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**Ab initio** nonequilibrium quantum transport and forces with the real-space projector augmented wave method

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We present an efficient implementation of a nonequilibrium Green’s function method for self-consistent calculations of electron transport and forces in nanostructured materials. The electronic structure is described at the level of density functional theory using the projector augmented wave method to describe the ionic cores and an atomic orbital basis set for the valence electrons. External bias and gate voltages are treated in a self-consistent manner and the Poisson equation with appropriate boundary conditions is solved in real space. Contour integration of the Green’s function and parallelization over $k$ points and real space makes the code highly efficient and applicable to systems containing several hundreds of atoms. The method is applied to a number of different systems, demonstrating the effects of bias and gate voltages, multiterminal setups, nonequilibrium forces, and spin transport.

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I. INTRODUCTION

Electron transport across nanostructured interfaces is important in a range of different areas, including nanoelectronics, organic photovoltaics, and electrochemistry. First-principles modeling of electron transport at the nanoscale has so far mostly been applied to molecular junctions consisting of molecules contacted by metallic electrodes. However, more recent applications also include graphene nanoribbons and semiconducting and metallic nanowires, and bulk tunneling junctions for magnetoresistance and electrochemical applications. The rapid developments in these areas toward atomic-scale control of interface structures, and the continuing miniaturization of electronics components makes the development of efficient and flexible computational tools for the description of charge transport at the nanoscale an important endeavor.

The vast majority of first-principles electron transport studies have been based on density functional theory (DFT) within the local density (LDA) or generalized gradient approximations (GGA). This approach is in principle unjustified because the eigenvalues of the effective Kohn-Sham Hamiltonian do not represent the true quasiparticle energy levels. In particular, for tunneling junctions the energy gap between the highest occupied states and lowest unoccupied states is too small and this can lead to an overestimation of the conductance. More accurate calculations, including self-interaction corrections and more recently the many-body GW approximation yield conductance values in better agreement with experiments. On the other hand, the nonequilibrium Green’s function (NEGF)-DFT approach often provides a satisfactory qualitative description and its computational simplicity makes it a powerful tool for addressing nonequilibrium properties of complex systems. It should be mentioned that the formal problems associated with the use of DFT for transport are overcome by time-dependent DFT (TDDFT) which allows for an, in principle, exact description of the (longitudinal) current due to an externally applied field. However, it has been recently found that the standard TDDFT exchange-correlation potentials do not yield any improvement over the NEGF-DFT in terms of accuracy in predicting conductance.

In addition to the electronic current, it is of interest to model the forces acting on the atoms under nonequilibrium conditions, i.e., under a finite bias voltage. Such forces ultimately determine the stability of current-carrying molecular devices, but can also be exploited to deliberately control the motion of single molecules by, e.g., injecting electrons into the molecular orbitals using a scanning tunneling microscope (STM).

In this paper we present the implementation of the NEGF-DFT method in the grid-based projector augmented wave (GPAW) electronic structure code. In GPAW the electronic states can be described either on a real-space grid or using an atomic orbital basis set. For the NEGF calculations, the Green’s function is expanded in the atomic orbital basis while the Poisson equation is solved in real space. Contour integration and sparse matrix techniques, together with parallelization over both $k$ points and real space is exploited for optimal efficiency. Although the basic elements of our implementation are not new and have been described in earlier papers, the possibility of applying a general gate and/or finite bias voltage, the use of multiple leads, and inclusion of nonequilibrium forces on the ions provides a flexible and efficient computational platform for general purpose modeling of charge transport at the nanoscale and should be of interest to a large and growing community.

This paper is organized as follows. In Sec. II the transport model and formalism are introduced. In Sec. III we describe the numeric details including the complex contour integration technique used to obtain the nonequilibrium electron density from the Green’s function, the use of sparse matrix methods, and the real-space solution of the Poisson equation. A number of illustrative applications are presented in Sec. IV. In Sec. V the computational details of these examples are listed, and Sec. VI is reserved for a brief summary.

