Electronic friction for a slow impurity in an inhomogeneous metallic system

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Abstract

The stopping power of a metal for a slow structureless impurity varies linearly with projectile velocity. We show that the coefficient of this linear behavior (friction coefficient) is determined exactly in a static ensemble Kohn-Sham scheme, by extension of the Shifted Fermi Surface procedure originally derived for an homogeneous jellium. We prove that the friction coefficient is determined only by local characteristics of the system. The error incurred when adding a spurious non-local contribution is illustrated for a simple 1D model.

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I. INTRODUCTION

The stopping power of a free (homogeneous) electron gas (FEG) for a structureless atom or ion varies linearly with projectile velocity in the limit of low velocities (see, e.g., Ref. 1–4). This result has been proved at the level of both linear and non-linear theories of stopping. The coefficient of this linear behavior is referred to as the friction coefficient and the associated process as electronic friction. Such a behavior has been observed in experiments for ion stopping in metals that are expected to mimic a FEG in this respect. Examples are for H, D and He ions in Al, H and He ions in Au and Cu, or H and He ions in Al, Zn and Au. It has also been observed in grazing collisions of ions on surfaces, like H and He ions on the Al(111) surface.

One important field in which electronic friction might play an important role, is that of surface chemistry, as a candidate for energy dissipation by adsorbates. It is crucial to assess its importance for such processes as atomic and molecular adsorption, diffusion on surfaces, range of hot atoms following molecular dissociation at surfaces, etc. Then, the adsorbate is immersed into a strongly non homogeneous electronic density which raises questions as to the relevance of values of friction coefficients obtained from calculations based on the FEG model. It is, therefore, desirable to develop methodologies that can make use of present day band structure codes to evaluate the friction coefficient. Such a task has been endeavored by Trail et al., based on the formulation proposed by Hellsing and Persson. Results have been obtained for H and D on Cu(111) or for H₂ on Cu(111) and N₂ on Ru(0001). However, some questions are raised by the latter results. Firstly, the H/Cu results are divergent at a certain value of the atom-surface distance. Secondly, the friction coefficient for N₂/Ru may reach values an order of magnitude larger than usually obtained from the FEG model.

In the present work, we reexamine the formulation of the friction coefficient determination from first principles in order to ensure that we get a consistent first order approximation in the projectile velocity. Our approach is based on Density Functional Theory (DFT) and extends the methodology developed for the homogeneous FEG to the inhomogeneous case.

Atomic units are used throughout.
We consider a medium composed of electrons moving in a periodic array of fixed nuclei. We are interested in the projectile energy loss due to the electronic state perturbation by the projectile motion, the role of the projectile frozen-lattice interaction (i.e., with the lattice nuclei) being trivial. The “external” potential, in which the electrons move, is composed of two parts: one, $V_{\text{lat}}$, due to the fixed nuclei and the other, $V_{eP}(t)$, to the moving, structure-less, projectile. As the projectile position with respect to the lattice, $R_P$, is time-dependent, the friction coefficient is a function of $R_P$. We suppose that the static problem, with the projectile at rest, has already been solved and restrict our study to the limit when the projectile velocity goes to zero, since the friction coefficient is associated with the first order approximation to the stopping in its power series expansion as a function of the projectile velocity. In principle, the density of the system, as a function of time, may be determined through the time-dependent Kohn-Sham (KS) scheme of the Time Dependent Density Functional Theory (TDDFT, for a general presentation, see e.g., Ref. [15]). However, in the low velocity limit, it is not necessary to resort to TDDFT as we show below.

Our strategy is as follows. In the present section we study the behavior of the system in the adiabatic limit. In the next section we show that the non-adiabatic corrections do not contribute to the friction coefficient.

Let us start with the projectile at rest in the lattice frame without interaction with the electrons. We could first solve the static problem by turning on the projectile-electron interaction and, subsequently, set the slow projectile motion. However, in the adiabatic limit, we may as well first set the projectile motion, without interaction with the electrons, and determine the transformation from the lattice to the projectile frame. We then turn on the projectile-electron interaction adiabatically. The latter procedure is justified since in the adiabatic limit the final state is independent of the evolution path.

A. Transformation to the projectile frame

We first consider the projectile moving without interaction with the electrons. We assume that the projectile velocity, $v$, is constant so that the transformation from the lattice to the projectile frame is Galilean. Let $\mathbf{r}_i$ be the position vector of electron $i$ with respect to an
origin fixed in the lattice frame and \( \mathbf{r}_i^P \) w.r.t. the projectile. The N-electron wave-function \( \Psi_T(\mathbf{r}_1, ..., \mathbf{r}_N) \) becomes in the projectile frame (discarding in this section, for short, the energy phases that play here no role):

\[
\Psi_P(\mathbf{r}_1^P, ..., \mathbf{r}_N^P; \mathbf{R}_P(t)) = e^{-i \sum_{j=1}^{N} \mathbf{v} \cdot \mathbf{r}_j^P} \Psi_T(\mathbf{r}_1, ..., \mathbf{r}_N)
\]  

with \( \mathbf{r}_i = \mathbf{r}_i^P + \mathbf{R}_P(t) \). The density in the target frame, and in the absence of projectile-target interaction, is exactly known through a static Kohn-Sham scheme. This static Kohn-Sham scheme is transformed into the projectile frame by a similar Galilean transformation involving the multiplication of each KS orbital in the lattice frame, \( \varphi_i^T \), by \( \exp\{-i \mathbf{v} \cdot \mathbf{r}_i^P\} \).

