\) is admittedly an important tool for studying magnetism on surfaces. Recent experimental advances using this technique allow the investigation of complex magnetic structures (frustrated antiferromagnets, spin spirals, skyrmion lattices, etc.).\(^{15–18}\) Spin-polarized scanning tunneling spectroscopy (SP-STS) has recently been used to find inversion of spin polarization above magnetic adatoms\(^{19–21}\) and the effect has been explained theoretically.\(^{22}\) Furthermore, SP-STS is useful to study atomic magnetism,\(^{23}\) many-body effects on substrate-supported adatoms,\(^{24}\) or magnetic interactions between adatoms\(^{25}\) as well. The effect of differently magnetized surface regions on SP-STS has also been reported,\(^{26,27}\) and the role of tip effects on SP-STS has also been highlighted.

Our work is concerned with the presentation of an efficient simulation method for SP-STS based on first-principles electronic structure data. We extend our atom-superposition-based method\(^{28,31}\) in the spin-polarized Tersoff-Hamann framework\(^{32}\) for simulating SP-STS by including the bias-dependent background and tip-derivative terms into the calculated differential conductance following Passoni \textit{et al.}\(^{7}\). The method is computationally cheap and it can be applied using results of any \textit{ab initio} electronic structure code. The main advance of our tunneling model is the inclusion of the tip electronic structure, which is neglected in Refs. 32 and 33, and it is only taken into account in a model way in Ref. 7. Our method, based on first-principles calculation of the tip electronic structure, enables to study tip effects on the SP-STS spectra. Taking a prototype frustrated hexagonal antiferromagnetic system, a Cr monolayer on Ag(111) in a noncollinear magnetic 120° Néel state, we simulate differential conductance tunneling spectra and magnetic asymmetries to illustrate the applicability of our method, and we analyze the contributing terms. Note that a three-dimensional (3D) approach to STS has been presented recently that is applicable to nonmagnetic systems only, and it takes into account the symmetry of the tip states but not the electronic structure of the tip apex.\(^{34}\) Our model is also a 3D approach in the sense that we sum up contributions from individual transitions between the tip apex atom and each of the surface atoms assuming the one-dimensional (1D) Wentzel-Kramers-Brillouin (WKB) approximation for electron tunneling processes in all these transitions, thus we call it a 3D WKB approach.

The paper is organized as follows. The atom-superposition theoretical model of SP-STS is presented in Sec. II. As an application, we investigate the surface of one monolayer (ML) Cr on Ag(111) in Sec. III. We simulate differential conductance...
tunneling spectra and magnetic asymmetries with two tip models, and we analyze the contributing terms to $dI/dV$. Moreover, we show simulation results of bias dependent two-dimensional (2D) differential conductance and qualitatively correct effective spin-polarization maps following a constant current contour above the surface, corresponding to a standard experimental setup. Our conclusions are found in Sec. IV. Finally, in Appendix, we report the 1D WKB theory of STS and give alternative expressions for the $dI/dV$.

II. THEORETICAL MODEL OF ATOM-SUPERPOSITION SP-STS

The 1D WKB theory for nonmagnetic STS is a well established approach,5,35 see Appendix. Here, we extend it to spin-polarized systems and adapt it to an atom superposition framework, which enables a computationally inexpensive calculation of tunneling properties based on first-principles electronic structure data.

In magnetic STM junctions, the total tunneling current can be decomposed into non-spin-polarized (TOPO) and spin-polarized (MAGN) parts,32,33,36,37

$$I_{\text{TOTAL}} = I_{\text{TOPO}} + I_{\text{MAGN}}.$$  \hspace{1cm} (1)

Following the spin-polarized Tersoff-Hamann model32 and its adaptation to the atom superposition framework,31,33 the magnetic contribution to the simple expression of the differential conductance at a given energy is proportional to the scalar product of the tip and sample magnetic density of states (DOS) vectors, $m_{\alpha}(E)$ and $m_{\alpha}(E)$, respectively,

$$\frac{dI_{\text{MAGN}}}{dU}(E) \propto m_{\alpha}(E)m_{\alpha}(E).$$  \hspace{1cm} (2)

Thus the spin-polarized parts of $dI/dV$ can similarly be calculated within the 1D WKB approximation as reported in Appendix, just replacing $n_{\alpha}(E)n_{\alpha}(E)$ by $m_{\alpha}(E)m_{\alpha}(E)$.

We formulate the tunneling current, the differential conductance, and their TOPO and MAGN parts within the atom superposition framework following Ref. 31. Here, we assume that electrons tunnel through one tip apex atom, and we sum up contributions from individual transitions between this apex atom and each of the surface atoms assuming the 1D WKB approximation for electron tunneling processes in all these transitions. The tunneling current at the tip position $R_{\text{TIP}}(x,y,z)$ and at bias voltage $V$ is given by

$$I_{\text{TOTAL}}(x,y,z,V) = I_{\text{TOPO}}(x,y,z,V) + I_{\text{MAGN}}(x,y,z,V),$$  \hspace{1cm} (3)

where the TOPO and MAGN terms are formally given as

$$I_{\text{TOPO}}(x,y,z,V) = \int_0^V \frac{dI_{\text{TOPO}}}{dU}(x,y,z,U,V)dU,$$  \hspace{1cm} (4)

$$I_{\text{MAGN}}(x,y,z,V) = \int_0^V \frac{dI_{\text{MAGN}}}{dU}(x,y,z,U,V)dU.$$  \hspace{1cm} (5)

The integrands are the so-called virtual differential conductances,

$$\frac{dI_{\text{TOPO}}}{dU}(x,y,z,U,V) = \varepsilon^2 \frac{e^2}{h} \sum_{\alpha} e^{-2\epsilon(U,V)d_{\alpha}(x,y,z)}$$
$$\times n_{\alpha}(E_F^T + eU - eV)n_{\alpha}^*(E_F^S + eU),$$  \hspace{1cm} (6)

$$\frac{dI_{\text{MAGN}}}{dU}(x,y,z,U,V) = \varepsilon^2 \frac{e^2}{h} \sum_{\alpha} e^{-2\epsilon(U,V)d_{\alpha}(x,y,z)}$$
$$\times m_{\alpha}(E_F^T + eU - eV)m_{\alpha}^*(E_F^S + eU).$$  \hspace{1cm} (7)

Here, $\varepsilon$ is the elementary charge, $h$ the Planck constant, and $E_F^T$ and $E_F^S$ the Fermi energies of the tip and the sample surface, respectively. $\varepsilon^2 e^2/h$ ensures that the $dI/dU$ is correctly measured in the units of $A/V$. $\epsilon$ has been chosen to 1 eV, but its actual value has to be determined comparing simulation results to experiments. The sum over $\alpha$ corresponds to the atomic superposition and has to be carried out, in principle, over all surface atoms. Convergence tests, however, showed that including a relatively small number of atoms in the sum provides converged $dI/dU$ values.29 The tip and sample electronic structures are included into this model via projected DOS (PDOS) onto the atoms, i.e., $n_{\alpha}(E)$ and $n_{\alpha}^*(E)$ denote projected charge DOS onto the apex and the $\alpha$th surface atom, respectively, while $m_{\alpha}(E)$ and $m_{\alpha}^*(E)$ are projected magnetization DOS vectors onto the corresponding atomic spheres. They can be obtained from collinear or noncollinear electronic structure calculations.31 In the present work, we determine the noncollinear PDOS for the sample surface and we use a collinear PDOS for a model CrFe tip.22