II. METHOD

The transport model is shown in Fig. 1. Following the standard approach, the system is divided into left and right electrodes and the central scattering region (see the detailed description in the caption). The Hamiltonian of the system...
The whole system can be subject to an external bias or condition for the Poisson equation, is also obtained from the electrode employing periodic boundary conditions in all directions. The Hartree is periodic and can be obtained from a ground-state DFT calculation of atoms is sandwiched between two semi-infinite electrodes. Periodic structure of interest (e.g., a molecule) and part of the electrode is calculated self-consistently in the presence of such external fields.

is given by (all notation related to PAW methodology is consistent with earlier GPW papers)

$$\tilde{H} = -\frac{1}{2} \nabla^2 + \tilde{v}_{\text{eff}} + \sum_{a \text{i, j} \text{label the PAW projector functions of a given atom. Using a (nonorthogonal) atomic orbital basis set, the Hamiltonian can be written in the following generic form:}}$$

$$\begin{bmatrix} H_{LL} & H_{LC} & 0 \\ H_{CL} & H_{CC} & H_{CR} \\ 0 & H_{RC} & H_{RR} \end{bmatrix}.$$ (2)

The “on-site” Hamiltonian matrices of the electrodes, $H_{LL}$ (left) and $H_{RR}$ (right), and the coupling matrices $H_{LC}$ and $H_{RC}$, can be obtained from a homogeneous bulk calculation. If a bias voltage $V$ is applied, the matrices corresponding to the left and right electrodes should be shifted by $eV$ relative to each other, e.g., $H_{LL} \rightarrow H_{LL} + eV S_{LC}$ and $H_{LL} \rightarrow H_{LL} + eV S_{RL}$, where $S$ denotes the overlap matrix. We assume that there is no coupling between basis functions belonging to different electrodes. This assumption can be always satisfied by making the scattering region large enough.

The retarded Green’s function is written as

$$G^r(\varepsilon) = \left[ (\varepsilon - H_{CC}) - \Sigma^r_{L}(\varepsilon) - \Sigma^r_{R}(\varepsilon) \right]^{-1}.$$ (3)

The self-energies $\Sigma^r_{L/R}$ represent the coupling to the electrodes and are obtained using the efficient decimation technique.31

The lesser Green’s function is written as

$$G^s(\varepsilon) = G^r(\varepsilon)\Sigma^s(\varepsilon)G^a(\varepsilon) + (1 + G^r(\varepsilon)\Sigma^r(\varepsilon)G^a(\varepsilon))(1 + \Sigma^a G^r).$$ (4)

The latter term is nonzero for truly bound states and vanishes for states acquiring any width.

The pseudodensity matrix (for the pseudowave in the PAW framework) is the integral of $G^r$,

$$D = \frac{1}{2\pi i} \int_c \left[ G^r(z) - G^a(z) \right] dz + \frac{i}{2\pi} \int_{E_f + eV/2}^{E_f - eV/2} G^r(\varepsilon) d\varepsilon$$

$$- 2\pi \theta i \sum_{\varepsilon_i} G^r(\varepsilon_i).$$ (5)

Here $\theta = k_B T$ and $C$ is a contour for the integral to be discussed further in Sec. III.

The nonequilibrium electron density is obtained as

$$\tilde{n}(\mathbf{r}) = \sum_{\nu \mu} D_{\nu \mu}(\mathbf{r})^* \Phi_{\mu}(\mathbf{r}) + \sum_a \tilde{n}_a^a,$$ (6)

where $\Phi_{\nu}$ is an atomic orbital basis function and $\tilde{n}_a^a$ is the atomic pseudocore density. As is standard in the PAW formalism a tilde indicates a smooth quantity as opposed to an all-electron quantity. The smooth charge density is given by

$$\tilde{\rho}(\mathbf{r}) = \tilde{n}(\mathbf{r}) + \sum_a \sum_{\ell m} Q_{\ell m}^a \tilde{\gamma}_{\ell m}^a(\mathbf{r}),$$ (7)

where $Q_{\ell m}^a$ are multipole moments and $\tilde{\gamma}_{\ell m}^a(\mathbf{r})$ is a so-called shape function. The last term is the contribution to the charge density coming from the positively charged nuclei.