As the KS orbitals are Bloch functions:

\[
\varphi_{k,i}^P(\mathbf{r}; \mathbf{R}_P(t)) = e^{-i \mathbf{v} \cdot \mathbf{r}_i^P} \varphi_{k,i}^T(\mathbf{r}) = e^{i \mathbf{v} \cdot \mathbf{R}_P} e^{i (\mathbf{k} - \mathbf{v}) \cdot \mathbf{r}} u_{k,i}(\mathbf{r})
\]  

where \( u_{k,i} \) is a periodic function\(^{22}\). In the absence of a gap close to the Fermi surface:

\[
e^{-i \mathbf{v} \cdot \mathbf{r}} \varphi_{k,i}^T(\mathbf{r}) = \varphi_{k-v,i}^T(\mathbf{r}) + e^{i (\mathbf{k} - \mathbf{v}) \cdot \mathbf{r}} \mathbf{v} \cdot \nabla_k u_{k,i}(\mathbf{r}) + 0(v^2)
\]

\[
= \varphi_{k-v,i}^T(\mathbf{r}) + 0(v)
\]  

(3)

When substituting \( \varphi_{k-v,i}^T \) for \( \varphi_{k,i}^T \) (disregarding the common time dependent phase factor \( \exp\{i \mathbf{v} \cdot \mathbf{R}_P\} \)), we introduce a shifted KS scheme in the target frame, i.e. a Fermi distribution shifted by \(-\mathbf{v}\) (SFS). It can be shown that this shifted KS scheme is correct to first order in the velocity. Firstly, the total number of states within the shifted Fermi surface is conserved up to first order in \( v \). Indeed, for vanishingly small \( v \), and in the absence of a gap in the vicinity of the Fermi surface:

\[
\int \mathbf{d}k \rho(\mathbf{k} - \mathbf{v}) = \int \mathbf{d}k \rho(\mathbf{k}) + \int \mathbf{d}k_F (-\mathbf{v} \cdot \hat{\mathbf{k}}_F) \rho(\mathbf{k}_F) + 0(v^2)
\]  

(4)

where \( \rho \) is the density of levels and the volume integrals are over the volume bounded by the unshifted Fermi surface. Since the Fermi surface is symmetric with respect to the origin of the Brillouin zone, the quantity \((-\mathbf{v} \cdot \hat{\mathbf{k}}_F) \mathbf{d}k_F \) is exactly compensated by the same term for \( \mathbf{k} = -\mathbf{k}_F \). Secondly, the density is also exact to first order in \( v \).

\[
n^P(\mathbf{r}) = \int_{\text{FS}} \mathbf{d}k \left| \varphi_{k-v,i}^T(\mathbf{r}) + e^{i (\mathbf{k} - \mathbf{v}) \cdot \mathbf{r}} \mathbf{v} \cdot \nabla_k u_{k,i}(\mathbf{r}) \right|^2 + 0(v^2)
\]

\[
= \int_{\text{FS}} \mathbf{d}k \left\{ \left| \varphi_{k-v,i}^T(\mathbf{r}) \right|^2 + 2 \Re \left[ u_{k,i}^*(\mathbf{r}) \mathbf{v} \cdot \nabla_k u_{k,i}(\mathbf{r}) \right] \right\} + 0(v^2)
\]  

(5)

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The symmetry with respect to the origin of the Brillouin zone implies that $u^*_k, i = u_{-k, i}$ so that the second term in the integral gives a zero contribution when integrated over $k$, which yields:

$$n(r) = \int_{\text{SFS}} \, dk \left| \phi^T_{k, i}(r) \right|^2 + O(v^2)$$

(6)

In other terms, the SFS prescription gives the correct density up to first order in $v$ which means that the corresponding KS scheme is exact to first order in the velocity. We may re-write the previous expression for short, as:

$$n^P(r_P; R_P(t)) = n^P_{KS}(r_P; R_P(t)) = \sum_i F_i \left| \phi^T_i(r) \right|^2 + O(v^2)$$

(7)

where $F_i$ corresponds to the Fermi distribution shifted by $-v$.

As expression (7) shows, the time dependent KS solution in the projectile frame can be formulated in terms of an ensemble KS procedure in the lattice frame. When $v \ll k_F$ and for a given orientation of $k$, the occupation of orbital $\varphi_k$ is 1 for $k \leq k_m$ and zero for $k > k_m$, where $k_m = k_F \hat{k} - v$. The Fermi surface being a surface of constant energy, this entails that the population decreases as the energy increases as required for the validity of the Hohenberg-Kohn theorem for ensembles. Then, the latter theorem proves, in the non-degenerate case, that the exact state of the system in the projectile frame is also described by an ensemble in the lattice frame, univocally defined from the K-S ensemble since the latter determines both the ensemble and the density.

