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Exact time-dependent exchange-correlation potential in electron scattering processes

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We identify peak and valley structures in the exact exchange-correlation potential of time-dependent density functional theory that are crucial for time-resolved electron scattering in a model one-dimensional system. These structures are completely missed by adiabatic approximations which consequently significantly underestimate the scattering probability. A recently-proposed non-adiabatic approximation is shown to correctly capture the approach of the electron to the target when the initial Kohn-Sham state is chosen judiciously, and is more accurate than standard adiabatic functionals, but it ultimately fails to accurately capture reflection. These results may explain the underestimate of scattering probabilities in some recent studies on molecules and surfaces.

Electron scattering is one of the most fundamental processes in physics, chemistry, and biology. Electrons constantly collide with other electrons and nuclei in chemical reactions and physical processes, from molecular electronics [1] to strong-field processes [2]. Some experimental techniques directly utilize electron scattering, such as transmission or scanning electron microscopy [3], to investigate surface structures. Radiation damage caused by low-energy electron scattering from DNA highlights its relevance for biomolecules [4]. Despite the importance of electron scattering processes, its theoretical description remains challenging: electron scattering is a highly-correlated many-body problem and generally requires treatment beyond perturbation theory.

Time-dependent density functional theory (TDDFT) [5] is the most widely used first-principles approach to study real-time many-electron dynamics. This is due to its favorable system-size scaling, which arises because it maps the correlated electron dynamics to that of the non-interacting Kohn-Sham (KS) system, evolving in a single-particle potential. All many-body effects are hidden in the single-particle exchange-correlation (xc) potential, which in practice must be approximated. TDDFT with approximate xc potentials has been successfully applied to interpret and predict electron dynamics in a range of situations [6], in addition to predicting linear response and spectra, which is the regime it is mostly known for. Indeed, it has been applied to compute elastic electron-atom scattering cross-sections by means of linear-response theory [7], and recently also applied to real-time non-perturbative calculations of proton-methane scattering [8] and of electron wavepacket scattering from graphene [9].

However, TDDFT with approximate xc potentials fails to even qualitatively reproduce the true dynamics in some applications to non-linear time-resolved dynamics [10]. In principle, the exact xc potential \( v_{xc} \) at time \( t \) functionally depends on the history of the density \( n(x,t') < t \), the initial interacting many-body state \( \Psi_0 \), and the choice of the initial KS state \( \Phi_0 \). In reality, almost all calculations today use an adiabatic approximation that inputs the instantaneous density into a ground-state xc functional. Recent studies on exactly-solvable model systems [11-14] reveal that large non-adiabatic features can appear in the exact xc potential that are missing in the approximations. How accurate is TDDFT with the currently available approximate xc potentials for electron scattering? Given the dearth of alternative practical ab initio methods for this problem and its relevance in a wide range of situations of interest today, it is crucial to assess the reliability of TDDFT approximations for scattering.

To this end, we study a model system of electron-Hydrogen (e-H) scattering that can be solved numerically exactly, and show that the exact xc potential develops non-adiabatic peak and valley structures that are dominantly responsible for causing scattering. Standard functional approximations lack these structures and severely underestimate the scattering probability. Although a recently-proposed non-adiabatic orbital-dependent functional significantly improves the dynamics in the approach of the electron to the target for a judiciously chosen initial KS state, it also ultimately fails to accurately scatter. We identify the term in the exact xc potential crucial for the scattering, as an explicit functional of the many-body density-matrix and KS orbitals. Our results stress the need to develop explicit density or orbital-functionals for this term, and suggest that, generally, adiabatic TDDFT tends to underestimate the scattering and energy-transfer in real systems [8,9].