The effective potential is found as

$$\tilde{v} = \tilde{v}_{\text{coul}} + \tilde{v}_{\text{xc}} + \sum_a \tilde{v}_a^a,$$ (8)

where the Coulomb potential satisfies the Poisson equation $\nabla^2 \tilde{v}_{\text{coul}} = -4\pi \tilde{\rho}$, while $\tilde{v}_{\text{xc}}$ and $\tilde{v}_a^a$ are the exchange-correlation potential and zero potential, respectively. $\tilde{v}_a^a$ is a parameter chosen to smoothen $\tilde{v}$ and which vanishes outside the augmentation sphere of atom $a$.31

To obtain self-consistency we thus have the iteration process $D \rightarrow \rho \rightarrow \tilde{V}_{\text{eff}} \rightarrow H \rightarrow D \rightarrow \ldots$. After convergence the current can be calculated by

$$I(V) = \frac{1}{\pi} \int_{-\infty}^{\infty} [f_L(\varepsilon) - f_R(\varepsilon)] \text{Tr} \left[ \Gamma_L(\varepsilon) G^r(\varepsilon) \right] d\varepsilon,$$ (9)

where $\Gamma_L/R(\varepsilon) = i(\Sigma^r_L/R(\varepsilon) - \Sigma^r_{L/R}(\varepsilon))$. For a derivation of the current formula we refer the reader to Ref. 34 (orthogonal basis) or Ref. 35 (nonorthogonal basis).

The nonequilibrium force is obtained from the derivative of the total energy with respect to atomic positions. In the PAW framework, the total energy is written

$$E = \tilde{E} + \sum_{a} (E_a^a - \tilde{E}_a^a),$$ (10)

with

$$\tilde{E} = \sum_{\nu \mu} \rho_{\nu \mu}(\Phi_{\mu}) + \frac{1}{2} \int \tilde{\rho}(\mathbf{r}) \tilde{\rho}(\mathbf{r})^* d\mathbf{r} d\mathbf{r}$$

$$+ \sum_a \int \tilde{n}(\mathbf{r}) \tilde{v}_a^a(\mathbf{r}) d\mathbf{r} + E_{\text{xc}}[\tilde{n}].$$
The force can be obtained as

$$ F^q = -\frac{\partial E}{\partial R^q}, $$

where

$$ \frac{\partial E}{\partial R^q} = \sum_{\nu\mu} \frac{\partial E}{\partial \rho_{\nu\mu}} \frac{\partial \rho_{\nu\mu}}{\partial R^q} + \sum_{\nu\mu} \frac{\partial E}{\partial T_{\nu\mu}} \frac{\partial T_{\nu\mu}}{\partial R^q} + \sum \int \frac{\delta E}{\delta \tilde{n}(r)} \frac{d\tilde{G}^q(r)}{dR^q} dr + \int \frac{\delta E}{\delta \tilde{v}^q(r)} \frac{d\tilde{v}^q(r)}{dR^q} dr + \sum_{h,c} \frac{\partial E}{\partial D^q_{hc}} \frac{\partial D^q_{hc}}{\partial R^q}. $$

We note that the expression given above does not include the recently discussed Berry phase contributions to the nonequilibrium force.\(^{24}\)

III. NUMERICAL DETAILS

A. Contour integration technique

The contour for the Green’s function integral in Eq. (5) is shown in Fig. 2. The retarded and lesser Green’s functions are integrated along the path AB (see Fig. 2) in the upper half plane and the path EF closely above the real axis in the bias window.\(^{29,36}\)