The system we are considering here involves degeneracies, so its ensemble is not determined univocally by the density. However, the energy remains a functional of the density and any quantity that can be expressed in terms of the density is also univocally defined. These properties are the only ones required for the validity of the following discussion.

As a conclusion, the Galilean transformation from the target to the projectile frame of the exact N-electron state yields, to first order in the velocity, an ensemble of states which are defined in the (static) target frame. In other terms, we have transformed the time-dependent problem in the projectile frame into a static one in the target frame, at the expense of using an ensemble defined by the Shifted Fermi Surface (SFS).
B. Projectile-target interaction in the adiabatic limit

1. Adiabatic evolution

Let $\Psi_j(t_i)$ be the exact solution of a time dependent problem when the system is in the eigenstate $j$ of the Hamiltonian at the initial time $t = t_i$.

$$H(t_i)\Psi_j(t_i) = E_j(t_i)\Psi_j(t_i)$$

(8)

If the evolution is adiabatic, the system remains in the eigenstate $j$ of $H(t)$ at any time, i.e., it evolves in such a way that the wave function is at any time the solution of a static problem. Evolution from two different initial states cannot lead to the same state at time $t$: adiabatic evolution is reversible. For simplicity, the derivation below assumes that the eigenenergies are not degenerate. Accordingly, the function $\Psi_j(t)$ is determined at any time $t$ (save for a time-dependent phase factor) by the energy $E_j(t)$, i.e., the evolution of the system state is determined by the knowledge of $dE_j(t)/dt$. This one to one correspondence between energy and state is central to our derivation below.

The same correspondence prevails for the evolution of an ensemble, which may be defined by the density operator:

$$\hat{\rho}(t) = \sum_j p_j |\Psi_j(t)\rangle\langle\Psi_j(t)|$$

(9)

where the weights $p_j$ are constant in time. The energy is given by:

$$E(t) = \text{Tr}\{\hat{\rho}(t) H(t)\} = \sum_j p_j E_j(t)$$

(10)

Again, the system is at all times characterized by quantities that are the solution of a static problem. Furthermore, when the $p_j$’s satisfy the same condition as that for the validity of the HK theorem for ensemble, the ensemble is determined at any time by its energy.

2. Application to the present problem

In the present problem, the adiabatic evolution is associated with that of the projectile-lattice relative position, i.e. with the variable $R_P$. What remains to be derived is the KS scheme associated with this adiabatic evolution. To this end, we switch on adiabatically the projectile-target interaction by writing the external potential as:

$$v_{\text{ext}}(r, t) = \lambda(t) V_{eP}[r, R_P(t)] + V_{\text{lat}}$$

(11)
The function $\lambda(t)$ is an arbitrary function that varies from 0 to 1 in the time interval $[t_i, t_f]$. Saying otherwise, we introduce through the function $\lambda(t)$ a fictitious evolution starting from an initial state where the projectile and electrons do not interact. Although we could keep $R_P$ fixed in this process, we conserve in (11) its time-dependence. As can be seen below, the conclusions of the present section are independent of the presence or absence of the latter time-dependence. We assume that it is always possible to choose the arbitrary function $\lambda(t)$ so that $d\lambda/dt$ is small enough, at any time, to ensure adiabaticity. The adiabatic approximation corresponds to a zero order approximation in $d\lambda/dt$ (see, e.g., 17).

Our aim is to determine the KS orbitals, at each time, when the system evolves adiabatically. To this end, we calculate the energy variation in the interval $dt$:

$$\frac{dE(t)}{dt} = \text{Tr}\{\hat{\rho}(t) \frac{dH}{dt}\}$$

$$= \int d\mathbf{r} n(\mathbf{r}, t) \left[ \frac{d\lambda}{dt} V_{eP} + \lambda(t) \mathbf{v} \cdot \nabla_{R_P} V_{eP} \right]$$

(12)

where $n$ is the exact density at $t$. According to the discussion of section II B 1, the KS orbitals must be such that they give the exact energy variation (12) in the limit of vanishing $d\lambda/dt$.

We suppose that, at a given time $t^0$, we have determined the KS orbitals for the ensemble defined by the distribution $G_i$ and study the time evolution in the interval $[t^0, t^0 + dt]$. Consider the orbitals $\varphi^a_i(\mathbf{r}, t)$ evolving adiabatically from the KS orbitals at $t^0$ for the same ensemble, i.e., assuming that $G_i$ is independent of time:

$$n^a(\mathbf{r}, t) = \sum_i G_i |\varphi^a_i(\mathbf{r}, t)|^2$$

(13)

where the orbitals $\varphi^a_i$ are the solution of a static problem

$$H_{KS} \varphi^a_i(\mathbf{r}, t) = \varepsilon^a_i(t) \varphi^a_i(\mathbf{r}, t)$$

(14)

with

$$H_{KS} = -\frac{1}{2} \nabla^2 + \lambda(t) V_{eP}(\mathbf{r}; R_P) + v_n(\mathbf{r}; [n^a(\mathbf{r}, t)])$$