The Hamiltonian of our one-dimensional two-electron model system reads (we use atomic units throughout unless otherwise stated): \( \hat{H}(x_1, x_2) = \sum_{i=1,2} \left( -\frac{1}{2} \frac{\partial^2}{\partial x_i^2} + v_{ext}(x_i) \right) + W_{xc}(x_1, x_2) \), where \( W_{xc}(x_1, x_2) = \frac{1}{\sqrt{(x_1-x_2)^2+1}} \) is the soft-Coulomb interaction [15] and \( v_{ext}(x) = -\frac{1}{\sqrt{(x+10)^2+1}} \) is the external potential that models the H atom located at \( x = -10.0 \)
a.u. The soft-Coulomb interaction has proven extremely useful for analysis and predictions in strong-field physics as well as in density functional theory as it captures the essential physics of real atoms and molecules [16].

The initial interacting wavefunction is taken to be a spin-singlet, with spatial part

$$\Psi_0(x_1, x_2) = \frac{1}{\sqrt{2}} (\phi_H(x_1)\phi_{WP}(x_2) + \phi_{WP}(x_1)\phi_H(x_2))$$

where $\phi_H(x)$ is the ground-state hydrogen wavefunction. $\phi_{WP}(x)$ is an incident Gaussian wavepacket,

$$\phi_{WP}(x) = (2\alpha/\pi)^{1/4} e^{-\alpha(x-x_0)^2 + i p(x-x_0)}$$

with $\alpha = 0.1$, representing an electron at $x_0 = 10.0$ a.u. approaching the target atom with a momentum $p = -1.5$ a.u. in our first example. The full time-dependent Schrödinger equation $i\partial_t \Psi(x_1, x_2, t) = \hat{H}(x_1, x_2)\Psi(x_1, x_2, t)$ can be solved numerically exactly for this system, and we plot the resulting density, $n(x, t) = \int |\Psi(x, x_2, t)|^2 dx_2$, as the black lines in the upper panel for different time-slices in Fig. [1][17]. The incident wavepacket collides with the target electron at around 0.24 fs, after which, a major part of the wavepacket is transmitted while some is reflected back. The black lines in upper panels of Fig. [2] show the number of electrons $N_R$ ($N_T$) in the reflection (transmission) region, calculated by integrating $n(x, t)$ over the region $x > -5.0$ a.u. ($x < -15.0$ a.u.); notice that the incoming electron leaves the target partially ionized. At this incident momentum, the scattering is inelastic, evident in density oscillations in the target after the incoming electron has well passed (evident in the movies in the supplementary information [17]); later we shall consider the case of low-energy elastic scattering. The length of the simulation box was 200 a.u. and the results are converged over this time period with respect to box size and boundary conditions (absorbing and hard-wall). Shown also in both these figures are the results from TDDFT approximations (red and blue lines), neither of which yield reflection nor the dynamics correctly. To understand why, we now consider the xc potential for an exact TDDFT calculation and compare this with the approximate potentials.

A TDDFT calculation starts with the specification of the initial KS state, for which there is considerable freedom. For initial ground-states, the natural choice is the KS ground-state, but for a general initial state $\Psi(0)$, one may pick any initial KS state $\Phi(0)$ that has the same density and first time-derivative of the density as that of the interacting system [13]. Recently the impact of this choice on the xc potentials has been studied [13][19].

We consider two natural choices of initial KS state for the scattering problem. The first one is the Slater determinant, which, for our spin-singlet state involves one doubly-occupied spatial orbital,

$$\Phi_0^{(1)}(x_1, x_2) = \phi_0(x_1)\phi_0(x_2)$$

with $\phi_0(x) = \sqrt{\frac{n_0(x)}{2}} \exp \left[ i \int x \frac{j_0(x')}{n_0(x')} \, dx' \right]$, where $n_0$ and $j_0$ are the initial density and current density of the interacting system respectively. On the one hand, as a Slater determinant, it is in keeping with usual KS approach; on the other hand it has a very different form than the physical state. The second initial KS state we consider has the scattering form of Eq. (1), with two orbitals, and to reproduce $n_0$ and $\partial_t n_0$ of Eq. (1) we must in fact take

$$\Phi_0^{(2)}(x_1, x_2) = \Psi_0(x_1, x_2).$$

Each of these KS states is a valid initial state for the KS evolution, for which the exact xc potentials $v_{xc}[n; \Psi_0, \Phi_0^{(1)}](x, t)$ and $v_{xc}[n; \Psi_0, \Phi_0^{(2)}](x, t)$ can be...
found that reproduce the exact density. This can be done by using the global fixed-point iteration method of Ref. [20]. (For the single-spatial orbital case, it can also be obtained more simply by inverting the TDKS equation [11]).