For the retarded Green’s function we use Gaussian quadrature by which a precision corresponding to a \(2^{10} = 1024\) integral in the complex energy plane. The coordinates of the indicated points are as follows: A (\(E_{\text{min}},0\)), where \(E_{\text{min}}\) is less than all the eigenenergies of the system, which is usually taken as \(\mu_L - 100 eV\) since we only calculate the valence electrons states; B (\(\mu_L + mk_BT, \Delta\)), where \(m\) satisfies \(e^{-m} \approx 0\) (a typical value of \(m\) is 10), \(k_B\) is the Boltzmann constant, \(T\) is the electron temperature, \(\Delta\) is normally between \(\epsilon_0\) and \(\epsilon_{a+1}\), where \(\epsilon_i = (2i - 1)\pi\) (with \(i\) a positive integer) is a pole of the Fermi-Dirac distribution function, thus the singulars below \(\Delta\) should be counted when summing up the residues; C (\(\mu_L - mk_BT, \Delta\)); D (\(\mu_R + mk_BT, \Delta\)); E (\(\mu_L - mk_BT, \eta\)), \(\eta\) is infinitesimal to avoid the inversion divergence; F (\(\mu_R + mk_BT, \eta\)).

![FIG. 2. (Color online) The contour used for the Green’s function integral in the complex energy plane. The coordinates of the indicated points are as follows: A (\(E_{\text{min}},0\)), where \(E_{\text{min}}\) is less than all the eigenenergies of the system, which is usually taken as \(\mu_L - 100 eV\) since we only calculate the valence electrons states; B (\(\mu_L + mk_BT, \Delta\)), where \(m\) satisfies \(e^{-m} \approx 0\) (a typical value of \(m\) is 10), \(k_B\) is the Boltzmann constant, \(T\) is the electron temperature, \(\Delta\) is normally between \(\epsilon_0\) and \(\epsilon_{a+1}\), where \(\epsilon_i = (2i - 1)\pi\) (with \(i\) a positive integer) is a pole of the Fermi-Dirac distribution function, thus the singulars below \(\Delta\) should be counted when summing up the residues; C (\(\mu_L - mk_BT, \Delta\)); D (\(\mu_R + mk_BT, \Delta\)); E (\(\mu_L - mk_BT, \eta\)), \(\eta\) is infinitesimal to avoid the inversion divergence; F (\(\mu_R + mk_BT, \eta\)).](image)

The force can be obtained as

$$ F^q = -\frac{\partial E}{\partial R^q}, $$

where

$$ \frac{\partial E}{\partial R^q} = \sum_{\nu\mu} \frac{\partial E}{\partial \rho_{\nu\mu}} \frac{\partial \rho_{\nu\mu}}{\partial R^q} + \sum_{\nu\mu} \frac{\partial E}{\partial T_{\nu\mu}} \frac{\partial T_{\nu\mu}}{\partial R^q} + \sum \int \frac{\delta E}{\delta \tilde{n}(r)} \frac{d\tilde{G}^q(r)}{dR^q} dr + \int \frac{\delta E}{\delta \tilde{v}^q(r)} \frac{d\tilde{v}^q(r)}{dR^q} dr + \sum_{h,c} \frac{\partial E}{\partial D^q_{hc}} \frac{\partial D^q_{hc}}{\partial R^q}. $$

We note that the expression given above does not include the recently discussed Berry phase contributions to the nonequilibrium force.\(^{24}\)

B. Sparse matrix handling

Because the matrix inversion cost scales as \(N^3\), where \(N\) is the dimension of the matrix, the matrix inversion turns out to be the main computational cost for large systems. Hence a sparse matrix method is implemented to obtain the Green’s function.