The transmission probability for electrons tunneling between states of atom $\alpha$ on the surface and the tip apex is of the simple form,

$$T(E_F^S + eU,V,d_{\alpha}(x,y,z)) = e^{-2\epsilon(U,V)d_{\alpha}(x,y,z)}.$$  \hspace{1cm} (8)

This corresponds to a spherical exponential decay of the electron wave functions. Here, $d_{\alpha}(x,y,z) = |R_{\text{TIP}}(x,y,z) - R_{\alpha}|$ is the distance between the tip apex and the surface atom labeled by $\alpha$ with position vector $R_{\alpha}$. Assuming an effective rectangular potential barrier between the tip apex and each surface atom, the vacuum decay $\kappa$ can be written as

$$\kappa(U,V) = \frac{1}{h} \sqrt{2m\left(\phi_S + \phi_T + eV - eU\right)},$$  \hspace{1cm} (9)

where the electron’s mass is $m$, $\hbar$ is the reduced Planck constant, and $\phi_S$ and $\phi_T$ are the average electron work function of the sample surface and the local electron work function of the tip apex, respectively. The method of determining the electron work functions is reported in Ref. 31. $\kappa$ is treated within the independent orbital approximation,33,38,39 which means that the same spherical decay is used for all type of orbitals. The interpretation of our simulation results with quantitative reliability compared to experiments has to be taken with care due to this approximation. However, extension of our model to take into account orbital dependent vacuum decay following Chen’s work40 is planned in the future, which is relevant for a more advanced description of tunneling between directional orbitals.
Moreover, in our model, the electron charge and magnetization local density of states above the sample surface in the vacuum, \( n_{\text{LDOS}} \) and \( m_{\text{LDOS}} \), respectively, are approximated by the following expressions:

\[
n_{\text{LDOS}}(x,y,z,E_F + eU) = \sum_\alpha e^{-2\kappa(U)d_\alpha(x,y,z)} n_\alpha^S(E_F + eU),
\]

\[
m_{\text{LDOS}}(x,y,z,E_F + eU) = \sum_\alpha e^{-2\kappa(U)d_\alpha(x,y,z)} m_\alpha^S(E_F + eU),
\]

with

\[
\kappa(U) = \frac{1}{\hbar} \sqrt{2m(\phi_S - eU)}.
\]

Note that the exact LDOS can be obtained by explicitly calculating the decay of the electron states into the vacuum taking their orbital symmetry into account as well, not via such a simple 3D WKB model. Our approach, however, has computational advantages as discussed in Ref. 31.

Similarly to the tunneling current, the physical differential conductance can be decomposed into TOPO and MAGN parts and it can be written at the tip position \( R_{\text{tip}}(x,y,z) \) and at bias voltage \( V \) as

\[
dI_{\text{TOTAL}}(x,y,z,V) = \frac{dI_{\text{TOPO}}}{dV}(x,y,z,V) + \frac{dI_{\text{MAGN}}}{dV}(x,y,z,V),
\]

where the contributions are given as [see Eq. (A10) in Appendix]

\[
dI_{\text{TOPO}}(x,y,z,V) = \frac{dI_{\text{TOPO}}}{dU}(x,y,z,V,V) \]

\[
+ B_{\text{TOPO}}(x,y,z,V) + D_T^{\text{TOPO}}(x,y,z,V),
\]

\[
dI_{\text{MAGN}}(x,y,z,V) = \frac{dI_{\text{MAGN}}}{dU}(x,y,z,V) \]

\[
+ B_{\text{MAGN}}(x,y,z,V) + D_T^{\text{MAGN}}(x,y,z,V).\]

Thus we formulated all components of the differential conductance in spin-polarized tunnel junctions within the atom superposition framework using first-principles electronic structure of the sample and the tip. Note that all \( dI/dV \) expressions in Eq. (A22) in Appendix can similarly be calculated within our 3D WKB approach.

\[
I(x,y,z,V) \text{ and } dI/dV(x,y,z,V) \text{ can be calculated at (x,y,z) grid points of a three-dimensional (3D) fine grid in a finite box above the surface. The recipe for simulating SP-STM images based on the 3D current map is given in Ref. 31. Here, we focus on the simulation of } dI/dV \text{ spectra. From the 3D differential conductance map, data can be extracted to be directly comparable to experiments. For example, a single-point spectrum corresponds to a fixed (x_0, y_0, z_0) tip position, and two-dimensional (2D) spectra can also be obtained, where the image resolution is determined.}
\]
by the density of \((x,y)\) grid points. There are usually two ways to define a 2D differential conductance map.\(^5\) The first method fixes the tip height at \(z = Z_{\text{stab}} = \text{const}\) and scans the surface, \(dI/dV(x,y,Z_{\text{stab}},V)\). The second option measures \(dI/dV\) on a constant current contour, \(I_{\text{TOTAL}} = I_{\text{stab}} = \text{const}\), which is the widely used method in experiments. Simulation of this can be done in two steps: first, we calculate the 3D current map with the given bias voltage \(V_{\text{stab}}\) and at the second step, we determine the height profile of a constant current contour \(z(x,y,V_{\text{stab}},I_{\text{TOTAL}} = I_{\text{stab}})\), using logarithmic interpolation.\(^3\) \(V_{\text{stab}}\) and \(I_{\text{stab}}\) are the tunneling parameters, and they stabilize the tip position at the height of \(z(x,y,V_{\text{stab}},I_{\text{TOTAL}} = I_{\text{stab}})\) above the \((x,y)\) sample surface point. The 2D differential conductance map on the constant current contour is then given by \(dI/dV(x,y,z(x,y,V_{\text{stab}},I_{\text{TOTAL}} = I_{\text{stab}}),V)\), where the \(V\) dependence is obtained by sweeping the bias voltage range using a lock-in technique in experiments.\(^5\) Recently, experimental efforts have been made to extract the TOPO component of the tunneling current,\(^4\) and measure spectroscopic data on such constant current contours, i.e., at \(I_{\text{TOPO}} = \text{const}\).\(^2\) According to Ref. 15, a constant tunneling transmission enables an easier interpretation of measured 2D spectroscopic data. We believe that a constant TOPO current contour is closer to this constant tunneling transmission criterion than a constant total current contour due to the appearance of spin dependent effects in the latter one. On the other hand, the calculation of any current contour is simple within our 3D WKB approach.\(^3\)