The black solid lines in the lower panels of Figs. 1 and 3 show the snapshots of the exact xc potentials $v_{xc}[n; \Psi_0, \Phi_0^{(1)}](x, t)$ and $v_{xc}[n; \Psi_0, \Phi_0^{(2)}](x, t)$ respectively, demonstrating remarkable features of the xc potentials in the scattering process [17]. In the $\Phi_0^{(1)}$ case (Fig. 1), $v_{xc}[n; \Psi_0, \Phi_0^{(1)}](x, t)$ develops dynamical peak and step structures throughout the scattering process, beginning from when the incident wavepacket approaches the atom’s electron. The exact xc potential for $\Phi_0^{(2)}$ (Fig. 3) has no structure at very early times, but displays a large peak structure behind the center of the target during the approach (at around $t = 0.24$ fs), and complicated structures after the electron reaches the interaction region. In fact for both choices of KS initial state, a series of peaks and valleys is evident in the interaction region, that are largely responsible for the scattering, as we will now argue.

We consider propagation under two approximations in which these structures are absent. The first is the adiabatic local density approximation (ALDA), based on the one-dimensional uniform gas [21]. The density and xc potential are shown as the red lines in Figs. 1 and 3 for the two choices of the initial state [17]; see also Fig. 2 for the $N_R$ and $N_T$ values. In both cases, the ALDA dynamics fails to even qualitatively capture the reflection; for $\Phi_0^{(1)}$, the electron density develops unphysical spurious oscillating structures in the interaction region, while for $\Phi_0^{(2)}$, although these oscillating structures are absent, and the earlier dynamics is better reproduced than the $\Phi_0^{(1)}$ case (until about $t = 0.36$ fs), the ALDA xc potential too smoothly tracks the instantaneous density throughout. In both cases, ALDA, with its simple local dependence on the instantaneous density, cannot develop the peaks and valleys of the exact xc potential, and yields far reduced reflected density and reduced energy loss [22] to the target.

One might be tempted to attribute the poor performance of ALDA for scattering simply to the incorrect fall-off of its xc potential, however similar results are obtained with adiabatic exact exchange AEXX (shown in Fig. 3), so getting the asymptotics correct is not enough to get good dynamics [17]. The next approximation arises from considering first the decomposition of the exact xc potential into kinetic (T) and interaction (W)

![FIG. 2. (color online). Number of electrons in the reflection region $N_R$ (left panel) and transmission region $N_T$ (right panel) for the exact (black solid), ALDA ($\Phi_0^{(1)}$: red dashed, $\Phi_0^{(2)}$: red solid), $v_{xc}^{ALDA}(\Phi_0^{(1)}$: blue dashed, $\Phi_0^{(2)}$: blue solid), AEXX ($\Phi_0^{(1)}$: equal to $v_{xc}^{ALDA}$ (blue dashed), $\Phi_0^{(2)}$: cyan solid) for the two different momenta $p = -1.5$ (upper panels) and $p = -0.3$ (lower panels).](image)

![FIG. 3. Same as Fig. 1 but for the initial KS state $\Phi_0^{(2)}$.](image)
terms, \(v_{xc} = v_c^T + v_{xc}^W\) \cite{12 13 23}, where

\[
v_{xc}^T(x, t) = \int \frac{1}{4n(x'', t)} \left( \frac{d}{dx'} - \frac{d}{dx''} \right) \left( \frac{d^2}{dx'^2} - \frac{d^2}{dx''^2} \right) (\rho_1(x', x'', t) - \rho_{1,s}(x', x'', t)) |x' = x''| dx',
\]