We define a quenching layer as a slab whose left side has no overlap with the right side due to the finite cutoff in the range of the atomic orbitals. Hence an overlap or Hamiltonian matrix can be split into several blocks, with each block representing the on-site values of a quenching layer or the coupling between
two adjacent quenching layers. Note that quenching layers here are
different from the principal layers used in the transport
framework, where the latter is supposed to be repeatable as
well.
Physical quantities such as density or transmission are
often determined by fairly few blocks of a matrix. To see
this consider the simple example of a two-probe system. In
this case, the scattering region is divided into five quenching
layers. Figure 3 shows the sparse matrix structure of the overlap
region are from electrode calculations and are always
fixed.
First we discuss how to obtain the real-space pseudodensity
matrix which can be obtained by a projection of the pseudodensity
matrix as in Eq. (6). We see that if the states $\Phi_i(r)$ and $\Phi_j(r)$
no overlap, the contribution from the pseudodensity
matrix is zero, i.e., the white blocks in Fig. 3(a) do not affect
the density. So when we calculate the density matrix from the
integral of the Green’s function, only the blue and green blocks in
the Green’s function matrix [Fig. 3(a)] are necessary. There are
really two different parts, because two types of Green’s
functions are involved when calculating the density matrix:
the equilibrium part and the nonequilibrium part. We need
the blue and green blocks of the retarded Green’s function
for the former and of the Keldysh Green’s function for the
latter. Through Eq. (4) and the finite extent of the self-energy
matrix, which is only nonzero in the principle layers close to
the electrodes, we see that the blue and green blocks in Fig. 3
in the Keldysh Green’s function matrix can be obtained from
only the red blocks of the retarded Green’s function [Fig. 3(c)].
So when we do the matrix inversion to calculate the retarded
Green’s function by Eq. (3), the red blocks in Fig. 3(a) are
necessary for energy points on the path EF in Fig. 2, and the
blue and green blocks in Fig. 3(c) are necessary for energy points
on the other path segments in Fig. 2.
We can also see that the red and orange blocks in Fig. 3(b)
are needed to calculate the density of states (DOS) by
$\text{DOS}(\epsilon) = -\frac{1}{\pi} \Im [\text{Tr} G(\epsilon^+) S]$, and the pink blocks in Fig. 3(b)
are needed to calculate the transmission function $T(\epsilon) =
\text{Tr} [\Gamma_L(\epsilon) G(\epsilon^+) \Gamma_R(\epsilon) G(\epsilon^+)]$.
The formulas below provide a quick solution for the
necessary. Here we just consider this particular matrix
(shown in Fig. 3) as an example to show how the method
works. A general formalism, which works for arbitrary number
of electrodes and arbitrary number of principal layers in each
electrode, is introduced in Ref. 39.
First, the central block $N$ in Fig. 3(b) of the retarded Green’s
function can be solved through the equations
\[
Q^L_{1} = L_{22}^{-1}, \quad Q^L_{2} = (L_{11} - L_{12} Q^L_{2} L_{21})^{-1}, \quad (15)
\]
\[
Q^R_{1} = R_{22}^{-1}, \quad Q^R_{2} = (R_{11} - R_{12} Q^R_{2} R_{21})^{-1}, \quad (16)
\]
\[
N = \left( M - \sum_{J=L,R} J_{12} Q^J_{12} J_{21} \right)^{-1}, \quad (17)
\]
where $L_{ij}$ and $R_{ij}$ are the blocks shown in Fig. 3(a) rep-
resenting the matrix $eS - H_{CC} - \Sigma^L_{1}(\epsilon) - \Sigma^R_{0}(\epsilon)$. Then, for the
remaining blocks of the retarded Green’s function matrix, we
have to iterate the formulas
\[
G^L_{i,j} = Q^L_{i,j} (I - L_{i,j-1} G^L_{i-1,j}),
\]
\[
G^L_{i,j} = (I - G^L_{i-1,j} R^L_{i,j}) Q^L_{i,j},
\]
\[
G^R_{i,j} = -Q^R_{j,i} L_{i,j-1} G^R_{i-1,j},
\]
\[
G^R_{i,j} = -G^R_{j,i-1} L_{i,j-1} Q^R_{j,i}, \quad (j < i), \quad (18)
\]
where $G^L_{i,j}$ is the block from the central part to electrode $L$
and $G^L_{i,0}$ is $N$ in Eq. (18), the blocks from the central part
to electrode $R$ have a similar solution. For all the required blocks,
a quick solution can be obtained using a combination of the
recursive formulas (18). If we denote the number of quenching
layers by $n$, the computational cost is roughly given by $n$ times
the cost of an inversion operation plus $4n$ times the cost of
matrix multiplication.39