(15)

and $v_n$ includes the Hartree potential, the lattice potential and the exchange correlation potential of the static problem for the KS ensemble defined by the occupations $G_i$. The question is now: what is the relation between these adiabatic KS orbitals and the KS orbitals associated with the exact adiabatic evolution of the system? Introduce the energy functional:

$$E_v[n] = \sum_i G_i \varepsilon_i(t; [n]) - \int d\mathbf{r} n(\mathbf{r}, t) v_{xc}(\mathbf{r}; [n(\mathbf{r}, t)])$$
\begin{equation}
- \frac{1}{2} \int \int \mathrm{d}r \, \mathrm{d}r' \frac{n(r, t) n'(r', t)}{|r - r'|} + E_{xc}[n] \tag{16}
\end{equation}

The functional $E_v[n]$ gives the exact energy of the static ensemble with density $n$, when the $\varepsilon_i$'s, the exchange-correlation potential $v_{xc}$ and the energy $E_{xc}$ are those of the static KS ensemble defined by the occupations $G_i$. At $t^0$, $n = n^a = n^0$ and $E_v[n^0] = E(t^0)$ where $n^0$ and $E(t^0)$ are the exact density and energy. The question is: what is the relation between $E_v[n^a]$ and $E(t)$ at $t = t^0 + dt$? We split the variation of $E_v[n^a]$ over the interval $dt$ into two terms. The first one corresponds to the variation of $n^a$:

\begin{equation}
\frac{\mathrm{d}E_v[n^a]}{\mathrm{d}t} \bigg|_{\lambda V_{eP}} = \int \mathrm{d}r \left. \frac{\mathrm{d}E_v[n^a]}{\mathrm{d}n^a} \right|_{n^a = n^0(r, t^0)} \frac{\mathrm{d}n^a(r, t)}{\mathrm{d}t} \tag{17}
\end{equation}

This term is identically zero because at $t = t^0$ the energy functional is stationary around the exact density $n^0(r, t^0)$. The second term corresponds to the explicit dependence on $\lambda(t) V_{eP}$ in (16) for a fixed $n^a$:

\begin{align*}
\frac{\mathrm{d}E_v[n^a]}{\mathrm{d}t} \bigg|_{n^a} &= \sum_i G_i \frac{\mathrm{d}\varepsilon_i^a(t)}{\mathrm{d}t} \\
&= \int \mathrm{d}r n^0(r, t^0) \left[ \frac{\mathrm{d}\lambda}{\mathrm{d}t} V_{eP} + \lambda(t) \mathbf{v} \cdot \nabla_{R_P} V_{eP} \right] \tag{18}
\end{align*}

which is identical to (12). As a conclusion, at $t^0 + dt$ the $\varphi^a$ orbitals are still the KS orbitals for the exact adiabatic evolution when the ensemble is defined by the same time independent distribution $G_i$. This result may be applied to the evolution over the whole interval $[t_i, t_f]$. Now, for $t = t_i$, i.e., in the absence of projectile target interaction ($\lambda(t) = 0$), the distribution $G_i$ is exactly given by the SFS prescription, $G_i = F_i$, $\forall i$. This means that the solution of the problem at $t = t_f$ ($\lambda(t) = 1$) is also given by the SFS prescription.

We are led to the following conclusion: the density, in the adiabatic limit, can be determined, at each position of the projectile, using a static ensemble KS scheme in which both the projectile and lattice potential are fixed. The fact that the energy is stationary with respect to an arbitrary variation of the density around its exact value (see Eq. (17) plays a central role in reaching this conclusion. The SFS Kohn-Sham scheme is an exact scheme for the determination of the system density for an impurity traveling through an inhomogeneous medium under adiabatic conditions.
III. FRICTION COEFFICIENT

The force acting on the projectile is:

$$\nabla_{R_P} E(R_P) = - \int dr \ n(r; R_P) \ \nabla_{R_P} V_e P(r; R_P)$$  \hspace{1cm} (19)

which involves only the derivative of $V_e P$ since, as for (17), the derivative of the energy functional with respect to density is zero.

When the density in (19) is the static density $n_0(r; R_P)$, the force is the static force, i.e. the limit when the projectile velocity $v$ goes to zero. When the projectile moves with a small but finite velocity $v$, the associated density is $n_v(r; R_P)$ and a dissipative process takes place which corresponds to friction. The friction coefficient for motion along the direction $\hat{v}$ is defined by:

$$F_{\hat{v}} = \lim_{v \to 0} \frac{1}{v} \left\{ \int dr \ \Delta n(r; R_P) \ \hat{v} \cdot \nabla_{R_P} V_e P(r; R_P) \right\}$$  \hspace{1cm} (20)

where

$$\Delta n(r; R_P) = n_v(r; R_P) - n_0(r; R_P)$$  \hspace{1cm} (21)

A. Non-adiabatic corrections

When $n_v$ is calculated in the adiabatic SFS approximation of section II B, the dependence of $n_v$ on $v$ arises entirely from the SFS. However, we have to check whether non-adiabatic corrections can be neglected when $n_v$ is calculated to first order in $v$.