(5)

\[
v_{xc}^W(x, t) = \int dx'' \int n_{xc}(x', x'', t) \frac{\partial}{\partial x''} W_{xc}(|x' - x''|) dx',
\]

(6)

where \(n_{xc}\) is the xc hole, defined via the pair density as \(P(x', x, t) = N(N - 1) \sum_{\sigma_1, \cdots, \sigma_N} \int |\Psi(x'\sigma_1, x\sigma_2, x_3\sigma_3 \cdots x_N\sigma_N, t)|^2 dx_3 \cdots dx_N = n(x, t)(n(x', t) + n_{xc}(x', x, t))\), and \(\rho_1\) and \(\rho_{1,s}\) are the spin-summed one-body density matrices for the interacting system and KS system respectively. For a wide range of dynamics of two-electron systems (driven dynamics to local excitations, charge-transfer dynamics, field-free dynamics of non-stationary states), \(v_{xc}^T\) has been found to exhibit larger non-adiabatic features than \(v_{xc}^W\) and was the main origin of step and peak structures \cite{12 13}. We find here that this is also true for scattering, where \(v_{xc}^T\) largely comprises the complicated peak and valley structures in the exact xc potential; this is plotted as the green dotted line in Figs. 1 and 3 (Note that the choice of \(\Phi_0^{(2)}\) makes \(v_{xc}^T\) zero at very early times).

The exact decomposition Eq. (5)–(6) motivates the \(v_{xc}^S\) approximation to the xc potential \cite{13}, defined as

\[
v_{xc}^S(x, t) = \int dx'' \int n_{xc}^S(x', x'', t) \frac{\partial}{\partial x''} W_{xc}(|x' - x''|) dx',
\]

(7)

where \(n_{xc}^S(x', x'', t)\) is the xc hole of the KS system. That is, \(v_{xc}^S(x, t)\) replaces all quantities on the right of Eq. (5)–(6) with their single-particle KS values. This approximation yields an orbital-dependent functional, that generally has spatial- and time- non-local dependence on the density, since it depends instantaneously on the density, each of which has a time-non-local dependence on the density, and on the KS initial state. In the general case it includes correlation but it reduces to the time-dependent (TD) EXX approximation when the KS state is a Slater determinant, and in the special case of two-electrons in the same spatial orbital, TDEXX reduces to AEXX, depending only on the instantaneous density. So, propagating \(\Phi_0^{(1)}\) with \(v_{xc}^S\) is equivalent to AEXX. Although this is self-interaction free and has correct asymptotic behavior, this also displays spurious oscillation in the density from early on as was seen in ALDA, and does not improve the time-resolved reflection probabilities (blue line in Figs. 1 and 2) \cite{17}. With the choice of \(\Phi_0^{(2)}\), \(v_{xc}^S\) reproduces the exact xc potential and dynamics very well at earlier times, as is evident in Fig. 3 and the approach of the electron to the target is very well-captured. Actually \(v_{xc}^S\) is a very good approximation to \(v_{xc}^W\), however without \(v_{xc}^T\) the dynamics begins to deviate from the exact during the scattering process, and ultimately it also gives reduced reflection and reduced energy loss to the target \cite{17}. In fact, propagation with the exact \(v_{xc}^W = v_{xc}^T - v_{xc}^S\), for either initial state \(\Phi_0^{(1)}\) or \(\Phi_0^{(2)}\), gives reduced reflection and energy loss (Data not shown here. For \(\Phi_0^{(2)}\), propagation with \(v_{xc}^W\) yields results very similar to that with \(v_{xc}^S(x, t)\)). Thus \(v_{xc}^T\) is the key component to reproduce scattering correctly, and approximations which lack a good model for this term will fail to even qualitatively capture the reflection.