C. Fixed boundary conditions
The electronic potential of the metal electrodes will usually
be very efficiently screened so that after only a few atomic
layers into the electrodes we can assume that the potential is
equal to the equilibrium potential plus a possible constant
bias potential, which is normally eV/2 for the cathode
and $-eV/2$ for the anode (V stands for the bias voltage).
We shall apply open boundary conditions (in contrast to, say,
periodic ones) where the bias is applied by fixing the potential
values at the boundaries before solving the Poisson equation.40
This procedure also allows for a net charge in the scattering
region, in which case the perturbation of the electron potential
into the electrodes will of course be somewhat more long
ranged.
The Poisson equation $\nabla^2 \psi_{\text{coul}} = -4\pi \rho$ is solved in reciprocal
space in the $x$ and $y$ directions while it is solved in real
space in the $z$ coordinate, i.e., in the the transport direction.
Mathematically we have
\[
\left( \frac{d^2}{dz^2} - \tilde{G}^2 \right) \tilde{\psi}_{\text{coul}}(z, \tilde{G}) = -4\pi \tilde{\rho}(z, \tilde{G}), \quad (19)
\]
where $\tilde{G}$ is the vectors of the two-dimensional (2D) real grids
used for the Fourier transformation.
Equation (19) is solved by the sparse matrix linear equation
subroutine provided by the LAPACK package. An advantage
of this Poisson equation solution is the good parallelization
behavior. The 2D Fourier transformations are independent for
the different real-space slices, and the linear equations (19)
can be solved independently for different $\tilde{G}$ vectors.

IV. RESULTS
A. Quantum capacitor
We consider a simple capacitor system consisting of two
semi-infinite Na electrodes separated by a vacuum gap (see
the upper part of Fig. 4). When a bias voltage is applied to
the system, electrons accumulate or deplete on the Na(100)
surfaces. According to the classic parallel plate capacitor
model, the surface charge should be
\[
Q_{\text{cl}} = e_0 V A / d, \quad (20)
\]
where $\varepsilon_0$, $A$, $d$ are the vacuum permittivity, area cross section of the unit cell, and the gap distance, respectively. We integrate the induced charge density in real space and obtain the net charge accumulation $Q = 0.45e$, which is close to the result of classic theory $Q_{cl} = 0.55e$. One difference is that, in the quantum theory, the charge decays from the surface into the bulk in an oscillatory fashion (Friedel oscillations), instead of being localized exactly on the surface as assumed in the classic model. We also note that at a distance of about four layers from the surface the values for both the potential and the charge are very close to their bulk values. Hence this calculation confirms the screening approximation, namely, that a few layers away from the surface or scattering region the potential has reached its bulk value. Finally, we note that the relatively high bias voltage of $\approx 5$ V is possible in the present case where no current is flowing. On the other hand, the nonequilibrium states determining the current flow become increasingly difficult to calculate accurately for larger bias values due to the insufficient integral of the Keldysh Green’s function in the bias window. As a consequence, electron transport calculations are typically possible or reliable up to bias voltages of around 2–3 V, depending on the transparency of the junction.

B. Nonequilibrium forces

The calculation of nonequilibrium forces is in principle a delicate problem involving nonconservative components. For highly conducting molecular bridges an instability may occur which involves the Berry phase of the wave function. The description of such phenomena is beyond the scope of the usual NEGF + DFT framework, but in simpler cases, in particular, in cases with no or little current flow, the force expressions (11) and (12) still apply. As an example, we show here a nonequilibrium force calculation for a Au/N2/Au junction, where we can see the tendency toward molecular dissociation under a bias voltage. The structure (see the upper part of Fig. 5) is relaxed under zero bias until the maximum force is below 0.01 eV/Å. When a positive bias is applied, electrons are redistributed over the molecule due to the electric field. Consequently, the two nitrogen atoms start to repel each other due to increased Coulomb repulsion which weakens the bond. The actual quantity of charge transfer to the molecule, which is about 0.01e for 1-V bias voltage on this system, shifts up the molecular energy spectrum, i.e., the energy levels follow the chemical potential of the left electrode (see the middle panel of Fig. 5). The force is mainly occurring only between the two nitrogen atoms, while there is no force induced between the electrode atoms and the N2 molecule. Equivalently, a negative bias voltage shifts down the levels and pull electrons out of the N2 molecule, and leads to an attractive force between the two nitrogen atoms.