From this, the magnetic asymmetry can similarly be calculated on appropriate constant current contours as described in the previous paragraph. Using Eq. (13), and the fact that the magnetic contribution for the AP tip magnetization direction \(dI_{\text{MAGN}}^{\text{AP}}/dV\) equals \(-dI_{\text{MAGN}}^{\text{BP}}/dV\), since the tip magnetization PDOS vector \(m_{\text{T}}(E)\) changes sign at all energies compared to the P tip magnetization direction, the differential conductances take the following form:

\[
dI_{\text{P}}/dV(x,y,z,V) = dI_{\text{TOPO}}/dV(x,y,z,V) + dI_{\text{MAGN}}^{\text{AP}}/dV(x,y,z,V),
\]

\[
dI_{\text{AP}}/dV(x,y,z,V) = dI_{\text{TOPO}}/dV(x,y,z,V) - dI_{\text{MAGN}}^{\text{BP}}/dV(x,y,z,V).
\]

Thus the magnetic asymmetry can be expressed as the fraction of the MAGN and TOPO differential conductances from Eqs. (14) and (15) as

\[
A(x,y,z) = \frac{dI_{\text{P}}/dV(x,y,z,V) - dI_{\text{AP}}/dV(x,y,z,V)}{dI_{\text{P}}/dV(x,y,z,V) + dI_{\text{AP}}/dV(x,y,z,V)}.
\]

This is an important quantity since it is related to the vacuum spin polarization of the sample in a simple way:

\[
A_{\text{diff}}/dU(x,y,z,V) = \frac{dI_{\text{P}}/dU(x,y,z,V) - dI_{\text{AP}}/dU(x,y,z,V)}{dI_{\text{P}}/dU(x,y,z,V) + dI_{\text{AP}}/dU(x,y,z,V)}.
\]

This is the correct magnetic asymmetry expression based on the physical differential conductances that can be obtained from experiments. However, a magnetic asymmetry can similarly be defined taking the virtual differential conductances from Eqs. (6) and (7):

\[
A_{\text{diff}}/dU(x,y,z,V) = \frac{P_T(E_F^+P_S(x,y,z,E_F^+ + eV) = \text{ESP}(x,y,z,V).}
\]

This is an important quantity since it is related to the vacuum spin polarization of the sample in a simple way:

\[
A_{\text{diff}}/dU(x,y,z,V) = \frac{dI_{\text{MAGN}}^{\text{AP}}/dU(x,y,z,V) - dI_{\text{MAGN}}^{\text{BP}}/dU(x,y,z,V)}{dI_{\text{MAGN}}^{\text{AP}}/dU(x,y,z,V) + dI_{\text{MAGN}}^{\text{BP}}/dU(x,y,z,V)}.
\]

By simulating differential conductance spectra above a magnetic surface with parallel (P) and antiparallel (AP) tip magnetization directions with respect to a predefined direction (usually the magnetization direction of a chosen surface atom is taken), the so-called magnetic asymmetry can be defined.\(^2\) In our case, this quantity can be calculated at all considered positions of the tip apex atom, i.e., at all \((x,y,z)\) grid points within our finite box above the surface,

\[
A(x,y,z,V) = \frac{dI_{\text{P}}/dV(x,y,z,V) - dI_{\text{AP}}/dV(x,y,z,V)}{dI_{\text{P}}/dV(x,y,z,V) + dI_{\text{AP}}/dV(x,y,z,V)}.
\]

This is an important quantity since it is related to the vacuum spin polarization of the sample in a simple way:

\[
A_{\text{diff}}/dU(x,y,z,V) = \frac{dI_{\text{MAGN}}^{\text{AP}}/dU(x,y,z,V) - dI_{\text{MAGN}}^{\text{BP}}/dU(x,y,z,V)}{dI_{\text{MAGN}}^{\text{AP}}/dU(x,y,z,V) + dI_{\text{MAGN}}^{\text{BP}}/dU(x,y,z,V)}.
\]
As $dI/dU(V,V)$ is one component of $dI/dV(V)$ according to Eq. (16), we will compare them and also the magnetic asymmetry expressions in Eqs. (27)–(31), in order to estimate the error one makes when neglecting the background and tip-related components of $dI/dV(V)$ for a given combination of a complex magnetic surface and a magnetic tip. On the other hand, we will calculate qualitatively correct bias-dependent 2D effective spin-polarization maps following a constant current contour.

It has to be noted that the presented method can also be applied to study nonmagnetic systems, where all magnetic contributions equal zero and the corresponding topographic STS spectra can be simulated. Of course, in this case, the magnetic asymmetry is zero.

III. RESULTS AND DISCUSSION

In order to demonstrate the reliability and capabilities of our model for simulating SP-STS on complex magnetic surfaces, we consider a sample surface with noncollinear magnetic order. One ML Cr on Ag(111) is a prototype of frustrated hexagonal antiferromagnets. Due to the geometrical frustration of the antiferromagnetic exchange interactions between Cr spin moments, its magnetic ground state is a noncollinear 120° Néel state. In the presence of spin-orbit coupling, two types of Néel states with opposite chiralities can form, and one of them is energetically favored.

We performed geometry relaxation and electronic structure calculations based on the density functional theory (DFT) within the generalized gradient approximation (GGA) implemented in the Vienna ab initio simulation package (VASP). A plane wave basis set for electronic wave function expansion together with the projector augmented wave (PAW) method has been applied, while the exchange-correlation functional is parametrized according to Perdew and Wang (PW91). For calculating the fully noncollinear electronic structure, we used the VASP code as well, with spin-orbit coupling considered.

We model the Cr/Ag(111) system by a slab of a five-layer Ag substrate and one monolayer Cr film on each side, where the surface Cr layers and the first Ag layers underneath have been fully relaxed. A separating vacuum region of 14.6 Å width in the surface normal (z) direction has been set up between neighboring supercell slabs. The average electron work function above the surface is $\phi_S = 4.47$ eV. We used an $11 \times 11 \times 1$ Monkhorst-Pack (MP)50 k-point grid for calculating the projected electron DOS (PDOS) onto the surface Cr atoms in our ($\sqrt{3} \times \sqrt{3}$) magnetic surface unit cell.

The energy-dependent charge and magnetization PDOS, $n_S(E)$ and $m_S(E)$, respectively, are shown in Fig. 1. We obtained these quantities from noncollinear calculations. The spin quantization axis of each surface Cr atom is chosen to be parallel to their magnetic moment direction, and $m_S(E)$ is the projection of the magnetization PDOS vector $m_S(E)$ to this direction. Except the spin quantization axes of the three different Cr atoms in the magnetic surface unit cell, their electronic structure is the same. We can interpret the results in terms of the commonly used spin-up (↑, majority) and spin-down (↓, minority) channels with respect to the atomic spin quantization axis, where $n(E) = n^\uparrow(E) + n^\downarrow(E)$, and $m(E) = n^\uparrow(E) - n^\downarrow(E)$. It is seen that the majority spin PDOS dominates over the minority spin PDOS below $E^\uparrow_S + 0.54$ eV, while $m_S(E) < 0$ above $E^\uparrow_S + 0.54$ eV. This implies a spin polarization reversal at this particular energy.