Deviations from the adiabatic approximation correspond to inelastic transitions between adiabatic states due to the variation of $R_P(t)$. Let us evaluate the contribution of these inelastic transitions to (19) when $R_P(t)$ varies from $R_0$ to $R_1$. We study the evolution of a KS orbital equal to the adiabatic orbital $\varphi_{k_0,i_0}$ for $R_P(t_0) = R_0$. We set $\hat{v} \cdot R_P(t) = Z = vt$ and, to simplify our notations, hereafter only mention explicitly the dependence on $Z$. For $Z > Z_0$, the KS orbital becomes $\psi_{k_0,i_0}$, which we express as:

$$\psi_{k_0,i_0}(Z) = \sum_{k',j} c_{k',j}(Z) \varphi_{k',j}(Z) \ \exp \left\{ - \frac{i}{v} \int_{Z_0}^{Z} dZ' \varepsilon_{k',j}(Z') \right\}$$  \hspace{1cm} (22)

with the initial condition $\psi_{k_0,i_0}(Z_0) = \varphi_{k_0,i_0}(Z_0)$. Using the time-dependent Schrödinger
corresponding contribution to the force \( (19) \) at \( Z \) and the integration is inside the unshifted Fermi surface \((FV)\). Using \((24)\) and keeping only the first order term:

\[
S(Z_1) = \sum_j \int_{FV} dk \int dk' \Re \left[ \langle \varphi_{k,i}(Z_1) | \frac{dV_{eP}(r; Z)}{dZ} \right|_{Z=Z_1} |\varphi_{k',j}(Z_1) \rangle
\]

\[
\int_{Z_0}^{Z_1} dZ \langle \varphi_{k',j}(Z) | \frac{d}{dZ} \varphi_{k,i}(Z) \rangle \exp \left\{ -\frac{i}{\nu} \int_{Z_0}^{Z} dZ' [\varepsilon_{k,i}(Z') - \varepsilon_{k',j}(Z')] \right\} \]

(27)

Remembering that \( \varphi_{k,i} \) is an eigenfunction of the static \((Z \text{ fixed})\) Schrödinger equation, we obtain readily:

\[
\langle \varphi_{k,i}(Z) | \frac{dV_{eP}(r; Z)}{dZ} | \varphi_{k',j}(Z) \rangle = (\varepsilon_{k,i} - \varepsilon_{k',j}) \langle \varphi_{k,i}(Z) | \frac{d}{dZ} \varphi_{k',j}(Z) \rangle
\]

(28)
The matrix element in the r.h.s. depends only on properties of the adiabatic functions $\varphi_{k,i}$ and not on the velocity. However, from the preceding discussion, we know that inelastic contributions are only appreciable if $(\varepsilon_{k,i} - \varepsilon_{k',j})$ is of the order of $v$. Furthermore, the integration over $k'$ in $(27)$ for a given $k$ can be transformed into an integration over $\varepsilon_{k',j}$. The transformation from one integration variable to the other involves only properties of the static electronic structure and is independent of the velocity. Again, the range of the integration over $\varepsilon_{k',j}$ is of order $v$ around $\varepsilon_{k,i} = \varepsilon_{k',j}$. As a consequence, the quantity $S$ is of order $v^2$.

Until now we have only considered the first order approximation to the $Z$-dependent problem as defined in $(24)$. However, it can be easily verified that each increase in the perturbative order introduces an additional integration over the energy of intermediate states and, therefore, an additional factor of $v$. As a consequence, the non-adiabatic contributions to the force on the projectile are of order $v^2$ at least, which means that they do not contribute to the friction coefficient.

We conclude that both $n_0$ and $n_v$ (to first order in $v$) can be determined exactly by the adiabatic SFS-KS scheme. So the latter scheme provides an exact procedure to determine the friction coefficient.

B. Corollary

The SFS-KS scheme relies only on local properties of the system, i.e., the determination of the density $n_v$ only requires information on the electronic state for a fixed value of $R_P$. It does not require information on the variation of any electronic quantity with $R_P$. Consequently, any alternative to $(20)$ for the evaluation of the friction coefficient must satisfy the same condition. This provides a powerful tool to evaluate the validity of procedures aiming at the evaluation of $\mathcal{F}_F$.

We may apply this condition to analyze the procedure of Trail et al.\textsuperscript{9,11,12} (see also Ref. 13). The latter authors use the expression of the friction coefficient proposed by Hellsing and Persson\textsuperscript{10} (see also Ref. 12).

\begin{align}
\mathcal{F}_{HP} = 2\pi k_F^2 \int d^2k_F \int d^2k_F' \left| \int d\mathbf{r} [\varphi_{k,F}^- (\mathbf{r})]^* \varphi_{k,F}^+ (\mathbf{r}) \right|^2 \\
\left. \mathbf{v} \cdot \nabla_{R_P} |_{n} v_{KS} (\mathbf{r}, R_P; [n]) \right|^2
\end{align}