C. Electrostatic gate control

One way of controlling the current flow through a nanoscale conductor is by electrostatic gating. This has been
FIG. 5. (Color online) The bias voltage effect on a nitrogen molecule between two gold electrodes. Upper panel: The atomic structure of the system. The arrows represent the directions of the atomic forces generated by the bias voltage. Middle panel: PDOS (partial density of states) of the nitrogen molecule at 0- and 1-V bias voltage. The Fermi level is located at zero and the red dashed lines show the location of the bias window in the 1-V case. Lower panel: The magnitude of the nonequilibrium atomic forces as a function of bias voltage.

demonstrated experimentally for graphene, where a metal-insulator transition was induced by gating,
and for single-molecule junctions where the individual electronic levels were moved in energy relative to the Fermi level of the source/drain electrodes. At the single molecule scale, the gate-molecule coupling is to a large extent determined by the device geometry with the screening effect playing an important role. For numerical simulations, the typical method of applying a gate is to add an external potential \( v_g(r) \) to the effective potential of the system:

\[
\text{\( \tilde{v}(r) = \tilde{v}_0(r) + v_g(r) \).} \]

We now consider the prototypical Au/benzene-1,4-dithiolate/Au (Au-BDT-Au) junction (see the upper panel of Fig. 6) subject to three different gate potentials, \( v_g(z) \) (see the middle panel of Fig. 6). We note that the structure of the Au/BDT junction is presently being debated. However, for our purpose the simple model makes the sense.

In the lower part of Fig. 6 we plot the resulting effective gate potential \( \Delta v(z) = \tilde{v}_s(z) - \tilde{v}_0(z) \), where the subscript \( s \) denotes self-consistency, and the superscript 0 denotes zero gate voltage. Due to the screening in the metal, the effective gate potentials only affect the molecule region, and the narrower gate potential is seen to be less influenced by the self-consistency because it does not induce considerable charge transfer at the metal surfaces—a charge transfer that otherwise tends to reduce the gate effect on the molecule.

We note that the gate efficiency factor, \( \alpha = \Delta v(z) / v_g(z) \), for these three potentials are about 0.8, 0.6, and 0.4 in the molecular region, with the larger efficiency obtained for the more localized gate potential. The value of 0.4 obtained for the most delocalized potential is fairly close to an experimental study of the Au-BDT-Au system where an efficiency factor of 0.25 was reported.

In the following we illustrate how the gate voltage can be used to tune the conductance of a molecular junction. It has recently been shown that the transport through the molecule anthraquinone is strongly suppressed due to destructive quantum interference occurring close to the Fermi level when the molecule is connected to metallic electrodes. The quantum interference leads to a dip in the transmission function inside the energy gap between the highest occupied (HOMO) and lowest unoccupied (LUMO) molecular orbitals. Hence a large on-off ratio is expected when shifting the molecular energy levels by an external gate voltage. The upper panel of Fig. 7 shows the molecule connected to two gold fcc(111) surfaces through Au-S bonds. The effective potential with the gate voltage \(-2\) V is shown in the middle panel. We see that the potential of the central part of the anthraquinone molecule is shifted less than the potential for the outer parts of the molecule. This is due to the fact that different parts of the molecule polarize differently as a consequence of the detailed electronic structure. The HOMO is, for example, mainly localized at the connecting wires. The lower panel of Fig. 7 shows the change of transmission coefficient when a gate voltage of \(-2\) V is applied. Due to the characteristic
interference dip in the transmission a large on-off ratio of about a factor of 1000 is achieved. Note that the relatively poor gate efficiency of around 0.1 is due to Fermi-level pinning of the HOMO.