In our model, the vacuum local density of states (LDOS) is obtained by the superposition of spherically decaying electron states according to the independent orbital approximation. Above a complex magnetic surface, the spin-up and spin-down notations are meaningless since there is no global spin quantization axis. Instead, we can consider the charge and magnetization (vector) character of the LDOS obtained from the PDOS, as defined in Eqs. (10) and (11) for $n_{LDOS}$ and $m_{LDOS}$, respectively. Above a surface Cr atom with lateral position $(x_0, y_0)$, both vacuum LDOS behave the same way as the corresponding PDOS, thus the spin polarization vector in vacuum $P_{LDOS}(x_0, y_0, z, E) = n_{LDOS}(x_0, y_0, z, E)/n_{LDOS}(x_0, y_0, z, E)$ equals the one obtained from the PDOS, i.e., $P_{LDOS}(x_0, y_0, z, E) = m_{LDOS}(x_0, y_0, z, E)$.

FIG. 1. (Color online) Projected charge and magnetization DOS of the surface Cr atom of the sample Cr/Ag(111), the tip apex Cr atom of the Cr/Fe(001) tip, and the flat tip.

$P_{LDOS}(x_0, y_0, z, E)$ will vary due to the different atomic spin quantization axes for all three Cr atoms in the magnetic surface unit cell and the considered vacuum decays. $n_{LDOS}$ will, however, remain qualitatively unchanged. The lateral variation of $m_{LDOS}$ will result in a position dependent vacuum spin polarization vector of the sample surface. This quantity multiplied by the tip spin polarization vector results in the effective spin polarization, defined in Eq. (29), which will be simulated later.

Dependence of the tunneling spectra on the tip electronic structure can be studied by considering different tip models. In this work, we compare spectra and magnetic asymmetries measured by a magnetic CrFe tip and an electronically flat magnetic tip. The electronic structure data of the CrFe tip apex was taken from Ref. 22, where the tip was modeled as a single Cr apex atom on the Fe(001) surface. Ferriani et al. furthermore reported that an antiferromagnetic coupling of the Cr adatom to the Fe(001) surface is energetically preferred, and the vacuum spin polarization is fairly constant at around $+0.8$ in the energy range $[E^F_L - 1 \text{ eV}, E^F_L + 1 \text{ eV}]$.22
The local electron work function above the tip apex is assumed to be \( \phi_T = 4.5 \) eV, that has been used to obtain the energy-dependent vacuum decay in Eq. (9). The charge and absolute magnetization PDOS of the Cr apex atom, \( n_T(E) \) and \( m_T(E) \), respectively, are shown in Fig. 1. We obtain qualitative correspondence to the PDOS of the sample Cr atom. However, due to the different surface orientation and the different local environment of the Cr/Ag(111) surface and CrFe(001) tip apex Cr atoms, the sample and tip Cr PDOS are quantitatively different. Concerning magnetic properties, we find a spin polarization reversal at \( E_T^F + 0.7 \) eV. On the other hand, there is no energy-dependent vacuum spin polarization reversal observed in Ref. 22. Ferriani et al. analyzed this effect in detail for an Fe atom on top of the Fe(001) surface, and they found a competition between majority and minority spin states with different decays into the vacuum. Such an orbital-dependent vacuum decay is not included in our model at the moment, but work is in progress to consider such effects.

The electronically flat magnetic tip has been modeled based on the electronic structure of the Cr apex (PDOS) of the CrFe tip. The charge and absolute magnetization PDOS, \( n_T(E) \) and \( m_T(E) \), respectively, have been averaged in the \( [E_T^F - 2 \text{ eV}, E_T^F + 2 \text{ eV}] \) range. We obtained \( n_T = 1.33/\text{eV} \) and \( m_T = 1.06/\text{eV} \), also shown in Fig. 1. Thus the spin polarization is \( P_T = m_T/n_T = +0.8 \). In this case, the tip-derivative term of the differential conductance \( D_T(V) \) is zero, since \( \delta m_T(E)/\delta E = \delta m_T(E)/\delta E = 0 \). The vacuum decay can be modeled using Eq. (9), where \( \kappa(U,V) \) has an explicit \( V \)-dependence, and we assume that \( \theta_T = \theta_s \). Alternatively, a simpler model for \( \kappa(U,V) \) can be considered without \( V \)-dependence as in Eq. (12). In this case, the background term of the differential conductance \( B(V) \) is zero, since the tunneling transmission does not depend on the bias voltage, and the physical differential conductance equals the virtual differential conductance, i.e., \( dI/dV(V) = dI/dU(U,V) \). On the other hand, by assuming a \( V \)-dependent vacuum decay \( \kappa(U,V) \), \( B(V) \) is not zero and it contributes to the total differential conductance, i.e., \( dI/dV(V) = dI/dU(U,V) + B(V) \).

Figure 2 shows the bias dependence of the total tunneling current \( I(V) \), calculated using Eq. (3), at the position \( z = 3.5 \) Å above the surface Cr atom probed with the CrFe tip having parallel (P) magnetization direction compared to the underlying surface Cr atom. Positive current means tunneling from the tip to the sample surface, whereas the current is negative in the opposite direction. We find that the absolute value of the current is higher in the negative bias range compared to the positive range. This is due to the surface and tip electronic structures. The sample occupied PDOS combined with the tip unoccupied PDOS is greater than the sample unoccupied PDOS combined with the tip occupied PDOS, see Fig. 1. Performing a numerical differentiation of \( I(V) \) with respect to \( V \), we obtain the differential conductance at this particular tip position. As can be seen this is extremely noisy, and a smoothing procedure should be applied to it before further analysis. Alternatively, the differential conductance can be calculated using Eq. (16), implemented within the atom superposition approach. Figure 2 shows that \( dI/dV \) obtained this way (black curve) is a smooth function that fits precisely to the noisy numerical derivative of the current. There is more discussion about avoiding the numerical differentiation of the tunneling current in determining the \( dI/dV \), e.g., in Ref. 11.