(29)
where \( v_{KS} \) is the full KS potential, \( k_F \) the Fermi momentum and \( \varphi^+ \) (resp. \( \varphi^- \)) satisfies outgoing (resp. ingoing) boundary conditions. From the derivation of (29) in Ref. 10 (or the alternative derivation in Ref. 12), it is not clear that it consists in a first order approximation in \( v \), though when the impurity is moving through an homogeneous jellium, it can be proved\(^\text{19}\) that (29) is equivalent to (20) to first order in \( v \). A key property, in the latter proof, is the invariance by translation of the free electron gas state in the absence of an external potential. For the inhomogeneous case such an equivalence has not been established. At any rate, in the latter case, and in view of the previous discussion, it is clear that the derivative of \( v_{KS} \) with respect to \( R_P \) in (29) must be carried out for a constant density, otherwise it would introduce a non-local contribution (associated with the variation of \( n \) when the projectile moves). However, the authors of Ref. 9,11–13 use a finite difference method to calculate the derivative of \( v_{KS} \): they determine the KS potential for the projectile at rest and for two different values of \( R_P \), say \( R_P + \delta R_P \) and \( R_P - \delta R_P \). In so doing, they include a term associated with the variation of the function \( v_{KS} \) with \( R_P \). The latter term, being non-local, introduces an error in the evaluation of the friction coefficient.

IV. ILLUSTRATION WITH A 1D MODEL

The previous discussion has raised two questions. Firstly, how important is the error due to the non-local term in the evaluation of the friction coefficient? Secondly, is expression (29) valid for the inhomogeneous case? To answer these questions, we have built a simple 1D model amenable to an exact numerical solution. Full details are given in the Ancillary material\(^\text{19}\). As we are concerned with general principles, our model needs not represent a real system, although we keep speaking of “electrons”. Though units are here irrelevant, since we are only interested in relative results, we keep using atomic units so that the “electrons” have a mass of 1. The interaction between two particles located at \( z \) and \( z' \) is described by a screened regularized coulomb potential:

\[
v_c^\pm(z, z') = \pm e^{-\alpha|z-z'|}/[|z-z'| + \epsilon]
\]

\( v_c^\pm \) between electrons and \( v_c^- \) between electrons and “positive” particles). In the numerical application, we have used \( \alpha = 2 \) and \( \epsilon = 0.1 \). We solve the problem at the level of the Hartree approximation. The Fermi momentum for a 1D paramagnetic homogeneous jellium
The interaction between an “electron” at $z$ and the impurity at $Z_P$ is represented by the potential $v_c^-(z,Z_P)$ with $\alpha = 4$ and $\epsilon = 0.1$. This potential supports one bound singly occupied state with energy -1.275. For all results given below, the projectile is located at $Z_P = 0$. Calculation of the density in the presence of the impurity (even for $v = 0$) requires to confine the system into a box because the Hartree potential behaves as $\sin(2k_F^0z + \gamma_{1,2})/z$ far from the impurity. We have used a finite interval $z \in [-5, 5]$. This does not invalidate our conclusions since we are interested in the difference between two expressions of the friction coefficient for a given system. We have checked that our conclusions are independent of the box size and the implementation of the contour conditions.
FIG. 2: Density in the presence of the background potential (31) with $z_1 = -0.5$, $z_2 = 0.5$, $z_s^1 = 1$ and (a) $z_s^2 = 2$ or (b) $z_s^2 = 1000$. The density has been divided by the density of a uniform jellium with $z_s = 1$.

In a first step, we assume that the impurity, the step potential and the associated background of positive charges move together against a uniform jellium with $z_s = z_s^2$. The friction coefficient can then be calculated in three different ways:

(i) From the 1D expression equivalent to (20):

$$F_{(i)} = \lim_{v \to 0} \frac{1}{v} \int_{-\infty}^{+\infty} dz \left[ n_v(z) - n_0(z) \right] \frac{d}{dZ_P} \left[ v_{eP}(z - Z_P) + v_{\text{step}}(z - Z_P) + v_{+}(z - Z_P) \right]$$

(ii) It is easily shown\textsuperscript{19} that it may be also calculated from the energy loss or gain associated with reflection and transmission by the potential. When $v < k_F^2$:

$$F_{(ii)} = \frac{1}{\pi} \left[ (k_F^1)^2 - (k_F^2)^2 + (k_F^1 + k_F^2)^2 P_r^F \right]$$

where $P_r^F$ is the reflection probability at the Fermi level for the static case. This very simple expression gives us a good check on the accuracy of our SFS calculations.
TABLE I: Friction coefficient when the impurity moves together with the background potential. Results in the first two columns are for double occupancy of the bound state and in the last two columns for single occupancy.

| $z_1$ ; $z_2^2$ | 0.75 ; 1 | 1 ; 1.5 | 1 ; 2 | 1 ; 5 |
|-----------------|---------|---------|-------|-------|
| $F_{(i)}$ (Eq. 32) | 0.576 | 0.472 | 0.545 | 1.06  |
| $F_{(ii)}$ (Eq. 33) | 0.573 | 0.466 | 0.546 | 1.07  |
| $F_{(iii)}$ (Eq. 34) | 0.576 | 0.476 | 0.560 | 1.12  |

(iii) From the 1D version of (29):

$$F_{(iii)} = 2\pi \int dk \int dk' \left| \int dz [\varphi_{k'}(z)]^* \varphi_k(z) \right|^2 \frac{d}{dZ_P} v_{KS}(z - Z_P; n(z, Z_P)) \delta(\varepsilon_F - \varepsilon_k) \delta(\varepsilon_F - \varepsilon_{k'})$$  \hspace{1cm} (34)

As explained above, the derivative with respect to $Z_P$ is evaluated for a constant density $n$, i.e., the quantities entering (34) must be local.