D. Spin transport

In this section, we investigate the nonequilibrium-driven magnetic transition in the spin transport in a zigzag graphene nanoribbon (ZGNR) which is proposed in Ref. 52 based on tight-binding calculations. The ZGNR’s edge is spin polarized and it has an antiferromagnetic spin configuration if its number of atomic layers is even.53 A gap of about 1 eV is opened between the different spin states and makes the ZGNR a semiconductor. It was noticed by Denis et al. that the ZGNR’s magnetic ordering is killed when the external bias voltage exceeds the size of the gap.52 Here we reproduce this result for the graphene/ZGNR/graphene system shown in the upper panel of Fig. 8, where a ZGNR (nn = 8) is sandwiched between two semi-infinite graphene flakes. Under zero bias the PDOS of the central C atom along the ZGNR edge shows two peaks above and below the Fermi level, corresponding to the different spin directions (the middle panel in Fig. 8). The distance between the two peaks is about 1.6 eV, and equals the band gap. When this bias voltage reaches 1.0 V, the current starts to increase (see Fig. 9). At bias voltage 2.0 V the edge magnetic moment disappears very abruptly, and the current starts to increase even faster.

Interestingly, in the tight-binding calculations presented in Ref. 52, both the magnetic moment and the current show a very abrupt feature at the bias threshold, while in our calculation, the current increases rather smoothly. This can be explained by the nonequilibrium potential in the ab initio calculation leads to a rehybridization and broadening of the spectral peaks (see the lower panel of Fig. 8). We also note that in our calculation the disappearance of the magnetic moment is due to the two
FIG. 10. (Color online) Left: The C\textsubscript{60}-6-terminal structure and real-space DOS at the Fermi level. Right: The averaged effective potential projected onto a plane cutting through the C\textsubscript{60} molecule.

Stoner peaks moving into the bias window being half occupied, which is different from the complete band collapse in Ref. 52; this is because our ZGNR is not long enough. We can see from the lower panel of Fig. 8 that the gap shrinks more for the C atom further from the contact.

E. Multiterminal transport

The expression for the Green’s function of the scattering region Eq. (3) can be straightforwardly extended to a multiterminal situation

\[
\mathcal{G}^r(\epsilon) = \left[ \epsilon S - H_{CC} - \sum_j \Sigma_j(\epsilon) \right]^{-1},
\]

where \(J\) is the index of the terminals. In contrast to the situation for a two-probe calculation, a zero boundary condition is applied for the effective potential for a multiterminal system, and buffer atoms are used to represent semi-infinite leads. This approach to multiterminal transport has been previously investigated in Ref. 54. It should be noted that the self-energy of a lead has to be “rotated” by an orthogonal transformation when the lead is not along either the \(x\), \(y\), or \(z\) axes.

As an example, we consider a C\textsubscript{60} molecule connected to six linear carbon atomic chains. Figure 10 (left) shows the projected DOS in real space evaluated at the Fermi level. The coverage suggests that the scattering states are itinerant in the whole system and the contact between the carbon chain and the C\textsubscript{60} molecule is transparent. Figure 10 (right) shows a 2D averaged potential in a plane cutting through the C\textsubscript{60} molecule.

A matrix indicating the transmission at the Fermi level between the different leads is shown in Fig. 11. The matrix index notation represents the lead number as shown in Fig. 10 (left). In particular, the diagonal corresponds to backscattering, i.e., it gives the reflection probability. We can see that electrons are more easily transmitted between leads opposing each other, whereas the transmission decreases if the electron has to turn an angle during the scattering process. This intuitive phenomenon can be explained by the quantum interference of the different partial waves. For the straightforward scattering, the quantum phases are the same for all the paths passing through the C\textsubscript{60} molecule, and the electron therefore attains the greatest transmission probability.\textsuperscript{51}

V. COMPUTATIONAL DETAILS

For completeness we list the key input parameters and CPU timings for the examples presented in this paper in Table I.

VI. SUMMARY

We have described the implementation of the NFGF + DFT transport method in the GPAW code and illustrated its performance through application to a number of different molecular junctions. The electronic structure is described within the PAW methodology which provides all-electron accuracy at a computational cost corresponding to pseudopotential calculations. The Green’s functions are represented in a basis set consisting of atomiclike orbitals while the Poisson equation with appropriate boundary conditions is solved in real space. The
code is parallelized over $k$ points and real-space domains and sparse matrix techniques are applied for maximal efficiency. The flexibility of the method was illustrated through examples demonstrating electron transport under finite bias voltage, effect of electrostatic gating, spin transport, nonequilibrium forces, and multiterminal leads.

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