We obtain more information about the \( dI/dV \) by analyzing its components, the virtual differential conductance \( dI/dU(U,V) \), the background term \( B(V) \), and the tip-derivative term \( D_T(V) \). We find that \( dI/dU(U,V) \) differs less than 10% compared to \( dI/dV \) in the bias range \([-0.01 \text{ V}, +0.01 \text{ V}] \), i.e., practically at the common Fermi level of tip and sample. This means that the virtual differential conductance approximation for the \( dI/dV \) (also known as the LDOS approximation) is not sufficient except at zero bias, where they are identical, \( dI/dV(0) = dI/dU(0,0) \). Moreover, one can recognize that most part of the \( dI/dV \) peak structure is already included in the \( dI/dU(U,V) \) term, which is qualitatively similar to the charge PDOS of the surface Cr atom of the sample, \( n_S(E) \), see Fig. 1. Apart from this, the peak structure of \( D_T(V) \), calculated via Eqs. (23) and (24), clearly shows up in the \( dI/dV \), particularly pronounced at high bias voltages. The reason is the rapidly changing tip electronic structure in these energy regions, see Fig. 1. The features from \( dI/dU(U,V) \) and \( D_T(V) \) are transferred to the physical differential conductance, since \( B(V) \), calculated via Eqs. (21) and (22), is smooth compared to the other two components in the whole bias range. Moreover, we find that \( B(V) \) is a monotonous function of the bias voltage, and it is nearly proportional to \( I(V) \) as has been reported earlier for different levels of STS theories. The proportionality function \( B(V)/I(V) \) is plotted in the inset of Fig. 2. It can be seen that its sign is in agreement with Ref. 7 and it has a nontrivial bias dependence. This is essentially due to the extra \( 1/\kappa(U,V) \) factor in the energy integration of the background term, Eqs. (21) and (22), compared to the tunneling current expression. The \( B(V)/I(V) \) function could, in principle, be calculated at different tip-sample distances \( z \) and could be compared to analytical expressions denoted by \( f(z,V) \) reported in Ref. 7. The comparison is,
we focus on the comparison of SP-STS spectra by reversing the tip magnetization, similarly as before. We find that \( dI_P/dV > dI_{AP}/dV \) below \( V = +0.54 \) V, and \( dI_P/dV < dI_{AP}/dV \) above \( V = +0.54 \) V. In our previous work, we identified the effective spin polarization \( \left[ P_F(E)P_g(E) = n_F(E)n_g(E)/n_T(E)n_0(E) \right] \) responsible for this effect. This is the decisive factor for determining the sign of the magnetic contribution to \( dI/dV \) at energy \( E \) in the improved SP-STS model presented in Sec. II as well. The magnetic part of the physical differential conductance is given in Eq. (15). Since the tip is electronically flat and the vacuum decay does not depend on the bias voltage \( V \) for the dotted curves, \( dI_{MAGN}/dV(V) = dI_{MAGN}/dU(V,V) \). Thus the sign change of \( dI_{MAGN}/dV \) occurs at the sign change of \( dI_{MAGN}/dU(V,V) \), i.e., at the reversal of the sample spin polarization vector at 0.54 eV above the sample Fermi level. see also Fig. 1. For the flat magnetic tip and the assumed bias-dependent vacuum decay (dashed curves), we find that \( dI_P^T/dV > dI_{AP}^T/dV \) below \( V = +0.5 \) V, and \( dI_P^T/dV < dI_{AP}^T/dV \) above \( V = +0.5 \) V, i.e., the sign change of the magnetic component is slightly shifted toward zero bias. The reason is the nonzero background term \( B_{MAGN}(V) \) due to \( \kappa(U,V) \) and \( \kappa(U) \) are assumed using Eqs. (12) and (9), respectively. The vertical dotted line at \( -1.2 \) V shows the bias position of the identified STS peak coming from the electronic structure of the CrFe tip.

However, not straightforward due to two reasons. First, the analytical expressions were reported based on the 1D WKB approximation, whereas our model is a 3D atomic superposition approach based on WKB, which results in an effective transmission coefficient, different from the 1D WKB transmission. Note that a 3D approach to STS with another position approach based on WKB, which results in an integrated quantities \( B_{MAGN}(V) \) and \( D_{MAGN}(V) \) terms. The role of the effective spin polarization is more complicated, since, apart from the \( dI_{MAGN}/dU(V,V) \) term, it appears in the \( dI/dV \) expression through the bias-integrated quantities \( B_{MAGN}(V) \) and \( D_{MAGN}(V) \). For the tip magnetization, \( dI_P^T/dV \) is the same as the black solid curve in Fig. 2, and its contributions are also shown there. In Fig. 3, we observe more changes of the relative height of the \( dI_P^T/dV \) and \( dI_{AP}^T/dV \) spectra measured with the CrFe tip than with the flat tip. These include the sign changes of the magnetic part of the spectra, similarly as before. We find that \( dI_P^T/dV > dI_{AP}^T/dV \) in the bias interval \([-1.04 \, \text{V}, +0.49 \, \text{V}] \), and a reversed relation is obtained in the complementary bias regime. Comparing the spectra to the ones measured with the flat magnetic tip, we see that they are qualitatively closer to the \( \kappa(U,V) \) model used for the flat tip due to the presence of the background terms. Moreover, the individual features coming from the sample and the tip electronic structures can be assigned. In our case, we identify the peak at \(-1.2 \) V, indicated by a vertical dotted line in Fig. 3, coming from the CrFe tip electronic structure since it is missing from the spectra calculated with the flat tip. All other features are related to the sample electronic structure as they appear in the spectra measured with the flat tip.

The relative heights of the differential conductance tunneling spectra \( dI_P^T/dV \) and \( dI_{AP}^T/dV \) can also be determined.