The equivalence between (34) and (32) or (33) is due to the fact that, under the present conditions, the transformation of the system state from one position of the full external potential to another one involves merely a translation (see Ref. 19 for more details).

In Table I we summarize the results obtained for some values of $(z_1; z_2^2)$. Results are the same for the three expressions of the friction coefficient within the accuracy of the calculations. This gives us confidence in our numerical procedures.

We now consider the case of real interest for our model: that of the impurity moving alone, against the step-potential and background of positive charges, i.e., an impurity moving within an inhomogeneous jellium. The calculations only differ from the previous ones by dropping $v_{\text{step}}$ and $v_+$ in (32) and noting that the dependence of $v_{KS}$ on $Z_P$ in (34), for a constant density, arises entirely from $V_{eP}$. Also, (33) is no longer valid. Results are given in Table II.

The first observation is that (34) is no longer equivalent to (32). Therefore, the expression of Hellsing and Persson does not provide the correct first order in $v$ for the energy loss of an impurity in an inhomogeneous medium. The error may be quite appreciable, as shown by our model. Note that the trivial difference mentioned above between the expressions
TABLE II: Same as Table I when the impurity is moving with respect to the fixed background potential. The last line gives the results obtained by inclusion in (34) of the spurious non-local term when the derivative of the Kohn-Sham potential is calculated along (35).

| $z_1$ : $z_2$ | 0.75 : 1 | 1 : 1.5 | 1 : 2 | 1 : 5 |
|-------------|---------|--------|------|------|
| From Eq. 32 | 0.458   | 0.321  | 0.440| 0.71 |
| From Eq. 34 | 1.944   | 2.36   | 2.59 | 4.41 |
| $\Delta v$ from (35) in (34) | 0.404   | 0.254  | 0.379| 0.484|

for a moving and fixed background potential precludes a numerical error as the cause of this discrepancy. The fact that our 1D model may be qualified as unrealistic cannot infirm the relevance of our conclusion since a single counterexample is sufficient to disprove an assertion.

We evaluate now the error incurred when taking into account the change in the potential function in the expression of the potential derivative, as done in the finite difference procedure of Trail et al.\textsuperscript{9,11,12} In place of the derivative of the KS potential for a constant density in (34) we use the expression:

$$\Delta v = \left\{ v_{KS}(z; Z_P + h; [n_{Z_P+h}]) - v_{KS}(z; Z_P - h; [n_{Z_P-h}]) \right\} / 2h$$ (35)

In actual calculations we have used $h = 0.01$. The two KS potentials in (35) are the result of a calculation where the projectile is at $Z_P + h$ and $Z_P - h$ respectively. For that reason, (35) includes a non-local contribution coming from the modification of the potential function when moving the projectile from $Z_P + h$ to $Z_P - h$. Results are given on the third line of Table II. They differ strongly from the previous ones (second line in Table II), up to nearly an order of magnitude. This difference is not due to the simple algorithm used in calculating the derivative as in (35). If we use the same algorithm in expression (34), while keeping the density constant, results agree with those given on the second line of Table II. Changing $h$ (within reasonable bounds) does not change either the conclusion. This demonstrates the significant error introduced by the functional derivative of the Kohn-Sham potential with respect to density. Note that the third line of Table II should not be compared with the first one since it corresponds to an incorrect calculation of expression (34), the latter being intrinsically incorrect for an inhomogeneous system, as verified above. The two errors being
totally unrelated, there is no reason why they should compensate each other. The fact that
the results of the third line in Table II are closer to those of the first one must be considered
as merely accidental.

V. CONCLUSION

Starting from first principles, we have shown that the friction coefficient is, in principle,
exactly determined by an ensemble Kohn-Sham procedure, the ensemble being defined by
a Shifted Fermi Surface. We have shown that it depends only on local properties of the
impurity/target system in the sense that its determination does not require information on
the variation of the system density with impurity position. It may be of interest to check
whether this conclusion could also be reached for other applications in which the low energy
behavior is determined through an adiabatic approach.

To our knowledge, this important constraint has been overlooked. As a consequence
erroneous values of the friction coefficient have been obtained. This is the case, for example,
in the work of Trail et al.\textsuperscript{9}. In addition, the latter authors use expression (29) of Hellsing and
Persson for the friction coefficient, which, though correct for an impurity moving through
an homogeneous system, is not valid for the inhomogeneous case.

That the error may be dramatic is illustrated by the divergence found by Trail et al.\textsuperscript{11,12}. They show, in their analysis, that the divergence is caused by the derivative of the system
spin with respect to $R_P$. The latter quantity is basically non-local (locality being used
here with the meaning defined above) and, therefore, the associated contribution to friction
is spurious. The latter authors wrongly attribute the divergence to a breakdown of the
adiabatic approximation. However, if the evolution for $v \to 0$ is not adiabatic, this means
that there is a discontinuity in the evolution of the system. In the case studied by Trail et
al, no such discontinuity exists. So, the divergence can only be the result of an error in the
evaluation of the friction coefficient, as confirmed by our analysis.