Can the relevant structures in \(v_{xc}^S\) be modeled by any adiabatic approximation? To answer this, we consider the best adiabatic approximation possible, the adiabatically exact (AE). This is defined as the exact ground-state xc potential evaluated on the instantaneous density, i.e., \(v_{xc}^{AE} = \rho_{ex}^{\sigma=\sigma_0} - \rho_{ex}^{\sigma=\sigma_1} - \rho_{ex}^{\sigma=\sigma_2}\), where \(\rho_{ex}^{\sigma=\sigma_0} = \rho_{ex}^{\sigma=\sigma_1} = \rho_{ex}^{\sigma=\sigma_2}\) is the external (KS) potential for interacting (non-interacting) electrons whose ground state has density \(n\). By inversion of the ground-state KS equation, \(\rho_{ex}^{\sigma=\sigma_0} = \nabla^2 \sqrt{n} / (2 \sqrt{n})\) up to a constant, while to find \(v_{xc}^{AE} = \rho_{ex}^{\sigma=\sigma_0} - \rho_{ex}^{\sigma=\sigma_1} - \rho_{ex}^{\sigma=\sigma_2}\), an iterative method \cite{24} was employed. In Fig. 4, we show snapshots of \(v_{xc}^{AE}\) evaluated on the exact instantaneous density along with the AE kinetic contribution \(v_{xc}^{AE,T} = v_{xc}^{AE} - v_{xc}^{AE,W}\). We see that

\[v_{xc}^{AE,T,}\text{ although not structure-less, misses the dominant structures of the exact kinetic contribution }v_{xc}^{S}\text{; these are truly non-adiabatic features and are essential to capture the scattering even qualitatively.}\]

We consider now the case of elastic scattering by reducing the incoming momentum to \(p = -0.3\) a.u. such that the energy is lower than the lowest excitation of the target (which is about \(\omega = 0.4\) a.u.), so that inelastic channels are closed. Again neither ALDA nor \(v_{xc}^{S}\) even qualitatively capture the scattering dynamics, for either choice of initial state, as is clear from the lower
panels in Fig. 2 (a movie is given in the supplementary information [17]). Yet, Refs. 7 show that good scattering cross-sections can be extracted from the TDDFT linear response formalism, when using standard approximations. With a formally exact theory, the time-domain picture should agree with the time-independent picture for elastic scattering in the long-time limit [25], however this is not the case for approximate TDDFT. The time-resolved picture presents scattering as a fully non-equilibrium problem, where the system starts far from a ground-state and so from the very start the xc functional is evaluated on a system far from a ground-state. This is quite unlike using the same functional in a linear response calculation where it is evaluated on densities close to a ground-state; typical adiabatic approximations work much better in the latter. As the present work has shown, non-adiabaticity beyond the adiabatic approximation and beyond what is contained in \( v_T^{xc} \) is required to give even qualitatively accurate time-resolved dynamics. The situation is similar to that for field-free dynamics of a system in a superposition state: one can find adiabatic functionals which yield good predictions for excitation energies and so can accurately predict the period of its density-oscillations, but when the time-resolved dynamics is run from a superposition state, the oscillation period of the time-dependent dipole can deviate significantly [12, 26].

In summary, we have analyzed the exact xc potentials for a model system for inelastic and elastic scattering. We showed the importance of choosing the initial KS state appropriately, and revealed how and why why ALDA and EXX cannot reproduce the correct scattering dynamics. Although the recently proposed non-adiabatic approximation \( v_T^{xc} \) greatly improves the dynamics up to the time of interaction, ultimately it also fails to capture the scattering accurately. The peak and valley structures in the kinetic component of the exact xc potential \( v_T^{xc} \) are missing in all these approximations, and this work suggests the urgent need for reasonable density- or orbital-functional approximations to this term (Eq. 5) to improve the reliability of TDDFT to describe time-resolved scattering processes. Similar trends hold for a model electron-Helium+ system, and also for triplet scattering. With regards to realistic three-dimensional systems, the one-dimensional model here obviously cannot describe contributions from channels where the electron scatters “around” the target, and the effect of the peak and valley structures uncovered here could be buffered when many electrons and vibronic effects are considered, but nevertheless the lack of these in approximations likely yield underestimated scattering probabilities in realistic systems.

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