**FIG. 3.** (Color online) Comparison of simulated single point differential conductance tunneling spectra following Eq. (16), probed with the flat magnetic tip and the model CrFe tip, 3.5 Å above a surface Cr atom. Parallel (P) and antiparallel (AP) tip magnetization directions are set relative to the underneath surface Cr atom. For the flat magnetic tip, two different vacuum decays, \( \kappa(U) \) and \( \kappa(U,V) \) are assumed using Eqs. (12) and (9), respectively. The vertical dotted line at \(-1.2 \) V shows the bias position of the identified STS peak coming from the electronic structure of the CrFe tip.
from the magnetic asymmetry, Eq. (25). Let us compare the magnetic asymmetries calculated from the spectra in Fig. 3 using the two magnetic tips. Moreover, for the CrFe tip we compare the asymmetry expressions defined in Eqs. (27)–(31), in order to estimate the error one makes when neglecting the background and tip-related components of $dI/dV(V)$. Figure 4 shows the calculated magnetic asymmetry functions at $z = 3.5$ Å above a surface Cr atom. It can be seen that $A_{\text{Flat, }U}(V)$ and $A_{\text{Flat, }U}(V)$ (dashed curves) behave qualitatively similarly. In addition, $A_{\text{Flat, }U}(V)$ is greater than $A_{\text{Flat, }U}(V)$ in almost the full studied bias range. Moreover, these two magnetic asymmetries are within 5% relative difference in the bias range $[-0.23$ $V$, $+0.31$ $V]$. Considering the CrFe tip, the experimentally measurable magnetic asymmetry $A_{\text{CrFe},dI/dV}(V)$ (black solid curve) is qualitatively different from the two asymmetry functions calculated with the flat tip, e.g., it has a richer structure at positive bias voltages. More importantly, it has an extra sign change occurring at $-1.04$ $V$ apart from $+0.49$ $V$. These correspond to the height changes of $dI^P/dV$ and $dI^AP/dV$ relative to each other in Fig. 3. Let us estimate the error of the magnetic asymmetry when neglecting the background and the tip-derivative terms. According to Eq. (28), $A_{\text{CrFe},dI/dU}(V)$ (curve with symbol “o”) considers the virtual differential conductances only. It is within 10% relative error compared to $A_{\text{CrFe},dI/dV}(V)$ in the bias range $[-0.65$ $V$, $+0.1$ $V]$. However, its sign does not correspond to $A_{\text{CrFe},dI/dV}(V)$ in the bias intervals $[-2$ $V$, $-1.04$ $V]$ and $[+0.49$ $V$, $+0.54$ $V]$. Adding the background term $B(V)$ to $dI/dU(V,V)$ results in an improved differential conductance expression, and $A_{\text{CrFe},dI/dU+B}(V)$ (curve with symbol “*”), defined in Eq. (30), behaves qualitatively similarly to $A_{\text{CrFe},dI/dU}(V)$ above $-0.65$ $V$. However, its sign change is shifted to $+0.45$ from $+0.54$ $V$. Additionally, a sign change in the negative bias range occurs at $-1.62$ $V$. Close to the sample Fermi level, $A_{\text{CrFe},dI/dU+B}(V)$ is within 10% relative error compared to $A_{\text{CrFe},dI/dV}(V)$ in a decreased bias range of $[-0.34$ $V$, $+0.1$ $V]$. Finally, by adding the tip-derivative term $D_T(V)$ to $dI/dU(V,V)$, $A_{\text{CrFe},dI/dU+D_T}(V)$ (curve with symbol “x”), defined in Eq. (31), shows the most closely related shape to $A_{\text{CrFe},dI/dV}(V)$. Furthermore, it is also quantitatively close to the physical magnetic asymmetry as its sign changes occur at $-1.01$ and $+0.5$ $V$, and it is within 10% relative error compared to $A_{\text{CrFe},dI/dV}(V)$ in an increased bias interval $[-0.90$ $V$, $+0.45$ $V]$. Summarizing this paragraph, the contribution of all three terms to the $dI/dV(V)$ according to Eq. (16) is needed to define the physical magnetic asymmetry, that should be comparable to experiments.

Our method presented in Sec. II also enables one to simulate two-dimensional (2D) $dI/dV$ and magnetic asymmetry maps in high spatial resolution above the surface that can be compared to results of SP-STS experiments. Such experiments are routinely performed while the tip follows a constant TOTAL current contour, see, e.g., Ref. 51. Figure 5 illustrates this capability of our method, where we used the flat tip model with tip magnetization direction $M_T$ parallel to the $(1/2, \sqrt{3}/2)$ direction (i.e., the magnetization direction of the surface Cr atom at the bottom left corner of the scan area, see top left part of Fig. 5). Moreover, $\kappa(U,V)$, Eq. (9) has been used for the vacuum decay. By choosing $V_{\text{stab}} = +1$ $V$, we calculate the 3D TOTAL current map in a box above the surface. From this 3D data we extract the current contour of $I_{\text{TOTAL}} = 54$ nA, which is around 3.5 Å above the sample surface and has a corrugation of 4.2 pm. This contour, $\chi(x,y,V_{\text{stab}} = +1$ $V, I_{\text{TOTAL}} = 54$ nA) is plotted in the bottom left part of Fig. 5. The apparent height of the Cr atom with parallel magnetic moment to the tip is lower than those of the other two Cr atoms in the magnetic surface unit cell. This has been explained in a previous work.\textsuperscript{31} The surface scan area and the magnetic unit cell are shown in the top left part of Fig. 5, indicated by a black-bordered rectangle and a yellow (light gray) rhombus, respectively. For calculating the differential conductance-related 2D maps, we vary the vertical position $z$ of the tip apex following the constant current contour shown in the bottom left part of Fig. 5. Thus spin-resolved $dI/dV$ and magnetic asymmetry maps can be simulated at different $V$ bias voltages corresponding to experiments. As an example, $dI/dV(x, y)$ and the effective spin polarization ESP($x, y$), see Eq. (29), are shown in the middle and right columns of Fig. 5, respectively, calculated at bias voltages $V = +0.5$ $V$ (top) and $V = +0.6$ $V$ (bottom). We chose these voltages close to the spin polarization reversal of the sample surface at 0.54 eV above its Fermi level, see Fig. 1, and Ref. 31. Indeed, the reversal of the 2D $dI/dV$ map at $V = +0.6$ $V$ compared to $V = +0.5$ $V$ can clearly be seen. While the SP-STM image at $+1$ $V$ and the $dI/dV$ map at $+0.6$ $V$ show the same type of contrast, the $dI/dV$ signal is inverted for $+0.5$ $V$. Since $P_T = +0.8$ is constant in the full energy range, this effect is due to the surface electronic structure. At $+0.6$ $V$ bias, all surface Cr spin polarization vectors point opposite to their local magnetic moment directions,\textsuperscript{31} and since $P_T = +0.8$ is set with respect to the $(1/2, \sqrt{3}/2)$ direction ($M_T$), the leading term of the magnetic differential conductance, $dI_{\text{MAG}}/dU(V,V)$ is negative above the surface Cr atom with parallel magnetic moment to the tip. Moreover,
FIG. 5. (Color online) (Top left) Surface geometry of 1 ML Cr on Ag(111). The Cr and Ag atoms are denoted by spheres colored by green (medium gray) and purple (dark gray), respectively, while the magnetic moments of individual Cr atoms are indicated by (red) arrows. The \( \sqrt{3} \times \sqrt{3} \) magnetic unit cell is denoted by yellow (light gray) color. The surface Cr positions are denoted by “x.” (Bottom left) Constant current contour about 3.5 Å above the surface with \( I_{\text{TOTAL}}(V_{\text{stab}} = +1 \, \text{V}) = 54 \, \text{nA} \) calculated with the flat magnetic tip using \( \kappa(U, V) \), Eq. (9). The tip magnetization direction \( (M_{\text{TIP}}) \) is indicated by an arrow. (Middle column) Simulated 2D differential conductance maps \( dI/dV(x, y, V = +0.5 \, \text{V}) \) (top middle: min. 15.8, max. 16.5 nA/V), and \( dI/dV(x, y, V = +0.6 \, \text{V}) \) (bottom middle: min. 22.9, max. 23.6 nA/V), while the tip is following the constant current contour at the bottom left of the figure. Minimum (MIN) and maximum (MAX) values are indicated. (Right column) Simulated effective spin polarization (ESP) maps on the same current contour following Eq. (29), ESP(x, y, V = +0.5V) (top right), and ESP(x, y, V = +0.6V) (bottom right). Black contours correspond to zero ESP, and the regions with positive (+) and negative (−) ESP are indicated. The surface magnetic unit cell is drawn by a yellow (light gray) rhombus on each 2D map.