Another conclusion can be drawn with respect to the evaluation by Luntz et al.\textsuperscript{20} of
the “local approximation for friction” (LDAF)\textsuperscript{21}. In the latter approximation, the friction
coefficient is evaluated as a weighted average of the friction coefficient for the impurity in
a homogeneous electron gas having the local density at each point of the inhomogeneous
system. The calculations of Luntz et al.\textsuperscript{20}, based on (29) and (35), are incorrect for the
inhomogeneous case and, therefore, the difference they find with respect to the LDAF cannot be considered as an evaluation of the error incurred when using the LDAF.

Finally, we may remark that the evaluation of friction with present day band structure codes requires an SFS calculation since, up to now, no well founded alternative exists.

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1 P. M. Echenique, F. Flores, and R. H. Ritchie, Solid State Phys. 43, 229 (1990).
2 P.M. Echenique and M.E. Uranga, in Interaction of Charged Particles with Solids and Surfaces, edited by A. Gras-Marti et al. (Plenum, New-York, 1991), p. 39–71.
3 P.M. Echenique, I. Nagy, and A. Arnau, Int. J. Quantum Chem.: Quantum Chem. Symp. 23, 521 (1989).
4 A. Salin, A. Arnau, P. M. Echenique, and E. Zaremba, Phys. Rev. B 59, 2537 (1999).
5 D. Primetzhofer, S. Rund, D. Roth, D. Goebl, and P. Bauer, Phys. Rev. Lett. 107, 163201 (2011).
6 S.N. Markin, D. Primetzhofer, M. Spitz, and P. Bauer, Phys. Rev. B 80, 205105 (2009).
7 G. Martínez-Tamayo, J.C. Eckardt, G.H. Lantschner, and N.R. Arista, Phys. Rev. A 54, 3131 (1996).
8 H. Winter, Phys. Rep. 367, 387 (2002).
9 J. R. Trail, M. C. Graham, and D.M. Bird, Comp. Phys. Comm. 137, 163 (2001).
10 B. Hellsing and M. Persson, Phys. Scripta 29, 360 (1984).
11 J. R. Trail, M. C. Graham, D.M. Bird, M. Persson, and S. Holloway, Phys. Rev. Lett. 88, 166802 (2002).
12 J. R. Trail, D.M. Bird, M. Persson, and S. Holloway, J. Chem. Phys. 119, 4539 (2003).
13 A. C. Luntz and M. Persson, J. Chem. Phys. 123, 074704 (2005).
E. Runge and E.K.U. Gross, Phys. Rev. Lett. 52, 997 (1984).

E.K.U. Gross, J.F. Dobson, and M. Petersilka, in Density Functional Theory, Springer series “Topics in Current Chemistry”, edited by R.F. Nalawajski (Springer, Heidelberg, 1996); M.E. Casida in Recent developments and applications of modern density functional theory, edited by J.M. Seminario (Elsevier, Amsterdam, 1996).

R. M. Dreizler and E.K.U. Gross, Density Functional Theory (Springer-Verlag, Berlin, 1990).

A. Messiah, Quantum Mechanics (North-Holland, Amsterdam, 1962), chapter XVII.

B.H. Bransden and C.J. Joachain, Quantum Mechanics (Longman, Harlow, 1989), section 9.4.

See Ancillary material for a proof of the equivalence between (20) and (29) for the homogenous FEG (Note1.pdf) and for full details on the solution of the 1D model (Note2.pdf).

A. C. Luntz, I. Makkonen, M. Persson, S. Holloway, D. M. Bird, and M. S. Mzielinski, Phys. Rev. Lett. 102, 109601 (2009).

J.I. Juaristi, M. Alducin, R. Díez-Muiño, H.F. Busnengo, and A. Salin, Phys. Rev. Lett. 100, 116102 (2008).

To conserve periodicity, an array of projectiles is required, as always done in band structure calculations. In spite of this, we always refer below, for simplicity, to the projectile.

If degeneracies occur, as is obviously the case in metallic systems, the evolution of the subspaces associated with a given energy follows the same trend as in the non-degenerate case (see, e.g., Ref. 17, section XVII-II). The relation between degenerate substates at different times must be done by continuity. This is trivial if the various substates differ in symmetry, i.e., one or more observables, commuting with the Hamiltonian, allow to distinguish the various substates. Even if the associated eigenvalues are time dependent, this allows to prescribe univocally the adiabatic correspondence between various times. All KS calculations for an impurity in a solid are periodic (periodic array of impurities, supercell for surfaces, etc.). Then, the KS eigenstates are Bloch functions characterized by a $k$ value, which allows to establish the required correspondence in time.

This result is compatible with a deformation of the Fermi surface in the evolution. The change in $k$ for a given state, in the adiabatic evolution, is determined by continuity and the inversion of (2) and (3) ensures the correspondence, to first order in $v$, at any $t$ between the shifted and a corresponding “unshifted” surface. Note that the latter does not correspond to the solution for the projectile at rest since the KS equations have been solved for a shifted Fermi surface. This
introduces a difference of order $v$ in the potential which is the origin of friction. Obviously, the SFS solution is not merely the Galilean transform of the solution with the projectile at rest.