The sign of \( dI_{\text{MAGN}}/dU(V, V) \) changes to positive above the other two Cr atoms in the magnetic unit cell. This results in the minimal total \( dI/dV(x, y) \) above the Cr atom at the bottom left corner of the scan area (22.9 nA/V, magnetic moment parallel to tip), whereas above the other two Cr atoms \( dI/dV \) is maximal (23.6 nA/V, magnetic moment not in line with tip). This happens even though the topographic differential conductance is higher above the Cr atom that is lower-lying on the constant current contour. Similarly, the case of +0.5 V is reversed, since all surface Cr spin-polarization vectors point along their local magnetic moment directions and the maximal total \( dI/dV(x, y) \) is achieved above the Cr atom at the bottom left corner of the scan area (16.5 nA/V, magnetic moment parallel to tip), whereas above the other two Cr atoms \( dI/dV \) is lower (16.0 nA/V, magnetic moment not in line with tip). The minimal \( dI/dV = 15.8 \, \text{nA/V} \) is obtained above the midpoint of the lines connecting two \( dI/dV \) maxima. If we introduce the notation of \( dI^P/dV(x, y) \) for the above calculated differential conductances with P parallel to the indicated \( M_{\text{TIP}} \) in Fig. 5, then the antiparallel tip orientation is denoted by \( A \), and \( dI^{AP}/dV(x, y) \) can similarly be calculated. For the very same reason as discussed, a reversed tip magnetization direction would result in a reversed \( dI^{AP}/dV \) map concerning the heights above the nonequivalent magnetic Cr atoms. Thus, at +0.6 V, the difference between \( dI^P/dV(x, y) \) and \( dI^{AP}/dV(x, y) \) is minimal and negative above the bottom left Cr atom in the scan area, and maximal and positive above the other two Cr atoms, while the opposite is true at +0.5 V. These explain qualitatively well the simulated ESP(x, y) maps, see the right column of Fig. 5. The ESP(x, y) = 0 contour acts as a border between surface regions with positive and negative ESP at the given bias. Note that the sign of the tip spin polarization has a crucial effect on the ESP(x, y) map. Reversing the sign of \( PT \) compared to the \( M_{\text{TIP}} \) direction would result in a reversed ESP(x, y) map.

We suggest that by applying our method to magnetic surfaces, two-dimensional \( dI^P/dV(x, y) \), \( dI^{AP}/dV(x, y) \), and magnetic asymmetry \( A(x, y) \) maps can be constructed on appropriate current contours at arbitrary V bias, corresponding to SP-STS experiments. Similarly, an ESP(x, y) map can be simulated. We stress again that the ESP cannot simply be obtained from experimental magnetic asymmetry due to the presence of the background and tip-derivative terms. By explicitly considering the tip electronic structure in our SP-STS model based on experimental information, it would help in a more reasonable interpretation of experimentally measured tunneling spectra, magnetic asymmetries, and effective spin polarization.

IV. CONCLUSIONS

We presented an efficient simulation method for spin-polarized scanning tunneling spectroscopy based on first-principles electronic structure data within our atom...
superposition framework\textsuperscript{31} by including the bias-dependent background and tip-derivative terms into the differential conductance formula following Passoni et al.\textsuperscript{7} We showed that our simulated data can be related to standard experimental setups. Taking the tip electronic structure into account, the effect of a richer variety of electronic structure properties can be investigated on the tunneling transport within the indicated approximations (atom superposition, orbital-independent spherical vacuum decay). The method is computationally cheap and it can be applied based on the results of any \textit{ab initio} electronic structure code. Taking a prototype frustrated hexagonal antiferromagnetic system, a Cr monolayer on Ag(111) in a noncollinear magnetic 120° Néel state, we simulated differential conductance tunneling spectra and magnetic asymmetries to illustrate the applicability of our method, and we analyzed the contributing terms. We found that the features of the tunneling spectra are coming from the virtual differential conductance and tip-derivative terms, and the background term is proportional to the tunneling current. We showed evidence that the tunneling spectra and the related magnetic asymmetries are sensitive to the tip electronic structure and to the vacuum decay. We also demonstrated a simulation method for 2D $dI/dV$, magnetic asymmetry, and qualitatively correct effective spin polarization maps above a complex magnetic surface following a constant current contour. Finally, we pointed out that the magnetic asymmetry obtained from experiments can not simply be related to the sample spin polarization due to the presence of the background and tip-derivative terms.

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**APPENDIX: THEORY OF STS WITHIN 1D WKB**

We report the formulation of the tunneling current and the differential conductance in the framework of the one-dimensional (1D) WKB approximation, which has been used in our atom superposition approach in Sec. II. Assuming elastic tunneling, the non-spin-polarized part of the tunneling current at zero temperature is given by\textsuperscript{5,35}

\[
I(V,d) = C \int_{E_F^S}^{E_F^S} T(E,V,d) n_T(E) n_S(E) dE,
\]

where $V$ is the bias voltage, $d$ is the tip-sample distance, $C$ is an appropriate constant, $E_F^S$ is the Fermi energy of the sample surface, $e$ is the elementary charge, $T$ is the tunneling transmission coefficient, while $n_T(E)$ and $n_S(E)$ are the tip and sample densities of states, respectively. Performing a change of variable from $E$ to $U$ using $E = E_F^S + eU$, the tunneling current reads

\[
I(V,d) = Ce \int_0^V T(E_F^S + eU,V,d) n_T(E_F^S + eU) \times n_S(E_F^S + eU) dU. \tag{A2}
\]

The applied bias voltage $V$ in the tunnel junction defines the difference between tip and sample Fermi levels, $E_F^T = E_F^S + eV$. Using this, the energy dependence of $n_T(E)$ can be rewritten related to the tip Fermi level $E_F^T$, and the tunneling current can be reformulated as

\[
I(V,d) = Ce \int_0^V T(E_F^T + eU,V,d) n_T(E_F^T + eU - eV) \times n_S(E_F^S + eU) dU. \tag{A3}
\]

We denote the integrand by the formal quantity,

\[
\frac{dI}{dU}(U,V,d) = CeT(E_F^T + eU,V,d) n_T(E_F^T + eU - eV)
\]

\[
\times n_S(E_F^S + eU), \tag{A4}
\]

called virtual differential conductance. The tunneling current can then be expressed as

\[
I(V,d) = \int_0^V \frac{dI}{dU}(U,V,d) dU. \tag{A5}
\]

The physical differential conductance can be obtained as the derivative of the tunneling current with respect to the bias voltage. This can formally be written as

\[
\frac{dI}{dV}(V,d) = \frac{dI}{dU}(V,d) \bigg|_{V=V}\left(\frac{\partial}{\partial V}\frac{dI}{dU}(U,V,d)\right) dU, \tag{A6}
\]

or using Eq. (A4) as

\[
\frac{dI}{dV}(V,d) = CeT(E_F^T + eU,V,d) n_T(E_F^T + eU - eV)
\]

\[
+ Ce \int_0^V \frac{\partial}{\partial V'} T(E_F^S + eU,V,d',\frac{\partial}{\partial V'} T(E_F^S + eU,V,d') \times n_T(E_F^T + eU - eV') \right|_{V'=V} n_S(E_F^S + eU) dU. \tag{A7}
\]

This is a known formula in the literature.\textsuperscript{5,35} If the tip electronic structure is assumed to be energetically flat, i.e., $n_T(E) = n_T$, which is still a widely used approximation in the recent literature, then the $V$ dependence of $n_T(E_F^T + eU - eV)$ disappears, i.e., $n_T(E_F^T + eU - eV) = n_T$, and the differential conductance becomes

\[
\frac{dI}{dV}(V,d) = CeT(E_F^S + eU,V,d) n_T(E_F^T + eU) n_S(E_F^S + eU)
\]

\[
+ Cen_T \int_0^V \frac{\partial T}{\partial V}(E_F^S + eU,V,d) n_S(E_F^S + eU) dU. \tag{A8}
\]

Here, the second term is the so-called background term, which is a monotonous function of the bias voltage.\textsuperscript{35} Going beyond the assumption of the electronically flat tip by incorporating
the tip electronic structure in the differential conductance expression, the effect of the tip can be studied on the tunneling spectra. The explicit energy dependence of $n_T(E)$ can be calculated from first principles or can be included in a model way. Following Eq. (A7), the differential conductance can be reformulated as

$$\frac{dI}{dV}(V,d) = CeT(E^S_F + eU,V,d)n_T(E^T_F)n_S(E^S_F + eV) + Ce \int_0^V \frac{\partial T}{\partial V}(E^S_F + eU,V,d)n_T(E^T_F + eU - eV)n_S(E^S_F + eU)dU + Ce \int_0^V T(E^S_F + eU,V,d)\frac{\partial n_T}{\partial V}(E^T_F + eU - eV)n_S(E^S_F + eU)dU. \quad (A9)$$

Using that $\partial n_T(E^T_F + eU - eV)/\partial V = -\partial n_T(E^T_F + eU - eV)/\partial U$, the differential conductance at bias voltage $V$ can be written as a sum of three terms:

$$\frac{dI}{dV}(V,d) = \frac{dI}{dU}(0,V,d) + B(V,d) + D_T(V,d) \quad (A10)$$

with

$$\frac{dI}{dU}(0,V,d) = CeT(E^S_F,V,d)n_T(E^T_F - eV)n_S(E^S_F), \quad (A15)$$

$$B(V,d) = Ce \int_0^V \frac{\partial T}{\partial U}(E^S_F + eU,V,d)n_T(E^T_F + eU - eV)n_S(E^S_F + eU)dU, \quad (A12)$$

$$D_T(V,d) = -Ce \int_0^V T(E^S_F + eU,V,d)\frac{\partial n_T}{\partial U}(E^T_F + eU - eV)n_S(E^S_F + eU)dU. \quad (A13)$$

Here, $B(V,d)$ is the background term usually considered in recent STS theories, and $D_T(V,d)$ is a term containing the energy derivative of the tip density of states (DOS), which is rarely taken into account for practical STS calculations and analyses of experimental STS data.

It can be shown that an alternative expression for the differential conductance can be derived using integration by parts,

$$\frac{dI}{dV}(V,d) = \frac{dI}{dU}(0,V,d) + B(V,d) + B_2(V,d) - D_S(V,d) \quad (A14)$$

with

$$\frac{dI}{dU}(0,V,d) = CeT(E^S_F,V,d)n_T(E^T_F - eV)n_S(E^S_F), \quad (A15)$$

$$B_2(V,d) = Ce \int_0^V \frac{\partial T}{\partial U}(E^S_F + eU,V,d)n_T(E^T_F + eU - eV)n_S(E^S_F + eU)dU, \quad (A16)$$

$$D_S(V,d) = -Ce \int_0^V T(E^S_F + eU,V,d)\frac{\partial n_T}{\partial U}(E^T_F + eU - eV)n_S(E^S_F + eU)dU. \quad (A17)$$

This way another background term, $B_2(V,d)$ enters the differential conductance formula, and the energy derivative of the sample DOS appears in the term $D_S(V,d)$. The average of the two $dI/dV$ expressions can also be formed as

$$\frac{dI}{dV}(V,d) = \frac{1}{2} \left[ \frac{dI}{dU}(0,V,d) + \frac{dI}{dU}(V,d) \right] + B(V,d) + \frac{1}{2}B_2(V,d) + \frac{1}{2}[D_T(V,d) - D_S(V,d)], \quad (A18)$$

which gives a third alternative form for the differential conductance within the 1D WKB approximation. On the other hand, by subtracting Eq. (A14) from Eq. (A10), one gets

$$0 = \left[ \frac{dI}{dU}(V,d) - \frac{dI}{dU}(0,V,d) \right] - [B_2(V,d) - D_T(V,d) - D_S(V,d)]. \quad (A19)$$

This is trivial since $B_2(V,d) - D_T(V,d) - D_S(V,d)$ is related to the partial derivative of $dI/dU(U,V,d)$ with respect to $U$:

$$B_2(V,d) - D_T(V,d) - D_S(V,d) = Ce \int_0^V \frac{\partial}{\partial U} \frac{dI}{dU}(U,V,d)dU \equiv \frac{dI}{dU}(V,d) - \frac{dI}{dU}(0,V,d). \quad (A20)$$

From the three equivalent $dI/dV$ formulas in Eqs. (A10), (A14), and (A18), the calculation of Eq. (A10) needs the least mathematical operations, thus, we adopted this formula to our atom superposition approach in Sec. II in order to simulate STS spectra based on electronic structure data calculated from first principles.
Finally, note that using the transmission function in Eq. (8), and the given form of the vacuum decay in Eq. (9), the derivative of the transmission probability with respect to $U$ is obtained as

$$\frac{\partial T}{\partial U}(E_F^s + eU,V,d) = 2me\frac{\partial}{\partial U} \frac{T(E_F^s + eU,V,d)}{\hbar^2\kappa(U,V)} = -2\frac{\partial}{\partial V} \frac{T(E_F^s + eU,V,d)}{\hbar^2\kappa(U,V)}. \quad (A1)$$

Here, we also considered the bias-derivative of the transmission, Eq. (20). Therefore, for this particular transmission function, $B_2(V,d) = -2B(V,d)$, and the $dI/dV$ can be expressed as

$$\frac{dI}{dV}(V,d) = \frac{dI}{dU}(V,V,d) + B(V,d) + D_T(V,d) = \frac{dI}{dU}(0,V,d) - B(V,d) - D_S(V,d)$$

$$= \frac{1}{2} \left[ \frac{dI}{dU}(0,V,d) + \frac{dI}{dU}(V,V,d) \right] + \frac{1}{2}[D_T(V,d) - D_S(V,d)] \quad (A2)$$

This formulation helps the better understanding of the structure of the differential conductance, and its contributing terms, and could prove to be useful for extracting information about the tip and sample electronic structures from experimentally measured spectra in the future.